The large dimer interface is predominantly hydrophobic in character and buries \( \sim 22\% \) of the overall surface of \( \gamma \)Nap1(5); it is therefore unlikely that \( \gamma \)Nap1 exists in a monomeric state except for perhaps under the most dilute conditions (6). All Nap1 family members have a C-terminal acidic domain (CTAD) of varying length that is not required for histone binding and chromatin assembly, and a variable N-terminal tail of unknown function (3).

Nap1 binds all four core histones as well as the linker histone H1 (8–10). Nap1-mediated nucleosome formation \textit{in vitro} is characterized by the transfer of a \((H3/H4)_2\) tetramer onto DNA, followed by the incorporation of \(H2A/H2B\) dimer (8). Nakagawa \textit{et al.} (11) have qualitatively shown that the affinity of the \((H3/H4)_2\) tetramer for DNA exceeds its affinity for Nap1, and that the affinity of \(H2A/H2B\) for a DNA-bound \((H3/H4)_2\) tetramer (a tetrasure) is greater than its affinity for Nap1.

Our understanding of the thermodynamics of chaperone-histone interactions in general, and of Nap1-histone interactions in particular is very limited. Qualitative data have been obtained from gel-shift and pull-down experiments. Depending on input stoichiometries, \(\gamma\)Nap1-histone complexes appear on gels shifts in at least two distinct complexes, which complicates the interpretation of these data (9). Pull-down experiments are notoriously difficult to quantitate. This has resulted in difficulty interpreting results especially regarding the contributions of histone tails and \(\gamma\)Nap1 tails on \(\gamma\)Nap1-histone interaction (9).

We and others (10, 12) have previously found that \(\gamma\)Nap1 promotes nucleosome disassembly under certain conditions, and that it is capable of exchanging histones and their variants into nucleosomes. The latter function is of particular interest, because it is currently not well understood how histone variants and replacement histones (non-allelic isoforms of major type histones with distinct amino acid sequence, expression patterns, and chromosomal locations) are incorporated into chromatin in a replication-independent manner. Furthermore, these two noncanonical chaperone functions require the CTAD (9, 10). The simplest model that explains the diverse roles of \(\gamma\)Nap1 is that the chaperone prevents the incorrect binding of histones to DNA or other histones, thereby promoting formation of canonical chromatin. This model of \(\gamma\)Nap1 function would also allow for histone exchange through establishing an equilibrium between \(\gamma\)Nap1-bound and nucleosomal histones. This model predicts a thermodynamic link between the nucleosome and \(\gamma\)Nap1. Thus, a quantitative and systematic exploration of the interactions of \(\gamma\)Nap1 with histones not only forms the basis for our understanding of the many functions of \(\gamma\)Nap1, but will also provide information on the extreme upper limits of nucleosome thermodynamic constants.
**EXPERIMENTAL PROCEDURES**

**Reagents**—Wild-type, mutant, and tail-less *Xenopus laevis* histones (amino acids 14–118 for H2A, 24–122 for H2B, 27–135 for H3, and 20–102 for H4) histones were prepared as described (12, 13). The H2B(T112C) and H4(E63C) were prepared according to Ref. 14. Cysteine-to-alanine mutagenesis of yNap1 was done using QuikChange mutagenesis. The resulting triple mutant, C200A/C249A/C272A, further referred to as (yNap1<sub>tr</sub>), leaves only the Cys-414 position available for labeling. Complex formation between H2A/H2B dimer and yNap1 was tested using native gel electrophoresis. All mutant yNap1 constructs interact with H2A/H2B dimer in the same manner as wild-type yNap1, as checked by gel-shift. Both Alexa 488 and 546 were purchased from Invitrogen, and labeling was done as described (14). Biotin polyethylene oxide iodoacetamide was purchased from Sigma-Aldrich and attached to the proteins via the same protocol as the Alexa dyes.

**Fluorescence Titrations**—Fluorescence was measured using either an AVIV model ATF105 or a Horiba Jobin Yvon Fluorolog-3 spectrophotometer. Labeled protein was added to both the sample and the reference cuvette, with non-labeled protein added to the sample cuvette and buffer added to the reference. Varying incubation times (2–15 min) confirmed that the fluorescence signal had reached equilibrium. The normalized fluorescence change was determined by Equation 1,

\[
\text{Norm.f.c}_{\text{obs}} = \frac{R_\text{obs} - R_t}{R_t - R_i} \quad \text{(Eq. 1)}
\]

where \(R_{\text{obs}}\) is the observed ratio of the fluorescence signal (sample cuvette signal/reference cuvette signal), \(R_t\) is ratio of the fluorescence initial, and \(R_i\) is the ratio of the fluorescence final or where saturation is reached. While the magnitude of the signal change was constant for each experiment, it varied from 10 to 30% between different experiments (i.e. labeled yNap1 binding H2A/H2B versus labeled H2A/H2B binding to yNap1) and with the label used (546 or 488 Alexa). We also monitored the normalized fluorescence ratio of protein titrated into its corresponding binding partner in either buffer or 5 mM guanidium-HCl.

The presence of guanidium-HCl did not alter the initial signal (pre-addition of the binding partner) nor did the signal change with the addition of \(\mu\)M concentrations of the binding partner in guanidium HCl.

**Biotin Pull-down Experiments**—Biotin-tagged yNap1 was used to pull Alexa-labeled histones out of solution to confirm that we were monitoring binding under low protein concentrations. Trace amounts (5–30 × 10<sup>-9</sup> M) of Alexa-labeled protein were mixed with either biotin-tagged protein at concentration 5–10<sup>-fold</sup> above the observed \(K_d\) or buffer and incubated for 5–20 min. The reaction mix was added to BioMag streptavidin (Qiagen) in a 100-fold excess and incubated for 30 min, while shaking in a 96-well plate. The plate was then placed on a 96-well magnet (Qiagen) and incubated for 15 min. Once the beads were pulled-down, the buffer was removed and checked for total fluorescence. Beads were then washed extensively, and the wash was also checked for fluorescence.

**Data Analysis**—Affinity measurements were done using concentrations of labeled protein (P) that were at least 5–10<sup>-fold</sup> less than the \(K_d^{\text{pp}}\). \(K_d^{\text{pp}}\) was determined by fitting Equation 2 derived from Reaction Scheme 1 to the normalized f.c.,

\[
P + nL \rightleftharpoons P(L)_n
\]

REACTION SCHEME 1

\[
\text{Norm.f.c.} = \frac{L^{nH}_\text{f.c.}}{L^{nH}_\text{f.c.} + K_d^{\text{app}}}
\]

where \(L\) is the total concentration of protein titrated, \(n_L\) is the Hill coefficient, and \(K_d^{\text{app}}\) is the apparent dissociation constant. The \(n_L\) was assumed to be one unless the data dictated otherwise. When the \(n_L\) was determined not to be equal to one, the linearized form of Equation 2 was used as Equation 3,

\[
\log\left[\frac{f}{1 - f}\right] = n_L \log[L] + b
\]

where \(f\) is equal to the normalized f.c. divided by the normalized f.c.<sub>max</sub>. A dimerization model (Reaction Scheme 2) was also used to determine if this could explain the observed cooperativity.

REACTION SCHEME 2

\[
P + 2L \rightleftharpoons P(L)_2
\]

In this model, the normalized f.c. is still described by Equation 2 except \(n_L = 1\) and \(L\) is replaced by \(\langle L\rangle_2\), where \(\langle L\rangle_2\) is defined in Equation 4.

\[
\langle L\rangle_2 = \frac{L_1 - L}{2}
\]

In Equation 4, \(L\) is total concentration of protein titrated, and \(L\) is calculated in Equation 5,

\[
L = \frac{-K_1 + \sqrt{(K_1)^2 + 8L_1K_1}}{4}
\]

where \(K_1\) is the dimerization constant, and \(L_1\) is total concentration of protein titrated. Stoichiometries were determined by fluorescence titrations with labeled protein concentration increased to >10<sup>-fold</sup> higher the \(K_d^{\text{pp}}\). The fluorescence ratio was plotted as a function of the ratio of protein titrated to labeled protein. Under these conditions, the protein ratio at which the normalized f.c. leveled off is equal to the stoichiometry. All data analysis was performed using either Kaleidagraph or Graphpad Prism.

**RESULTS**

**A Quantitative Assay to Study Nap1-Histone Interactions**—To determine the binding affinity of yNap1 to histones we developed a series of fluorescence binding assays. yNap1 contains four cysteines in the structured region of yNap1. These were changed to alanine (yNap1<sub>tr</sub>), while the cysteine that is located in the unstructured CTAD was labeled with either Alexa-546 or -488. With the exception of Cys-272 (at the tip of \(\beta_4\)), these residues are not conserved among the various members of the family. The addition of unlabeled H2A/H2B dimer
to a preparation of yNap1m labeled with Alexa-488 (yNap1m*) resulted in a decrease in fluorescence (Fig. 1A). To exclude the possibility that either mutation or derivatization of Cys-414 have an impact on the affinity of yNap1 for histones, we measured the interaction of an H2A/H2B(T112C) dimer in which H2B had been labeled with Alexa-546. H2BT112 is located in the well-defined αC-helix and is solvent-exposed; the attachment of a fluorescent label to this position is not expected to disrupt the structure of the histone dimer (15). Addition of the unlabeled protein (in this case wild-type yNap1) resulted in a change in fluorescence (Fig. 1).

The quantitative analysis of yNap1-histone interactions is complicated by the oligomeric nature of all interaction partners. The minimal unit of yNap1 is very likely a homodimer, but we cannot completely exclude a monomer-dimer equilibrium at the low concentrations used in some of the assays (7). H2A/H2B very likely exists exclusively as a heterodimer under physiological conditions (16), whereas the (H3/H4)4 tetramer may dissociate into two half-tetramers (consisting of a histone fold dimer of one copy each of H3 and H4) under certain conditions, although this has not been investigated systematically (17, 18). When either histone complex is refolded, it elutes as an H2A/H2B dimer and (H3/H4)2 tetramer, respectively (13). For ease of comparison, all numbers given here assume a single chain of yNap1, and one histone fold dimer for either H2A/H2B or H3/H4, unless otherwise stated.

The labeled protein was kept at ~0.5–0.05 nM (or 10-fold below the Kₐ) while the binding partner was titrated to ~10⁻⁹ to 10⁻⁷ M. The ratio of these titrations to a titration of equivalent buffer volumes is plotted in Fig. 1B. The fluorescence change is reversed with the addition of high concentrations of salt (>850 mM final concentration, data not shown). Fig. 1B shows the normalized fluorescence change as a function of either H2A/H2B dimer titrated into Alexa-546-labeled yNap1 (yNap1m*) or wild-type yNap1 titrated into Alexa-546 labeled H2A/H2B (T112C) dimer. The two curves (assuming yNap1 as a monomer) superimpose almost exactly, clearly demonstrating the validity of our assay. At concentrations above 10⁻⁷ M, a second phase was observed that did not plateau at the maximal concentrations testable (~10⁻⁴ M, not shown). This second phase is consistent with gel-shift experiments, where multiple higher shifts were observed at similar concentrations (9). We assume that the higher order gel-shifts and the second phase of fluorescence seen at high concentrations are due to nonspecific interactions between histones and the chaperone.

To confirm that the first change in fluorescence was indeed due to the interaction between histones and yNap1, we used biotin-tagged histones at concentrations above the first plateau (10-fold above the Kₐ; ~100 nM) and 200-fold lower than the second transition to pull-down (yNap1m*)-Alexa-546 (10 nM). Under these conditions, we were able to reduce the amount of unbound (yNap1m*)-Alexa-546 to undetectable levels (data not shown), confirming that the first phase of fluorescence change indeed represents histone binding by yNap1.

One yNap1 Dimer Binds Two H2A/H2B Dimers with High Affinity—The change in fluorescence of H2A/H2B (T112C)-Alexa-488 (<5-fold the Kₐ) was measured as a function of yNap1 (Fig. 1C and Table 1). The observed change in fluorescence followed a cooperative dependence on the concentration of yNap1 added. The data were plotted as a function of yNap1 concentration assuming either a yNap1 monomer or yNap1 dimer (Table 1). The Kₐ and Hill coefficient (n_H) was calculated from the fit of a binding isotherm (Equation 2) to fluorescence change (f.c.) (Fig. 1B). The Kₐ of H2A/H2B dimer for a yNap1 monomer was determined to be 1.3 ± 0.1 nM with a n_H of 2.3 ± 0.3, and as expected, the Kₐ was ~2-fold lower (Kₐ = 0.7 ± 0.1 nM, n_H = 2.6 ± 0.3) when yNap1 was fit as a dimer (Table 1).

The fit to quenching data obtained from the converse experiment (i.e. labeled yNap1 and unlabeled H2A/H2B dimer, Fig. 1 and Table 1) resulted in a Kₐ of 1.2 ± 0.1 × 10⁻⁷ M and a n_H of 1.9 ± 0.4 (Fig. 1B). Taken together these data indicate yNap1 has a high affinity for the H2A/H2B dimer and that neither the point mutations in H2B or yNap1, nor the addition of fluorescent labels to defined regions in either binding partner has a
The Cooperative Nature of the H2A/H2B-yNap1 Interaction Is Dependent on the Ionic Strength and on the Histone Tails—Given the highly charged character of yNap1 and histone H2A/H2B dimer, we hypothesized that the affinity of histones for yNap1 would decrease with increasing ionic strength. We also wanted to investigate the effect of ionic strength on the cooperative nature of histone binding. We measured the affinity of yNap1 to H2A/H2B at 0.15, 0.35, and 0.55 M total ionic strength (Tables 1 and 2). While the \( K_d \) increased from 1.3 ± 0.1 to 7.8 ± 1.5 \( \times 10^{-9} \) M upon increasing the ionic strength from 0.15 to 0.35 and 0.55 M, surprisingly the cooperativity also changed from a Hill coefficient \( (n_H) \) of ~2 to 1 (Fig. 3A and Tables 1 and 2). It has been proposed that the histone tails contribute to the interaction with yNap1 (9). To quantitate this contribution we measured the affinity of tail-less H2A/H2B to yNap1m* -Alexa-546 (\(<5\)-fold \( K_d^{pp} \)). Because tail-less histones are notoriously sticky at low ionic strength, especially when fluorescently labeled, all subsequent measurements were performed at 0.35 M NaCl (Table 2). The data for the interaction of tail-less H2A/H2B with yNap1 were best described by a cooperative model with a \( K_d^{pp} \) of 2.1 ± 0.2 \( \times 10^{-9} \) M and a \( n_H \) of 1.7 ± 0.2. Thus, these histone constructs exhibit cooperativity under conditions where wild type histones bind in a non-cooperative manner (Fig. 3B, inset). We conclude that the tails of H2A and H2B contribute negatively to the interaction with the chaperone, and also are responsible for preventing cooperative binding under these conditions.

**Two Copies of H3/H4 Bind One yNap1 Dimer with High Affinity**—To determine the affinity of yNap1 to (H3/H4), we titrated \((yNap1^{m*})\)-Alexa-546 (\(<5\)-fold \( K_d^{pp} \)) with refolded (H3/H4) tetramer. The \( K_d^{pp} \) was 20.0 ± 0.7 nM with a \( n_H \) of 1.5 ± 0.1 when the concentration of H3/H4 was calculated as a H3/H4 heterodimer, and half that when the concentration was calculated assuming a (H3/H4)2 tetramer (Table 2). This measurement is in excellent agreement with the value obtained from Alexa-546-H3/H4(E63C) (Table 2), further validating our assay. As with H2A/H2B, the removal of the histone tails of H3 and H4 resulted in an increased affinity for yNap1 \( K_d^{pp} \) of 4.0 ± 0.2 \( \times 10^{-9} \) M and Table 2).

The stoichiometry of the H3/H4-yNap1 complex was measured as described above (Fig. 4A, inset). This experiment clearly shows that one H3/H4 histone-fold dimer binds one yNap1 monomer. If the predominant form of yNap1 under these conditions.

**TABLE 2**

| Protein titrated | \( K_d^{pp} \) \( \times 10^{-9} \) M | Hill coefficient | Labeled protein |
|-----------------|-----------------|-----------------|----------------|
| H2A/H2B         | 7.8 ± 0.4        | n.a.*           | yNap1m*        | 7.5 |
| Tailless H2A/H2B| 2.1 ± 0.2        | 1.7 ± 0.2       | yNap1m*        | 7.5 |
| H2A.B8/H2B      | 4.2 ± 0.5        | n.a.            | yNap1m*        | 7.5 |
| H2A.Z/H2B       | 7.4 ± 0.4        | n.a.            | yNap1m*        | 7.5 |
| H3/H4-tetramer  | 10.0 ± 0.6       | 1.4 ± 0.1       | yNap1m*        | 7.5 |
| Dimer           | 20.2 ± 0.7       | 1.5 ± 0.1       | yNap1m*        | 7.5 |
| Tailless H3/H4-tetramer | 4.0 ± 0.2 | 1.5 ± 0.1 | yNap1m*        | 7.5 |
| Dimer           | 8.8 ± 0.7        | 1.3 ± 0.1       | yNap1m*        | 7.5 |
| H3(H113A)/H4-tetramer | 7.9 ± 0.4 | 1.3 ± 0.1 | yNap1m*        | 7.5 |
| Dimer           | 16 ± 0.7         | 1.3 ± 0.1       | yNap1m*        | 7.5 |
| H1              | 3.0 ± 0.1        | 1.4 ± 0.1       | yNap1m*        | 7.5 |
| H2A             | 3.5 ± 0.5        | n.a.            | yNap1m*        | 7.5 |
| H3              | >10              | n.a.            | yNap1m*        | 7.5 |
| Archaic Histones| d.n.b.           | n.a.            | yNap1m*        | 7.5 |
| DNA             | 1.6 ± 0.2        | n.a.            | H3/H4(E63C)    | 7.5 |
| H2A/H2B         | 7.8 ± 1.5        | n.a.            | yNap1m*        | 7.5 |
| H2A.Z/H2B       | 21.6 ± 1.8       | n.a.            | yNap1m*        | 7.5 |
| yNap1 - Monomer | 20.1 ± 1.4       | n.a.            | H3/H4(E63C)    | 7.5 |
| Dimer           | 10.1 ± 0.7       | n.a.            | yNap1m*        | 7.5 |

* n.a., not applicable.
Thermodynamics of Nap1-Histone Interactions

FIGURE 3. Cooperativity is dependent on both the ionic strength and histone tails. A, Hill plot of H2A/H2B binding yNap1 at −0.15 M (closed circles) and −0.35 M (open squares) of same data in plots) ionic strength. The slopes of these data result in a nH of 1 ± 0.1 and 2.4 ± 0.2, respectively. B, normalized fluorescence change as a function of histone dimer. Closed circles are tail-less dimers, and open squares are major type histone dimer. Inset of B is the Hill plot of the tail-less (closed circles) and major type (open squares) dimer. The slopes of these data result in an nH of 1.5 ± 0.1 and 1.0 ± 0.1, respectively.

FIGURE 4. The interaction of yNap1 and DNA with histone tetramer. A, normalized fluorescence change as a function of histone tetramer binding to (yNap1m*)-Alexa-546 (closed circles), tail-less (H3/H4), (open squares), and H3(H113A)/H4 (plotted as tetramer) (open squares). Inset of A is the stoichiometry of (H3/H4) to yNap1 dimer. (H3/H4) was titrated against 1 × 10⁻⁷ M (yNap1m*)-Alexa-546 to result in a stoichiometry of 1 histone tetramer to 1 yNap1 dimer. B, normalized fluorescence change as a function of DNA binding to (H3/H4(E63C))²-Alexa-488 (10⁻⁶ M). Buffer conditions were 20–50 mM Tris, pH 7.5, 1 mM dithiothreitol, 0.1 mg/ml bovine serum albumin, and 300 mM NaCl.

ure the affinity of (H3/H4)² tetramer for DNA, we used Alexa-546-(H3/H4(E63C))² (<5-fold the Kd²) and titrated in a 146-base pair DNA fragment (Fig. 4B). These data were fit to a simple binding isotherm with a Kd² of 1.6 ± 0.2 × 10⁻⁹ M. This is ~5-fold tighter than the interaction of (H3/H4)² tetramer with yNap1, and translates into a ΔΔG of ~1.9 kcal/mol. While we cannot say at this point if the (H3/H4)²-DNA interaction is a biologically relevant tetrasome, our results clearly indicate that (H3/H4)² interactions with DNA are more favorable than the interaction with the chaperone.

yNap1 Does Not Distinguish between Major Type Histones and Their Variants—We measured the affinities of complexes containing variants of H2A to yNap1 (Table 2). A dimer of H2A.Z-H2B or H2A. Bbd/H2B has an almost identical Kd app compared to major type H2A/H2B dimer. Together with the similarities in yNap1 affinity for H2A/H2B and H3/H4, our results lead us to conclude that yNap1 recognizes the architecture of the basic histone fold that is common to H3/H4 and H2A/H2B (major type or variant H2A) rather than specific amino acids.

This interpretation is supported by our finding that yNap1 interacts with H2A (refolded in the absence of H2B) with high affinity (Table 2), while H3 binds with very low affinity. H2A is able to form a homo-dimer, most likely through a histone fold dimer-like assembly, whereas H3 alone forms aggregates (as shown by sedimentation equilibrium; data not shown). We were unable to detect any change in signal of (yNap1m*)-Alexa-546 upon addition of homodimeric hMFB, an archaeal histone that binds and compacts DNA (19), indicating that no complex is formed under these conditions.

At Least Two Molecules of Linker Histone H1 Bind yNap1 with High Affinity—yNap1 removes linker histone from chromatin (21), but the affinity of H1 to nucleosomes has been estimated to be at least low nm (22, 23). We measured the affinity of H1 to yNap1 by titrating H1 against yNap1 (yNap1m*)-Alexa-546 (<5-fold the Kd²). These data were best described by a cooperative model with an Kd² of 3.0 ± 0.1 × 10⁻⁹ M and a nH of 1.4 ± 0.1 (Fig. 5). Under stoichiometric conditions (yNap1m*)-Alexa-546 > 10-fold the Kd² we observed the typical linear increase, but it did not level out completely at higher ratios (Fig. 5, inset). The intersection of the two slopes occurs at two H1 histones per yNap1 dimer, but the fact that a second less steep slope is observed suggests an additional mode of H1 interaction with yNap1.

5 M. Resch and K. Luger, unpublished results.
strongly suggest that γNap1 exists as a homodimer (3, 5, 20), consistent with earlier biophysical analyses (6), implying that one γNap1 dimer binds two H2A/H2B dimers or one (H3/H4)2 tetramer (or two H3/H4 heterodimers), respectively.

Earlier in vitro GST pull-down and gel-shift assays concluded that γNap1 has a slight preference for H3/H4; however, this interaction was not observed in in vivo pull-down experiments (9, 10). Our quantitative analysis shows that γNap1 binds H3/H4 with slightly lower affinity than H2A/H2B, irrespective of the ability of H3 to form a heterotetramer via a four-helix bundle structure. In both cases, the basic histone tails had a negative effect on the interaction with γNap1. This counterintuitive result may be explained by an entropic penalty upon binding. While we do not know what effect, if any, post-translational modifications of histones have on the interaction with γNap1, it is intriguing to speculate that the affinity for γNap1 may be regulated by modification of histone tails (21). In light of our finding that complexes of H3/H4 and H2A/H2B bind with very similar affinities, it is not surprising that complexes refolded with histone H2A variants are not distinguished through differences in binding affinities. In contrast, the archaeanal histone Hmfb does not interact with γNap1. This histone-fold dimer resembles eukaryotic histone fold dimers in that it has a basic pi and binds DNA, but it lacks the histone tails and secondary structure elements outside of the histone fold (22, 23).

γNap1 acts as a chaperone for the linker histone H1 in vitro and is capable of removing histone H1 from nucleosomal arrays (24). The affinity of the linker histone H1 to DNA has been estimated to be in the low nm range (25, 26). We find that one γNap1 dimer binds two molecules of linker histone H1 with high affinity; however, additional H1 molecules appear to bind with high affinity at higher H1:γNap1 ratios. Saccharomyces cerevisiae does not have stoichiometric amounts of linker histone; however, we find that metazoan Nap1 binds histones with comparable affinities to γNap1, and we expect the same to be true for linker histones (data not shown).

The binding of all histones to γNap1 is cooperative in nature with the exception of full-length histone dimers and their variants at high ionic strength. We consider at least three possible explanations for this observation. (i) γNap1 or histones could have a monomer to dimer $K_d$ (or dimer to tetramer $K_d$) close to the concentrations we are working at; (ii) γNap1 undergoes a conformational change after the initial histone binding event; or (iii) the histone fold dimers interact with each other upon interaction with a γNap1 dimer. We should be able to observe differences in the binding of γNap1 to histones in scenarios (i) and (iii) by changing the protein being titrated or by studying mutations in histones that interfere with histone-histone interaction. To consider the effects of a γNap1 monomer-dimer transition on the affinities, we worked with several models; however, none were able to adequately explain our data. The possibility that the interaction between the histone fold dimers is responsible for the observed cooperativity is intriguing given that the H2A/H2B dimer binds with a Hill coefficient of 1, whereas the (H3/H4)$_2$ tetramer binds with a Hill coefficient of 2 at 300 mm NaCl. A mutant H3(H113A)/H4 complex which remains in the dimeric state had little effect on
the cooperativity but did alter the affinity. However, yNap1 titration into labeled H3/H4 is not cooperative, while the titration of H3/H4 is cooperative (see Reaction Scheme 2). While the fact that changing the protein being titrated changes the n_H is consistent with the hypothesis that histone-histone interaction is responsible for the observed cooperativity; this hypothesis is not supported by the H3(H113A) mutation. It is possible that yNap1 facilitates the formation of (H3-H4)_2 tetramer and that H3(H113A) does not alter the thermodynamics for tetramer formation enough to change the apparent cooperativity of binding. It is also possible that other histone-histone interactions exist. It is likely that the observed variability in cooperativity cannot be explained by one single the mechanisms presented here. Therefore, histone identity, histone tails, and ionic strength not only change the observed cooperative nature of binding but potentially the mechanism of cooperativity.

It has been proposed that the acidic C-terminal domain (CTAD) of yNap1 helps neutralize the charge of the histones (9). Consistent with this, it has been previously shown that a version of yNap1 lacking the CTAD fails to disassemble nucleosomes in vitro (10). We find that while the effects of removing either the CTAD or the less basic N-terminal tail have only moderate effects on the affinity, the removal of both affects histone binding synergistically. This makes sense in light of the structure of yNap1, where both tails emerge at the concave underside of the dome-shaped molecule (3).

The mechanism by which histone chaperones assemble and disassemble chromatin is largely unknown. The first assumption in a completely thermodynamic mechanism of Nap1-mediated chromatin assembly is that the (H3/H4)_2 tetramer must have a higher affinity for DNA than for yNap1, as shown here. While this does not rule out other more complex mechanisms for chromatin assembly by yNap1, this is consistent with a thermodynamic model. Our data suggest that very small changes in binding energy may have large effects on chromatin/nucleosome structure.

Our working hypothesis is that yNap1 serves to prevent non-correction histone interaction and that only correct biologically relevant structures can compete with yNap1 for histones. Consistent with this we have measured the interaction of H2A/H2B dimer with DNA as ~40 nM.5 The working model stipulates that yNap1 maintains a very low free histone pool to prevent incorrect binding. This model also suggests that the localization of yNap1 or other chaperones to either complexes such as p300 (21) or to the membrane as observed with Arabidopsis thaliana tNAP1 (27) could alter the saturation of nucleosomes on DNA. Alternatively, the possibility of a more direct interaction between yNap1 and the nucleosome, such as the tails of yNap1 interacting with an exposed surface of histone to promote histone release, cannot be excluded.

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