Carbocations from dibenz[a,j]anthracene and
dibenz[a,h]anthracene, their methylated derivatives, and oxidized
metabolites; a stable ion and DFT study

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Contribution for the commemorative issue in honor of Prof. Ted Sorensen

Abstract
Parent dibenz[a,j]anthracene DB[a,j]A 1 is protonated (in TfOH/SO$_2$ClF) to give a 1:1 mixture of meso-protonated arenium ions 1aH$^+$ and 1bH$^+$. The 7-methyl- DB[a,j]A 2 is protonated (in FSO$_3$H/SO$_2$ClF or TfOH/SO$_2$ClF) at the unsubstituted meso position (C-14) to give 2H$^+$, and the 7,14-dimethyl derivative 3 is ipso-protonated (in FSO$_3$H/SO$_2$ClF) at C-14 to give 3H$^+$. Experimental and/or GIAO-NMR derived $\Delta$δ$^{13}$C values, as well as changes in the computed NPA charges, were used to derive charge delocalization maps for the protonated carbocations derived from parent and methylated DB[a,j]A and DB[a,h]A. DFT and GIAO-DFT were also employed to model the bay-region anti-diol-epoxides (DEs), their derived carbocations, and model covalent adducts. The higher tumorigenic potencies of DB[a,h]A and its methylated derivatives as compared to those of DB[a,j]A are reflected in relative ease of carboxation formation from the DEs. Preference for anti stereochemistry in the covalent adducts derived from DB[a,h]A increases with increasing steric crowding at the bay-region.

Keywords: Dibenz[a,j]anthracene, dibenz[a,h]anthracene, carbocations via protonation, bay-region carbocations, charge delocalization mapping, DFT and GIAO-DFT

Introduction
Polycyclic aromatic hydrocarbons (PAHs) are widespread environmental contaminants that are metabolically activated to form dihydrodiols, that upon further epoxidation, generate diol-epoxides (DEs) as ultimate carcinogens.$^{1-4}$ Reactive diol-epoxides could rapidly intercalate into DNA to form non-covalent complexes, from which benzylic carbocations could form in an acid-
catalyzed ring opening step. The resulting delocalized carbocations are capable of forming covalent adducts with the nucleotides on DNA and RNA. The PAHs that possess bay- and fjord-region(s) are especially potent DNA alkylating agents, and numerous studies have underscored the importance of benzylic carbocations as intermediates in the metabolism of bay- and fjord-region PAHs.

Extensive structure-activity relationship data exist for the benz[a]anthracene (BA) skeleton, and stable ion studies have also been carried out to model the carbocations derived from various substituted BAs. In comparison, limited comparative bioactivity data are available for isomeric dibenzanthracenes (DBAs), namely DB[a,j]A, DB[a,h]A and DB[a,c]A (Fig 1), and their methylated derivatives.

DB[a,j]A is less tumorigenic than DB[a,h]A. It is epoxidized in mammalian liver systems at C-3/C-4 (M-region) as a minor metabolite which is enzymatically hydrated to 3,4-trans-dihydrodiol and further epoxidized to 3,4-diol-1,2-epoxide (see also Fig 2). The trans-DE enantiomers were shown to form covalent adducts with DNA.

**Figure 1.** Isomeric DBA’s.

Formation of the 5,6-dihydrodiol (K-region) has also been reported in the metabolism of DB[a,j]A, as well as other more polar metabolites which were attributed to the bis-dihydrodiols. The 3,4-dihydrodiol is the major metabolite of DB[a,h]A, with the 1,2-dihydrodiol identified as the minor metabolite. The 3,4-dihydrodiol was found be highly tumorigenic (see also Fig 2). The bay-region DE derived from DB[a,h]A formed covalent adducts with DNA. The 7-nitro-DB[a,h]A, with a buttressed nitro group at the meso-position, is also metabolized to the 3,4- and 10,11-dihydrodiols which formed covalent DNA adducts.

As for DB[a,c]A, it is an active tumorigen that formed the 10,11-, 3,4- and the 1,2-dihydrodiols in metabolic studies. Several studies have dealt with the synthesis of these dihydrodiols.

Limited comparative mouse skin assays are available on the methylated DBAs. Whereas introduction of methyl group(s) into the meso-position(s) at C-7 and C-14 increased tumorigenicity in the case of DB[a,h]A and DB[a,j]A (see Fig 2), DB[a,c]A and its 9,14-dimethyl-derivative were both found to be weakly active as tumor initiator. The X-ray
structure of 7,14-dimethyl-DB[a,j]A has been determined, showing that methyl introduction at C-7/C-14 causes severe skeletal distortion.\textsuperscript{19}

As part of broader study on stable ion study of large methylene-bridged PAHs, low temperature protonation of DB[a,c]A and DB[a,h]A were included for comparison.\textsuperscript{18} In agreement with bromination and nitration, in both cases protonation occurred at the meso-region.\textsuperscript{20} In the present study we have used a combined experimental NMR and GIAO-DFT study to examine the geometries, relative energies, charge delocalization modes and relative aromaticity in model ring-protonated carbocations derived DB[a,h]A and DB[a,j]A and their methylated derivatives (compound 1-4 and compounds 9, 11, and 14 in Fig 2). We have also used DFT to examine the corresponding anti-DEs, their ring opening to benzylic carbocations, and covalent adduct formation (quenching) with a model nucleophile.
Figure 2. Dibenz [a,j]anthracenes and dibenz[a,h]anthracenes, their bay-region dihydrodiols and diol-epoxides metabolites, benzylic carbocations via bay-region epoxide ring opening, and model ring protonated carbocations.
Results and Discussion

Protonation study of dibenz[a,j]anthracene (1) and its methylated derivatives (2-4) (Scheme 1; Charts 1-3 and S1; Tables S1-S3; Figures S1-S4).

The $^1$H and $^{13}$C NMR assignments for the neutral dibenzanthracenes 1, 2, and 3 are gathered in Chart S1.

Low temperature reaction of parent DB[a,j]A 1 with FSO$_3$H/SO$_2$ClF gave rather broad NMR signals, likely due to competing oxidation to form the radical cation. Protonation with the less oxidizing CF$_3$SO$_3$H/SO$_2$ClF produced improved NMR spectra, with less signal-broadening (Figure S1), which indicated the formation of carbocations 1$aH^+$ and 1$bH^+$ in 1:1 ratio. Based on DFT (Table S1) 1$bH^+$ is more stable than 1$aH^+$ by 1.8 kcal/mol.

Protonation of 2 in CF$_3$SO$_3$H or in FSO$_3$H/SO$_2$ClF gave carbocation 2$aH^+$ (>95% by NMR) by protonation at C-14 (see Figure S2). DFT computes 2$bH^+$ to be only 1.6 kcal/mol less stable (Table S1).

Protonation of the carcinogenic dimethyl-derivative 3 in FSO$_3$H/SO$_2$ClF gave the carbocation 3$aH^+$ by ipso-attack at C-14 (~95% by NMR; see Figures S3 and S4). Anisotropic shielding of the ipso methyl protons in 3$aH^+$ is noteworthy, appearing at 1.61 ppm (the other methyl is at 3.51 ppm). This shielding effect is analogous to the previously reported examples of ipso-protonated methyl- and ethyl-BA carbocations.$^9,^{10}$ Presence of tiny methyl signals at 3.83 and 1.77 ppm appear consistent with minor formation of 3$bH^+$ (<5%).

Relief of steric crowding by protonation at C-14 can be inferred from DFT for compound 4 (Table S1), showing 4$bH^+$ to be 6.3 kcal/mol more stable. Preference for ipso attack at C-14 becomes more pronounced in the dimethyl-derivative 3, where 3$aH^+$ (observed) is computed to be 9.1 kcal/mol more stable.

Table S2 addresses the angles between various rings from the optimized structures for the DBAs and their protonation carbocations. Whereas parent 1 and its carbocations are planar, methylation at C-7 induces slight deviations (up to 14°). The situation changes in the C-7/C-14 dimethyl-derivative 3 and its carbocations, where the structure becomes significantly more distorted out of planarity (for example 16° angle between AC and CE rings in 3). The correspondence between the reported X-ray structure$^20$ and the computed gas phase structure seems poor (Table S3)! The ipso-protonated carbocations 3$aH^+$/3$bH^+$ remain buckled. Parent 9 is planar but its meso-methylated derivatives and their carbocations are severely distorted, with 14 showing an AE angle of 40°! The degree of bucking diminishes upon ipso-protonation (as in 14$H^+$).
**Scheme 1.** Protonation of dibenz[a,j]anthracenes.

Charts S2-S3 address the relative aromaticity in DBAs 1-4 and their carbocations by using NICS(1)zz.21 There are minor variations in relative aromaticity in the angular rings, with methyl introduction, as distortions forces them out of planarity, so that angular rings become slightly more aromatic. Although NICS(1)zz implies that the meso-protonated center rings are no longer aromatic, observation of anisotropic shielding of the meso-methyl protons argues that strong ring current effects still operate and the systems are therefore overall aromatic.

The experimental and GIAO-NMR data for the resulting carbocations (at the 6-31G(d) and 6-311+G(d,p) levels), along with their NPA-derived charge delocalization maps are gathered in Charts 1-4 for comparison (for relative energy data see Table S1).

NMR chemical shifts computed at the augmented basis-set are typically circa 10-15 ppm more deshielded and appear to correspond more closely with the available experimental $^{13}$C NMR values. Charts 5a and 5b summarize the GIAO-NMR data for the carbocations derived from meso-protonation of DB[a,h]A which were computed at two different basis-sets for comparison (no parallel experimental NMR data could be obtained on this set due to lack of availability of the authentic compounds). The charge delocalization maps derived from $\Delta\delta^{13}$C values correspond closely with those derived based on changes in the NPA-charges (cation minus neutral).

A major difference between the overall charge delocalization modes in the DBA carbocations, as revealed from this work, and the previously studied alkyl-BA carbocations 9,10 is that in the BA carbocations a more distinct anthracenium ion character exists, whereas with DBA
carbocations the delocalization path is less extended and less homogeneous. This could stem from increased twisting of the framework in the DBA carbocations.

**DFT Study of the benzylic carbocations via epoxide ring opening (Figure 2, Chart 6; Table S2).**

In the context of the present stable ion study and for comparison, the bay-region DEs 5-8 derived from DB[a,j]A compounds 1-4 and the bay-region DEs (10, 12, 13 and 15) derived from DB[a,h]A (compounds 9, 11, and 14), and their derived carbocations (see further) were computed by DFT. Using a model methanol quenching experiment, relative energies of the syn and anti methyl ethers were also computed for comparison. Table S4 summarizes the relative energy data.

The NPA-derived charge delocalization maps for the benzylic carbocations 5\(^+\) and 10\(^+\) derived from the anti-DEs of DB[a,j]A 1 and DB[a,h]A 9 were also determined and are sketched in Chart 6, showing that the positive charge is extensively delocalized into the anthracene moiety. The 6/7 positions in 5\(^+\) and 5/13/14 positions in 10\(^+\) (using numbering system in Fig 1) are sites that are potentially susceptible to attack by nucleophile!

Effect of steric congestion in the bay-region on the relative ease of carbocation formation can be deduced by comparing the relative energies of isomeric carbocations 7\(^+\) and 6\(^+\) (formation of 7\(^+\) being more favorable by 1.7 kcal/mol). In the case of 5\(^+\), the methoxy-adduct with anti stereochemistry is favored by 1.5 kcal/mol. This preference almost vanishes in the case of 7\(^+\), for which adducts 7a/7b have nearly the same energies. This is also true in the case of adducts 8a/8b via carbocations 8\(^+\). The bay-region carbocations 10\(^+\), 12\(^+\), 13\(^+\), and 15\(^+\) (via the anti-DEs of DB[a,h]A) are more easily formed as compared to carbocations 5\(^+\)-7\(^+\) (via the anti-DEs of DB[a,j]A). The computed relative preference for carbocation formation via the DEs in DB[a,j]A increases with methyl introduction at C-14 and becomes quite notable with the dimethyl-derivatives (~ 4 kcal/mol) (see Table S4). The preference for anti stereochemistry in the adduct formed via 10 is larger than that via 5, and the same observation is made for 12 versus 6. Interestingly, whereas relative preference for the syn and anti adduct formation is diminished in the case of anti-DE derived from DB[a,h]A by introduction of methyl at C-14, preference for anti-stereochemistry increases for the anti-DE derived from DB[a,h]A (for 7 and 8 the syn/anti energy differences are 0.2 and 0.1 kcal/mol respectively, whereas they are 1.6 and 1.3 kcal/mol respectively for 13 and 15).

The higher tumorigenic potencies of DB[a,h]A and in particular the meso-methylated derivatives relative to the corresponding DB[a,j]A compounds seem to correlate with the present computational data, showing stronger tendencies for carbocation formation from the anti-DEs in the former group.

Interestingly, whereas anti/syn stereoselectivity for covalent adduct formation with methanol (as a model reaction) practically vanishes with increasing steric congestion in DB[a,j]A, anti-selectivity for adduct formation increases with increasing steric crowding for the anti-DEs of DB[a,h]A.
Chart 1. Experimental $^1$H NMR, GIAO-derived $^{13}$C NMR and NPA-derived charges for the carbocations derived from 1 (Δδ$^{13}$C values and NPA change of charges relative to those of the corresponding hydrocarbons in parentheses; a-d denote interchangeable assignments; nd = not determined).
Chart 2. Experimental and GIAO-derived NMR data and computed NPA-derived charges for the carbocations derived from 2 (Δδ^{13}C values and NPA change of charges relative to those of the corresponding hydrocarbons in parentheses).
Chart 3. Experimental and GIAO-derived NMR data and computed NPA-derived charges for the carbocations derived from 3 (Δδ\(^{13}\)C values and NPA change of charges relative to those of the corresponding hydrocarbons in parentheses).
Chart 4. GIAO-derived NMR and computed NPA-derived charges for the carbocations derived from 4 (GIAO $\Delta\delta^{13}$C values and NPA change of charges relative to those of the corresponding hydrocarbons in parentheses).
Chart 5a. GIAO-derived NMR data for the carbocations of dibenz[a,h]anthracenes by B3LYP/6-31G(d) (GIAO $\Delta^13C$ values relative to those of the corresponding hydrocarbons in parentheses).
Chart 5b. GIAO-derived NMR data for the carbocations of diben[\(a,h\)]anthracenes by B3LYP/6-311+G(d,p)//B3LYP/6-31G(d) (GIAO \(\Delta\delta^{13}\)C values relative to those of the corresponding hydrocarbons in parentheses).

Chart 6. NPA-derived charge delocalization map for the benzylic carbocations 5\(^+\) and 10\(^+\) via anti diol-epoxides 5 and 10 by B3LYP/6-31G(d) (changes of charges relative to the corresponding diol-epoxides in parentheses; Arrows identify positions of highest positive charge that could potentially be target for attack by nucleophile).
Experimental Section

General Procedures. The $^1$H and $^{13}$C NMR spectra of the neutral substrates were recorded on a 400 MHz instrument. Low temperature NMR studies of the carbocations were performed on a 500 MHz instrument at the specified temperatures. FSO$_3$H and CF$_3$SO$_3$H were freshly distilled twice in all-glass apparatus under inert atmosphere of nitrogen. SO$_2$ClF was prepared according to a modified procedure of Prakash et al.$^{22}$ Compounds (1)$^{23,24}$ and the methyl-derivatives (2$^{24}$ and 3$^{25}$) were small gift samples, kindly donated by Prof. R. G. Harvey (Univ of Chicago Ben May Institute).

Carbocation generation in FSO$_3$H or CF$_3$SO$_3$H/SO$_2$ClF. The substrate (5-10 mg) was placed in a 5 mm NMR tube. Into the NMR tube, which was cooled in a dry-ice acetone bath (-78 ºC), SO$_2$ClF (0.5 ml) was condensed. FSO$_3$H or CF$_3$SO$_3$H (2-3 drops) was then added under a nitrogen atmosphere at dry ice-acetone temperature. The resulting colored solution was efficiently mixed (vortex mixer). Finally a few drops (2-3 drops) of cold CD$_2$Cl$_2$ were added to the NMR tube. The resulting solution was once again mixed (vortex) prior to NMR study. The color of the studied carbocation solutions were dark-red except for 1 in CF$_3$SO$_3$H/SO$_2$ClF (dark-blue).

Quenching experiments. Cold solutions of the carbocations in NMR tubes were poured into ice-sodium bicarbonate. The resulting mixture was extracted with CH$_2$Cl$_2$ and dried (MgSO$_4$). After evaporation of the solvent, the residue was analyzed by $^1$H NMR, which confirmed the recovery of skeletally intact PAHs in all cases.

Computational protocols. Structures were optimized using a C$_1$ molecular point group by the density function theory (DFT) method at B3LYP/6-31G(d) level using the Gaussian 03 package.$^{26,27}$ All computed geometries were verified by frequency calculations to have no imaginary frequencies. Energies are summarized in Tables S1 and S4. Natural population analysis (NPA)-derived charges were computed at the same level. NMR chemical shifts were calculated by the GIAO$^{28}$ method at the B3LYP/6-311+G(d,p)/B3LYP/6-31G(d) and B3LYP/6-31G(d)/B3LYP/6-31G(d) level. NMR chemical shifts were referenced to TMS (GIAO magnetic shielding tensor = 184.1 and 189.8 ppm in TMS; these values are related to the GIAO isotropic magnetic susceptibility for $^{13}$C), calculated with molecular symmetry of T$_d$ at the same level of theory. Nuclear independent chemical shifts (NICS)$^{29}$ were calculated by the GIAO method by B3LYP/6-311+G(d,p)/B3LYP/6-31G(d). The "out-of-plane" components of the NICS tensors were calculated at 1 Å above the ring centers by rotation the planes of the rings orthogonal to $z$-axis, which denotes as NICS(1)$_{zz}$. The ring centers were defined as the simple average of Cartesian coordinates for all the carbons in the ring. Planes of the rings were calculated by least-square methods using coordinates of the all carbons in the rings.
Supplementary Information Available

Chart S1 (NMR assignments for 1-3); Charts S2-S3 (NICS data); Tables S1 and S4 (energy data); Tables S2-S3 (angles between rings); Figures S1-S4 (selected NMR spectra for the carbocations).

Acknowledgements

This work was partially supported by the NCI of NIH (2R15CA078235-02A1).

References


