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Atmospheric Chemistry of Gas-phase Polycyclic Aromatic Hydrocarbons: Formation of Atmospheric Mutagens

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The atmospheric chemistry of the 2- to 4-ring polycyclic aromatic hydrocarbons (PAH), which exist mainly in the gas phase in the atmosphere, is discussed. The dominant loss process for the gas-phase PAH is by reaction with the hydroxyl radical, resulting in calculated lifetimes in the atmosphere of generally less than one day. The hydroxyl (OH) radical-initiated reactions and nitrate (NO$_3$) radical-initiated reactions often lead to the formation of mutagenic nitro-PAH and other nitro-polyaromatic compounds, including nitrobenzopyranones. These atmospheric reactions have a significant effect on ambient mutagenic activity, indicating that health risk assessments of combustion emissions should include atmospheric transformation products. — Environ Health Perspect 102(Suppl 4):117–126 (1994).

Key words: polycyclic aromatic hydrocarbons, nitro-polycyclic aromatic hydrocarbons, gas-phase reactions, mutagens, hydroxyl radical, nitrate radical

Introduction

Polycyclic aromatic hydrocarbons (PAH) and certain nitro-polycyclic aromatic hydrocarbons (nitro-PAH) are emitted into the atmosphere from combustion sources (1,2). For many years, research concerning the health implications of PAH emissions has been conducted (3), and more recently the nitro-PAH have been studied (2,4–6). Ten years ago a warning was given that the health effects of not only the directly emitted carcinogenic and mutagenic PAH and nitro-PAH, but also of their atmospheric transformation products, need to be assessed (7,8).

The last decade has brought a gradual realization that the gas-phase atmospheric chemistry of the PAH has a significant impact on the mutagenic activity of ambient atmospheres, both for vapor phase and particle-associated mutagens. The majority of ambient nitro-PAH now are thought to be formed in the atmosphere from the gas-phase reactions of the PAH with four rings or less (9–14). Atmospheric reactions generally produce products of increased polarity (15–18). Recently this has been shown to account for the trend of increased polarity seen in the direct-acting mutagenicity of ambient particles in comparison with, for example, diesel particles (19,20). Based largely on work conducted at the Statewide Air Pollution Research Center, University of California, Riverside, over the past 10 years, our current knowledge of the atmospheric reactions and lifetimes of the gas-phase PAH, their formation of mutagenic products, and the contributions of these products to the mutagenic activity of ambient atmospheres will be discussed.

Phase Distribution of PAH and PAH-derivatives in the Atmosphere

The PAH, nitro-PAH, and other polycyclic aromatic compounds (PAC) present in the atmosphere are distributed between the gas and particle phases, with this gas- or particle-phase distribution depending mainly on the liquid-phase vapor pressure of the PAH or PAC at the temperature of the ambient air parcel containing them (21). As discussed by Bidleman (21) and Pankow and Bidleman (22), organic compounds with liquid-phase vapor pressures greater than 10$^{-6}$ Torr at the ambient air temperature will exist, at least partially, in the gas phase in the atmosphere. The subcooled liquid vapor pressures of the 2- to 4-ring PAH are greater than or equal to 10$^{-6}$ torr at 298 K, and ambient air measurements (23–28) have shown that the 2- to 4-ring PAH, as well as the 2-ring nitro-PAH, are largely gas-phase species.

Table 1 shows the measured ambient air concentrations of a series of PAH and nitro-PAH collected on Teflon-coated glass fiber filters compared with the total ambient air concentrations from collections on filters, polycarbonate foam, and Tenax solid adsorbents (29). These and other data (24,26,28) show that the 4-ring PAH fluoranthene and pyrene are mainly (greater than or equal to 90%) adsorbed to the filters and not collected on the polycarbonate foam adsorbent located downstream from the filters (30).

Laboratory Studies of Atmospheric Reactions of PAH and Formation of Nitro-PAH

As for other classes of organic compounds, the gas-phase PAH and nitro-PAH can undergo wet and dry deposition (16,21), photolysis, and gas-phase reactions with OH radicals, NO$_3$ radicals, and O$_3$ (16,29–32). For the gas-phase PAH, dry deposition is
Table 1. Particle-associated (filter) and total atmospheric (sum of filter and solid adsorbent) concentrations of polycyclic aromatic hydrocarbons and nitro-polycyclic aromatic hydrocarbons in a daytime sample collected at El Camino Community College, Torrance, CA, February 25, 1986, 6:00 A.M. to 6:00 P.M. (24).

| Compound          | Molecular weight | Filter | Σ (Filter + solid adsorbent) |
|-------------------|------------------|--------|-----------------------------|
| PAH               |                  |        |                             |
| Naphthalene       | 128              | 0.45   | 3300                        |
| Phenanthrene      | 178              | 0.33   | 78                          |
| Anthracene        | 178              | 0.03   | 6.1                         |
| Fluoranthene      | 202              | 0.47   | 8.0                         |
| Pyrene            | 202              | 0.80   | 8.0                         |
| Benzenopyrene     | 252              | 0.59   | 0.6                         |
| Perylene          | 252              | 0.16   | 0.2                         |
| Nitro-PAH         |                  |        |                             |
| 1-Nitronaphthalene| 173              | 0.05   | 3.0                         |
| 2-Nitronaphthalene| 173              | 0.006  | 2.9                         |
| 3-Nitrobiphenyl   | 199              | 0.03   | 6.0                         |
| 9-Nitroanthracene | 223              | 0.05   | 0.05                        |
| 2-Nitrofluoranthene| 247              | 0.28   | 0.3                         |
| 1-Nitropyrene     | 247              | 0.04   | 0.04                        |
| 2-Nitropyrene     | 247              | 0.04   | 0.04                        |

PAH, polycyclic aromatic hydrocarbon.

Table 2. Room temperature rate constants, k, for the gas-phase reactions of OH radicals, NO₃ radicals, and O₃ with PAH and nitro-PAH.

| k (cm³ molecule⁻¹ s⁻¹) for reaction with | OH⁵ | NO₃⁶ | O₃⁷ |
|------------------------------------------|-----|------|-----|
| PAH                                      |     |      |     |
| Naphthalene                              | 2.16 x 10⁻¹¹ | 3.6 x 10⁻¹² | <2 x 10⁻¹⁹ |
| 1-Methylnaphthalene                      | 5.3 x 10⁻¹¹ | 7.7 x 10⁻¹² | <1.3 x 10⁻¹⁹ |
| 2-Methylnaphthalene                      | 5.2 x 10⁻¹¹ | 1.08 x 10⁻¹² | <4 x 10⁻¹⁹ |
| Biphenyl                                 | 7.2 x 10⁻¹² | <5 x 10⁻¹⁰ | <2 x 10⁻¹⁹ |
| 2,3-Dimethylnaphthalene                  | 7.7 x 10⁻¹² | 1.55 x 10⁻¹² | <4 x 10⁻¹⁹ |
| Acenaphthene                             | 1.0 x 10⁻¹⁰ | (4.6 x 10⁻¹³ + 1.7 x 10⁻²⁷ [NO₂]) | <5 x 10⁻¹⁹ |
| Acenaphthylene                           | 1.1 x 10⁻¹⁰ | 5.5 x 10⁻¹² | 5.5 x 10⁻¹⁶ |
| Fluorene                                 | 1.3 x 10⁻¹¹ |          |     |
| Phenanthrene                             | 3.1 x 10⁻¹¹ |          |     |
| Anthracene                               | 1.3 x 10⁻¹⁰ |          |     |
| Fluoranthene                             |          | 5.1 x 10⁻²⁰ [NO₂] |     |
| Pyrene                                   |          | 1.6 x 10⁻²⁷ [NO₂] |     |
| Nitro-PAH                                |          |          |     |
| 1-Nitronaphthalene                       | 5.4 x 10⁻¹² | 3.0 x 10⁻¹⁰ | <6 x 10⁻¹⁹ |
| 2-Nitronaphthalene                       | 5.6 x 10⁻¹² | 2.7 x 10⁻¹⁰ | <6 x 10⁻¹⁹ |
| 2-Methyl-1-nitronaphthalene              | 8.3 x 10⁻¹² | {1.1 x 10⁻¹⁴ + 2.7 x 10⁻²⁹ [NO₂]} | <3 x 10⁻¹⁹ |

PAH, polycyclic aromatic hydrocarbon.⁵ Taken from Atkinson (29).⁶ Taken from Atkinson (31); NO₃ concentrations in molecule cm⁻³ units.⁷ Taken from Atkinson (32) except as indicated.⁸ Taken from Atkinson et al. (44). At 298 K, estimated from data obtained at 325 K.

expected to be of little importance (33), and based on the wash-out ratios measured by Ligocki et al. (34), wet deposition is expected to be of minor importance as an atmospheric loss process for gas-phase PAH. After the origin of each reactive species is noted, the individual gas-phase reaction processes are discussed below.

Ozone concentrations in the unpolluted troposphere are typically about 7 x 10¹¹ molecule cm⁻³ (30 ppb mixing ratio at ground level) (35). The photolysis of O₃ in the troposphere to yield the O(¹D) atom leads to the formation of the OH radical:

O₃ + hv (λ<320 nm) → O₂(¹Δg) + O(¹D)

O(¹D) + H₂O → 2 OH
O(¹D) + M → O(³P) + M
(M = N₂, O₂, and CO₂)

From consideration of the estimated emissions of CH₃CCl₃ into the atmosphere, the atmospheric concentrations of CH₃CCl₃, and a knowledge of the rate constant for reaction of CH₃CCl₃ with the OH radical (the major tropospheric loss process for CH₃CCl₃), Prinn et al. have obtained an annually, seasonally and diurnally, averaged global tropospheric OH radical concentration of 8 x 10¹³ molecule cm⁻³ (36).

The NO₃ radical is formed from the sequence of reactions

NO + O₃ → NO₂ + O₂
NO₂ + O₃ → NO₃ + O₂
NO₃ + NO → 2 NO₂

NO₃ + hv → NO₃ + O(³P)

Because of the rapid photolysis of the NO₃ radical (with a photolysis lifetime at solar noon of approximately five sec) and the rapid reactions of NO with O₃ and of the NO₂ radical with NO (37), NO₃ radical concentrations are low during daylight hours. In the presence of NO₂ and O₃, NO₃ radical concentrations generally increase over continental areas after sunset (38,39). Based on the available data, Atkinson (31) suggested an average NO₃ radical concentration in the lower troposphere during nighttime hours of approximately 5 x 10¹⁰ molecule cm⁻³ (approximately 20 ppt) over continental areas. Over marine areas, NO₃ radical concentrations are lower [approximately 0.25 ppt at 3 km altitude near Hawaii (40)], as expected because of the low NO₂ concentrations (40,41).

The reactions of the 2- to 4-ring PAH and 2-ring nitro-PAH that may be important in the atmosphere have been studied experimentally under laboratory conditions (9,11–14,19,20,42–59). The kinetic and product data from these studies, combined with our understanding of the atmospheric chemistry of other classes of organic compounds (30,32), allow a reasonably consistent, though still incomplete, understanding of the atmospheric chemistry of the 2- to 4-ring PAH and 2-ring nitro-PAH. The individual reaction processes are discussed below.

Reactions of Gas-phase PAH and Nitro-PAH with the OH Radical

The rate constants for the gas-phase reactions of the OH radical with PAH and nitro-PAH are given in Table 2. Only for naphthalene, biphenyl, and phenanthrene have studies been conducted by more than one research group, and the rate constants given in Table 2 for these three PAH are the recommended values of Atkinson (29). As discussed by Atkinson (29), the OH radical reactions with the PAH and PAH-derivatives proceed by two reaction pathways: OH radical addition to the aromatic ring to form an initially
energy-rich hydroxycycloheptadienyl-type radical and OH radical interaction with the substituent groups, either through H-atom abstraction from C-H bonds or OH radical addition to >C = C< bonds. For example, for acenaphthene, the reactions are

![Diagram of OH + CH2=CH2 reaction](attachment:image)

and for acenaphthylene the reactions are

![Diagram of OH + CH=CH reaction](attachment:image)

For the alkyl-substituted PAH such as 1- and 2-methylnaphthalene, 2,3-dimethyl-naphthalene, and (probably) acenaphthene, the reaction pathway involving OH radical addition to the aromatic ring dominates under atmospheric conditions (29). For those PAH containing unsaturated cyclopenta-fused rings (acenaphthene and acenaphthenylene), OH radical addition to the cyclopenta-fused ring >C = C< bond may be significant.

The products of these OH radical-initiated reactions are not well understood. The observed products of the OH radical-initiated reactions (in the presence of NOx) of naphthalene and biphenyl are hydroxy- and nitro-PAH (52). The yields of the naphthols are 7 and 4% for 1- and 2-naphthol, respectively, significantly higher than the 1- and 2-nitronaphthalene yields of approximately 0.3% each (52). Similarly, the yield of 2-hydroxybiphenyl from biphenyl is 20% (also with much lower amounts of 3- and 4-hydroxybiphenyl being produced), while the single nitro-derivative is observed is 3-nitrobiphenyl in approximately 5% yield (52).

The specific nitro-PAH isomers formed from the gas-phase OH radical-initiated reactions of naphthalene (52), 1- and 2-methylnaphthalene (11), biphenyl (12,52),acenaphthene (12), acenaphthylene (12), fluorene (58), phenanthrene (12), anthracene (12), acenaphthenylene (55), fluoranthene (9,14), and pyrene (9,14), as well as their product yields, are given in Table 3. It should be noted that the nitrofluoranthenes and nitropyrenes formed from the gas-phase reactions of fluoranthene and pyrene have sufficiently low vapor pressures that they condense onto particles in the atmosphere, and at least for the 4-ring PAH, particle-phase nitro-PAH are formed from gas-phase PAH precursors.

The available product data for the mono-cyclic aromatic hydrocarbons and biphenyl (52,60,61) indicate that the nitroarene product yields do not extrapolate to zero at low NOx concentrations and that the nitroarene formation yields determined under laboratory conditions (Table 3) may be applicable to ambient atmospheric conditions (52,60,61). The nitroarene product formation yields are low in all cases, ranging from less than or equal to 0.2 to 5%, and as noted above, the hydroxy-PAH yields for naphthalene and biphenyl are a factor of approximately 5 to 10 higher than the nitro-PAH yields. It is important to note that the majority of the OH radical-initiated reaction products of the PAH remain unidentified. While there are uncertainties about the reaction mechanisms, a recently postulated mechanism (60) that is consistent with our product data (9,14) is shown below for the reaction of the OH radical with fluoranthene in the presence of NOx.

![Diagram of OH + C18H12 reaction](attachment:image)

The nitroarenes formed from the OH radical-initiated reactions of the PAH (Table 3) are often isomers distinct from
Reactions of Gas-phase PAH and Nitro-PAH with the NO₃ Radical

Naphthalene and the alkyl-substituted naphthalenes are observed to react in N₂O₅−NO₃−NO₂-air mixtures, in which NO₃ radicals are generated by the thermal decomposition of N₂O₅:

\[ N₂O₅ \rightarrow NO₃ + NO₂. \]

The disappearance rates of the naphthalenes relative to those of alkenes such as propene and trans-2-butene in these reaction mixtures as a function of the NO₂ concentration indicate that the PAH-loss processes are kinetically equivalent to reaction with N₂O₅ (48,52−54,57). The experimental data (57) show that the reaction of naphthalene in N₂O₅−NO₃−NO₂ air mixtures occurs by the initial addition of the NO₃ radical to the aromatic rings to form a nitratocyclohexadienyl-type radical, which then either decomposes back to reactants or reacts exclusively with NO₂.

\[ \text{products, including nitrates} \]

The measured rate constant \( k_{obs} \) for reaction with the NO₃ radical is

\[ k_{obs} = \left\{ \frac{(-d[PAH])/dt}{[PAH][NO₃]} \right\} = k_a k_b [NO₂]/k_b. \]  

[1]

For those PAH containing substituent groups, a parallel reaction pathway involving NO₂ radical reaction with the substituent group(s) also can occur (12,54) in addition to NO₃ radical addition to the aromatic ring. For example, the reaction for acenaphthene is

\[ \text{and for acenaphthene, NO₃ radical addition to the cyclopenta-fused } C=C \text{ bond is the dominant reaction pathway (54)} \]

and for acenaphthene, 1-nitronaphthalene, 2-nitronaphthalene, and 2-methyl-1-nitronaphthalene are (1.66 ± 0.13) \times 10^{-4} \text{ sec}^{-1}, (1.28 ± 0.10) \times 10^{-4} \text{ sec}^{-1}, and (1.3 ± 0.4) \times 10^{-4} \text{ sec}^{-1}, respectively (13). Nitro-PAH that are particle-associated under atmospheric conditions may be fully or partially protected from photolysis (13,67−70).

Calculated Atmospheric Lifetimes of Gas-phase PAH and Nitro-PAH

The photolysis and reaction rate data given above can be combined with the ambient radiation flux and the ambient concentrations of OH and NO₃ radicals, NO₂ and O₃ to allow the estimation of the lifetimes of the PAH and nitro-PAH with respect to each of these tropospheric loss processes. These calculated lifetimes are given in Table 4. For the PAH not containing cyclopenta-fused rings, the dominant tropospheric loss process is by reaction with the OH radical, with calculated lifetimes of one day or less (note that OH radical reaction only occurs during daylight hours). For the PAH containing cyclopenta-fused rings such as acenaphthene and acenaphthylene react with NO₃ radicals at a significant rate. The reaction pathway involving NO₂ radical addition to the fused rings of the PAH is not a significant tropospheric loss process for any of the gas-phase PAH. PAH having unsaturated cyclopenta-fused rings, such as acenaphthylene,acenaphthylene, and cyclopenta[cd]pyrene, react, or are expected to react, with O₃ at a significant rate.

In contrast to O₃ and the OH radical, which are ubiquitous at reasonably consistent (on a day-to-day level) ambient concentrations (28,35,36), the ambient concentrations of the NO₃ radical in the lower troposphere over continental areas exhibit large variations, with the mixing ratios ranging from less than 2 to 430 ppt (71). The ambient tropospheric concentration of the NO₃ radical at any given time (during nighttime) and place must be viewed as uncertain by a factor of at least 10. A good approximation is that the dominant tropospheric removal process for the PAH is by daytime reaction with the OH radical, leading to lifetimes of approximately 8 hr or less.

As seen from the rate-constant data given in Table 2 and the calculated lifetimes in Table 4, the presence of the nitro substituent group in the nitroarenes leads to a marked decrease in their reactivity toward the OH radical. To date, kinetic and product studies have been carried out only for three gas-phase, fused-ring nitroarenes (13,56), and photolysis will be the dominant tropospheric
removal process for these compounds, with calculated lifetimes of approximately 2 hr.

**Evidence from Ambient Data for Transformations of Gas-phase PAH and Mutagen Formation**

The recent ambient air measurement study of Arey et al. (28) provided clear evidence for the reactions of the volatile PAH with the OH radical, with the nighttime/daytime concentration ratios exhibiting a linear correlation with the OH radical reaction rate constant (Figure 1). From an estimate of the nighttime dilution rate provided by the daytime/nighttime ratio of 3-nitrobenzyl [a nitro-PAH believed to be formed only in the atmosphere from the daytime reaction of benzyl with the OH radical in the presence of NO₃ (52)], an average 12-hr daytime OH radical concentration of 2.2 × 10^6 molecule cm⁻³ (during August) was derived, uncertain to at least a factor of 2 (28). This estimated OH radical concentration in an urban area is similar to the annually averaged global tropospheric 12-hr daytime OH radical concentration of 1.6 × 10^6 molecule cm⁻³ (36) and provides very strong evidence that the gas-phase PAH do react in the troposphere.

Furthermore, the specific isomers of the nitro-PAH and nitro-PAC observed in ambient air suggest that they are formed in the atmosphere through the gas-phase reactions of the 2- to 4-ring PAH (9,13,19,24–26,28,49,52,55,58,59,72–78). Thus, ambient air contains nitro-PAH isomers distinct from the PAH electrophilic nitration products reported in direct emissions. The nitro-PAH isomers not formed from electrophilic nitrations are observed, however, in laboratory simulations of the atmospheric reactions of the PAH, providing strong evidence for atmospheric formation of nitro-PAH. For example, Figure 2 shows a combined gas chromatography-mass spectrometry single ion trace (GC-MS SIM) for the m/z 247 nitro-PAH (nitrofluoranthenes, nitropyrenes, and nitroacenaphthenynes) present in an extract of a diesel exhaust particle sample and in an extract of an ambient air sample collected on filters. Figure 2 clearly shows that the ambient air particle sample contains several other nitrofluoranthenes and nitropyrenes in addition to the 1-nitropyrene expected to be a direct emission (as shown by the GC-MS SIM trace for the diesel exhaust particle sample in Figure 2). Furthermore, the additional nitrofluoranthenes (in particular the 2-nitrofluoranthene) and nitropyrenes are precisely those nitrofluoranthenes and nitropyrenes formed from the gas-phase OH radical-initiated reactions of fluoranthene and pyrene. [A further small peak on the GC-MS trace that elutes between 8-nitrofluoranthene and 4-nitropyrene is a nitroacenaphthenylene formed from the OH radical-initiated reaction of acephenanthrylene (55).]

We have observed that the 2-nitrofluoranthene concentration in ambient air samples consistently is higher than the directly emitted 1-nitropyrene concentration (9,10,24,26), showing the importance of atmospheric transformations of the 4-ring PAH with respect

| PAH | OH lifetime | NO₃ lifetime | O₃ lifetime | Photolysis lifetime |
|-----|-------------|--------------|-------------|-------------------|
| Naphthalene | 8.0 hr | 1.5 years | > 80 days | |
| 1-Methyl naphthalene | 3.3 hr | 250 days | > 125 days | |
| 2-Methyl naphthalene | 3.3 hr | 180 days | > 40 days | |
| 2,3-Dimethyl naphthalene | 2.3 hr | 125 days | > 40 days | |
| Biphenyl | 2.0 days | > 105 years | > 80 days | |
| Acenaphthene | 1.7 hr | 1.2 hr | > 30 days | |
| Acenaphthylene | 1.6 hr | 6 min | > 43 min | |
| Fluorene | 1.1 days | | | |
| Phenanthrene | 5.6 hr | | | |
| Anthracene | 1.3 hr | | | |
| Fluoranthene | = 3.5 hr | = 85 days | | |
| Pyrene | = 3.5 hr | = 30 days | | |
| 1-Nitronaphthalene | 2.7 days | 18 years | > 28 days | 1.7 hr |
| 2-Nitronaphthalene | 2.6 days | 20 years | > 28 days | 2.2 hr |
| 2-Methyl-1-nitronaphthalene | > 1.7 days | 4.2 days | > 55 days | 2.1 hr |

*a* For a 12-hr average OH radical concentration of 1.6 × 10^6 molecule cm⁻³ (36). *b* For a 12-hr average nighttime NO₃ radical concentration of 5 × 10^6 molecule cm⁻³ (31) and an NO₂ concentration of 2.4 × 10^11 molecule cm⁻³. *c* For a 24-hr average O₃ concentration of 7 × 10^11 molecule cm⁻³ (35). *d* For an average 12-hr daytime NO₂ photolysis rate of J(NO₂) = 5.2 × 10⁻³ sec⁻¹. *e* Using estimated OH radical reaction rate constant of 5 × 10⁻¹¹ cm³ molecule⁻¹ sec⁻¹ based on rate constant correlation with ionization potential (48).

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**Figure 1.** Plot of the average nighttime/average daytime concentrations of gas-phase PAH against their OH radical reaction rate constants. Data from Glendora, California, during August 1986 (29).
Figure 2. GC-MS SIM traces for the molecular ions of the nitrofluoranthenes (NF) and nitropyrenes (NP) in extracts from diesel particles and ambient particles (24) collected on filters in Torrance, California.

Table 5. Comparison of measured and calculated ambient nitroarene concentrations at Glendora, California, in August 1989 (13).

| Nitroarene          | Measured | Calculated |
|---------------------|----------|------------|
| 1- + 2-Nitronaphthalene | 4.7      | 8.0        |
| 3-Nitrobenzidine     | 1.0      | 1.2        |
| 2-Nitrofluoranthene  | 0.27     | 0.41       |
| 2-Nitropyrene        | 0.012    | 0.040      |

and the measured or estimated ambient PAH and OH radical concentrations (28), there is strikingly good agreement between the calculated and measured nitroarene concentrations at this site (Table 5). Only the nitronaphthalenes are expected to be present in direct emissions such as diesel exhaust, and the predicted concentrations for the nitronaphthalenes, which were slightly higher than the observed concentrations, suggest that atmospheric formation of these species dominates over their direct emission, at least for this site at the time of the measurements.

**Contribution of PAH Transformation Products to Ambient Direct-acting Mutagenicity**

It has been known for many years that extracts of ambient air particles are carcinogenic (79) and mutagenic (80–87). Using the microsuspension modification of the Ames Salmonella typhimurium assay (88), we have measured (89) the direct-acting (in the absence of microsomal activation) mutagenicity of extracts of ambient air samples collected on Teflon-coated glass fiber filters (particle phase) and polyurethane foam (PUF) plugs (semivolatile vapor phase). Figure 3 shows mutagrams, plots of mutagenic activity against the HPLC fraction number with increasing HPLC fraction number corresponding to increasing polarity, of the vapor-phase and particle-phase extracts. This direct-acting ambient air mutagenicity cannot be due to the PAH themselves, because the PAH require microsomal activation for expression of their mutagenicity. The nitro-PAH are strong, direct-acting mutagens (2) and elute in the HPLC fraction 4 for the HPLC program used by Harger et al. (Figure 3) (89). For the samples collected and tested for mutagenic activity shown in Figure 3, the total direct-acting mutagenicity in the vapor-phase PUF plug sample was actually higher than that in the particle-phase filter sample (210 revertants m⁻³ for the vapor-phase sample versus 160 revertants m⁻³ for the particle-phase sample) (89), showing the potential importance of vapor-phase mutagens in the atmosphere.

While the vapor-phase sample contained approximately 50% of the overall mutagenicity in the nitro-PAH-containing fraction 4, the majority (94%) of the mutagenicity in the particle-phase sample was due to compounds more polar than the nitro-PAH (Figure 3) (89). Mutagrams of particle extracts using the
standard Ames plate incorporation assay also show profiles with more activity in the more polar fractions (17,18,59). From the measured concentrations of nitrofluoranthenes and nitropyrenes in several particular samples and their mutagenic activities (in the standard assay), it was calculated that the nitrofluoranthenes and nitropyrenes contributed less than or equal to 10% of the direct-acting mutagenicity of these extracts (26,91). Furthermore, the direct-acting mutagenicity of a series of ambient air filter samples collected at seven sites in California did not correlate with the PAH concentrations but rather with the 2-nitropyrene concentrations (26). Because 2-nitropyrene is formed in the atmosphere from the OH radical-initiated reaction of gas-phase pyrene, the remainder of the ambient air direct-acting mutagenicity may be associated with the OH radical reaction products of organic compounds and may be due to the mutagenicity of the 2- to 4-ring PAH reaction products other than the nitro-PAH. For example, it is interesting that the nitro-PAH account for 5% or less of the products of the gas-phase OH radical-initiated reactions of the 2- to 4-ring PAH (Table 3) and 10% or less of the ambient air particle phase, direct-acting mutagenicity.

Environmental chamber studies of the gas-phase, OH radical-initiated reactions of naphthalene, fluorene, and phenanthrene have been carried out (19,20,58,59), with 2000- to 4000-L volume gas samples being collected from the chamber for HPLC fractionation with subsequent mutagenicity testing (using the microsuspension modification of the standard plate incorporation assay) and chemical analysis by GC-MS. The mutagrams obtained from these chamber OH radical-initiated reactions of naphthalene, fluorene, and phenanthrene are shown in Figure 3.

For the naphthalene and fluorene reactions, the mutagrams exhibit profiles in which the majority of the activity is in fraction 4, which contains nitro-PAH. Chemical analyses showed the presence in the HPLC fraction 4 of 1- and 2-nitronaphthalene from naphthalene (19) and 1-, 2-, 3-, and 4-nitrofluorene (with 3-nitrofluorene being the dominant isomer) from fluorene (19,58). Use of the mutagenic activities of the nitronaphthalenes and nitrofluorenes (19) showed that the nitronaphthalenes, in particular 2-nitronaphthalene, accounted for approximately 90% of the activity of fraction 4 of the naphthalene reaction products and that the nitrofluorenes, in particular 3-nitrofluorene, accounted for approximately 75% of the activity of fraction 4 of the fluorene reaction products (19). The nitronaphthalenes and nitrofluorenes are present in the atmosphere mainly in the gas phase (24,28,58), and these volatile nitro-PAH contribute to the observed vapor-phase fraction 4 mutagenicity. It is expected that the nitronaphthalenes and methyl-nitronaphthalenes are significant contributors to the observed vapor-phase fraction 4 mutagenicity (89) because 2-nitronaphthalene accounted for approximately 13% of the activity of fraction 4 of the vapor phase sample shown in Figure 3 and the methyl-nitronaphthalenes are abundant in southern California ambient air (11,24,92).

In contrast to the mutagenicity profiles from the naphthalene and fluorene reactions, the majority of the mutagenic activity from the phenanthrene reaction products resides in fraction 6, an HPLC fraction more polar than the nitro-PAH and therefore is generally similar to the particle-phase ambient air mutagenicity profile. Chemical analysis showed the presence of the mutagenic 2-nitro-6H-dibenzo[\(b,d\)]pyran-6-one (Structure I) and 4-nitro-6H-dibenzo[\(b,d\)]pyran-6-one (Structure II) in this mutagenic fraction 6 of the phenanthrene reaction products.

Based on the mutagenic activities of these two nitro dibenzopyranones in the microsuspension assay (19,89), the 2-isomer accounted for all of the mutagenicity in fraction 6 of the phenanthrene reaction products (20,59). Moreover, 2- and 4-nitrodibenzopyranone were observed in both the gas and particle phases (but mainly in the particle phase) in ambient air samples collected in southern California (20,59). The nitrodibenzopyranones were also found in the National Institute of Standards and Technology Standard Reference Material 1649 urban dust collected in Washington, DC (59), as well as in ambient air samples collected in Boise, Idaho, and Philadelphia, Pennsylvania (J. Lewtas and M. G. Nishioka, personal communication).

For four particle and vapor-phase samples on which we have conducted HPLC fractionation with mutagenicity testing of the individual fractions, the 2-nitrodibenzopyranone accounts for essentially all of the mutagenic activity in fraction 6 of both the vapor-phase and particle-phase samples (59). Moreover, the 2-nitrodibenzopyranone accounted for approximately 20% of the total direct-acting mutagenicity in the microsuspension assay of the crude extract of a Riverside, California, ambient air particle sample (20).

In addition to the 2- and 4-nitrodibenzopyranones, seven nitro-PAH lactones (tentatively identified as methyl-nitrodibenzopyranones (molecular weight 255) and nitrophenanthropyranones (molecular weight 265)) have been tentatively identified by GC-MS in an extract from ambient particulate samples collected in Riverside, California (20). Thus,
although the proportion of the ambient activity attributable to the nitro dibenzo pyransones or to any individual compound or class of compound will be dependent upon the assay system used ([19]; J Lewtas, MG Nishioka, personal communication), it is likely that nitro-PAH lactams formed in the atmosphere will prove to be an important class of ambient mutagens.

Note added in proof. Recent kinetic (93) and product (94) studies show that the hydroxycyclohexadienyl radicals formed from OH radical addition to benzene, toluene, and the xylenes react with both O₂ and NO₃, with the O₂ reaction dominating under tropospheric conditions. However, Atkinson et al. (95) have shown that the rate constant for reaction of the NO₂⁺-naphthalene adduct, formed by addition of the NO₂ radical to naphthalene, with NO₃ is >2.5 × 10⁶ higher than that for reaction of the adduct with O₂ at 298 K. The ratio of the rate constants for the corresponding reactions of NO₂ and O₂ with the OH-naphthalene adduct may be expected to be similar to those for the NO₂⁺-naphthalene adduct. The NO₂ reactions with the OH-naphthalene and NO₃⁺-naphthalene adducts may then dominate in urban and rural air masses (95).

The rate constants given in Table 2 for the reactions of phenanthrene and anthracene with the OH radical are supersedes those recently measured by Kwok et al. (96) of 1.3 × 10¹¹ cm³ molecule⁻¹ sec⁻¹ for both phenanthrene and anthracene at 296 ± 2 K. Additionally, Kwok et al. (96) obtained rate constants for the NO₂ radical and O₂ reactions with phenanthrene of [1.2 × 10⁻¹³ + 7.0 ± 10⁻¹⁸ NO₂] cm³ molecule⁻¹ sec⁻¹ and 4 × 10⁻¹⁹ cm³ molecule⁻¹ sec⁻¹, respectively, at 296 ± 2 K.

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