Size of the cell nucleus and its effect on the chromatin structure in living cells.

Artem K. Efremov,1,2,∗ Ladislav Hovan,3 and Jie Yan2
1Institute of Systems and Physical Biology, Shenzhen Bay Laboratory, Shenzhen, China 518132
2Mechanobiology Institute, National University of Singapore, Singapore 117411
3School of Pharmaceutical Sciences, University of Geneva, Geneva, Switzerland 1211
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DNA-architectural proteins play a major role in organization of chromosomal DNA in living cells by packaging it into chromatin, whose spatial conformation is determined by an intricate interplay between the DNA-binding properties of architectural proteins and physical constraints applied to the DNA by a tight nuclear space. Yet, the exact effects of the cell nucleus size on DNA-protein interactions and chromatin structure currently remain obscure. Furthermore, there is even no clear understanding of molecular mechanisms responsible for the nucleus size regulation in living cells. To find answers to these questions, we developed a general theoretical framework based on a combination of polymer field theory and transfer-matrix calculations, which showed that the nucleus size is mainly determined by the difference between the surface tensions of the nuclear envelope and the endoplasmic reticulum membrane, as well as the osmotic pressure created by cytosolic macromolecules on the cell nucleus. In addition, the model predicted the existence of a previously unknown link between the cell nucleus size and stability of nucleosomes, providing new insights into the potential role of nuclear organization in shaping the cell response to environmental cues.

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I. INTRODUCTION

Homeostasis and biological functioning of living cells rely on sophisticated synergistic cooperation between multiple molecular subsystems that must coexist with each other in a tight and highly crowded cellular space. This makes the problem of space allocation to each of the cell components one of the most important for intracellular organization, especially taking into account that many of the subcellular systems have very different requirements to the surrounding microenvironment needed for their proper operation. However, at the present time, there is still no clear understanding of molecular mechanisms responsible for size regulation of the most of cellular organelles, including even the major ones, such as the cell nucleus [1–4].

Indeed, while many models with different levels of detail have been developed in the past to describe regulation of the cell volume [5–7], no similar analogue has been created so far to address the problem of the nucleus size control. As a result, such a very well-known phenomenon as correlation between the cell and nucleus volumes under a broad range of experimental conditions still remains poorly understood [8–11]. Furthermore, lack of the knowledge of the main molecular mechanisms responsible for the nucleus size regulation makes it hard to fully comprehend potential effects of various environmental factors in shaping nuclear organization.

Specifically, based on recent experimental studies it has been suggested that by modulating DNA-binding properties of architectural proteins, cells may be able to actively respond to a wide range of environmental cues via changing the condensation level of various parts of DNA and switching between different gene expression patterns [12–18]. Furthermore, it has been found that such a mechanism not only plays the central role in cell response to chemical signals [12–14], but also to extracellular mechanical forces as well [16–25]. In the latter case, rearrangement of the chromatin structure and changes in gene transcription have been found to be tightly associated with alternations in the nucleus size and shape [18, 21, 25]. Yet, despite a profound role of mechanical forces in regulation of the nuclear organization, the exact molecular mechanisms underlying it still remain unclear.

The main difficulty in understanding the role of mechanical forces in shaping the chromatin structure comes from the fact that the latter is predominantly determined by a tight interplay between several key factors, whose exact contributions, however, cannot be easily quantified

* Corresponding author e-mail: artem@szbl.ac.cn
in experimental studies. Namely, physical constraints applied by the nuclear envelope (NE) to DNA results in appearance of highly crowded microenvironment inside the cell nucleus, whose behaviour in a large part is driven by electrostatic and volume-exclusion interactions between DNA and proteins [26–32]. The strength of these interactions is determined by nucleoplasmonic concentrations of DNA and proteins, which are dictated by the nucleus size. As it is hard to reproduce similar unique microenvironment in in vitro studies with the same DNA and protein composition, the main understanding of the role of such physical interactions as well as their play with mechanical forces in chromatin organization currently can be gained only by means of theoretical modelling.

Unfortunately, extreme complexity of this task and strong limitations of the existing computational methods do not allow to run molecular dynamics simulations for long DNA molecules comparable in size to the cell genome. Even the most advanced modern mathematical models, which heavily rely on coarse-grained description of chromatin, can currently predict an approximate conformation of DNA molecules no longer than \( \sim 100 \text{ Mbp} \) in length [33–45], – about two orders of magnitude smaller than the typical size of chromosomal DNA in human cells (\( \sim 6.2 \text{ Gbp} \)) [46]. And while these models have provided many interesting insights into molecular mechanisms underlying large-scale chromatin organization, it should be noted that due to their coarse-grained nature, they have a number of limitations.

First, they do not take into consideration one of the major physical forces being involved in chromatin organization – electrostatic DNA-DNA and protein-DNA interactions, which make a significant contribution into the total free energy of chromatin, dominating over all other energy contributions made by the rest of physical and chemical factors. As a result, the role of intracellular ionic environment in chromatin organization still remains poorly understood despite experimental studies showing that it plays an important role in shaping the chromatin structure [26–30, 32, 47].

Second, use of extend segments of the chromatin fiber as elementary modelled units makes it nearly impossible to take into consideration formation and dissociation of individual nucleoprotein complexes from DNA as well as their transition between various states, – processes which play the central role in governing genome-wide changes in chromatin organization in response to environmental cues [12–14, 17, 18, 48]. Thus, capacity of the existing models to address in detail the core property of chromatin – its ability to dynamically reorganize in response to signals received from surrounding microenvironment, is compromised by application of the coarse-grained approach that lacks insights into behaviour of individual nucleoprotein complexes.

Finally, it should be noted that the possible number of ways nucleoprotein complexes, such as nucleosomes, can be positioned on the chromosomal DNA is extremely large (\( \gg 10^{12000000} \)). Since none of the existing models describing chromatin organization can deal with so many possible DNA-protein conformations, it seems that the above problem cannot be solved on the basis of currently used theoretical methods, and an alternative approach needs to be developed.

In this study, by utilizing elements of polymer field theory and statistical physics, we have constructed a general theoretical framework that allows one to address these issues and predict the size of the cell nucleus and the conformation of chromosomal DNA, taking into account DNA-protein interactions and the main physical forces responsible for nuclear organization. As a result, it has been found that the nucleus size in the general case is predominantly determined by a tug-of-war between the osmotic pressure exerted by cytosolic macromolecules on the NE and the difference between the surface tensions of the NE and ER membrane. In addition, the model showed the existence of a previously unknown link between the chromatin structure and the nucleus size, revealing a new potential molecular pathway, which may contribute to cell mechanosensing.

II. METHODS

In this work, we used the previously developed theoretical framework based on transfer-matrix calculations that allows one to compute the grand partition function of DNA in the presence of DNA-binding proteins [49–51], extending it to include in consideration physical forces discussed in Introduction. While all mathematical details can be found in Appendices A–L, here we will only outline the central idea of the study and the main model assumptions.

Packaging of long chromosomal DNA into a tiny nuclear space in eukaryotic cells is mainly done with the help of special DNA-architectural proteins, histones. These proteins assemble into positively charged octamer complexes that wrap negatively charged DNA around themselves, leading to formation of compact nucleosomes. Nucleosome assembly is performed via help of histone-binding chaperones, such as Asf1 or Nap1, which are involved in transportation of H2A-H2B and H3-H4 histone dimers into the cell nucleus [52, 53]. Correspondingly, in the model we considered two types of chaperones: 1) \( c_1 \) chaperones that can either be in empty / unloaded state (\( c_1 u \)) or histone-bound state (\( c_1 b \)), in which they form a complex with a H2A-H2B dimer, and 2) \( c_2 \) chaperones that also can be either in unloaded state (\( c_2 u \)) or histone-bound state (\( c_2 b \)), but this time forming a complex with a H3-H4 dimer, see schematic Figure 1(a). Since experimental data suggest that some histone-binding chaperones, such as Nap1, shuttle between the nucleus and cytosol in eukaryotic cells [54], in our model it was assumed for simplicity that both histone-loaded and unloaded chaperones \( c_1 \) and \( c_2 \) can move through nuclear pores without any restriction.

In the model, the cell nucleus is represented by a spher-
ically shaped NE enclosing nucleoplasm of $V_{\text{nucl}}$ volume. The nucleus itself is submerged into the cell cytosol of $V_{\text{cyto}}$ volume, which may have an arbitrary shape. Cytosol in the model serves as a buffer solution cushioning changes in the nucleoplasmic concentrations of unloaded and histone-bound chaperones that may result from rearrangement of the chromatin structure.

Besides chaperones, the cytosol also buffers any changes in the nucleoplasmic levels of Na$^+$, K$^+$ and Cl$^-$ ions, which have a major effect on the strength of DNA-histone interactions and chromatin organization [28, 30, 32]. Microelectrode measurements as well as radiographic and extractive analysis suggest that these monovalent ions can rather freely move between the cell nucleus and cytosol [55, 56]. Furthermore, their cytosolic concentrations are usually kept at nearly constant levels by transmembrane ion pumps and ion channels to maintain electroneutrality of intracellular environment and at the same time to counterbalance osmotic pressure created by the cell metabolites and proteins onto the cell membrane [6, 7, 57]. Following previously published experimental and theoretical studies [5, 57, 58], the total cytosolic concentrations of monovalent Na$^+$ and K$^+$ ions on one hand, and Cl$^-$ ions and negatively charged cell metabolites on the other hand were both put equal to $c_{\text{ions}} = 150$ mM in all our calculations.

As for the chromosomal DNA, it was modelled as a set of polymers constrained inside the cell nucleus, whose total number ($Q$) matches the number of chromosomes in the studied eukaryotic cells (in human cells $Q = 46$), with the total length of DNA pieces ($L = \sum_{u=1}^{Q} L_u$) being equal to the total size of the cell genome. Here $L_u$ is the length of the $u$th DNA polymer comprising the $u$th chromosome. Thus, to describe chromatin organization in human cells the total length of all DNA polymers was set equal to $L = 6.2$ Gbp in all calculations.

Three-dimensional conformations of the DNA polymers were represented in the model by $R^{(u)}(s_u)$, $u = 1, \ldots, Q$ functions, where $s_u \in [0, L_u]$ is the arc length along the $u$th DNA polymer and $R^{(u)}(s_u)$ is a three-dimensional Euler rotation matrix assigned to each point residing on the contour of the $u$th chromosome, which is a function of the arc length, $s_u$. This matrix indicates orientation of the DNA backbone at the corresponding point with respect to the global coordinate system, $(x_0, y_0, z_0)$, such that the unit vector $x^{(u)}(s_u) = R^{(u)}(s_u)x_0$ resulting from the rotation of $z_0$-axis of the global coordinate system via Euler matrix $R^{(u)}(s_u)$ is tangential to the DNA backbone; whereas, the other two unit vectors, $y^{(u)}(s_u) = R^{(u)}(s_u)y_0$ and $z^{(u)}(s_u) = R^{(u)}(s_u)z_0$, are normal to the DNA backbone, keeping track of the DNA twist angle, see schematic Figure 1(b).

Nucleosomes formed on the chromosomal DNA were represented in the model by solenoid-like structures schematically shown in Figure 1(b), whose geometry matches that of DNA wrapped around histone octamers in nucleosome complexes [59, 60].

To find out how confinement of DNA inside the cell nucleus as well as electrostatic interactions between the system components affect the chromatin conformation, we used statistical physics approach by calculating the partition function of the system, $Z$. Indeed, while living cells usually operate far from thermodynamic equilibrium, previous studies suggest that DNA interaction with proteins still can be accurately described by equilibrium thermodynamics since assembly of nucleoprotein complexes on DNA is frequently governed by DNA-binding affinities and positioning entropies of the proteins [61–65]. Furthermore, it has been shown in previous theoretical studies that methods of statistical physics seem to correctly predict positions of nucleosomes on the chromosomal DNA near transcription start sites as well as on the global genomic scale [66–70], suggesting that they can be used to provide important insights into organization of chromosomal DNA in living cells.

In the general case, the partition function, $Z$, of the system of DNA polymers described above equals to (see Appendix A for details):

$$Z = \sum_{\text{DNA-protein}} \prod_{u=1}^{Q} \frac{1}{n_j} \int_{V_{\text{nucl}}} \frac{Z^{in}_{dr_{jk}}}{\Lambda_j^3} \times$$

$$\times \prod_{u=1}^{Q} \left[ \int_{V_{\text{cyto}}} d\mathbf{r}_j \langle \mathcal{S}\mathbf{R}^{(u)}(\mathbf{r}) e^{-\beta E_{\mathbf{r}}(\mathbf{R}^{(u)})} \rangle \right]$$

$$\times e^{-\frac{\beta}{2} \int_{V_{\text{cyto}}} d\mathbf{r}_j \int_{V_{\text{nucl}}} d\mathbf{r}_{j'} \rho_{ij}(\mathbf{r}_j)(\mathbf{r}-\mathbf{r}')(\mathbf{r}_{j'})}$$

(1)

Here $\sum_{\text{DNA-protein}}$ is the sum over all possible ways, in which nucleosomes can be positioned on chromosomal DNA. $\prod_{u=1}^{Q}$ is the product over all types of particles (ions, proteins, cell metabolites, etc.) diffusing inside the cell, which are enumerated by index $j$, $n_j$ is the total number of particles of type $j$ inside the cell. $\prod_{u=1}^{Q}$ is the product over all particles of the same type, $j$, whose positions in space are described by the radius vectors $\mathbf{r}_{jk}$. Integration $\int_{V_{\text{cyto}}} d\mathbf{r}_j$ in the above formula is performed over the volume $V_{\text{cyto}}$ accessible to particles of type $j$. In the case of ions, small proteins and cell metabolites, $V_{\text{cyto}}$ corresponds to the union of the cell nucleus and cytosol volumes: $V_{\text{cyto}} = V_{\text{nucl}} \cup V_{\text{cyto}}$; whereas, in the case of macromolecular complexes that cannot move through nuclear pores into the nucleoplasm, we have $V_{\text{cyto}} = V_{\text{cyto}}$. $\Lambda_j$ and $Z^{in}_{dr_{jk}}$ are thermal de Broglie wavelength and the partition function describing inner degrees of freedom of the $j$th type of particles, respectively. $\prod_{u=1}^{Q}$ is the product over all chromosomes confined inside the nucleus, whose 3D conformations are represented by $\mathbf{R}^{(u)}$ functions. $\mathbf{r}_0^{(u)}$ is the radius vector describing position of the starting end of the DNA polymer comprising the $u$th chromosome. Integrals $\int \mathcal{S}\mathbf{R}^{(u)}$ are Feynman-like path integrals along the contours of the respective DNA polymers ($u = 1, \ldots, Q$), which are calculated over all possible chromosome conformations. $E_{\mathbf{r}}(\mathbf{R}^{(u)})$ is the energy of the DNA polymer comprising the $u$th chromosome, which incorporates elastic deformations of the DNA as well as its interaction with DNA-binding proteins (histones), see details in
Appendices A-B. In the general case, $E_w$ energy is determined by the 3D conformation of the $u^{th}$ chromosome, which is described by $R^{(u)}(s_{nu})$ function. $\beta = 1/k_B T$ is the Boltzmann constant and $T$ is temperature of the surrounding environment. $U_{sp}\left(r, r'\right) = \frac{1}{\varepsilon r}$ is the core part of the electrostatic potential, where $\varepsilon$ is the permittivity of intracellular media, which in the model was put equal to water permittivity. Finally, $\rho_e(r)$ is the distribution of electrical charges inside the cell, which is described by Eq. (A4) (Appendix A). More details regarding the above mathematical expression as well as treatment of the volume-exclusion effect can be found in Appendices A, L.

To calculate the partition function given by Eq. (1), in this study we employed the mean field approach by considering randomly fluctuating electrostatic field $\psi$ created by electrically charged ions, metabolites and DNA, based on which it can be shown that Eq. (1) can be reduced to the following simple formula (see Appendix A for details):

$$
\ln Z = \ln Z_{\psi sp} - \beta \sum_j n_j \mu_{j\psi sp}
$$

(2)

Where $Z_{\psi sp}$ and $\mu_{j\psi sp}$ are the partition functions of DNA and electrochemical potential of the $j^{th}$ type of particles in the presence of the stationary phase electrostatic field, $\psi_{sp}$, which solves the following functional equation (see Appendix A):

$$
\frac{1}{4\pi \beta r^2} \frac{\delta \ln Z_\psi}{\delta \psi(r)} + 2q_e \varepsilon_{\text{Gou}} \sinh[\beta q_e \psi(r)] - \varepsilon \Delta \psi(r) = 0
$$

(3)

Here $r$ is the radial distance measured from the center of the cell nucleus. $q_e = 1.6 \cdot 10^{-19}$ C is the elementary charge. $Z_\psi$ is the partition function of DNA interacting with histone octamers in the presence of electrostatic field $\psi$, see Appendices B-D. $\frac{\ln Z_\psi}{\rho e(r)}$ is the functional derivative of the DNA partition function, $Z_{\psi}$, with respect to the electrostatic field, $\psi$, see Appendix E.

To solve Eq. (3), electrostatic field $\psi(r)$ was represented in our study in a form of the following Fourier-Bessel expansion series:

$$
\psi(r) = \frac{\psi_0}{2} \left[ 1 - \tanh \left( \frac{r-R_{\text{nucl}}}{w} \right) \right] + \sum_{n=1}^{n_{\text{max}}} \psi_n j_0 \left( \frac{\pi n r}{R_{\text{nucl}}} \right)
$$

(4)

Where $R_{\text{nucl}}$ is the radius of the cell nucleus. $n_{\text{max}}$ is a number of Fourier-Bessel modes used in calculations (in this study, $n_{\text{max}} = 18$). As for $R_0$, $w$, $\psi_0$, $\psi_1$, ..., $\psi_{n_{\text{max}}}$, these are the model fitting parameters, whose values were determined by Nelder-Mead simplex algorithm [71], which was used to minimize deviation of the left side of Eq. (3) from zero, see Appendix A for details.

Finally, to calculate the partition function of chromosomal DNA, $Z_{\psi}$, in an arbitrary potential field $\psi$, we used a discretized semiflexible polymer chain model in which DNA is partitioned into smaller straight segments, whose size is much smaller than the bending persistence length of DNA ($A = 50$ nm [72, 73]), see schematic Figure 1(c). Since the elastic bending and twisting energies of DNA can be described based on relative orientations of neighbouring DNA segments, it is then possible to utilize transfer-matrix calculations to obtain the partition function of DNA, see Appendices B-E. As a result, it can be shown that up to a non-essential fixed prefactor, $Z_{\psi}$ can be represented in the following form:

$$
Z_{\psi} \propto \prod_{u=1}^{Q} \left[ UL^{N_u-1} Y \right]
$$

(5)

Where $N_u$ is the total number of segments in a discretized polygonal chain representing the DNA polymer comprising the $u^{th}$ chromosome. $U$ and $Y$ are boundary condition vectors that depict physical states of the DNA end segments in each chromosome. $L$ is a transfer-matrix, which characterizes DNA segments’ interactions with DNA-binding proteins and surrounding potential field $\psi$, as well as describes local bending and twisting elasticities of DNA.

The main advantage of Eq. (5) is that the value of each matrix product, $UL^{N_u-1} Y$, in this equation is mainly determined by the dominant eigenvalue ($\lambda_{\text{max}}$) of the transfer-matrix $L$ when $N_u \gg 1$, which is the case for long DNA molecules found in eukaryotic cells. Specifically, it can be shown that $Z_{\psi} \approx C_0 \lambda_{\text{max}}^{N_u}$, where $N_{\text{tot}} = \sum_{u=1}^{Q} N_u$ is the total number of DNA segments in all of the chromosomes, and $C_0$ is a constant. Since both $C_0$ and the dominant eigenvalue $\lambda_{\text{max}}$ can be found with the help of standard techniques, such as the power iteration method [74] (also see Appendices E, I), this makes it possible to calculate the partition function of an arbitrarily long DNA in the presence of DNA-binding proteins in solution and electrostatic interactions between all of the system components – a task, which cannot be handled by any of the existing theoretical methods.

By taking derivatives of the partition function logarithm defined by Eq. (2) with respect to various model parameters, such as the volume of the cell nucleus or histones binding energy to DNA, etc., it is possible to find the mean pressure created by DNA on the NE, the average DNA conformation and occupancy by nucleosomes, DNA distribution inside the nucleus, and many other quantities characterizing the physical state of chromosomal DNA and the cell nucleus. The full list of these quantities and formulas that were applied to calculate them can be found at the end of Appendix I. The rest of the model parameters as well as their values used in the calculations are listed in supplementary Table I.

The source code of the programs that were used to compute the results presented below can be downloaded from http://www.artem-efremov.org website.

Finally, it should be noted that while here we have only described the model approach to study of DNA organization in nuclei of living cells, the same method, with
III. RESULTS

To test Eq. (5), we first performed calculations for a simple non-linear system – a mechanically stretched bare DNA confined inside a nano-channel in the absence of DNA-DNA interactions, comparing obtained results to Metropolis-Monte Carlo (MMC) simulations, see Figure 2. From the figure it can be seen that the transform-matrix approach based on Eq. (5), where $\psi$ field was set to an infinite cylindrical potential describing the walls of the nano-channel, accurately depicts DNA behaviour up to the point where the channel radius becomes comparable to the size of DNA segments in the discretized polymer model. These results suggest that Eq. (5) is able to accurately predict behaviour of DNA molecules in highly non-linear potential fields.

Next, we applied the developed theoretical framework to description of a more complex process – DNA packaging in a viral particle in the presence of electrostatic DNA-DNA and ion-DNA interactions. By varying the size of the viral capsid, it has been found that the free energy of a DNA-loaded viral particle, $G$, changes almost as an inverse cubic function of the capsid radius (i.e., $G \propto R_{\text{vir}}^{-2.8}$), deviating from this trend only when the particle size reaches the value comparable to the DNA bending persistence length (50 nm [72, 73]). At this point the slope of the free energy curve becomes flatter, which may seem to be counter-intuitive as one expects that at this point DNA will start to apply more pressure on the capsid wall due to accumulation of the DNA elastic bending energy, see Figure 3(a). However, additional calculations performed for a polymer with zero bending and twisting elasticities showed nearly the same results [see dashed curves in Figure 3(a)], suggesting that the elastic bending and twisting energies of DNA make rather negligible contribution into the total free energy of a viral particle.

Indeed, by estimating other energy terms, such as electrostatic DNA-DNA and ion-DNA interactions as well as DNA-DNA volume exclusion effect, it has been found that the vast majority of the viral particle free energy comes from electrostatic interactions between the system components, which are by $\sim 1 - 2$ orders of magnitude stronger than the DNA-DNA volume-exclusion effect, see Figure 3(c) and Appendix L. This result is in good agreement with previous experimental studies showing that DNA-DNA electrostatic interactions make the dominant contribution to the viral particle energy [75].

Calculations further demonstrated that the mean electrostatic field inside viral particles is rather uniform. At the same time the absolute value of the electrostatic field appeared to rapidly increase with decreasing size of the viral capsid, $R_{\text{vir}}$, see Figures 3(d-e). At $R_{\text{vir}} = 50$ nm the electrostatic potential was found to reach the value of $\sim 20$ mV, at which the energy associated with movement of monovalent ions between the inner space of the capsid and outside environment becomes comparable to the thermal energy ($1 k_{\text{B}} T \approx 27$ e-mV). Beyond this point positively charged ions start to accumulate inside the capsid, enhancing electrostatic screening effect of DNA-DNA interactions. As a result, the system free energy grows slower at smaller values of the capsid size, explaining unusual behaviour of the energy curves at $R_{\text{vir}} \leq 50$ nm shown in Figure 3(a).

To compare the above results to experimental data, we next calculated derivatives of the ions and DNA free energy terms with respect to the capsid volume to estimate pressures generated by these components on the wall of the viral capsid, see Figure 3(b). From the figure it can be seen that overall behaviour of the pressure curves seems to be similar to that of energy curves displayed in Figure 3(a), with the only difference being much steeper slopes of the pressure curves.

Previous experimental studies indicate that the amount of pressure applied by viral DNA to the capsid wall can potentially reach a very high value of $\sim 10 - 40$ atm [76, 77], see the green rectangle in Figure 3(b) marking experimentally measured ranges of the pressure inside $\lambda$-phage particles and their sizes. By looking at Figure 3(b), it can be seen that the theoretically predicted DNA and net pressure curves pass exactly through the center of the experimentally measured range, suggesting that our model provides accurate description of the physical properties of packed viral DNA.

To further test the model, we calculated the correlation between unit vectors tangent to the DNA backbone as well as the root mean squared distance (RMSD) between two points residing on DNA as functions of the genomic distance along the DNA, see Figures 3(g-h). It has been found that the DNA conformation becomes more and more ordered with decreasing size of the viral capsid, and starting from $R_{\text{vir}} \approx 50$ nm the correlation function takes the form of damped oscillations [see Figure 3(g)], indicating that DNA becomes folded into a coil-like structure, as illustrated in schematic Figure 3(f). This result is in good agreement with existing experimental data [78, 79] and theoretical simulations [80]. RMSD plot shown in Figure 3(h) further confirms this observation, suggesting that our model provides rather accurate description of DNA packaging in viral particles.

Interestingly, RMSD graphs shown in Figure 3(h) look very similar to those reported in a recent study of polymer molecules confined in 2D spaces, indicating strong similarity between polymers’ behaviour in 2D and 3D cases, see ref. [81].

Altogether, the above results demonstrate that the developed model correctly describes behaviour of DNA confined in a tight space, making it suitable for study of DNA packaging in nuclei of living cells as described in Methods. In this work, we used this approach to gain understanding of molecular mechanisms responsible for regulation of the cell nucleus size and its potential downstream ef-
fects on the chromatin structure, such as changes in the strength of DNA-protein interactions.

In the general case, the nucleus size is determined by mechanical equilibrium corresponding to a net zero pressure acting on the nuclear envelope (NE). The latter includes the pressure created by chromosomal DNA (pDNA) as well as the osmotic pressure resulting from the gradients of electrically charged ions and metabolites across the NE (pions), which develop due to the mean negative electrostatic potential of the cell nucleus with respect to the cytoplasm, see schematic Figure 4(e). Cytosolic macromolecular compounds that cannot move through nuclear pores into the nucleus, have also been previously shown to be involved in generation of outside osmotic pressure on the NE, pmacro [82, 83]. In addition, it has been suggested that lamin proteins, which polymerize into a dense lamina network underlying the NE [84, 85], may be responsible for the nucleus growth during the cell cycle in metazoan cells [86, 87]. On the other hand, proteins binding to the membrane of endoplasmic reticulum (ER) have been found to slow down this process [88]. As a result, it has been hypothesized that the NE and ER may be involved in a tug-of-war for the shared lipid membrane [1], which from a physical point of view can be described in terms of opposite pressures, pNE and pER, acting on the NE created by the surface tensions of the NE and ER membrane, σNE and σER, see Appendix K for details.

Summing all the above pressures, it is easy to obtain the following formula for mechanical equilibrium of the NE:

\[ p_{\text{DNA}} + p_{\text{ions}} + p_{\text{macro}} + p_{\text{NE}} + p_{\text{ER}} = 0 \]  

(6)

It should be noted that previously it has been proposed that the cell cytoskeleton may also exert mechanical forces on the nucleus, affecting its size and shape [89]. However, experimental measurements demonstrate that a typical mechanical stress developed by the cell cytoskeleton is generally small (∼ 20 – 1000 Pa [11, 90]) in comparison to other pressures acting on the NE (see results below). Furthermore, it has been shown that disruption of the actin cytoskeleton by cytochalasin D does not influence osmolarity-induced change in the cell nucleus size [83]. Altogether, these results indicate that while the cell cytoskeleton may develop strong enough forces to affect the cell nucleus shape, it is unlikely to be involved in the nucleus size / volume regulation via generating pressure on the NE. Though, under certain conditions, it may have an indirect effect on the nucleus size / volume, affecting its size and shape, such as changes in the relative contributions of various cell components into the total positive and negative pressures acting on the NE [Figure 4(b) and Figure S1(b)] it can be concluded that neither DNA, nor ions make a substantial contribution in a wide range of the model parameters. Only in the case when the nucleus is compressed strong enough, the model predicts that DNA starts to produce sufficiently large repulsing force counteracting the applied pressure, which is in good agreement with existing experimental studies showing that DNA does not directly influence the nucleus size, rather setting its minimum possible value [10, 82, 93].

Thus, in the most cases, an approximate equilibrium radius of the nucleus, Rnucl, can be found as a unique real-valued solution of the following cubic equation that can be derived from Eq. (6) (see Appendix K for details):

\[ p_{\text{NE}} + p_{\text{ER}} + p_{\text{macro}} = 0 \Rightarrow \frac{4\Delta \sigma}{R_{\text{nucl}}} = \frac{2\eta_{\text{macro}}k_BT}{V_{\text{cell}}} - 3\frac{\sigma_{\text{NE}}}{\zeta_{\text{ne}}} - \frac{\sigma_{\text{ER}}}{\zeta_{\text{er}}}, \]  

(7)

and thus

\[ R_{\text{nucl}} = \sqrt[3]{-\frac{2}{3} + \sqrt{\frac{2}{3} + \frac{27}{4} \frac{\eta_{\text{macro}}k_BT}{V_{\text{cell}}} - 3\frac{\sigma_{\text{NE}}}{\zeta_{\text{ne}}} - \frac{\sigma_{\text{ER}}}{\zeta_{\text{er}}}}}. \]  

(8)

Where \( \Delta \sigma = \sigma_{\text{ER}} - \sigma_{\text{NE}} \) is the difference between the surface tensions of the NE and ER membrane. \( V_{\text{cell}}^{\text{osm}} = V_{\text{nucl}} + V_{\text{cyto}} \) is the osmotically active volume of a cell, which typically occupies ∼ 70% of the total cell volume, \( V_{\text{cell}} ^ { \text{osm} } \approx 70\% \cdot V_{\text{cell}} \) [11, 94]. \( \eta_{\text{macro}} \) is the total number of macromolecules in the cell cytosol. Taking into account that most of the macromolecules are either sufficiently large proteins or protein complexes that cannot move through nuclear pores, it is natural to expect that their number is approximately proportional to the total number of proteins in a living cell:

\[ \eta_{\text{macro}} \approx \zeta_{\text{ne}} n_{\text{pro}} = \zeta_{\text{cell}} V_{\text{cell}}. \]  

Here \( \zeta \) is a proportionality coefficient (0 < \( \zeta < 1 \)), and \( c_{\text{pr}} \) is the average protein concentration in living cells: \( c_{\text{pr}} = 2.7 \cdot 10^6 \text{ proteins/\mu m}^3 \) [95]. Finally, q and p coefficients in Eq. (8) are:

\[ q = -\frac{3}{4} \frac{\sigma_{\text{NE}}}{\zeta_{\text{ne}}} \]  

and

\[ p = \frac{3\eta_{\text{macro}}k_BT}{16\pi\Delta \sigma} = \frac{3\zeta_{\text{cell}} V_{\text{cell}} k_BT}{16\pi \Delta \sigma}. \]

We then used Eq. (7) to estimate \( \Delta \sigma / \zeta \) ratio based on experimentally measured volumes of different types of cells and their nuclei [8, 9, 11], see Figure 4(d). From the figure it can be seen that \( \Delta \sigma / \zeta \) ratio changes almost linearly with the cell volume in all considered cases. As a result, by fitting each of the data set shown in Figure 4(d) to a linear function and then substituting this function into Eq. (8), a nearly perfect fit of the experimentally measured correlations between the cell and nucleus volumes could be achieved, see Figure 4(e). Fitting of the experimental data to the full model described by Eq. (6) led to very similar results [Figure 4(e)], indicating robustness of Eq. (7)-Eq. (8).
Interestingly, the calculated values of $\Delta \sigma / \zeta$ ratio appeared to be of the order of $\sim 20 - 40$ mN/m [Figure 4(d)], which is close to the elastic modulus of the NE measured in isolated nuclei of Xenopus laevis oocytes ($28 \pm 8$ mN/m [82]). This result suggests that the nuclear lamina network likely plays a major role in governing the nucleus size regulation in Xenopus oocytes, – a model prediction, which is in good agreement with previous experimental studies [86, 87].

By varying $\Delta \sigma / \zeta$ ratio in Eq. (8), we have also found the difference between the surface tensions of the NE and ER membrane that has to be maintained by a living cell in order to retain the nucleus size at a specific value, see Figure S1(c). It can be seen from the figure that bigger nuclei require considerably larger difference between the surface tensions of the NE and ER membrane, which corresponds to a high surface free energy density of the order of $\Delta \sigma / \zeta \approx 1 - 10$ kBT/nm$^2$. This observation further stresses importance of the nuclear lamina network as a potential power source driving the cell nucleus growth both throughout the cell cycle [86, 87] and during the cell differentiation process [96], see also Discussion below.

Since electrostatic interactions seem to play a major role in DNA organization, we next checked how the size of the cell nucleus affects its mean electrostatic potential with respect to the cytoplasm. From the results shown in Figures 5(a-b) it can be seen that this potential experiences a rapid increase with the decreasing size of the nucleus, reaching the value of several millivolts, which is within the experimentally measured range of the nuclear potential in Xenopus oocytes, MDCK cells and isolated murine pronuclei ($\sim$ from $-10$ mV to 0 mV [56, 97, 98]).

As can be seen from comparison of the solid curves in Figure 5(f) plotted by using Eq. (9) and the data points obtained by fitting of the force-extension curves shown in Figure 5(d), the above formula accurately predicts behaviour of the binding free energy of histone octamers to DNA as a function of the nucleus size.

Interestingly, from Eq. (9) it can be seen that not only the electrical charges of histones are important for stabilization of nucleosomes by the nuclear electrostatic potential, but also electrical charges of histone-binding chaperones responsible for their transportation. Indeed, simple model calculations based on Eq. (9) show that depending on the electrical charge of histone-binding chaperones, a living cell can switch between two regimes: 1) nucleus size-sensing, in which the binding free energy of histone octamers to DNA depends on the nucleus size, and 2) nucleus-size insensitive, in which the binding free energy of histone octamers to DNA is independent of the nucleus size, indicating existence of a previously unknown potential regulatory pathway of nucleosome stability.

Altogether, the above results suggest that contingent on physical properties of histone-binding chaperones, living...
ing cells may have very diverse response to changes in the nucleus size in terms of the chromatin structure, whose reorganization may be driven by variation in the average strength of the nuclear electrostatic potential.

### IV. DISCUSSION

In this study, we have developed a general theoretical framework aimed at description of DNA packaging in viral particles and living cells, which allowed us to address the question of nuclear organization by taking into account main physical forces contributing to it. As a result, it has been found that the nucleus size in eukaryotic cells is predominantly determined by a tug-of-war between the osmotic pressure created by cytosolic macromolecules and the NE and the difference between the surface tensions of the NE and ER membrane, see Figure 4 and Figure S1. This finding is in good agreement with previous experimental studies showing that cytosolic macromolecules and lamins play a central role in the nucleus size regulation in metazoan cells [3, 82, 83, 86, 87, 101, 102].

It should be noted that although there may exist other molecular mechanisms which might contribute to the nucleus size regulation, it seems that the tug-of-war between cytosolic macromolecules, NE and ER will still play the central role in this process. For example, it is known that yeasts do not have lamin analogues, and thus their nuclei do not possess a lamina network. Because of that, these cells have to rely on alternative mechanisms to regulate volumes of their nuclei, for instance, through accumulation of nuclear proteins, which may be used by cells to build up internal osmotic pressure on the NE counterbalancing the outside pressure created by cytosolic macromolecules [4]. In addition, despite the lack of lamins, yeasts may create a difference in the surface tensions of the NE and ER membrane, $\Delta \sigma$, by maintaining a slightly distinct lipid composition of the NE and ER membrane [103, 104]. In both cases, the equilibrium nucleus size still will be described by the same Eq. (6)-Eq. (8), in which $n_{\text{macro}}$ needs to be simply replaced by $n_{\text{macro}} - n_{\text{nuclear}}$, where $n_{\text{nuclear}}$ is the total number of macromolecules inside the cell nucleus.

Recently, it has been proposed that the cell cytoskeleton may also contribute to regulation of the nucleus size in mammalian cells via application of mechanical forces to the NE through nesprin proteins connecting it to actin filaments [105]. However, it should be noted that independent experimental studies show that the actin network typically generates rather small mechanical stress ($\sim 20 - 1000$ Pa [11, 90]) in comparison to pressures developed by other cell components on the NE (see Figure 4). Thus, while mechanical forces created by the cell cytoskeleton are strong enough to induce changes in the nucleus shape, it seems unlikely that they may cause considerable perturbations in the nucleus volume, leading instead only to small fluctuations of the nucleus size of the order of tens to hundreds of nanometers, which indeed have been observed in recent experimental studies [106, 107].

On the other hand, the role of the cell cytoskeleton in regulation of the average level of lamin A/C and nucleocytoplasmic shuttling of histone-modifying enzymes has been documented in several previous studies [18, 89, 91, 92], suggesting that it may have rather an indirect effect on the nucleus size by modulating $p_{\text{NE}}$ and $p_{\text{DNA}}$ terms in Eq. (6).

Interestingly, existing experimental data indicate that abnormal expression of lamins as well as their defective localization frequently result in aberrant nuclear sizes and anomalies in the nuclear-to-cell volume ratio (i.e., N/C ratio), both of which strongly correlate with development of human-related diseases, such as certain types of cancers [108–110]. Based on this and other observations, it has been previously suggested that there may exist a unique connection between the nucleus size, the condensation state of chromatin and the transcription level of various genes, leading to hypothesis of the nucleus size-dependent regulation of gene transcription [15, 18, 26, 29, 93, 111]. However, possible molecular mechanisms that might be involved in such regulation remain unclear.

To gain insights into the above problem, we used the developed theoretical approach to predict a potential effect of the nucleus size on the chromatin structure. As a result, it has been found that by influencing the average density of negatively charged DNA, the nucleus size determines the mean electrostatic potential of the cell nucleus with respect to the cytoplasm, which in turn defines stability of nucleosomes as well as the average DNA occupancy by them, see Figures 5, S2 and S3. Since in a recent experimental study it has been shown that stability of nucleosomes bound to promoter regions has a strong impact on the transcriptional level of the downstream genes [112], these results point to a possibility that model-predicted changes in nucleosome stability and the average DNA occupancy level in response to the nucleus size variations may have a profound effect on the global gene expression profile.

Indeed, in ref. [93, 111], it has been found that the nucleus size and N/C ratio strongly correlate with the total amount of RNA molecules in living cells as well as their synthesis rate by RNA polymerase II. This observation is further supported by a recent experimental study in which it has been demonstrated that application of external mechanical pressure on NIH 3T3 cells leads to decrease in the nucleus volume accompanied by condensation of chromatin and subsequent decrease in transcriptional activity of RNA polymerase II [18]. And while it has been suggested that such downregulation of RNA polymerase II takes place due to shuttling of MRTF-A transcription cofactor to the cell cytoplasm [18], simple estimations based on our model indicate that this phenomenon may actually have a deeper physical origin.

Namely, based on the results presented in Figure 5(f), it can be shown that pressure-induced changes in the nu-
ucleus volume measured in ref. [18] lead to $4 - 5 \times k_BT$ increase in the binding free energy of histone octamers to DNA. As suggested by our recent theoretical study [65], this amount of free energy may be enough to tip the balance in DNA-binding competition between transcription factors and histone octamers in favour of nucleosome formation, resulting in release of transcription factors from the chromosomal DNA and global downregulation of gene transcription. Thus, our model predicts that the observed correlation between N/C ratio and RNA synthesis rate in living cells may be caused by physical mechanisms related to stabilization of nucleosomes in a nucleus size-dependent manner, providing new insights into this interesting and important phenomenon.

It should be noted that the above increase of $4 - 5 \times k_BT$ in the binding free energy of histone octamers to DNA in response to pressure application to living cells is caused by a rather mild reduction of the cell nucleus volume by $\sim 20 - 30\%$ [18]. At the same time, it has been previously reported that nuclei of living cells experience significantly larger volume increase by $\sim 100 - 200\%$ (i.e., by 2 – 3 times) during G1 phase of the cell cycle [87, 113, 114], which, according to our model, leads to a profound drop in the binding free energy of histone octamers to DNA by $\sim 8 - 11 \times k_BT$. Therefore, expansion of nuclei observed during G1 phase may have a considerable impact on the condensation state of chromatin and the average synthesis rate of RNA, which may help cells to faster produce and double all their cytoplasmic content in preparation for cell division.

Similarly, the nucleus size-dependent regulation of nucleosome stability may also play an important role during cell differentiation process. Indeed, as cells undergo differentiation, they need to synthesize a large amount of proteins to fulfil their new biological functions. Thus, one might expect that differentiated cells on average require more intensive production of multiple RNAs in comparison to stem cells, which, according to ref. [93, 111] and our study, can be achieved by increase in the nucleus size and/or N/C ratio. This hypothesis is supported by experimental findings showing that differentiation of human embryonic stem cells is indeed accompanied by expansion of their nuclei [96], which may be potentially caused by nuclear accumulation of lamins A/C – a phenomenon that has been found to take place not only in differentiating human cells, but also in cells from other animal species [96, 115, 116].

Interestingly, among different proteins that constitute the nuclear lamina network, lamin A has the strongest effect on the nuclear lamina architecture and its elastic properties [47, 85]. Furthermore, it has been revealed that lamin A is expressed in living cells in a mechanosensitive way, enhancing matrix-directed stem cell differentiation [117]. This observation together with our findings raise an interesting possibility of a previously unknown pathway of gene transcription regulation. Namely, from the results published in ref. [25, 117], it follows that the cell shape and/or mechanical properties of the surrounding environment have a profound effect on the expression level of lamin A, which, in turn, has a strong influence on the elastic properties of the nuclear lamina network and therefore the surface tension of the NE [47]. The latter, according to our study, determines the nucleus size and, as a result, nucleosome stability, which may have a global impact on gene transcription profiles.

Existence of such a molecular link between expression of lamin A and the global gene transcription profile is supported by a recent study [17]. In this work, it has been demonstrated that incubation of mouse fibroblasts on micropatterned substrates causes nuclear reprogramming in the absence of any exogenous reprogramming factors, which is accompanied by downregulation of lamin A and changes in 3D chromatin organization, with a systematic progression of cells from mesenchymal to stem cell transcriptome.

Remarkably, our model suggests that such a nucleus size-dependent regulation may in fact be quite selective, involving only a certain set of genes instead of the whole cell genome. Specifically, the model shows that stabilization of nucleosomes in a nucleus size-dependent manner is strongly affected by electrical charges of histone-binding chaperones, see Figure 6. Thus, by using histone-binding chaperones or chaperone-related cofactors with different electrical charges, living cells may be able to have several alternative regulatory pathways, some of which might be nucleus size-sensing, while others not, making it possible for cells to precisely fine-tune stability of nucleosomes at specific locations on the chromosomal DNA in response to nucleus size variations. Such a biological function of histone-binding chaperones and their cofactors can be performed in cooperation with histone-modifying epigenetic factors [118], and/or importins and exportins, which regulate nucleocytoplasmic trafficking of proteins [119], including histone-binding chaperones [53, 120, 121] and lamins [101].

Furthermore, from Eq. (9) it can be seen that not only the nuclear electrostatic potential and electrical charges of histones and chaperones contribute to the binding free energy of histone octamers to DNA, but also the loading ratios of histone-binding chaperones, indicating that molecular pathways responsible for histone synthesis and degradation may as well affect nucleosome stability.

Presence of nuclear ion channels adds an extra level of regulation to the aforementioned pathways. Indeed, while some experimental studies suggest that nuclear pore complexes and nuclear ion channels serve as passive elements allowing free diffusion of electrically charged molecules and ions between the cell cytosol and nucleoplasm [55, 56], other point to a possibility of a more specialized role of K$^+$-selective channels in modulation of the nuclear electrostatic potential in certain types of living cells [98]. In the latter case, experimental measurements show that K$^+$-selective channels contribute to the formation of a greater nuclear electrostatic potential ($\sim -10 \text{ mV}$) [98] in comparison to cells in which ions can freely move between the cell cytoplasm and nucleoplasm.
As a result, the nucleus size-dependent stabilization of nucleosomes will be stronger pronounced in cells expressing K⁺-selective NE channels, leading to more significant changes in chromatin.

Overall, the above results indicate that living cells may utilize multiple pathways to fine-tune the chromatin structure in response to environmental cues. However, it should be noted that all these pathways operate on top of a more general physical background created by the molecular processes discussed in our study, which make a major contribution into chromatin organization.

It should be noted that versatility of the developed theoretical approach allows one to incorporate into the model all of the aforementioned mechanisms for a better understanding of their effect on chromatin, warranting future study. Furthermore, with a few minor modifications, the model can be also used to investigate the role of other types of interactions in shaping the chromatin landscape, such as HP1-dependent cross-linking of nucleosomes [122, 123] and formation of contacts between the nuclear lamina network and chromosomal DNA [124, 125]. This can provide valuable information about phase-separation processes taking place inside nuclei of living cells, such as self-organization of hetero- and euchromatin domains, making it possible to better comprehend their role in regulation of the cell response to various environmental cues.

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VI. AUTHOR CONTRIBUTIONS

L.H. and A.K.E. made the computer program used for model calculations. Y.J. supported the study, providing useful suggestions. A.K.E. designed the research, derived formulas, carried out computations, analyzed the data and wrote the paper.
FIG. 1. Semiflexible polymer model of chromosomal DNA packaging in the nucleus of a eukaryotic cell. (a) Main protein complexes involved in histone transportation and nucleosome formation. In the model, nucleosomes assemble on DNA from H2A-H2B and H3-H4 histone dimers, which are transported by histone-binding chaperones schematically indicated as c1 and c2 in the figure. Specifically, by picking up newly synthesised histone dimers in the cell cytoplasm, chaperones switch from an unloaded state (c1u or c2u) to a histone-bound state (c1b or c2b, respectively). Then by moving through nuclear pores, chaperones enter the cell nucleus, where they can deposit histone dimers onto the chromosomal DNA via nucleosomes assembly. This way, by shuttling between the cell nucleus and cytoplasm, histone-binding chaperones help living cells to maintain a constant level of soluble histones in the nucleoplasm. In addition to histone-binding chaperones, nuclear pores also allow free diffusion of ions and small metabolites between the cell nucleus and cytosol, helping to maintain stability of the ionic microenvironment in the nucleoplasm. (b) Chromosome conformations. In the model, conformations of chromosomes were described by triplets of unit vectors \((x^{(u)}(s_u), y^{(u)}(s_u), z^{(u)}(s_u))\), \(u = 1, \ldots, Q\), with each triplet indicating the orientation of the respective \(u^{th}\) DNA molecule with reference to the fixed coordinate system, \((x_0, y_0, z_0)\), at a point corresponding to the arc length \(s_u \in [0, L_u]\) along the DNA. Here \(L_u\) is the length of the \(u^{th}\) DNA molecule. (c) Discretized polymer model of DNA. In the partition function calculations, each DNA polymer was represented by a polygonal chain comprised of small straight segments, each of which was treated as a rigid body with an attached local Cartesian coordinate frame, \((x_j, y_j, z_j)\), representing the DNA segment orientation in space with respect to the fixed global coordinate system, \((x_0, y_0, z_0)\). More details regarding description of DNA and nucleoprotein complexes can be found in Appendices A-B.
FIG. 2. DNA mechanical stretching in a nano-channel. The graph demonstrates force-extension curves of bare DNA in nano-channels of different radii. Data points shown in the figure were calculated by using Metropolis-Monte Carlo algorithm applied to a discretized DNA polymer comprised of \( N = 200 \) segments of \( b = 5 \) nm size; whereas, solid curves were obtained by using transfer-matrix calculations based on Eq. (5). Good agreement between the two methods can be seen from the plot.
FIG. 3. DNA packaging in viral particles. (a) Contributions made by DNA (blue curve) and ions (red curve) into the total free energy of a viral particle (black curve) as functions of the capsid size. Data points shown in the graph are the results of transfer-matrix calculations; whereas, lines indicate smoothing spline interpolation. Solid curves correspond to the case of a semiflexible DNA with the bending and twisting persistence lengths of $A = 50$ nm and $C = 95$ nm, respectively; whereas, dashed curves demonstrate results obtained for a freely-joint polymer chain of the same length but with zero bending and twisting elasticities between the polymer segments. All energy values in the graph are shown up to an additive constant, which is not essential for model calculations. (b) Pressures generated by DNA and ions on the walls of a viral particle as functions of the capsid size. Green rectangle indicates the area corresponding to experimentally measured values for λ-phage particles [76, 77]. (c) Electrostatic and volume-exclusion DNA interaction energies per single DNA base-pair as functions of the capsid size. (d) Distribution of the electrostatic potential in viral particles of different sizes. Each data point shown in the panel represents the corresponding curve displayed in panel (d). Solid line demonstrates interpolation of the data points by a smoothing spline. (f) Schematic picture of DNA packaging in a viral particle. Results shown in panels (g-h) suggest that DNA assumes a coil-like conformation inside the capsid. (g) and (h) The average correlation function between a couple unit vectors tangent to the DNA backbone and the root mean squared distance (RMSD) between two points residing on DNA as functions of the genomic distance between the vectors and points, respectively. Mathematical details regarding the formulas used in all the above calculations can be found in Appendices I-J, L. In all computations, the DNA length was 14 µm, corresponding to the size of EMBL3 λ-phage genome of 41.5 kbp [76], with the size of DNA segments in the discretized polymer model being equal to $b = 3.4$ nm.
FIG. 4.  Cell nucleus size regulation.  (a) Pressures generated on the NE by DNA, ions, macromolecules, and the surface tensions of the NE and ER membrane. The yellow curve representing the osmotic pressure created by macromolecules is plotted for the case of \( n_{macro} = 10^{10} \) molecules; whereas, the magenta curve demonstrates the case of the NE and ER membrane surface tension difference of \( \Delta \sigma = 20 \) mN/m. In transfer-matrix calculations, the total length of DNA was set equal to 2.1 m, corresponding to the size of human genome of \( \sim 6.2 \) Gbp [46]. The length of bare DNA segments in the discretized polymer model was \( b = 3.4 \) nm.  (b) Relative contributions of different cell components to the total positive / negative pressure acting on the NE. Results shown in panels (a) and (b) were obtained for a cell with the total volume of \( V_{cell} = 8000 \) \( \mu \)m\(^3\). Smaller size cells (\( V_{cell} = 4000 \) \( \mu \)m\(^3\)) show very similar behaviour, see Figure S1.  (c) Correlation between the nucleus and cell volumes. Solid curves indicate fitting of experimental data from ref. [8, 9, 11] either to the full model described by Eq. (6) or to the reduced model described by Eq. (7). In the former case, the fitting was done by varying a single model parameter – the difference between the surface tensions of the NE and ER membrane, \( \Delta \sigma \); whereas in the latter case, the linear fitting of \( \Delta \sigma / \zeta \) ratio shown in panel (d) was used as an input in Eq. (8). Inset displays a large scale view of the fitting of experimental data from ref. [8].  (d) \( \Delta \sigma / \zeta \) ratio as a function of the nucleus size. Data points show values calculated by using Eq. (7) based on experimental data displayed in panel (c). Solid lines indicate data points fitting to a linear function. Inset demonstrates a large scale view of the fitting of data points obtained based on experimental data from ref. [8].  (e) Schematic picture of cellular components exerting pressure on the NE. The size of the cell nucleus in the general case is determined by mechanical equilibrium described by Eq. (6), which involves a tight interplay between the pressures generated by DNA (\( p_{DNA} \)), ions (\( p_{ions} \)) and macromolecules (\( p_{macro} \)). In addition, it has been previously hypothesised that NE and ER, which share the same bilayer lipid membrane, may be involved in a tug-of-war with each other via indirect membrane-binding competition between lamins and reticulons [1] that can be described in terms of \( p_{NE} \) and \( p_{ER} \) pressures associated with the surface tensions of the NE and ER membrane.
FIG. 5. Nuclear electrostatic potential, and correlation between the nucleus size and nucleosome stability. (a) Distribution of the electrostatic potential in nuclei of different sizes. The plot shows results obtained for a cell with the total volume of $V_{\text{cell}} = 8000 \, \mu\text{m}^3$. Smaller size cells ($V_{\text{cell}} = 4000 \, \mu\text{m}^3$) demonstrate a very similar nuclear electrostatic potential distribution, see Figure S2. (b) Volume-averaged value of the electrostatic potential in nuclei of different sizes. Data points indicate values calculated based on the curves shown in panel (a) and Figure S2; whereas, solid lines demonstrate smoothing spline interpolation. (c) Occupancy fraction of chromosomal DNA by nucleosomes as a function of the cell nucleus size. Data points represent results of transfer-matrix calculations; whereas, solid lines show smoothing spline interpolation. (d) and (e) Force-extension and force-DNA occupancy fraction curves obtained by mechanical stretching of a small part of chromosomal DNA (2 $\mu$m in length) in nuclei of different sizes. Solid curves on panels (d) and (e) demonstrate results of transfer-matrix calculations performed on a cell with the total volume of $V_{\text{cell}} = 8000 \, \mu\text{m}^3$ (see also Figure S3 for $V_{\text{cell}} = 4000 \, \mu\text{m}^3$ case). Dashed curves in panel (d) display results of the force-extension curves’ fitting to the previously published model of a mechanically stretched DNA [51]. (f) Binding free energy of histone octamers to DNA as a function of the cell nucleus size. Data points shown in the graph are calculated based on the force-extension curves’ fitting displayed in panel (d) and Figure S3(a). Solid curves indicate predictions of the binding free energy of histone octamers by Eq. (9).
FIG. 6. Role of the electrical charge of chaperones in regulation of the histones’ binding free energy to DNA. The plot demonstrates results of calculations based on Eq. (9) performed for different electrical charges of histone-binding chaperones and cell nucleus sizes. For the sake of simplicity, in calculations it was assumed that H2A-H2B-binding chaperones as well as H3-H4-binding chaperones, both have similar electrical charges: $q_{c1u} = q_{c2u}$. Dashed curve shown in the graph indicates model predictions based on estimation of the chaperone electrical charge ($q_{c1u} = q_{c2u} = -28.2 q_e$, where $q_e = 1.6 \times 10^{-19}$ C is the elementary charge), which was obtained from experimental measurements of relative histone concentrations in the cytosol and nucleoplasm of HeLa cells [126], see Appendix A for details. This curve corresponds to the results shown in Figures 4, 5. It can be seen from the plot that the binding free energy of histone octamers to DNA may either change with the size of the cell nucleus, or remain insensitive to nucleus size variations, depending on the absolute value of the electrical charge of histone-binding chaperones.


