MOLECULAR UNDERSTANDING OF MUTAGENICITY USING POTENTIAL ENERGY METHODS

Progress Report

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PROGRESS REPORT

A. Project Overview

The objective of our work, for many years, has been to elucidate on a molecular level, at atomic resolution, the structures of DNAs modified by highly mutagenic aromatic amines and hydrocarbons. We have given particular attention to the carcinogens 2-aminofluorene (AF) (Ia, figure 1) and its N-acetyl derivative, 2-acetylaminofluorene (AAF) (Ib, figure 1). Activated forms of both compounds bind to DNA, and the most prominent adducts are those that arise from substitution at position 8 of guanine (IIa and b, Figure 1). The mutagenic specificity of these compounds has been explored by a number of groups. As controls, we have also studied their less mutagenic chemically related analogs, and unmodified DNAs. The ultimate purpose of this undertaking is to obtain an understanding of the relationship between DNA structure and mutagenicity.

The underlying hypothesis is that DNA replicates with reduced fidelity when its normal right-handed B-structure is altered, and one result is a higher mutation rate. This change in structure may occur normally at a low incidence, for example by the formation of hairpin loops in appropriate sequences, but it may be enhanced greatly after covalent modification by a mutagenic substance. Other mechanisms may also lead to mutation after covalent modification, for example the operation of induced error-prone repair pathways. Ultimately it is DNA structure, mediated by replication and repair enzymes, that determines whether a mutation will take place.

The methods that we use to elucidate structures are computational, but we keep in close contact with experimental developments, and have been able to incorporate the first data from NMR studies in our calculations. The reason why computational approaches to structure generation are so important in this area is that x-ray and spectroscopic studies have not succeeded in producing atomic resolution views of mutagen and carcinogen-oligonucleotide adducts. The NMR method cannot alone yield molecular views, though it has recently done so in combination with our computations. The specific methods that we employ are minimized potential energy calculations using the torsion angle space molecular mechanics program DUPLEX (developed and written by Dr. Brian E. Hingerty) to yield static views. Molecular dynamics simulations, with full solvent and salt, of the important static structures are carried out with the program AMBER; this yields mobile views in a medium that mimics the natural aqueous environment of the cell as well as can be done with current available computing resources.

In order to obtain the most reliable structures possible, we are continuously working to advance the state-of-the-art on the computational front. Many of our efforts involve new methods for dealing with the multiple-minima problem; this impediment prevents the unambiguous identification of the global minimum energy conformation, or even of all important local energy minima. DUPLEX was developed to reduce the number of conformational variables that must be minimized at the same time in comparison to the larger number (3n - 6, where n = the number of atoms) used in cartesian space.
molecular mechanics. We vary only the torsion angles in our program, and hold bond lengths, bond angles, and out-of-plane-movements of aromatic moieties fixed. The number of flexible variables in a deoxynucleoside monophosphate (building block of a DNA helix) is thus reduced to just 12, including the termini and sugar puckers (Figure 1). We feel that very little is sacrificed by this strategy in determining the structure of modified DNA, as torsion angles changes are very large, in comparison to those of the other variables, while great advantage is derived in optimization capability.

We have also been developing new strategies for searching conformation space and building DNA duplexes from favored subunit structures. We are thus able to compute structures de novo by conformational searches. Included among these techniques is a novel approach for locating and ranking in order of energy structures with unusual hydrogen bonding patterns or denaturation at specified sites. Neither model building, with its attendant bias by the modelers aesthetics, nor experimental input is required in our approach. If, however, high resolution NMR data is available in the form of interproton distances, then these distances can be incorporated as constraints in first stage minimizations, to aid us in locating a structure that agrees with experiment. These constraints are released in the terminal minimizations, to produce unconstrained energy-minimized structures that are consistent with the NMR data.

In addition, we are collaborating with a colleague, Dr. Tamar Schlick (Chemistry Department and Courant Institute of Mathematics, New York University), with the goal of interfacing her powerful new algorithms for potential energy minimization and molecular dynamics simulation with DUPLEX. Details of this methodological approach are given in the publication by Schlick, et al. (1990).

B. Variation of the Structure of AF- and AAF-modified DNA with Sequence. (Theoretical Studies With No Experimental Input).

1. Studies on Unmodified DNA:

In a pilot project, we implemented a build up technique for polynucleotides akin to one devised for prediction of polypeptide structure from the amino acid building blocks by Scheraga and coworkers (for a review of the polypeptide work, see Gibson and Scheraga, 1988). This technique could not be applied to polynucleotides until the advent of supercomputers because the deoxynucleoside monophosphate building blocks of DNA (Figure 1) have many more torsional angles that must be optimized simultaneously than do amino acids. Briefly, we made large conformational searches for the "dimer" deoxynucleoside building blocks d(CpG) and d(GpC), and then combined these minima to produce energy minimized structures for the single strand trimer: d(CpGpC). These structures were then employed to search for minimum energy conformations of the duplex d(CpGpC).d(GpCpG). At each stage, large numbers of trials were necessary because of the combinatorial nature of the searches. It is of interest that the two lowest energy forms of the trimer duplex were B and Z DNA helices, which agreed with the experimental preferences of the alternating G-C sequence.
These two computed duplex trimers, built to the energy-minimized dodecamer level, yielded full length B and Z helixes. Thus, these structures were obtained with no experimental input other than the force field parameters. In addition, many novel double-stranded structures of higher energy were located in the duplex trimers. As our knowledge of genome structure and function grows, some of them may prove to play a role in particular circumstances. A full account of this work has been published (Hingerty, et al., 1989).

2. Studies on Modified DNA:

a. AF- and AAF-modified single-stranded DNA; We have also used our buildup techniques to study AF- and AAF-modified oligomers, following a strategy similar to that employed for unmodified DNA. The problem is much more difficult here due to the added flexibility at the amine-base linkage (Figure 1). Yet intensive conformational studies, involving thousands of trials, have been completed. An abbreviated form of this study has been presented at the 1992 meeting of the American Association for Cancer Research: Miao, Y.-S., Hingerty, B.E., Broyde, S. and Shapiro, R., "Mutagenesis by 2-Aminofluorene (AF) and 2-Acetylaminofluorene (AAF): Implications from DNA Single-strand Conformations", Proc. Amer. Assn. Cancer Res. 33, 142 (1992). A full-length manuscript is being prepared for publication.

In this work we studied the effects of AF- and AAF-modification at C-8 of guanine in deoxydinucleoside monophosphates (dimers) of all possible sequences. Further, modified alternating CGC trimers were built up from the CG and GC dimers, and the pentamers GC^AFCG and GC^AAFCG were constructed from the trimers by embedding the trimers in B-DNA. The number of trials run was the following: For dimers, 4,897 trials for all cases except for the AF- and AAF-modified CG and GC sequences. These cases were selected for a pilot test with 31,104 additional trials run for each. In general, the larger search did not prove worthwhile. The new minima found to 5.5 kcal/mol by the extended search in no case included a global minimum, and (with one exception) contributed in total less than 1.5% to the conformation population.

The exception came up in the case of CpG^AAF, in which the long search produced the second most stable minimum, 0.76 kcal/mol above the global one, contributing 6% of the total mix. It was of the predominant syn-guanine, AAF-base stacking type, but contained syn-cytosine as well. A number of structures of higher energy with syn-cytosine were also located by the longer searches, but missed by the shorter ones. This occurred because the longer searches included a syn starting conformation for cytosine but the shorter ones did not. It may be worthwhile to include syn pyrimidines in future searches, but apart from this, it may be concluded that the expansion of trials in the long search did not produce results worth the expenditure of computer time.

For trimers, 59,568 trials were run for AF-modification, and 87,572 for AAF-modification. Full details of the methods are provided
in the Ph.D. thesis of Y.-S. Miao, New York University, 1992, and are to be published.

Conformation of Dimers: \( d-\text{cAF}_{pN} \) AND \( d-\text{GAAF}_{pN} \) (\( N = A, G, C, T \)): The global minima and most of the low energy structures for all eight of the 5'-modified dimers studied fall into a single conformational class. Guanine is syn, and stacks at an angle with its 3'-neighboring base, while AF or AAF has close contacts with the bottom or side of the neighboring sugar. The plane of the guanine ring is close to perpendicular with that of the amide bond, and the fluorene ring plane is also at a sharp angle with that of the amide bond. This circumstance creates a wedge-shaped binding pocket. The 3' nucleotide residue binds within the concave surface of this pocket, and the amine H-H or acetyl protrudes from the less-hindered convex side.

Two subclasses can be defined, depending on whether the neighboring base is a pyrimidine or purine. In the former case, the pyrimidine forms a triangle with the sides of the pocket, and stacks with both guanine and, to a lesser extent, fluorene. This is illustrated for the global minimum of \( d\text{GAF}_{pC} \) in Figure 2. If the neighbor is a purine, it interacts only with guanine (see the global minimum of \( d\text{GAAF}_{pA} \), Figure 3.

Conformation of Dimers: \( d-\text{NpGAAF} \) (\( N = A, G, C, T \)): A single class of conformation predominates in this series, representing the global minima and most of the low-energy conformers of all four dimers. The fluorene ring participates in a near parallel-stack with the neighboring base, and guanine contacts the side (and top in some cases) of the 5'-sugar. The backbone is right handed, with, however, uncommon angles for the C-O bonds \( \phi' \) and \( \phi \) (\( \beta \) and \( \gamma \)).

The modified guanine is syn in these conformers, but some variation can be seen in the glycosyl-bond orientation of the neighbor, with both syn and anti-conformers prominent. The global minimum for \( \text{CpGAAF} \) (Figure 4) has anti-cytosine, which stacks with the middle and distal fluorene rings (the stack is the same for the anti conformers of A and G). The global minima for \( \text{ApGAAF} \) (Figure 5) and \( \text{GpGAAF} \) have the unmodified purine syn, and it stacks with the middle and proximal fluorene rings (the stack is the same for the syn conformer of C). \( \text{TpGAAF} \) is an exception to this pattern. The global minimum T-anti conformer (no syn-T was observed) stacks with the proximal fluorene ring.

Conformation of Dimers: \( d\text{NpGAF} \) (\( N = A, G, C, T \)): The dimer class in which the 3'-residue is modified by aminofluorene shows the most diverse behavior. It is the only group in which the identity of the neighboring base has a profound effect on the nature of the most stable structure(s). Two major subgroups can be defined: one has near parallel base-fluorene stacking, with guanine in a low syn (-20°) glycosyl orientation. the other has guanine anti, with base-base stacking that varies remarkably from one conformer to another.

The global minimum for \( d\text{TpGAF} \) (Figure 6) illustrates the base-fluorene stacking subgroup. The backbone is left-handed (Z-type), and guanine stacks loosely on the phosphate. This kind of structure occurs for neighboring C, A, and G as well, but occurs 0.6-1.0 kcal/mol above the minimum. The right handed base-fluorene stacked
conformer that dominated the NpG<sub>AAF</sub> series can be found here as well, but falls 2.2-3.7 kcal/mol above the global minimum, in the four dimers.

The guanine-anti, base-base stacked subgroup is diverse on its own, with the global minima for dCpG<sub>AF</sub>, dApG<sub>AF</sub> and dGpG<sub>AF</sub> showing different features. The minimum for dCpG<sub>AF</sub> (Figure 7) has a right-handed backbone, but with gauche α (ε) and trans ψ (γ) values. The base-base stack has only partial overlap, and is at an angle of about 25°, while the fluorene stacks on the top of the 5'-sugar and on phosphate.

The dApG<sub>AF</sub> co-global minimum (Figure 8) has a B-DNA backbone, but with ω (α) at 215° (trans), and the glycosyl angle of both bases near 165°. The fluorene ring stacks on the backside of the 5'-sugar and on phosphate. The bases have a near-parallel stack. However each uses the opposite face to that used in B-DNA for the stack. In terms of their hydrogen-bonding sites, they have essentially changed places. Further, the glycosyl bond of one purine is rotated 60° with respect to the other so that the amino group of A is over N-1 of G. If this should occur in single- or double-stranded DNA, it could possibly lead to a mutagenic two-base inversion, or a transition.

The dGpG<sub>AF</sub> global minimum is illustrated in Figure 9. It displays a right-handed backbone, but with ω (α) at 215° (trans), and the glycosyl angle of both bases near 165°. The fluorene rings stacks on the backside of the 5'-sugar and on phosphate. The orientation of the bases with respect to one-another is novel, however, with the carbonyl group of one G over the amine of the other (and vice-versa) in reverse Watson-Crick alignment. If this structure were feasible in larger fragments of DNA, its potential for mutagenic miscoding could possibly be significant.

Summary of Dimer Studies: Comparison with Experiment:

(1) Fluorene-Base Stacking vs. Base-Base Stacking: In earlier literature, these were often considered as two mutually exclusive and competing options. In fact, in the low energy minima, both guanine and fluorene generally find stacking partners; one selects the neighboring base, the other the neighboring sugar or phosphate. Fluorene-base stacking is commonly near parallel, while base-base stacking will sometimes be at an angle greater than 20°. In such cases, greater stabilization may result from optimizing the fluorene-sugar contacts than by achieving a good base-base interaction.

(2) AAF Conformations vs. AF Conformations: The principal effect of the acetyl group appears to be the destabilization of anti conformations of guanine, and to a lesser extent, conformations on the borderline of syn and anti. In the 5'-substituted dimers, the most stable AF-modified conformations involved syn guanine, and quite similar ones were favored in the AAF series. In the 3'-substituted dimers, the most stable conformations involve anti or low syn guanine. Such structures are disfavored in the AAF series, and the most stable conformations have syn guanine. In the AF series, syn guanine structures are of higher energy.
(3) 5'-Substitution vs. 3'-Substitution: One conformational type predominates in the low energy structures for the 5'-substituted adducts of both AAF and AF: right handed, with syn-guanine, imperfect base-base stacking, and fluorene to 3'-sugar contacts. The AAF-substituted 3'-adducts primarily displayed good base-fluorene stacking, with syn-guanine in contact with the 5'-sugar. The AF-substituted 3'-adducts were more diverse, showing a variety of right-handed low energy structures with guanine anti, good to poor base-base stacking and fluorene to 5'-sugar contacts. Substituted 3'-adducts primarily displayed good base-fluorene stacking, with syn guanine in contact with the 5'-sugar. The AF-substituted 3'-adducts were more diverse, showing a variety of right-handed low energy structures with guanine anti, good-to-poor base-base stacking and fluorene to 5'-sugar and phosphate contacts. The NpG$_{AF}$ series also featured a left-handed, low energy conformation with good base-fluorene stacking and guanine-phosphate contacts.

(4) Effect of Neighboring Base. This was important only in the 3'-AF substituted series, where TpG$_{AF}$ had a left-handed base-fluorene stacked form as global minimum, and the remainder displayed differing right-handed forms with base-base stacking.

(5) Comparison to Experiment. dApG$_{AF}$ and dApG$_{AAF}$ were examined by NMR and CD (Santella, et al. 1980; Leng, et al., 1980). The authors concluded that fluorene-base stacking was more significant in the former than the latter. A detailed NMR study of dCpG$_{AF}$ has been carried out by Evans and Levine (1988), who found strong cytosine-fluorine stacking, with guanine syn, and the guanine C(8) to amine N bond ($\alpha'$) near 90°. Our results agree with these experimental ones, though individual conformers that match the specific sugar pucker and C-5' to C-4' ($\gamma$) torsional angles suggested by Evans and Levine have not yet emerged in our study.

Trimer Conformations: dCpG$_{AF}$pC and dCpG$_{AAF}$pC: In the case of the AF-modified trimers, the global minimum (Figure 10) and the majority of low energy trimers fall into a single class. This has a backbone considerably rearranged from that present in a normal B-DNA stack. Near-parallel base-base stacking takes place between cytosines (1) and (3), with the amino of each over the carbonyl of the other. Syn guanine has looped out and is in contact with the edges of the two cytosines, and the fluorene ring stacks on the back of the 3'-end sugar and on its phosphate. In this compact structure, the terminal 5'-OH and 3'-OH have come close to one another, with the O to O distance measured at 4.82 Å. (The normal distance from a 5'-O to the 3'-O of the same sugar in B-DNA is about 4.33 Å). A structure of this type could possibly form a loop in single stranded DNA.

Several other quite different conformations could also be found at relatively low energy. One of them, which involves an anti ($\lambda = 170^\circ$) guanine to cytosine (3) to cytosine (1) stack, with fluorene to sugar (1) contacts, is illustrated in Figure 11. The guanine to cytosine (3) stack shows normal alignment, but the cytosine to cytosine stack is reversed, as described for the global minimum.
The AAF-modified trimer showed two co-global minima, both of which had the same general shape and stacking interactions as the major AF-modified class described above. The two co-minima differed from one another in several of their backbone angles, however. One of them is pictured in Figure 12. Most of the other low energy AAF-modified conformers were in that general class. The OH-ends of these structures again were about 4.8 Å from one another, with the potential for forming a loop. One alternative conformer type (energy 1.08 kcal/mol above the global minimum) is pictured in Figure 13. It exhibits a syn guanine to cytosine (1) to cytosine (3) stack, with fluorene in contact with the underside of sugar (3). Normal B or Z single-stranded structures, with conventional C-G-C stacking, were not found at energies up to 3.5 kcal/mol in either the AF or AAF-modified structures.

AF- and AAF- Modified Pentamers. To explore whether the global minima for CpGpApc and CpGpAfpCpG could be incorporated into larger DNA oligomers, a limited number of trials was run in which dGMP was added to the 3' and 5' ends of the AF trimer in Figure 10, and the resulting structure was minimized. The same procedure was followed with the AAF trimer in Figure 12. The most stable pentamer conformers were produced when the added monomer fragments were started in B-DNA geometry. They are illustrated for dGpCpGpAfpCpG in Figure 14 and for dGpCpGpAfpCpG in Figure 15. The above AF-modified conformer was 3.5 kcal/mol more stable than the conformer with B-DNA geometry (GAF anti), while the above AAF-modified pentamer was 13.7 kcal/mol more stable than the corresponding B-DNA conformer(GAfpCpG syn). An AAF-modified pentamer in the geometry of Figure 14 was also feasible. Its energy was slightly higher (1.6 kcal/mol) than the one in Figure 15.

The AF-modified pentamer is noteworthy in that it contains a G1-G5-C2-C4 stack. The C4 end is "capped" by the binding pocket formed by modified G3 and AF. G3 contacts the edge of C4 (and to a lesser extent, C2), while the distal ring of AF stacks onto the backside of the C4-sugar. This situation resembles that of the parent trimer. The 5'-0 of G1 is now 9.36 Å from the 3'-0 of G5. (The distance from 0-5' of one residue in B-DNA to 0-3' of the 3'-neighbor is about 10.5 Å) and they are oriented in the same direction. The pentamer thus spans the approximate distance used by a dimer in a DNA chain. Further buildup should be possible, but has not yet been attempted.

The AAF-modified pentamer (Figure 15) contains G1-C2-C4 parallel stack. G1 and C2 overlap in a normal Watson-Crick manner, while the C2-C4 stack has reverse Watson-Crick geometry. As in the AF case above, this stack is capped by the G3-AAF pocket. The proximal AAF ring has a partial overlap (the angles between the planes is 22°) with C4, while the distal ring contacts the sugar of C4. G5 is out of the stack, in contact with those areas of G1 and C2 that would line the major groove in B-DNA.

Thus, these studies show that modification by AF or AAF perturbs the structure of short single stranded DNA oligomers. Among the novelties observed are sequence inversion, reverse Watson-Crick stacking arrays, and the formation of compact loops with out-of
sequence stacking. If such features should persist in larger fragments of single-stranded DNA, they could provide a molecular basis for substitution, base-inversion and deletion mutants.

In an intensive continuation of this work that is now in progress, we are combining the single stranded trimers dCpGApC and dCpGApC with dGpCpG to generate energy minimized duplexes. The number of trials needed approaches the 100,000 range for these energy minimized duplex trimer searches.

b. AAF-Modified Double-stranded DNA: In this study, minimized potential energy calculations were employed to locate and evaluate energetically a number of different models for DNA modified at carbon-8 of guanine by AAF, using very limited searches. Three different duplex nonamer sequences were investigated. In addition to syn guanine models which have some denaturation and carcinogen-base stacking in certain cases, and a Z-DNA model, we have found two new types of structures in which guanine remains syn and the AAF is placed in the minor groove of a B-DNA helix. One type features Hoogsteen base pairing between the modified guanine and a protonated cytosine, with a sharply bent helix. The other (here termed the "wedge" model because the aromatic amine is wedged into the minor groove) can maintain a single hydrogen bond between O6 of the modified guanine and N3 of protonated cytosine, with much less deformation of the helix, and close Van der Waals contacts between the AAF and the walls of the minor groove. Both types of structure (as well as the related forms produced by deprotonation of cytosine) are energetically important in all three sequences examined. The wedge-type minor groove model, which is most favored except in alternating G-C sequences, had been previously observed in combined NMR and computational characterization of an aminofluorene (AF) modified guanine opposite adenine in a DNA duplex undecamer (described below). This work has been published (Shapiro, et al., 1989).

C. Studies on the Structures of Modified Double-Stranded Oligonucleotides, Employing NMR Data.

1. An AF modified duplex 11-mer with adenine opposite the lesion site: molecular view of a mismatch mutation: A combination of NMR data from the Patel-Grunberger laboratories at Columbia University and our calculations have produced the first combined experimental and theoretical structure of a DNA-aromatic amine adduct. Furthermore, an entirely new type of structure for this adduct was found: The aromatic amine is wedged into the minor groove of a DNA duplex in a manner that minimizes the exposure to solvent of the hydrophobic moiety. Only the edges of the aromatic rings are exposed. This orientation is achieved by positioning the modified guanine in a syn conformation, rotated approximately 180° in the glycosyl linkage from its normal anti domain. An unusual hydrogen bond from NH-1 of protonated adenine to O-6 of guanine stabilizes the structure at acidic (but not neutral or basic) pH. Adenine was selected as partner for the modified guanine in the duplex because AF modification often leads to GC --> TA transversion mutations (see section B3bii of our accompanying grant proposal). One plausible mechanism for these genetic changes would be the formation of GAF to A mismaps during
replication, as in our duplex. This work has been published (Norman, et al., 1989).

2. AF-modified DNA duplex 11-mers with hypoxanthine, guanine or cytosine opposite the lesion site: As experimental data had suggested that G to G mispairs may also be involved in AF mutagenesis, the Patel-Grunberger group prepared the above modified 11-mer with guanine opposite the AF-modified guanine. The NMR data was difficult to interpret in this case, however, so the analog with hypoxanthine (which lacks the guanine amino group) opposite \( \text{G}^{\text{AF}} \) was prepared as well. It afforded data amenable to interpretation, and this data was of value in interpreting the results from the G to G mismatch. We have computed molecular views of both structures, which agree with the data. They reveal that AF is again situated in the minor groove of B-DNA, in an orientation that is very similar to that found in the A to G AF mismatch (Figure 16). A weak hydrogen bond to O-6 of the modified guanine from H-1 of hypoxanthine adds stability to the structure. This bond is absent in the G AF to G mispair. A manuscript describing this work is in preparation.

We have also computed, without experimental input, a molecular view of an analogous structure in a duplex in which a normal C partner is opposite AF-modified G (Broyde, et al., 1990).

3. An AAF modified duplex nonamer with deoxycytidine opposite the lesion site: We have solved the structure of an AAF-modified duplex nonamer with the usual Watson-Crick sequence: cytosine opposite the modified guanine. This work was carried out in collaboration with Prof. Thomas Krugh and coworkers at the University of Rochester, who carried out the NMR investigation. We employed Krugh’s measured interproton distances to guide our conformational searches with DUPLEX, and followed this with unconstrained molecular dynamics studies with the program AMBER, which includes solvent and salt. A major and minor structure were delineated. In both, the modified guanine is syn, rather than anti, which is normal in B-DNA. The guanine is displaced from its normal base-stacked position and does not hydrogen bond to the opposite cytosine. The major conformer features AAF-base stacking, while the minor one places the AAF in a position in which it protrudes from the minor groove, and lacks stacking interactions with adjacent bases. The DNA is bent in both structures.

A video of the dynamics trajectory (169 picoseconds) of the major conformer is included. The interproton distances during the trajectory are consistent with the NMR data. Thus the view is a realistic one of the solution mobility of this conformer, but the time frame was too short, due to computational limitations, to reveal a transition between the major and minor conformers. The time span required for this change would be orders of magnitude larger than our trajectory. Preliminary accounts of this work have been published (Broyde, et al., 1991; O’Handley, et al., 1992), and a fully detailed manuscript is in preparation.

An abnormal syn guanine thus appears to be an important feature of the conformation of AF- and AAF-modified DNAs. If it is not repaired by an error-free mechanism in vivo, then it could be
misinformational at the replication fork, and lead to a mutation. Some of our ideas on mutagenic conformations are presented in Broyde, et al., (1990).

4. The structure of a DNA duplex hexamer cross-linked by mitomycin: Mitomycin is a clinically useful antibiotic and antitumor agent, and a mutagen. This substance can form a monoadduct to the N-2 position of guanine or a diadduct to two appropriately positioned guanines on opposite strands. NMR studies by Prof. D. Patel and co-workers (Columbia University) on a di-adduct in a hexamer duplex synthesized by Prof. M. Tomasz and co-workers (Hunter College, City University) produced interproton distances. The data was used in our computations to model the structure.

Two structures were calculated that were compatible with the NMR data and indistinguishable experimentally. They differed in the conformation of the saturated ring of mitomycin, and may actually interconvert in solution. In both structures, the mitomycin was situated in a B-DNA duplex with a widened minor groove; the mitomycin resides in closer proximity to one of the two DNA strands. The end base pairs are not hydrogen-bonded. The cross-link is thought to be responsible for the anti-tumor action of mitomycin: presumably it interferes with replication by preventing the unwinding of DNA. Details of this work are given in Norman, et al., (1990).

5. Structures of the trans adducts formed at N-2 of guanine in DNA by (+) and (-) anti-benzo[a]pyrene diol epoxides (BPDE): It has been known for over a decade that (+) anti BPDE is tumorigenic while its (-) anti mirror image is not (Conney, 1982). The trans-anti major adduct formed by reaction of the (+) isomer with the N2-position of guanine is believed to be responsible for the carcinogenicity of the parent compound, while the mirror image adduct formed by the (-) reactant appears much less active in that respect (Conney, 1982). Thus, this pair of adducts offers an ideal system to explore structure-function relationships.

We have solved structures for these two adducts. Indeed, we predicted that the structural difference resides in opposite orientations of the pyrenyl moiety in the B-DNA minor groove. The long axis is oriented toward the 5' end of the modified strand in the (+) adduct, and toward the 3' end of the modified strand in the (-) adduct (Singh, et al., 1991).

Subsequently, experimental interproton distances for a DNA duplex 11-mer became available from synthetic efforts by Geacintov and co-workers (NYU) and the NMR studies of Patel and his colleagues (Columbia). We employed the data in computational studies to determine the structure of this adduct. The results verified fully our predictions concerning minor groove orientation (Cosman, et al., 1992; de los Santos et al., 1992).

6. Structure of the cis adduct formed at N-2 of guanine in DNA by (+) anti-benzo[a]pyrene diol epoxide (BPDE): This adduct is a minor product of the tumorigenic BPDE isomer. It is formed by cis, rather than trans, opening of the epoxide ring (see Figure 1 in Singh, et al., 1991). The structure of a DNA duplex 11-mer
containing this adduct has now been determined by the same collaboration given above for the BPDE trans-anti adducts. The result, illustrated in Figure 17, places the benzo[a]pyrene moiety in an intercalated position between adjacent base pairs. It displaces the modified guanine from its normal base-stacked position within the helix, and base-pairing by this guanine is ruptured. The guanine retains a glycosyl conformation within the anti range, but of a value 60° different from its usual one in B-DNA.

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Structure, numbering scheme, and variable conformational angle designations for d(CpG). The dihedral angles A-B-C-D are defined as follows: \( \chi, O1'-C1'-N1-C6(pyr); O1'-C1'-N9-C8(pur); \psi, C3'-C4'-C5'-O5'; \varphi, P-O3'-C3'-C4'; \psi', C4'-C5'-O5'-P; \omega', O5'-P-O3'-C3'; \omega, C5'-O5'-P-O3'. \) The angle A-B-C-D is measured by a clockwise rotation of D with respect to A, looking down the B-C bond. A ecliping D is 0°. Sugar puckering in the calculations is defined by the pseudorotation parameter P. An additional flexible torsion angle, Me, employed for the thymine methyl group is defined as C6-C5-C-H. IUPAC torsion angle designations are given in parentheses. In the IUPAC convention 180° is added to \( \chi \).
LOWEST ENERGY CONFORMERS

Figure 2.
\( d(G\text{-}AF\text{-}pC) \)

Figure 3.
\( d(G\text{-}AAF\text{-}pA) \)

Figure 4.
\( d(CpG\text{-}AAF) \)

Figure 5.
\( d(ApG\text{-}AAF) \)
**Figure 10.**
\[ \Delta E = 0 \text{ kcal/mol} \]

**Figure 11.**
\[ \Delta E = 1.48 \text{ kcal/mol} \]

**Figure 12.**
\[ \Delta E = 0.05 \text{ kcal/mol} \]

**Figure 13.**
\[ \Delta E = 1.08 \text{ kcal/mol} \]
Figure 14.
d(GpCpG-AF-pCpG Lowest Energy Conformer

Figure 15.
d(GpCpG-AAF-pCpG Lowest Energy Conformer
Figure 16. Structures of AF modified DNA duplex
11-mers computed with NMR measured interproton
distances obtained from Patel-Grunberger laboratories
(Columbia University). (a) AF-G...I (b) AF-G...G
Figure 17. Structure of the (+) cis-anti adduct of BPDE in a duplex 11-mer, computed with the aid of NMR data obtained in the laboratory of D. J. Patel (Columbia).
PUBLICATIONS


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