Site-Specific Mutagenesis in *Escherichia coli* by Bulky Exocyclic Amino-Substituted Guanine and Adenine Derivatives in Double-Stranded or Gapped Plasmids

Ki-Young Moon, Ph.D.

Department of Clinical Pathology, and Bioindustry and Technology Research Institute, Kwangju Health College, Gwangju, Korea

**Purpose:** 7-Bromomethylbenz[a]anthracene is a known mutagen and carcinogen. The mutagenic potency of its two major DNA adducts, i.e., N²-(benz[a]anthracen-7-ylmethyl)-2’-deoxyguanosine (b[qa]²G) and N²-(benz[a]anthracen-7-ylmethyl)-2’-deoxyadenosine (b[qa]²A), as well as the simpler benzylated analogs, N²-benzyl-2’-deoxyguanosine (bn²G) and N²-benzyl-2’-deoxyadenosine (bn²A), were determined in *E. coli*.

**Materials and Methods:** Double-stranded and gapped plasmid vectors were used to determine the mutagenicity of b[qa]²G, b[qa]²A, bn²G and bn²A in *E. coli*. The four, suitably protected, bulky exocyclic amino-substituted adducts were incorporated into 16-base oligodeoxyribonucleotides, in place of normal guanine or adenine residues, which form part of the ATG initiation codon for the lacZ’ complementation gene. The site-specifically modified oligodeoxyribonucleotides were then incorporated into double-stranded plasmids, which contained uracil residues in the complementary strand in the vicinity of the initiation codon. The uracil residues lead to the creation of a gap in the complementary strand due to the actions of *E. coli* uracil-DNA glycosylase and AP endonuclease. Following the transfection of these plasmid vectors into *E. coli* strain GP102, a lacZ alpha complementing version of the parent strain AB1157, their propensity to induce mutation was investigated.

**Results:** The percentages of mutant colonies produced by the four modified nucleosides, in both the double-stranded and gapped plasmid vectors, were not significantly different from those produced by the unmodified plasmids. The mutagenicities of the b[qa]²G and b[qa]²A were extremely low, and a totally unexpected result, whereas, those of the bn²G and bn²A were undetectable.

**Conclusion:** In this *E. coli* site-specific mutagenesis system, these bulky aralkylated adducts exhibited no significant mutagenicities, either with or without SOS induction. (Cancer Research and Treatment 2003;35:75-80)

**Key Words:** Mutagenesis, 7-Bromomethylbenz[a]anthracene, Gapped plasmid, Bulky exocyclic amino-substituted adducts, SOS induction, *E. coli*

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**INTRODUCTION**

Exposure to low levels of chemical carcinogens in the environment is essentially unavoidable, and their damage to cellular DNA has been recognized in the initiation of carcinogenic processes through mutagenic mechanisms. It is difficult to evaluate how any one carcinogen adduct can alter the structure, or coding properties, of the damaged DNA, as carcinogens can react with a variety of sites on DNA bases, and with several bases simultaneously. Therefore, the synthetic methods for the preparation of oligodeoxyribonucleotides, containing a single specific carcinogen-modified base at defined sites, have been developed, and play key roles in the investigations of individual carcinogen-DNA adducts, and how they induce mutagenic potency in biological systems (1,2).

The environmental pollutant 7-bromomethylbenz[a]anthracene (7-BrMeBA) is a known mutagen and carcinogen (3-7). The exocyclic N² and N⁶ amino groups of deoxyguanosine and deoxyadenosine in DNA are major sites of aralkylation. N²-(benz[a]anthracen-7-ylmethyl)-2’-deoxyguanosine (b[qa]²G) and N⁶-(benz[a]anthracen-7-ylmethyl)-2’-deoxyadenosine (b[qa]⁶A) are the major modified deoxyribonucleosides produced in DNA by this carcinogen.

This study was based on synthesized oligodeoxyribonucleotides, containing a single carcinogen-modified base at a specific location. To determine the mutagenic potencies of these bulky adducts, and that of the simpler benzylated analogs, i.e., N²-benzyl-2’-deoxyguanosine (bn²G) and N⁶-benzyl-2’-deoxyadenosine (bn⁶A).
The mutagenicities of the major DNA adducts produced by 7-BrMeBA, when the modified base was incorporated into plasmid vectors, was examined, and compared, in E. coli. The mutagenicities of these adducts were also compared to those of the simpler benzylated derivatives of guanine and adenine, i.e., bm'G and bm'A, in order to observe the effects of increasing the size of the exocyclic amino group substituents on mutagenic potency. This was performed to allow us to broaden our understanding on how substituted structure can influence the mutagenicity of bulky exocyclic amino-substituted adducts. Even though 7-BrMeBA is a known carcinogen, this is the first evaluation of the mutagenicity induced by the site-specific incorporation of its major adducts in E. coli.

This study now reports the site-specific mutagenesis induced by four aralkylated nucleosides in E. coli.

**MATERIALS AND METHODS**

The restriction enzymes, purchased from New England Biolabs, Beverly, MA, were used in the buffer supplied by the manufacturer, unless specified. The enzymes, and all other reagents used in this study, were described previously (11). The 2'-Deoxyguanosine and 2'-deoxyadenosine were purchased from Sigma Chemical Co., St. Louis, MO. Most other reagents and solvents were purchased from Aldrich Chemical Co., Inc., Milwaukee, WI. The 7-bromomethylbenz[a]anthracene was prepared as described previously (12). The N'-benzyl-2'-deoxyguanosine, N'-benzyl-2'-deoxyadenosine, N'-benz[a]anthracen-7-ylmethyl)-2'-deoxyguanosine, N'-benz[a]anthracen-7-ylmethyl)-2'-deoxyadenosine and the bulky aralkylated adducts, containing the 16-base oligodeoxyribonucleotides, were prepared as described previously (10).

1) **Synthesis and purification of oligonucleotides**

The 16-base oligodeoxyribonucleotides were prepared in an Applied Biosystems, Inc., Model 380B DNA synthesizer, and purified by polyacrylamide gel electrophoresis.

2) **Preparation of adduct containing plasmids**

Pairs of oligonucleotides were incorporated into the E. coli mutagenesis plasmid, pGP10 (Fig. 3), using the detailed procedure described previously (11,13  15). In these constructions, the vector DNA was digested with BglII restriction enzyme to remove the insert DNA sequence, as enclosed in the box, illustrated in Fig. 3. The major fragment produced during the BglII digestion was isolated, and recircularized by ligating pairs of oligonucleotides, as illustrated in Fig. 3, into the plasmid where the insert had been. Following the recircularization, the covalently closed plasmids were isolated.

3) **Mutagenesis experiments in E. coli**

The E. coli strain, GP102 (15), is a lacZ alpha complementing version of the parent strain, AB1157 (16). Electroporation competent GP102 cells were prepared, and transformed with adduct containing, or control, plasmids, according to the procedure described previously (13). In experiments where SOS induction was required, 100 ml of cells were grown to their mid log phase, as previously described (13), and then
pelleted by centrifugation at 1,600 g for 5 min at room temperature. The resulting cell pellet was suspended in 25 ml of 
“E” salts (17), transferred to a 100 mm sterile culture dish, with 
no lid, and irradiated with 75 J/m² of 254 nm light from a 
model UVGL-55 mineralight® lamp (UVP, Inc., San Gabriel, 
CA). This exposure had previously been determined to give a 
fifty percent survival of the GP102. The cells were re-pelleted 
under the same conditions, resuspended in 100 ml of fresh LB 
broth and incubated, with shaking, at 37°C for 30 min. The 
electroporation competent cells were then prepared, and trans-
formed with the plasmids, as described previously (13). 
Following the electroporation and recovery, the bacteria were 
plated onto media containing X-gal and IPTG, and incubated 
as described in previously (15). The plates were then scored 
for blue and white colonies.

4) Sequencing of mutant plasmids

In order to identify the different kinds of lacZ inactivating mutations that occurred, the adduct containing and control 
plasmids, the white colonies, were isolated and subcultured. 
The plasmid DNA was isolated, and the region surrounding the 
lacZ initiation codon sequenced using a dye terminator cycle 
sequencing kit (Perkin Elmer, Foster City, CA), with an 
Applied Biosystems 373A DNA sequencer (Foster City, CA).

RESULTS

1) Mutagenicity in double-stranded plasmids

A cassette plasmid permits monitoring of the mutagenicity of 
the carcinogen-modified bases in E. coli when they are 
positioned in the ATG initiation codon of the β-galactosidase 
gene (Fig. 3). This plasmid contains a nonfunctional lacZ' 
complementation gene, where the gene promoter (P) is separated 
from the coding sequence (Lac) by a 0.95-kilobase insert. This insert is flanked by two recognition sequences for the 
restriction enzyme, BspMI, and treatment of the plasmid 
with this enzyme permits the removal of the insert sequence. 
The recircularization of the plasmid is accomplished by ligating 
the partially complementary 16-base sequence, created by the 
BspMI digestion, to the stick ends of the major DNA fragment. 
The joining of the promoter, to the lacZ' coding sequence by a 
16-base oligonucleotide containing the ATG initiation codon, 
results (Fig. 3). This system was used to measure the muta-
ticities of the βn²G, βn²A, βGα²G and βGα²A residues that had been incorporated in place of the normal 2'-deoxyguanosine and 2'-deoxyadenosine of the ATG initiation codon. 
The site-specific mutagenesis induced by these bulky adducts 
was monitored using the sectored colony assay system, as 
previously described (11). The adduct-induced mutations inacti-
uate the gene, which led to the formation of blue and white 
colored colonies when the transformants from an E-comple-

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Table 1. Relative transformation efficiency (RTE) from double-
stranded plasmids in wild type E. coli

<table>
<thead>
<tr>
<th>Adduct</th>
<th>RTE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>G or A</td>
<td>100b</td>
</tr>
<tr>
<td>βn²G</td>
<td>93.8±17.0</td>
</tr>
<tr>
<td>βn²A</td>
<td>107.2±24.3</td>
</tr>
<tr>
<td>bGα²G</td>
<td>84.6±10.8</td>
</tr>
<tr>
<td>bGα²A</td>
<td>116.7±5.0</td>
</tr>
</tbody>
</table>

bMean±SD of four experiments. This value is defined as 100 for each experiment. The actual number of colonies ranged between 1500 and 4400.
menting version of the *E. coli* strain, AB 1157, are grown on media containing 5-bromo-4-chloro-3-indolyl β-D-galactoside (X-gal). In the absence of a mutation, blue colonies are produced (11). Table 1 shows the percentage of colonies produced by the double-stranded plasmids in wild type *E. coli*. There was no effect on the transformation efficiency, whether the double-stranded plasmids contained these bulky adducts, or not, indicating that none of these bulky adducts impedes the DNA replication. The percentages of mutant colonies produced by the modified plasmids were not significantly different from those produced by the unmodified plasmids (Fig. 4). The mutagenicities of the b[ga’G] and b[ga’A] were extremely low, which was totally unexpected. This study aimed to quantify the mutagenic potencies induced by these bulky adducts since 7-BrMeBA is a known mutagen and carcinogen. No mutagenicity for the bn’G and bn’A were detectable (Fig. 4).

2) Mutagenicity in gapped plasmids

The modified oligodeoxynucleotides were incorporated into double-stranded plasmids containing uracil residues, which led to the creation of a gap in the complementary strand due to the action of the *E. coli* uracil-DNA glycosylase. The *E. coli* containing the uracil-DNA glycosylase (UDG) and AP endonu-

3) SOS induction

SOS induction is commonly used for bulky adducts to observe increases in the mutagenicity (18). To test whether SOS induction had any effect on mutagenicities of the bulky adducts, it was induced by irradiating *E. coli* with UV light (75 J, 254 nm). However, the UV pretreatment of these adducts in *E. coli* showed no distinguishable change in the mutagenicities compared to that of the control (Fig. 5). The SOS induction was considered to have no measurable effects on the mutagenicities of the adducts.

4) Mutations in *E. coli*

Following the transformation, with the control and adduct containing plasmids, ten white colonies were chosen. The plasmid DNA was isolated and sequenced. The majority of the identified mutations were most likely artifacts of the plasmid manipulation. In the case of b[ga’G], one G-to-C and one G-to-T transversions, and in the case of b[ga’A], a single A-to-T transversion, were observed at the site of the adduct. No base changes in, or near, the initiation codon could be reliably attributed to the presence of either the bn’G or bn’A.

**DISCUSSION**

The primary objective of this study was to examine the site-specific mutagenicity in *E. coli* of the major DNA adducts produced by the aralkylating agent, 7-(bromomethyl)benz[a]anthracene, i.e., b[ga’G] and b[ga’A], when the modified base was incorporated into either a double-stranded or gapped plasmid vector. We then compare these data to those for the analogous, but simpler benzylation derivatives of guanine and adenine, to observe the effect of increasing the size of the exocyclic amino group substituent on the mutagenic potency, and how a substituted structure can influence the mutagenicity of bulky exocyclic amino-substituted adducts. Unfortunately,
however, the four aralkylated bulky adducts revealed no mutagenic potencies (Fig. 4 and 5), even though 7-bromomethylbenz[a]anthracene is a known carcinogen (4–7). It has been assumed that the notable low frequency of mutations induced by adducts, in site-specific mutagenesis studies, resulted from the efficient repair of the adduct. The lack in a mutation frequency observed for the C[4]G, C[4]A, C[6]G and C[6]A residues might be as a result of repairs. However, this seems an unlikely explanation since active excision repairs would not be expected on gapped plasmids containing four aralkylated bulky adducts (13,15).

Several studies have shown marked differences in the mutagenic potencies of the exocyclic DNA adducts [e.g., 1,N -ethenodeoxyguanosine, 2,N -ethenodeoxyguanosine, 3,N -ethenodeoxyctydine and 1,N -ethenodeoxadenosine] produced by chloroethylen oxide and chloroacetalddehyde, which are reactive metabolites of the human carcinogen, vinyl chloride, between E. coli and mammalian cells (19–21). 1,N -Ethenodeoxyadenosine, one of the four exocyclic DNA adducts, was highly mutagenic in mammalian cells, but only marginally mutagenic in E. coli (19). 1,N -ethenodeoxyguanosine was highly mutagenic in E. coli, but only weakly mutagenic in mammalian cells. For 3,N -ethenodeoxyctydine, a converse relationship was observed (20). These data strongly suggest that the site-specific mutagenesis of C[4]G, C[4]A, C[6]G and C[6]A should be examined in human cells.

The sequence context has profoundly influenced the mutagenic potencies of some bulky adducts in site-specific mutagenesis studies (22–25). This sequence context effect may be the reason why the bulky aralkylated adducts showed no significant mutagenicities in E. coli.

CONCLUSIONS

In this E. coli site-specific mutagenesis system, none of the aralkylated adducts exhibited any significant genotoxicity or mutagenicity. A SOS induction had no measurable effect on the mutagenicity of these adducts. The low genotoxicity and mutagenicity of these adducts may be either intrinsic to their structure, or as a result of a sequence context effect, in this system. Further studies are in progress to elucidate the mechanism by which these aralkylated adducts exert no mutagenic effect in E. coli. Other studies will also be required to determine the mutagenicity of these same adducts in human cells.

REFERENCES

22. Rodriguez H, Loechler EL. Mutational specificity of the (+) anti-diol epoxide of benzo[a]pyrene in a sup F gene of an

