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Research Article

# Structure and biological activity of pathogen-like synthetic nanomedicines

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## Abstract

Here we characterize the structure, stability and intracellular mode of action of DermaVir nanomedicine that is under clinical development for the treatment of HIV/AIDS. This nanomedicine comprises pathogen-like pDNA/PEIm nanoparticles (NPs) having the structure and function resembling spherical viruses that naturally evolved to deliver nucleic acids to the cells. Atomic force microscopy demonstrated spherical 100 – 200 nm NPs with a smooth polymer surface protecting the pDNA in the core. Optical absorption determined both the NP structural stability and biological activity relevant to their ability to escape from the endosome and release the pDNA at the nucleus. Salt, pH and temperature influence nanomedicine shelf-life and intracellular stability. This approach facilitates the development of diverse polyplex nanomedicines where the delivered pDNA-expressed antigens induce immune responses to kill infected cells.

**From the Clinical Editor:** The authors investigated DermaVir nanomedicine comprised of pathogen-like pDNA/PEIm nanoparticles with structure and function resembling spherical viruses. DermaVir delivery of pDNA expresses antigens that induce immune responses to kill HIV infected cells.

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**Key words:** Vaccine; Immunotherapy; Immunity; DermaVir

Nanomedicines developed for the induction of immune responses comprise the formulation of plasmid DNA (pDNA) into nanoparticles (NPs), and thus targeting antigen presenting cells of the immune system.<sup>1–3</sup> It is established that negatively charged pDNA spontaneously forms NPs with positively

charged polymers.<sup>4–8</sup> Linear polyethylenimine (PEI) has been developed as a synthetic delivery system for pDNA<sup>1,8–10</sup> and can act as a proton sponge thus protecting the pDNA from endosomal degradation. We have modified the PEI with sugar residues and obtained a novel mannobiosylated linear polyethylenimine (PEIm) to target the pDNA-encoded antigens to antigen-presenting cells required for the induction of potent immune responses.<sup>11</sup> The surface mannose residues on the pDNA/PEIm NPs mimic the surface of pathogens that are recognized by antigen-presenting cells.<sup>12,13</sup> In comparison with viral vectors, synthetic NPs are attractive for pDNA-based vaccine development due to their excellent safety features and the absence of vector-induced immune responses that makes them suitable for repeated administration and the boosting of immunity. The safety, immunogenicity and preliminary efficacy of the pDNA/PEIm nanomedicine that we studied (DermaVir) have been reported in SIV-infected primate and HIV-infected human subjects.<sup>14–18</sup> These polyplex nanomedicines offer a promising new treatment paradigm for the treatment of diverse

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diseases, including cancer, allergic and chronic infectious diseases. However, before polyplex nanomedicine products approach the global market, a number of issues need to be comprehensively addressed beyond clinical efficiency and safety, such as the relationship of their physical structure and biological function.

In this study we investigated how the physicochemical parameters of a polyplex nanomedicine influence its biological activity and clarify the relationship between NP structure, nanomedicine shelf-life and cellular mode of action. First, we characterized the geometry of the DermaVir nanomedicine (DV) and its active pharmaceutical ingredient, a large pDNA encoding 15 HIV antigens, by direct observation made by atomic force microscopy (AFM) and confirmed these findings by dynamic light scattering (DLS). These results showed that the large pDNA alone does not form NPs and the extent of NPs' compaction is formulation dependent. Because the variable degree of compaction of the pDNA might influence the biological activity, we characterized the stability of these nanomedicines by using optical absorption spectroscopy (hyperchromicity). We found that hyperchromicity measurement can be effectively used to determine the effect of the primary physical variables (pH, salt concentration, temperature) on the nanomedicine stability and "aging" (property drifts in time due NP reorganization). Finally, we correlate the biological activity of the nanomedicine to the physical methods of characterization. Having measurements to routinely characterize the properties of nanomedicine formulations is required for efficient development of these materials for the prevention and treatment of major chronic diseases.<sup>9</sup>

## Methods

All experiments were repeated at least once, more preferably twice. Three parallel samples were used for UV-vis and particle size measurements, and 5 samples were used for biological activity assays. Uncertainties were estimated as 1 standard deviation (SD) from the mean from multiple measurements.

### Preparation of NPs

pDNA: pLWXu1 1 mg/mL solutions encoding 15 HIV antigens (12.5 kbp).<sup>19</sup> PEIm: 13.6 mM solution of linear 22 kDa polyethylenimine containing 3% grafted mannobiose, pKa =  $8.3 \pm 0.022$  (in vivo-jetPEI-Man, PolyPlus Transfection, Illkirch, France).<sup>20</sup> Different pDNA and PEIm lots were manufactured. 10 mmol/L triethanolamine buffer and triethanolamine-hydrochloride (TEA/HCl, Sigma, Budapest, Hungary) containing 10w/w% Mannitol (TEAM) was prepared at pH  $7.5 \pm 0.2$ , or as indicated.

DV1 formulation was prepared by diluting the pDNA with 6 volume glucose solution (10 w/w%) and PEIm at a PEIm/pDNA molar equivalent (N/P ratios) of 4 (unless otherwise indicated). DV2 formulation was prepared by diluting the pDNA with 6 volume TEAM buffer and PEIm at a PEIm/pDNA molar equivalent (N/P ratios) of 4 (unless otherwise indicated). N/P ratio is calculated by dividing the nitrogen concentration of the PEIm solution with the phosphorous concentration of the pDNA solution.

### SDS decomplexation of NPs

1,000 times molar excess of sodium dodecyl sulfate (SDS, Sigma) solution was added to the prepared NPs and incubated 1 hour at room temperature ( $23 \pm 2^\circ\text{C}$ ). The spectra of the SDS-treated NPs were normalized with the spectra of the SDS in the used concentration.

### Nuclease resistance of the NPs

NPs were incubated with endonuclease enzyme (DNase I, New England BioLabs, Ipswich, Massachusetts), at  $37^\circ\text{C}$  for 15 minutes. Control pDNA was digested for 5 minutes at  $37^\circ\text{C}$ . Sample-enzyme ratio was chosen according to the manufacturer's instructions. Reaction was terminated by adding EDTA solution to the samples. After digestion SDS decomplexation of the NPs was performed as described above and evaluated by AGE as described below.

### AFM

Measurements were made using a commercial AFM (Bioscope I with Nanoscope IV controller, Veeco Instruments, Santa Barbara California). A 7  $\mu\text{L}$  drop of the NP solution was placed on freshly cleaved mica or APS-mica for 3 minutes, then washed with ultra-pure, deionized water to remove salts and dried in a gentle stream of argon. AFM imaging was made in air in the tapping mode. Silicon cantilevers were used (OMCL by Olympus, Tokyo, Japan), which have a nominal spring constant of 42 N/nm, resonant frequency of 300 kHz and nominal tip radius of 7 nm. Image processing was done using the Nanoscope Imaging software (Veeco) and the NIH ImageJ software (available at <http://rsb.info.nih.gov/ij>). For sampling consistency, all height and width measurements were performed along 2 cross-sections of each molecule or NP. We report the average data of the measurements made on 40 to 80 adsorbed NPs. Measurements showed little change in the nanomedicine morphology with the surface interaction (APS or mica).

### Agarose gel electrophoresis (AGE)

Overall, 250 ng pDNA was loaded into 1% agarose gel (containing ethidium bromide) and TAE buffer pH 8.3 (Invitrogen, Carlsbad, California). For the evaluation of supercoiled content, NIH ImageJ software was used.

### Hyperchromicity (Hc %)

Hc % was determined by UV-vis spectrophotometry, validated according to ICHQ2R guidelines, performed on 30  $\mu\text{g/mL}$  pDNA in triplicate (repeatability is within 2%), using quartz cuvette in a calibrated Jasco V-630 instrument. Full spectra were recorded from 190 to 1100 nm with 5 nm step width. The calculation of Hc percentage was the following:

$$100 \times (A_{260\text{NP}} - \sum A_{260\text{components}}^*) / (\sum A_{260\text{components}})$$

$$* \sum A_{260\text{components}} = A_{260\text{pDNA}} + A_{260\text{PEIm}} + A_{260\text{formulation solvent}}$$

### Particle size measurements

Particle size measurements were performed from 10  $\mu\text{g/ml}$  pDNA samples using Brookhaven ZetaPALS instrument equipped with quartz cuvette. Parameters: 3 runs; temperature: 25°C.

### Biological activity

Biological activity of DermaVir nanomedicine was determined by quantifying the expressed proteins in the supernatant by p24 ELISA as previously described.<sup>9,20</sup>

### Zeta potential analysis

Zeta potential analysis was performed from 10  $\mu\text{g/ml}$  pDNA samples using a Brookhaven ZetaPALS instrument equipped with “four side clear” plastic cuvette. Parameters: 5 runs, 10 cycles; temperature: 25°C.

### Complement activation assay

In vitro complement activation assay for the quantification of serum S-protein-bound C terminal complex (SC5b-9) was performed as described earlier.<sup>21</sup> The test samples and control compounds were incubated with human sera for 45 minutes at 37°C. After terminating the reaction samples were tested for SC5b-9 levels using the TCC ELISA kit (Quidel Co., San Diego, California), following the manufacturer’s instructions. For positive control Zymosan, which is known to induce high activation of the complement system<sup>22</sup> was used.

### Statistical analysis

To assess the significance of the results, Student’s *t*-test was performed on selected data.

## Results

We examined 2 nanomedicine formulations prepared by the same mixing procedure: DV1 and DV2 in glucose and in TEAM buffer, respectively (see Methods section). Some physicochemical properties of the 2 formulations are summarized in Table 1. We found that minor changes in formulation present major differences in structure and biological activity.<sup>9</sup>

### Characterization of pDNA nanomedicine morphology

We determined the structure of the nanomedicine with 2 independent methods because NP size is one of the critical parameters in our conceptual model of pathogen-like particle

delivery. Figures 1, A–D show AFM images of the pDNA and the NPs found in both DV1 and DV2 formulations. The uncondensed pDNA strands are uniformly distributed and form a network structure typical of solutions containing long overlapping semiflexible polymer chains (Figure 1, A). The NPs, in both cases, have a roughly spherical shape having a diameter in the range between 100 nm to 200 nm, but the NPs in the DV2 formulation have rather smooth surface (Figure 1, B). In contrast, the DV1 formulation exhibits hair-like protrusions of presumably uncompacted pDNA, hinting that these structures might be less stable (Figure 1, C, D). The minimum pDNA width is about 6 nm, indicating that this part of the pDNA is uncoated. At other places, the DNA is thicker (about 10 nm) suggesting that PEIm coats the pDNA (Figure 1, D). Similar structures are seen when such pDNA complexes are induced to become uncomplexed by molecular additives so that DV1 is apparently a less stable nanomedicine formulation.<sup>9,19</sup> We amplify on this interpretation below for the different morphologies of DV1 and DV2 NPs. Direct AFM measurements of polyplexes between pDNA and poly(amido amine) synthetic polyelectrolytes has directly revealed that these complexes predominantly involve a single pDNA molecule, although some larger clusters containing a few pDNA molecules are also observed to occur.<sup>23</sup> We expect the same behavior to occur in polyplexes and indeed find evidence for smaller and larger clusters in our DLS measurements.

To further validate the estimates of the nanomedicine size, we performed DLS measurements in aqueous solution (see Methods). Consistent with the AFM data, the NP diameter estimated from the experimental diffusion coefficient and the Stokes-Einstein relationship was found to be between 100 nm and 200 nm for both formulations (Table 1). Both the AFM and DLS measurements indicated a bimodal size distribution with NP sizes near 50 nm to 90 nm and 170 nm to 260 nm (Figure 1, E).

Next, we studied the size of the NPs using DLS for variable pDNAs lengths, percentage of supercoiled content of the plasmid and pDNA and PEIm component ratios. For each variable, we demonstrated the formation of 70 nm- to 300 nm-size NPs. These results indicate that the length, the supercoiled content of the pDNA and the molar ratio of the pDNA and PEIm component (N/P ratio, see Methods) does not substantially affect NP size (Supplementary Tables 1, 2 and 3). We also investigated NPs made with different molecular weight PEIm’s and with PEI and showed that neither of these changes was related to the particle size (Supplementary Table 4). These results further demonstrated the robust capacity of these cationic polymers to condense the pDNA and form NPs.

Both DLS and AFM measurements indicate that these pathogen-like nanomedicines have shapes and dimensions similar to those of viruses,<sup>24,25</sup> but the different formulations exhibit a somewhat different fine structure. These variations in NP structure strongly suggest that the strength of the associations between the pDNA and synthetic polymer component is different in the 2 formulations, even though the overall size of the NPs remained constant.

### Characterization of NP formation

Although a fundamental measurement of the binding constant between the pDNA and synthetic polymers is possible through

Table 1  
Physicochemical properties of DV1 and DV2 nanomedicines

Property	DV1	DV2
pH	3.5	7.5
Conductivity ( $\mu\text{S/cm}$ )	135	75
Particle size ( $D_{\text{eff}} \pm \text{SD nm}$ )*	130 $\pm$ 7	133 $\pm$ 5
Hyperchromicity (Hc) % <sup>†</sup>	13.7	23.7

\* Particle size measured by DLS method.

<sup>†</sup> The calculation of Hc % is described in Methods section.

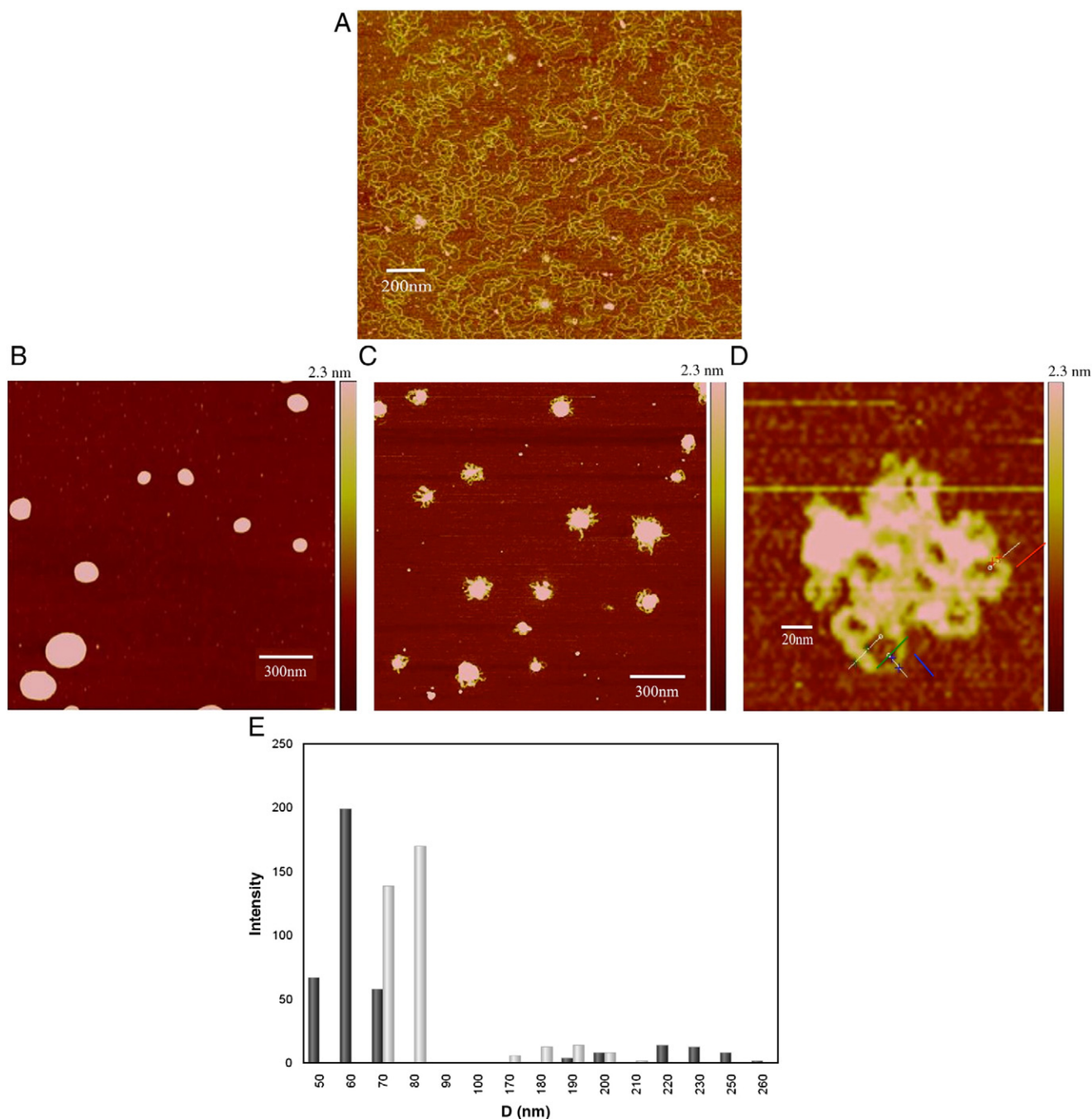


Figure 1. Investigation of the particle sizes of pDNA/PEIm NPs. **(A)** AFM image of pDNA in the absence of PEIm on mica surface. The pDNA strands form a random mesh typical of polymer solutions of long overlapping chains. **(B)**, AFM images of DV2 on mica. The NPs have smooth interfaces. **(C)**, AFM image of DV1 on mica. Hair-like protrusions of presumably uncompacted DNA are clearly visible. **(D)**, Magnified AFM image of the DV1. The minimum pDNA width is 6 nm, indicating that this part of the pDNA is uncoated DNA. At other places the pDNA is thicker (about 10 nm), suggesting that PEIm coats the pDNA. **(E)** Representative particle size distributions of DV1 and DV2 measured by DLS; ■—DV1 and □—DV2.

competitive binding measurements,<sup>26,27</sup> it is desirable for clinical product formulation development to have a more facile method. A qualitative indication of this binding phenomenon can be obtained from absorption spectroscopy measurements. After adding PEIm to the pDNA we have observed an increase of absorbance — without any shift — in the whole spectral range, including the wavelength range far from pDNA absorption, i.e.,

$\lambda \geq 320$  nm (Figure 2, A). This Hc phenomenon is defined by the increase of optical density of a solution caused by changes of associative interactions of chemical species (see Methods for calculation of Hc%) and this measurement is particularly sensitive to the configuration of the base units of DNA because the proximity of these polarizable species gives rise to shifts in the optical absorption. Hc measurements are thus routinely used

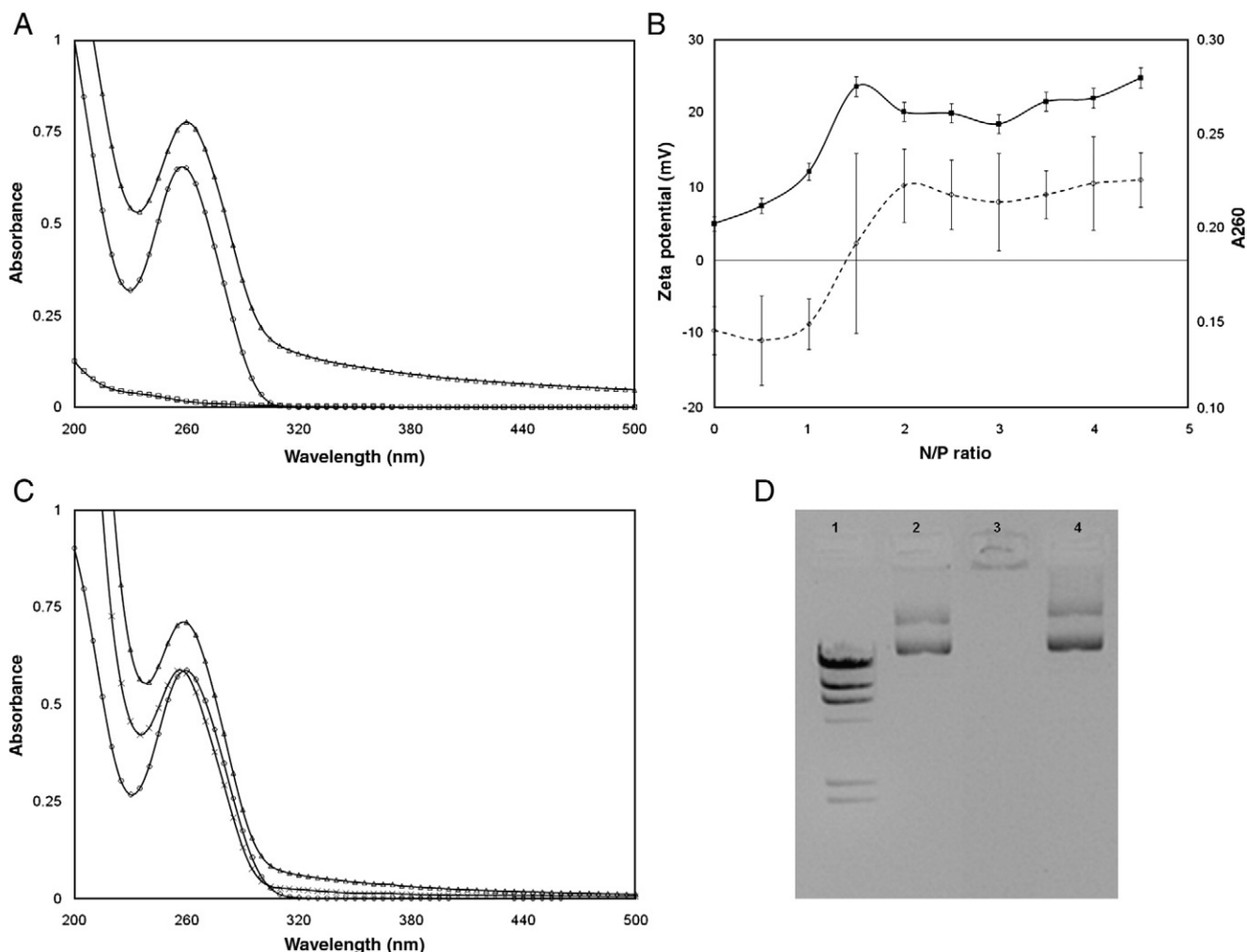


Figure 2. Spectrophotometric and zeta potential investigation of pDNA/PEIm NPs (DV2). (A), UV-VIS-spectra of pDNA/PEIm NP and its components; pDNA (O), PEIm (□), pDNA/PEIm NP (Δ). (B), Titration of pDNA (10  $\mu\text{g/ml}$ ) with PEIm by UV-spectrophotometry (■) and zeta potential analysis (○). N/P = 0 sample contains only pDNA. (C), UV-VIS-spectra of pDNA (O), pDNA/PEIm NP before SDS treatment (Δ) and pDNA/PEIm NP after SDS treatment (×). SDS absorbance was subtracted from the spectrum. (D), Residual gel electrophoresis of the pDNA (lane #2) and the NP before (lane #3) and after (lane #4) SDS treatment.

to characterize DNA double helix formation (“hybridization”) in solution and these previous studies of DNA–DNA and DNA–NP interaction are informative about the interpretation that underlies this type of change in pDNA solution absorption spectra.<sup>28</sup> For investigation of Hc in nanomedicine formulations, we focused on solution absorbance changes at 260 nm. We found that titration of the pDNA (P) with PEIm (N) causes continuous absorbance increase from molar N/P ratio of 0.5 to 1.5 (Figure 2, B). Above N/P ratio of 2, where the pDNA is completely condensed with the PEIm, the absorbance remained constant, indicating that no further interactions evolved between the components. As PEIm has baseline absorbance, the measured Hc can be related to the forming interactions between the pDNA and the PEIm. This might be a general phenomenon for NPs because this type of measurement has also been utilized to characterize the binding of DNA to single wall carbon nanotubes.<sup>28</sup>

The observation of the Hc was confirmed with both zeta potential titration (Figure 2, B) and AGE (Supplementary Figure 1). Above N/P ratio of 2, zeta potential remained

constant and positive and the NP showed essentially no migration on the gel (Supplementary Figure 1). Once the pDNA is totally condensed, the addition of PEIm in excess caused no changes in particle size either (Supplementary Table 3). These results suggest that the percentage Hc correlates with the number of bonds formed between the phosphate and the amine groups of the NPs’ components. The correlation was confirmed by the decomplexation of the NPs. After the decomplexation of the NPs with SDS, the Hc decreased to the absorbance of the pDNA (Figure 2, C). In the decomplexed NPs, the pDNA migrated similarly to the control pDNA, and the NPs that were not treated with SDS did not migrate (Figure 2, D).

We also investigated DV1 and DV2 formulations and found that Hc of DV1 is smaller than DV2 (Table 1), suggesting a smaller degree of association between the components. This observation is consistent with AFM images presented above and confirm that Hc provides a complementary tool to the NP size measurements for pDNA nanomedicine characterization. It is

noteworthy that the differences in the degree of association between the DV1 and DV2 formulations upon treatment of competing SDS could not be measured in freshly prepared samples by agarose gel migration (data not shown). Besides giving information on the degree of association between the pDNA and PEIm components of the nanoparticles, Hc measurement can also be utilized to describe the kinetics of the dissolution of the NPs to form freely suspended pDNA and PEIm molecules (see below). We note that the measurement of true binding constants of the complex formation by Hc, or other binding equilibrium measurement, requires the existence of equilibrium where Hc would then be constant. It is evident that the NPs can sometimes exist in a metastable state that can evolve over long periods. Hc still provides an indicator of the nature and evolution of this complex state, even if it has no simple interpretation in terms of a simple binding constant. This circumstance is actually a common physical situation and the discussion of this type of non-equilibrium binding process has been discussed extensively recently in the binding of quantum dots to collagen layers.<sup>29</sup>

#### *Influence of temperature on NP stability*

Temperature is an important factor in clinical contexts and in this study we showed additional evidence that the DV2 formulation is substantially more stable than DV1. We investigated the variation of Hc as a function of time at 3 different temperatures (4°C, 25°C and 37°C). We saw that Hc for the DV1 drifted in time as the NPs structure evolved (Figure 3, A). In contrast, the Hc values for the DV2 sample were stable in time (Figure 3, B). The increase of Hc of the pDNA nanomedicine reflects the increase of absorbance of the free pDNA especially at higher temperatures (Figure 3, C). Absorbance of the free pDNA saturated within a week; however, the Hc of DV1 did not reach that saturation. The stability of the DV1 formulation, as inferred from the Hc measurements, increased upon cooling (Figure 3, A). A similar trend was apparent in the pDNA that relates to the organization of the pDNA bases within the pDNA nanomedicine.<sup>30,31</sup> The Hc of the DV2 formulation remains constant over a period of 8 days (Figure 3, C). These stability data is consistent with the AFM measurements discussed above and suggest that the PEIm forms a robust complex with the pDNA in the DV2 formulation that shields the pDNA from the solution environment, whereas some of the pDNA in the DV1 formulation is in a state similar to the free pDNA in solution where the pDNA is potentially susceptible to degradation. Consequently, Hc provides a general parameter for “strength” of binding between the pDNA and PEIm and the evolution of the NPs, essential information in relation to controlling the stability and activity of the nanomedicine. The irregular structures characterizing the DV1 sample (Figure 1, C) were found to become progressively more disorganized with increasing “aging” time, and after a week, no well-defined NPs were detected. A similar NP structure evolution was not observed in the DV2 sample, whose structure remained essentially invariant over a period of several weeks (data not shown).

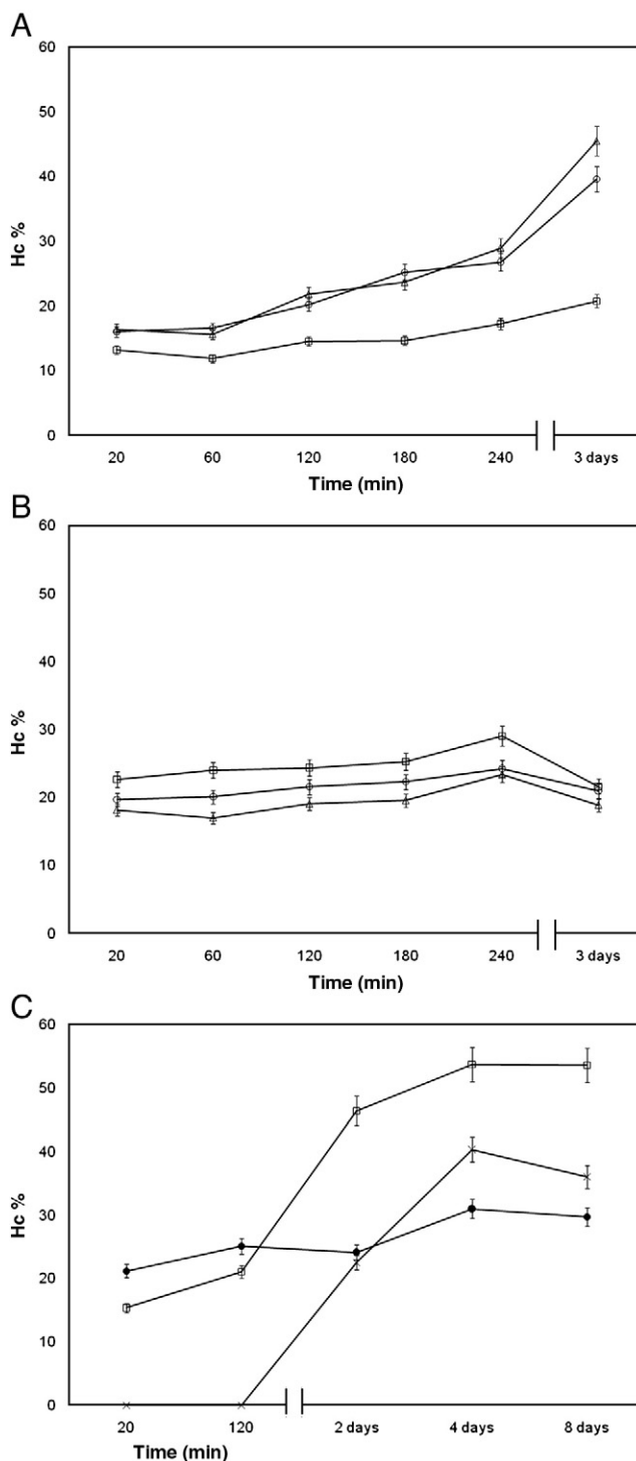


Figure 3. Investigation of pDNA/PEIm NP stability. (A), Hc of DV1 measured in different time points after NP preparation; (□) 4°C, (○) 25°C, (Δ) 37°C. (B), Hc of DV2 measured in different time points after NP preparation; (□) 4°C, (○) 25°C, (Δ) 37°C. (C), Hc of DV1 (□), DV2 (●) and control pDNA (×) measured in different time points after incubating at 37°C.

#### *Influence of pH and salt concentration on NP stability*

Next, we used Hc measurements to assess the effect of pH and salt (NaCl) concentration on NP structure. These are the

most obvious variables in the nanomedicine formulation and understanding the sensitivity of nanomedicine to these variables is crucial for the control of the nanomedicine efficacy. As the 2 studied formulations DV1 and DV2 differed in both their pH and conductivity (Table 1), we investigated how these basic parameters affect the stability of nanomedicine through Hc measurements. First, we examined the effect of pH because this variable should strongly influence the strength of the ionic interactions between the 2 polyelectrolytes (pDNA and PEIm).

We found that the Hc of the pDNA nanomedicine is directly associated with the pH of the solvent (Figure 4, A); as the pH increases, the Hc of the nanomedicine also increases. We also studied the effect of salt concentration (ionic strength) on the

nanomedicine. The pDNA was supplemented with (0.1, 1, 3, 6 or 10) molar equivalents NaCl (calculated on the phosphate concentration of the pDNA) before mixing with the PEIm solution. Increasing the ionic strength decreased Hc (Figure 4, B). Consistent results were obtained in the Hc vs. pH measurements (Figure 4, A) we obtained similar results, shown in Table 1: for DV1 formulation 13% (for pH 3.5) and for DV2 formulation 21% (for pH 7.5). The inverse correlation between Hc and NaCl concentration is also consistent with the lower Hc observed for DV1 than DV2 (Table 1) because the conductivity of the DV1 formulation is higher than for DV2 (Table 1). These results demonstrate that the morphology and property differences of the DV1 and DV2 formulations are the result of the different pH and ionic strength that determine the different degree of association of the components in the NPs and suggest that both of these parameters of the nanomedicine are important for stability.

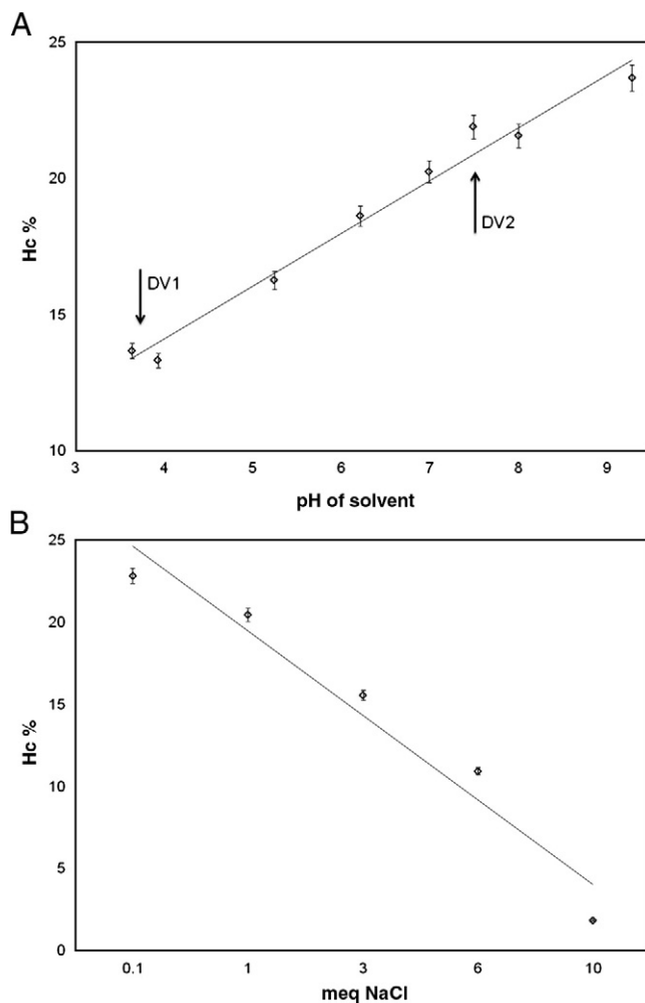


Figure 4. Effect of the ionic environment on the hyperchromicity of the pDNA/PEIm NPs. (A), Linear correlation between the pH of the solvent and the Hc% of pDNA/PEIm NPs. NP formulations were prepared using the same pDNA and PEIm at N/P ratio of 4 in different pH TEAM solutions. The pH of the DV1 and DV2 are marked with the arrows. ( $y = 1.9403x + 6.3354$ ;  $r^2 = 0.98$ ) (B), Inverse correlation between the NaCl concentration of pDNA solution and the measured Hc% of NP. Meq on the x-axis stands for molar equivalents calculated on the phosphate concentration of pDNA ( $y = -5.1448x + 29.761$ ;  $r^2 = 0.95$ ).

#### Comparison of biological activities of NP formulations

The biological activity of pDNA nanomedicine depends on the expression of the antigens encoded in the pDNA.<sup>12</sup> We found that DV2 has significantly higher biological activity than DV1 (Figure 5, A). This efficient gene expression correlates with the Hc trend as well as the higher nuclease resistance of DV2 in comparison with DV1 formulation (Table 1; Figure 5, B). Consistent with the AFM data (Figure 1) the nuclease resistance assay demonstrates that the PEIm “envelope” protects the pDNA condensed in the DV2 in contrast to DV1 where the PEIm not perfectly covers the pDNA. Nucleases are present in both extra- and intracellular space to protect the organism from DNA; therefore, we conclude that more pDNA of DV2 can enter the nucleus in contrast with DV1 where some of the pDNA is digested. This result makes intuitive sense if we consider the mechanism of action depicted on a hypothetical model on Figure 5, C. After cellular uptake, the NPs escape endosomal degradation with the help of the proton-sponge effect of the PEIm.<sup>10,32</sup> The protonable nitrogens of the PEIm backbone are able to buffer the proton-rich environment of the endosome, thus protecting the pDNA inside the NP. It is evident that the stability of the NPs in the endosome having a rather low pH requires the NPs to be relatively stable and requires the pDNA to be protected to avoid degradation. Our AFM, Hc and nuclease resistance results suggest that the NPs of the DV1 formulation might not resist the intracellular degradation process as efficiently as in the DV2 formulation. The chance of the uncompacted pDNA to become degraded in the lysosome is also evidently higher in the latter formulation. Our model is supported by recent findings showing that co-localization of PEI and the pDNA in the nuclei is required for gene expression.<sup>28,33,34</sup>

One of the advantages of the “pathogen-like” nanomedicine in comparison with viral vectors is that DermaVir is suitable for repeated administration to keep boosting the immune responses required to fight chronic infectious diseases, like HIV/AIDS.<sup>11,12,14</sup> We investigated the activation of the innate immune system with DermaVir because PEI was shown to activate the complement system.<sup>35,36</sup> DV1 and DV2 were incubated with 3 different human sera and surprisingly we found no complement activation by DermaVir nanomedicine

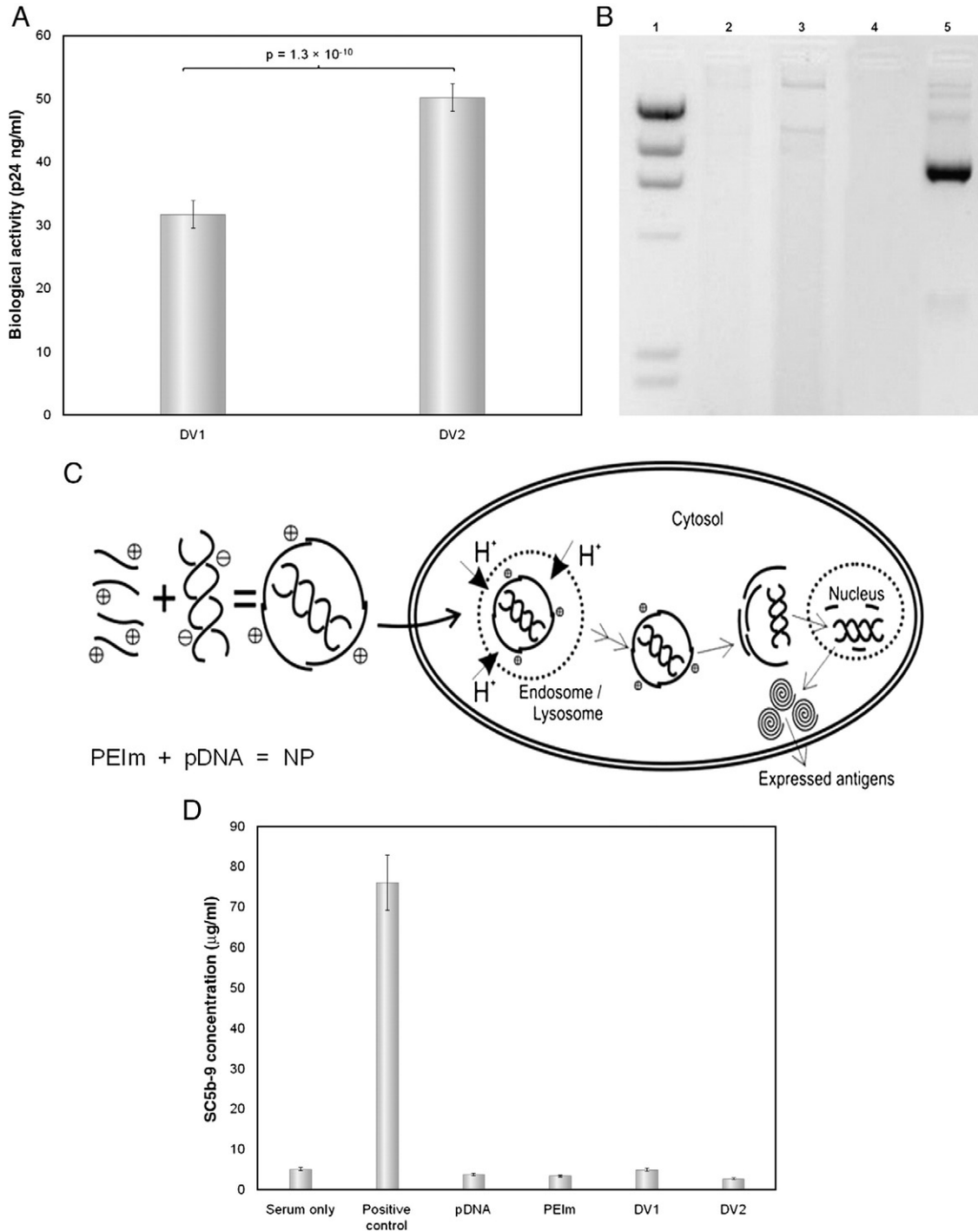


Figure 5. Effect of the hyperchromicity on gene expression. **(A)**, Biological activity of DV1 and DV2. **(B)**, Nuclease resistance of DV2 in comparison with DV1 formulation. Lanes: #1: marker; #2: DV1 digested with aspecific endonuclease and decomplexed; #3: DV2 digested with specific endonuclease and decomplexed; #4: pDNA digested with aspecific endonuclease; #5 control, untreated pDNA. **(C)**, Hypothetical model of pDNA nanomedicine uptake and antigen expression exemplified with the DermaVir HIV nanomedicine candidate. When Hc is optimal, the nanomedicine escapes from endosomal degradation, loosens in the cytosol and the pDNA reaches the nucleus, where the encoded several antigens are expressed. **(D)**, Complement activation of DV1, DV2, pDNA and PEIm. Positive control is Zymosan.<sup>22</sup>

(Figure 5, D). It is noteworthy that activation of the innate immunity by DermaVir cannot be completely excluded because of the high variation of multiple human sera on complement activation.<sup>37,38</sup>

## Discussion

We have established by both AFM and DLS measurements that the size of the pDNA nanomedicine falls in the size range of

viruses that naturally evolved to deliver genetic materials to cells (Figures 1, A–E). This led us to a strategy of directly engineering synthetic pathogen-like NPs to optimize the pDNA delivery to cells. Uptake of larger (>400 nm) sized particles and bacteria generally occurs by different cellular transport mechanisms, such as macropinocytosis and phagocytosis, whereas smaller-sized particles and viruses usually translocate into the cell by receptor-mediated endocytosis.<sup>24,25</sup> It was surprising to us that basic nanomedicine parameters like the molar ratio of the components, the supercoiled form content and the length of the pDNA polymer molecular weight did not appreciably influence the particle size of the NPs (Supplementary Tables 1, 2, 3 and 4). These facts were crucial in setting our nanomedicine design parameters. Once the particle size was in the optimal range to ensure the receptor-mediated entry into target cells, sufficient stability was required to support the release of the pDNA from the endosome to the nucleus to efficiently express antigens in the cell.<sup>39</sup>

We found that the biological activity of the pDNA nanomedicines is dependent on their inherent structure and binding between the pDNA and PEIm, as determined by AFM and Hc, respectively. There are competing requirements of having sufficient stability to escape endosomal degradation after cellular uptake and to remain intact all the way to arrive in the nucleus, where the function of NPs requires unraveling in the nuclear environment.<sup>27,34</sup> The strength of the complex formation must be carefully controlled to satisfy all these requirements and to set the degree of association of the pDNA and PEIm components.

The evaluation of Hc is based on the “constant state” of the NPs when both the N/P ratio and the particle size are fixed. The measurable Hc is linked to the degree of association with the NP. Hc should be interpreted as a parameter describing the degree of interactions between the negatively charged phosphate oxygen of pDNA and the positively charged nitrogen (secondary amine) of PEIm. Because these interactions are driven by electrostatic forces, Hc is sensitive to ionic strength and pH (Figure 4). These findings are confirmed by isothermal calorimetric (IC) titration and surface plasmon resonance (SPR) measuring the interaction between immobilized DNA and free PEI in the presence and absence of NaCl.<sup>27</sup>

The linear correlation between the Hc and the bulk pH of the nanomedicine is the result of the protonation state of its components. Theoretically, in low pH the secondary amines of the PEIm and the phosphates of the pDNA are protonated; therefore, few bonds can be formed between them. At a high pH, the pDNA becomes deprotonated and makes more interactions with the PEIm. According to Eliyahu et al, the degree of ionization of the amino group is highly dependent on the local pH (surface pH), which may differ from the bulk pH by 2 to 4 pH units.<sup>40</sup> Because measuring the local pH is not feasible, we operated with the bulk pH and Hc. We have established that the 6 to 7.5 optimal bulk pH value corresponding to a higher local surface pH value is optimal for both the thermodynamic stability and the biological activity.<sup>9</sup> The instrumental analysis methods like AFM, IC or SPR are powerful for the physical characterization of NPs, but their validation is problematic; therefore, these methods are not suitable for the quality control of clinical nanomedicinal products.

Based on our results we concluded that the use of Hc as the measure of the degree of association of the pDNA and PEIm

components is a powerful method for the determination of the NPs' inherent structure that affects the biological activity and the stability of the NPs. The approach described here should guide the clinical development of nanomedicines for the prevention and treatment of infectious, neoplastic and immunological diseases.

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## Appendix A. Supplementary data

Supplementary materials related to this article can be found online at [doi:10.1016/j.nano.2011.07.013](https://doi.org/10.1016/j.nano.2011.07.013).

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