Constructing the constraints: The main DNA-histone interactions are hydrogen bonds that involve 28 distinct DNA phosphates [4]. However, the rigid base-pair model does not contain the phosphates explicitly. Here we determine their positions with respect to the middle-frame of the corresponding basepair (bp) steps. The position vector of each phosphate along the DNA with respect to the corresponding middle-frame (see S1 Fig) can be written as

\[ \mathbf{r}_P = a_i \mathbf{d}_1^m + b_i \mathbf{d}_2^m + c_i \mathbf{d}_3^m \quad i = 3, 5. \]  

In this equation \( \mathbf{d}_1^m, \mathbf{d}_2^m \) and \( \mathbf{d}_3^m \) are the basis vectors of the middle-frame, and \( a, b \) and \( c \) are the components of the relative position vector along these basis vectors, with the subscripts 3 and 5 indicating the phosphates on the 3' and 5' strands respectively. S2 Fig shows the distribution functions of \( a, b \) and \( c \) for all the phosphates in two nucleosome crystal structures: NCP147 [16] and NCP601L [38].

It can be seen that the two crystal structures give rise to similar distributions, and each distribution has a sharp peak. Calculating the average of each distribution for both crystal structures, we obtain

\[ \langle a_3 \rangle = \langle a_5 \rangle = -0.30 \text{ nm} \]
\[ \langle b_3 \rangle = -\langle b_5 \rangle = 0.89 \text{ nm} \]
\[ \langle c_3 \rangle = -\langle c_5 \rangle = -0.05 \text{ nm} \]  

The standard deviation of all the distribution functions is about \( \sigma \approx 0.05 \text{ nm} \), comparable to the error in the crystallography experiments [19]. Therefore, within the accuracy of the experimental data, it can be assumed safely that all of the phosphates are firmly attached to the middle-frames and their relative positions are given by the Eqs. (2) and (3).

To identify the strongly bound phosphates in the nucleosomal DNA, we looked at the crystallographic B-factor for all the phosphates in the NCP147 structure [16]. Each local minimum in the B-factor corresponds to a strongly bound phosphate. This gives rise to 14 distinct nucleosome binding sites, each containing two bound phosphates [41]. Starting from the NCP147 crystal structure, we calculated the positions of the bound phosphates from Eqs. (2) and (3) and we assumed that they are all fixed in their places. Since the preferred orientations of the bound middle-frames are in principle affected by the DNA sequence, we do not use the crystal structure to constrain them. Ideally one would derive the elasticities of those constraints from a collection of many nucleosome crystal structures but the currently available data make such an approach unattainable [42]. Instead we allow for a pre-relaxation of the system before fixing the bound middle-frames permanently. The pre-relaxation was performed using the standard Monte Carlo simulation with a homogeneous DNA sequence, to remove any sequence-dependent bias. The configurational Monte Carlo moves during the pre-relaxation course are designed such that they keep the positions of the bound phosphates fixed, but allow for the free rotation of the bound middle-frames. After the pre-relaxation we put each of the bound middle-frames in its preferred orientation, and we kept it fixed during all subsequent Monte Carlo simulations.

Nucleosome binding free energies: It is known that nucleosome assembly in vitro is initiated by binding the H3-H4 tetramer to the DNA molecule. It is only after the reduction of the salt concentration that the H2A-H2B dimers bind to the tetramer to form a full nucleosome. Therefore it is expected that the apparent in vitro nucleosome free energies are mainly determined by the interaction of the DNA with the histone tetramer [1, 43]. To
account for this, we model the tetramer by allowing the DNA to only attach to the ±2.5, ±1.5 and ±0.5 binding sites. S3 Fig shows the binding free energy difference between pairs of DNA molecules (of varying lengths, sometimes even substantially shorter than 147 bp). We calculate the binding free energy accounting for all possible positions of the tetramer on a given DNA molecule. The following 22 pairs were studied: c1/c2, c1/c3, d1/d2, d1/d3, d1/d4, d1/d5, c1/e2, e1/e3 [1], TG/TG-T, TG/TR-5, TG/TRGC [39], TG/ANISO, TG/TTT, TG/NOTA, TG/EXAT, TG/EXGC, TG/IAT, TG/IGC, TG/END, TG/ANNA, TG/34 and TG/20 [40]. The root-mean-square deviation between our model prediction and the data is 1.2 $k_B T$. This can be compared to the prediction of the computational nucleosome model by de Pablo and coworkers [22] where a subset of our pairs was studied. The root-mean-square deviation of those 9 data points is worse, 2.1 $k_B T$. This does not necessarily mean that their model is worse, as they considered the free energy of the full nucleosome (at a fixed position). In fact, linear least squares gives for our model $\Delta \Delta G_{\text{model}} = 1.50 \Delta \Delta G_{\text{exp}} - 0.88 k_B T$ whereas the full octamer prediction in Ref. [22] shows a much steeper dependence: $\Delta \Delta G_{\text{model}} = 2.20 \Delta \Delta G_{\text{exp}} - 2.04 k_B T$.

**Nucleosome positioning in vivo:** We calculated the energy landscape for a single nucleosome on yeast chromosome I and compared the distances between in vivo mapped nucleosomes [25] to local minima in our landscape. A histogram of the distances from the positions of all the 1293 mapped nucleosomes to the nearest local minimum as predicted by our model (more precisely, the smallest energy within a window of 11 bp length centered around each mapped nucleosome) is given in S11 Fig A (red rectangles). About 60 percent of the nucleosomes lie within the range of one bp around a local minimum. As a comparison we show also the prediction from a probabilistic model trained on in vitro data (blue rectangles) [27] where about 43 percent of these nucleosomes are within one bp around a local minimum. We found also about 60 percent agreement when restricting this analysis to the 769 mapped nucleosome on top of genes, S11 Fig B. If the nucleosomes were randomly positioned (e.g. if the action of chromatin remodellers would overrule sequence preferences) one would find that only $3 \times 100/11\% \approx 27\%$ of the nucleosomes to be within one bp around a local minimum. As we find a much higher fraction, this demonstrates that – even in vivo – a large fraction of nucleosomes is rotationally positioned and that our model is capable of capturing this effect to a substantial extent.

However, a precise comparison with the in vivo data not only requires an accurate nucleosome positioning energy landscape, but it also needs to take into account exclusion between nucleosomes that compete for the same DNA substrate. This can be taken care of using statistical physics approaches [44-46], allowing us to calculate the nucleosome density profile along the DNA. Assuming that nucleosomes interact with each other via a hard-core potential, the nucleosome density on a DNA molecule, $\rho$, as a function of the nucleosome starting point $x$, satisfies the Percus equation [47]:

$$
\mu = E(x) + \ln \rho(x) - \ln \left[ 1 - \int_{x}^{x+\sigma} \rho(x') \, dx' \right] + \int_{x-\sigma}^{x} \frac{\rho(x')}{1 - \int_{x'}^{x+\sigma} \rho(x''') \, dx'''} \, dx'
$$

where $\sigma = 147\, \text{bp}$ is the nucleosome footprint, $E(x)$ is the elastic energy landscape and $\mu$ is the chemical potential, i.e. the free energy gain by the reservoir when a nucleosome unbinds from the DNA. Eq. (4) can be solved numerically [48] for the nucleosome density. The effective energy can then be calculated from

$$
E_{\text{eff}}(x) = -k_B T \ln \rho.
$$
For small chemical potentials, nucleosome positioning is mainly governed by DNA elasticity and thus the elastic and effective energy landscapes are similar. As the chemical potential increases, alternating regions with high and low nucleosome density appear along the DNA molecule. We calculated the effective energy for the YAL002W gene, which resides on chromosome I between positions 143709 and 147533, see Fig 2B and S10 Fig. To diminish end effects, we included the two 1000 bp sequences which flank the gene at its two sides on yeast chromosome I. S10 Fig shows in blue the elastic energy landscape and in red the effective energy landscape for \( \mu = 80 k_B T \) which is about 10 \( k_B T \) higher than the average elastic energy of random sequences, a value consistent with the findings in Ref. [46]. 25 distinct high-density regions can be easily identified along the gene, where the effective energy landscape is low, a number equal to the number of experimentally mapped nucleosomes [25], shown in S10 Fig as vertical lines. The experimental nucleosome positions fall typically into the local minima of the effective energy landscape, especially for the nucleosomes at the central part of the gene, where there is an impressive agreement between the model and the experimental data. At the terminal regions, on the other hand, we did not find a strong correlation between the experimental data and the model predictions. This discrepancy could reflect the presence of other DNA binding proteins that may affect nucleosome positioning in vivo [46].

S2 Interpretation of the positioning rules for the model nucleosome

We used the Monte Carlo simulation in the configuration space to find the averaged DNA structure in our model nucleosome. S8 Fig shows the averaged degrees of freedom for the NCP147 DNA sequence as obtained in the model (solid curves, blue), in comparison with the crystal structure [16] (dashed curves, red). As can be seen, the rotational degrees of freedom are clearly correlated with the crystal structure. In particular, the model captures the periodic variations in roll. However, the large variations in shift and rise are not reproduced by the model, and the peaks in the slide are underestimated. Despite these shortcoming our model is successful in predicting the positioning rules that are discussed here in more detail.

The emergence of the nucleosome positioning rules in our model is connected with DNA bending. In the rigid base-pair model the DNA bending is expressed in terms of two perpendicular bending modes known as tilt and roll. Tilt corresponds to the bending of a dinucleotide step over its backbone, while positive and negative rolls are defined as bending toward the major and minor grooves respectively. As mentioned above, bending the DNA into a super-helical configuration results in periodic oscillations of roll and tilt in our model nucleosome, see S8 Fig. These oscillations have a period of 10 bp and a phase difference of 2.5 bp approximately. Roll has a minimum at the minor groove bending sites, while tilt changes sign from positive to negative. The opposite occurs at the major groove bending sites, where roll reaches its maximum value and tilt goes from negative values to positive values.

We found that the nucleosome positioning rules in our model either make DNA locally softer with respect to bending, or help the DNA to intrinsically bend into the correct "direction" at the minor groove and major groove bending sites. As a typical example, S9 Fig shows the occurrence frequencies of two tetranucleotides along the nucleosomal DNA, namely TTAA and AGCT.

As can be seen, TTAA prefers the minor groove bending sites while AGCT prefers the major groove bending sites. This can be understood by looking at the tilt and roll elastic parameters in the model (S4 Fig). Although the TA step has a relatively large positive
intrinsic roll, it is the softest step with respect to bending. On the other hand AA and TT steps are rather rigid, but they have the lowest intrinsic roll after GC, and also have significantly high negative and positive intrinsic tilts respectively. Therefore the motif TTAA is suitable for the minor groove bending sites. Considering the motif AGCT, one can see in S4 Fig that the GC step is one of the most resistant steps towards bending. In addition AG/CT steps are rather stiff. Nevertheless AGCT occurs at the major groove binding sites because AG/CT steps have large positive intrinsic roll and the highest negative and positive intrinsic tilts respectively, so the intrinsic bending of AGCT is compatible with the DNA bending at these locations.

Similar situations occur for CG, CC, and GG steps, as these steps can come together as neighbours of the GC step, and they all provide a favorable intrinsic bending for DNA at the major groove bending sites (S4 Fig). Furthermore, CG is the softest step with respect to bending after TA. As another minor groove example, the AT step has low intrinsic roll and zero intrinsic tilt, and is much stiffer compared to TA. At high enough temperatures, it appears in the Mutation Monte Carlo simulation because it can act as a bridge between an AA or a TA step at the left, and a TT or a TA step at the right.

The above examples illustrate how the nucleosome positioning rules can be understood.

Supplemental references

43. Dong F, van Holde KE. Nucleosome positioning is determined by the (H3-H4)2 tetramer. Proc Natl Acad Sci USA. 1991;88: 10596-10600.
Fig. S1: **Base-pair step together with its corresponding midframe.** The red spheres represent the phosphates whose positions with respect to the middle frame are given by Eqs. (2) and (3) in S1 Text.
Fig. S2: **DNA phosphate positions inside nucleosome crystal structures.** The distribution functions of a, b and c as defined in Eq. (2) of S1 Text, for all the phosphates in the NCP147 [16] (red) and NCP601L [38] (green) crystal structures.
Fig. S3: Predicted and experimental binding free energy. Each point corresponds to a pair of DNA molecules, 22 pairs in total: c1/c2, c1/c3, d1/d2, d1/d3, d1/d4, d1/d5, e1/e2, e1/e3 [1], TG/TG-T, TG/TR-5, TG/TRGC [39], TG/ANISO, TG/TTT, TG/NOTA, TG/EXAT, TG/EXGC, TG/IAT, TG/IGC, TG/END, TG/ANNA, TG/34 and TG/20 [40]. The dashed line corresponds to perfect agreement. The root-mean-square deviation between our model prediction (the tetramer free energy; see S1 Text for detail) and the experimental data is $1.2k_BT$. 
Fig. S4: (A) The tilt stiffness versus the intrinsic tilt and (B) the roll stiffness versus the intrinsic roll for the ten distinct dinucleotide steps in our model. For the remaining six steps, the bending parameters are simply obtained by the inversion transformation, which changes the sign of the intrinsic tilt and keeps other parameters unchanged.
Fig. S5: **Dependence of dinucleotide distributions on effective temperature.** Probability distribution of the AA step, obtained by the MMC for three different temperatures: $T = 600$ K (blue), $T = 100$ K (green) and $T = 21$ K (red).
Fig. S6: A color-map of the frequencies for all 16 dinucleotide steps as a function of the position. The distributions are obtained in a Mutation Monte Carlo simulation at temperature 100 K.
Fig. S7: **Nucleosome positioning rules.** (A) Fraction of dinucleotides AA/AT/TA/TT and separately CC/CG/GC/GG at each position along the nucleosome model found in 10 million high affinity sequences produced by MMC at 100 K. The model recovers the basic nucleosome positioning code. (B) Same as (A) but on top of 1200 randomly generated coding sequences (produced by sMMC). The same periodic signals are found albeit with a smaller amplitude.
Fig. S8: **Comparison between model and crystal structure.** The averaged degrees of freedom for NCP147 DNA sequence as obtained in the model (solid curves, blue), in comparison with the crystal structure (dashed curves, red) [16].
Fig. S9: The occurrence frequencies of TTAA (triangles, red) and AGCT (dots, green) as obtained in an unconstrained Mutation Monte Carlo simulation at 100 K. The solid and dashed vertical lines indicate minor and major groove bending sites respectively.
Fig. S10: **Translational positioning of nucleosomes.** The effective energy landscape with $\mu = 80 \ kT$ (red curves), the elastic energy landscape (blue curves) and the experimentally mapped nucleosomes [25] (vertical black lines) along the YAL002W yeast gene. The elastic energy is shifted down by $30 \ kT$ for clarity. The top panel shows the landscapes over the entire gene. Each of the remaining panels zooms into a 765 bp long portion of the gene. All of the experimentally mapped nucleosome positions fall into local minima. In addition, the corresponding minima are quite deep in the central region of the gene.
Fig. S11: **Rotational positioning of nucleosomes.** (A) The histogram of the distances between 1293 experimentally mapped nucleosomes [25] on yeast chromosome I and the nearest local minima in the theoretical energy landscape (red rectangles). As a comparison we show also the prediction from a probabilistic model trained on in vitro data (blue rectangles) [27]. (B) The distance histogram as defined in (A) for 769 nucleosomes on yeast chromosome I which are located on the genes. The two histograms are quite similar. In both cases, 60 percent of the experimental nucleosome positions lie within the range of one bp around a local minimum in the theoretical energy landscape.
Fig. S12: **Evidence for multiplexing in two eukaryotic genomes.** Normalized Fourier amplitudes of the distribution of the synonymous codons for all 20 amino acids along nucleosomes on top of genes (purple curve) and of the distribution of the corresponding trinucleotides along nucleosomes outside genes (blue curve) for *S. cerevisiae* (left) and *S. pombe* (right). The peaks at spatial periodicity 13 corresponds to a 10 bp periodic signal. In most cases the height of this peak is larger for the non-coding case, evidence for multiplexing of genetic and mechanical information.