Altered gene expression correlates with DNA structure

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We examined the participation of triplex DNA structure in gene regulation using a poly(dG)-poly(dC) sequence as a model. We show that a poly(dG)-poly(dC) sequence, which can adopt an intramolecular dG.dG.dC triplex under superhelical strain, strongly augments gene expression when placed 5' to a promoter. The activity of this sequence exhibits a striking length dependency: dG tracts of 27–30 bp augment the expression of a reporter gene to a level comparable to that observed with the polyoma enhancer in mouse LTK- cells, whereas tracts of 35 bp and longer have virtually no effect. A supercoiled plasmid containing a dG tract of 30 bp competes in vivo for a trans-acting factor as revealed by reduction in the reporter gene transcription driven by the (dG)29 promoter of the test plasmid, while dGs of 35 bp and longer in the competition plasmid failed to compete. In purified supercoiled plasmid DNA at a superhelical density of -0.05, dG tracts of 32 bp and longer form a triplex, whereas those of 30 bp and shorter remain double-stranded under a PBS solution. These results suggest that a localized superhelical strain can exist, at least transiently, in mouse LTK- cells, and before being relaxed by topoisomerases this rapidly induces dG tracts of 35 bp and longer to adopt a triplex preventing the factor from binding. Thus, these data suggest that a poly(dG)-poly(dC) sequence can function as a negative regulator by adopting an intramolecular triple helix structure in vivo.

[Key Words: Transcriptional regulation, negative supercoiling; poly(dG)-poly(dC) sequence; dG.dG.dC triple helix; transcription factor]

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It has been assumed that transcriptional regulation is governed by proteins that interact with DNA through recognition of specific base sequences. Is DNA always a passive participant in its own utilization by providing sequences to be recognized by proteins? Because DNA is a flexible molecule and can form various structures other than the Watson–Crick B-form structure (non-B form DNA structure), DNA could utilize this potential to participate actively in biological reactions involving DNA. When introduced in the proper context, an anomalous structure, such as curved DNA, affects gene expression in Escherichia coli [Bracco et al. 1989].

Sequences residing in regulatory regions of genes are often found to be preferential targets, either in purified supercoiled plasmid DNA or chromosomal DNA in cells, for reaction with chemical probes and endonucleases such as S1, which preferentially attack single-stranded DNA [Larsen and Weintraub 1982; Kohwi-Shigematsu et al. 1983; Bode et al. 1986; Kohwi-Shigematsu and Nelson 1988]. The hyper-reactivity of these DNA regions to these probes suggests the formation of non-B form DNA structures in cells. Although the various types of sequences confer the non-B form DNA structure under superhelical strain, many of the S1-sensitive or chemical probe-reactive sequences have been identified as homopurine–homopyrimidine sequences.

For example, these homopurine–homopyrimidine sequences have been mapped in the regulatory regions of Drosophila heat shock genes [Mace et al. 1983; Siegfried et al. 1986], human c-myc gene [Boles and Hogan 1987], adenovirus major later promoter [Goding and Russell 1983], human e-globin gene [Allan et al. 1983], and chicken β-globin gene [Larsen and Weintraub 1982; Kohwi-Shigematsu et al. 1983, Nickol and Felsenfeld 1983; Kohwi 1989]. These homopurine–homopyrimidine tracts are often found to bind sequence-specific factors, for example, a GA repeat element upstream of the Drosophila heat shock gene [Gilmour et al. 1989] and another type of a homopurine–homopyrimidine tract upstream of the human c-myc gene [Davis et al. 1989]. Also, an erythrocyte–specific factor, BGP1, which binds to a poly(dG) tract in the 5'-flanking region of the chicken adult β-globin gene, has been reported [Lewis et al. 1988], the binding of which requires (dG) as a minimum recognition length [Clark et al. 1990]. These factors bind to the corresponding homopurine–homopyrimidine sequences in the linear DNA fragments where altered structures cannot be formed.

To test whether DNA that forms a non-B-form DNA structure under superhelical strain plays a role in gene regulation by changing its structure, we chose the poly(dG)-poly(dC) sequence as a model. At a physiolog...
ical pH and in the presence of Mg	extsuperscript{2+}, a poly(dG)-poly(dC) sequence in supercoiled DNA folds sharply into halves at the center, forming a poly(dG)-poly(dG)-poly(dC) triple helix (abbreviated as dG.dG.dC triplex), with both halves of the poly(dG) tract and the 3' half of the poly(dC) tract leaving the 5' half of the poly(dC) tract base unpaired (Kohwi and Kohwi-Shigematsu 1988, Kohwi 1989) (see Fig. 4, below). This structure, which is distinct from the similar structure, consists of a protonated poly(dC)-poly(dG)-poly(dC) triplex (abbreviated as dC.dG.dC triplex) and single-stranded 5' half of the dG tract that is commonly known as H-DNA, which occurs only at acidic pH without Mg	extsuperscript{2+} [Lyamichev et al. 1987, Kohwi-Shigematsu and Kohwi 1988]. Recently, we reported that under a given ionic condition and superhelicity, there is a critical length requirement of the dG tract for stabilization of the intramolecular dG.dG.dC triplex formation (Kohwi-Shigematsu and Kohwi 1991). Therefore, if there is a biological activity associated with the dG tract in vivo, this activity may exhibit a striking length dependency if the triplex formation is allowed in cells. Because the triplex formation requires superhelical strain, the present experiment addresses whether there is unconstrained superhelicity at any given time in eukaryotic cells.

Here, we report three independent observations regarding dramatic length dependency for poly(dG)-poly(dC) sequences: (1) effects on gene expression, (2) binding by a trans-acting factor in vivo, and (3) in vitro dG.dG.dC triplex formation. The results of each of these experiments identified a critical, sharp transition point in the length of the dG tracts. Taken together, these observations suggest that there is a structural transition for the poly(dG)-poly(dC) sequences that occurs in vivo and is responsible for altered gene expression.

Results

Length-dependent effects on gene expression

To examine whether poly(dG)-poly(dC) sequences have any biological activity on gene expression, various lengths of the [dG]n tracts were inserted 23 bp from the 5' end of the herpes simplex virus thymidine kinase (TK) promoter. This promoter is a 236-bp PvuII-HincII fragment containing all three elements of the TK promoter (McKnight 1982), in addition to its 5' cap site. This was cloned next to the bacterial chloramphenicol acetyltransferase (CAT) gene as shown in Figure 1A. These plasmids were transfected into mouse LTK− cells. We found that poly(dG)-poly(dC) sequences located 259 bp from the cap site augmented gene expression, and this enhancing activity depended critically on the length of the dG tracts.

The insertion of poly(dG)-poly(dC) sequences of 27 bp (data not shown), 29, and 30 bp produced a strong transcriptional augmenting activity on CAT expression comparable to the polyoma enhancer (Fig. 1B, lanes 5, 6). Surprisingly, the activity dropped dramatically for the dG tract of 33 bp (data not shown) and remained very low for increasing dG lengths of 35 and 44 bp (Fig. 1B, lanes 7, 8). The maximum transcriptional augmenting activity was seen with dG tracts of 29 and 30 bp, but a significant level of activity was also seen with the dG tract of 21 bp (Fig. 1B, lane 4). A dG length shorter than 16 bp had virtually no effect (data not shown). The transcriptional augmenting activity of the dG tracts requires the TK promoter (Fig. 1B, lane 9) and is independent of the orientation of the dG tracts (Fig. 1B, lanes 6, 15).

If a trans-acting factor binds to the dG tract, the effect of the length of the dG tract on the regulation of CAT expression could be due to an alteration of the stereo-
specific alignment between the factors binding to a specific position within a dG tract and the TK promoter. To examine this, we changed the original distance of 23 bp between the dG tracts and the TK promoter by approximately one-half a helical turn by deleting 4 bp. This deletion had no effect on the strong transcriptional augmenting activity of the 29-bp dG tract nor did it improve the very weak activity of the 35-bp dG tract (Fig 1B, lanes 11–14). Therefore, the transcriptional augmenting activity of dG tracts does not display the DNA turn dependence seen in the SV40 DNA early promoter (Takahashi et al. 1986). The length-dependent effect of poly[dG] tracts on gene expression cannot be explained by the deletion of the long dG tracts in LTK− cells. This was confirmed by sequence analysis on the plasmid DNAs that contain various lengths of dG tracts extracted from the LTK− cells and amplified in bacteria. The maximal deletion frequency was ~6% for the 44-bp dG tract and <1% for the 35-bp dG tract (data not shown).

Enhancers are generally known to augment gene expression from distance in an orientation-independent manner (Schöler and Gruss 1984 and references therein). The effect by the dG tracts was abolished when a 600-bp sequence was inserted between the dG tracts and the TK promoter (data not shown). Thus, the dG tracts act as upstream activating elements rather than enhancers, even though the dG tracts can influence transcription in an orientation-independent manner as an enhancer.

**In vivo competition and gel mobility-shift assay**

We studied whether the strong transcriptional augmenting activity of poly[dG]–poly[dC] sequences of certain lengths was the result of the binding of a trans-acting factor to the sequence. If there were such a trans-acting factor, it should be possible to titrate out the factor by adding excess amounts of a competitor plasmid that contains poly[dG]–poly[dC] sequences (Schöler and Gruss 1984). In parallel, we asked whether the structure of the target sequence influences binding by the factor. Our hypothesis is that a long, double-stranded dG tract may be preferred over a short tract for binding; however, when it exceeds a critical length where a triple helix structure is formed, the binding of the factor is inhibited. For the in vivo competition experiment, the transfection employing calcium phosphate was avoided because the intramolecular dG.dG.dC triplex formation is facilitated by divalent cations including Ca2+ (Kohwi 1989). We employed a modified DEAE–Dextran transfection protocol as described in Materials and methods. To obtain the CAT activity of the uncompeteted test plasmid to the level of at least 10% conversion of unacetylated chloramphenicol to the acetylated form, 1–2 μg of the test plasmid DNA was used. Our modified DEAE–dextran transfection procedure gave the most reproducible results with a total of 10 μg of DNA with pUC18 plasmid DNA as the carrier DNA. Under this condition, the level of CAT activity did not vary >5% among different cell cultures transfected with a given DNA sample within the same experiment.

We cotransfected [dG]29TKCAT plasmid (the test plasmid of 3.7 kb, 2 μg) and plasmid DNA containing only [dG]n, where n = 13, 18, 21, 29, 30, 35, and 44 bp (the competitor plasmids of 2.9 kb, 8 μg) as control competitor plasmids, we employed pUC18 vector plasmids alone (negative control) and pUC18, which contains only the TK promoter (positive control), because various protein factors have been reported to bind to this promoter region (for review, see Johnson and McKnight 1989). Data from a representative experiment employing 5-molar excess of competitor plasmid are shown in Figure 2A. The CAT activity of the test plasmid was not affected by cotransfecting with pUC18 plasmid (Fig. 2A, lane 1), whereas it was reproducibly reduced by 30 ± 5% when cotransfected with the TK promoter-containing plasmid (Fig. 2A, lane 2). The CAT activity of the test plasmid was also reduced by 30 ± 5% when cotransfected with the competitor plasmid containing dG tracts of 29 or 30 bp without the TK promoter (Fig. 2A, lanes 7, 8). The percent inhibition of CAT activity again correlates with the length of the dG tract in the competitor plasmid. The percent inhibition increased with increasing length of dG tracts up to 30 bp, and it decreased dramatically beyond 35 bp (Fig. 2A, lanes 4–10). The plasmid containing both TK promoter and [dG]30 competed more efficiently than the plasmid containing either sequence alone (Fig. 2A, lane 3). The PyTKCAT plasmid containing the polymer enhancer (the PvuII-4 fragment) also competed to a moderate extent (Fig. 2A, lane 11). By increasing the ratio of competitor plasmid DNAs to the test plasmid, a maximum of 50–60% inhibition was seen with the plasmid containing [dG]29–30 tracts. However, when higher ratios were used, the level of inhibition was subject to more variation among independent experiments. Nevertheless, it should be emphasized that all of our experiments showed efficient competition with dG tracts up to 30 bp and abrupt loss of competing activity for the dG tracts of 35 bp and longer. This transition point in length between 30 and 35 bp that distinguishes the competition activity is similar to that showing a dramatic difference in transcriptional enhancement. These results also support our hypothesis that a trans-acting factor that binds to the poly[dG]–poly[dC] tract exists in mouse LTK− cells and that the binding is affected by the DNA structure of the tract.

A gel mobility-shift assay with whole-cell extract (Eul et al. 1989) of mouse LTK− cells also suggested that there is a factor that binds specifically to poly[dG]–poly[dC] tracts (Fig. 2B). After mixing the whole-cell extract obtained from the mouse L cells and an end-labeled, linear DNA fragment containing a dG tract of 30 bp, we detected a band of slower migration. The intensity of this band decreased when the extract included either 10- or 30-molar excess of unlabeled linear pUC18 containing dG tracts of 21, 30, and 35 bp (Fig 2B, lanes 4–9); however, addition of pUC18 alone (Fig. 2B, lanes 2, 3) or the presence of the TK promoter (Fig. 2B, lanes 10, 11) had no effect. When 30-fold molar excess of competition plasmids with dG of 21, 30, or 35 bp was used, the band nearly vanished. Because we used linear rather than su-
percoiled DNA, it is not surprising that no transition point in length that distinguishes the binding potential was observed in this experiment. Supercoiled DNA could not be used in this in vitro band-shift assay with whole-cell extract because of the topoisomerase activity. The enzyme activity is difficult to eliminate by a simple fractionation of the cell extract by chromatography. It apparently requires an extensive purification of the dG tract-binding protein before successfully removing the topoisomerase activity.

Length requirement for dG.dG.dC triplex formation

We have reported previously that either an intramolecular dG.dG.dC or dC+dG.dC triplex formation depends on the length of poly[dG]=poly[dC] sequence (Kohwi-Shigematsu and Kohwi 1991). The condition that we employed previously was 50 mM Na+ concentration at a bacterial superhelical density of −0.05. To examine whether the length dependency that we observed for the effect of the sequence on gene expression and in vivo competition reflects the triplex formation beyond a certain length of the sequence, we performed the DNA structural analysis of various lengths of poly[dG]=poly[dC] sequences in PBS. Plasmid DNAs containing dG stretches of 21, 25, 27, 28, 29, 30, 32, 35, and 44 bp were constructed, and their triplex-forming potentials were examined under two slightly different ionic conditions: (1) 100% PBS [137 mM NaCl, 2.68 mM KCl, 8 mM Na2HPO4, 1.47 mM KH2PO4, pH 7.4] with 2 mM Mg2+, and (2) 80% of this solution (with 20% H2O). The structure was detected by reacting plasmid DNAs with chloroacetalddehyde [CAA], an unpaired DNA base-specific chemical probe, followed by the chemical cleavage method (Kohwi and Kohwi-Shigematsu 1988 and references therein). When a poly[dG]=poly[dC] tract forms a dC.dG.dC triplex, cytosines over the 5' half of the dC tract that are not involved in the triplex formation are unpaired and are reactive with CAA. Thus, the dC.dG.dC triplex formation can be demonstrated by the cleavages of DNA at these cytosines, as shown by the appearance of new bands within sequencing ladders. Our results showed that in 100% PBS, a distinct transition point in lengths exists between 30 and 32 bp that separates double-stranded (up to 30 bp, Fig. 3B, lanes 1–6) versus triple helix structure (32 bp and longer, Fig. 3B, lanes 7–9) under the bacterial superhelical density of −0.05. This result is well correlated with the length-dependent CAT activity for dG tracts (Fig. 1B). In 80% PBS, a slightly lower ionic condition, the transition point in lengths occurred between 27 and 28 bp (Fig. 3B, lanes 7–9) under the bacterial superhelical density of −0.05. This result is well correlated with the length-dependent CAT activity for dG tracts (Fig. 1B). In 80% PBS, a slightly lower ionic condition, the transition point in lengths occurred between 27 and 28 bp (Fig. 3A, lanes 3, 4); that is, tracts of 28 bp and longer form a dC.dG.dC triplex, whereas those of 27 bp and shorter form a double strand. We conclude that there is a sharp transition point in lengths of dG tracts between the triplex and B-form DNA structure under physiological ionic strength and at bacterial superhelical density. An increase of only two dG residues is sufficient to induce the triplex formation. The exact length at which the transition occurs is influenced by a slight difference in ionic strength.

Discussion

To examine the structure-function relationship of sequences that have the potential to adopt non-B-form DNA structures, we employed poly[dG]=poly[dC] sequences of various lengths as a model system. The formation of the intramolecular dG.dG.dC triplex is depen-
Gene regulation by the dG.dG.dC triplex

Our results strongly suggest that non-B-form DNA structures may play an active role in gene expression. On the basis of our data, a possible mechanism by which this sequence may affect gene expression is shown in Figure 4. In this model, an intramolecular triplex formed by longer dG tracts in vivo prevents a trans-acting factor from binding and abolishing transcriptional enhancement. Altered gene expression could also be caused by the triplex formation disrupting local chromatin structure, for example, by changing nucleosome positioning. There are sequence-specific factors reported that bind to homopurine–homopyrimidine sequences, including dG tracts, located within regulatory regions of various genes (Lewis et al. 1988; Davis et al. 1989; Gilmour et al. 1989). Under superhelical strain, these types of sequences form intramolecular triple helices (for review, see Wells et al. 1988) and other triplex-related DNA structures, including an intramolecular double triplex (Kohwi-Shigematsu and Kohwi 1991). Therefore, the results obtained with the model poly(dG)–poly(dC) sequence could be applied to further understand the role of homopurine–homopyrimidine sequences in gene regulation.

An intramolecular dG.dG.dC triplex can form only under superhelical stress; therefore, one must consider whether such supercoiling occurs in biological systems. In E. coli and yeast systems, it has been shown that active transcription of a gene induces local negative supercoiling 5' and positive supercoiling 3' of the gene (Liu and Wang 1987; Giaever and Wang 1988; Wu et al. 1988), as was shown in topoisomerase-defective mutant cells. In eukaryotes, the potential also exists to generate superhelical stress by the local removal of histone octamers. Recent in vitro experiments with the fibroin gene have shown that negative supercoiling plays an important role in eukaryotic transcription (Hirose and Suzuki 1988) and factor binding (Mizutani et al. 1991). Also, in vitro transcription of a topologically unconstrained template has been shown to transiently induce a B–Z transition in a region of alternating purines and pyrimidines as a result of transcription-induced negative supercoiling (Dröge and Nordheim 1991).

In our assay system we propose that sufficient local supercoiling is generated 5' of the CAT gene to rapidly induce long dG tracts (33 bp and longer) to fold into a triple helix before topoisomerases relax the DNA. The critical length of poly(dG) tracts that separates the biological activity of this sequence may vary depending on the chemical and superhelical environments at specific sites in chromosomal DNA. Our results show that a slight difference in the ionic concentration leads to a reduction in the length of the poly(dG)–poly(dC) sequence that can adopt the intramolecular triplex by 4 bp. The superhelical environment at specific points upstream of an active gene may vary at different times during the transcription process. Therefore, it may be possible for relatively short tracts of homopurine–homopyrimidine sequences to fine-tune the level of transcription by making a rapid transition between B- and non-B-form DNA structures.

To prove that the dG.dG.dC triplex structure plays a role in gene regulation, it is important to detect the structure formation inside cells. Recently, by direct chemical probing, we demonstrated that this structure forms in E. coli cells transfected with the dG tract-containing plasmid DNAs in a dG-length and superhelicidependent fashion (Y. Kohwi et al. 1991). Unfortunately, in situ probing of the DNA structure in mouse LTK- cells did not succeed due to the low amount of the transfected DNAs in these cells. The experiment was not successful even after attempting to amplify the signal for the DNA sequencing ladder by electroblotting the poly-
Figure 4. A model for negative regulation by poly(dG)-poly(dC) sequence. A model was postulated that the trans-acting factor (solid region) fails to bind to poly(dG)-poly(dC) sequence when the DNA folds into halves to form an intramolecular dG.dG.dC triplex. This model is consistent with the results that a 2- to 5-bp difference in length of poly(dG)-poly(dC) sequence confers a dramatic variation in biological activities.

acrylamide gel onto a nylon membrane, followed by hybridization with RNA probes. Amplification of DNA with the primer-extension technique is unsuitable for our particular system, because polymerases terminate at the dG tracts (Baran et al. 1991). Therefore, we are currently studying the structure formation employing a plasmid DNA that can replicate to a high copy number in mouse LTK- cells.

On the basis of our results correlating the length dependency for the poly(dG)-poly(dC) sequence activity in vivo and the dG.dG.dC formation in vitro, we speculate that non-B-form DNA-forming sequences in the regulatory regions participate actively in gene regulation, either positively or negatively, by making a structural transition in response to the local negative supercoiling.

Materials and methods

Chemicals

CAA was purchased from Fluka and doubly distilled (boiling point, 78–80°C). The purified CAA was aliquoted in small fractions and stored at –20°C until used. Hydrazine, formic acid, and piperidine were purchased from Sigma.

Plasmid DNA

The expression vector (pTKCAT) in pUC13 used in the transfection assay contains a 236-bp PvuII [5’ end]–HincII [3’ end] fragment of the herpes simplex virus TK promoter placed in the PstI and HindIII sites in the polylinker region. One of the two PstI sites (the 3’ site) shown in Figure 1A is the internal site in the TK promoter located 22 bp upstream of the HincII site. The pTKCAT contains the CAT gene and an AATAAA sequence for the polyadenylation signal (Linney and Donerly 1983). Varying lengths of poly(dG) tract were inserted into the Smal–ScaI sites in the expression vector by the G-C tailing method employing terminal transferase to generate [dG]n TKCAT. n is the number of dG residues (for a detailed protocol, see Kohwi-Shigematsu and Kohwi 1991). The PyTKCAT construct contains a 191-bp PvuII-4 fragment [PyFl10] of the polyoma enhancer sequence placed 5’ of the TK promoter and the CAT gene (Linney and Donerly 1983). This PvuII-4 fragment has a tandem duplication of 54 bp having a single-point mutation in each repeat (Fujimura et al. 1981). The (dG)n (–4 bp)TKCAT plasmid was constructed by deleting 4 bp between dG tracts and the TK promoter by first restricting the (dG)n TKCAT with XbaI and HincII and then filling in the ends with Klenow large fragment and ligating.

DNA transfection and the CAT assay

A total of 10 µg of DNA [8 µg of pUC18 vector and 2 µg of test DNA (described in the legend to Fig. 1B)] was transfected as follows: The DNA was resuspended in 200 µl of buffer A [20 mM HEPES (pH 7.4–7.6) and 140 mM NaCl]. This was mixed with 200 µl of buffer A containing 20% dimethylsulfoxide (DMSO) (Sigma), 1 mg/ml of DEAE–dextran from Pharmacia (stock solution: 40% DMSO, 2 mg/ml of DEAE–dextran in buffer A). Because there were no Mg2+ ions present in the solution, the plasmid DNAs had no dG.dG.dC triplex formed before they were introduced into the cells. The resulting DNA mixture was added to cells in a 10-cm culture dish and incubated for 30 min at room temperature. The DNA mixture was aspirated and washed once with buffer A and replaced with complete media, and the dish was placed in a CO2 incubator with 5% CO2. This
is a simple, one-step transfection procedure that gives excellent reproducibility and transfection efficiency for mouse LTK- cells. The high level of reproducibility was confirmed by cotransfecting with the human β-galactosidase gene driven by the human β-actin promoter and measuring the β-galactosidase activities of the cell extracts (Oshima et al. 1990). CAT assays were performed after 2 days, and the conversion was quantitated as described previously (Gorman et al. 1982).

In vivo competition

A total of 10 μg of DNA containing 2 μg of test plasmid (pGm TKCAT) and 8 μg of competitor plasmids without the CAT gene [pUC18 plasmid, pUC13 containing the TK promoter, the TK promoter and (dG)_{13}, (dG)_n, where n = 13, 18, 21, 30, 35, and 44], and the 200-bp PvuII fragment of the polyoma enhancer were transfected at 1 : 5 molar ratio, as described in the legend to Fig. 2.

Gel mobility-shift assay

The gel mobility-shift assay with whole-cell extract of LTK- cells was performed according to Eul et al. (1989). A radiolabeled DNA fragment containing the dG_{30} [0.1 μg] was mixed with poly[d(I-C)] oligomer (1 μg), mouse LTK- cell extracts (5 μl containing 1–5 μg of protein), and HEPES buffer [final concentration, 10 mM HEPES [pH 7.9], 1 mM DTT, 50 mM KCl, 2.5 mM MgCl₂, 10% glycerol], either with HindIII-digested competitor DNA containing various lengths of dG tracts or with no competitor in a total volume of 20 μl, as described in Figure 2B. The mixture was incubated at 30°C for 20 min. The DNA–protein mixtures were then loaded onto a 4% native polyacrylamide gel and run at 4°C. The gel was dried and exposed to an X-ray film.

Mapping of CAA-modified sites

CAA-modified sites were determined as described in Kohwi and Kohwi-Shigematsu (1988). Supercoiled plasmid DNA containing varying lengths of poly(dG) tracts was reacted with 2 μl of CAA in 100-μl reaction volume of either 100% PBS or 80% PBS at 37°C for 1 hr. CAA-modified DNAs were made into 0.3 M [Na⁺] and precipitated successively with 3 volumes of ethanol. Two micrograms of DNA was digested with HindIII, and the 3' end was labeled with Klenow large fragment of DNA polymerase I. The end-labeled DNA was cleaved further at a distal site with the second restriction enzyme, BglII. The DNA fragment containing the poly[dG]–poly[dC] sequence was isolated from a native polyacrylamide gel, and the DNA was eluted from the gel in an elution buffer [0.5 M [Na₅]OAc, 1 mM EDTA, 0.1% SDS] at 42°C overnight. The DNA isolated from a large segment of acrylamide gel was cleaned further by Geneclean II (BIO-101) and was then subjected to hydrazine or formic acid reaction followed by piperidine reaction as described in Maxam and Gilbert (1977). The chemically cleaved DNA was heat denatured and loaded onto an 8% urea-denaturing polyacrylamide gel.

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