# The Distribution of the Hydroxyl Radical in the Troposphere

By Jack Fishman Paul J. Crutzen

Department of Atmospheric Science Colorado State University Fort Collins, Colorado



### Reachbutton as me Hydraxyl Radical in the Troposphere

ing s Bodis Brittmahn Brock and size m

- Populition I among a California, Sanana 1990 - Amain Stitut I, niwar ang 1991 - California Antonao



### THE DISTRIBUTION OF THE HYDROXYL RADICAL IN THE TROPOSPHERE

by

Jack Fishman

and

Paul J. Crutzen

Preparation of this report has been financially supported by Environmental Protection Agency Grant No. R804921-01

Department of Atmospheric Science Colorado State University Fort Collins, Colorado January, 1978

Atmospheric Science Paper No. 284

THE DESTRICTION OF THE REPORTS ADDRESS OF THE TROPOSYMENE

and the man of the manufactory

Bud peak rinkwithily supported by

Gram Roy R964921-C

because of Amorphysics in Sciences

Februari all schutte abarrated

Sharolol , malikal 170%

James 1977

ACCURATE CONCERNMENT STORE SALES

#### Abstract

A quasi-steady state photochemical numerical model is developed to calculate a two-dimensional distribution of the hydroxyl (OH) radical in the troposphere. The diurnally, seasonally averaged global value of OH derived by this model is  $3 \times 10^5$  cm<sup>-3</sup> which is several times lower than the number computed previously by other models, but is in good agreement with the value inferred from the analysis of the tropospheric distribution of methyl chloroform. Likewise, the effects of the computed OH distribution on the tropospheric budgets of ozone and carbon monoxide are not inconsistent with this lower computed value.

One important result of this research is the detailed analysis of the distribution of tropospheric ozone in the Southern Hemisphere. Our work shows that there is a considerable difference in the tropospheric ozone patterns of the two hemispheres and that through the analysis of the likely photochemistry occurring in the troposphere, a significant source of tropospheric ozone may exist in the Northern Hemisphere due to carbon monoxide oxidation. Future research efforts will be devoted to the meteorological dynamics of the two hemispheres to try to distinguish if these physical processes are similarly able to explain the interhemispheric differences in tropospheric ozone.

As a result of the lesser amount of ozone found in the Southern Hemisphere troposphere, our calculations do not indicate that there should be substantially higher concentrations of OH in that hemisphere as had been previously speculated (Crutzen and Fishman, 1977; Singh, 1977b).

ii

By performing a sensitivity study on the budgets of tropospheric ozone and carbon monoxide, our model results indicate that there should be very low background concentrations of the nitrogen oxides  $(NO \text{ and } NO_2)$  in the troposphere. In addition, a relatively fast heterogeneous removal rate for several gases (e.g., hydrogen peroxide and nitric acid) is most likely necessary to obtain an understanding of the overall photochemistry occurring in the troposphere.

### Table of Contents

Page

Abstra	ct					
Acknow	ledgements					
I.	INTRODUCTION					
II.	THE PHOTOCHEMISTRY OF ODD HYDROGEN RADICALS 4					
III.	CALCULATION OF ODD HYDROGEN RADICAL NUMBER DENSITIES 12					
IV.	INPUT DATA					
V.	TROPOSPHERIC OH AND AN EXAMINATION OF THE EFFECTS OF CERTAIN MODEL PARAMETERS ON ITS DISTRIBUTION					
VI.	TROPOSPHERIC RESIDENCE TIME OF METHYL CHLOROFORM 61					
VII.	THE PRODUCTION RATE OF METASTABLE ATOMIC OXYGEN 64					
VIII.	SUMMARY AND CONCLUSIONS					
References						

#### Acknowledgements

We wish to thank Susan Solomon of the Department of Chemistry, University of California, Berkeley, for her help in the analysis of the Southern Hemisphere tropospheric ozone data which played an integral part in the formulation of this report. Susan worked with us at the National Center for Atmospheric Research (NCAR) in Boulder, Colorado, as a UCAR (University Corporation for Atmospheric Research) Fellowship Participant during the Summer of 1977. We also acknowledge the cooperation of Russ Dickerson and Donald Stedman, both of the Department of Chemistry at the University of Michigan, who made available to us some of their preliminary data for comparison in this report. The clerical competence of Julie Wilson is also much appreciated.

The computations for this study were performed at the NCAR Computing Facility in cooperation with the Air Quality Division of NCAR.

This research was partially funded by Environmental Protection Agency Grant R804921-01.

#### I. INTRODUCTION

The abundance of the hydroxyl (OH) radical in the troposphere is one of the primary factors to be considered in the determination of the removal rate of many trace gases in this region. In addition to attacking naturally-occurring species such as methane, OH reactions likewise provide the major sink for many anthropogenically released compounds. In particular, most of the chlorine contained in many commonly used chlorocarbon chemicals (e.g., methyl chloroform, perchloroethylene, etc.) is believed to be removed in the troposphere through reaction sequences initiated by hydroxyl radical attack. If these chemicals did diffuse into the stratosphere, ultraviolet radiation could photolyze them where free chlorine atoms (products of the photolysis) could cause catalytic destruction of ozone through the sequence (e.g., see Stolarski and Cicerone, 1974; Molina and Rowland, 1974; Crutzen et al., 1977):

 $C1 + 0_{3} \rightarrow C10 + 0_{2}$   $C10 + 0 \rightarrow C1 + 0_{2}$   $0_{3} + 0 \rightarrow 20_{2}.$ 

This discussion focuses on the global distribution of OH in the troposphere and its relevance to the fates of several industrially released chemicals. Whereas previously accepted estimates of average OH concentration have been on the order of 10 to 30 x  $10^5$  cm<sup>-3</sup> (see Singh, 1977a), the results computed by the model described in this study calculated average OH concentrations on the order of 3 x  $10^5$  cm<sup>-3</sup>. This number is consistent with the calculations derived in consideration of the CO budget (Crutzen and Fishman, 1977), and the tropospheric methyl chloroform (CH<sub>3</sub>CCl<sub>3</sub>)

distribution (Singh, 1977a). Although budget considerations of the above gases support the lower OH concentrations presented in this study, direct computation using available photokinetic information had always suggested the higher OH concentrations (e.g., Levy, 1972, 1973; Chang and Wuebbles, 1976).

The lower computed hydroxyl radical number densities are partially the result of some new information about quantum yields and reaction rate kinetics which influence OH directly. The experiments by Cox <u>et al</u>. (1976a), Sie <u>et al</u>. (1976) and Chan <u>et al</u>. (1977) have shown that the primary removal process of OH in the atmosphere,

$$CO + OH \rightarrow CO_2 + H$$
,

exhibits a pressure dependence. In particular, Chan <u>et al</u>. (1977) give a reaction rate constant at lower tropospheric pressures ( $\sim$ l atm) which is more than twice that which was previously accepted (Hampson and Garvin, 1975). Furthermore, recent laboratory studies (Arnold <u>et al</u>., 1977; Moortgat <u>et al</u>., 1977) also have shown that the quantum yield of metastable atomic oxygen,  $O(^{1}D)$  from ozone photolysis between 300 and 320 nm decreases with lower temperature. Previously, calculations had been carried out with quantum yield data found at room temperature (e.g., Moortgat and Warneck, 1975). In the following sections, the effects of these new data as well as some other factors which influence OH radical number density in the troposphere will be examined. The composition of the troposphere being considered in this study does not include the photochemistry of hydrocarbons other than methane and the products of methane oxidation. Future modelling efforts tentatively will include these more complex

#### II. THE PHOTOCHEMISTRY OF ODD HYDROGEN RADICALS

The odd hydrogen radicals in the troposphere consist primarily of the hydroxyl (OH) radical and the perhydroxyl (HO<sub>2</sub>) radical. These reactants exist only during daytime since they are formed either by photolysis or by reactions involving very short-lived products of photolysis. The odd hydrogen radical-producing reactions considered in these calculations are:

> R1  $H_2O + O(^1D) \rightarrow 2$  OH R2  $H_2 + O(^1D) \rightarrow H + OH$ R3  $CH_4 + O(^1D) \rightarrow CH_3 + OH$ R5  $CH_3OOH + h\nu \rightarrow CH_3O + OH$ R8  $CH_3O + O_2 \rightarrow CH_2O + HO_2$ R7a  $CH_2O + h\nu \rightarrow CHO + H$ R4  $H_2O_2 + h\nu \rightarrow 2$  OH

and R6 HNO<sub>3</sub> +  $h_{\nu} \rightarrow NO_2$  + OH.

Upon examination of the above reactions, two important points must be emphasized. First of all, the production of formyl radicals, CHO, and hydrogen atoms, H, are considered as production of perhydroxyl radicals since the recombination of these species with an oxygen molecule is assumed to proceed instantaneously:

R31 HCO +  $O_2 \rightarrow CO + HO_2$ , and

R30 H +  $O_2$  + M  $\rightarrow$  H $O_2$  + M.

Secondly, except for the first three reactions which involve the metastable oxygen atom,  $O(^{1}D)$ , the remaining chemical reactions involve reactants whose formation requires the consumption of an odd hydrogen radical. For example, the formation of nitric acid, HNO<sub>3</sub>, takes place only through

the reaction:

R16 OH + NO<sub>2</sub> (+M)  $\rightarrow$  HNO<sub>2</sub> (+M).

Thus, it appears that the actual amount of odd hydrogen radical production in the troposphere is most dependent on the amount of  $O(^{1}D)$  present. In turn, the amount of metastable oxygen atoms in the troposphere is controlled by the photolysis of ozone:

R22 
$$0_3 + h\nu \rightarrow 0_2 + 0(^1D), \lambda < 330 \text{ nm}.$$

To determine the production of metastable oxygen atoms, the following must be known:

- 1. ozone concentration,
- 2. absorption cross section data of 0,
- 3. quantum yield data between 300 and 320 nm, and
- 4. integrated ozone column density.

A detailed examination of these quantities will be presented later in this report.

Loss of odd hydrogen radicals in the troposphere occurs by the reactions:

R21  $CH_{3}OOH + OH \rightarrow CH_{3}O_{2} + H_{2}O$ , R32  $OH + OH \rightarrow H_{2}O + O$ , R19  $HNO_{3} + OH \rightarrow NO_{3} + H_{2}O$ , R18  $CH_{4} + OH \rightarrow CH_{3} + H_{2}O$ , R17  $OH + HO_{2} \rightarrow H_{2}O + O_{2}$ , R20  $HO_{2} + HO_{2} \rightarrow H_{2}O_{2} + O_{2}$ , R20a  $HO_{2} + CH_{3}O_{2} \rightarrow CH_{3}OOH + O_{2}$ , and R16  $OH + NO_{2}(+M) \rightarrow HNO_{3}(+M)$ .

Of the above reactions, only R32 and R17 yield products which cannot readily reform odd hydrogen radicals and the loss of radicals due to the latter reaction is two to three orders of magnitude greater than the loss created by R32 in the troposphere. For this reason, R32 can be excluded from the reaction sequence without altering the results. This is not to say, however, that the other reactions are not important in the determination of OH abundance. An example of this is the recycling of OH through nitric acid formation and photolysis:

R16 OH + NO<sub>2</sub>(+M)  $\rightarrow$  HNO<sub>3</sub>(+M) and

R6 HNO<sub>3</sub> +  $h\nu \rightarrow OH + NO_2$ .

Near the ground where the  $HNO_3$  photolysis is relatively low, the above cycle acts as a sink for hydroxyl radicals since heterogeneous removal of nitric acid in this region is a more dominant process than photodissociation. However, in the upper troposphere, where the prescribed amount of  $NO_2$  is less, where heterogeneous processes play a less important role, and where the photolysis rate of  $HNO_3$  is increased, this cycle comes much more into balance and the amount of OH returned by nitric acid photodissociation can even exceed the OH sink brought about by  $HNO_3$  formation. A comprehensive discussion of the effects of heterogeneous processes on the odd hydrogen and odd nitrogen distribution in the troposphere is given by Fishman (1977).

Finally, the last set of reactions which goes into the calculation of tropospheric odd hydrogen radicals are the conversion reactions which are important for determining the ratio of hydroxyl to perhydroxyl radicals. The two reactions which convert  $HO_2$  to OH involve nitrogen oxide and ozone:

R14 HO<sub>2</sub> + NO  $\rightarrow$  OH + NO<sub>2</sub> and R15 HO<sub>2</sub> + O<sub>3</sub>  $\rightarrow$  OH + 2 O<sub>2</sub>. Conversion of OH to HO2 occurs through:

R9a OH + CO  $\rightarrow$  H + CO<sub>2</sub> R9b OH + CO + M  $\rightarrow$  CO<sub>2</sub> + H + M R11 OH + O<sub>3</sub>  $\rightarrow$  HO<sub>2</sub> + O<sub>2</sub> R13 OH + CH<sub>2</sub>O  $\rightarrow$  CHO + H<sub>2</sub>O R10 OH + H<sub>2</sub>  $\rightarrow$  H + H<sub>2</sub>O, and R12 OH + H<sub>2</sub>O<sub>2</sub>  $\rightarrow$  HO<sub>2</sub> + H<sub>2</sub>O.

In the mid-latitude, lower troposphere where anthropogenic emissions determine the amount of carbon monoxide and nitrogen oxides in the air, the ratio of the concentrations of these reactants are most important in the determination of the OH to  $HO_2$  ratio.

A schematic representation of the chemical processes affecting tropospheric odd hydrogen radicals is shown in Figure 1. The rate constants governing these reactions are summarized in Table 1.



ODD HYDROGEN PHOTOCHEMISTRY

## TABLE 1

## Reactions and Rate Constants

	Reaction	Rate constant	Reference
R1	$H_2O + O(^1D) \rightarrow 2 OH$	2.3(-10)	NASA (1977)
R2	$H_2 + O(^1D) \rightarrow H + OH$	9.9(-11)	NASA (1977)
R3	$CH_4 + O(^1D) \rightarrow CH_3 + OH$	1.3(-10)	NASA (1977)
R4	$H_2O_2 + hv \rightarrow 2 \text{ OH}$	j <sub>4</sub>	Molina <u>et</u> <u>al</u> . (1977)
R5	$CH_3OOH + h\nu \rightarrow CH_3O + OH$	$j_5 = j_4$ (assumed)	
R6	$HNO_3 + hv \rightarrow NO_2 + OH$	j <sub>6</sub>	Johnston and Graham (1973)
R7a	$CH_2O + h\nu \rightarrow CHO + H$	3.7(-5)	Calvert <u>et</u> <u>al</u> . (1972)
R7b	$CH_2O + hv \rightarrow H_2 + CO$	1.1(-4)	Calvert <u>et</u> <u>al</u> . (1972)
R8	$CH_3O + O_2 \rightarrow CH_2O + HO_2$	1.6(-13) exp (-3300/T)	NASA (1977)
R9a	$CO + OH \rightarrow CO_2 + H$	1.4(-13)	Hampson and Garvin (1975)
R9b	$CO + OH + M \rightarrow CO_2 + H + M$	7.3(-33)	Chan <u>et al</u> . (1977)
R10	$H_2 + OH \rightarrow H_2O + H$	1.8(-11) exp (-2330/T)	Hampson and Garvin (1975)
R11	$0_3 + 0H \rightarrow 0_2 + H0_2$	1.5(-12) exp (-1000/T)	NASA (1977)
R12	$H_2O_2 + OH \rightarrow H_2O + HO_2$	1.0(-11) exp (-750/T)	NASA (1977)
R13	$CH_2O + OH \rightarrow H_2O + CHO$	3.0(-11) exp (-250/T)	NASA (1977)
R14	$\text{NO} + \text{HO}_2 \rightarrow \text{NO}_2 + \text{OH}$	8.0(-12)	Howard and Evenson (1977)
R15	$0_3 + H0_2 \rightarrow 2 0_2 + OH$	1.0(-13) exp (-1525/T)	NASA (1977)
R16	$NO_2 + OH(+M) \rightarrow HNO_3(+M)$		NASA (1977)
R17	$HO_2 + OH \rightarrow H_2O + O_2$	5.1(-11)	Burrows <u>et</u> <u>al</u> . (1977)

# TABLE 1 (cont'd)

	Reaction	Rate constant	Reference
R18	$CH_4 + OH \rightarrow CH_3 + H_2O$	2.4(-12) exp (-1710/T)	NASA (1977)
R19	$HNO_3 + OH \rightarrow NO_3 + H_2O$	8.0(-14)	NASA (1977)
R20	$HO_2 + HO_2 \rightarrow H_2O_2 + O_2$	5.0(-11) exp (-500/T)	NASA (1977)
R20a	$CH_{3}O_{2} + HO_{2} \rightarrow CH_{3}OOH + O_{2}$	$k_{20a} = k_{20}$	assumed
R21	$CH_{3}OOH + OH \rightarrow CH_{3}O_{2} + H_{2}O$	$k_{21} = k_{17}$	assumed
R22	$0_3 + h\nu \rightarrow 0_2 + 0(^1D)$	See text	
R23	$O(^{1}D) + M \rightarrow O + M$	5.0(-11)	NASA (1977)
R24	$NO + O_3 \rightarrow NO_2 + O_2$	2.1(-12) exp (-1450/T)	NASA (1977)
R25	$NO_2 + hv \rightarrow NO + O$	j <sub>25</sub>	Hampson and Garvin (1975)
R26a	$H_2O_2 \rightarrow heterogeneous$ removal	2.0(-5) - 2.4(-6) x f(z)	See text
R26b	CH <sub>3</sub> 00H → heterogeneous removal	$2.0(-5) - 2.4(-6) \times f(z)$	See text
R27	$CH_3CCl_3 + OH \rightarrow H_2O + CH_2CCl_3$	3.5(-12) exp (-1562/T)	NASA (1977)
R28	$0 + 0_2 + M \rightarrow 0_3 + M$	6.6(-35) exp (-510/T)	Hampson and Garvin (1975)
R29	$CH_3 + O_2 + M \rightarrow CH_3O_2 + M$	2.6(-31)	Hampson and Garvin (1975)
R30	$H + O_2 + M \rightarrow HO_2 + M$	6.7(-33) exp (290/T)	Hampson and Garvin (1975)
R31	$CHO + O_2 \rightarrow CO + HO_2$	6.0(-12)	NASA (1977)
R32	$OH + OH \rightarrow H_2O + O$	1.0(-11) exp (-550/T)	NASA (1977)
R33	$CH_3O_2 + NO \rightarrow CH_3O + NO_2$	$k_{33} = k_{14}$	Assumed

### TABLE 1 (cont'd)

Table 1. Reactions and reaction rate constants used in this study.

Most rate constants are as recommended for a recent Chloro-fluoromethane Assessment Workshop (NASA, 1977).

 $f(z) = \exp \left[-0.46 \text{ (k-5)}\right], k > kms = 1.0, k \le 5 \text{ kms}.$ Numbers in parenthesis indicate to powers of ten.

Units for unimolecular rate constants are  $s^{-1}$ ; bimolecular are  $cm^3$  mol.<sup>-1</sup> $s^{-1}$ ; trimolecular are  $cm^3$  mol.<sup>-1</sup> $s^{-1}$ .

#### III. CALCULATION OF ODD HYDROGEN RADICAL NUMBER DENSITIES

Because of the highly reactive nature of these radicals, it is assumed that each of them is in a photo-stationary state. By doing so, the assumption is made that  $d[OH]/dt = d[HO_2]/dt = 0$ . If we let P be the photochemical production rate of X (where X represents either OH or HO<sub>2</sub>) and D the destruction rate of X, the above assumption permits

$$\frac{d[X]}{dt} = P - [X] \cdot D = 0.$$
<sup>(1)</sup>

Furthermore, we couple the individual expressions of (1) for OH and  $HO_2$  which produces the following set of equations:

$$P_1 + P_2[HO_2] - D_1[OH] = 0$$
 and (2)

$$P_3 + P_4[OH] - D_2[HO_2] - D_3[HO_2]^2 = 0.$$
 (3)

In the above equations,  $P_1$  is the set of reactions which produce OH without consuming a perhydroxyl radical;  $P_2$  consists of the reactions which convert HO<sub>2</sub> to OH;  $D_1$  is the destruction rate of OH;  $P_2$  and  $P_4$  are the counterparts of  $P_1$  and  $P_2$ , but for the HO<sub>2</sub> radical;  $D_2$  and  $D_3$  are the destruction frequencies of HO<sub>2</sub>.

For these particular calculations, the following relations are defined for the various terms in Eqs. (2) and (3):

$$P_{1} = [o(^{1}D)] \cdot \{2 \cdot [H_{2}O] \cdot k_{1} + [H_{2}] \cdot k_{2} + [CH_{4}] \cdot k_{3}\}$$
  
+ 2 \cdot [H\_{2}O\_{2}] \cdot j\_{4} + [CH\_{3}OOH] \cdot j\_{5} + [HNO\_{3}] \cdot j\_{6}  
$$P_{2} = [NO] \cdot k_{14} + [O_{3}] \cdot k_{15}$$

$$\begin{split} \mathtt{D}_{1} &= [\mathtt{HO}_{2}] \cdot \mathtt{k}_{17} + [\mathtt{NO}_{2}] \cdot \mathtt{k}_{16} + [\mathtt{CH}_{3}\mathtt{OOH}] \cdot \mathtt{k}_{21} \\ &+ [\mathtt{CH}_{4}] \cdot \mathtt{k}_{18} + [\mathtt{HNO}_{3}] \cdot \mathtt{k}_{19} + [\mathtt{CO}] \cdot \{\mathtt{k}_{9a} + [\mathtt{M}] \, \mathtt{k}_{9b}\} \\ &+ [\mathtt{O}_{3}] \cdot \mathtt{k}_{11} + [\mathtt{CH}_{2}\mathtt{O}] \cdot \mathtt{k}_{13} + [\mathtt{H}_{2}] \cdot \mathtt{k}_{10} + [\mathtt{H}_{2}\mathtt{O}_{2}] \cdot \mathtt{k}_{12} \\ \mathtt{P}_{3} &= [\mathtt{H}_{2}] \cdot [\mathtt{O}(^{1}\mathtt{D})] \cdot \mathtt{k}_{2} + [\mathtt{CH}_{3}\mathtt{O}] \cdot [\mathtt{O}_{2}] \cdot \mathtt{k}_{8} + 2 \cdot [\mathtt{CH}_{2}\mathtt{O}] \cdot \mathtt{j}_{7a} \\ \mathtt{P}_{4} &= [\mathtt{CO}] \cdot \{\mathtt{k}_{9a} + [\mathtt{M}] \cdot \mathtt{k}_{9b}\} + [\mathtt{O}_{3}] \cdot \mathtt{k}_{11} + [\mathtt{H}_{2}] \cdot \mathtt{k}_{10} \\ &+ [\mathtt{H}_{2}\mathtt{O}_{2}] \cdot \mathtt{k}_{12} \\ \mathtt{D}_{2} &= [\mathtt{NO}] \cdot \mathtt{k}_{14} + [\mathtt{O}_{3}] \cdot \mathtt{k}_{15} + [\mathtt{CH}_{3}\mathtt{O}_{2}] \cdot \mathtt{k}_{20a} + [\mathtt{OH}] \cdot \mathtt{k}_{17} \\ \mathtt{D}_{3} &= 2 \cdot \mathtt{k}_{20} . \end{split}$$

The expression for [OH] in Equation (2):

$$[OH] = \frac{P_2 + P_2[HO_2]}{D_1}$$
(2a)

can be substituted into Equation (3) giving:

$$- (P_3 + \frac{P_4P_1}{D_1} + (D_2 - \frac{P_4P_2}{D_1})[HO_2] + D_3[HO_2]^2 = 0$$
(3a)

where  $[HO_2]$  can be solved directly using the standard quadratic formula. Once  $[HO_2]$  is computed explicitly, [OH] is readily found from Eq (2a). Solutions to Eqs. (2a) and (3a) are iterated five times which produces differences between successive iterations comparable to the accuracy of the computer.

### IV. INPUT DATA

To compute the global distribution of odd hydrogen radicals, input distributions of several trace constituents must be known; these gases include:

- 1. ozone,
- 2. water vapor,
- 3. carbon monoxide, and
- 4. nitrogen oxides.

The distribution of tropospheric ozone and water vapor are important since the amount of each of these species determines the amount of initial production of radicals through the sequence of reactions

R22 
$$O_3 + hv \rightarrow O(^1D) + O_2, \lambda < 320 \text{ nm}$$
  
and R1  $O(^1D) + H_2O \rightarrow 2 \text{ OH}.$ 

The amount of CO and NO are important since the relative concentrations of these gases determine the OH to  $HO_2$  ratio through the conversion reactions:

R9a(b) OH + CO(+M)  $\rightarrow$  CO<sub>2</sub> + H(+M)

(R30  $H + O_2 + M \rightarrow HO_2 + M$ )

and R14 HO<sub>2</sub> + NO  $\rightarrow$  OH + NO<sub>2</sub>.

Unfortunately, only the global distribution of CO had been published in the literature (Seiler, 1974) before this research was initiated. Data for tropospheric ozone in the Northern Hemisphere were available from Chatfield and Harrison (1977b) and Northern Hemisphere water vapor content was given in U.S. Standard Atmosphere Supplement (1966). The latter set of data did not yield enough temporal resolution and was revised in later calculations using the climatological data available at the National Center for Atmospheric Research. At present, a good representation of tropospheric  $NO_x$  still does not exist. A complete description of the input data used for the model calculations is given in the following sections.

Until recently, a good description of the distribution of tropospheric ozone with respect to latitude, altitude, and season had not been published. Although no specific observational program was ever established to make such a data set available, the efforts of Chatfield and Harrison (1977a, 1977b) produced a comprehensive analysis of existing ozonesonde data taken by the North American Ozonesonde Network (primarily in the 1960's). Thus, we have used the data given in Chatfield and Harrison (1977b) as our input for the calculations described previously.

Because a comprehensive analysis of tropospheric ozone in the Southern Hemisphere was not available prior to this study, substantial effort was devoted to compiling such a data set. The primary sources for these data were <u>Ozone Data for the World</u>, a bimonthly publication of the Canadian Department of Transportation dating back to 1962 and an analysis of ozonesonde data from Aspendale by Pittock (1974). Whereas the Aspendale data consisted of over 400 ozonesondes, only about 170 soundings were available from other sites in the Southern Hemisphere. This fact indicates that the data to be presented may not be as representative of true seasonal and latitudinal averages as one would desire. Nevertheless, we contend that this data is probably the most comprehensive that exists and that the situation cannot be improved until a comprehensive sampling program in the Southern Hemisphere is developed.

Ozone Data for the World gives five Southern Hemisphere sites which have launched ozonesondes in the 1960's and 1970's in addition to

occasional soundings available from ships. The data analyzed in this study primarily consisted of soundings taken from the following stations: Canton Island (3°S), La Paz (16°S), Aspendale (38°S), Christchurch (43°S), and Syowa (69°S). The Aspendale data previously had been analyzed by Pittock (1974) and his monthly and altitude distribution has been used as part of our analysis of the entire Southern Hemisphere. Because of the sparse nature of the data available from the remaining stations, all soundings were classified into threemonth (seasonal) groups to minimize the effects of any one anomalous profile.

The meridional distribution of ozone averaged over the entire year is depicted in Fig. 2. The tropospheric ozone distributions representative of the months of January and July are presented in Figs. 3a and 3b, respectively. Note that the units of these analyses are expressed in volume mixing ratio concentrations and that volume mixing ratios greater than 100 parts per billion (ppbv) are assumed to be stratospheric air.

Figure 4a is presented to show the summer-winter ozone profile difference in the mid-latitudes for the Northern Hemisphere. Of primary importance is the fact that 20 to 30% more ozone is present below seven kilometers in the summer even though stratospherictropospheric exchange processes are more prevalent in the winter (e.g., see Danielsen, 1968). This suggests that more photochemical production of ozone in the lower troposphere may take place at these latitudes in the summer. Although a similar phenomenon takes place at 15°N, it is not as clearly seen at 5°N (Fig. 4b). The reason for this phenomenon is difficult to assess without looking at both the



MEPUDIONAL OZONE DISTRIBUTION

•

Fig. 2



Fig. 3a





Fig. 4a



Fig. 4b

meteorological dynamics and photochemistry in much detail. The analysis of the tropospheric ozone budget presented in Section V does support the hypothesis that ozone may be produced photochemically in Northern Hemisphere mid-latitudes (Fishman and Crutzen, 1977) and transported to lower latitudes, but one must examine troposphere-stratosphere exchange properties as a function of both season and latitude to determine what effect this mechanism would have on the data shown in Figs 3a and 3b. Future research efforts will be devoted to obtaining a better picture of the meteorological dynamics in order to resolve what tropospheric ozone distribution one might expect considering stratosphere-troposphere exchange processes only.

The seasonal variation in the Southern Hemisphere mid-latitudes (40°S) is shown in Fig. 5. Unlike the variation in the Northern Hemisphere mid-latitudes (Fig. 4a), more ozone is observed in the winter than in the summer at this latitude which suggests that mixing may play the predominant role in determining the tropospheric ozone distribution in all seasons. If photochemical production were the dominant source of ozone in this region, one might expect more tropospheric ozone in the summer when this mechanism is enhanced. However, the meteorological dynamics of the Southern Hemisphere must be examined in detail in future research to determine if this seasonal variation of tropospheric ozone is consistent with large scale transport processes in this hemisphere. Another feature of these profiles is that the total seasonal variation in the Southern Hemisphere midlatitudes is less than in the Northern Hemisphere. This point has previously been brought forth by Pittock (1974) who compared his data with Dütsch (1970).



Fig. 5

Figure 6 depicts the interhemispheric difference in the midlatitude tropospheric ozone profiles for respective summer seasons. Particularly noteworthy in this comparison is the fact that ozone concentrations and the shapes of these profiles above 10 kms are nearly identical. On the other hand, the Northern Hemisphere average summer ozone concentrations are 15 to 20 ppb greater (30-70%) than the corresponding concentrations in the Southern Hemisphere between one and eight kms.

Figure 7 shows that when the ozone data are averaged over the entire year, we observe more ozone in the Northern Hemisphere troposphere at low latitudes. At  $5^{\circ}$ , Northern Hemisphere concentrations are 7 to 13 ppb greater (30-100%) between 1 and 8 kms whereas the profiles converge at 12 kms. Similarly, at  $15^{\circ}$ , we find 5 to 12 ppb more (20-65%) ozone in the Northern Hemisphere mid-troposphere even though there are slightly higher concentrations in the Southern Hemisphere above 10 kms.

Lastly, we point out that the similarity of ozone concentrations in the upper troposphere at all corresponding latitudes of the two hemispheres makes the differences observed between the low and middle troposphere of the two hemispheres more credible. If these hemispheric differences are real (and not merely the product of inadequate data or faulty analysis), they suggest that there are important differences in the photochemical and/or physical processes taking place in these hemispheres. These points will be discussed in the next section.







### V. TROPOSPHERIC OH AND AN EXAMINATION OF THE EFFECTS OF CERTAIN MODEL PARAMETERS ON ITS DISTRIBUTION

Whereas the amount of ozone is directly responsible for determining the amount of odd hydrogen radicals produced in the troposphere, we recall that the relative amounts of carbon monoxide and nitrogen oxides ( $NO_x$ ) primarily determine the ratio of OH to HO<sub>2</sub> through the reactions:

> R9a(b) OH + CO(+M)  $\rightarrow$  H + CO<sub>2</sub>(+M), (R30 H + O<sub>2</sub> + M  $\rightarrow$  HO<sub>2</sub> + M), and R14 HO<sub>2</sub> + NO  $\rightarrow$  OH + NO<sub>2</sub>.

The most comprehensive analysis of CO has been carried out by Seiler (1974); his data are used as input for the current study. Figure 8 depicts four tropospheric CO profiles representative of mid- and low latitudes in both hemispheres. Particularly noteworthy is the uniform decrease in surface and low altitude CO concentrations as one progresses southward. At low altitudes, the amount of CO at 45<sup>o</sup>N is more than three times larger than the concentration at 45<sup>o</sup>S; this factor decreases to two in the middle troposphere whereas equal concentrations are found at the mid-latitude tropopause height (12 km) in the two hemispheres.

At  $15^{\circ}$ , we note that there is 70% more CO near the surface in the Northern Hemisphere than in the Southern Hemisphere but that at 6 kms, less than a 10% difference is observed. Recalling that the hemispheric difference of ozone at these latitudes is 20-65% at these altitudes, one could estimate that the calculated amount of OH at these two latitudes would be quite comparable; this speculation is borne out by the calculated tropospheric average concentrations:  $5.1 \times 10^5$  cm<sup>-3</sup>




at  $15^{\circ}N$  vs.  $4.5 \times 10^{5}$  cm<sup>-3</sup> at  $15^{\circ}S$ . Using the fact that there is more OH at  $15^{\circ}N$  than at  $15^{\circ}S$ , one would expect a greater difference in OH at  $5^{\circ}N$  and  $5^{\circ}S$  since the CO differences are smaller (0-25%) and the ozone differences are larger (30-100%) than at  $15^{\circ}$ . The calculations indicate a 40% difference:  $5.8 \times 10^{5}$  cm<sup>-3</sup> at  $5^{\circ}N$  and  $4.1 \times 10^{5}$  cm<sup>-3</sup> at  $5^{\circ}S$ . [Note that for all the latitudes discussed above (and for all latitudes  $15^{\circ}N$  and southward), the concentration of  $NO_{x}$  at every latitude is the same for any corresponding altitude.]

The above analysis is presented to show why we should not be surprised by the fact that there is slightly more OH in the Northern Hemisphere  $(3.3 \times 10^5 \text{ cm}^{-3})$  than in the Southern Hemisphere troposphere  $(2.9 \times 10^5 \text{ cm}^{-3})$ . This finding is contrary to the analysis of Crutzen and Fishman (1977) who hypothesized that the average OH concentration should be 2.7 times larger in the Southern Hemisphere because of less CO in this hemisphere, and Singh (1977b) who analyzed the tropospheric distribution of methyl chloroform to suggest that this ratio be between 1.6 and 3.0. The primary reason for the discrepancy noted here can be directly attributed to the difference in the amount of tropospheric ozone in the two hemispheres.

Figure 9 shows the diurnally and monthly averaged OH distribution in the troposphere. Although the highest average OH concentration is found in the Southern Hemisphere (15<sup>°</sup>S at the surface), a much stronger intrahemispheric gradient exists both latitudinally and vertically resulting in comparable amounts of OH in both hemispheres. To see how the computed OH distributions vary seasonably, Figs. 10a, b, c, and d are presented. As expected, more OH is computed for each respective hemisphere's summer season (about three times higher than



Fig. 9







Fig. 10b



Fig. 10c



Fig. 10d

what is found in the winter season). The high values found above 9 kms during the summer season at high latitudes can be attibuted to the very high concentrations of stratospheric ozone and the fact that these latitudes are in sunlight throughout most of the day at this time of the year.

Figures 10e, f, g, and h are presented for comparison with the few daytime OH measurements that are available. Davis et al. (1976) have observed OH number densities between 3 and 8 x  $10^6$  cm<sup>-3</sup> in October, 1975, during daytime clean air conditions between 21°N and 32°N in the mid- and upper troposphere. The values computed (Fig. 10h) in this study are about five times lower than Davis et al. (1976) have observed. Near the surface, Perner et al. (1976) measured between 0 and 7 x  $10^6$  cm<sup>-3</sup> in Julich, Germany (51°N), in late summer and early autumn. Unfortunately, their detection limit was 4 x  $10^6$  cm<sup>-3</sup>, and no OH was observed in most cases. Their finding that OH generally remains below this level is not inconsistent with our results. Lastly, the preliminary results of Campbell (1977), taken near the surface at Pullman, Washington (47°N), in August and October, 1977, range between 2 and 8 x  $10^5$  cm<sup>-3</sup> which are in relatively good agreement with this study's calculations. Unfortunately, none of the observations have concurrent measurements of ozone, water vapor, carbon monoxide, nitric oxide, nitrogen dioxide and incoming solar radiation -- all of which are quite variable and have an important effect on the modelderived OH. Therefore, it is difficult to assess the model's ability to simulate OH number densities accurately based on these few measurements and its is necessary to employ other considerations to determine the validity of the output of the model.



Fig. 10e



Fig. 10f



Fig. 10g



Fig. 10h

The most striking difference between the two hemispheres is the difference in the vertical gradients of OH. Figure 11 shows the yearly averaged OH profiles for both hemispheres which clearly depicts this difference. Although the Southern Hemisphere's lower troposphere values are considerably higher, the profiles cross at an altitude near 2 kms at which point more OH is seen in the Northern Hemisphere's mid- and upper troposphere. These higher values can be attributed to the higher tropospheric ozone concentrations found in this region. Again, it is interesting to note the very similar values found in the upper troposphere. Above 9 kms, OH begins to increase due to the enhanced ozone levels at most latitudes. The slight increase in Northern Hemisphere OH compared to Southern Hemisphere upper troposphere OH values is brought about by the slightly warmer temperatures in this region. Because conventional radiosonde dew point data were not available for these altitudes, water vapor concentrations were computed from relative humidity information (U.S. Standard Atmosphere Supplement, 1966) in the upper troposphere. Although the relative humidity was equal in both hemispheres, the warmer temperatures in the Northern Hemisphere resulted in a higher H20 number density. Thus, OH production via Rl is enhanced, yielding slightly higher OH concentrations.

Although much more can be said about the model-derived OH results presented in Figs. 9, 10, and 11, we conclude that such a discussion would be speculative at this time since the results of our computations are quite sensitive to model assumptions. Therefore, we feel that the focus of this discussion should be to examine some of the key factors which had to be prescribed in the calculations and the sensitivity of the results to these assumptions.



Because of the relatively few observations of the nitrogen oxides  $(NO_x = NO + NO_2)$  in the "clean" troposphere, the uncertainty surrounding the prescription of the concentrations of these gases must be examined in detail. Noxon (1977) presents the argument that the column abundance of NO<sub>2</sub> in the troposphere is less than 5 x  $10^{14}$  cm<sup>-2</sup> but that a true value of the column density on a global scale still is not known. Three points which hinder the assessment of true global values are:

- 1. the definition of "unpolluted" air;
- the influence of polluted air on average tropospheric values; and
  - the observation of NO<sub>2</sub> tropospheric column densities below detectable limits.

It therefore becomes quite difficult to assign tropospheric NO and  $NO_2$  profiles that are representative of global averages and we must investigate the effects of these prescribed concentrations on both the computed OH number densities and the overall photochemistry taking place in the troposphere. Lastly, we point out that since Noxon's (1977) measurements were carried out very close to sunrise and sunset, his  $NO_2$  column densities are probably the same as the  $NO_2$  column density since the NO to  $NO_2$  ratio,

$$\frac{\text{NO}}{\text{NO}_2} = \frac{j_{25}}{[O_3] \cdot k_{24}}$$

is much less than unity. The remainder of the discussion focuses on the NO  $_{\rm X}$  distribution in the troposphere, keeping in mind that the above ratio is used in the computations.

Figure 12 shows the tropospheric NO  $_{\rm X}$  profiles prescribed in the model calculations which have been presented to this point. Curve A is the NO distribution assumed for "clean" tropospheric air and is prescribed



throughout the Southern Hemisphere and southwards of  $15^{\circ}$  in the Northern Hemisphere. Curve B is the profile prescribed at  $45^{\circ}$ N and northward; intermediate profiles are assumed between  $15^{\circ}$ N and  $45^{\circ}$ N. The integrated NO<sub>x</sub> column density is 2.1 x  $10^{14}$  cm<sup>-2</sup> for A and 4.6 x  $10^{14}$  cm<sup>-2</sup> for B.

In Fig. 13, we show the sensitivity of calculated OH to assumed tropospheric NO<sub>x</sub> distributions. Plotted in this figure are average Northern Hemispheric profiles for July at noon (refer to Fig. 10g) using the following NO<sub>y</sub> distributions:

Profile I: NO<sub>x</sub> is prescribed according to Fig. 12 (Standard Model);

Profile II:  $NO_x$  shown in Fig. 12 is doubled everywhere; Profile III:  $NO_x = 0.1$  ppb  $(10^{-10} v v^{-1})$  at every point; Profile IV:  $NO_x = 0.001$  ppb  $(10^{-12} v v^{-1})$  at every point.

Doubling the amount of  $NO_x$  increases the average OH in the Northern Hemisphere by 20% from 3.3 to 3.9 x  $10^5$  cm<sup>-3</sup>. By letting the  $NO_x$  mixing ratio equal 0.1 ppb (Profile III), the total amount of tropospheric  $NO_x$ is increased by a factor of 6.5 over the amount of tropospheric  $NO_x$ used to generate Profile I; although only slightly more  $NO_x$  is prescribed at the surface, a fifty-fold increase in  $NO_x$  is seen in the low-latitude upper troposphere. The net result of this relatively high prescription of  $NO_x$  is that the average Northern Hemisphere OH is increased to  $9.5 \times 10^5$  cm<sup>-3</sup> (192% increase over Profile I). The implications of such an  $NO_x$  distribution in the troposphere will be discussed later. Lastly, by assuming a tropospheric  $NO_x$  mixing ratio of 0.001 ppb everywhere, the OH distribution depicted by Profile IV is produced. Thus, it is clear from Fig. 13 that a wide variation of tropospheric OH can be calculated



Fig. 13

from the different prescriptions of  $NO_x$ . In addition to the relatively few observations of NO and  $NO_2$ , consideration of the tropospheric budgets of other gases (namely, CO and  $O_3$ ) must be used in the determination of the "correct" tropospheric  $NO_x$  distribution. Although little is known about the latitudinal and height distributions of NO and  $NO_2$ , the budget analyses do give a crude estimate of how much total  $NO_x$  in the troposphere is necessary to be consistent with our present understanding of these budgets.

Table 2 summarizes some of the photochemical calculations in the Northern Hemisphere for the four  $NO_x$  distributions described above. In general, the more  $NO_x$  present enhances not only the concentration of OH, but the amount of carbon monoxide oxidized [from CO + OH(+M)  $\rightarrow$  $CO_2$  + H(+M)] and the amount of methane oxidized (CH<sub>4</sub> + OH  $\rightarrow$  CH<sub>3</sub> + H<sub>2</sub>O). Particularly noteworthy is the fact that both CO and CH<sub>4</sub> oxidation lead to the production of ozone in the troposphere (see Crutzen, 1974; Fishman and Crutzen, 1977), and upper limits on the photochemical production of tropospheric ozone can readily be determined.

Catalytic production of ozone initiated by OH attack on carbon monoxide occurs through the reaction sequence:

> R9a(b) C0 + OH(+M)  $\rightarrow$  CO<sub>2</sub> + H(+M) R30 H + O<sub>2</sub> + M  $\rightarrow$  HO<sub>2</sub> + M R14 HO<sub>2</sub> + NO  $\rightarrow$  OH + NO<sub>2</sub> R25 NO<sub>2</sub> + hv  $\rightarrow$  NO + O R28 O + O<sub>2</sub> + M  $\rightarrow$  O<sub>3</sub> + M CO + 2 O<sub>2</sub> + hv  $\rightarrow$  CO<sub>2</sub> + O<sub>3</sub>

Based on what we presently know about tropospheric photochemistry, the only reaction in this chain that may not possibly occur nearly 100%

# TABLE 2

Sensitivity of Model Calculations to Prescribed NO concentrations for the Northern Hemisphere  $\mathbf{x}$ 

Profile	I Standard	$\frac{\text{II}}{\text{Double NO}_{x} \text{ of I}}$	NO <sub>x</sub> = 0.1 ppb	$\frac{IV}{NO_{x}} = 0.001 \text{ ppb}$
Tropospheric $NO_x$ Column Density (10 <sup>14</sup> cm <sup>-2</sup> )	3.4	6.8	22.9	0.2
Average OH $(10^5 \text{ cm}^{-3})$	3.3	3.9	9.5	2.9
CO Oxidation $(10^{11} \text{ mol cm}^{-2} \text{ s}^{-1})$	2.4	3.0	8.5	1.8
CH <sub>4</sub> Oxidation ( $10^{11}$ mol cm <sup>-2</sup> s <sup>-1</sup> )	0.4	0.6	1.0	0.4
Upper Limit $0_3$ Production $(10^{11} \text{ mol cm}^{-2} \text{ s}^{-1})$	3.5	4.4	10.9	2.8
0 <sub>3</sub> Efficiency Production Factor	0.50	0.75	0.92	0.03
Tropospheric O <sub>3</sub> Production by Photochemistry (10 <sup>11</sup> mol cm <sup>-2</sup> s <sup>-1</sup> )	1.8	3.3	10.0	0.1
Tropospheric O <sub>3</sub> Destruction by Photochemistry (10 <sup>11</sup> mol cm <sup>-2</sup> s <sup>-1</sup> )	1.2	1.2	a <b>1.4</b> a	1.2
Net Photochemical $0_3$ Production $(10^{11} \text{ mol cm}^{-2} \text{ s}^{-1})$	+0.6	+2.1	+8.6	-1.1

Estimated  $0_3$  Destruction at the Ground (10<sup>11</sup> mol cm<sup>-2</sup> s<sup>-1</sup>) 0.4-0.8

of the time is R14. In other words, we can assume that every H produced from R9 will become HO<sub>2</sub> and that every NO<sub>2</sub> produced through R14 will photolyze to give atomic oxygen which, in turn, will find an oxygen molecule to make ozone.

An examination of Fig. 1 and Table 1 shows that the following reactions involving HO, are most likely to take place in the troposphere:

R14 NO + HO<sub>2</sub>  $\Rightarrow$  NO<sub>2</sub> + OH, R15 O<sub>3</sub> + HO<sub>2</sub>  $\Rightarrow$  2 O<sub>2</sub> + OH, R17 OH + HO<sub>2</sub>  $\Rightarrow$  H<sub>2</sub>O + O<sub>2</sub>, R20 HO<sub>2</sub> + HO<sub>2</sub>  $\Rightarrow$  H<sub>2</sub>O<sub>2</sub> + O<sub>2</sub>, and R20a CH<sub>3</sub>O<sub>2</sub> + HO<sub>2</sub>  $\Rightarrow$  CH<sub>3</sub>O<sub>2</sub>H + O<sub>2</sub>.

Using representative tropospheric concentrations and atmospheric conditions, we can look at which of the above reactions occurs most frequently; these are summarized in Table 3. The numbers show that even in an atmosphere in which  $NO_x$  concentrations are very low (<0.1 ppb), the NO + HO<sub>2</sub> reaction determines the fate of the HO<sub>2</sub> radical. However, it is clear that the percentage of HO<sub>2</sub> that follows the Rl4 path depends on the assumed NO concentration, the computed radical concentrations (OH, HO<sub>2</sub> and CH<sub>3</sub>O<sub>2</sub>) and the background ozone concentration. On the other hand, because of the extremely fast value of k<sub>14</sub> (Howard and Evenson, 1977), it is also quite evident that the other HO<sub>2</sub> reactions do not determine the path of HO<sub>2</sub> until NO<sub>x</sub> mixing ratios go below 0.01 ppb. Note that our model-assumed NO<sub>x</sub> profile (Fig. 12) does prescribe NO<sub>x</sub> mixing ratios less than this value above an altitude of 4 kms.

A similar examination of ozone production from methane oxidation can be made, but the conclusion is the same: the amount of ozone produced TABLE 3

Competition of Reactions Involving the HO2 Radical for Representative Tropospheric Conditions

Reaction	Reactant Concentration	Rate Constant	Rate of Reaction	Fraction
$HO_2 + NO \rightarrow OH + NO_2$	1x10 <sup>9</sup> cm <sup>-3</sup> (v0.04 ppb)	8x10 <sup>-12</sup>	$8 \times 10^{-3}$	0.89
$HO_2 + O_3 \rightarrow OH + 2 O_2$	1x10 <sup>12</sup> cm <sup>-3</sup> (~40 ppb)	4x10 <sup>-16</sup>	4x10 <sup>-4</sup>	0.05
$HO_2 + OH \rightarrow H_2O + O_2$	$4 \times 10^6 \text{ cm}^{-3}$	$5 \times 10^{-11}$	$2 \times 10^{-4}$	0.02
$HO_2 + HO_2 + H_2O_2 + O_2$	$4x10^8 \text{ cm}^{-3}$	8x10-13	$3x10^{-4}$	0.03
$CH_3O_2 + HO_2 + CH_3O_2H + O_2$	2x10 <sup>8</sup> cm <sup>-3</sup>	8x10 <sup>-13</sup>	$1 \times 10^{-4}$	0.01

from methane oxidation is dependent on the background concentration of the nitrogen oxides. Although the methane oxidation chemistry is more complex than CO oxidation (see Crutzen, 1974), we will assume in this study that no more than 2.5 ozone molecules can be produced from each  $CH_4$  + OH reaction. The ratio of O<sub>3</sub> produced to  $CH_4$  oxidized is most dependent of the photolytic path of formaldehyde (CH,0). If CH,0 photodissociates completely to H + CHO (R7a), as many as four ozone molecules from each methane molecule oxidized could be produced. However, using the dissociation rates shown in Table 1 for j<sub>7a</sub> and j<sub>7b</sub>, only 2.5 ozone molecules are produced from the OH attack on CH, . For the four cases given in Table 2, CO oxidation dominates  $CH_{\underline{\lambda}}$  oxidation by at least a factor of 5; therefore, the contribution to the upper limit of the amount of ozone produced by CO oxidation is at least twice as great as the ozone produced from CH, oxidation. Thus, the quantity presented in Table 2 under the category of " $O_3$  Production (Upper limit)" is the sum of "CO Oxidation" and 2.5 times "CH, Oxidation."

From the actual model results, one can determine the true "ozone efficiency production factor" for each of the four  $NO_x$  input distributions. For the examples presented here, this factor ranges from 0.03 where the  $NO_x$  mixing ratio is 0.001 ppb, to 0.92 where a 0.1 ppb  $NO_x$  mixing ratio is prescribed everywhere. Thus, it is important to note that both the upper limit of the amount of ozone production and the efficiency of the ozone production is increased when more  $NO_x$  is present. An important consequence of these calculations is that the amount of tropospheric ozone which is likely to be produced from photochemistry (~1 x  $10^{12}$ mol cm<sup>-2</sup> s<sup>-1</sup>) when the  $NO_x$  mixing ratio of 0.1 ppb (or greater) everywhere in the troposphere is greatly out of line with previous Competition of Reactions Involving the HO2 Radical for Representative Tropospheric Conditions

TABLE 3

Reaction	Reactant Concentration	Rate Constant	Rate of Reaction	Fraction
$10_2 + NO \rightarrow OH + NO_2$	1x10 <sup>9</sup> cm <sup>-3</sup> (~0.04 ppb)	$8 \times 10^{-12}$	$8 \times 10^{-3}$	0.63
$10_2 + 0_3 \div 0H + 2 0_2$	$1 \times 10^{12} \text{ cm}^{-3} (040 \text{ ppb})$	$4 \times 10^{-16}$	$4x10^{-4}$	0.03
$10_{2} + 0H + H_{2}0 + 0_{2}$	$4x10^{6}$ cm <sup>-3</sup>	$5 \times 10^{-11}$	$2x10^{-4}$	0.02
$10_{2} + H0_{3} + H_{2}0_{3} + 0_{3}$	$3 \times 10^8 \text{ cm}^{-3}$	$8 \times 10^{-12}$	$2x10^{-3}$	0.19
$H_{3}O_{2} + HO_{2} + CH_{3}O_{2}H + O_{2}$	$2 \times 10^8 \text{ cm}^{-3}$	$8 \times 10^{-12}$	$2x10^{-3}$	0.13

from methane oxidation is dependent on the background concentration of the nitrogen oxides. Although the methane oxidation chemistry is more complex than CO oxidation (see Crutzen, 1974), we will assume in this study that no more than 2.5 ozone molecules can be produced from each  $CH_{L}$  + OH reaction. The ratio of O<sub>3</sub> produced to  $CH_{L}$  oxidized is most dependent of the photolytic path of formaldehyde (CH20). If CH20 . photodissociates completely to H + CHO (R7a), as many as four ozone molecules from each methane molecule oxidized could be produced. However, using the dissociation rates shown in Table 1 for  $j_{7a}$  and  $j_{7b}$ , only 2.5 ozone molecules are produced from the OH attack on  $CH_{\rm A}$ . For the four cases given in Table 2, CO oxidation dominates  $\mathrm{CH}_{\mathrm{A}}$  oxidation by at least a factor of 5; therefore, the contribution to the upper limit of the amount of ozone produced by CO oxidation is at least twice as great as the ozone produced from  $CH_{\underline{\lambda}}$  oxidation. Thus, the quantity presented in Table 2 under the category of "0 $_3$  Production (Upper limit)" is the sum of "CO Oxidation" and 2.5 times "CH, Oxidation."

From the actual model results, one can determine the true "ozone efficiency production factor" for each of the four  $NO_x$  input distributions. For the examples presented here, this factor ranges from 0.03 where the  $NO_x$  mixing ratio is 0.001 ppb, to 0.92 where a 0.1 ppb  $NO_x$  mixing ratio is prescribed everywhere. Thus, it is important to note that both the upper limit of the amount of ozone production and the efficiency of the ozone production is increased when more  $NO_x$  is present. An important consequence of these calculations is that the amount of tropospheric ozone which is likely to be produced from photochemistry ( $v1 \ge 10^{12}$ mol cm<sup>-2</sup> s<sup>-1</sup>) when the  $NO_x$  mixing ratio of 0.1 ppb (or greater) everywhere in the troposphere is greatly out of line with previous estimates of the tropospheric ozone budget (Fishman and Crutzen, 1977; Chameides and Stedman, 1977; Fabian and Pruchniewicz, 1977). Furthermore, the amount of ozone destroyed photochemically does not vary appreciably from 1 x  $10^{11}$  mol. cm<sup>-2</sup> s<sup>-1</sup> since this quantity is primarily fixed by the amount of ozone lost through R1 and does not depend on any concentrations other than those of ozone and water vapor. Thus, an efficient ozone destruction mechanism presently not considered in any tropospheric ozone budget must dominate the ozone sink if so much tropospheric ozone is produced when an 0.1 ppb NO<sub>x</sub> mixing ratio is present everywhere in the troposphere. Presently we cannot speculate on such a destruction process and therefore conclude that such a high NO<sub>x</sub> mixing ratio probably does not exist in the troposphere.

On the other hand, the NO<sub>x</sub> distribution depicted in Fig. 12 does yield a tropospheric ozone budget which can be said to be in reasonable agreement with what is believed to the tropospheric ozone budget. In addition, these NO<sub>x</sub> concentrations are consistent with the tropospheric NO<sub>x</sub> column density observations of Noxon (1977) and the surface NO observations of Drummond (1977) for "clean" air in Wyoming.

Similarly, a tropospheric ozone budget for the Southern Hemisphere can be derived. In these calculations, the  $NO_x$  concentrations depicted by Curve A in Fig. 12 is prescribed at all latitudes (see Table 4). Comparing the photochemistry occurring in both hemispheres using this particular set of model-derived results, two very general conclusions are made:

> the net photochemical production of ozone in the Northern Hemisphere troposphere may be comparable or even greater than the amount of ozone coming through the tropopause (Danielsen and Mohnen, 1977; Fabian and Pruchnienicz, 1977); and

# TABLE 4 STATES STATES TO TABLE 4

# Model-Derived Tropospheric Budgets

	Northern Hemisphere	Southern Hemisphere
	Conservation of the second	
Tropospheric NO Column Density $(10^{14} \text{ cm}^{-2})^{\text{X}}$	3.4	2.1
Average OH $(10^5 \text{ cm}^{-3})$	3.3	2.9
CO Oxidation	2.4	1.3
CH <sub>4</sub> Oxidation	0.4	0.4
Ozone Production (Upper Limit)	3.5	2.4
Ozone Production Efficiency Factor (Nondimensional)	0.50	0.31
Tropospheric Ozone Produced by Photochemistry	1.8	0.7
Tropospheric Ozone Destroyed by Photochemistry	1.2	0.6
Net Photochemical Ozone Production	+0.6	+0.1
Ozone Destruction at the Ground (from Fabian and Pruchniewicz, 1977)	0.7	0.4

Note: All units are  $10^{11}$  mol cm<sup>-2</sup> s<sup>-1</sup>, unless otherwise noted.

### 2. there seems to be less net photochemical production of ozone taking place in the Southern Hemisphere.

Thus, even if the transport of ozone in both hemispheres were comparable, one would expect to find more ozone in the Northern Hemisphere due to the enhanced photochemical production. Furthermore, the difference in the photochemical production is primarily due to CO oxidation rather than  $CH_{\Delta}$  oxidation.

Lastly, in this section, we present the global CO budget originally given by Seiler (1974) but updated by the photochemical calculations of our present model (refer to Table 5). Recalling Seiler's CO budget, we note that there was a large degree of uncertainty in the amounts of CO produced and destroyed photochemically:  $15-40 \times 10^{14} \text{ g yr}^{-1}$  produced from methane oxidation and  $19-50 \times 10^{14} \text{ g yr}^{-1}$  destroyed by OH-attack. Furthermore, we observe that these mechanisms greatly dominate all other source and sink terms in Seiler's budget.

Using our model-derived results, on the other hand, the amount of CO produced from methane oxidation is only about half as large as the contribution from Seiler's anthropogic estimate. In addition, we note that the number we give under the "CH<sub>4</sub> + OH" leading in Table 3 (3.3 x  $10^{14}$  g yr<sup>-1</sup>) is an upper limit to the amount of CO produced photochemically since we must assume that every CH<sub>4</sub> oxidized results in a CO molecule produced. This assumption may not be valid if some of the peroxides or the formaldehyde produced as intermediate products of methane oxidation are rapidly removed by heterogeneous processes. Since the methane concentration is well known everywhere in the troposphere (Ehhalt, 1974), photochemical production of carbon monoxide can be increased only if the tropospheric concentrations of OH are increased.

# TABLE 5

### Global Carbon Monoxide Budget

(from Seiler, 1974, and modified by this study)

Sources  $(10^{14} \text{ g yr}^{-1})$ Global Northern Southern Hemisphere Hemisphere 1.0 6.4 5.4 Anthropogenic Oceans 1.0 0.4 0.6 0.2 0.6 0.4 Forest Burning 0.6 0.4 0.2 Oxidation of Hydrocarbons  $CH_{4} + OH$  (this study) 3.3 1.6 1.7 8.3 3.6 11.9 Total Sinks  $(10^{14} \text{ g yr}^{-1})$ 4.5 1.5 3.0 Soil Uptake 0.2 1.1 0.9 Into Stratosphere 8.9 5.0 13.9 CO + OH (this study) 19.5 12.8 7.7 Total Net

-7.6 (Budget does not balance)

However, for every OH present, more than five times CO is destroyed by OH-attack than is produced via methane oxidation. Thus, the imbalance shown in Table 5 would become worse if the average amount of OH were increased and that a revision of Seiler's budget must be considered. In future research efforts, we will examine the possibility that CO can be released by soils under certain conditions and that the amount of carbon monoxide produced by the oxidation of natural and anthropogenic hydrocarbons may be higher than Seiler had estimated.

Another fundamental unknown in atmospheric chemistry is the rate of removal of atmospheric gases by processes other than gas-phase reactions. In the present study, three gases are included in the model whose concentrations are not prescribed and which may be removed more efficiently by heterogeneous reactions than by homogeneous ones. Of these species  $(HNO_3, H_2O_2, and CH_3O_2H)$ , the tropospheric fate of hydrogen peroxide,  $H_2O_2$ , is the most critical in the calculation of the OH number density.

The method adopted for heterogeneous removal of these gases is similar to the one prescribed by Fishman and Crutzen (1977). A removal rate ( $s^{-1}$ ) is defined for the lowest five kms of the troposphere and then falls off exponentially above that level to 12 kms according to the formula:

$$k(z)_{HET} = k_{HET}^{o} \exp \left[-0.46(z-5)\right], z > 5 \text{ km},$$
  
 $k(z)_{HET} = k_{HET}^{o}, z \le 5 \text{ km},$ 

where k(z) is the heterogeneous removal rate at altitude z, and  $k_{HET}^{o}$  is the prescribed heterogeneous removal rate below 5 kms. The exponential decay with height reproduces the observations of Davidson <u>et al</u>. (1966) for removal of radioactive debris.

The prescription of  $k_{HET}^{o}$  does influence the model calculations and the sensitivity of the computations to this parameter is presented in Figures 14, 15, and Table 6. As expected, a faster heterogeneous removal rate depletes the abundance of both  $H_2O_2$  and OH since hydrogen peroxide acts as a reservoir for odd hydrogen species in the lower atmosphere. The primary problem to be resolved is to choose the correct value of  $k_{HET}^{o}$  for the model calculations. The three values discussed here are  $2 \times 10^{-5} \text{ s}^{-1}$  which was used for the "standard" model calculations (Profile A in Figures 14 and 15),  $5 \times 10^{-6} \text{ s}^{-1}$  (Profile B), and  $2.4 \times 10^{-6} \text{ s}^{-1}$ (Profile C). Unfortunately, observations of hydrogen peroxide in the unpolluted atmosphere do not exist and it is therefore impossible to tune this removal rate (which is the primary loss mechanism for  $H_2O_2$ ) to measurements. Thus, other considerations must be taken into account for the proper prescription of  $k_{HET}^{O}$ .

The most widely investigated gas whose fate is primarily controlled by heterogeneous processes is sulfur dioxide. In prior studies, the residence times computed for SO<sub>2</sub> vary by more than an order of magnitude from five days (Junge, 1960) to less than half a day (Eliassen and Saltbones, 1975; Metham, 1950). These values yield removal rates of 2.3 x  $10^{-6}$  s<sup>-1</sup> and 2.3 x  $10^{-5}$  s<sup>-1</sup>, respectively. Whereas the above studies used emission data and measurements to derive residence time, Henmi <u>et al</u>. (1977) compute SO<sub>2</sub> residence times over the eastern United States using climatological data of the mixing layer height, precipitation information and the consideration of microphysical cloud processes to produce a "scavenging" coefficient between 1 and 4 x  $10^{-5}$  s<sup>-1</sup> (SO<sub>2</sub> residence times between 6 and 24 hours). Thus, since any of the three removal rates summarized in Table 6 could be used justifiably in the calculations, other considerations must be examined.



Fig. 14

FD

SENSITIVITY OF MODEL-DERI



#### VI. TROPOSPHERIC RESIDENCE TIME OF METHYL CHLOROFORM

Because of increased usage of methyl chloroform  $(CH_3CCl_3)$  as an industrial solvent, this chemical may become a significant source of chlorine in the stratosphere. According to Crutzen <u>et al</u>. (1977), if the production of  $CH_3CCl_3$  continues to increase at its current rate (~15% per year), depletion of ozone in the stratosphere due to the chlorine contained in methyl chloroform will exceed the perturbations to the ozone layer by fluorochloromethanes by the end of this century (assuming that the latter's production rates remained fixed at 1975 levels). These estimates, however, are quite sensitive to how much methyl chloroform is removed in the troposphere through OH-attack and thus, it is quite important to know the  $CH_3CCl_3$  removal rate in the troposphere.

Calculations of the methyl chloroform tropospheric residence time before 1977 ranged between 1 and 3 years, and thus it was thought that OH-attack in the troposphere was an efficient enough mechanism to prevent much of it from reaching the stratosphere. However, some recent studies of the global CH<sub>3</sub>CCl<sub>3</sub> distribution and its budget (e.g. Singh, 1977b; Chang and Penner, 1977) indicate that a tropospheric residence time on the order of 10 years is more reasonable. This value is consistent with the 10 year residence time computed in this study employing the global OH distribution described earlier.

A summary of the calculations of methyl chloroform residence times is presented in Table 7. With the exception of the study by  $Cox \ \underline{et} \ \underline{al}.$  (1976b), the rate constant for  $CH_3CCl_3 + OH \rightarrow CH_2CCl_3 + H_2O$ is very close to 1 x  $10^{-14} \text{ cm}^3 \text{ mol}.^{-1} \text{ s}^{-1}$  (at 265°K) for the calculations.

# TABLE 7

# Methyl Chloroform Residence Time Calculations

Study	<sup>k</sup> сн <sub>3</sub> сс1 <sub>3</sub> + он	OH (Method of Calculation)	Residence Time
Yung et al. (1975)	8.9x10 <sup>-15</sup> (265 <sup>о</sup> к)	$12 \times 10^5$ (computed)	3 yrs
Cox <u>et al</u> . (1976b)	2.8x10 <sup>-14</sup>	$10 \times 10^5$ (assumed)	l yr
Chang and Wuebbles (1976)	1.4x10 <sup>-14</sup> (288 <sup>o</sup> K)	17x10 <sup>5</sup> (computed)	l yr
Neely and Plonka (1976)	1.0x10 <sup>-14</sup> (265 <sup>°</sup> K)	9x10 <sup>5</sup> (inferred)	3 yrs
Singh (1977b)	9.6x10 <sup>-15</sup> (265 <sup>°</sup> K)	4x10 <sup>5</sup> (inferred)	8 yrs
Chang and Penner (1977)	$[1.1 \times 10^{-14}]$	3x10 <sup>5</sup> (inferred)	11 yrs
This study	9.6x10 <sup>-15</sup> (265 <sup>°</sup> K)	3x10 <sup>5</sup> (computed)	10 yrs

Note: k is in units of  $cm^3 mol.^{-1} s^{-1}$ ;  $\overline{OH}$  is in  $cm^{-3}$ .

# TABLE 6

# Effect of Heterogeneous Removal on Model Calculations

	А	В	С
$k_{HET}^{o}$ (s <sup>-1</sup> )	2.0x10 <sup>-5</sup>	5.0x10 <sup>-6</sup>	$2.4 \times 10^{-6}$
OH (hemispheric annual average)	3.3x10 <sup>5</sup>	4.8x10 <sup>5</sup>	5.7x10 <sup>5</sup>
$H_2O_2$ (1 km)	0.34 ppb	1.40 ppb	2.46 ppb
$H_{2}O_{2}$ (10 km)	0.23 ppb	0.62 ppb	0.78 ppb

in Chastern St.

Justification for employing the relatively fast removal rate used to compute Profile A once again comes from the consideration of global tropospheric budgets. In addition to creating a greater imbalance in the tropospheric carbon monoxide budget by allowing more OH to be present, slower heterogeneous removal rates are inconsistent with the modelprescribed NO<sub>x</sub> distribution. Recalling that the average NO<sub>x</sub> column abundance in the Northern Hemisphere is 3.4 x  $10^{14}$  mol. cm<sup>-2</sup> (see Table 2, Column I) for these calculations, a tropospheric residence time for NO. can be found if the source strength were known. Anthropogenic input and lightning are believed to be the major sources of  $NO_v$  in the troposphere (Chameides et al., 1977; Söderlund and Svenssen, 1976), each contributing a flux in the Northern Hemisphere of about 1 x  $10^{10}$  mol. cm<sup>-2</sup> s<sup>-1</sup>. If we assume an average NO<sub>x</sub> flux in the Northern Hemisphere of 2 x  $10^{10}$ mol.  $cm^{-2} s^{-1}$ , the resultant NO<sub>2</sub> residence time is  $\sim 2 \times 10^4$  s ( $\sim 5$  hours) producing a removal rate for NO<sub>x</sub> of 5 x  $10^{-5}$  s<sup>-1</sup>. The current model's OH distribution can remove NO<sub>x</sub> at a rate no faster than 2 x  $10^9$  mol. cm<sup>-2</sup>  $s^{-1}$ , which means that another NO<sub>x</sub> removal process must be taking place at a rate four times faster than gas-phase chemistry. Even the sharp vertical gradient depicted for  $NO_x$  in Fig. 12 and the large globally averaged (and probably unrealistic) NO deposition velocity of 2 cm s  $^{-1}$ still demands that another heterogeneous process is removing NO, at a rate greater than  $10^{10}$  mol. cm<sup>-2</sup> s<sup>-1</sup>. Therefore, in light of the above discussion, it does not seem unreasonable that  $SO_2^{}$ ,  $NO_2^{}$  and all other tropospheric gases that are susceptible to scavenging processes in the atmosphere (such as  $HNO_3$ ,  $H_2O_2$ , and  $CH_3O_2H$ ) should be removed rapidly and that  $k_{HET}^{o} = 2 \times 10^{-5} \text{ s}^{-1}$  is a realistic value. Obviously, more global scale measurements of NO, NO $_2$ ,  ${}^{\rm H}_2{}^{\rm O}_2$ , and HNO $_3$  are needed to verify this contention.
For some reason, Chang and Wuebbles (1976) used the rate constant expression at  $288^{\circ}$ K to calculate tropospheric residence times. Evaluating their rate Arrhenius expression at  $265^{\circ}$ K yields a constant of  $8.3 \times 10^{-15}$  cm<sup>3</sup> mol.<sup>-1</sup> s<sup>-1</sup> and a residence time of 2 years. It is interesting to note that the OH calculated directly by previous photo-kinetic models (e.g., Yung <u>et al</u>., 1975 and Chang and Weubbles, 1976) compute OH values that are considerably higher than those computed in this study. The fact that we have used the faster rate constant for the CO + OH reaction given by Chan <u>et al</u>. (1977) may be responsible for lowering the amount of tropospheric OH by nearly a factor of two.

On the other hand, the OH abundance in the troposphere inferred from the various analyses of the methyl chloroform budget (with the exception of the study by Neely and Plonka, 1976) support the lower OH values computed in this study. Furthermore, if there existed an additional methyl chloroform sink in the troposphere other than OH-attack, then the inferred OH abundance would be even lower than the values presented in Table 7. Thus, we point out that the distribution of methyl chloroform and the analysis of its tropospheric budget are consistent with the OH distributions presented in this study and that the lower OH values computed directly are necessary to explain the global carbon monoxide and tropospheric ozone budgets in addition to being in better agreement with what is currently known about methyl chloroform. In summary, we conclude in this section that there is not enough OH in the troposphere to remove methyl chloroform efficiently and that it is likely that a substantial fraction of this chemical released to the atmosphere may find its way into the stratosphere.

VII, THE PRODUCTION RATE OF METASTABLE ATOMIC OXYGEN

Due to the fact that there are only a limited number of tropospheric measurements of OH, much of the discussion in Sections V and VI dealt with trying to reconcile the calculated OH distribution with the tropospheric budgets of ozone, carbon monoxide and methyl chloroform in order to determine the feasibility of such a distribution. Since global averages were used to compute OH in the model, it is felt that such an analysis is more practical than trying to compare local measurements with the calculations because of the likely variability of such observations.

Since the production of hydroxyl radicals in the troposphere primarily takes place through the reaction sequence:

> R22  $0_3 + hv \neq 0_2 + 0(^1D)$ , and R1  $H_2O + O(^1D) \neq 2$  OH,

comparison of the model-derived production rate with the measured production rate would provide a useful measure of the model's ability to simulate initial production of odd hydrogen radicals. Assuming that the amount of water vapor and ozone are known through observations, favorable comparison of the computed photolysis rate  $j_{22}$  with the measured value of  $j_{22}$  would indicate that the initial rate of formation of odd hydrogen radicals is being modelled correctly.

At the National Center for Atmospheric Research in Boulder, Colorado, Russ Dickerson, a graduate student under the direction of Donald Stedman of the Department of Chemistry at the University of Michigan, is currently measuring j directly. By placing a transparent bulb filled with nitrous oxide (N $_2$ O) and ozone in sunlight, the following reaction sequence takes place:

$$0_3 + hv \neq 0(^1D) + 0_2$$
, and  
 $0(^1D) + N_2O \neq 2 NO$   
or,  $\Rightarrow N_2 + 0_2$ .

By knowing the branching ratio of the latter two reactions, one can determine  $j_{22}$  by monitoring the amount of NO being produced. Note that there are other reactions taking place in the glass bulb, but that these are properly taken into account. A detailed description of the system is available from Dickerson.

The value of  $j_{22}$  computed in the model is calculated through the following expression:

$$\begin{array}{l} 320 \text{ nm} \\ j = \int Q(\lambda)\sigma(\lambda)\Phi(\lambda)d\lambda, \\ 290 \text{ nm} \end{array}$$

where  $Q(\lambda)$  is the solar flux (photons cm<sup>-2</sup> s<sup>-1</sup> nm<sup>-1</sup>),

 $\sigma(\lambda)$  is the ozone cross section (cm<sup>2</sup>),

and  $\Phi(\lambda)$  is the quantum yield of  $O(^{1}D)$  from  $O_{3}$  photolysis.

The effective solar flux at each level of the atmosphere in this wavelength interval is determined from the amount of radiation incident upon the top of the atmosphere, and subsequently from how much of that radiation is absorbed, scatterd, and reflected. The Rayleigh scattering and absorbing properties of the atmosphere are parameterized by an efficient scheme developed by Isaksen et al. (1976).

The input data used for these calculations are presented in Figures 16, 17 and 18. The incoming extraterrestrial radiation between 300 and 320 nm is shown in Figure 16 and is taken from Arvesen et al.





Construction of the accounters of the entropy of the basis swelt of the second states of the second seco





Fig. 18. (from Arnold et al., 1977)

(1969) with a spectral resolution of 0.4 nm. Figure 17 depicts the ozone absorption cross section between 300 and 320 nm (Moortgat and Warneck, 1975). Arnold et al. (1977) have developed an analytical expression for the temperature and wavelength dependence of the  $O(^{1}D)$  quantum yield between 300 and 320 nm. Figure 18 shows these values for three typical ground-level temperatures. The temperature dependence of the quantum yield can alter the calculated production rate of  $O(^{1}D)$  by as much as 20% over the range of tropospheric temperatures.

The model-derived photolysis rates are compared with Dickerson's measurements in Figure 19. The lightly dotted area is the modelderived photolysis rate plotted against the effective ozone column density. The scatter in the calculations exists because of the temperature dependence of the quantum yield and the parameterization of light scattering and absorption at various solar zenith angles. In general, the comparison of the calculated photodissociation rates with the measurements is quite good, and falls within the error bars of the observations for many cases. Overall, the calculated j<sub>22</sub> is higher than the measured value, especially when the sun is high in the sky (at relatively low values of effective ozone column density). In the particular model calculations shown in Figure 19, the earth's albedo is assumed to be zero and none of the incoming solar radiation is allowed to be attenuated by Mie scattering. In the actual calculations, however, this process was taken into account by assuming an average cloud height in the troposphere centered at 5 kms, and using the albedo data given as a function of latitude and month by Raschke et al. (1973). Note that by incorporating such processes into the calculations shown



Fig. 19

in Figure 19, the model-derived values would be lowered somewhat and might be in better agreement with Dickerson's measurements. In general, however, the model calculations do agree well enough with the measurements that we can conclude that the model is doing a reasonably good job of computing this important photolysis rate.

## VIII. SUMMARY AND CONCLUSIONS

Using the global tropospheric distributions of ozone, water vapor, carbon monoxide, temperature and the presently available chemical photokinetic information, we have developed a seasonally varying quasi-steady state numerical model to calculate the distribution of the hydroxyl radical in the troposphere. The resultant tropospheric OH concentration for the globe diurnally averaged over the entire year is approximately 3 x  $10^5$  cm<sup>-3</sup>, which is a factor of two to ten times lower than has been computed directly in previous photochemical models (see Neely and Plonka, 1976). However, this value is consistent with the analysis of methyl cholorform production history and its present global distribution (Singh, 1977a; Chang and Penner, 1977).

In addition to the better agreement with the tropospheric methyl chloroform budget, we showed in Section V that higher average OH values cause Seiler's (1974) carbon monoxide budget to be more out of balance than the lower OH concentrations. Regardless of which OH number densities are correct, an additional source of CO is required to balance Seiler's budget. Furthermore, we showed that the tropospheric budget of ozone may be closely related to the distribution of carbon monoxide and that CO oxidation may be a considerable source of ozone in the Norhtern Hemisphere troposphere. In fact, we conclude from our analyses that the lesser amount of tropospheric ozone seen in the Southern Hemisphere (compared to the Northern Hemisphere) may be a result of the smaller concentrations of CO in the Southern Hemisphere. Ongoing research is looking further into this phenomonen taking into account the possible differences in tropospheric-stratospheric exchange processes in the two hemispheres.

Since ozone enhances OH concentrations and carbon monoxide suppresses them (see Figure 1), the net effect of the CO and O<sub>3</sub> distribution serves to maintain a fairly equal value of OH in both hemispheres. This finding is contrary to the hypotheses of both Crutzen and Fishman (1977) and Singh (1977b) which suggested that OH levels in the Southern Hemisphere could be as much as three times larger than the values in the Northen Hemisphere.

Although we feel in these calculations that examination of the budgets of various trace gases is the best method of determining the validity of the OH distribution, comparison of the model-derived OH number densities with the scattered measurements does not universally refute or support the model results. Compared with our calculated OH values, the measurements of Davis et al. (1976) are several times higher, Campbell's (1977) preliminary observations are slightly lower, and the data of Perner <u>et al</u>. (1976) are too scattered and indeterminate to make a valid comparison. Furthermore, one must realize that a high variability is likely for any particular local OH measurement because of the high variation that exists for the parameters which affect OH directly (e.g., ozone, water vapor, etc.), and that the accuracy of the model cannot be assessed properly unless <u>all</u> important atmospheric parameters are well-known.

In Section VII, we compared our model-derived photolysis rate for the reaction

R22  $O_3 + hv \rightarrow O(^1D) + O_2$ ,

with the preliminary measurements of Dickerson. Since this photodissociation rate determines the rate of production of metastable atomic

oxygen, O(<sup>1</sup>D), which, through reaction with water vapor, serves as the primary source of OH in the troposphere, a valid comparison of our calculated photolysis rate with the measurements would be indicative of the model's ability to simulate this important rate in the atmosphere. The reasonably good comparison of Dickerson's measurements with our calculations is another sign which suggests that the model's results provide a reasonable approximation of the physical processes taking place in the atmosphere and that the model calculations may yield a fairly realistic picture of the tropospheric OH distribution.

Lastly, a large part of the discussion in this report focuses on the uncertainties involved in the calculations. In particular, the average distributions of the nitrogen oxides (NO and NO2) in the troposphere and the parameterization of heterogeneous removal processes in the atmosphere are two factors which greatly affect the calculated amounts of OH in the troposphere. This study suggests that very low background levels of NO and NO, must be present in the troposphere to yield a reasonable amount of production of tropospheric ozone. The presence of an NO, tropospheric column density greater than Noxon's (1977) upper limit of 5 x  $10^{14}$  molecules cm<sup>-2</sup> appears to enhance tropospheric photochemical production of ozone through carbon monoxide and methane oxidation to a very high level which exceeds the amount of ozone which can be destroyed in the troposphere by presently known destruction mechanisms. Likewise, through the analysis of various tropospheric budgets, this study shows that a relatively fast heterogeneous removal rate yielding tropospheric residence times of less than half a day for such species as SO2, NO2, HNO3, H2O2 and CH3O2H is not unrealistic in the troposphere. Until extensive measurements programs

of background levels of NO,  $NO_2$ ,  $HNO_3$  and  $H_2O_2$  are established and a good understanding of aerosol formation evolves, the incorporation of these parameters into numerical models will remain a source of uncertainty in photochemical calculations.

In conclusion, we feel that in addition to having developed a model which calculates a tropospheric OH distribution, we have provided a comprehensive analysis of the budgets of several trace gases in the troposphere which tends to support this distribution. We have examined some of the more important uncertainties in the model and note that it is not possible to minimize them until more measurements and a better understanding of microphysical processes are available. Lastly, we emphasize that this study should not be looked upon as the final compendium of tropospheric photochemistry, but rather as a first attempt to gain some insights into this complex and important field.

## References

Arnold, I., F.J. Comes, and G.K. Moortgat, 1977: Laser flash photolysis quantum yield of O(<sup>1</sup>D)-formation from ozone, Max Planck Institute for Chemistry, Mainz, preprint.
Arvesen, J.D., R.N. Griffin, Jr., and D.G. Pearson, Jr., 1969: Determination of extraterrestrial solar spectral irradiance from a research aircraft, <u>Appl. Optics</u>, <u>8</u>, 2215-2232.
Burrows, J.P., G.W. Harris and B.A. Thrush, 1977: The rates of reaction of HO<sub>2</sub> with HO and O studied by laser magnetic resonance, Dept. of Physical Chemistry, Cambridge, U.K.,

preprint.

- Calvert, J.G., J.A. Kerr, K.L. Demerjian and R.D. McQuigg, 1972: Photolysis of fromaldehyde as a hydrogen atom source in the lower atmosphere, Science, 175, 751-752.
- Campbell, M.H., 1977: Unpublished data, presented at the Fall, 1977, American Geophysical Union Meeting, December 5-9, 1977, San Francisco.
- Chameides, W.L., and D.H. Stedman, 1977: Tropospheric ozone: Coupling transport and photochemistry, <u>J. Geophys. Res.</u>, <u>82</u>, 1787-1794.
- Chameides, W.L., D.H. Stedman, R.R. Dickerson, D.W. Rusch and R.J. Cicerone, 1977: NO production in lightning, <u>J. Atmos.</u> <u>Sci.</u>, <u>34</u>, 143-149.
- Chan, W.H., W.M. Uselman, J.G. Calvert, and J.A. Shaw, 1977: The pressure dependence of the rate constant for the reaction  $HO + CO \rightarrow H + CO_2$ , <u>Chem. Phys. Ltrs.</u>, <u>45</u>, 240-243.

Chang, J.S., and J.E. Penner, 1977: Analysis of global budgets of halocarbons, submitted to Atmos. Env.

- Chang, J.S., and D.J. Wuebbles, 1976: A theoretical model of global tropospheric OH distributions, Proceedings of the Non-Urban Tropospheric Composition Symposium, Hollywood, Fla., jointly sponsored by the Americal Geophysical Union and the Americal Meteorological Society.
- Chatfield, R.B., and H. Harrison, 1977a: Troposperic ozone I: Evidence for higher background concentrations, <u>J. Geophys. Res.</u>, <u>82</u>, 5965-5968.
- Chatfield, R.B., and H. Harrison, 1977b: Tropospheric ozone II: Variations along a meridional band, <u>J. Geophys. Res.</u>, <u>82</u>, 5969-5978.
- Cox, R.A., R.G. Derwent, A.E.G. Eggleton and J.E. Lovelock, 1976a: Photochemical oxidation of halocarbons in the troposphere, <u>Atmos. Env.</u>, <u>10</u>, 305-308.
- Cox, R.A., R.G. Derwent, P.M. Holt and J.A. Kerr, 1976b: Photooxidation of methane in the presence of NO and NO<sub>2</sub>, <u>J. Chem. Soc.</u> <u>Faraday I</u>, <u>72</u>, 2044-2060.
- Crutzen, P.J., 1974: Photochemical reactions initiated by and influencing ozone in umpolluted tropospheric air, <u>Tellus</u>, <u>26</u>, 47-57.
- Crutzen, P.J., and J. Fishman, 1977: Average concentrations of OH in the Northern Hemisphere, and the budgets of CH<sub>4</sub>, CO, H<sub>2</sub> and CH<sub>3</sub>CCl<sub>3</sub>, <u>Geophys. Res. Ltrs.</u>, <u>4</u>, 321-324.
- Crutzen, P.J., I.S.A. Isaksen, and J.R. McAfee, 1977: The impact of the chlorocarbon industry on the ozone layer, <u>J. Geophys. Res.</u>, in press.

- Danielsen, E.F., 1968: Stratospheric-tropospheric exchange based on radioactivity, ozone, and potential vorticity, <u>J. Atmos. Sci.</u>, 25, 502-518.
- Danielsen, E.F., and V.A. Mohnen, 1977: Project Dustorm report: Ozone measurements and meteorological analysis of tropopause folding, J. Geophys. Res., in press.
- Davidson, B., J.P. Friend, and H. Seitz, 1966: Numerical models of diffusion and rainout of stratospheric radioactive materials, <u>Tellus</u>, <u>18</u>, 301-315.
- Davis, D.D., W. Heaps, and T. McGee, 1976: Direct measurements of natural tropospheric levels of OH via an aircraft-borne tunable dye laser, Geophys. Res. Ltrs., 3, 331-333.
- Drummond, J., 1977: Atmospheric measurements of nitric oxide using chemiluminescence, Ph.D. Dissertation, Dept. of Physics, Univ. of Wyoming, Laramie.
- Dütsch, H.U., 1970: Two years of regular ozone soundings over Boulder, Colo., National Center for Atmospheric Research Technical Note, NCAR-TN-10, Boulder, Colo., 441 pp.
- Ehhalt, D.H., 1974: The atmospheric cycle of methane, <u>Tellus</u>, <u>26</u>, 58-70.
- Eliassen, A., and J. Saltbones, 1975: Decay and transformation rates of SO<sub>2</sub>, as estimated from emission data, trajectories and measured air concentrations, <u>Atmos. Env.</u>, <u>9</u>, 425-430.
- Fabian, P., and P.G. Pruchniewicz, 1977: Meridional distribution of ozone in the troposphere and its seasonal variations, <u>J. Geophys.</u> Res., 82, 2063-2073.

Fishman, J., 1977: A numerical investigation of the meteorological and photochemical processes which influence tropospheric ozone and other trace constituents, Ph.D. Dissertation, Dept. of Earth and Atmos. Sci., Saint Louis University, St. Louis, Mo., 114 pp.

Fishman, J., and P.J. Crutzen, 1977: A numerical study of tropospheric photochemistry using a one-dimensional model, <u>J. Geophys.</u>, Res., 82, 5897-5906.

- Hampson, R.F., and D. Garvin, 1975: Chemical kinetic and photochemical data for modelling atmospheric chemistry, NBS Tech Note 866, U.S. Dept. of Commerce, 112 pp.
- Henmi, T., E.R. Reiter, and R. Edson, 1976: Residence time of atmospheric pollutants and long-range transport, Dept. of Atmos. Sci., Colorado State Univ., Ft. Collins, 78 pp.
- Howard, C.J., and K.M. Evenson, 1977: Kinetics of the reaction of HO<sub>2</sub> with NO, <u>Geophys. Res. Ltrs.</u>, <u>4</u>, 437-440.
- Isaksen, I.S.A., K.Midtbö, J. Sunde, and P.J. Crutzen, 1976: A simplified method to include molecular scattering and reflection in calculations of photon fluxes and photodissociation rates, Rept. No. 20, Institute of Geophysics, University of Oslo, Norway, 31 pp.
- Johnston, H.S., and R.A. Graham, 1973: Gas-phase ultraviolet absorption spectrum of nitric acid vapor, <u>J. Phys. Chem.</u>, <u>77</u>, 62-63.

Junge, C.E., 1960: Sulfur in the atmosphere, <u>J. Geophys. Res.</u>,

66, 227-237.

Levy, H., 1972: Photochemistry in the lower troposphere. <u>Planet.</u> Space Sci., 20, 919-935.

- Levy, H., 1973: Photochemistry of minor constituents in the troposphere, Planet. Space Sci., 21, 575-591.
- Meetham, A.R., 1950: Natural removal of pollution from the atmosphere, Quart. J. Roy. Meteor. Soc., 80, 96-99.
- Molina, L.T., S. Schinke, and M.J. Molina, 1977: Ultraviolet absorption spectrum of hydrogen peroxide, <u>Geophys. Res. Ltrs.</u>, <u>4</u>, 580-582.
- Molina, M.J., and F.S. Rowland, 1974: Stratospheric sink for chlorofluoromethanes: Chlorine atom-catalysed destruction of ozone, Nature, 249, 810-811.
- Moortgat, G.K., and P. Warneck, 1975: Relative O(<sup>1</sup>D) quantum yields in the near UV photolysis of ozone at 298 K, <u>Z. Naturforsch</u>, 30a, 835-844.
- Moortgat, G.K., E. Kudszus, and P. Warneck, 1977: Temperature dependence of O(<sup>1</sup>D)-formation in near U.V. photolysis, Max Planck Institute of Chemistry, Mainz, preprint.
- NASA, 1977: Chlorofluoromethane Assessment Workshop Report, Goddard Space Flight Center, Maryland.
- Neely, W.B., and J.H. Plonka, 1976: An estimation of the time averaged hydroxyl radical concentration in the troposphere, Dow Chemical Co., Midland, Mich., preprint.

Noxon, J.F., 1977: Tropospheric NO<sub>2</sub>, <u>J. Geophys. Res.</u>, in press.
Perner, D., D.H. Ehhalt, H.W. Pätz, E.P. Roth, and A. Volz, 1976:
OH-radicals in the lower troposphere, <u>Geophys. Res. Ltrs.</u>, <u>3</u>, 466-468.

- Pittock, A.B., 1974: Ozone climatology, trends and the monitoring problem, Conf. on Structure, Composition and Seasonal Circulation of the Upper and Lower Atmosphere and Possible Anthropogenic Perturbations, Melbourne, Australia, 455-466.
- Raschke, E., T.H. Vonder Haar, W.B. Bandeen, and M. Pasternak, 1973: The annual radiation balance of the earth-atmosphere system during 1969-1970 from Nimbus-3 measurements, <u>J. Atmos. Sci.</u>, <u>30</u>, 341-364.

Seiler, W., 1974: The cycle of Atmospheric CO, <u>Tellus</u>, <u>26</u>, 116-135.
Sie, B.K.T., R. Simonitis, and J. Heicklen, 1976: The reaction of
OH with CO, Int. J. Chem. Kin., 8, 85-88.

- Singh, H.B., 1977a: Atmospheric halocarbons: evidence in favor of reduced average hydroxyl radical concentration in the troposphere, <u>Geophys. Res. Ltrs.</u>, <u>4</u>, 101-104.
- Singh, H.B., 1977b: Preliminary estimation of average HO concentrations in the Northern and Southern hemispheres, <u>Geophys. Res.</u> <u>Ltrs.</u>, <u>4</u>, 453-456.
- Söderlund, R., and B.H. Svensson, 1976: The global nitrogen cycle, <u>Nitrogen, Phosphorus and Sulfur--Global Cycles</u>, edited by B.H. Svensson and R. Söderlund, SCOPE Rept. No. 7 and Ecological Bulletin No. 22, Swedish National Science Research Council, Stockholm.
- Stolarski, R.S., and R.J. Cicerone, 1974: Stratospheric chlorine: a possible sink for ozone, Can. J. Chem., 52, 1610-1615.
- U.S. Standard Atmosphere Supplements, 1966: U.S. Govt. Printing Office, Washington, D.C.

Yung, Y.K., M.B. McElroy, and S.C. Wofsy, 1975: Atmospheric halocarbons: a discussion with emphasis on chloroform, <u>Geophys. Res.</u> Ltrs., 2, 397-399.

D BLIOGRAPHIC DATA 1. Report No.	10			
TSHEET CSU-ATSP-284	2.	3. Recipient	's Accession No.	
4. Title and Subtitle	l	5. Report D.	ate	
The Distribution of the Hydroxyl Rad	lical in the	January	, 1978	
rioposphere		0.		
7. Author(s) Jack Fishman and Paul J. Crutzen			8. Performing Organization Rept. No.	
9. Performing Organization Name and Address		10. Project/	Task/Work Unit No.	
Colorado State University		11 Contract	Grant No.	
Fort Collins, Colorado 80523		EPA-R80	4921-01	
12. Sponsoring Organization Name and Address		13. Type of	Report & Period	
Environmental Protection Agency	matom:	Covered	Donort	
Research Triangle Park, North Carol	na 27711	Project	Keport	
		14.	6	
15. Supplementary Notes				
a two-dimensional distribution of the hydroxyl	uumerical model (OH) radical i	is develope in the tropos	d to calculate phere. The s 3 x $10^5$ cm <sup>-3</sup>	
<pre>diurnally, seasonally averaged global value of which is several times lower than the number co is in good agreement with the value inferred fr distribution of methyl chloroform. Likewise, on the tropospheric budgets of ozone and carbon lower computed value. One important result of this research is to of tropospheric ozone in the Southern Hemispher considerable difference in the tropospheric ozone that through the analysis of the likely photoch significant source of tropospheric ozone may ex- carbon monoxide oxidation. 17. Key Woras and Document Analysis. 170. Descriptors Tropospheric photochemistry Hydroxyl radical Ozone Budget Carbon Monoxide Budget</pre>	OH derived by omputed previou com the analysi the effects of a monoxide are the detailed an re. Our work s one patterns of temistry occurr tist in the Nor	alysis of the tro the computed not inconsis alysis of th hows that th the two hem ing in the t thern Hemisp	models, but pospheric OH distributio tent with this e distribution ere is a ispheres and roposphere, a here due to	
<pre>diurnally, seasonally averaged global value of which is several times lower than the number cd is in good agreement with the value inferred fr distribution of methyl chloroform. Likewise, for on the tropospheric budgets of ozone and carbon lower computed value.</pre>	OH derived by omputed previou com the analysi the effects of a monoxide are the detailed an re. Our work s one patterns of memistry occurr tist in the Nor	alysis of the tro the computed not inconsis alysis of th hows that th the two hem ing in the t thern Hemisp	models, but pospheric OH distributio tent with this e distribution ere is a ispheres and roposphere, a here due to	
<pre>diurnally, seasonally averaged global value of which is several times lower than the number co is in good agreement with the value inferred fr distribution of methyl chloroform. Likewise, to on the tropospheric budgets of ozone and carbon lower computed value.</pre>	OH derived by omputed previou om the analysi the effects of a monoxide are the detailed an re. Our work s one patterns of memistry occurr tist in the Nor	alysis of the tro the computed not inconsis alysis of th hows that th the two hem ing in the t thern Hemisp	models, but pospheric OH distributio tent with this e distribution ere is a ispheres and roposphere, a here due to	
<pre>diurnally, seasonally averaged global value of which is several times lower than the number co is in good agreement with the value inferred fr distribution of methyl chloroform. Likewise, for on the tropospheric budgets of ozone and carbon lower computed value.</pre>	OH derived by omputed previou com the analysi the effects of a monoxide are the detailed an re. Our work s one patterns of memistry occurr tist in the Nor	alysis of the tro the computed not inconsis alysis of th hows that th the two hem ing in the t thern Hemisp	models, but pospheric OH distributio tent with this e distribution ere is a ispheres and roposphere, a here due to	
<pre>diurnally, seasonally averaged global value of which is several times lower than the number cd is in good agreement with the value inferred fr distribution of methyl chloroform. Likewise, it on the tropospheric budgets of ozone and carbon lower computed value.</pre>	OH derived by omputed previou com the analysi the effects of a monoxide are the detailed an ce. Our work s one patterns of temistry occurr tist in the Nor	asly by other s of the tro the computed not inconsis alysis of th hows that th the two hem ing in the t thern Hemisp	models, but pospheric OH distributio tent with this e distribution ere is a ispheres and roposphere, a here due to	
<pre>diurnally, seasonally averaged global value of which is several times lower than the number cd is in good agreement with the value inferred fr distribution of methyl chloroform. Likewise, it on the tropospheric budgets of ozone and carbon lower computed value.</pre>	OH derived by omputed previou com the analysi the effects of a monoxide are the detailed an re. Our work s one patterns of memistry occurr tist in the Nor	alysis of the tro the computed not inconsis alysis of th hows that th the two hem ing in the t thern Hemisp	models, but pospheric OH distributio tent with this e distribution ere is a ispheres and roposphere, a here due to	
<pre>diurnally, seasonally averaged global value of which is several times lower than the number co is in good agreement with the value inferred for distribution of methyl chloroform. Likewise, for on the tropospheric budgets of ozone and carbon lower computed value.</pre>	OH derived by omputed previou com the analysi the effects of a monoxide are the detailed an re. Our work s one patterns of memistry occurr tist in the Nor	alysis of the tro the computed not inconsis alysis of th thows that th the two hem ing in the t thern Hemisp	models, but pospheric OH distributio tent with this e distribution ere is a ispheres and roposphere, a here due to	
<pre>diurnally, seasonally averaged global value of which is several times lower than the number co is in good agreement with the value inferred fn distribution of methyl chloroform. Likewise, on the tropospheric budgets of ozone and carbon lower computed value.</pre>	OH derived by omputed previou om the analysi the effects of a monoxide are the detailed an re. Our work so one patterns of memistry occurr tist in the Nor	alysis of the tro the computed not inconsis alysis of th hows that th the two hem ing in the t thern Hemisp	models, but pospheric OH distributio tent with this e distribution ere is a ispheres and roposphere, a here due to	
<pre>diurnally, seasonally averaged global value of which is several times lower than the number co is in good agreement with the value inferred fn distribution of methyl chloroform. Likewise, is on the tropospheric budgets of ozone and carbon lower computed value.</pre>	OH derived by omputed previou com the analysi the effects of a monoxide are the detailed an re. Our work so one patterns of memistry occurr tist in the Nor	this model i asly by other s of the tro the computed not inconsis alysis of th hows that th the two hem ing in the t thern Hemisp	models, but pospheric OH distributio tent with this e distribution ere is a ispheres and roposphere, a here due to	
<pre>diurnally, seasonally averaged global value of which is several times lower than the number co is in good agreement with the value inferred fn distribution of methyl chloroform. Likewise, for on the tropospheric budgets of ozone and carbon lower computed value.</pre>	OH derived by omputed previou com the analysi the effects of a monoxide are the detailed an re. Our work s one patterns of memistry occurr tist in the Nor	s of the tro the computed not inconsis alysis of th thows that th the two hem ing in the t thern Hemisp	models, but pospheric OH distributio tent with this e distribution ere is a ispheres and roposphere, a here due to 21. No. of Pages	
<pre>diurnally, seasonally averaged global value of which is several times lower than the number co is in good agreement with the value inferred fn distribution of methyl chloroform. Likewise, to on the tropospheric budgets of ozone and carbon lower computed value. One important result of this research is to of tropospheric ozone in the Southern Hemispher considerable difference in the tropospheric ozon that through the analysis of the likely photocol significant source of tropospheric ozone may ex- carbon monoxide oxidation. 17. Key woras and Document Analysis. 17a. Descriptors Tropospheric photochemistry Hydroxyl radical Ozone Budget Carbon Monoxide Budget 17b. Identifiers/Open-Ended Terms 17c. COSATI Field/Group 18. Availability Statement</pre>	OH derived by omputed previou com the analysi the effects of a monoxide are the detailed an re. Our work so one patterns of memistry occurr tist in the Nor	Class (This Class (This Class (This	models, but pospheric OH distributio tent with this e distribution ere is a ispheres and roposphere, a here due to 21. No. of Pages 82 22. Price	

•

	1
	1