Enhancement of DNA Flexibility in Vitro and in Vivo by HMGB Box A Proteins Carrying Box B Residues[†]

Nadia T. Sebastian, ^{‡,§} Emily M. Bystry, ^{§,II} Nicole A. Becker, ^{II} and L. James Maher III*, ^{II}

Department of Chemistry, Creighton University, 2500 California Place, Omaha, Nebraska 68178, and Department of Biochemistry and Molecular Biology, College of Medicine, Mayo Clinic, 200 First Street, Southwest, Rochester, Minnesota 55905

Received December 12, 2008; Revised Manuscript Received January 16, 2009

ABSTRACT: HMGB proteins are abundant non-histone components of eukaryotic chromatin. The biological function of DNA sequence-nonspecific HMGB proteins is obscure. These proteins are composed of one or two conserved HMG box domains, each forming three α-helices that fold into a sequence-nonspecific DNA-binding module recognizing the DNA minor groove. Box A and box B homology domains have subtle sequence differences such that box B domains bend DNA strongly while DNA bending by isolated box A domains is weaker. Both box A and box B domains preferentially bind to distorted DNA structures. Here we show using DNA cyclization kinetics assays in vitro and *Escherichia coli* DNA looping assays in vivo that an isolated HMG box A domain derived from human HMGB2 folds poorly and does not enhance apparent DNA flexibility. Surprisingly, substitution of a small number of cationic residues from the N-terminal leader of a functional yeast box B protein, Nhp6Ap, confers the ability to enhance DNA flexibility. These results demonstrate important roles for cationic leader amino acids in HMGB folding, DNA interaction, and DNA bending.

HMGB¹ sequence-nonspecific DNA binding proteins are abundant non-histone proteins in eukaryotic cells (I-3). These small proteins are composed of one [e.g., Saccharomyces cerevisiae Nhp6Ap (4-8) or Drosophila melanogaster HMG-D (9, 10)] or two [e.g., vertebrate HMGB1, HMGB2 (II-15), and S. cerevisiae Hmo1p (I6)] repeats of a conserved \sim 80-residue "HMG box" motif. Highly charged flanking sequences are also often present (3). Thus, HMGB proteins are a subset of the proteins that carry one or two domains homologous to the ancestral HMG box. HMGB proteins accumulate to greater than micromolar concentrations in eukaryotic cells such that there is approximately one HMGB protein for every 5-10 nucleosomes in chromatin.

Among the striking features of HMGB proteins are their ability to induce strong DNA bending and, consequentially, their preference for binding to distorted DNA structures such

as cruciforms (11, 16-18) and chemical lesions (19-23). DNA bending in these complexes can exceed 90°. In this sense, HMGB proteins are functionally (though not structurally) reminiscent of the prokaryotic nucleoid protein HU (24). There is evidence that HU functions by inducing a flexible hinge in DNA (25, 26), and HMGB proteins appear to alter DNA structure through formation of hinges with enhanced flexibility (27-29).

The biological functions of HMGB proteins are not understood. Initial insight into HMGB function came from pioneering studies by Johnson and co-workers (6, 7, 12). These authors found that HMGB proteins could substitute for prokaryotic HU proteins in transactions involving tightly bent DNA, such as recombination (12). These observations were extended to living Escherichia coli where defects associated with loss of the HU nucleoid protein were suppressed by HMGB expression (7). There is also evidence that HMGB proteins facilitate the DNA binding of some sequence-specific DNA binding transcription factors and recombinases (reviewed in ref 3). In some cases, this HMGB participation appears to be catalytic (30). The Nhp6A protein of budding yeast has been shown to enhance transcription activation at least in part through facilitation of Gal4p binding to DNA (6). Similarly, lethal hypoglycemia in newborn mice lacking HMGB1 may be due to defective glucocorticoid receptor binding to DNA (31). DNA bending to facilitate protein-protein interactions may be an important role of HMGB proteins in the function of enhancesomes (32, 33). On the other hand, HMGB proteins participate as subunits of the FACT complex required for efficient transcription of chromatin templates in yeast (34). In addition, the chromatin of yeast cells lacking the two-box HMGB protein Hmo1p

[†] Supported by the Mayo Foundation and NIH Grant GM 75965 to L.J.M.

^{*} To whom correspondence should be addressed. Telephone: (507) 284-9041. Fax: (507) 284-2053. E-mail: maher@mayo.edu.

[‡] Creighton University.

[§] These authors contributed equally to this work.

II Mayo Clinic.

 $^{^1}$ Abbreviations: BSA, bovine serum albumin; box A, N-terminal high-mobility group homology domain; box B, C-terminal high-mobility group homology domain; EDTA, ethylenediaminetetraacetic acid; FACT, facilitator of chromatin transactions; HEPES, 4-(2-hydroxyethyl)-1-piperazineethylsulfonic acid; HMG, high-mobility group; HMGB, high-mobility group B protein family; HMGB1, mammalian member 1 of the HMGB protein family; HMGB2, mammalian member 2 of the HMGB protein family; HPLC, high-performance liquid chromatography; IPTG, isopropyl β -D-1-thiogalactopyranoside; J factor, concentration of one DNA end in the vicinity of the other; LB, Luria-Bertani broth; Ni-NTA, Ni^2+-nitrilotriacetic acid; PDB, Protein Data Bank; TE, Tris-HCl, EDTA buffer; TLC, thin layer chromatography.

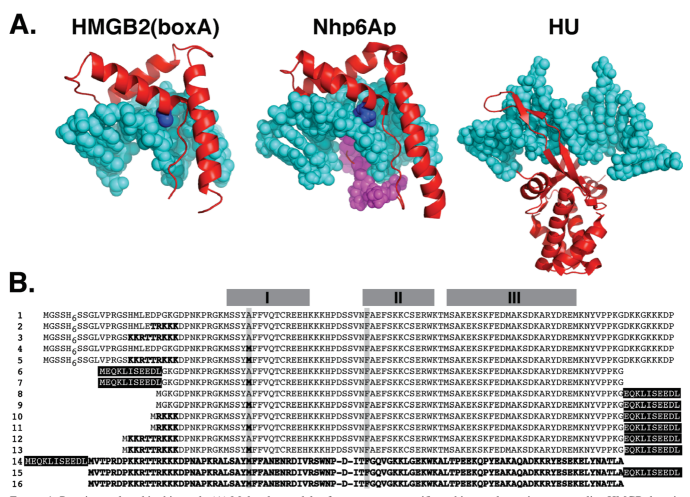


FIGURE 1: Proteins analyzed in this work. (A) Molecular models of sequence-nonspecific architectural proteins: mammalian HMGB domain HMGB2(box A) [left, PDB entry 1ckt (21)], yeast HMGB protein Nhp6Ap [center, PDB entry 1j5n (4)], and *E. coli* HU [right, PDB entry 1p51 (25)]. Bent DNA is rendered as cyan spheres, and protein is shown as a red ribbon. Potential intercalating residues in the HMGB proteins are colored blue. The cationic N-terminal leader of Nhp6Ap is colored magenta. (B) Sequence alignments of wild-type (WT) and chimeric HMGB proteins. Helical regions I—III are indicated above. Potential intercalating residues are indicated by vertical shading. Residues derived from HMGB2(box A) and Nhp6Ap are shown in plain text and bold text, respectively. The c-Myc epitope tag decapeptide is indicated by black rectangles. Proteins 1–5 and 16 were tested in vitro for their ability to increase the *J* factor of 200 bp DNA fragments. Proteins 6–15 were tested in *E. coli* for their ability to complement the *lac* repression defect of *hupA/B*Δ cells.

becomes sensitive to micrococcal nuclease digestion (35). These results suggest multiple roles for HMGB proteins in chromatin structure and function (36).

We became interested in HMGB proteins after observing that the activator dependence of in vitro transcription from plasmid DNA templates in HeLa cell nuclear extract was not consistent with the expectations of DNA looping over short distances (37). We subsequently confirmed that mammalian nuclear extracts are rich in a heat-resistant activity that dramatically enhances the apparent flexibility of DNA (38). This work also shows that recombinant HMGB proteins could substitute for at least some of this activity. We hypothesized that HMGB proteins increase the apparent flexibility of DNA in regions not packaged with histone octamers. Recent studies using single-molecule force spectroscopy confirmed that apparent DNA persistence length decreases substantially in the presence of low concentrations of HMGB proteins, while higher HMGB concentrations can produce a more rigid protein-DNA filament (28, 29).

To study apparent DNA physical properties in vivo, we have recapitulated classic experiments of Record (39–41) and Müller-Hill (42–46) involving DNA looping in the *E. coli lac* operon. Our work has confirmed that the physical

properties of DNA measured in vivo suggest enhanced bending and twisting flexibilities relative to in vitro expectations and that nucleoid proteins modify these properties (47, 48). We have recently shown that certain eukaryotic HMGB proteins can complement the loss of bacterial HU (49). Thus, effects on in vivo DNA looping in *E. coli* can provide a sensitive bioassay for HMGB function.

The HMG box sequence encodes three α -helices that fold into an L-shaped structure (Figure 1A). This module interacts with the DNA minor groove to both widen the groove and allow partial intercalation of one or more amino acid side chains between stacked base pairs (3). The combination of these perturbations can cause strong DNA bending away from the engaged DNA surface. HMG box A of human HMGB2 (Figure 1A, left) is shown bound to a DNA segment distorted by platinum cross-linking (21). Also illustrated in Figure 1A (center) is the structure of the yeast single-box protein Nhp6Ap complexed with unmodified DNA (4). Putative intercalating residues are colored blue. Importantly, Nhp6Ap also contains a strongly cationic leader sequence (Figure 1A, center, magenta) disposed in the major groove to asymmetrically neutralize the compressed sugar-phosphate backbone. There is evidence that this interaction contributes

significantly to DNA bending (50). Thus, sharp DNA bending by HMGB proteins can involve minor groove widening, amino acid wedging, and asymmetric charge neutralization. Favorable electrostatic forces have been shown to dominate other interactions in both DNA binding and bending by sequence-nonspecific HMGB proteins (50). It is noteworthy that strong and sequence-nonspecific bending by the prokaryotic nucleoid protein HU involves minor groove interactions, but within a completely different scaffold (Figure 1A, right).

Though similar in sequence and structure, it has been noted that subtle features of the "A box" domain of the two-box HMGB1 protein distinguish it from the "B box" domain (3). The A box domain is distinct in the shape and orientation of helix I, and the identity of potential intercalating residues. Isolated A box domains are reportedly less effective at DNA bending (12). The sequence of a His6-tagged version of human HMGB2(box A) is illustrated as construct 1 in Figure 1B. Whereas the intercalating phenylalanine in helix II is present in both box A and box B domains of two-box HMGB proteins, the alanine in helix I does not intercalate strongly (3). Furthermore, box A and box B domains are distinguished by the relative lengths and geometries of helices I and II, with helix I being short and straight in box A, but bent in box B (3). The single-box HMGB protein Nhp6Ap conforms to the box B family and is shown as construct 16 in Figure 1B. Nhp6Ap is distinguished from HMGB2(box A) by the identity of the intercalating residue in helix I (methionine in Nhp6Ap), a longer helix I, and a highly cationic leader sequence.

In this study, we apply in vitro and in vivo assays to understand what minimal features of an HMG B box confer on an A box the ability to enhance apparent DNA flexibility.

MATERIALS AND METHODS

Bacterial Strains and Gene Disruptions. The hupA and hupB genes in parental E. coli strain FW102 (51) were disrupted as described previously (47, 52). Diagnostic PCR amplification following conjugation and selection was used to confirm strain deletion status and the presence of the proper looping assay episomes.

Expression and Purification of HMGB Proteins. Plasmid pJ583 encodes human HMGB2(box A) cloned in-frame with a His₆ tag in plasmid pET-15b (Novagen). Mutations were introduced using the QuickChange site-directed mutagenesis kit (Stratagene). Plasmids encoding HMGB2(box A) and variants were transformed into E. coli BL21-Gold (Stratagene), and 6 mL overnight cultures were prepared from single colonies in LB medium containing carbenicillin. Five milliliters of overnight culture was washed with fresh LB to remove secreted β -lactamase, and the cells were used to seed a 250 mL LB culture. Cells were grown at 37 °C with shaking until the culture reached a cell density corresponding to an OD_{600} of 0.6. IPTG was added to a final concentration of 1 mM, and cells were grown at 37 °C for 3 h with shaking, pelleted by centrifugation at 6000g, and stored at -80 °C. The induced cell pellet was then resuspended in 10 mL of binding buffer (Novagen) per 250 mL of pelleted cells and sonicated on ice with 15 s pulses separated by 1 min cooling intervals. The lysate was then subjected to centrifugation at 15000g for 20 min at 4 °C, and the supernatant was recovered. His6-tagged protein was purified using Ni-NTA agarose resin (Qiagen) per the manufacturer's recommendations. Briefly, washed Ni-NTA agarose resin was added in a 1:4 (v:v) ratio to the lysate, the mixture gently rotated at 4 °C for 1 h, and the resin collected by centrifugation at 2000 rpm for 2 min at 4 °C in a clinical centrifuge. Beads were washed three times with wash buffer (10 mL), and protein was eluted from the resin twice by gentle rotation in 10 mL of elution buffer for 5 min at 4 °C. The eluted sample volume was reduced to 2 mL using 5000 molecular weight cutoff centrifugal cartridges (Vivaspin), and proteins were dialyzed in 3500 molecular weight cutoff cassettes (Slidea-Lyzer, Pierce) at 4 °C against 1 L of buffer containing 20 mM HEPES (pH 7.5), 100 mM KCl, 1 mM EDTA, and 1 mM DTT, followed by a second dialysis against the same buffer containing 5% glycerol. Proteins were then purified by reverse phase HPLC on a Jupiter C18 column [250 mm \times 21.2 mm, 15 μ m (Phenomenex, Torrance, CA)] in a 0.1% TFA/water mixture with a 50 min gradient from 10 to 70% B, where B was 80% aqueous acetonitrile with 0.1% TFA. Lyophilized proteins were resuspended and characterized by mass spectrometry.

HMGB Protein Characterization in Vitro. CD spectra were recorded using a Jasco 810 spectropolarimeter in continuous mode, collecting measurements every 1 nm between 200 and 260 nm at 295 K with an averaging time of 5 s. Protein sample concentrations were 2–10 μ M in 20 mM Tris-HCl (pH 8.0), 30 mM KCl, and 10 mM MgCl₂ with or without addition of 400 μ M (base pairs) duplex DNA. All buffer and nucleic acid contributions to ellipticity were subtracted after data collection, and raw ellipticity data were converted to mean residue ellipticity (MRE).

Fluorescence anisotropy measurements were used to characterize DNA-protein interactions and the fraction of active protein. A 22 bp fluorescein-labeled duplex DNA probe was created by the annealing of oligonucleotides LJM-1100 (5'-AGT₂GAG₄ACT₃C₃AG₂C-F, where F represents fluorescein) and LJM-1101 (5'-GC₂TG₃A₃GTC₄TCA₂CT-3'). Fluorescein labeling efficiency was confirmed by comparing absorbance measurements at 495 and 260 nm. Ten nanomoles of cDNA strands was mixed in 48.5 μ L of TE buffer (pH 8.0); 1.5 μL of 1 M NaCl was added, and the sample was annealed (92 °C for 10 min, 65 °C for 5 min, 37 °C for 1.5 h, and 22 °C for 30 min). The resulting duplex was purified by electrophoresis through a 12% native polyacrylamide gel in $0.5 \times$ TBE buffer. The fluorescent probe duplex was cut from the gel after detection over a TLC indicator plate under ultraviolet illumination and eluted overnight in 300 mM NaOAc (pH 4.5), precipitated from ethanol, suspended in TE buffer, and quantitated. Fluorescence anisotropy measurements for equilibrium dissociation constant estimation were performed in 96-well microtiter plates with 200 μ L binding reaction mixtures containing fluorescent probe (5 nM) and protein dilutions in 20 mM Tris-HCl (pH 8.0), 30 mM KCl, 10 mM MgCl₂, and 100 μ g/mL BSA. Protein dilutions were prepared in binding buffer with 10% glycerol and 100 µg/mL BSA. The Safire2 instrument (TECAN) was controlled with X-Fluor4. Data were fit to eq 1 to estimate the equilibrium dissociation constant.

$$\theta = \frac{A_{\text{obs}} - A_{\text{min}}}{A_{\text{max}} - A_{\text{min}}} \times \frac{P_{\text{t}}}{K_{\text{d}} + P_{\text{t}}}$$
(1)

where A indicates sample anisotropy, subscripts obs, min, and max represent observed, minimum, and maximum, respectively, $P_{\rm t}$ is the total concentration of added protein, and $K_{\rm d}$ is the fit equilibrium dissociation constant. $P_{\rm t} \sim P_{\rm free}$ under these binding conditions of large protein excess over fluorescent DNA probe. Estimation of the protein active fraction employed similar methods, except titration was performed with a fluorescent duplex DNA probe concentration of 250 nM ($\gg K_{\rm d}$), and the discontinuity in the titration plot indicated the point of protein/DNA concentration equivalence.

J Factor Estimate by Ligase-Mediated DNA Cyclization Assay. Plasmids pJ823, pJ825, pJ827, pJ829, pJ831, and pJ833 contain intrinsically straight ~200 bp sequences and were the kind gifts of A. Vologodskii (53). Probe sequences were amplified by PCR using primers LJM-3222 (5'-G₃TA₂-CGC₂AG₃T₄-3') and LJM-3223 (5'-TGTGAGT₂AGCTCAC-TCAT₂AG₂-3') such that an additional \sim 100 bp was included upstream and downstream of the HindIII sites defining the probe. Polymerase chain reaction mixtures included plasmid template (20 ng), primers (400 nM), BSA (100 µg/mL), Taq DNA polymerase buffer (Invitrogen), MgCl₂ (4 mM), dNTPs (200 μ M each), 50 μ Ci of [α -³²P]dATP, and 3.2 units/ μ L Taq DNA polymerase (Invitrogen). Cycle parameters were as follows: 3 min at 98 °C; 30 cycles of 30 s at 94 °C, 30 s at 55 °C, and 45 s at 72 °C; 5 min at 72 °C; held at 4 °C. PCR products were treated with 50 ng/µL proteinase K in the presence of 5 mM EDTA, 10 mM Tris-HCl (pH 8.0), and 0.5% SDS at 37 °C for 1 h. The DNA was extracted with a 1:1 phenol/chloroform mixture and precipitated from ethanol. The DNA was digested overnight at 37 °C with 40 units of HindIII (New England Biolabs) in the presence of 100 µg/mL BSA. Digested product was loaded onto a 5% native polyacrylamide (29:1 acrylamide:bisacrylamide) gel and visualized by exposure of the wet gel to BioMAX XR film. The desired \sim 200 bp DNA product was cut from the gel and eluted overnight at 22 °C in 50 mM NaOAc buffer (pH 7.0). The final DNA concentration was obtained by determining the fraction of total original radiolabel present in the purified product, multiplying this fraction by the total concentration of dATP in the labeling reaction, and dividing this result by the number of dA residues in each duplex DNA product.

DNA cyclization reactions (10 μ L) were performed with 10 nM DNA probe and the indicated concentration of HMGB protein in T4 DNA ligase buffer [20 mM Tris-HCl (pH 8.0), 30 mM KCl, 100 μ g/mL BSA, 1.8 mM ATP, and 10 mM MgCl₂], and T4 DNA ligase [20–50 units/ μ L (New England Biolabs)] at 22 °C. Reactions were stopped at the indicated times by adjusting the EDTA concentration to 20 mM. DNA was purified using QIAQuick PCR clean up columns (Qiagen). DNA was eluted in 30 μ L of elution buffer, and 15 μ L samples were loaded onto 5% (29:1 acrylamide: bisacrylamide) native polyacrylamide gels. After electrophoresis in 0.5× TBE buffer, gels were dried and exposed to storage phosphor screens. Circular DNA products were identified in preliminary experiments by their resistance to *Bal*31 exonuclease (New England Biolabs).

Image quantitation was performed with ImageQuant. Intensities of the monomer circle (MC) product and the sum of linear dimers and all other ligation products (LD) were determined for each time point and corrected for background signal, and their ratio was plotted versus time. The J factor was determined using eq 2:

$$J = 2M_0 \lim_{t \to 0} (MC/LD)$$
 (2)

where M_0 is the initial DNA concentration and the indicated limit is the *y*-axis intercept of the plot. Conditions of time, ligase concentration, and initial DNA concentration were empirically determined for each DNA probe and HMGB protein so that labeled starting material was not significantly depleted and linear fits were obtained.

Cloning and Expression of HMGB Proteins in Bacteria. HMG protein expression constructs were created by inserting purified PCR products into plasmid pJ1035, a modified version of pLX20 (51) containing a promoter for a moderate level of protein expression and an inactive *lac* operator. In addition, a 10-amino acid c-Myc epitope tag (EQKLISE₂DL) was added either to protein N-termini (constructs 6 and 7) or to C-termini (constructs 8-13) to facilitate analysis of accumulation. To create plasmid construct 8 (pJ1491), forward primer LJM-3314 (GCTCTAGA₂TG₃TA₃G₂AGAC₄-A₂C) and reverse primer LJM-3318 [CGA₂GCT₂GCAGC-TA*CAGATCT*₂*CT*₂*CAGA*₃*TA*₂*GT*₅*GT*₂*C*AC₂T₃G₃AG₂A₂C (c-Myc tag in bold italics)] were used to amplify the portion from M1 to G83 of HMGB2(box A). Plasmid construct 6 (pJ1493) was created using forward primer LJM-3315 (GCT-CTAGA₂TG*GA*₂*CA*₅*CT*₂*AT*₃*CTGA*₂*GA*₂*GATCTG*G₂TA₃G₂-AGAC₄A₂C) and reverse primer LJM-3327 (CGA₂GCT₂-GCAGCTA₂C₂T₃G₃AG₂A₂CGTA₂T₅C). Plasmid construct 10 was created by PCR amplification using forward primer LJM-3621 (GCTCTAGA₂TGAGA₃GA₂GA₂GA₂GA₂AC₄A₂CA₂G-C₂G) and reverse primer LJM-3318. Plasmid construct 12 was created with forward primer LJM-3620 (GCTCTAGA₂-TGA₂GA₂GAGA₂C₂ACTAGA₃GA₂GA₂GA₂GA₂CA₂CA₂GC₂G) and reverse primer LJM-3318. All PCR products were inserted between XbaI and HindIII restriction sites of pJ1035. Site-directed mutagenesis (Stratagene) employed primers LJM-3613 (CA₄TGTC₂TCGTACATGT₂CT₂CGTGCAG-AC) and LJM-3614 (GTCTGCACGA₂GA₂CATGTACGA-G₂ACAT₄G) to introduce the A17M mutation in constructs 7, 9, 11, and 13 from constructs 6, 8, 10, and 12, respectively. Accumulation of epitope-tagged HMGB proteins was monitored by Western blotting of E. coli extracts using primary antibody op10T (CalBiochem) as previously described (49).

In Vivo DNA Looping Assay. DNA looping constructs were based on plasmid pJ992, created by modifications of pFW11-null (51) as previously described (47–49). Reporter gene assays were performed as previously described (49).

RESULTS

Experimental Strategy. The experimental strategy for comparing activities of single-HMGB box chimeric proteins is shown in Figure 2. In vitro analyses of DNA flexibility enhancement were performed as shown in panel A. Purified His₆-tagged HMGB protein was mixed with a radiolabeled sample of 200 bp intrinsically straight duplex DNA containing *Hind*III cohesive termini (53) and incubated with T4

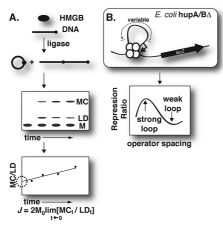


FIGURE 2: Experimental strategy. (A) In vitro measurement of HMGB protein enhancement of apparent DNA flexibility by ligase-catalyzed DNA cyclization kinetics. The top panel shows the HMGB protein was incubated for various times with \sim 200 bp radiolabeled DNA (M) in the presence of T4 DNA ligase to produce a mixture of intramolecular monomer circles (MC) and products derived from linear dimers (LD). The middle panel is the electrophoretic analysis. The bottom panel shows quantitation and extrapolation estimating the J factor. (B) The top panel shows in vivo measurement of HMGB protein complementation of the *lac* repression loop defect in $hupA/B\Delta$ E. coli cells. Lac operator spacing was varied between 75 and 85 bp in an E. coli hupA/B deletion strain that is disabled in lac repression looping. The bottom panel shows the strength of DNA repression looping as a function of operator spacing, and HMGB protein expression is monitored by the repression ratio.

DNA ligase for different times to generate a mixture of labeled products (Figure 2A, top). These DNAs were separated by native polyacrylamide gel electrophoresis and quantitated by storage phosphor technology (Figure 2A, middle). The ratio of monomer circles (MC) to all other species, including (and derived from) linear dimers (LD), is extrapolated to time zero to estimate the J factor [Figure 2A, bottom (54)]. The J factor, in turn, expresses the effective local concentration of one terminus of the 200 bp DNA with respect to the other, a parameter that should increase with an increase in longitudinal DNA flexibility. The J factor also measures DNA twist because cohesive termini must be in proper rotary alignment for ligation. Our simulations of ring closure kinetics in the presence of HMGB proteins (L. Czapla, E. Bystry, W. Olson, and L. J. Maher, manuscript in preparation) have shown that J factors can be influenced by protein-induced changes in both DNA twist and DNA bending at the site of the bound protein. In addition, we have shown that bound HMGB proteins cause enhanced anisotropic DNA flexibility in the protein-DNA complex (27). This combination of effects (changes in DNA twist, DNA bending, and anisotropic DNA flexibility) increases the J factor associated with the ensemble of DNA complexes present during the ligation reaction.

As shown in Figure 2B, in vivo studies of single-HMGB box chimeric proteins were performed by expression of the epitope-tagged proteins in E. coli. The experimental strains harbored F' episomes engineered to carry a lacZ reporter gene driven by a promoter flanked by two lac operators (47). The upstream operator strongly binds lac repressor, while the downstream operator is weaker. Operator spacing is varied so that the stability of the *lac* repression loop (Figure 2B, top) varies with DNA looping energy. This relationship can be monitored

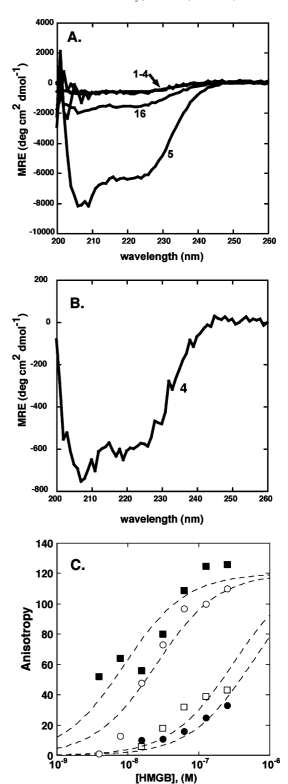


FIGURE 3: HMGB protein characterization in vitro. (A) Analysis of protein folding by circular dichroism spectroscopy showing mean residue ellipticity (MRE) as a function of wavelength for protein constructs 1-5 and 16. (B) Rescaled plot of CD data for construct 4 showing a weak α-helical folding signature. (C) Example HMGB protein DNA binding data obtained by fluorescence anisotropy. Raw anisotropy values are fit to a simple binding isotherm to estimate the equilibrium dissociation constant as described in Materials and Methods: construct 1 (\bullet), construct 2 (\square), construct 3 (\bigcirc), and construct 5 (■).

as the ratio of induced to uninduced β -galactosidase activity as shown schematically in the bottom panel of Figure 2B.

Table 1: In Vitro Characteristics of HMGB Chimeric Proteins

			$\boldsymbol{\Theta}_{222}{}^{c}$		anisotropy d		
construct ^a	length ^b	pI	without DNA	with DNA	$K_{\rm d}$ (nM)	% active	J factor $(nM)^e$
1	116	9.69	-614	-711	570 ± 40	nd^g	1.5 ± 0.7
2	116	9.9	-619	-398	268 ± 27	nd^g	9.6 ± 3.2
3	116	10.1	-730	-311	31 ± 6	40	28.2 ± 4.7
4	116	9.69	-600	-660	103 ± 20	nd^g	2.2 ± 0.6
5	116	10.1	-6354	-2567	11 ± 2	100	14.0 ± 2.5
16	93	9.72	-1585	-1488	10 ^f	40	8.2 ± 0.7

^a See Figure 1. ^b Amino acid residues. ^c MRE by CD (deg cm² dmol⁻¹). ^d Fluorescence anisotropy measurements as described in Materials and Methods. K_d estimates (average \pm standard error) were based on three determinations. ^e Determined for a 200 bp DNA fragment (intrinsic J factor of 1.5 \pm 0.3 nM) as described in Materials and Methods. ^f Determined previously from pooled data. ^g Not determined because of insufficient DNA affinity.

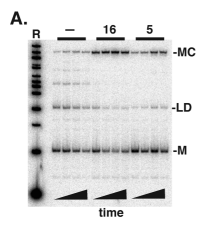
In Vitro Characterization of Single-HMGB Box Chimeric *Proteins.* HMGB protein variants 1–5 and 16 (Figure 1B) were expressed in E. coli and purified. The proteins represent HMGB2(box A) (construct 1), Nhp6Ap (construct 16), and chimeras differing in terms of the identity of the primary intercalating residue in helix I (constructs 4 and 5) and the amount of basic leader sequence substituted from Nhp6Ap (constructs 2, 3, and 5). Proteins were initially characterized by circular dichroism spectroscopy to monitor folding in the absence of DNA (Figure 3 and Table 1). Nhp6Ap (construct 16) exhibited a moderate ellipticity signal consistent with α -helix formation in the absence of DNA (Figure 3A). The corresponding α-helical signal from construct 5 (seven cationic leader substitutions and methionine as the primary intercalating residue) was particularly strong (Figure 3A). Interestingly, signals from constructs 1-4 were all weak (Figure 3A), though clearly α -helical (Figure 3B). Surprisingly, CD signals did not substantially change in the presence of excess DNA (Table 1), suggesting that the various similar HMG box proteins strongly differed in their folding stabilities.

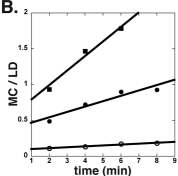
DNA binding was further characterized by fluorescence anisotropy measurements. Example data are shown in Figure 3C, with compiled results listed in Table 1. Equilibrium dissociation constants for DNA binding ranged from 10 nM (Nhp6Ap, construct 16) to >500 nM [HMGB(box A), construct 1] with affinities decreasing as follows: $16 \sim 5 > 3 > 4 > 2 > 1$ (Table 1). In the cases where strong DNA binding activity was detected, the active protein fraction was measured by binding titration in the presence of DNA duplex concentrations much higher than K_d and found to be in the range of 40-100% (Table 1). Overall, these data suggest that both intrinsic protein folding stability and DNA interaction are strongly correlated with the protein isoelectric point (Table 1).

In Vitro Enhancement of Apparent DNA Flexibility by Single-HMGB Box Chimeric Proteins. Results of in vitro cyclization kinetics experiments are shown in Figure 4. Examples of electrophoretic data fitting results for J factor assays of DNA probe alone (solid line) or in the presence of 40 nM HMGB proteins 16 and 5 (Figure 1B) are shown in panels A and B of Figure 4. Estimated J factors are given in Figure 4C and Table 1. Consistent with previous studies (53), the naked 200 bp DNA duplex is characterized by a Jfactor of 1.5 nM. This DNA flexibility is significantly enhanced by incubation with 40 nM construct 16 (Nhp6Ap) but not by incubation with human HMGB2(box A), even when the primary intercalating residue of helix I is changed from alanine to methionine (Figure 4C, constructs 1 and 4). Interestingly, certain chimeras between Nhp6Ap and HMGB2-(box A) exhibited activities at least as strong as that of Nhp6Ap. For example, substitution of the leader residues of HMGB2(box A) (construct 1) with a small number of the corresponding cationic residues from Nhp6Ap (Figure 4C, constructs 2 and 3) resulted in strong enhancement of apparent DNA flexibility, independent of whether the primary intercalating residue of helix I was alanine (constructs 2 and 3) or methionine (construct 5). These results strongly correlate with the predicted protein isoelectric point and observed DNA binding affinity, indicating that a cationic leader is sufficient to endow any of these HMGB2(box A) variants with the ability to enhance apparent DNA flexibility. Asymmetric phosphate neutralization of the compressed major groove (as in Nhp6Ap) may play a role in this effect.

In Vivo Characterization of Single-HMGB Box Chimeric *Proteins.* The results of in vitro experiments showed surprisingly strong effects of N-terminal cationic leader sequences on protein folding, DNA binding, and enhancement of DNA flexibility by HMG domains. We wished to rule out the possibility that these effects were due to artifacts of protein purification and/or refolding. The same chimeric single-HMG box proteins were therefore evaluated in vivo for their abilities to complement the *lac* repression looping defect observed in living hupA/BΔ E. coli cells (47, 48). Accumulation of epitope-tagged HMGB constructs 6–15 (Figure 1B) was first assessed by Western blotting as shown in Figure 5. Panel A of Figure 5 indicates that epitope tagging of the N-terminus supports an increased level of HMGB protein accumulation (constructs 6 and 8 are otherwise identical). The exposure in panel B of Figure 5 is shorter, allowing visualization of the comparable accumulation levels of constructs 6, 7, and 10–13. Panel C of Figure 5 compares accumulation of constructs 14 and 15. Importantly, although construct 15 (C-terminally epitope-tagged Nhp6Ap) accumulates to the lowest level of all tested HMGB constructs, even this level of accumulation has previously been shown to be sufficient for strong complementation of the $hupA/B\Delta$ repression looping defect (49). Thus, all of the tested HMGB proteins and chimeras accumulate within E. coli cells at levels with the potential to complement the $hupA/B\Delta$ defect.

lac repression looping in hupA/B Δ E. coli cells was assessed for operator spacings in the range of 75–85 bp. The results are shown in Figure 6. Top panels show repression ratio data (ratio of induced to uninduced β-galactosidase activity). Higher peak values indicate favorable DNA loop geometries. Middle and bottom panels display β-galactosidase output (E') in the presence (middle panel) or absence (bottom panel) of IPTG inducer. E' normalizes the raw reporter gene signal in a given genetic strain background to the output from an unlooped promoter lacking an upstream lac operator in that same strain. Higher E' values





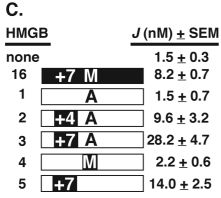


FIGURE 4: In vitro assay of DNA flexibility enhancement by HMGB proteins and chimeras. (A) Example data from a T4 DNA ligase cyclization assay for 200 bp DNA probe in the absence (-) and presence of 40 nM HMGB constructs 16 and 5 (see Figure 1). (B) Graphical analysis and extrapolation to estimate J factor. Data for DNA probe alone (\bigcirc), construct 16 (\bigcirc), and construct 5 (\blacksquare) are shown. (C) Summary data for in vitro measurement of effects of HMGB proteins and chimeras on the J factor of 200 bp DNA. The residue at potential intercalating position 17 (alanine vs methionine) and upstream cationic chimeras (RKKK "+4" vs KKRTTRKKK "+7") are highlighted schematically (see Figure 1B). Black and white regions indicate sequences derived from Nhp6Ap and HMGB2(box A), respectively.

indicate more gene expression (less energetically favorable DNA repression loops). Figure 6A compares repression in WT and $hupA/B\Delta$ genetic backgrounds. Shaded regions highlight the increased promoter "leakiness" in cells lacking HU (white symbols).

Graphs B-F of Figure 6 compare complementation of the hupA/B∆ looping defect by various single-HMG box chimeric proteins. The bottom panels of Figure 6 (reporter activity under repressed conditions) are especially informative. Box A derived from HMGB2 could not complement the $hupA/B\Delta$ looping defect, regardless of epitope tag



FIGURE 5: In vivo expression of HMGB proteins and chimeras. Accumulation of the indicated proteins (see Figure 1B) was analyzed by Western blotting. (A) Exposure comparing expression of HMGB2(box A) variants carrying C-terminal Myc tags (constructs 8 and 9) vs an N-terminal tag (construct 6) or vector control (v). (B) Shorter exposure showing expression levels of other HMGB2(box A)-Nhp6Ap chimeric HMGB proteins. (C) Comparison of expression levels of the Nhp6A protein carrying an N-terminal (construct 14) or C-terminal (construct 15) epitope

location (Figure 6B). In contrast, both N- and C-terminally epitope-tagged versions of yeast Nhp6Ap were active in complementation, with the more strongly expressed Nterminally epitope-tagged construct 14 being particularly effective (Figure 6C). Regardless of epitope tag location, replacement of alanine with methionine as the primary intercalating residue in helix I of HMGB2(box A) did not confer complementation activity (Figure 6D). When the Nhp6Ap contribution to the HMGB2(box A) construct was limited to four basic amino acids in the leader (RKKK), complementation depended on the identity of the primary intercalating amino acid in helix I. A protein with alanine in this position (Figure 6E, construct 10) had no detectable function. In contrast, the combination of RKKK leader and primary intercalating methionine (Figure 6E, construct 11, note bottom panel) supported complementation even though the methionine substitution was ineffective when tested alone (Figure 6D, construct 9). Finally, as observed for DNA flexibility enhancement in vitro (Figure 3), substitution of Nhp6Ap leader sequence KKRTTRKKK endowed the otherwise nonfunctional HMGB2(box A) construct with the ability to fully complement DNA looping in $hupA/B\Delta$ E. coli, independent of whether alanine or methionine was in the primary intercalating position of helix I (Figure 6F).

DISCUSSION

We are interested in understanding features of HMG boxes A and B that distinguish their abilities to enhance apparent DNA flexibility in vitro and in vivo. Discussion has focused on differentiating characteristics of helix I and the loop between helices I and II (3, 10). For example, the Drosophila single-HMG box B protein HMGD has been carefully analyzed (10). A sequence-nonspecific HMGB protein, HMGD resembles yeast Nhp6Ap in its bulky methionine residue at the primary intercalating position of helix I, where HMGB(box A) has alanine at this position. Mutation of methionine to alanine in HMGD somewhat reduced the thermal stability of the protein (2 °C) and its DNA binding affinity (6-fold) (10). Importantly, these authors found that residues near the primary intercalating amino acid in helix I also played roles in DNA binding and bending. These results clearly point to the importance of the primary intercalating residue for tuning function. On the other hand, these mutations of the primary intercalating residue of HMGD caused relatively subtle effects on DNA binding and bending (10). In fact, detailed thermodynamic analysis indicates that it is electrostatic interactions that drive sequence-nonspecific

FIGURE 6: Complementation analysis of the *lac* repression looping defect in $hupA/B\Delta$ cells by expression of HMGB proteins and chimeras. Top panels show the repression ratio. The shaded reference regions illustrate the differences in repression ratio behavior for WT and mutant $E.\ coli.$ The shaded region is replicated in all top panels for reference. The bottom panels show reporter gene expression (E') in the presence (middle) or absence (bottom) of IPTG. The shaded regions are replicated in all bottom panels for reference to contrast reporter gene expression in WT and $hupaA/B\Delta\ E.\ coli$ cells.

HMGB proteins onto DNA (50, 55). In addition, results of fluorescence resonance energy transfer experiments showed that the positively charged leader sequence of yeast Nhp6Ap (Figure 1A, magenta spheres; Figure 1B, construct 16) contributes significantly to DNA bending (50). Strongly cationic sequences are found N-terminal to helix I in Nhp6Ap, and C-terminal to helix III in HMGD and HMGB1/ 2. The HMG box fold places both N-terminal and C-terminal sequences in approximately the same position near the major groove of DNA where both release of condensed counterions and asymmetric phosphodiester charge neutralization could enhance binding affinity and DNA bending. We therefore wished to compare the functional significance of (i) a primary intercalating residue in helix I and (ii) a cationic leader sequence for HMGB function. We have studied HMGB2(box A) as an example.

We previously showed that budding yeast Nhp6Ap (a member of the box B family of HMGB domains) could complement loss of HU protein in an E. coli assay of DNA looping in the *lac* operon, while human HMGB2(box A) could not (49). This result suggested an experimental approach to understanding functional features distinguishing Nhp6Ap from HMGB2(box A). In this study, we constructed a small number of chimeric proteins by substituting the primary intercalating methionine residue of Nhp6Ap helix I and /or cationic segments of the Nhp6Ap leader with corresponding sequences in HMGB2(box A). Our in vitro and in vivo results show that substitution of cationic leader residues of Nhp6Ap can determine protein folding stability and DNA binding affinity. In fact, a nine-amino acid segment (seven cationic residues) of the Nhp6Ap leader is sufficient to confer full activity on HMGB2(box A), whereas substitution of HMGB2(box A) with the primary intercalating methionine of Nhp6Ap makes a detectably favorable contribution only when a smaller number of Nhp6Ap cationic leader residues is included.

The similarity of our results obtained in vitro and vivo suggests a remarkably dominant effect of leader cationic residues in HMGB folding, DNA binding, and DNA bending. The cationic leader (and thus the higher protein isoelectric point) correlates with more stable protein folding and stronger DNA binding. It has previously been shown that truncation of the basic residues of Nhp6Ap reduces binding affinity and the level of DNA bending (50). These authors estimated that each ionic contact in the major groove contributes $\sim 6^{\circ}$ of DNA bending (50). Our data are consistent with these results. Sequence-nonspecific HMGB proteins asymmetrically neutralize DNA charges at their random DNA binding sites, with a corresponding contribution to DNA bending at each site. Analogy can therefore be drawn between this natural DNA bending strategy and the ability of sequence-specific DNA binding proteins with engineered basic domains to induce directed DNA bending at specific binding sites (56-59).

ACKNOWLEDGMENT

We thank A. Vologodskii and J. Kahn for materials and advice related to cyclization kinetics assays, J. Peterson and the Mayo Clinic Proteomics Research Center for assistance with protein purification, W. and B. Owen and R. Weinshilboum for assistance with fluorescence anisotropy, and L. Sikkink and M. Ramirez-Alvarado for assistance with CD spectroscopy.

REFERENCES

- 1. Crothers, D. M. (1993) Architectural elements in nucleoprotein complexes. *Curr. Biol.* 3, 675–676.
- 2. Travers, A. A., Ner, S. S., and Churchill, M. E. A. (1994) DNA chaperones: A solution to a persistence problem. *Cell* 77, 167–160
- Thomas, J. O., and Travers, A. A. (2001) HMG1 and 2, and related 'architectural' DNA-binding proteins. *Trends Biochem. Sci.* 26, 167–174.

- 4. Allain, F. H., Yen, Y. M., Masse, J. E., Schultze, P., Dieckmann, T., Johnson, R. C., and Feigon, J. (1999) Solution structure of the HMG protein NHP6A and its interaction with DNA reveals the structural determinants for non-sequence-specific binding. EMBO J. 18, 2563–2579.
- 5. Masse, J. E., Wong, B., Yen, Y. M., Allain, F. H., Johnson, R. C., and Feigon, J. (2002) The S. cerevisiae architectural HMGB protein NHP6A complexed with DNA: DNA and protein conformational changes upon binding. J. Mol. Biol. 323, 263-284.
- 6. Paull, T. T., Carey, M., and Johnson, R. C. (1996) Yeast HMG proteins NHP6A/B potentiate promoter-specific transcriptional activation in vivo and assembly of preinitiation complexes in vitro. Genes Dev. 10, 2769-2781.
- 7. Paull, T. T., and Johnson, R. C. (1995) DNA looping by Saccharomyces cerevisiae high mobility group proteins NHP6A/ B. J. Biol. Chem. 270, 8744-8754.
- 8. Yen, Y. M., Wong, B., and Johnson, R. C. (1998) Determinants of DNA binding and bending by the Saccharomyces cerevisiae high mobility group protein NHP6A that are important for its biological activities. Role of the unique N terminus and putative intercalating methionine. J. Biol. Chem. 273, 4424-4435.
- Churchill, M. E., Jones, D. N., Glaser, T., Hefner, H., Searles, M. A., and Travers, A. A. (1995) HMG-D is an architecture-specific protein that preferentially binds to DNA containing the dinucleotide TG. EMBO J. 14, 1264–1275.
- 10. Klass, J., Murphy, F. V. t., Fouts, S., Serenil, M., Changela, A., Siple, J., and Churchill, M. E. (2003) The role of intercalating residues in chromosomal high-mobility-group protein DNA binding, bending and specificity. Nucleic Acids Res. 31, 2852-2864.
- 11. Bianchi, M. E., Beltrame, M., and Paonessa, G. (1989) Specific recognition of cruciform DNA by nuclear protein HMG1. Science 243, 1056-1059.
- 12. Paull, T. T., Haykinson, M. J., and Johnson, R. C. (1993) The nonspecific DNA-binding and -bending proteins HMG1 and HMG2 promote the assembly of complex nucleoprotein structures. Genes Dev. 7, 1521–1534.
- 13. Pil, P. M., Chow, C. S., and Lippard, S. J. (1993) High-mobilitygroup 1 protein mediates DNA bending as determined by ring closure. Proc. Natl. Acad. Sci. U.S.A. 90, 9465-9469.
- 14. Weir, H. M., Kraulis, P. J., Hill, C. S., Raine, A. R. C., Laue, E. D., and Thomas, J. O. (1993) Structure of the HMG box motif in the B-domain of HMG1. EMBO J. 12, 1311-1319.
- 15. Wolffe, A. P. (1999) Architectural regulations and Hmg1. Nat. Genet. 22, 215-217.
- 16. Kamau, E., Bauerle, K. T., and Grove, A. (2004) The Saccharomyces cerevisiae high mobility group box protein HMO1 contains two functional DNA binding domains. J. Biol. Chem. 279, 55234-55240.
- 17. Grasser, K. D., Teo, S. H., Lee, K. B., Broadhurst, R. W., Rees, C., Hardman, C. H., and Thomas, J. O. (1998) DNA-binding properties of the tandem HMG boxes of high-mobility-group protein 1 (HMG1). Eur. J. Biochem. 253, 787-795.
- 18. Zimmerman, J., and Maher, L. J., III. (2008) Transient HMGB protein interactions with B-DNA duplexes and complexes. Biochem. Biophys. Res. Commun. 371, 79-84.
- 19. He, Q., Ohndorf, U. M., and Lippard, S. J. (2000) Intercalating residues determine the mode of HMG1 domains A and B binding to cisplatin-modified DNA. Biochemistry 39, 14426-14435.
- 20. Jung, Y., and Lippard, S. J. (2003) Nature of full-length HMGB1 binding to cisplatin-modified DNA. Biochemistry 42, 2664–2671.
- 21. Ohndorf, U. M., Rould, M. A., He, Q., Pabo, C. O., and Lippard, S. J. (1999) Basis for recognition of cisplatin-modified DNA by high-mobility-group proteins. Nature 399, 708-712.
- 22. Takahara, P. M., Rosenzweig, A. C., Frederick, C. A., and Lippard, S. J. (1995) Crystal structure of double-stranded DNA containing the major adduct of the anticancer drug cisplatin. Nature 377, 649-
- 23. Churchill, M. E., Changela, A., Dow, L. K., and Krieg, A. J. (1999) Interactions of high mobility group box proteins with DNA and chromatin. *Methods Enzymol.* 304, 99–133. Swinger, K. K., and Rice, P. A. (2004) IHF and HU: Flexible
- architects of bent DNA. Curr. Opin. Struct. Biol. 14, 28-35.
- 25. Swinger, K. K., Lemberg, K. M., Zhang, Y., and Rice, P. A. (2003) Flexible DNA bending in HU-DNA cocrystal structures. EMBO J. 22, 3749-3760.
- 26. Van Noort, J., Verbrugge, S., Goosen, N., Dekker, C., and Dame, R. T. (2004) Dual architectural roles of HU: Formation of flexible hinges and rigid filaments. Proc. Natl. Acad. Sci. U.S.A. 101, 6969-6974.

- 27. Zhang, J., McCauley, M. J., Maher, L. J., III, Williams, M. C., and Israeloff, N. E. (2009) Mechanism of DNA flexibility enhancement by HMGB proteins, Nucleic Acids Res. (in press)
- 28. McCauley, M. J., Zimmerman, J., Maher, L. J., III, and Williams, M. C. (2007) HMGB binding to DNA: Single and double box motifs. J. Mol. Biol. 374, 993-1004.
- 29. McCauley, M., Hardwidge, P. R., Maher, L. J., III, and Williams, M. C. (2005) Dual binding modes for an HMG domain from human HMGB2 on DNA. Biophys. J. 89, 353-364.
- 30. Jayaraman, L., Moorthy, N. C., Murthy, K. G., Manley, J. L., Bustin, M., and Prives, C. (1998) High mobility group protein-1 (HMG-1) is a unique activator of p53. Genes Dev. 12, 462-472.
- 31. Calogero, S., Grassi, F., Aguzzi, A., Voigtlander, T., Ferrier, P., Ferrari, S., and Bianchi, M. E. (1999) The lack of chromosomal protein HMG1 does not disrupt cell growth but causes lethal hypoglycaemia in newborn mice. Nat. Genet. 22, 276-280.
- 32. Ellwood, K. B., Yen, Y. M., Johnson, R. C., and Carey, M. (2000) Mechanism for specificity by HMG-1 in enhanceosome assembly. Mol. Cell. Biol. 20, 4359-4370.
- 33. Mitsouras, K., Wong, B., Arayata, C., Johnson, R. C., and Carey, M. (2002) The DNA architectural protein HMGB1 displays two distinct modes of action that promote enhanceosome assembly. Mol. Cell. Biol. 22, 4390-4401.
- 34. LeRoy, G., Orphanides, G., Lane, W. S., and Reinberg, D. (1998) Requirement of RSF and FACT for transcription of chromatin templates in vitro. Science 282, 1900-1904.
- 35. Lu, J., Kobayashi, R., and Brill, S. J. (1996) Characterization of a high mobility group 1/2 homolog in yeast. J. Biol. Chem. 271, 33678-33685.
- 36. Travers, A. A. (2003) Priming the nucleosome: A role for HMGB proteins? EMBO Rep. 4, 131-136.
- 37. Ross, E. D., Keating, A. M., and Maher, L. J., III. (2000) DNA constraints on transcription activation in vitro. J. Mol. Biol. 297,
- 38. Ross, E. D., Hardwidge, P. R., and Maher, L. J., III. (2001) HMG proteins and DNA flexibility in transcription activation. Mol. Cell. Biol. 21, 6598-6605.
- 39. Bellomy, G., Mossing, M., and Record, M. (1988) Physical properties of DNA in vivo as probed by the length dependence of the lac operator looping process. Biochemistry 27, 3900-3906.
- 40. Law, S. M., Bellomy, G. R., Schlax, P. J., and Record, M. T., Jr. (1993) In vivo thermodynamic analysis of repression with and without looping in lac constructs. Estimates of free and local lac repressor concentrations and of physical properties of a region of supercoiled plasmid DNA in vivo. J. Mol. Biol. 230, 161-173.
- 41. Mossing, M. C., and Record, M. T., Jr. (1986) Upstream operators enhance repression of the lac promoter. Science 233, 889-892.
- 42. Kramer, H., Amouyal, M., Nordheim, A., and Müller-Hill, B. (1988) DNA supercoiling changes the spacing requirements of two lac operators for DNA loop formation with lac repressor. EMBO J. 7, 547-556.
- 43. Kramer, H., Niemoller, M., Amouyal, M., Revet, B., von Wilcken-Bergmann, B., and Müller-Hill, B. (1987) lac repressor forms loops with linear DNA carrying two suitably spaced lac operators. EMBO J. 6, 1481–1491.
- 44. Muller, J., Oehler, S., and Müller-Hill, B. (1996) Repression of lac promoter as a function of distance, phase and quality of an auxiliary lac operator. J. Mol. Biol. 257, 21-29.
- 45. Oehler, S., Amouyal, M., Kolkhof, P., von Wilcken-Bergmann, B., and Müller-Hill, B. (1994) Quality and position of the three lac operators of E. coli define efficiency of repression. EMBO J. 13, 3348–3355.
- 46. Oehler, S., Eismann, E. R., Kramer, H., and Müller-Hill, B. (1990) The three operators of the lac operon cooperate in repression. EMBO J. 9, 973-979.
- 47. Becker, N. A., Kahn, J. D., and Maher, L. J., III. (2005) Bacterial repression loops require enhanced DNA flexibility. J. Mol. Biol. *349*, 716–730.
- 48. Becker, N. A., Kahn, J. D., and Maher, L. J., III. (2007) Effects of nucleoid proteins on DNA repression loop formation in Escherichia coli. Nucleic Acids Res. 35, 3988-4000.
- 49. Becker, N. A., Kahn, J. D., and Maher, L. J., III. (2008) Eukaryotic HMGB proteins as replacements for HU in E. coli repression loop formation. Nucleic Acids Res. 36, 4009-4021.
- 50. Dragan, A. I., Read, C. M., Makeyeva, E. N., Milgotina, E. I., Churchill, M. E., Crane-Robinson, C., and Privalov, P. L. (2004) DNA binding and bending by HMG boxes: Energetic determinants of specificity. J. Mol. Biol. 343, 371-393.

- 51. Whipple, F. W. (1998) Genetic analysis of prokaryotic and eukaryotic DNA-binding proteins in *Escherichia coli*. *Nucleic Acids Res.* 26, 3700–3706.
- Datsenko, K. A., and Wanner, B. L. (2000) One-step inactivation of chromosomal genes in *Escherichia coli* K-12 using PCR products. *Proc. Natl. Acad. Sci. U.S.A.* 97, 6640–6645.
- Vologodskaia, M., and Vologodskii, A. (2002) Contribution of the intrinsic curvature to measured DNA persistence length. *J. Mol. Biol.* 317, 205–213.
- Hagerman, P. J., and Ramadevi, V. A. (1990) Application of the method of phage T4 DNA ligase-catalyzed ring-closure to the study of DNA structure. I. Computational analysis. *J. Mol. Biol.* 212, 351–362.
- 55. Stros, M. (2001) Two mutations of basic residues within the N-terminus of HMG-1 B domain with different effects on DNA supercoiling and binding to bent DNA. *Biochemistry* 40, 4769–4779.

- Strauss-Soukup, J. K., and Maher, L. J., III. (1998) Electrostatic effects in DNA bending by GCN4 mutants. *Biochemistry* 37, 1060– 1066
- Hardwidge, P. R., Kahn, J. D., and Maher, L. J., III. (2002) Dominant effect of protein charge rather than protein shape in apparent DNA bending by engineered bZIP domains. *Biochemistry* 41, 8277–8288.
- McDonald, R. J., Kahn, J. D., and Maher, L. J., III. (2006) DNA bending by bHLH charge variants. *Nucleic Acids Res.* 34, 4846– 4856.
- McDonald, R. J., Dragan, A. I., Kirk, W. R., Neff, K. L., Privalov,
 P. L., and Maher, L. J., III. (2007) DNA bending by charged peptides: Electrophoretic and spectroscopic analyses. *Biochemistry* 46, 2306–2316.

BI802269F