# The use of $NO_y$ , $H_2O_2$ , and $HNO_3$ as indicators for ozone- $NO_x$ -hydrocarbon sensitivity in urban locations

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Abstract. Correlations are presented between model predictions for O<sub>3</sub>-NO<sub>x</sub>-hydrocarbon sensitivity and afternoon concentrations of four "indicator species": NO<sub>y</sub>, O<sub>3</sub>/(NO<sub>y</sub> - NO<sub>x</sub>), HCHO/NO<sub>v</sub>, and H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub>. The indicator species correlations are based on a series of photochemical simulations with varying rates of anthropogenic and biogenic emissions and meteorology. Hydrocarbon-sensitive chemistry in models is shown to be linked to afternoon NO<sub>v</sub> > 20 ppb,  $O_3/(NO_y - NO_x) < 7$ , HCHO/ $NO_y < 0.28$ , and  $H_2O_2/HNO_3 < 0.4$ . Lower  $NO_y$  and higher ratios correspond with NO<sub>x</sub>-sensitive ozone. The correlation between NO<sub>x</sub>-hydrocarbon sensitivity and indicator species remains, even when model emission rates and hydrocarbon/NO<sub>x</sub> ratios are changed by a factor of 2. Methods are developed for evaluating the goodness of fit between model NO<sub>x</sub>-hydrocarbon sensitivity and indicator values. Ozone chemistry is also analyzed in terms of fundamental properties of odd hydrogen, and theoretical criteria for the transition between NO<sub>x</sub>- and hydrocarbon-sensitive regimes are derived. A theoretical correlation between  $O_3$  and  $H_2O_2 + NO_y$  -  $NO_x$  is developed as a way to extend rural  $O_3$ - $NO_y$  correlations into urban locations. Measured indicator values during pollution events in Los Angeles, Atlanta, and rural Virginia are used to illustrate the range of observed values under different environmental conditions.

#### Introduction

The relationship among ozone,  $NO_x$ , and reactive organic gases (ROG) in polluted environments represents a major uncertainty in terms of both science and public policy. It is generally known that for certain conditions the rate of ozone formation will increase with increasing  $NO_x$  and will be insensitive to ROG, while for other conditions the rate of formation will increase with increasing ROG and will be unchanged (or perhaps even decrease) with increasing  $NO_x$ . However, the question of whether peak ozone concentrations in a specific location are sensitive to  $NO_x$  or to ROG has proven elusive.

A major problem for the study of ozone-NO<sub>x</sub>-ROG sensitivity has been the inability to gain evidence based on direct measurements rather than theoretical calculations. Evaluations of ozone-NO<sub>x</sub>-ROG sensitivity have relied on results of threedimensional dynamical/photochemical models such as the Urban Airshed Model (UAM). These evaluations are difficult because they depend on assumptions, e.g., about emission rates, that are highly uncertain [Fujita et al., 1992] and because predicted sensitivity cannot be tested empirically. Model accuracy is assessed by comparing predicted and observed ozone and occasionally other species [Tesche et al., 1990]. Even when the model application appears successful in comparison with measured ozone there may still be considerable doubt about the accuracy of the model prediction for ozone-NO<sub>x</sub>-ROG sensitivity. Because similar quantities of ozone can be produced in ROG-sensitive and NO<sub>x</sub>-sensitive environments, it is possible for a simulation to return accurate

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Paper number 94JD02953. 0148-0227/95/94JD-02953\$05.00 predictions for ozone and still err in its prediction for sensitivity. In recent years, model assessment has been improved by including comparisons with measured concentrations of primary ROG species [Tesche et al., 1990]. Since primary species concentrations are strongly affected by micrometeorology, vertical mixing rates, and on-site sources, there is still room for doubt concerning model sensitivity predictions.

An alternative approach to determining ozone sensitivity is to identify individual species or species ratios that consistently assume different values under conditions of NO<sub>x</sub>-sensitive and ROG-sensitive ozone. If these types of "indicator species" can be identified, then ozone-NO<sub>x</sub>-ROG sensitivity could be determined directly from measurements rather than from models. Comparisons between model predictions and measured values for the indicator species would also provide a test of the accuracy of model sensitivity predictions.

Milford et al. [1994] have established a link between ozone, sensitivity and total reactive nitrogen  $(NO_y=NO_x+HNO_3+peroxyacetylnitrates+$  alkyl nitrates). In the study by Milford et al., low values of afternoon  $NO_y$  (<12 ppb) were consistently associated with  $NO_x$ -sensitive ozone, and high values of  $NO_y$  (>25 ppb) were associated with ROG-sensitive ozone, based on sensitivity predictions from four different model applications to cities and regions in the United States. The potential link between sensitivity and  $NO_x$  demonstrated by Milford et al. [1994] is especially useful because laboratory techniques for measurement of  $NO_y$  have been extensively developed, tested, and applied [Trainer et al., 1993; Fehsenfeld et al., 1987].

The indicator species approach to ozone sensitivity has two significant drawbacks. First, correlations between sensitivity and indicator species (e.g., NO<sub>y</sub>) may shift significantly when changes in initial emission rates are made. This type of shift would suggest that the indicator-based estimates for ozone

sensitivity still depend on model assumptions rather than provide a fully independent estimate. Second, the theoretical results that link ozone sensitivity with an indicator species provide no basis for testing the validity of the theory. Whereas the original determination of sensitivity through photochemical modeling required an essentially untestable faith in the accuracy of model ozone-NO<sub>x</sub>-ROG sensitivity, the use of indicator species such as NO<sub>y</sub> appears to require a similarly untestable faith in the accuracy of the model-based indicator species correlation.

This paper extends the indicator-species concept of Milford et al. [1994] to include three additional empirical indicators for NO<sub>x</sub>-sensitive versus ROG-sensitive ozone: the ratios  $O_3/(NO_y - NO_x)$ , HCHO/ $NO_y$ , and  $H_2O_2/HNO_3$  (or  $H_2O_2/NO_y$ ). Along with NO<sub>y</sub>, these species provide four semi-independent indicators for ozone sensitivity. The HCHO/NO, and H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub> ratios involve species that are more difficult to measure than NO<sub>v</sub>, but measurement techniques exist [Kleindeinst et al., 1988; Hering et al., 1988; Lee et al., 1993, 1994], and they provide a link with ozone sensitivity that relates to model assumptions in a different way than NO<sub>v</sub> does. In addition, the identification of four separate indicators for ozone sensitivity provides for a more rigorous evaluation because the individual indicators must be consistent with one another. This provides for a limited validation of the theory through a comparison of simulated and observed correlations between the species. Sensitivity correlations and correlations between indicator species are shown for photochemical model applications to the Lake Michigan region and the northeast corridor of the United States. [Sillman et al., 1993] with a wide range of initial assumptions. Measurements during air pollution events in Los Angeles, Atlanta, and rural Virginia show that the indicator species assume different values in different photochemical environments.

# **Overview of Relevant Chemistry**

Although the chemistry of urban ozone formation is well known [e.g., National Research Council (NRC), 1991], some aspects of chemistry that relate to ozone sensitivity and the indicator species are not well known and are worth summarizing. The ozone-producing reaction sequence is almost always initiated by the reaction of a hydrocarbon with OH,

(R1) 
$$RH + OH \xrightarrow{[O_2]} RO_2 + H_2O$$

followed by reactions of RO2 and HO2 radicals with NO,

(R2) 
$$RO_2 + NO \xrightarrow{[O_2]} R'CHO + HO_2 + NO_2$$

(R3) 
$$HO_2 + NO \xrightarrow{[O_2]} OH + NO_2$$

(For example, propane  $(C_3H_8)$  reacts via (R1) to form the  $RO_2$  radical  $C_3H_7O_2$  and then via (R2) to form propional dehyde  $(C_2H_5CHO)$ . Most hydrocarbons have more complex reaction pathways, but the general pattern of RH to  $RO_2$  via (R1) and to an intermediate carbonyl via (R2) accounts for most hydrocarbon reactions.)

The conversion of NO to NO<sub>2</sub> results in the production of an ozone following photolysis of NO<sub>2</sub>. Since the radical reactions occur rapidly, the hydrocarbon-OH reaction represents the rate-limiting step for the sequence. The intermediate aldehyde product R'CHO (replaced by a ketone or

dicarbonyl for many hydrocarbon species) may undergo further ozone-producing reactions, initiated either by reaction with OH or by photolysis.

At moderately high  $NO_x$  concentrations ( $NO_x > 0.3$  ppb), reactions (R2) and (R3) represent the dominant reaction pathway for  $RO_2$  and  $HO_2$  radicals, and ozone production rates are roughly proportional to reaction (R1). Consequently, the rate of ozone production is roughly proportional to the summed rate of the hydrocarbon + OH reactions (R1) along with the analogous reaction with CO,

(R4) 
$$CO + OH \xrightarrow{[O_2]} HO_2 + CO_2$$

In most urban environments and in the polluted rural environments of Europe and the eastern United States there is an abundant supply of  $NO_x$ , CO, and hydrocarbons to fuel the ozone-producing reactions. The rate of ozone production and the division into  $NO_x$ -sensitive and ROG-sensitive photochemical regimes is linked to the abundance of the OH radical and the odd hydrogen cycle.

Odd hydrogen in polluted environments is most conveniently viewed as the sum of OH,  $HO_2$ , and  $RO_2$  radicals rather than simply OH +  $HO_2$  [Kleinman, 1986, 1991; Sillman et al., 1990b; Sillman, 1991]. With this definition, odd hydrogen is conserved by reactions (R1)-(R4), which also provide the major pathways for interconversion of OH,  $HO_2$ , and  $RO_2$ . The major sources of odd hydrogen are the photolysis of ozone and subsequent reaction of  $O(^1D)$  with water vapor and the photolysis of aldehydes and other intermediate ROG species. The important sinks of odd hydrogen are

(R5) 
$$HO_2 + HO_2 \rightarrow H_2O_2 + O_2$$

(R6) 
$$RO_2 + HO_2 \rightarrow ROOH + O_2$$

(R7) 
$$OH + NO_2 \rightarrow HNO_3$$

Formation of peroxyacetylnitrate (PAN) is also a significant sink for odd hydrogen in some urban environments.

The division into NO<sub>x</sub>-sensitive and ROG-sensitive photochemical regimes is determined by the relative size of (R5)-(R7) and their role as odd hydrogen sinks. When formation of nitric acid (R7) represents the major sink for odd hydrogen, then the equation of odd hydrogen sources and sinks demonstrates that OH must decrease with increasing NO<sub>x</sub>. OH will also increase slightly with increasing ROG, reflecting the role of the latter as sources of odd hydrogen. The rate of ozoneproducing reactions, being proportional to (R1), increases rapidly with increasing ROG and decreases with increasing NO<sub>x</sub>. Alternatively, when formation of peroxides (reactions (R5) and (R6)) represents the major sink for odd hydrogen, then the concentration of HO<sub>2</sub> radicals is fixed by the size of the odd hydrogen source and is independent of NO<sub>x</sub>. Because the odd hydrogen loss rate is proportional to  $(HO_2)^2$ , the  $HO_2$ concentration also shows little sensitivity to ROG even when hydrocarbons form a significant source of odd hydrogen. The concentration of OH is governed by the interconversion of OH, HO<sub>2</sub>, and RO<sub>2</sub> (reactions (R1)-(R4)). OH increases with increasing NO<sub>x</sub> (due to (R3)) and decreases with increasing ROG (due to (R1)). The rate of ozone production increases with increasing NO<sub>x</sub> but is insensitive to ROG, as increases in ROG coincide with decreases in OH.

Sillman et al. [1990b] have shown that reactions (R1)-(R7) lead to the following equation for OH, assuming that OH, HO<sub>2</sub>, and RO<sub>2</sub> are in photochemical steady state with one another

based on reactions (R1)-(R4) and odd hydrogen sources and sinks are in similar steady state:

$$S_{\rm H} = k_7 [NO_2] [OH] + \{k_5 (\frac{k_4 [CO] + k_1 [RH]}{k_3 [NO]})^2 + k_6 \frac{k_4 [CO] + k_1 [RH]}{k_3 [NO]} \frac{k_1 [RH]}{k_2 [NO]} \} [OH]^2$$
(1)

where  $S_{\rm H}$  represents the source of odd hydrogen from photolysis of ozone and aldehydes. The description of ROG-sensitive and  $NO_x$ -sensitive regimes follows directly from this equation, assuming that production of ozone is proportional to the rate of reaction (R1).

A link between ozone-NO<sub>x</sub>-ROG sensitivity and the ratio H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub> can be deduced from this simplified chemistry because both NO<sub>x</sub>- versus ROG-sensitive ozone and the H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub> ratio are dependent on the relative magnitude of (R5) and (R7). A similar link can be deduced for the ratio of organic peroxides (ROOH) to HNO<sub>3</sub>, associated with the relative magnitude of (R6) and (R7). The well-known dependence of ozone-NO<sub>x</sub>-ROG sensitivity on the ROG/NO<sub>x</sub> ratio can also be deduced from this representation. A complete derivation is shown in the appendix.

Kleinman [1991, 1994] has described a similar transition between "NO<sub>x</sub>-limited" and "radical-limited" photochemical regimes associated with season (summer versus winter) and demonstrated its link with peroxide concentrations. Their results suggest that moderately polluted environments in the eastern United States have NOx-limited chemistry in summer and radical-limited chemistry in winter, associated with high and low peroxide concentrations, respectively. The NO<sub>x</sub>limited and radical-limited regimes of Kleinman [1991] are defined by the relative size of the source of odd hydrogen (S<sub>H</sub>) relative to the NO<sub>x</sub> source (S<sub>N</sub>). The radical-limited regime is associated with conditions during nighttime and winter when OH-driven photochemistry virtually ceases due to lack of sunlight and H<sub>2</sub>O. This is fundamentally different from ROGsensitive urban photochemistry, which includes high OH and significant photochemical production of O<sub>3</sub>, but there are direct parallels between the two. The relation between Kleinman's radical-limited regime and the urban ROG-sensitive regime is discussed further in the appendix.

# **Simulation Methods and Scenarios**

Modeling results are based on simulated air pollution events in the northeast corridor (June 15, 1988) and Lake Michigan regions (August 2, 1988) [Sillman et al., 1993]. Both these events featured O<sub>3</sub> in excess of 150 ppb and meteorology characterized by high temperatures (303°-306°K daytime) and water vapor (mixing ratio of 0.016), light winds, and restricted vertical mixing. Simulation methods are described in detail elsewhere [Sillman et al., 1993]; a summary is given here. The simulations use the chemistry of Lurmann et al. [1986] with updated reaction rates [DeMore et al., 1992], added RO<sub>2</sub> + HO<sub>2</sub> reactions [Jacob and Wofsy, 1988] and chemistry of isoprene and related species based on the work of Paulson and Seinfeld [1992]. Photolysis rates are based on those of Madronich [1987]. An aerosol optical depth of 0.68 was assumed, representing moderately polluted conditions typical of the eastern United States [Flowers et al., 1969]. assumptions in the photolysis rate calculation included clear skies, a total O<sub>3</sub> column of 325 Dobson units (DU), surface albedo of 0.15, and single-scattering albedo of 0.75. Anthropogenic emissions are based on the National Acid Precipitation Assessment Program (NAPAP) 1980 [Environmental Protection Agency (EPA), 1986], and biogenic emissions are derived from data by Lamb et al. [1985] with land use data by Matthews [1983]. Deposition velocities over land were as follows: O<sub>3</sub> and NO<sub>2</sub>, 0.6 cm s<sup>-1</sup>; NO, 0.1 cm s<sup>-1</sup>; HNO<sub>3</sub>, cm s<sup>-1</sup>; PAN, 0.25 cm s<sup>-1</sup>; and H<sub>2</sub>O<sub>2</sub>, 1.0 cm s<sup>-1</sup>. Deposition velocities over water were 0.05 cm s<sup>-1</sup> for all these species.

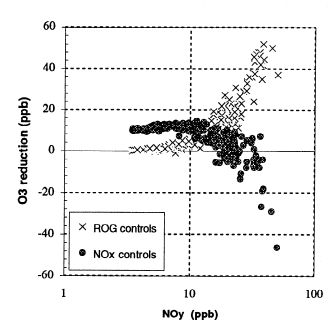
The model includes urban subsections with 20 x 20 km horizontal resolution over domains of approximately 300 x 400 km in combination with simulated regional-scale photochemistry on a spatial domain that includes most of the eastern United States [Sillman et al., 1990a]. A vertical grid structure with variable heights is used to approximate the effect of the ocean and Lake Michigan in suppressing vertical mixing. Results represent average concentrations for the mixed layer. The mixing height during the daytime was derived from vertical temperature profiles and represents regionwide averages, but the model vertical structure includes low (200 m) mixing heights over water with an adjustment to represent the extent of vertical dispersion of land-based emissions transported over water.

Results for each scenario are derived from a simulated base case and subsequent simulations with emission rates for either anthropogenic ROG or NO<sub>x</sub> reduced by 35% relative to emissions in the base case. In addition, a number of altered scenarios have been explored. These include (1) anthropogenic ROG emissions doubled from the inventory values (Lake Michigan), (2) anthropogenic ROG emissions cut in half from the inventory values (Lake Michigan), (3) zero biogenic emissions (northeast corridor), (4) height of the daytime mixed layer cut in half (northeast corridor), (5) biogenic emissions doubled from inventory values and height of the daytime mixed layer cut in half (northeast corridor).

The altered scenarios were designed to test the robustness of the indicator correlations and do not necessarily represent real events. However, it has been reported that emission rates for both anthropogenic and biogenic ROG may be twice as large as those reported in current inventories in the United States [Fujita et al., 1992; Geron et al., 1994]. Scenarios with ROG emissions lower than inventory values or with zero isoprene are unlikely to represent real conditions in the northeast corridor or Lake Michigan, but they may correspond to conditions in other environments (e.g., Europe or the southwestern United States).

# NO<sub>x</sub>-ROG Sensitivity and Indicator Species

The overall correlation between simulated O<sub>3</sub>-NO<sub>x</sub>-ROG sensitivity and potential indicator species will be illustrated for NO<sub>y</sub>, which was discussed extensively by Milford et al. [1994]. Figure 1a shows the correlation between simulated sensitivity and NO<sub>y</sub> for the Lake Michigan base case scenario. The figure shows several features that identify a good indicator species. There is a sharp delineation between NO<sub>x</sub>-sensitive locations (corresponding to NO<sub>y</sub> from 3 to 12 ppb), and ROG-sensitive locations (corresponding to NO<sub>y</sub> from 11 to 50 ppb). The overlap between NO<sub>x</sub>-sensitive locations and ROG-sensitive locations occupies a very narrow range of NO<sub>y</sub> values (11-12 ppb) and the number of locations with NO<sub>y</sub> in this range is a small fraction of the total model domain. In addition, the simulated NO<sub>y</sub> extend over a wide range and are not clustered near the transition between NO<sub>x</sub>- and ROG-sensitive locations.



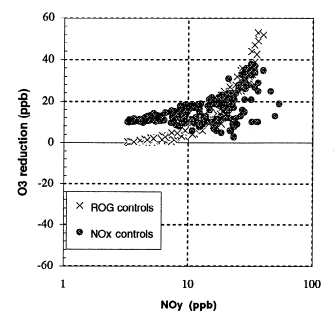


Figure 1a. Predicted reduction in peak  $O_3$  (in parts per billion, or ppb) resulting from a 35% reduction in the emission rate for anthropogenic ROG (crosses) and from a 35% reduction in the emission rate for  $NO_x$  (circles), plotted against  $NO_y$  (ppb) coincident with the ozone peak in the simulation for the Lake Michigan base case scenario [from Milford et al., 1994; Sillman et al., 1993].

Figure 1b. Predicted reduction in peak  $O_3$  (ppb) resulting from a 35% reduction in the emission rate for anthropogenic ROG (crosses) and from a 35% reduction in the emission rate for  $NO_x$  (circles), plotted against  $NO_y$  (ppb) coincident with the ozone peak in the simulation for the Lake Michigan scenario with doubled anthropogenic ROG [from *Milford et al.*, 1994].

When indicator values are located close to the  $NO_x$ -ROG transition point, the indicator-sensitivity correlation is likely to fall within the range of uncertainty associated with the model [e.g., Milford et al., 1993].

Results from the Lake Michigan scenario with doubled ROG emissions (Figure 1b) illustrate a less successful correlation between sensitivity and NO<sub>y</sub>. Here the range of NO<sub>y</sub> values associated with both NO<sub>x</sub>- and ROG-sensitive locations is much wider (16-32 ppb), and the number of locations with NO<sub>y</sub> within this overlap region represents a significant fraction (43%) of the entire domain. A comparison with Figure 1a also shows that the range of NO<sub>y</sub> associated with NO<sub>x</sub>-sensitive O<sub>3</sub> has expanded and the range associated with ROG-sensitive NO<sub>y</sub> has contracted in relation to the former scenario. Therefore NO<sub>y</sub> as an indicator apparently fails to account for the change in sensitivity that would result from a doubling of base case anthropogenic ROG.

The good correlation between sensitivity and NO<sub>y</sub> in Figure 1a and the comparatively poor correlation in Figure 1b can be summarized by recording the following parameters for each model scenario: scenario (a), the range of indicator values associated with NO<sub>x</sub>-sensitive locations (defined as locations where the simulated reduction in peak O<sub>3</sub> associated with reduced NO<sub>x</sub> exceeds the simulated reduction associated with reduced ROG by > 5 ppb); scenario (b), the range of indicator values associated with ROG-sensitive locations (defined similarly); and scenario (c), the "20% uncertainty fraction," equal to the fraction of the model domain with indicator values that are within 20% of the range of indicator values associated with the opposite NO<sub>x</sub>-ROG sensitivity. These parameters are recorded for each indicator and each scenario in Table 1 and will be the basis for identifying their usefulness. Results for each indicator will be discussed here.

NO<sub>y</sub>. The use of NO<sub>y</sub> as an indicator for sensitivity was discussed in detail by *Milford et al.* [1994] and has been illustrated in Figure 1. The two cases in Figure 1 illustrate both the positive and negative features of NO<sub>y</sub> as an indicator.

As shown in Table 1, most of the individual simulations show a strong correlation between sensitivity and NO, with a broad range of simulated NO<sub>v</sub> concentrations and a relatively narrow range overlap between NO<sub>x</sub>- and ROG-sensitive locations. Results from Milford et al. [1994] showed a similar strong correlation and similar ranges of transition between NO<sub>x</sub>- and ROG-sensitive NO<sub>v</sub>. However, the transition point between NO<sub>x</sub>- and ROG-sensitive locations tends to shift in response to changes in anthropogenic and biogenic ROG. Referring to Table 1, the transition occurs at ~10 ppb in simulations with little or no biogenic ROG (Lake Michigan base scenario and northeast corrridor scenario with zero isoprene). The northeast corridor base scenario has significantly more biogenic ROG than Lake Michigan and a higher transition (~20 ppb), and the northeast corridor scenario with doubled isoprene has a transition at ~35 ppb. Similarly, the Lake Michigan scenario with doubled anthropogenic ROG (Figure 1b) has a higher transition than the base scenario. By contrast the northeast scenario with a low mixed layer has a transition point (22 ppb) virtually identical with the base scenario.

The rationale for NO<sub>y</sub> as an indicator is based in part on the impact of stagnant meteorology on NO<sub>x</sub>-ROG sensitivity. Stagnant meteorology and associated high NO<sub>x</sub>, ROG, and NO<sub>y</sub> cause an increase in the photochemical lifetimes of NO<sub>x</sub> and ROG, with the result that an aging urban plume remains in the ROG-sensitive regime for a longer period of time. With more vigorous meteorological dispersion and lower NO<sub>x</sub>, ROG, and NO<sub>y</sub> an aging urban plume would rapidly become NO<sub>x</sub> sensitive [Milford et al., 1994]. The contrast between the northeast

Table 1. NO, and ROG-Sensitive Ranges for Indicator Species

Model Scenario	Indicator						
			O <sub>3</sub> O <sub>3</sub>		O3-40 ppb	O <sub>3</sub>	
	NO <sub>y</sub> , ppb	$NO_y - NO_x$ , ppb	NOy	NO <sub>y</sub> - NO <sub>x</sub>	$\frac{O_3-40 \text{ ppb}}{NO_y - NO_x}$	HNO <sub>3</sub>	
Lake Michigan Base C	Case Scenario						
(a) NO <sub>x</sub> range	4-11	3-9	7.2-17	9.5-19	4.8-8.8	12-45	
(b) ROG range	11-50	10-33	1.9-7.4	3.5-8.4	2.0-5.5	4.1-12	
(c) Uncertainty	0.17	0.12	0.27	0.13	0.49	0.15	
Lake Michigan Scenar	io with Doubled A	Anthropogenic ROG					
(a) NO <sub>x</sub> range	4-31	3-29	6.9-17	7.3-19	5.2-9.1	10-52	
(b) ROG range	16-52	13-45	3.8-7.8	4.8-8.7	3.4-6.4	6.7-15	
(c) Uncertainty	0.54	0.53	0.57	0.54	0.75	0.54	
Lake Michigan Scenari	io With Anthropos	genic ROG Reduced by	50%				
(a) NO <sub>x</sub> range	4-12	3-11	6.5-17	7.2-21	3.6-8.7	8.5-47	
(b) ROG range	14-35	9-26	2.2-6.4	3.6-8.9	1.8-4.9	4.0-12	
(c) Uncertainty	0.11	0.32	0.18	0.32	0.41	0.28	
Lake Michigan Base C	'ase Scenario. Ser	sitivity of O3 at Noon	1				
(a) NO <sub>x</sub> range	3-7	2-5	9.0-22	14-25	5.3-7.8	11-56	
(b) ROG range	9-65	6-18	0.9-7.0	3.1-11	0.7-4.9	3.5-16	
(c) Uncertainty	0.03	0.13	0.04	0.09	0.20	0.29	
Northeast Corridor Bas	se Case Scenario						
(a) NO <sub>x</sub> range	4-22	4-19	5.6-19	7.9-21	5.3-11	14-86	
(b) ROG range	16-34	11-23	3.4-7.1	6.2-8.6	4.1-5.6	9.6-16	
(c) Uncertainty	0.14	0.29	0.10	0.15	0.11	0.15	
Northeast Corridor Sce	enario With Zero I	Isoprene					
(a) NO <sub>x</sub> range	3-8	3-6	10-19	14-22	6.4-11	17-34	
(b) ROG range	10-30	7-21	3.0-11	5.7-13	1.5-7.4	7.3-20	
(c) Uncertainty	0.09	0.09	0.33	0.26	0.91	0.75	
Northeast Corridor Wit	h Mixed Laver Re	educed by 50%					
(a) NO <sub>x</sub> range	5-26	4-22	4.1-19	7.5-20	5.3-11	14-92	
b) ROG range	19-44	10-30	2.5-7.4	5.2-9.0	3.7-6.3	8.0-16	
(c) Uncertainty	0.17	0.36	0.15	0.20	0.18	0.18	
Northeast Corridor Wit	h Doubled Isopre	ne and Mixed Layer Ro	educed by 509	%			
(a) NO <sub>x</sub> range	5-38	5-31	3.0-19	6.1-19	4.5-11	11-180	
(b) ROG range	34-44	19-31	3.0-5.2	6.3-7.0	4.8-5.3	11-15	
(c) Uncertainty	0.02	0.15	0.04	0.07	0.12	0.04	

corridor base scenario and the scenario with a low mixed layer demonstrates the impact of stagnant meteorology. The base scenario is dominated by NO<sub>x</sub>-sensitive photochemistry, but the scenario with a low mixed layer has an extensive ROG-sensitive region. The difference in NO<sub>x</sub>-ROG sensitivity between these two scenarios is reflected by higher NO<sub>y</sub> in the latter, and the transition between NO<sub>x</sub>- and ROG-sensitive photochemistry occurs at virtually the same NO<sub>y</sub> in each (20-22 ppb). NO<sub>y</sub> as an indicator can also be explained in terms of ROG/NO<sub>x</sub> ratios if reactivity-weighted ROG remains relatively constant, as suggested by *Chameides et al.* [1992]. The changes in the transition point for NO<sub>y</sub> in simulations with different ROG are consistent with the well-known dependence of NO<sub>x</sub>-ROG sensitivity on ROG/NO<sub>x</sub> ratios.

The transition point for NO<sub>y</sub> also varies with the photochemical age of an air mass. Ozone formation is more likely to be sensitive to ROG in locations close to emission sources, and the associated transition between NO<sub>x</sub>- and ROG-sensitive chemistry is likely to occur at lower NO<sub>y</sub>. This effect can be illustrated by comparing ozone-NO<sub>y</sub>-ROG sensitivity early in the day with sensitivity associated with peak O<sub>3</sub>, usually in middle or late afternoon. As shown in Table 1, the sensitivity of O<sub>3</sub> at noon in the Lake Michigan base scenario correlates with NO<sub>y</sub>, but the transition point is shifted toward lower NO<sub>y</sub> (8 ppb versus 11 ppb for peak O<sub>3</sub>).

The sum of  $NO_x$  reaction products (=  $NO_y$  -  $NO_x$ ) can also be used as an indicator for sensitivity and is included in Table 1.  $NO_y$  -  $NO_x$  has an advantage over  $NO_y$  as an indicator because

Table 1. (continued)

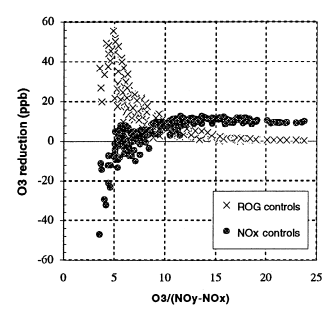
	Indicator					
M-1-1 C	НСНО	НСНО-5 ррь	H <sub>2</sub> O <sub>2</sub>	H <sub>2</sub> O <sub>2</sub> NO <sub>y</sub> - NO <sub>x</sub>	H <sub>2</sub> O <sub>2</sub> NO <sub>y</sub>	
Model Scenario	NOy	NOy	HNO <sub>3</sub>			
Lake Michigan Base (	Case Scenario					
(a) NO <sub>x</sub> range	0.29-0.67	0.79-1.9	0.37-2.3	0.27-0.97	0.22-0.82	
(b) ROG range	0.14-0.31	0.28-0.73	0.01-0.30	0.01-0.25	0.007-0.24	
(c) Uncertainty	0.30	0.16	0.03	0.11	0.16	
Lake Michigan scenar	io with doubled anth	ropogenic ROG				
(a) NO <sub>x</sub> range	0.35-0.76	0.55-2.0	0.25-2.8	0.17-1.0	0.16-0.89	
(b) ROG range	0.23-0.42	0.42-067	0.03-0.36	0.02-0.23	0.02-0.22	
(c) Uncertainty	0.64	0.55	0.34	0.38	0.40	
Lake Michigan scenar	io with anthropogenic	: ROG reduced by 50%				
(a) NO <sub>x</sub> range	0.20-0.66	0.60-1.9	0.26-2.4	0.21-1.1	0.18-0.84	
(b) ROG range	0.1023	0.26-0.56	0.03-0.29	0.02-0.22	0.01-0.16	
(c) Uncertainty	0.34	0.16	0.16	0.16	0.11	
Lake Michigan Base (	Case Scenario, SNoon	,				
(a) NO <sub>x</sub> range	0.39-0.82	1.1-2.8	0.67-2.2	0.31-1.1	0.27-1.0	
(b) ROG range	0.11-0.36	0.22-0.82	0.008-0.38	0.01-0.28	0.002-0.15	
(c) Uncertainty	0.17	0.04	0	0.06	0	
Northeast Corridor Ba	se Case Scenario					
(a) NO <sub>x</sub> range	0.26-1.2	0.55-2.2	0.43-6.1	0.24-1.6	0.22-1.5	
(b) ROG range	0.23-0.42	0.41-0.67	0.19-0.54	0.12-0.29	0.09-0.23	
(c) Uncertainty	0.57	0.18	0.08	0.08	0.02	
Northeast corridor sce	enario with zero isopr	ene				
(a) NO <sub>x</sub> range	0.26-0.52	0.92-1.7	0.61-1.5	0.48-1.2	0.39-1.1	
(b) ROG range	0.1738	0.37-0.83	0.10-0.64	0.08-0.50	0.05-0.45	
(c) Uncertainty	0.93	0.17	0.16	0.15	0.21	
Northeast Corridor Wi	ith Mixed Layer Redu	ced by 50%				
(a) NO <sub>x</sub> range	0.28-1.3	0.54-2.2	0.46-7.0	0.24-1.5	0.15-1.4	
(b) ROG range	0.25-0.52	0.36-0.77	0.12-0.55	0.08-0.35	0.05-0.33	
(c) Uncertainty	0.62	0.28	0.07	0.12	0.13	
Northeast Corridor Wi	th Doubled Isoprene d	and Mixed Layer Reduct	ed by 50%			
(a) NO <sub>x</sub> range	0.33-1.8	0.49-2.6	0.45-20.	0.20-1.6	0.15-1.5	
(b) ROG range	0.32-0.43	0.43-0.56	0.20-0.62	0.11-0.28	0.08-0.17	
(c) Uncertainty	0.09	0.04	0.04	0.05	0.01	

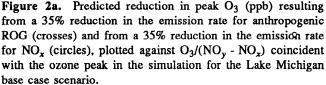
The range of indicator values associated with (a)  $NO_x$ -sensitive chemistry and (b) ROG-sensitive chemistry is given for each simulation, along with (c) the fraction of model indicator values associated with uncertain  $NO_x$ -ROG sensitivity. The  $NO_x$ -sensitive range is defined as the range of locations with simulated peak  $O_3$  in simulations with reduced  $NO_x$  lower than in simulations with reduced ROG by at least 5 ppb. The ROG-sensitive range is defined analogously. The uncertainty fraction is defined by the fraction of the model domain with indicator values within 20% of the indicator range associated with the opposite  $NO_x$ -ROG sensitivity.

 $NO_y$  field measurements in urban locations may be impacted by on-site  $NO_x$  emission sources. However, the correlation between sensitivity and  $NO_y$  -  $NO_x$  in models is somewhat worse than the correlation with  $NO_y$ , with a broader overlap between  $NO_x$ -sensitive and ROG-sensitive ranges (Table 1). Transition points for  $NO_y$  -  $NO_x$  are ~30% lower than the corresponding transition point for  $NO_y$  in these simulations.

The subsequent indicator ratios based on  $NO_y$  can all be used with  $NO_y$  -  $NO_x$  substituted for  $NO_y$  and with the same 30% change in transition values.

$$\frac{O_3}{NO_y - NO_x}$$
 and  $\frac{O_3 - 40ppb}{NO_y - NO_x}$ . The slope of  $O_3$  versus  $NO_y$  or  $O_3$  versus  $NO_x$  reaction products  $(NO_y - NO_x)$  has been used





frequently to analyze photochemistry of  $O_3$  in polluted rural areas [Sillman et al., 1990b; McKeen et al., 1991; Trainer et al., 1993]. The ratio  $(O_3 - 40 \text{ ppb})/(NO_y - NO_x)$  can be used from individual measurements to quantify the slope, where 40 ppb represents background  $O_3$  both in the current simulations and in the measured correlations between  $O_3$  and  $NO_y - NO_x$  of Trainer et al. [1993].

A rationale for either  $O_3/NO_y$  or  $(O_3 - 40 \text{ ppb})/NO_y$  as indicator for  $NO_x$ -ROG sensitivity can be derived from the analysis of odd hydrogen in the appendix. The transition from  $NO_x$ - to ROG-sensitive chemistry is associated with the ratio of sources of odd hydrogen  $(S_H)$  to a modified sum of  $NO_x$  reaction products  $(NO_y - NO_x + HNO_3)$  (equation (A6)). If the source of odd hydrogen is proportional to  $[O_3]$ , then the transition criterion given by (A6) is approximately correlated to the ratio  $O_3/(NO_y - NO_x)$ . Alternatively, if the source of odd hydrogen is proportional to the rate of production of ozone, then the transition criterion is equivalent to a constant slope of  $O_3$  versus  $NO_y - NO_x$ . The slope of  $O_3$  versus  $NO_y - NO_x$  has also been discussed in terms of production efficiencies for  $O_3$  [Liu et al., 1987; Lin et al., 1988].

Results for both indicators are shown in Figure 2 and Table 1. The performance of  $O_3/(NO_y - NO_x)$  as an indicator is comparable to  $NO_y$  with a well-defined transition between  $NO_x$ -sensitive chemistry  $(O_3/(NO_y - NO_x) > 9)$  and ROG-sensitive chemistry  $(O_3/(NO_y - NO_x) < 8)$ . In contrast to  $NO_y$ , the transition point for  $O_3/(NO_y - NO_x)$  shows relatively little variation among the model scenarios. For example, the transition points for  $O_3/(NO_y - NO_x)$  in the Lake Michigan base- and doubled-ROG scenarios (Figures 2a and 2b) are virtually identical. By contrast, the transition point for  $NO_y$  changed by a factor of 2 between the scenarios. Significant differences in the transition point for  $O_3/(NO_y - NO_x)$  only appear for the northeast corridor scenarios with zero isoprene (transition at  $O_3/(NO_y - NO_x) \cong 13$ ) and doubled isoprene

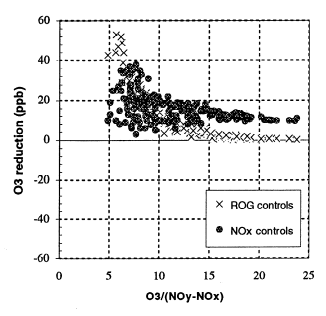


Figure 2b. Predicted reduction in peak  $O_3$  (ppb) resulting from a 35% reduction in the emission rate for anthropogenic ROG (crosses) and from a 35% reduction in the emission rate for  $NO_x$  (circles), plotted against  $O_3/(NO_y - NO_x)$  coincident with the ozone peak in the simulation for the Lake Michigan scenario with doubled anthropogenic ROG.

(transition at  $O_3/(NO_y - NO_x) \cong 6$ ). The correlation with sensitivity at noon also shows a different transition point  $(O_3/(NO_y - NO_x) \cong 13)$ . The ratio  $O_3/NO_y$  also performs well as an indicator in these simulations (transition point at  $O_3/NO_y = 6-7$ ), although  $O_3/(NO_y - NO_x)$  is correct in terms of theory. The ratio  $O_3/HNO_3$  also performs well.

The ratio  $(O_3 - 40 \text{ ppb})/(NO_y - NO_z)$  (Figure 3 and Table 1) is less successful as in indicator than  $O_3/(NO_y - NO_z)$ . Although

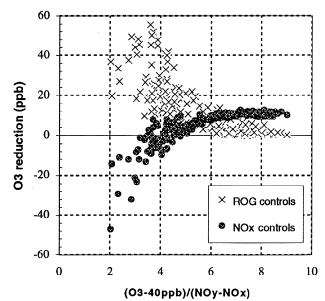


Figure 3. Predicted reduction in peak  $O_3$  (ppb) resulting from a 35% reduction in the emission rate for anthropogenic ROG (crosses) and from a 35% reduction in the emission rate for  $NO_x$  (circles), plotted against  $(O_3 - 40 \text{ ppb})/(NO_y - NO_x)$  coincident with the ozone peak in the simulation for the Lake Michigan base case scenario.

the transition point associated with (O<sub>3</sub> - 40 ppb)/(NO<sub>y</sub> - NO<sub>x</sub>) is generally consistent among the seven model scenarios, a large proportion of indicator values are located very close to the transition region. Consequently, the fraction of the model domain associated with or close to the region of overlap between NO<sub>x</sub>- and ROG-sensitive ranges is larger than that for  $NO_y$  or  $O_3/(NO_y - NO_x)$ . The ratio  $(O_3 - 40 \text{ ppb})/NO_y$  performs The ratio (O<sub>3</sub> - 40 ppb)/NO<sub>y</sub> is especially interesting because it is directly comparable to the AIRTRAK method [Johnson, 1984; Johnson et al., 1990; Blanchard et al., 1993], which is being investigated for regulatory use by the U.S. Environmental Protection Agency as a basis for identifying NO<sub>x</sub>- versus ROG-sensitive chemistry. Johnson et al. derived an indicator for NO<sub>x</sub>-ROG sensitivity equivalent to (O<sub>3</sub> - 40 ppb)/NO<sub>y</sub>, using results from smog chambers, and identified the transition point as  $(O_3 - 40 \text{ ppb})/NO_v = 4.09$ . The simulations here have a transition point at (O<sub>3</sub> - 40 ppb)/ $NO_y = 3-4$  but with considerable scatter.

Both  $O_3/(NO_y - NO_x)$  and  $(O_3 - 40 \text{ ppb})/(NO_y - NO_x)$  are associated in theory with rates of production for odd hydrogen. Consequently, it is likely that transition points will shift in simulations with different amounts of solar radiation of  $H_2O$ , both of which affect the odd hydrogen source term, and with temperature, which affects PAN.

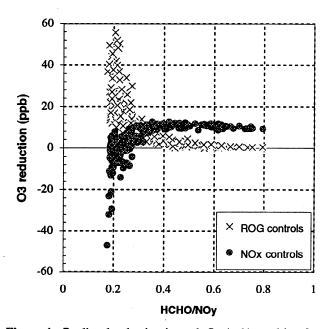
HCHO/NO<sub>y</sub>. Figure 4 shows the correlation between NO<sub>x</sub>-ROG sensitivity and the ratio HCHO/NO<sub>y</sub> at the time of the ozone peak. The ratio HCHO/NO<sub>y</sub> functions as a reactivity-weighted ROG/NO<sub>x</sub> ratio, since production of HCHO is roughly proportional to the summed rate of reactions of ROG with OH. Low HCHO/NO<sub>y</sub> is associated with ROG-sensitive ozone, a result that parallels the relation between ozone sensitivity and ROG/NO<sub>x</sub>. Other ROG-based indicators, including reactivity-weighted total ROG and ROG/NO<sub>x</sub> and ROG/NO<sub>y</sub> ratios, did not provide a good correlation with ozone-NO<sub>x</sub>-ROG sensitivity in

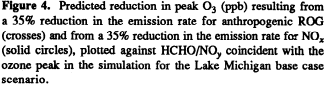
the simulations. The ratio  $HCHO/(NO_y - NO_x)$  can also be associated with  $NO_x$ -ROG sensitivity through the analysis of odd hy drogen, as was done for  $O_3/(NO_y - NO_x)$ , with the assumption that the odd hydrogen source is proportional to HCHO.

The correlation between sensitivity and HCHO/NO<sub>y</sub> is comparable with NO<sub>y</sub> in its consistency, although its usefulness is partially compromised by its relatively narrow range of values. The model predicts the crossover between NO<sub>x</sub>-sensitive and ROG-sensitive ozone occurs at HCHO/NO<sub>y</sub>  $\cong$  0.28, while the ROG-sensitive Chicago urban plume is predicted to have HCHO/NO<sub>y</sub>  $\cong$  0.20. In highly polluted ROG-sensitive environments the ratio appears to approach 0.15 as an asymptotic limit. The clustering of HCHO/NO<sub>y</sub> close to the transition point is reflected in the large fractions of the model domain associated with overlap between NO<sub>x</sub>- and ROG-sensitive ranges (Table 1).

An interesting feature of HCHO/NO<sub>y</sub> as an indicator is that the HCHO/NO<sub>y</sub> correlations represent the impact of changes in ROG emissions. HCHO/NO<sub>y</sub> can be combined with NO<sub>y</sub> to form an indicator ((HCHO + 5 ppb)/NO<sub>y</sub>) with a transition point that shows relatively small variations in response to changed model emission rates for ROG (see Table 1).

H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub>. The connection between NO<sub>x</sub>-ROG sensitivity and the ratio of hydrogen peroxide to nitric acid is stronger than for the other indicators in terms of both theory and simulation results. It was shown in section 2 and the appendix (Equation (A6)) that NO<sub>x</sub>-ROG sensitivity is linked to the relative rates of formation of peroxides and nitric acid and to their role as sinks for odd hydrogen. Simulation results (Figure 5 and Table 1) show a close correlation between H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub> and sensitivity for all model scenarios. The overlap between NO<sub>x</sub>- and ROG-sensitive ranges is consistently small in comparison with the range of simulated values for H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub>,





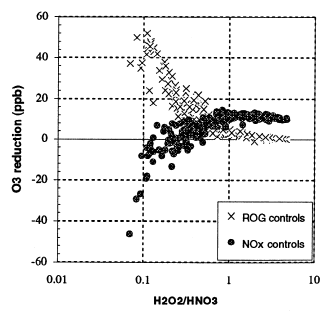


Figure 5. Predicted reduction in peak  $O_3$  (ppb) resulting from a 35% reduction in the emission rate for anthropogenic ROG (crosses) and from a 35% reduction in the emission rate for  $NO_x$  (solid circles), plotted against  $H_2O_2/HNO_3$  coincident with the ozone peak in the simulation for the Lake Michigan base case scenario.

and the "uncertainty fractions" are low in comparison with the other indicators. The transition from  $NO_{x^-}$  to ROG-sensitive chemistry occurs at  $H_2O_2/HNO_3 \cong 0.3$ -0.5 and shows relatively small variation among the model scenarios. The northeast corridor scenario with zero isoprene has a signficantly higher transition ratio (0.6), and the northeast scenarios all have slightly higher transition points than the Lake Michigan scenarios. In contrast to the previous indicators, the sensitivity correlation for  $H_2O_2/HNO_3$  for  $O_3$  at noon shows the same transition point as the correlations for peak  $O_3$ . Table 1 also shows indicator correlations for  $H_2O_2/NO_y$ , which might be used in place of  $H_2O_2/HNO_3$  if  $HNO_3$  measurements are unavailable.

Uncertainties. The extent of overlap between NO<sub>x</sub>-sensitive and ROG-sensitive ranges for indicator species and the comparison between indicator transition points in different model scenarios provides a partial representation of the uncertainties associated with the sensitivity-indicator correlations. The use of model-based correlations also involves additional uncertainties based on model assumptions or omissions. Some of these uncertainties will be discussed here.

All of the indicators include at least one species (HNO<sub>3</sub>) which has a high rate of surface deposition. Another species (H<sub>2</sub>O<sub>2</sub>) may form an aerosol in the presense of haze (C. Walcek. State University of New York at Albany, private communication, 1994). The indicator values reported here are therefore sensitive to assumed deposition and aerosol formation rates. The impact of deposition in the models can be seen by comparing transition points for the northeast corridor scenario with zero isoprene with transition points for other scenarios. The zero-isoprene scenario is unusual in that ROGsensitive chemistry persists in urban plumes for a period of more than 24 hours downwind from emission sources, allowing for much greater deposition of HNO<sub>3</sub> during transport associated with the transition region. Day-old urban plumes all have NO<sub>x</sub>-sensitive chemistry in the other scenarios. Table 1 shows that transition points in the zero-isoprene scenario are different from the other scenarios for every indicator, and that the difference is consistent with a higher rate of removal for

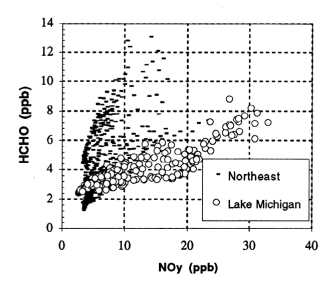


Figure 6. HCHO versus NO<sub>y</sub> (ppb) at 1800 LT in simulations for the northeast corridor (points) and the Lake Michigan airshed (circles).

HNO<sub>3</sub> at the transition point. Differences in model deposition rates also affect predicted cross-species correlations (below).

The impact of solar radiation,  $[H_2O]$  or temperature on indicator correlations has not been explored. At least one indicator  $(O_3/(NO_y - NO_x))$  is expected to vary significantly in response to changes in radiation and  $[H_2O]$ , which affect production rates for odd hydrogen. Temperature affects formation rates for PAN and therefore may also have an impact on the performance of indicators involving  $NO_y$ .

All model calculations are sensitive to uncertainties in chemical rate constants and stoichiometry, which generally cause a 20-30% uncertainty in resulting species concentrations [Milford et al., 1993].  $H_2O_2$  is especially vulnerable to reaction rate and mechanism uncertainties.  $H_2O_2$  is closely associated with model formation of organic peroxides  $RO_2$  +  $HO_2$  reactions (R6), which varies significantly among different photochemical mechanisms. In the current simulation the formation of organic peroxides via (R6) is approximately half the rate of formation of  $H_2O_2$ , but this simulated rate has not been compared with atmospheric measurements. Formation of  $H_2O_2$  and some organic peroxides (e.g.,  $HOCH_2OOH$ ) from reactions other than (R6) has not been included and adds further uncertainty to the interpretation of  $H_2O_2$  and organic peroxide concentrations.

## **Correlations Between Indicator Species**

The proposed connection between  $NO_x$ -ROG sensitivity and indicator species would be strengthened if the simulations could also be used to generate correlations between indicator species that could be tested against measurements. Unfortunately, the simulations predict no strong correlations among all the indicator species apart from the  $O_3$  -  $NO_y$  correlation explored by *Trainer et al.* [1993].

Figures 6 and 7 show simulated correlations between the species associated with indicator ratios, HCHO versus NO<sub>y</sub> and H<sub>2</sub>O<sub>2</sub> versus HNO<sub>3</sub>. Results illustrate the differences between the largely NO<sub>x</sub>-sensitive chemistry in the northeast corridor base case scenario and the largely ROG-sensitive chemistry in

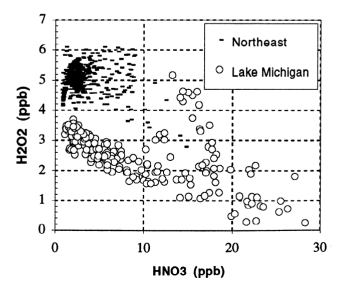


Figure 7. H<sub>2</sub>O<sub>2</sub> versus HNO<sub>3</sub> (ppb) at 1800 LT in simulations for the northeast corridor (points) and the Lake Michigan airshed (circles).

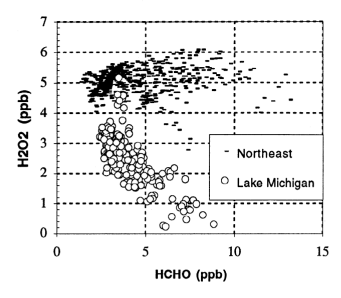


Figure 8. H<sub>2</sub>O<sub>2</sub> versus HCHO (ppb) at 1800 LT in simulations for the northeast corridor (points) and the Lake Michigan airshed (circles).

the Lake Michigan base case. HCHO increases versus  $NO_y$  with two distinct slopes, apparently corresponding to  $NO_x$ - versus ROG-sensitive chemistry. The distinct slopes may also represent the difference between photochemical evolution of combined anthropogenic and biogenic emissions over land versus evolution of urban plumes transported over water with little biogenic emission.  $H_2O_2$  and  $HNO_3$  anticorrelate with each other, but the northeast corridor simulation has higher  $H_2O_2$  and a weaker anticorrelation than the Lake Michigan simulation. The lack of a consistent correlation among these species is partially explained by their tendency to assume different correlative patterns in  $NO_x$ - versus ROG-sensitive conditions.

There is also no consistent correlation between the simulated  $H_2O_2$  and HCHO or between  $H_2O_2$  and  $O_3$  (Figures 8 and 9). However, there is an intriguing triple correlation among O<sub>3</sub>, H<sub>2</sub>O<sub>2</sub>, and NO<sub>2</sub> reaction products. In theory (see appendix) the sum H<sub>2</sub>O<sub>2</sub> + NO<sub>y</sub> - NO<sub>x</sub> represents the cumulative sink for odd hydrogen and may be expected to correlate with O3. A correlation of this type is predicted in the simulation for the northeast corridor (Figure 10a), where  $O_3$  versus  $H_2O_2 + NO_v$  -NO<sub>x</sub> shows a stronger correlation that O<sub>3</sub> versus NO<sub>y</sub> - NO<sub>x</sub> or O<sub>3</sub> versus HNO<sub>3</sub>. A weaker pattern appears in the Lake Michigan simulation (Figure 10b), although the correlation between O<sub>3</sub> and H<sub>2</sub>O<sub>2</sub> + NO<sub>y</sub> - NO<sub>x</sub> is again stronger than the correlation between O<sub>3</sub> and reactive nitrogen. A significant part of the poor correlation in Figure 9b is due to the change in surface deposition rates between the land and the lake shore in the model. The high  $H_2O_2 + NO_y - NO_x$  versus  $O_3$  all corresponds to locations associated with transport over Lake Michigan, where model deposition rates are low. The low ratios in Figure 10b represent locations over land and are similar to the ratios in the northeast corridor simulation (Figure 10a).

Although difficult to measure, the predicted  $H_2O_2 + NO_y - NO_x$  versus  $O_3$  and its interpretation in terms of odd hydrogen may represent an extension of the rural  $O_3$  versus  $NO_y - NO_x$  to urban environments. The viewpoint here is different from that of *Liu et al.* [1987], *Lin et al.* [1988], *McKeen et al.* [1991], and

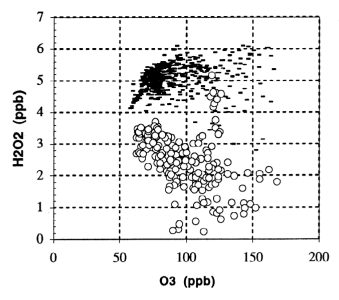


Figure 9. H<sub>2</sub>O<sub>2</sub> versus O<sub>3</sub> (ppb) at 1800 LT in simulations for the northeast corridor (points) and the Lake Michigan airshed (circles).

Trainer et al. [1993], who emphasize the role of  $NO_x$  as precursor for  $O_3$  and the slope  $O_3$  versus  $NO_y$  -  $NO_x$  as a representation of varying production efficiencies for  $O_3$ . The view presented here emphasizes  $O_3$  as a source for odd hydrogen, either directly or through association with intermediate hydrocarbons, and rates of formation for both peroxides and reactive nitrogen as limited by the size of the odd hydrogen source. The transition from  $NO_x$ - to ROG-sensitive chemistry is linked with the replacement of peroxides by HNO<sub>3</sub> as the dominant sink for odd hydrogen, and therefore by a decreasing ratio of  $O_3$  to reactive nitrogen.

# **Comparison With Observations**

In recent years, concurrent measurements of NO<sub>y</sub>, HCHO, H<sub>2</sub>O<sub>2</sub>, and HNO<sub>3</sub> have been made in Claremont, California, near Los Angeles (Bart Croes, State of California Air Resources Board, Sacramento, California, private communication, 1994), in Atlanta, Georga [Sillman et al., 1995], and at a rural site in Virginia [Jacob et al., 1995]. Complete analyses of field measurements will be provided in papers authored by the principal investigators, but results have been graciously provided. Because some results are preliminary, they have large uncertainties (+/- 25%) but they provide evidence for cross correlation among the indicator species.

As shown in Table 2, measured values during an event at Claremont, California are consistent with ROG-sensitive chemistry. These included very high (> 50 ppb) afternoon NO<sub>y</sub>, low (< 1 ppb)  $\rm H_2O_2$ , and an HCHO/NO<sub>y</sub> ratio of 0.16. The ROG-sensitive nature of ozone in Claremont is consistent with published modelling studies for the region [Milford et al., 1989], although model results also suggest that the chemistry becomes sensitive to NO<sub>x</sub> at downwind locations within greater Los Angeles.

Measured concentrations for Atlanta are more difficult to interpret because NO<sub>y</sub> was only availale from helicopter-based measurements 40 km downwind from Atlanta, while H<sub>2</sub>O<sub>2</sub> and HCHO were measured at a site near downtown. Helicopter-based

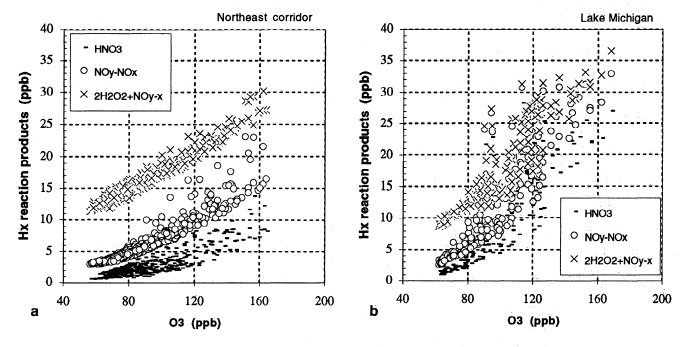


Figure 10. HNO<sub>3</sub> (points), NO<sub>y</sub> - NO<sub>x</sub> (circles) and  $H_2O_2$  + (NOy - NOx) (crosses) versus  $O_3$ , all in ppb at 1800 LT in simulations for (a) the northeast corridor and (b) the Lake Michigan airshed.

measurements at 600 m found ~12 ppb NO<sub>y</sub> coincident with peak O<sub>3</sub> (143 ppb), values which are consistent with models [Sillman et al., 1995]. Given the high emission rate for isoprene in Atlanta, the measured NO<sub>y</sub> and O<sub>3</sub>/NO<sub>y</sub> both suggest NO<sub>x</sub>-sensitive ozone. However, surface measurements of HCHO at South Dekalb, ~5 km from the helicopter measurements, were lower than would be expected in a NO<sub>x</sub>-sensitive environment and lower than those predicted by photochemical models [Sillman et al., 1995. Peroxides (3.5 ppb) were measured near downtown, but up to 50% of this total may represent organic peroxides other than H<sub>2</sub>O<sub>2</sub> [Sillman et al., 1995; Lee et al., 1993]. The resulting values for both HCHO/NO<sub>y</sub> and H<sub>2</sub>O<sub>2</sub>/NO<sub>y</sub> (see Table 2) suggest NO<sub>x</sub>-sensitive chemistry but lie close to the transition point. All four indicators show a large contrast between Claremont and

Atlanta, suggesting that chemistry is fundamentally different in the two locations.

Jacob et al. [1995] measured NO<sub>y</sub> and H<sub>2</sub>O<sub>2</sub> at a site in rural Virginia during September and October 1990. They reported a significant variation in H<sub>2</sub>O<sub>2</sub>/NO<sub>y</sub> ratios between the beginning and the end of the time period, which corresponded to a sharp decrease in solar radiation, H<sub>2</sub>O, and biogenic emission of isoprene at the start of autumn. The measured H<sub>2</sub>O<sub>2</sub>/NO<sub>y</sub> ratio on a day with high O<sub>3</sub> in early September, shown in Table 2, was higher than that in the Atlanta measurements and is consistent with predictions of NO<sub>x</sub>-sensitive O<sub>3</sub> in rural eastern North America [Sillman et al., 1990b; McKeen et al., 1991; Possiel et al., 1991]. As discussed by Jacob et al. [1995], the sharp drop in H<sub>2</sub>O<sub>2</sub>/NO<sub>y</sub> between September and October may be associated with a shift

Table 2. Measured Concentrations of Indicator Species

Species	Claremont, California, <sup>a</sup> June 25, 1987, 1300 LT	Atlanta, Georgia, <sup>b</sup> August 10, 1990, 1700 LT	Virginia, <sup>c</sup> September 12, 1990, 1500 LT
O <sub>3</sub>	200	(143)	75
NO <sub>ν</sub>	65	(12)	4
нсно	10.5	4	•••
H <sub>2</sub> O <sub>2</sub>	0.99	(3.5)	1.5
HNO <sub>3</sub>	20	•••	
O <sub>3</sub> /NO <sub>y</sub>	3.1	(12)	19
HCHO/NO <sub>v</sub>	0.16	(0.33)	•••
H <sub>2</sub> O <sub>2</sub> /NO <sub>y</sub>	0.017	(0.1-5-0.3)	0.38
H <sub>2</sub> O <sub>2</sub> /HNO <sub>3</sub>	0.05	,	

a From B. Croes (private communication, 1994).

b Ratios for Atlanta are based on helicopter observations of the plume 30 km southeast of Atlanta, coincident with peak O<sub>3</sub> at 600 m. HCHO and H<sub>2</sub>O<sub>2</sub> were observed at the Georgia Tech campus near downtown. From Sillman et al. [1995].

c From Jacob et al. [1995].

toward ROG-sensitive chemistry during the winter. *Kleinman* [1991] also discusses the seasonal shift from NO<sub>x</sub>-sensitive to ROG-sensitive chemistry in eastern North America and correlates it with changes in measured H<sub>2</sub>O<sub>2</sub>.

#### Conclusions

Model predictions for  $O_3$ -NO<sub>x</sub>-ROG sensitivity have been shown to correlate with simulated values for four indicator species: NO<sub>y</sub>,  $O_3/(NO_y - NO_x)$ , (HCHO-5 ppb)/(NO<sub>y</sub> - NO<sub>x</sub>), and  $H_2O_2/HNO_3$ , all coincident with the ozone peak. The derived correlations between ozone sensitivity and the indicator species change somewhat when model assumptions are altered, but the model correlations remain at least partially valid, even when model emission rates are stretched far beyond the range of currently accepted values. The sensitivity correlations for  $O_3/(NO_y - NO_x)$  and  $H_2O_2/HNO_3$  appear to be robust and are not affected by changes in model assumptions, although they may be affected by changes (e.g., solar radiation, and aerosol formation) not included here. The connection between these species and  $NO_x$ -ROG sensitivity has also been explained in terms of the fundamental chemistry of  $O_3$ ,  $NO_x$ , and ROG.

The correlations suggest that the indicator species may provide a powerful tool in assessing the relative effectiveness of ROG versus  $NO_x$  controls and also will provide an important test for the accuracy of ozone- $NO_x$ -ROG sensitivity as predicted by individual applications of urban models. The success of  $O_3/(NO_y - NO_x)$  as an indicator is especially important as a tool for model evaluation. At present, regulatory decisions in the United States are made on the basis of results of photochemical models that are evaluated in terms of their ability to simulate measured  $O_3$ . The results of this paper suggest that evaluation of model performances in comparison with measured peak  $O_3$  and concurrent  $NO_y - NO_x$  would provide a stronger basis for confidence in predicted  $NO_x$ -ROG sensitivity.

# Appendix: Derivation of O<sub>3</sub>-ROG-NO<sub>x</sub> Sensitivity in Terms of ROG/NO<sub>x</sub>, H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub>, and O<sub>3</sub>/NO<sub>y</sub> Ratios

The split between ROG-sensitive and  $NO_x$ -sensitive photochemistry can be derived in theory from the steady state equation for odd hydrogen radicals (equation (1)). Production of  $O_3$  is assumed equal to the summed rates of the OH + RH and OH + CO reactions (R1) and (R4). In addition, equation (1) will be modified to include formation of organic nitrates (including both PAN and alkyl nitrates) as a sink for odd hydrogen with the assumption that the rate of formation ( $P_{PAN}$ ) is directly proportional to production rate for  $O_3$ . By using the following nomenclature substitutions,

$$P_{O_4} = k_4 [CO][OH] + k_1 [RH][OH]$$
 (A1a)

$$f_{\text{PAN}} = \frac{P_{\text{PAN}}}{P_{\text{O}_1}} \tag{A1b}$$

$$\chi = \frac{k_1[RH]}{k_4[CO] + k_1[RH]}$$
 (A1c)

$$\psi_1 = \frac{[NO]}{[NO_*]} \qquad \psi_2 = \frac{[NO_2]}{[NO_*]}$$
(A1d)

$$k_{56} = \frac{k_5}{{\psi_1}^2 k_3^2} + \frac{k_6 \chi}{{\psi_1}^2 k_2 k_3}$$
 (A1e)

Equation (1) becomes

$$S_{\rm H} = \frac{\chi k_7 \Psi_2 [NO_x]}{k_1 [RH]} P_{\rm O_3} + k_{56} \frac{1}{[NO_x]^2} P_{\rm O_3}^2 + P_{\rm PAN} \quad (A2)$$

The transition between the  $NO_x$ - and ROG-sensitive regimes can be identified by equating partial derivatives for  $P_{O_3}$  with respect to a percentage increase in  $NO_x$  ( $\partial NO_x/NO_x$ ) or RH ( $\partial RH/RH$ ):

$$[RH] \frac{\partial P_{O_3}}{\partial [RH]} = [NO_x] \frac{\partial P_{O_3}}{\partial [NO_x]}$$
 (A3)

Equation (A3) will be substituted into (A2) with the assumption that both  $S_H$  and  $P_{PAN}$  can be expressed as functions of  $P_{O_*}$ ;

i.e., 
$$\frac{\partial}{\partial [RH]}$$
 and  $\frac{\partial}{\partial [NO_x]}$  are zero for these terms. The

analytical relation between PAN and  $O_3$  is discussed by Sillman et al. [1990b, 1991]. It will also be assumed that  $\chi$  is constant. The resulting substitution yields

$$P_{\rm O_3} = \frac{\chi k_7 \psi_2}{k_1 k_{56}} \frac{[\rm NO_x]^3}{[\rm RH]}$$
 (A4)

Equation (A4) can be substituted into the rate expressions for reactions (R5), (R6), and (R7) to obtain a criterion for the transition point between  $NO_x$ -sensitive and ROG-sensitive chemistry in terms of photochemical production rates for  $H_2O_2$ , ROOH, and  $HNO_3$  ( $P_{H_2O_2}$ ,  $P_{ROOH_3}$  and  $P_{HNO_3}$ ):

$$P_{\text{HNO}_3} = 2(P_{\text{H}_2\text{O}_2} + P_{\text{ROOH}})$$
 (A5)

Equation (A5) provides the rationale for the use of peroxide to nitric acid ratios as indicators for sensitivity with an apparent transition point at  $(H_2O_2 + ROOH)/HNO_3 = 0.5$ , where ROOH is assumed to represent organic peroxides formed via reaction (R6) only. This is equivalent to a transition point at  $H_2O_2/HNO_3 = 0.35$  based on the relative magnitude of reactions (R5) and (R6) in the simulations. The simulation results in Table 1 show transition points at  $H_2O_2/HNO_3 \cong 0.3$ -0.6 with the highest transition ratios in simulations with greatest photochemical aging. The higher transition ratios in aged air may be caused by the higher rate of dry deposition of  $HNO_3$  relative to  $H_2O_2$ .

Equation (A5) can also be used to obtain transition criteria in terms of  $O_3$ ,  $NO_y$ , and  $NO_x$ . Substitution of (A5) into (A2) yields

$$S_{\rm H} = 2P_{\rm HNO_3} + P_{\rm PAN} \tag{A6}$$

Sources of odd hydrogen  $(S_H)$  include photolysis of both  $O_3$  and aldehydes, where aldehyde concentrations tend to increase with increasing  $O_3$ . If it is assumed that the source of odd hydrogen is proportional to  $O_3$ , then the transition between  $NO_x$ - and ROG-sensitive chemistry should be associated with a fixed value for the ratio

$$\frac{O_3}{NO_y - NO_x + HNO_3}$$
 (A7)

In practical terms this suggests the use of either O<sub>3</sub>/(NO<sub>y</sub> - NO<sub>x</sub>) or O<sub>3</sub>/HNO<sub>3</sub> as an indicator for NO<sub>x</sub>-ROG sensitivity.

The use of the slope of  $O_3$  versus  $NO_y$  -  $NO_x$  as an indicator can also be justified by converting (A2) to differential form,

$$\Delta S_{\rm H} = \Delta (2P_{\rm H_2O_2} + 2P_{\rm ROOH} + P_{\rm HNO_3} + P_{\rm PAN})$$
 (A8)

where  $\Delta S_{\rm H}$  versus  $\Delta (P_{\rm HNO_3} + P_{\rm PAN})$  can be associated with the slope of O<sub>3</sub> versus NO<sub>y</sub> - NO<sub>x</sub> for an ensemble of measurements. In combination with the criteria defined by (A5), this suggests that an ensemble of points with ROG-sensitive chemistry (with

high  $P_{\mathrm{HNO_3}}/P_{\mathrm{H_2O_2}}$ ) will have a lower  $\Delta \mathrm{O_3}/\Delta(\mathrm{NO_y} - \mathrm{NO_x})$  than a  $\mathrm{NO_x}$ -sensitive ensemble with low  $P_{\mathrm{HNO_3}}/P_{\mathrm{H_2O_2}}$ . However, a low  $\Delta \mathrm{O_3}/\Delta(\mathrm{NO_y} - \mathrm{NO_x})$  can also be indicative of an ensemble of points in transition between  $\mathrm{NO_x}$ - and ROG-sensitive chemistry (with rapidly varying  $P_{\mathrm{HNO_3}}/P_{\mathrm{H_2O_2}}$ ) as well as an ROG-sensitive ensemble.

A link between  $NO_x$ -ROG sensitivity and the ratio  $ROG/NO_x$  can also be derived from equations (A2) and (A4) if it is assumed that the odd hydrogen source term increases with  $P_{O_3}$ . In this case, (A2) and (A4) combine to yield the following criterion for the transition from  $NO_x$ - to ROG-sensitive chemistry:

$$\frac{k_1[RH]}{[NO_x]} = \frac{2\chi k_7 \psi_2}{\partial S_H / \partial P_{O_1}}$$
 (A9)

This is the familiar result that the transition between  $NO_x$ -sensitive and ROG-sensitive  $O_3$  in polluted environments occurs at a fixed  $ROG/NO_x$  ratio, with a higher ratio corresponding to  $NO_x$ -sensitive  $O_3$ . The requisite assumption  $(S_H \sim PO_3)$  is valid only in the limiting case of high RH and  $NO_x$ . The product  $k_1[RH]$  in (A9) implicitly represents a reactivity-weighted sum of ROG because  $k_1[RH]$  represents the the sum of many individual RH + OH reactions [see *Chameides et al.*, 1992].

It is useful to compare this formulation of  $NO_x$ - and ROG-sensitive photochemistry with the  $NO_x$ - and radical-limited regimes of *Kleinman* [1991, 1994]. Kleinman's radical-limited regime represents a situation in which the source term for odd hydrogen  $(S_H)$  is smaller than the  $NO_x$  source  $(S_N)$ , i.e.,  $S_H < S_N$ , and is characterized by continually increasing  $NO_x$ , very low OH, and no sigifnicant photochemical production of  $O_3$ . Sources and sinks for  $NO_x$  are in equilibrium in Kleinman's calculation  $(S_N = P_{HNO_3} + P_{RN})$  unless  $S_H > S_N$ . In these terms, ROG-sensitive chemistry is associated with the criteria (from (A6))

$$S_{\rm H} < \xi S_{\rm N}$$
 (A10a)

$$\xi = 1 + \frac{P_{\text{HNO}_3}}{P_{\text{HNO}_3} + P_{\text{RN}}}$$
 (A10b)

The region where  $S_N < S_H < \xi S_N$  is of special interest for urban photochemistry, because it includes significant OH-driven chemistry and production of  $O_3$  but the rate of ozone production is sensitive to ROG rather than to  $NO_x$ . True  $NO_x$ -sensitive chemistry only occurs when  $S_H > \xi S_N$ . The inclusion of  $NO_x$ - and ROG-sensitive regimes as defined here appears to represent a significant addition to Kleinman's formulation. The  $NO_x$ - and ROG-sensitive regime defined here are also associated with the multiple steady state solution hypothesized by *Kleinman* [1994].

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