Protein Modification by Adenine Propenal

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Supporting Information

ABSTRACT: Base propenals are products of the reaction of DNA with oxidants such as peroxynitrite and bleomycin. The most reactive base propenal, adenine propenal, is mutagenic in Escherichia coli and reacts with DNA to form covalent adducts; however, the reaction of adenine propenal with protein has not yet been investigated. A survey of the reaction of adenine propenal with amino acids revealed that lysine and cysteine form adducts, whereas histidine and arginine do not. N′-Oxopropenyllysine, a lysine–lysine cross-link, and S-oxopropenyl cysteine are the major products. Comprehensive profiling of the reaction of adenine propenal with human serum albumin and the DNA repair protein, XPA, revealed that the only stable adduct is N′-oxopropenyllysine. The most reactive sites for modification in human albumin are K190 and K351. Three sites of modification of XPA are in the DNA-binding domain, and two sites are subject to regulatory acetylation. Modification by adenine propenal dramatically reduces XPA’s ability to bind to a DNA substrate.

INTRODUCTION

The major reactive aldehyde produced during peroxidation of polysaturated fatty acids, malondialdehyde (MDA), reacts with nucleophilic sites of DNA and protein to form oxopropenyl adducts. Reaction of MDA with DNA produces primarily 3-(2-deoxy-β-D-erythro-pentofuranosyl)pyrimido[1,2-a]purine-10(3H)-one (M1dG) and lesser amounts of 2′-oxopropenyl-deoxyadenosine (Figure 1). M1dG is mutagenic in vitro and in vivo, and 2′-oxopropenyl-deoxyadenosine alters DNA polymerase activity.2–4

The reaction of MDA with protein occurs predominantly at lysine residues under physiological conditions.5–9 Our understanding of the detailed chemistry and physiological consequences of MDA–protein adduct formation is rudimentary, as the exact targets and sites of adduction are largely unknown. However, a particularly interesting effect of MDA exposure in human cells is the inhibition of nucleotide excision repair (NER), which is associated with increased sensitivity to the mutagenicity of ultraviolet light and benzo[a]pyrene dihydrodiol epoxide.10 NER is the primary process for repair of bulky DNA adducts, and in humans, NER deficiencies result in xeroderma pigmentosum, a spectrum of disorders characterized by hypersensitivity to sunlight, neurological degeneration, and a greatly increased incidence of cancer.11 Although not verified experimentally, inhibition of NER by MDA was proposed to arise from loss of function of MDA-modified NER proteins.10 A potential MDA target is the NER scaffold protein, xeroderma pigmentosum complementation group A (XPA), which binds to the damaged DNA substrate via a C-terminal basic cleft comprising multiple lysine residues.12,13 XPA coordinates recruitment and assembly of the enzymes that repair the damaged DNA, and mutations in its DNA-binding domain are associated with the most severe clinical XP phenotypes.11

Oxopropenyl adducts to DNA and protein can also be formed through reaction with base propenal, a reactive aldehyde produced by oxidant-mediated hydrogen abstraction from the 4′-position of DNA bases (Figure 1).14–17 Base adducts are approximately 100-fold more reactive to DNA than MDA, and experiments in which Escherichia coli strains of varying membrane composition were exposed to oxidants demonstrated that base propenals are more important than MDA as a source of M1dG.18 Currently, very little is known about potential protein targets of base propenal addition; however, as a highly reactive form of DNA damage, base propenals are ideally situated to oxopropenylate DNA-binding proteins such as XPA. This has led us to hypothesize that base propenal-mediated protein damage contributes to toxicity and mutagenicity associated with oxidative or electrophile stress. To test this hypothesis, a comprehensive evaluation of base propenal’s ability to directly damage proteins is required. Adenine propenal is known to react with glutathione via Michael addition to form glutathionylepropenal and a glutathione–adenine propenal cross-link,19 but, to date, no detailed investigation of the reaction of any base propenal with amino acids or proteins has been reported.

Received: June 3, 2014
Published: September 11, 2014
One approach to evaluate the protein adduction potential of an electrophile is to identify all sites of modification and the structures of those modifications using a model protein. A frequently used target for this purpose is human serum albumin (HSA). Albumin is highly abundant and has affinity for a broad range of ligands, a relatively long half-life, and the potential for noninvasive sampling. Thus, it is an ideal candidate for the global qualitative and quantitative evaluation of exposure to electrophiles of endogenous or exogenous origin (the exposome). In fact, in a large multilaboratory comparison of biomarkers of oxidant generation, the level of carbonylated plasma protein (i.e., albumin) was one of only two markers that correlated with oxidant exposure and tissue pathology, and it was the only marker that was effective longer than 6 h after exposure. To exploit albumin’s potential as a biomarker, however, requires a more complete understanding of its major sites of adduction and determinants of their reactivity to a range of electrophiles. For example, whereas much attention has been directed to HSA’s highly reactive Cys-34 as a sensor of oxidant damage, the major site of oxopropenylation by MDA is Lys-525.7 Here, we report the application of a recently described mass spectrometric approach to identify adenine propenal modifications of HSA and XPA. Our data demonstrate that, like MDA, adenine propenal primarily reacts at protein lysine residues; however, the major site of attack on HSA is distinct from that of MDA. We also report that adenine propenal modifies lysine residues of XPA, including residues within its DNA-binding domain and residues reported to be sites of regulation via SIRT1-mediated deacetylation. Biophysical analysis confirms that modification of XPA by adenine propenal decreases its affinity for damaged DNA.

**EXPERIMENTAL PROCEDURES**

**Materials.** HSA was obtained from Abcam (Cambridge, MA). N-α-Acetylsine, trichloroacetic acid (TCA), dithiothreitol, iodoacetate (Sigma Ultra), and NaCNBH3 were obtained from Sigma-Aldrich (St. Louis, MO). Trypsin (gold) was obtained from Promega (Madison,
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Preparation of the Lysine Cross-Link (4). The sodium salt of MDA (100 mg, 1.06 mmol) was added to a solution of N-acetyllysine (200 mg, 1.06 mmol) in 5 mL of ammonium acetate buffer (pH 4.5). The reaction was then stirred at room temperature for 15 h. The reaction mixture was lyophilized, and the residue was purified by column chromatography (EtOAc/MeOH, 5:1) on silica gel (60 Å porosity, 40–63 μm particle size). NMR (D2O): δ (ppm) 13.0 (m, 4H), 1.59–1.61 (m, 6H), 1.73 (m, 2H), 1.88 (s, 6H), 3.22 (m, 2H), 3.30 (t, J = 7.3 Hz, 2H), 4.08 (m, 2H), 5.45 (m, 1H), 7.49 (d, J = 11.5 Hz, 1H), 7.56 (d, J = 11.5 Hz, 1H). 13C NMR (D2O): δ (ppm) 21.3, 21.8, 25.7, 30.2, 42.6, 53.7, 88.5, 158.9, 160.5, 177.5, 178.9. ESI-MS: [M+H]+, m/z 413.3. (See Supporting Information Figures S5–S7.)

HPLC Analysis of N-α-Acetyllysine and Adenine Propanol Reactions. N-α-Acetyllysine (various concentrations) was mixed with 10 mM adenosine propenal in 50 μL reaction mixtures containing 10 mM NaCl and 10 mM 4-morpholinepropanesulfonic acid (MOPS) (pH 6.5). At 1 h intervals, aliquots of each reaction mixture were injected onto a Waters HPLC system equipped with a 717 Plus autosampler, 1525 Binary HPLC pump, and 2996 photodiode array detector. Analyses were separated using a Phenomenex Polar-RP Synergy HPLC column, 75 cm × 2 mm with 4 μm particle size, at a flow rate of 0.6 mL/min. The mobile phase solvents were H2O + 0.05% acetic acid (v/v, solvent A) and 3:1 MeOH/ACN + 0.05% acetic acid (v/v, solvent B) delivered in a 19 min gradient consisting of the following: 0–10.5 min, 5–20% B; 10.5–12.5 min, 20–80% B; 12.5–16 min, 80% B; 16–19 min, 5% B. Data were analyzed using Empower software, and peak areas were manually integrated. Eluting peaks were collected and analyzed by MS (see below).

MS Analysis of the Reaction Products of Adenine Propanol and N-α-Acetyllysine. Reactions were performed as described above and analyzed using a Thermo Accela HPLC pump (Thermo Fisher Scientific) and HTS PAL autosampler (LEAP Technologies) in-line with a Thermo Quantum triple quadrupole mass spectrometer (Thermo Fisher Scientific). The Quantum was equipped with an electrospray source operated in Q1 full scan, positive ion mode. Q1 was scanned from 100–600 amu in 0.45 s. Analytes were chromatographically separated on a Synergi Polar-RP HPLC column (Phenomenex, 7.5 × 0.2 cm, 4 μm particle size) using the following gradient: 0–10.5 min, 5–20% B; 10.5–12.5 min, 20–80% B; 12.5–16 min, 80% B; 16–17 min, 5% B. The mobile phase solvents were H2O + 0.05% acetic acid (v/v, solvent A) and acetonitrile + 0.05% acetic acid (v/v, solvent B), and they were delivered at a flow rate of 0.6 mL/min. All data were acquired and analyzed using Thermo Xcalibur software.

Spectrophotometric Quantification of the Lysine–Lysine Cross-Link (4) in Reaction Mixtures of N-α-Acetyllysine and Adenine Propanol. Solutions containing known concentrations of synthetic 3 and 4 were used to measure extinction coefficients of each compound. The value obtained for both compounds (33 600 ± 200 M−1 cm−1) was the same within experimental error; however, the wavelengths of maximal absorbance were different (280 nm for 3 and 300 nm for 4) (Figure S1). Adenine propanol (10 mM) was incubated with increasing concentrations of N-α-acetyllysine (1, 2, 5, 10, 50, and 100 mM) in reaction mixtures containing 10 mM NaCl and 10 mM MOPS (pH 6.5). The mixtures were incubated for 2 h at room temperature and then diluted 1:1000 in H2O. The absorbance at 300 nm of each dilution was then obtained using a DU800 spectrophotometer (Beckman Coulter), and the experimental extinction coefficient of 4 was used to estimate its concentration in the reaction mixture. Note that, for reaction mixtures containing >10 mM N-α-acetyllysine, complete consumption of adenine propanol eliminates concerns of its contribution to the absorbance of the final reaction mixture. The adenine product does not absorb significantly at 300 nm (Figure S1). On the basis of its full UV spectrum (Figure S1), the extinction coefficient of 3 at 300 nm is estimated to be 7.8 × 105 M−1 cm−1. The presence of this product at a concentration equal to approximately 10% of the concentration of 4, as found in the reaction mixture, indicates that its contribution to the absorbance at 300 nm will be no more than 3%. Consequently, the absorbance at 300 nm can be used to estimate the concentration of 4 with reasonable accuracy.

Reaction of N-α-Acetyllysine with Adenine Propanol under Reducing Conditions. N-α-Acetyllysine (10 mM) was incubated with adenine propanol (10 mM) in 10 mM MOPS (pH 6.5) containing 10 mM NaCl in a 200 μL reaction mixture for 30 min. After incubation, 50 mM NaCNBH3 (in H2O) (Sigma-Aldrich) was added for an additional 30 min, and samples were then analyzed using LC-MS as described below.
Analysis of Reduced Intermediate (6) by LC-MS. LC-MS analysis was performed on the Thermo Quantum triple quadrupole mass spectrometer. Here, the instrument was operated in Q1 full-scan negative ion mode using electrospray ionization. Q1 was scanned over the range 100–400 amu over 0.5 s. Chromatography was carried out on a Syngery Max-RP column (Phenomenex, 7.5 × 0.2 cm, 5 μm particle size) using the following gradient: 0–5 min linear gradient from 2% B to 50% B, 5–7 min at 50% B, 7–10 min 50% B to 2% B. The mobile phase solvents were H2O + 0.05% formic acid (v/v, solvent A) and 3:1 CH3CN/MeOH + 0.05% formic acid (v/v, solvent B), and they were delivered at a flow rate of 400 μL/min. All data were acquired and analyzed using Thermo Xcalibur software. Using this chromatographic system, the reduced intermediate eluted at approximately 2.9 min.

Analysis of the Reaction of N-Acetylcysteine with Adenine Propenal. N-Acetylcysteine (200 μM) was combined with 50 μM adenine propenal in 50 mM sodium phosphate buffer, pH 7.4, and incubated for 6 h at 37 °C. Unreduced sample was immediately frozen. NaN3 (50 mM) (Sigma-Aldrich) was used to reduce samples as indicated. Analytes were separated on a Supelco Ascentis C18 column (50 × 2.1 mm, 3 μm) with buffers A and B comprising 0.1% formic acid in water and 0.1% formic acid in acetonitrile, respectively. The gradient was as follows: 1–3 min, 1% B; 3–7 min, 98% B; 7–11 min, 98% B; 11–17 min, 1% B. Samples were analyzed on a ThermoFinnigan TSQ Quantum mass spectrometer with ESI source interfaced to a ThermoFinnigan MS pump plus and autosampler plus (Thermo, San Jose, CA). N-Acetylcysteine-added species were analyzed with data-dependent scanning enabled in negative ion mode, scanning from m/z 150–650 over 0.5 s. The top two m/z peaks from each MS scan were fragmented with 10 eV collision energy for 0.5 s.

LC-MS/MS Analysis of HSA Modification by Adenine Propenal. HSA (15 μM) (Abcam, Cambridge, MA) was incubated with increasing amounts of adenine propenal (0.15, 0.38, 1.5, 3, and 7 mM), representing a molar excess of adenine propenal to HSA of 10, 25, 100, 250, and 500, respectively. Reactions were incubated for 6 h at room temperature in a buffer containing 10 mM NaCl and 10 mM MOPS (pH 6.5). Following incubation, samples were precipitated with 25% trichloroacetic acid (Sigma-Aldrich) on ice for 1 h, washed with cold acetone, dried, and reconstituted in 50 mM Tris (pH 8.0) containing 50% 2,2,2-trifluoroethanol (Acros Organics, Pittsburgh, PA). Samples were reduced with DTT (Sigma-Aldrich), carboximdemethy- lated with iodoaceticamide (Sigma-Aldrich), and diluted 5-fold with 100 μM Tris (to obtain a final solution containing 10% 2,2,2-trifluoroethanol). Following dilution, samples were digested with sequencing-grade trypsin (Promega, Madison, WI) overnight, acidified, and diluted in 0.1% formic acid. The resulting solutions of tryptic-generated peptides were loaded onto a capillary reverse-phase analytical column (360 μm o.d. × 100 μm i.d.) using an Eksigent NanoLC Ultra HPLC and autosampler. The 20 cm analytical column was directly packed into a laser-pulled emitter tip using Jupiter C18 reverse-phase medium (3 μm beads, 300 Å pore size, Phenomenex). Mobile phase solvents consisting of 0.1% formic acid in water (solvent A) and 0.1% formic acid in acetonitrile (solvent B) at a flow rate of 500 nL/min were used to elute the peptides. The 90 min gradient consisted of the following: 0–10 min, 2% B; 10–50 min, 2–35% B; 50–60 min, 35–95% B; 60–65 min, 95% B; 65–70 min 95% B; 70–90 min, 2% B. Mass analysis of eluting peptides was performed on an LTQ Orbitrap XL mass spectrometer (Thermo Scientific), equipped with a nano-electrospray ionization source with detection in positive ion mode. The LTQ Orbitrap was operated using a data-dependent method, enabling dynamic exclusion. Full scan (m/z 400–2000) spectra were acquired with the Orbitrap (resolution 60 000), and the five most abundant ions in each MS scan were selected for fragmentation in the LTQ. An isolation width of 2 m/z, 30 ms activation time, and collision energy normalized to 35% were used to generate MS2 spectra. Dynamic exclusion settings allowed for a repeat count of 2 within a 10 s duration, with exclusion duration time set to 15 s. For identification of modified peptides, tandem mass spectra were searched with SEQUEST (Thermo Fisher Scientific) against a human subset database created from the UniProtKB protein database (www.uniprot.org). Variable modifications of +57.0214 on Cys (carboximidemethylation), +15.9949 on Met (oxidation), and +54.0105 on Lys or Cys (oxopropanelylation) were included for database searching. Search results were assembled with Scaffold 3.0 (Proteome Software) using threshold filtering criteria consisting of 95% peptide probability to achieve a peptide false discovery rate estimate of ≤0.2%. Sites of modification were validated by manual interrogation of tandem mass spectra using Xcalibur 2.1 Qual Browser software. Calculations of mass errors (in ppm) of precursor ions mass analyzed in the Orbitrap were performed using the following equation: [(theoretical mass − observed mass)/theoretical mass] × 106. To obtain theoretical masses, the Protein Prospector MS-Product tool was used (v.5.10.4. UCSC). Analysis of cross-links was performed in a similar manner with inclusion of +36 on Lys for database searching.

Proteolytic Digestion of HSA Treated with Adenine Propenal for Lysine–Lysine Cross-Link (4) Identification. Purified HSA (0.75 μM) was incubated with increasing concentrations of adenine propenal in reaction mixtures containing 10 mM NaCl and 10 mM MOPS (pH 6.5) for 6 h at room temperature. Following incubation, the protein was heated at 95 °C for 10 min, cooled to room temperature, and then treated with 1 mg of Pronase (Calbiochem, Gibbstown, NJ) per mg of protein at 37 °C for 24 h. Pronase was inactivated by heating at 95 °C for 10 min, followed by cooling to room temperature, and treating with 1 μL of aminopeptidase M (Calbiochem, Gibbstown, NJ) at 37 °C for 24 h. A final incubation at 95 °C for 10 min inactivated the aminopeptidase M, and the samples were then cooled to room temperature for further purification.

Purification of Lysine–Lysine Cross-Links (4) and Analysis by MS. For use as an internal standard, [3H]-[15C]-4 was prepared by reaction of MDA with [3H]-lysine containing [14C]-lysine as a tracer and quantified using a method similar to that described for the [13C]-[15C]-lysine-lactam internal standard.28 For quantitation, the transition between m/z 341 and m/z 306 was monitored; it corresponds to the loss of one molecule of water and one molecule of ammonia. Portions of samples containing the standard were then concentrated using 1 cc Oasis HLB columns and dried under nitrogen. The purified samples were analyzed using a Thermo Scientific TSQ Vantage quadrupole mass spectrometer equipped with Xcalibur software for data collection and manipulation. All measured MDA-adduct signal in each sample was divided by the signal for the [3H]-[15C]-lysine–lysine cross-link standard to quantify experimental results. These methods will be published in detail elsewhere.

Proteomics Analysis of XPA Modification by Adenine Propenal. XPA was expressed and purified as described.30 To modify XPA, 3.2 μM of purified protein was incubated with 500X (1.6 mM) of adenine propenal for 6 h at room temperature in a buffer containing 10 mM NaCl and 10 mM MOPS (pH 6.5). Modifications were identified as described above for HSA.

Fluorescence Anisotropy DNA Binding Assay. A high-throughput fluorescence anisotropy assay was used to measure the DNA-binding activity of XPA.31,32 The substrate for this assay was a fluorescein-labeled Y-shaped ssDNA–dsDNA junction substrate containing 8 base pairs of duplex and 12 nucleotide noncomplementary ssDNA overhangs at both the 3′ and 5′ ends. XPA was modified with NHS–biotin as described previously.33 Modified or mock-treated XPA protein and the DNA substrate were diluted in binding buffer (20 mM NaCl, 10 mM Tris (pH 7.4), 2 mM MgCl2). The assay was run in a total volume of 4 μL. The fluorescence anisotropy was determined using a FL-221 spectrofluorometer (Jasco, Easton, MD) and is reported as the average of at least three independent experiments for each condition.
HEPES, pH 7.9, 75 mM KCl, 5 mM MgCl₂, 5% glycerol, 1 mM DTT. DNA substrate was added to a final concentration of 50 nM. Fluorescence anisotropy was measured using a Synergy H1 plate reader (λ<sub>Ex</sub> = 485 nm, λ<sub>Em</sub> = 528 nm). Binding measurements were acquired in triplicate for each DNA substrate. Apparent dissociation constants (<i>K<sub>d</sub></i>) were determined for each individual titration by plotting fluorescence anisotropy against protein concentration and fitting to a simple two-state binding model with KaleidaGraph (v4.03) software.

**RESULTS**

Adenine Propenal Modifies N-α-Acetyllysine, Forming N<sup>ε</sup>-Oxopropenyl-N-α-acetyllysine (3) and Lysine—Lysine Crosslink (4).
Cross-Links (4). Adenine propenal was reacted with $N\alpha$-acetyllysine, $N\alpha$-acetylhistidine, $N\alpha$-acetylariginine, or $N\alpha$-acetylcysteine in 10 mM NaCl buffered with 10 mM MOPS (pH 6.5). Reaction progress was monitored hourly by HPLC using a C18 reverse-phase column for product separation. Adenine propenal eluted at $\sim$8 min and displayed a maximum absorbance of 260 nm (Figures 2 and S1). Reaction with $N\alpha$-acetyllysine resulted in the appearance of three new chromatographic peaks at 1.0, 4.8, and 5.3 min, which were analyzed by UV/vis, NMR, and MS (Figures 2 and S1). The peak eluting at 1 min displayed a UV absorbance maximum at 260 nm and an $m/z$ of 136, corresponding to protonated adenine ([M + H]+) (1). The 4.8 min peak displayed a UV maximum at 280 nm. On the basis of NMR analysis, this product was identified as 3, and LC-MS analysis gave an $m/z$ of 243, which corresponds to the protonated species ([M + H]+). The 5.3 min peak, with a maximum absorbance at 300 nm and an $m/z$ of 413, which corresponds to the protonated species ([M + H]+), was identified as 4 by NMR analysis. The total amount of product formed was unaffected by variations in pH over a range of 6.5 to 9.5.

Chemical synthesis of 3 and 4 yielded standards that verified the product structure assignments. The molar extinction coefficients of the pure compounds ($\varepsilon = 33.6 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$ at 280 nm for 3 and $\varepsilon = 33.6 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$ at 300 nm for 4) were similar to that of adenine propenal ($\varepsilon = 32.5 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$ at 260 nm). HPLC analysis of reaction mixtures of $N\alpha$-acetyllysine and adenine propenal using diode-array detection enabled direct comparison of peak areas for each compound measured at its optimal wavelength (Figure 3). The essentially identical absorbance coefficients of 3 and 4 allowed peak areas to be used to estimate relative product quantity. These comparisons indicated that the major product formed from the reaction was 4, with much lower quantities of 3.

Variation of the amount of $N\alpha$-acetyllysine (from 2 to 100 mM) in reaction mixtures while holding the amount of adenine propenal (10 mM) fixed demonstrated that the amount of 3 formed increased with $N\alpha$-acetyllysine concentrations from 2 to 10 mM and then decreased somewhat at the higher concentrations. In contrast, the amount of 4 produced increased with $N\alpha$-acetyllysine concentrations up to 50 mM. On the basis of the relative peak area, the ratio of 4 to 3 in most reaction mixtures was approximately 10:1. Estimation of the concentration of 4 using absorbance at 300 nm indicated a maximum of $\sim$9 mM formed in reaction mixtures containing 50–100 mM $N\alpha$-acetyllysine (Figure S2). These results are consistent with complete consumption of adenine propenal under those conditions.

Our initial hypothesis for the production of 4 was that it arose by reaction of $N\alpha$-acetyllysine with 3, but this reaction did not occur under conditions comparable to those used for the reaction of adenine propenal with $N\alpha$-acetyllysine. An alternative
reaction scheme, in which an enaminoimine cross-link between adenine propenal and $N\alpha$-acetyllysine is first formed and then reacts with $H_2O$ to produce $3$ or another equivalent of $N\alpha$-acetyllysine to form $4$, is shown in Figure 4. If this scheme is correct, then our data indicate that, at the concentrations of $N\alpha$-acetyllysine and adenine propenal used here, reaction of the proposed intermediate with another molecule of $N\alpha$-acetyllysine is more favorable than its reaction with water. HPLC analysis with UV or in-line NMR was unable to detect the putative intermediate $5$ (Figure 4). Therefore, we attempted to trap it by reduction with NaCNBH$_3$. Adenine propenal was incubated with $N\alpha$-acetyllysine in $10$ mM NaCl/$10$ mM MOPS (pH 6.5) in the presence of NaCNBH$_3$. We utilized LC-MS analysis of the reaction mixture to detect ions with $m/z$ 360, which corresponds to the partially reduced intermediate ($6$). A peak corresponding to this mass eluted at $2.9$ min, consistent with the transient formation of the adenine propenal adduct to $N\alpha$-acetyllysine (Figure 5).

Reactions of adenine propenal with $N\alpha$-acetylhistidine and $N\alpha$-acetylarginine did not result in the formation of new chromatographic peaks. Reaction with $N\alpha$-acetylcysteine resulted in the formation of an oxopropenylated product, which displayed $m/z$ 216, corresponding to the deprotonated molecular ion, and its major fragmentation product, $m/z$ 86.9. Treatment of the reaction mixture with NaBH$_4$ yielded the partially reduced product ($m/z$ 218) and its major fragmentation product ($m/z$ 89) (Figure S3).

**Adenine Propenal Reacts Predominantly with Lys-214 and Lys-375 in HSA.** To ascertain adenine propenal’s ability to react with protein, we incubated increasing amounts of the electrophile with a fixed amount of purified HSA for $6$ h, and the samples were prepared for LC-MS/MS by trichloroacetic acid precipitation, resolubilization of the precipitated protein pellet, reduction and carbamidomethylation of cysteine residues, and proteolytic digestion with sequencing-grade trypsin. The resultant proteolytic peptides were analyzed via LC-MS/MS using an LTQ-Orbitrap Velos mass spectrometer with collision-
induced dissociation. Sequence coverage for unmodified HSA was found to be 90%, but exposure to increasing concentrations of adenine propenal reduced coverage to a minimum value of 62% (Table S1). HSA peptides were interrogated for the presence of a +$^{+4.0105}$ mass increase, as would be expected from the addition of an oxopropenyl group. To verify positively identified residues, data were manually examined using Xcalibur software. Extracted ion chromatograms were generated from precursor peptide masses, and theoretical masses calculated from Protein Prospector were compared to observed masses to calculate ppm mass error. For modified precursor peptides displaying a ppm mass error less than 10, MS/MS spectra were interrogated with Xcalibur software. At least 40% of each theoretical b- and y-product ion series was present in the observed product ion spectra of positively identified peptides, and 50% sequence coverage was obtained for each peptide validated. To validate the site of adduction, at least two ions within a product ion series with a +$^{+4.0105}$ adduction mass shift were required. Multiple lysine residues were found to be modified in adenine propenal-exposed HSA, including Lys26, Lys36, Lys161, Lys183, Lys214, Lys219, Lys236, Lys249, Lys257, Lys347, Lys375, Lys402, Lys413, Lys438, Lys549, Lys569, and Lys598. Residues verified to have a +$^{+4.0105}$ adduction mass were mapped to the crystal structure of HSA (PDB ID: 1AO6) and were found to be largely confined to the protein surface (Figure 6A). The 17 modified lysine residues represented 28% of the total lysines within the protein (Table 1). Although we determined that adenine propenal can also form oxopropenylated cysteine, searching for adducts on cysteine residues within HSA did not result in the identification of any modified sites. S-Oxopropenal-N-acetylcysteine was stable to treatment with TCA as well as DTT, which are reagents used for proteomic analysis, so any oxopropenylated cysteine residues on HSA-derived peptides should have survived the workup prior to mass spectrometry.\(^{34}\) Therefore, adenine propenal is acting as a lysine-specific modifier of HSA.

The use of HSA as a biomarker has primarily been described in the context of electrophilic modifications at Cys34.\(^{23}\) Adenine propenal did not modify Cys34, so we sought to identify the most reactive lysine residue(s) that might be used as a biomarker for modification by adenine propenal. We reduced the levels of adenine propenal relative to protein to a ratio of 10:1, which was the lowest ratio at which modification was observed. Using the interrogation methods described above to verify adduction sites, two lysine residues, K190 and K351, displayed the highest reactivity (Table 1). Mapping of those residues to the crystal structure of HSA revealed that both are located within domain II, a recognized binding region for many compounds (Figure 6A).\(^{35}\)

**Investigation of Cross-Link Formation in the Reaction of Adenine Propenal with HSA.** Decreased HSA sequence coverage observed with increasing concentrations of adenine propenal suggested that adducts impede protein digestion and/or fragmentation required for peptide identification. As lysine–lysine cross-links are formed from the reaction of adenine propenal with N-$\alpha$-acetyllysine, we hypothesized that at high concentrations of adenine propenal, lysine–lysine cross-links may be induced in HSA. To investigate the presence of cross-links, we interrogated previously acquired proteomics data for a +$^{+36}$ adduct mass, corresponding to an unreduced 3-carbon linkage between lysine residues.\(^{37}\) This analysis did not result in the identification of any lysine–lysine cross-links, which was not surprising considering the decreased sequence coverage and potential hindrance of peptide fragmentation by cross-links. Although cross-linked sites were not identified with these algorithm-based data searching methods, we manually interrogated the data to look for the presence of precursor peptides containing cross-linked residues. To predict lysine residues likely to participate in cross-link formation, we performed *in silico* analysis of the HSA crystal structure (1AO6). As a typical carbon–carbon bond is approximately 1.5 Å in length, we hypothesized that the 3-carbon cross-link would be ~5–7 Å long. Flexible, dynamic protein movements in solution may result in cross-link formation between lysines at a distance greater than 7 Å. We therefore examined the crystal structure to identify lysine residue pairs separated by a distance of less than or equal to 20 Å as potential sites of cross-link formation. Following identification of these residues, tryptic peptide sequences were determined from the primary sequence, and theoretical peptide masses containing the cross-link mass of +$^{+36}$ were calculated using Protein Prospector. We then thoroughly interrogated the data manually for the presence of ions of the theoretical precursor peptide masses. No evidence for the presence of cross-link-containing peptides was obtained.

As proteomics-based mass spectrometric analyses were unsuccessful, we employed a different analytical approach to search for lysine–lysine cross-links, in which modified HSA was degraded to the level of amino acids, and the resultant hydrolysate was analyzed by MS. This methodology employed selected reaction monitoring to identify analytes exhibiting a mass transition from m/z 329 to m/z 294, as exhibited by analysis of the lysine–lysine cross-link standard. Addition of a [${}^{3}$H]-[${}^{13}$C]-4 internal standard (m/z 341 to m/z 306 transition) enabled identification of analyte-containing fractions during purification by scintillation counting and quantification by stable isotope dilution. Application of this approach to HSA that had been incubated with adenine propenal resulted in the identification of 30 ± 12 ng cross-link/mg protein in vehicle-treated HSA and 57 ± 17 ng cross-link/mg protein in adenine propenal-treated HSA. These differences were not statistically significant, supporting the conclusion that adenine propenal did not induce intramolecular cross-links in HSA.

**Adenine Propenal Oxopropenylates XPA.** Having confirmed adenine propenal’s capacity to modify lysine residues in HSA, we investigated its reaction with XPA. Adenine propenal was incubated with XPA for 6 h at room temperature, and the samples were prepared for and analyzed by LC-MS/MS using the same methods as described for HSA. As for HSA, peptides were interrogated for the presence of a +$^{+4.0105}$ mass addition, and rigorous criteria were applied to peptide and site of adduction identification. The data revealed six modified residues, listed in Table 2. Of these, three of these, K145, K183, and K204, are

| residue | peptide |
|---------|---------|
| Lys-63 | PYSATAAAAATGGMANVK*AAPK |
| Lys-67 | AAPK*IIDTGGGFILEEEEQQK |
| Lys-86 | IIDTGGGFILEEEEQQK*IGK |
| Lys-145 | TEAK*QELYLK |
| Lys-183 | NPHHSQWGDME*KLYLK |
| Lys-204 | SLEVWSQALEEAK*EVR |

*Purified XPA was incubated with a 500-fold molar excess of adenine propenal and analyzed for lysine residues modified with +$^{+4.0105}$. The modified lysine in each peptide sequence is indicated with an asterisk.
Covalent modifications of DNA and proteins by electrophiles produced during oxidative stress have multiple consequences on cellular function, including mutagenesis, stress responses, cellular dysfunction, and ultimately cell death. All of these may contribute to pathological processes such as carcinogenesis, neurodegeneration, atherosclerosis, and chronic inflammation. However, elucidation of the exact mechanisms by which specific macromolecular damage leads to pathology requires a better understanding of the sites and nature of that damage.

The most frequently used biomarker of electrophile damage to protein during oxidative stress is the change in global protein carbonylation. Although useful as a general indicator that stress has occurred, this approach does not identify the exact type of modification or the modifying electrophile. Recent developments in highly sensitive MS-based proteomics techniques have allowed for expansion of this approach to the investigation of the specific types of modifications induced and the impact of these modifications on protein activity. The resultant increased ability to identify specific protein modifications induced by a single electrophile enables the search for biomarkers that distinguish between different mechanisms of oxidant damage.

The generation of adenine propenal occurs predominantly in intact DNA, suggesting that most of its targets will likely be DNA-associated proteins in the nucleus or mitochondria. Nevertheless, we initiated our investigation of adenine propenal’s protein-modifying profile, using HSA as the model protein. HSA’s reactivity with electrophilic agents has been studied extensively in the past, allowing us to place adenine propenal-dependent modifications in the context of those existing data. For example, teucrin A was found to preferentially modify Lys351 and Lys545, while 4-hydroxyxynenal modifies Lys195, Lys199, Lys233, and Lys525, and polycyclic aromatic hydrocarbon epoxides modify Lys137. As MDA has been shown to modify multiple lysine residues including Lys136, Lys174, Lys240, Lys281, and Lys525, and polycyclic aromatic hydrocarbon epoxides modify Lys137. As MDA and adenine propenal both produce oxopropanyl adducts on lysine residues, it is interesting to note that the major sites of adduction for the two electrophiles are different: Lys190 and Lys351 for adenine propenal versus Lys525 for MDA. These findings suggest that distinct HSA modification patterns may be used as markers for different electrophiles (Figure 6B). Clearly, the use of HSA as a biomarker of electrophile damage is attractive, given its high concentration in the blood and ease of sampling. However, as noted above, adenine propenal generation is largely confined to intact DNA, so it remains to be seen if enough of the

Table 3. Dissociation Constants for XPA Binding to Y-Shaped ssDNA–dsDNA Junction Substrate

| treatment           | $K_d$ (μM) |
|---------------------|------------|
| unmodified          | 0.30 ± 0.03|
| DMSO                | 0.31 ± 0.03|
| adenine propenal    | 1 ± 0.1    |
| NHS–biotin          | n.d.       |

“Dissociation constants ($K_d$) were determined by fitting fluorescence anisotropy data to a simple two-state binding model. $K_d$ values represent the mean ± SD of at least three titrations. The biotin-modification of XPA causes such a large loss of DNA-binding affinity that there is no significant change in fluorescence anisotropy; hence, a $K_d$ value could not be determined.
electrophile reaches the circulation to allow the modification of HSA to serve as a biomarker of exposure.

The impact of electrophiles has long been studied in the context of covalent modification of DNA and its potential to lead directly to mutations. However, adenine propenal produced in the nucleus also has the potential to modify DNA-associated proteins, including those involved in DNA repair. We report here that adenine propenal covalently modifies the NER protein XPA in vitro and that this modification leads to a reduction in XPA’s DNA-binding activity. Oxopropenylation of residues known to be acetylated further suggests that adenine propenal-dependent modification could result in failure of regulation of XPA’s activity through SIRT-1-dependent deacetylation. XPA is a critical component of the NER machinery, without which repair activity ceases. Mutations in XPA, particularly in the DNA-binding domain, are associated with severe XP clinical phenotypes. Thus, it is possible that decreased XPA-binding activity resulting from adenine propenal-dependent modification could result in reduced NER activity in cells and increased potential for mutagenesis and carcinogenesis. It should be noted that, in this model, the modification of repair proteins may have a synergistic effect on mutagenesis by preventing the repair of the original DNA lesion. In this case, even moderate reductions in the activity of the repair proteins have the potential to result in significant increases in genomic instability. Further analysis will be required to determine whether reactive base propenal species modify DNA repair proteins in cells under oxidative stress and to characterize the effects of modification on DNA repair activity in vivo.

**ASSOCIATED CONTENT**

1. Supporting Information
   UV/vis spectra of adenine propenal reactant and products with N-α-acetyllysine; quantification of lysine cross-link formation; MS spectrum of adenine propenal–cysteine adduct; XPA-junction substrate-binding data; COSY spectrum, HMBC, and HSQC of 3 and 4; and sequence coverage of HSA. This material is available free of charge via the Internet at http://pubs.acs.org.

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Funding
This work was supported by research and training grants from the National Institutes of Health [R37 CA087819 (L.J.M.), R01 ES1065561 (W.J.C.), DK020593 (J.A.O.), GM15431 (J.A.O.), and F32 CA159701-01 (S.C.S.)], S.M.S is supported by postdoctoral fellowship 119569-PF-11-271-01-DMC from the American Cancer Society. Partial support for the NMR and MS was provided by P30 ES00267. Funds for the 600 MHz spectrometer was provided from the National Institutes of Health (S10 RR019022), and we thank Dr. Don Stec for providing this service.

Notes
The authors declare no competing financial interest.

**ABBREVIATIONS**

MDA, malondialdehyde; M2dG, 3-(2′-deoxy-β-d-erythro-pentofuranosyl)-pyrimido[1,2-α]-purin-10(3H)-one; NER, nucleotide excision repair; XPA, xeroderma pigmentosum comple-
endogenous M1dG adduct in cellular DNA. J. Biol. Chem. 280, 25377–25382.

(19) Berhane, K., Widersten, M., Engstrom, A., Kozarich, J. W., and Mannervik, B. (1994) Detoxication of base propenals and other alpha, beta-unsaturated aldehyde products of radical reactions and lipid peroxidation by human glutathione transferases. Proc. Natl. Acad. Sci. U.S.A. 91, 1480–1484.

(20) Dalle-Donne, I., Scaloni, A., Giustarini, D., Cavarra, E., Tell, G., Lungarella, G., Colombo, R., Rossi, R., and Milzani, A. (2005) Proteins as biomarkers of oxidative/nitrosative stress in diseases: the contribution of redox proteomics. Mass Spectrom. Rev. 24, 55–99.

(21) Fanali, G., di Masi, A., Trezza, V., Marino, M., Fasano, M., and Ascenzi, P. (2012) Human serum albumin: from bench to bedside. Mol. Aspects Med. 33, 209–290.

(22) Kadiiska, M. B., Gladen, B. C., Baird, D. D., Germolec, D., Graham, L. B., Parker, C. E., Nyska, A., Wachsmann, J. T., Ames, B. N., Basu, S., Brot, N., Fitzgerald, G. A., Floyd, R. A., George, M., Heinecke, J. W., Hatch, G. E., Hensley, K., Lawson, J. A., Marnett, L. J., Morrow, J. D., Murray, D. M., Plastaras, J., Roberts, L. J., II, Rokach, J., Shigenaga, M. K., Sohal, R. S., Sun, J., Tice, R. R., Van Thiel, D. H., Wellner, D., Walter, P. B., Tomer, K. B., Mason, R. P., and Barrett, J. C. (2005) Biomarkers of oxidative stress study II: are oxidation products of lipids, proteins, and DNA markers of CCl4 poisoning? Free Radical Biol. Med. 38, 698–710.

(23) Nagumo, K., Tanaka, M., Chuang, V. T., Setoyama, H., Watanabe, H., Yamada, N., Kubota, K., Tanaka, M., Matsushita, K., Yoshida, A., Jinnouchi, H., Anraku, M., Kadowaki, D., Ishima, Y., Sasaki, Y., Otagiri, H., Yamada, N., Kubota, K., Tanaka, M., Matsushita, K., Yoshida, A., Jinnouchi, H., Anraku, M., Kadowaki, D., Ishima, Y., Sasaki, Y., Otagiri, M., and Maruyama, T. (2014) Cys34-cysteinylated human serum albumin is a sensitive plasma marker in oxidative stress-related chronic diseases. PLoS One 9, e85216.

(24) Shuck, S. C., Rose, K. L., and Marnett, L. J. (2014) Mass spectrometric methods for the analysis of nucleoside-protein cross-links: application to oxoprophenyl-deoxyadenosine. Chem. Res. Toxicol. 27, 136–146.

(25) Fan, W., and Luo, J. (2010) SIRT1 regulates UV-induced DNA repair through deacetylating XPA. Mol. Cell 39, 247–258.

(26) Johnson, F., Pillai, K. M., Grollman, A. P., Tseng, L., and Marnett, L. J., Bienkowski, M. J., Ram, A. P., Tseng, L., and Takeshita, M. (1984) Synthesis and biochemical activity of a new class of cytotoxic agents: N-(3-oxoprop-1-enyl)-substituted pyrimidines and purines. J. Med. Chem. 27, 954–958.

(27) Nair, V., Vietti, D. E., and Cooper, C. S. (1981) Degenerative chemistry of malondialdehyde. Structure, stereochemistry, and kinetics of formation of enaminals from reaction with amino acids. J. Am. Chem. Soc. 103, 3030–3036.

(28) Marnett, L. J., Bienkowski, M. J., Raban, M., and Tuttle, M. A. (1979) Studies of the hydrolysis of 14C-labeled tetraethoxyp propane to malondialdehyde. Anal. Biochem. 99, 458–463.

(29) Bernoud-Hubac, N., Davies, S. S., Boutaud, O., Montine, T. J., and Roberts, L. J., II. (2001) Formation of highly reactive gamma-ketoaldehydes (neuroketals) as products of the neuroprostane pathway. J. Biol. Chem. 276, 30964–30970.

(30) Sugitani, N., Shell, S. M., Soss, S. E., and Chazin, W. J. (2014) Redefining the DNA-binding domain of human XPA. J. Am. Chem. Soc. 136, 10830–10833.

(31) Shao, B., Pennathur, S., Pagani, A., Oda, M. N., Wittzum, J. L., Oram, J. F., and Heinecke, J. W. (2010) Modifying apolipoprotein A-I by a reactive metabolite A-I by malondialdehyde, but not by an array of other reactive carbonyls, blocks cholesterol efflux by the ABCA1 pathway. J. Biol. Chem. 285, 18473–18484.

(32) Shell, S. M., Hawkins, E. K., Tsai, M. S., Hlaing, A. S., Rizzo, C. J., and Chazin, W. J. (2013) Xeroderma pigmentosum complementation group C protein (XPC) serves as a general sensor of damaged DNA. DNA Repair 12, 947–953.

(33) Shell, S. M., Li, Z., Shkriabai, N., Kvaratskhelia, M., Brosey, C., Serrano, M. A., Chazin, W. J., Musich, P. R., and Zou, Y. (2009) Checkpoint kinase ATR promotes nucleotide excision repair of UV-induced DNA damage via physical interaction with xeroderma pigmentosum group A. J. Biol. Chem. 284, 24213–24222.

(34) Galligan, J. Unpublished results.