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# STRATOSPHERIC REACTIONS OF PEROXYNITRIC ACID

John R. Barker, Paula L. Trevor, Richard A. Kenley, Jenq-Sian Chang, John E. Davenport, Bosco Y. Lan, and Graham Black

> SRI International 333 Ravenswood Avenue Menio Park, 94025





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## ABSTRACT

PNA (HOONO<sub>2</sub>) reactions have been studied because of their potential importance in the atmosphere. Experiments have been performed to measure the reaction rate constants for the following reactions:

$$0 + HOONO_2 \rightarrow Products$$
 (A)

 $H + HOONO_2 \rightarrow Products$  (B)

$$OH + HOONO_{\sim} \rightarrow Products$$
 (C)

$$O_{a} + HOONO_{a} \rightarrow Products$$
 (D)

The results were:

 $(k_{A} \pm 1\sigma) = (7.0 \pm 12.2) \times 10^{-11} \exp(-3369 \pm 489/T) \text{ cm}^{3} \text{ s}^{-1}, 228 \text{ K} \le 297 \text{ K} \\ \sigma_{AB} = 5.97 \times 10^{-8}$   $(k_{B} \pm 2\sigma) = (2.46 \pm 0.35) \times 10^{-14} \text{ cm}^{3} \text{ s}^{-1}, 248 \text{ K} \le T \le 315 \text{ K}$   $(k_{C} \pm 2\sigma) = (4.0 \pm 1.0) \times 10^{-12} \text{ cm}^{3} \text{ s}^{-1}, 246 \text{ K} \le T \le 324 \text{ K}$ 

 $k_{\rm D} \le 5 \times 10^{-20} \text{ cm}^3 \text{ s}^{-1}$ , 278 K

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## ACKNOW LEDGMENTS

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#### I INTRODUCTION

Both  $HO_2$  and  $NO_2$  play crucial roles in the chemistry of the upper atmosphere. For several years, these two species were thought to react via a radical disproportionation reaction, although it was suggested<sup>1</sup> that a longer-lived complex could be formed according to reaction (1):

$$HO_2 + NO_2 + M \neq HOONO_2 + M$$
 (1)

The importance of reaction (1) has recently been verified by direct observation of  $HO_2NO_2$  by Fourier transform infrared spectroscopy.<sup>2</sup> Moreover, the rate of reaction (1) has been measured,<sup>3</sup> and its reverse reaction (-1) has been studied.<sup>4</sup> Also, RRKM theory has been applied to these data for reaction (-1) so that its rate can be reliably estimated as a function of both temperature and pressure.<sup>5</sup> In another publication, the effect of  $HO_2NO_2$  on tropospheric photochemical smog chemistry<sup>6</sup> was discussed.

A concerted effort is under way to investigate PNA and determine the rates of various loss mechanisms that affect its residence time in the stratosphere. In addition to the experimental studies reported on its thermal decomposition rate, results of investigations on optical absorption cross sections have been reported<sup>7-9</sup> that permit calculation of the photolysis rates appropriate for the atmosphere.

In our laboratory, reactions of PNA with several atmospheric species have been studied to determine bimolecular rate constants for the destruction of PNA. The rate constants for PNA reactions with O-atoms, H-atoms,  $O_3$ , and OH are reported here.

Although the primary reason for studying PNA reactions is their potential importance in the atmosphere, PNA also is an interesting chemical that is part of the fabric of H-N-O species. In this sense, it is a member of the homologous series of nitrogen acids (e.g., HONO,  $HONO_2$ ,  $HOONO_2$ ), as well as being the first member of a series of pernitrates (e.g.,  $HOONO_2$ ,  $CH_3OONO_2$ ,  $CH_3COONO_2$ ); in addition, it is a hydroperoxide (e.g., HOOH, HOOR,  $HOONO_2$ ). All these species are important for various reasons, and data are needed on their reactions. Such data can then be used as a basis for predicting reaction rate constants for species that have not, as yet, been studied experimentally.

Each of the Appendices to this report has been submitted for publication in a technical journal and is intended to stand alone. Appendix A describes a convenient preparation of relatively pure PNA, and Appendices B-E give detailed descriptions of experiments designed to measure the rate constant for PNA reactions with O-atoms, OH,  $O_3$ , and H-atoms.

## II SUMMARY AND CONCLUSIONS

This section summarizes the results of experimental work on the preparation of PNA and some of its reactions and discusses the implications of atmospheric chemistry.

#### A. Results

#### 1. Preparation of PNA

Nitration of  $H_2O_2$  with  $NO_2BF_4$  provides a convenient method for preparing pure PNA in relatively large quantities. Aqueous PNA decomposes both by molecular elimination of  $O_2$  and nitrite from the pernitrate anion and by acid-catalyzed decomposition of PNA molecules.

## 2. $O(^{3}P) + HOONO_{2} \rightarrow Products$

The rate constant was measured under O-atom-rich conditions as a function of temperature; the rate constant is given by  $(k \pm 1\sigma) = (7.0 \pm 12.2) \times 10^{-11} \exp(-3369 \pm 489/T) \text{ cm}^3 \text{ s}^{-1}$ , covariance  $\sigma_{AB} = 5.97 \times 10^{-8}$ . Nine temperatures were studied ranging from 228 K to 297 K, and the uncertainties given are one standard deviation, based on a weighted least-squares analysis of the data. The products of the reaction could not be observed, but they are thought to be OH +  $O_2$  +  $NO_2$ ; HNO<sub>3</sub> can be ruled out since it was not observed.

## 3. OH + HOONO<sub>2</sub> $\rightarrow$ Products

Rate constants were measured under PNA-rich conditions by the flash-photolysis-resonance-fluorescence method for nine temperatures ranging from 246 K to 324 K. The rate constant was determined to be

 $(k \pm 2\sigma) = (4.0 \pm 1.0) \times 10^{-12} \text{ cm}^3 \text{ s}^{-1}$ , independent of temperature. Products could not be observed, and several pathways are possible.

## 4. $O_3 + HOONO_2 \rightarrow Products$

This reaction was investigated at 278 K under static conditions with  $O_3$  in excess. The reaction rate was very slow, giving an upper limit to the rate constant of  $k \leq 5 \times 10^{-20} \text{ cm}^3 \text{ s}^{-1}$ . Products were not observed.

#### 5. $H + HOONO_{2} \rightarrow Products$

This reaction was studied under H-atom-rich conditions at seven temperatures ranging from 226 to 315 K, but at 226 K the rate was too fast to allow measurement of a rate constant and there were indications that wall reactions were important at temperatures lower than 248 K. For temperatures above 248 K, the rate constant was  $(k \pm 2\sigma) = (2.46 \pm 0.35)$  $\times 10^{-14}$  cm<sup>3</sup> s<sup>-1</sup>, independent of temperature. The experimental technique appears to be free of problems and a tentative explanation is proposed for the very low A-factor. Products of the reaction were not observed, although they are expected to be analogous to those resulting from the reactions of H-atoms with  $H_2O_2$  and  $CH_3OOH$ . Nitric acid was not observed and can be ruled out as a reaction product.

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#### B. Implications for Atmospheric Chemistry

The potential importance of  $HO_2NO_2$  is due to its ability to act as a reservoir for both  $HO_2$  and  $NO_2$ . If  $HO_2NO_2$  has a long lifetime, it will act as a radical trap and slow down the overall free radical chain mechanism; thus, ozone will probably not be destroyed as quickly. Acting as a reservoir,  $HO_2NO_2$  can decompose slowly to release a steady stream of  $HO_2$ radicals and  $NO_2$ , even in darkness. Thus, it can serve as a free radical

initiator as well as a trap. This unique role underlines its potential importance and makes it difficult to estimate its effects without detailed modeling calculations.

Although  $HO_2NO_2$  can be shown not to influence tropospheric photochemical smog,<sup>6</sup> it potentially has a much more profound effect in the stratosphere because the lower temperatures and pressures combine to give it a much longer lifetime with respect to unimolecular decomposition. For example, at 220 K and 50 torr, its lifetime is estimated to be nearly 8 months, but at 280 K and 10 torr, the lifetime is only about 45 minutes.<sup>4</sup>,<sup>5</sup>

Reactions with atmospheric species must also be considered. Depending on the nature of such reactions, the net result might be to retard destructive stratospheric chemistry even further. For example, the potentially important reaction with OH can result in the net loss of an  $HO_2$  radical as well as an OH radical:

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For the reaction of O-atoms, however, the net result may have less effect on the ozone layer:

These two reactions can be quite important because of the net chemical changes involved and because the shortened lifetime of  $HO_2NO_2$ can influence the extent of diurnal effects; a long lifetime tends to damp them out.

Ozone and H-atoms might also react with  $HO_2NO_2$  to cause chemical effects as well as to shorten the  $HO_2NO_2$  lifetime:

$$O_3 + HOONO_2 \rightarrow HO + 2O_2 + NO_2$$
 (22)  
H + HOONO\_2  $\rightarrow$  Products (23)

As written, the ozone reaction products would have the following effect:

$$HO_{2} + NO_{2} \xrightarrow{M} HOONO_{2}$$

$$O_{3} + HO_{2}NO_{2} \xrightarrow{} OH + 2O_{2} + NO_{2}$$

$$OH + O_{3} \xrightarrow{} O_{2} + HO_{2}$$

$$met: 2O_{3} \xrightarrow{} 3O_{2} \qquad (29)$$

Again, this would have less effect on atmospheric chemistry than destruction of OH or  $HO_2$  radicals.

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Appendix A

PREPARATION AND THERMAL DECOMPOSITION OF PERNITRIC ACID IN AQUEOUS MEDIA

(Richard A. Kenley, Paula L. Trevor, Bosco Y. Lan)

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## A. Introduction

Pernitric acid (PNA),  $HOONO_2$ , can be viewed either as an inorganic peracid, ROOH (where  $R = NO_2$ ), a mixed anhydride of two acids (HOOH and HNO<sub>3</sub>) or as the parent member of the family of peroxynitrates,  $ROONO_2$ (where R = H). As such, PNA can be expected to participate either in free radical or ionic reactions and it therefore affords the opportunity for some interesting kinetic and mechanistic investigations.

In 1911 D'An's and Friederich<sup>1</sup> reacted  $H_2O_2$  with  $N_2O_5$  and obtained a solution that liberated  $Br_2$  from  $Br^-$ . They suggested that the solution contained HOONO<sub>2</sub>, but did not offer definitive proof for the existence of PNA. Schwarz<sup>2</sup> reported similar observations in 1948.

In 1977 Niki et al.<sup>3</sup> studied the gas phase reaction of photochemically generated chlorine atoms with a mixture of  $H_2$ ,  $O_2$  and  $NO_2$  and obtained the infrared spectrum of a species to which they assigned the formula  $HO_2NO_2$ . Other authors<sup>4,5</sup> have reported analogous photochemical methods for generating HOONO<sub>2</sub> in the gas phase. Graham et al.<sup>6</sup> condensed  $H_2O_2$  in a 253°K trap and passed a mixture of 1%  $NO_2$  in  $N_2$  through the trap. They then degassed the trap under  $N_2$  flow to yield PNA vapor.

The foregoing methods for generating HOONO<sub>2</sub> give a product that is contaminated with impurities and side products. As batch-type methods the existing procedures are not suitable for continuous generation of PNA, such as might be required for gas phase kinetic studies in flow systems. Moreover, the methods necessarily yield gaseous PNA, thereby precluding the possibility of investigating some interesting solution phase chemistry.

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It is well-known<sup>7-10</sup> that direct nitration of organic hydroperoxides yields peroxynitrates. The reaction can be generalized as in equation (1),

$$ROOH + NO_2^{T}X^{T} \rightarrow ROONO_2 + HX$$
(1)

where R = alkyl, acyl, aroyl, and X = OH,  $NO_3$  or  $BF_4$ , etc. The original method of D'Ans and Friederich can be considered as an example of this basic process if the reagents are written in an appropriate fashion, i.e. as in equation (2)

 $HOOH + NO_{2}^{+}NO_{3}^{-}(=N_{2}O_{5}) + HOONO_{2} + HNO_{3}$ (2) Nitric acid can also be written as  $NO_{2}^{+}OH^{-}$  and nitryl fluoroborate as  $NO_{2}^{+}BF_{4}^{-}$  using this formalism.

In view of the foregoing we examined the potential utility of reaction (1) as a synthetic method for PNA. In this paper we describe the preparation of PNA in solution, and the suitability of flow methods for generating relatively pure PNA vapor from the reaction mixtures. We also report on some decomposition reactions of  $HOONO_2$ .

#### B. Experimental

#### Materials

 $NO_2BF_4$  (Alfa Ventron), 90%  $H_2O_2$  (FMC) and 70%  $HNO_3(Baker)$  were used as received.

## Safety Hazards

The procedures used for preparation and analysis of PNA are potentially dangerous and certain safety precautions should routinely be observed.

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We encountered one explosion in the course of our work. While PNA vapor was passed through the heated glass inlet system of the mass

spectrometer we raised the temperature of the inlet to 700°K. There was a sudden pressure increase in the mass spectrometer chamber followed by a violent explosion that shattered the inlet system. Fortunately the inlet was behind a safety shield and there were no injuries. In all subsequent work the inlet temperature was never raised above 600°K and there were no further incidents of this  $t_{12}$  (Note: Schwarz<sup>2</sup> also reported a violent explosion apparencial incidents of PNA vapor).

For PNA reaction mixtures made to evolve (1), see below, and allowed to stand at room temperature, gas evolves for and the temperature of the reaction mixture increased over the period of a few hours. Eventually, gas evolution became so vigorous that the liquid bumped out of the reaction flask (again see reference 2). This behavior was not observed for PNA prepared by methods (2) and (3). The possibility of detonations due to large exotherms recommends that the synthetic reactions not be run on a scale exceeding ca. 5g. PNA reaction mixtures prepared by methods (1) and (2) are highly acidic and are powerful oxidizing agents. Skin contact results in painful burns. The reaction mixtures exhibit a sharp odor characteristic of many peroxides. One of us (RK) has complained of respiratory problems apparently associated with repeated exposure to PNA and other peroxynitrate vapors. These materials should be handled in a hood or other well-ventilated area.

## Preparation of HO<sub>2</sub>NO<sub>2</sub> from 90% H<sub>2</sub>O<sub>2</sub> and 70% HNO<sub>3</sub> (Method 1)

A 2 g (2.2 x  $10^{-2}$  mol) sample of 70% HNO<sub>3</sub> was added to a 1 dram specimen vial (A. H. Thomas) equipped with a 10-mm magnetic stirring bar. The acid was chilled in a small ice bath and 0.9 g (2.4 x  $10^{-2}$  mol) of 90% H<sub>2</sub>O<sub>2</sub> was added dropwise via a capillary pipette with vigorous agitation over a 10 minute period.

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## Preparation of $HO_2NO_2$ from 90% $H_2O_2$ and $NO_2BF_4$ (Method 2)

A 2 g (5.3 x  $10^{-2}$  mol) sample of 90% H<sub>2</sub>O<sub>2</sub> was chilled and stirred as described above. Solid NO<sub>2</sub>BF<sub>4</sub>, 0.4 g (3 x  $10^{-3}$  mol) was added in approximately 5 to 10 mg portions using a spatula. Effervescence and evolution of brown fumes accompany addition of NO<sub>2</sub>BF<sub>4</sub>.

#### Preparation of HO<sub>2</sub>NO<sub>2</sub> in CH<sub>3</sub>CN (Method 3)

90%  $H_2O_2$ , 0.1 g (3 x 10<sup>-3</sup> mol) was dissolved in 1 g of  $CH_3CN$  and cooled to 0°C with stirring.  $NO_2BF_4$ , 0.2 g (1.5 x 10<sup>-3</sup> mol) was added as described in Method 2.

## Characterization of HO<sub>2</sub>NO<sub>2</sub>

Infrared spectra of PNA vapor in Ar carrier were obtained using a Perkin Elmer Model 457 spectrophotometer and either a Wilks 20-m variable pathlength cell or a 2.5-cm ID x 10-cm pathlength Pyrex flow-through cell. Observed absorption maxima at 1745, 1715, 1390, 1300, 940, and 800 cm<sup>-1</sup> compared favorably with literature values for  $HOONO_2$ .<sup>3</sup>,<sup>6</sup> Impurities in the sample included  $NO_2$  (1620 cm<sup>-1</sup>) and  $HNO_3$  (840 cm<sup>-1</sup>).

#### Procedures

Vapor phase samples of PNA for ir analysis were generated from liquid reaction mixtures as follows. A capillary pipette was used to transfer about 3 ml of reaction mixture from the specimen vial used as a reaction flash (and maintained at  $273^{\circ}$ K) to a 10-ml bulb equipped with an 0-ring joint and a vacuum stopcock leading to an ebullation tube. The bulb was connected via the 0-ring joint to the ir cell and allowed to warm to ambient temperature. Argon was admitted to the bulb via the stopcock at a rate of approximately 30 cm<sup>3</sup> s<sup>-1</sup>, bubbled through the reaction mixture, and used to carry PNA vapor through the cell.

Vapor samples for mass spectral analysis were handled similarly, except that the Argon flow was eliminated. Evolution of oxygen from decomposition of PNA in the reaction mixture was sufficient to sweep PNA vapors into the system. The mass spectrometer and inlet system have been described elsewhere.<sup>24</sup>

The evolution of oxygen from aqueous PNA solutions was monitored with a calibrated pressure transducer (Validyne model DP-7) connected via a three-way stopcock to a jacketed, 250-ml bulb equipped with magnetic stirring. To perform the experiments, 100.0 ml of buffer was added to the bulb and allowed to come to temperature while  $0_2$  was vigorously bubbled through the solution. Next, the  $0_2$  flow was stopped, 25 to 100  $\mu 1$ of a PNA reaction mixture were added via syringe, the stopcock closed and transducer output monitored with a strip chart recorder. For a 50  $\mu$ l aliquot of preparative method (2), assuming 100% yield of PNA, the initial PNA concentration in 100 ml of buffer is calculated to be 7.5 x 10<sup>-4</sup> M. For a 150 cm<sup>3</sup> gas volume, the final pressure of the system is calculated to be 9 torr, assuming complete decomposition to  $0_2$ . Consistent with these calculations, observed final pressures were typically 5 to 10 torr. Control experiments indicated significant oxygen evolution from excess  $H_2O_2$  for PNA decompositions using preparative method (2) and pH  $\leq$  4. To circumvent this problem, we adopted preparative method (3), which does not require a large excess of  $H_2O_2$ .

Oxygen was identified as the reaction product by allowing a sample of PNA to decompose in a sealed, degassed, tube and then analyzing the gas over the frozen solution by mass spectroscopy.

Nitrite evolution was followed by monitoring the absorption at 365 nm with a Perkin-Elmer model 554 spectrophotometer. The buffer solution was first equilibrated to the desired temperature and 50-100  $\mu$ l of PNA solution was then injected into the buffer and mixed well. The absorbance at 365 nm was monitored until completion of the reaction.

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#### C. Results and Discussion

## Preparation and Identification of PNA

We used three general procedures (see experimental section) for preparation of PNA at 273°K:

- (1) reaction of neat 90%  $H_2O_2$  with 70%  $HNO_3$ ,
- (2) reaction of neat 90%  $H_2O_2$  with  $NO_2BF_4$ , and
- (3) reaction of  $NO_2BF_4$  and 90%  $H_2O_2$  in  $CH_3CN$ .

Successful preparation of PNA was confirmed for methods (1) and (2) by warming the reaction mixtures to ambient temperature and passing a stream of Ar into the solutions and through a 10-cm pathlength ir cell or into a long-path cell. In both cases, the characteristic ir spectrum of PNA<sup>3</sup>,<sup>6</sup> was readily observed.

On the basis of the reported value of the  $HOONO_2$  absorption coefficient at 803 cm<sup>-1</sup>,<sup>11</sup> we calculate that concentrations of  $HOONO_2$  as high as 10<sup>-4</sup> M were generated in the Ar stream using either method (1) or method (2). For method (1), we also observed variable, but large amounts (often exceeding the amount of PNA) of  $HNO_3$  and  $NO_2$  in the Ar stream. Using method (2), however, we found that the  $HNO_3$  and  $NO_2$  levels were normally less than 10% of the PNA level for cases where the concentration of PNA was on the order of  $16^{-6}$  M.

For all three methods, PNA vapors could be continuously generated from a single reaction mixture for periods of up to 60 minutes. This suggested the possibility of developing a convenient flow method for

generation and analysis of gaseous PNA. Accordingly, we constructed a glass inlet system for admitting PNA/Ar mixtures to a mass spectrometer. The inlet was divided into two sections, one at ambient temperature and one heated to  $600^{\circ}$ K (<u>Caution</u>! See experimental details). We prepared PNA via method (2) and admitted PNA vapors from the reaction mixture into the inlet system. We similarly obtained reference spectra for possible impurities (NO<sub>2</sub> and HNO<sub>3</sub>), and the data are given in Table 1.

The 300°K PNA mass spectrum does not exhibit a molecular ion at m/e = 79. We attribute the peaks at m/e = 46, 30, 18, and 17, respectively, to the fragmentation products  $NO_2^{\bullet+}$ ,  $NO^{\bullet+}H_2O^{\bullet+}$ , and  $OH^{\bullet+}$ . The m/e = 32 peak may derive either from fragmentation of the PNA molecular ion or from  $O_2$  generated by decomposition of PNA in the reaction mixture (vide infra). At 600°K, the PNA mass spectrum shows the same mass ions as at 300°K; the relative intensities of the peaks differ, however. The thermal decomposition of PNA at 600°K should proceed as in reactions (3) and  $(4)^{7,11-17}$ 

$$HOO-NO_2 \stackrel{*}{\leftarrow} HOO \bullet + NO_2 \tag{3}$$

$$2HOO \bullet \rightarrow H_2O_2 + O_2 \tag{4}$$

 $H_2O_2$  would also decompose at this temperature via reactions (5) and (6)

$$H0\bullet + H00\bullet \rightarrow H_20 + O_2 \tag{6}$$

The expected products of the PNA decomposition are thus  $NO_2$ ,  $O_2$  and  $H_2O$ . The 600°K PNA spectrum shows enhanced peaks at m/e = 32, 18, 17, and 16, consistent with  $O_2$  and  $H_2O$  as decomposition products. Also for PNA the 30/46 mass ion ratio increases from 0.95 at 300°K to 3.4 at 600°K. By comparison, the mass spectrum of  $NO_2$  at 600°K shows a 30/46 mass ion ratio

Table l

MASS SPECTRAL DATA FOR PERNITRIC ACID (PNA), NO2 AND HNO3

Compound <sup>a</sup>	Inlet Temperature °K		Relativ	e Inter	isity at	m/e <sup>b</sup> .c			1
		16	17	18	28	30	32	34	46
PNA	300	0.56	0.31	0.53	đ	0.95	2.0	۲∽	1
PNA	600	1.73	2.0	5.3	р	3.4	3.0	р	I
HNO3	300	ð	Ð	e	0.24	1.16	0.17	в	T
4NO3	600	ں ا	e	a	0.20	1.20	0.10	a	~-1
NO2	600	Ð	a	a	0.10	4.1	e	e	1

<sup>a</sup>PNA prepared by method (2), see experimental details. HNO<sub>3</sub> prepared by

reaction of H<sub>2</sub>SO4 with NaNO3.

<sup>b</sup>Each mass spectrum is separately normalized to m/e = 46.

cRelative intensity  $\leq$  0.05 for m/e not listed.

d<sub>Relative</sub> intensity ≤0.20.

eRelative intensity ≤ 0.05.

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of 4.1 while HNO<sub>3</sub> exhibits a mass ion ratio of 1.18  $\pm$  0.02 at either 300 or 600°K. The value of 3.4 for PNA at 600°K is sufficiently close to the value of 4.1 for NO<sub>2</sub> to indicate the latter as a product of the decomposition. The presence of HNO<sub>3</sub> as an additional product (perhaps via wall reactions)<sup>13</sup> would account for the 30/46 ratio being somewhat lower for PNA than for pure NO<sub>2</sub>. Thus, the change on heating of the 30/46 mass ion ratio serves to distinguish PNA from either NO<sub>2</sub> or HNO3.

## Decomposition of PNA in Solution

PNA reaction mixtures continually evolve oxygen (identified by mass spectroscopy) over a period of several hours. We examined the kinetics of oxygen evolution by measuring the increase in pressure of a closed system containing the reaction mixture from preparative method (2). Semilog plots of pressure versus time were linear, and the corresponding observed first-order rate constant for oxygen evolution,  $k_{obs}$ , was 2.6 x  $10^{-5}$  s<sup>-1</sup> at 273°K and 1.2 x  $10^{-4}$  s<sup>-1</sup> at 293°K.

Two possible mechanisms can account for the evolution of oxygen from PNA solutions. In either mechanism PNA is assumed to participate in the acid-base equilibrium, reaction (7)

$$HOONO_2 \xrightarrow{k_7}_{k_7} H^+ + O_2 NOO^-$$
(7)

PNA decomposition could then proceed via either free radical or ionic reactions. In the latter case, pernitrate ion would undergo unimolecular elimination of  $O_2$  and nitrite, reaction (8)

$$0_2 NOO^{-} \xrightarrow{k_0} 0_2 + NO_2^{-}$$
(8)

In the free radical mechanism, reversible homolytic O-N bond scission, reaction (9)

$$HOO-NO_2 \xrightarrow{k_9} HOO \cdot + NO_2$$
(9)

is followed by hydroperoxyl radical disproprotionation, reaction (4). Oxygen and  $H_2O_2$  are thus the expected free radical reaction products. Any NO<sub>2</sub> formed in reaction (9) would hydrolyze to a mixture of nitrite and nitrate.<sup>18</sup>

In an attempt to differentiate between the possible mechanisms we investigated the pH dependence of the kinetics of oxygen evolution from decomposition of  $10^{-3}$  M PNA in aqueous buffer. Table 2 lists values of  $k_{obs}$  for PNA solutions at 283.6°K and various pH. From the table, it is evident that  $k_{obs}$  increases with increasing pH over the range investigated. The good agreement for  $k_{obs}$  values obtained in acetate buffer using both preparative methods (2) and (3) is taken as evidence that PNA was formed by the latter method. For PNA solutions prepared by method (3) and diluted in acetate buffer (pH = 4.7), nitrite formation was detected spectrophotometrically ( $\lambda_{max}$  = 365 nm). For two runs at 293°K, the observed first-order rate constant for production of nitrite was (22.5 ± 1.8) x  $10^{-3}$  s<sup>-1</sup>, which is in reasonable agreement with the rate constant for oxygen evolution at the same temperature, but higher pH.

To the extent that only reactions (7) and (8) are operative, applying the steady-state approximation to  $0_2 NOO^{-1}$  gives equations (10) and (11)

$$+d[0_{2}]/dt = [PNA]k_{2}k_{8}/([H^{\dagger}]k_{-2} + k_{8}) = [PNA]k_{obs}$$
(10)

$$(k_{obs})^{-1} = (k_7)^{-1} + [H^T](k_{-7}/k_7k_8)$$
(11)

## Table 2

OBSERVED FIRST-ORDER RATE CONSTANTS FOR OXYGEN EVOLUTION (kobs) FROM

10<sup>-3</sup> M PERNITRIC ACID IN AQUEOUS BUFFER AT 10.5°C <sup>a</sup>

Run	Preparative <sup>D</sup> Method	Buffer <sup>C</sup>	рН	$k_{obs}$ , $d_{s^{-1}} \times 10^3$
1 <sup>e</sup>	2	acetate	4.90	$39.2 \pm 2.9^{f}$
2	3	acetate	4.87	7.40
3	2	acetate	4.70	$7.06 \pm 0.75^{g}$
4	3	acetate	4.32	4.46
5	3	KH phthalate	3.80	3.21
6	3	KH phthalate	3.55	3.05
7	3	KH phthalate	3.28	2.47
8	3	KH phthalate	3.04	2.17
9	3	HC1	1.36	1.57

<sup>a</sup>Temperature control to  $\pm$  0.5° K.

<sup>b</sup>See experimental details.

<sup>C</sup>Buffer =  $5 \times 10^{-2}$  M, prepared according to standard formulations

(reference  $\underline{19}$ ) and made to ionic strength = 0.5 with NaClO<sub>4</sub>.

 ${}^{d}k_{obs} = 1n2/t_{l_2}$ , from semi-log plot of pressure vs. time.

<sup>e</sup>Temperature = 293°K.

f Average of three runs.

<sup>g</sup>Average of four runs.

Assuming the mechanism given by reactions (7) and (8) a plot of  $(k_{obs})^{-1}$  versus  $[H^+]$  will be linear. In actuality such a plot (Figure 1) shows pronounced downward curvature below pH = 5. This indicates  $O_2$  evolution by an additional mechanism, and raises the possibility that PNA decomposition occurs at least in part by reactions (9) and (4).

It is interesting to compare the aqueous chemistry of PNA with that of other peroxynitrates. We have shown that formation of PNA via reaction (1) is entirely analogous to formation of alkyl, acyl, and aroyl peroxynitrates. There are also some interesting parallels in terms of decomposition mechanism. Alkaline hydrolysis of peroxyacetyl nitrate (PAN),  $CH_3C(0)00NO_2$ , yields stoichiometric amounts of acetate, nitrite, and (singlet) molecular oxygen.<sup>20</sup>,<sup>21</sup> Mudd<sup>22</sup> observed first-order kinetics for production of nitrite from hydrolysis of PAN ( $k_{obs} = 2 \times 10^{-3} s^{-1}$  at pH = 7.6 and 300°K). On the basis of these results Stephens<sup>23</sup> postulated a mechanism for alkaline hydrolysis of PAN involving the intermediacy of pernitrate anion, i.e., reaction (10)

 $CH_{3}C(0)OONO_{2} + OH^{-} \rightarrow CH_{3}C(0)OH + O_{2}NOO^{-}$  (10)

followed by reaction (8). Our results are consistent with this mechanism although a comparison of our kinetics with those of Mudd suggests that reaction (3), if it is involved in PAN hydrolysis, is not rate-limiting.

#### D. Conclusions

#### CONCLUSIONS

Nitration of concentrated  $H_2O_2$  with  $NO_2BF_4$  at 273°K yields PNA. PNA solutions thus prepared provide a convenient source of relatively pure PNA vapor. The analysis of PNA by infrared spectrophotometry or by mass spectroscopy in flow systems permits the study of reactions of PNA with radicals or other reactive species. These investigations are in progress and will be reported separately.

Aqueous PNA decomposes to molecular oxygen and nitrite. Unimolecular decomposition of the pernitrate anion to  $O_2$  and  $NO_2^-$  seems a likely mechanism, though a free radical mechanism for decomposition of PNA may also be important.

Similarities in the hydrolytic chemistry of PNA and peroxyacetyl nitrate suggest that the pernitrate anion is an intermediate in both cases. PNA and peroyxacetyl nitrate both play important roles in the free radical chemistry of photochemical smog. One question that now **remains** is whether the ionic chemistry of these pernitrates in heterogeneous phases (aerosols, surfaces, etc.) is also of environmental concern.

#### ACKNOWLEDGEMENT

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# Appendix B

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 $O(^{3}P)$  + HOONO<sub>2</sub>  $\rightarrow$  PRODUCTS: TEMPERATURE-DEPENDENT RATE CONSTANT (Jenq-Sian Chang, Paula L. Trevor, and John R. Barker)

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#### A. Introduction

Both  $HO_2$  radicals and  $NO_2$  play crucial roles in the chemistry of the atmosphere. Thus, it is not surprising that pernitric acid (PNA), the product of reaction (1), has received a good deal of recent attention:<sup>1</sup>

$$HOO + NO_2 \stackrel{M}{\neq} HOONO_2$$
 (1)

Because PNA couples the  $HO_x$  and  $NO_x$  catalytic cycles, its fate is of great importance for developing accurate computer models of atmospheric chemistry.<sup>2</sup> Oxygen atoms are present in the upper stratosphere and their reaction with PNA could affect its atmospheric lifetime:

$$0 + HOONO_2 \rightarrow Products$$
 (2)

Although the primary impetus for studying PNA reactions is their potential importance in the atmosphere, PNA is also an interesting chemical species that is an important part of the fabric of H-N-O chemistry. In this sense, it is a member of the homologous series of nitrogen acids (i.e., HONO,  $HONO_2$ ,  $HOONO_2$ ), as well as being the first member of the series of pernitrates (i.e.,  $HOONO_2$ ,  $CH_3OONO_2$ ,  $CH_3COONO_2$ , etc.); in addition, it is a hydroperoxide (i.e., HOOH, HOOR,  $HOONO_2$ ). Each of these species may be important in atmospheric chemistry, as well as in other chemical systems, and data are needed on their reaction rate constants. Such data can then be used as a basis for predicting reaction rate constants for species that have not, as yet, been studied experimentally.

#### B. Experimental

Pernitric acid was prepared<sup>3</sup> by slowly and <u>carefully</u> dissolving 0.6 g of  $NO_2BF_4$  in 2.0 g of 90% hydrogen peroxide. The solution was stirred vigorously, while immersed in an ice bath, and then maintained at ~10°C; the PNA vapor that evolved was introduced into the reactor through a short

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capillary tube. This preparative technique reliably gives relatively large yields of PNA with variable levels of contaminants. The level of  $NO_2$  usually was found to be less than 10% of the PNA. Hydrogen peroxide, on the other hand, tended to be highly variable; in some batches it was not measurable, but in others it exceeded the concentration of PNA itself. Partly because of these impurities, the kinetics were studied under O-atom-rich conditions, so that effects on the rate constants due to secondary reactions were minimized.

Because of the slowness of the reaction between O and PNA, and because we wished to study the kinetics under O-atom-rich conditions, a 2-liter spherical stirred-flow reactor (SFR) was used, as shown in Figure 1. High purity helium (Liquid Carbonic) carried 1-5% molecular oxygen through a microwave discharge to produce  $O({}^{3}P)$  atoms. In the 1-meter flow tube<sup>4</sup> (prior to entry into the SFR),  $NO_2$  could be added to the gas stream and the O-atom concentration could be determined by titration; the endpoint was signaled by extinction of the greenish-yellow glow due to 0 + NO  $\rightarrow$  NO<sub>2</sub> + h $\nu$ .<sup>5</sup> A correction was applied to compensate for the formation of  $O_3$  in the SFR, since it reduced the atom concentration from that measured in the flow tube (see below). An alternative method for measuring the O-atom concentration was to measure the  $O_2$  flow rate, and then monitor the mass spectrometer signal at m/e = 32 to determine the fraction of  $O_2$  dissoc+ iated in the microwave discharge. Both methods gave the same results within experimental uncertainty. Additional experiments were performed in which NO<sub>2</sub> titration of O-atoms was carried out at three different points in the apparatus: (a) in the flow tube prior to entry into the SFR, (b) in the SFR, and (c) at the exit of the SFR. Measured O-atom concentrations were the same at all three points, indicating that the O-atom wall loss rate constant is very slow and can be neglected. At higher O-atom concentrations, homogeneous O, formation depletes the atom concentration, as discussed below.

The helium carrier gas-flow rate was measured with a calibrated rotameter and the other gas flows (except for PNA itself) were measured by timing the pressure rise in calibrated volumes. Pressure rises were measured with calibrated variable reluctance pressure transducers (Validyne) and the total pressure in the reactor was measured with a calibrated Wallace and Tiernan gauge.

The gas stream was "turbulently" introduced in the SFR (V = 2389 cm<sup>3</sup>) through an inlet port that consisted of a 2-cm stub of tubing with four orifices oriented tangentially to the surface of the SFR. Reactant gases (such as PNA) were introduced through capillary inlets (1 cm x 1 mm) at C and D, located at the top of the SFR. The effluent gases passed through an outlet tube that was similar to the inlet port.

Due to the design of the experiment, O-atoms were intimately mixed with the carrier gas prior to introduction into the SFR. Since wall loss of O-atoms was negligible, the O-atom concentration was uniform within the reactor. For pseudo first order conditions, the rate of disappearance of PNA is controlled by the O-atom concentration, which is initially uniform, but which could be locally depleted due to nonhomogeneous mixing of PNA and its subsequent reaction with O-atoms. Because mixing of PNA appears to be quite rapid compared to residence times in the SFR, it is unlikely that local depletion of O-atoms plays a role in the measurements. Moreover, the complete mixing tends to insure that PNA experiences the same residence time as the carrier gas.

The rate of mixing of PNA within the SFR was not measured directly, but even without turbulence, the diffusion mixing times were shorter than any of the nominal residence times employed in experiments. The diffusion coefficient can be estimated by assuming that helium and PNA o ohave hard-sphere radii of 2.6 A and 6 A, respectively, giving the binary

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diffusion coefficient  $D \approx 200/P_{\tau}$ , where  $P_{\tau}$  is the pressure in torr. For diffusion in three dimensions,  $\langle r^2 \rangle \approx 6 \text{ Dt}_d$  and thus  $t_d \approx (3V/4\pi)^{2/3}/(6D)$ , where V is the volume of the reactor. At room temperature and for a pressure of 3 torr,  $t_d \approx 0.2$  sec. All of the experiments reported were for residence times greater than 0.8 sec., sufficient time for diffusive mixing of PNA throughout the SFR. The turbulence in the SFR reduces the mixing time still further, very effectively giving well-stirred conditions even if the diffusive mixing times are somewhat longer than estimated. Experiments described in the next section were performed at 297K for residence times that varied from 0.853 sec. to 1.39 sec., and there was no measurable variation in the rate constant observed.

Concentrations of species in the reactor could be monitored by sampling through a small hole (~ 0.01 inch) drilled through a thin (~ 1 mm) Teflon plug into a modulated-molecular-beam-mass-spectrometer (MMB-MS), which has been described elsewhere.<sup>4,6</sup> Calibrations were performed daily for known concentrations of stable species that were introduced into the gas stream. The PNA concentration was determined by monitoring the intensity of m/e = 46 and comparing it to the intensity corresponding to a known concentration of  $NO_2$ . The relative sensitivities of PNA and  $NO_2$  were determined by diverting the flow of PNA through a heated glass inlet and measuring the  $NO_2$  produced in the thermal decomposition reaction, as described in great detail elsewhere.<sup>7</sup>

$$HO_2NO_2 \rightarrow HOO + NO_2$$

$$2 HOO \rightarrow H_2O_2 + H_2O$$
(-1)
(3)

For the present experiment, the PNA gave about six times as intense a signal at m/e = 46 as did the same concentration of NO<sub>2</sub>. The addition of O-atoms to the gas stream during these measurements removed NO<sub>2</sub> impurities, and also constituted an independent check on the purity of each batch of PNA that

was used. The mass spectrometer was calibrated for  $HNO_3$  with a sample of dry gas prepared by the reaction of  $NaNO_3$  with concentrated  $H_2SO_4$ , and it was found that the sensitivities for  $HNO_3$  and PNA are similar at m/e = 46. The presence of  $HNO_3$  in our batches of PNA would be evidenced by the presence of a residual mass spectral intensity at m/e = 46 when the PNA is diverted through the heated inlet and O-atoms are added to remove  $NO_2$ . There was never any indication of such a residual signal, allowing us to rule out the presence of  $HNO_3$ .

Typically, experiments were performed at 1-3 torr total pressure, and residence times in the reactor were  $\sim 0.8$  to 2 seconds. Wall reactions were effectively discouraged by use of halocarbon wax (Halocarbon Products Corporation).

For temperature control, the reactor was enclosed in a special jacket and cold nitrogen gas was passed through the jacket. The special jacket consisted of a layer of heavy aluminum foil, followed by a porous layer of copper wool that allowed circulation of the cold nitrogen gas. The entire assembly was then enclosed in a thick layer of rigid foam insulation. The temperature control was less than perfect at the lower temperatures, and adequate stability was maintained for only two to three kinetic runs at a time. The temperature was monitored by seven chromelconstantan thermocouples located at various points on the reactor surface; they showed a temperature uniformity of  $\pm 1$  K at 270 K and  $\pm 4$  K at 240 K.

#### C. Results and Discussion

The stirred flow technique is a venerable one<sup>8</sup> that is especially useful for measuring relatively slow reaction rates. For the SFR used in the present experiments, data analysis is carried out for the wellstirred steady-state conditions assumed to be present in the reactor.

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The average residence time, t, for gases in the reactor is related to the total flow rate  $F_T$ , the total concentration  $c_T$ , and the reactor volume V, by:  $c_T V$ 

$$t = \frac{c_T^v}{F_T}$$

The concentration of the i<sup>th</sup> component of the mixture is given by

$$c_{i} = \frac{F_{i}}{F_{T}} c_{T}$$

where  $F_i$  is the flow rate of the i<sup>th</sup> component and  $c_T$  is the total gas concentration as related to  $P_T$  by the ideal gas law.

For a general bimolecular reaction  $A + B \rightarrow$  Products, where B is maintained in great excess so that pseudo-first-order conditions hold, a simple steady-state analysis gives the following equation:

$$\frac{\left[A\right]_{0}}{\left[A\right]} = 1 + k_{II} \left[B\right]_{t}$$
(4)

where  $[A]_0$  and [A] are the concentrations of species A in the absence and in the presence of B, respectively; t is the residence time in the SFR, and  $k_{II}$  is the bimolecular rate constant for the reaction. Data for  $[A]_0 / [A]$  plotted versus the product t[B] should give a straight line of slope  $k_{II}$  and unit intercept.

As a pragmatic test of the SFR, a reaction with a known rate constant was investigated at room temperature. The reaction was:

$$0 + Cl_a \rightarrow Cl0 + Cl \tag{5}$$

and the observed room temperature rate constant is  $k_s = (3.5 \pm 1.0) \times 10^{-14} \text{ cm}^3 \text{ sec}^{-1}$ , compared to the literature value<sup>8</sup>  $k_s = (4.2 \pm 0.6) \times 10^{-14} \text{ cm}^3 \text{ sec}^{-1}$ . This test shows that the SFR gives absolute rate constants in good agreement with those in the literature and that systematic errors are probably no greater than  $\pm 30\%$ .

As mentioned above, the O-atom concentrations were often measured in the flow tube (prior to entry into the SFR) by  $NO_2$  titration, and thus a correction for ozone formation was applied to obtain the corresponding O-atom concentrations in the SFR. The correction was performed as part of the steady-state analysis for the entire reaction scheme:

 $F_{NO_2} \xrightarrow{V} 0$   $O + HOONO_2 \xrightarrow{V} OH + O_2 + NO_2$   $O + O_2 + M \xrightarrow{O} OH + O_2$  (2)

$$0 + 0_2 + M \longrightarrow 0_3 + M$$
 (6)

$$0 + 0_3 \xrightarrow{\qquad} 2 \ 0_2 \tag{7}$$

$$0 + 0H \longrightarrow 0_2 + H$$
 (6)

$$H + HOONO_2 \longrightarrow H_2 + O_2 + NO_2$$
(9)

$$0 + NO_2 \longrightarrow O_2 + NO$$
 (10)

The reaction rate constant  $k_9$  has been measured in the companion paper and has been found to be  $\approx 3 \times 10^{-14} \text{ cm}^3 \text{ s}^{-1}$  with only a slight temperature variation. If the fast secondary reactions (8) and (10) are grouped together with reaction (2) and a steady state analysis is performed, the steady-state oxygen atom concentration is given by

$$\left[O\right] = \frac{F_{NO_{2}}}{V} \left\{ \frac{1}{t} + k_{6} \left[O_{2}^{\gamma} \left[M^{\gamma} \left(1 + \frac{k_{7} \left[O\right]}{k_{7} \left[O\right] + t^{-1}}\right) + k_{2} \left[PNA^{\gamma} \left(3 + \frac{k_{9} \left[PNA^{\gamma} + t^{-1}\right]}{k_{9} \left[PNA^{\gamma} + t^{-1}\right]}\right) \right\}^{-1} \right]$$
(11)

where  $F_{NO_2}$  is the flow of  $NO_2$  necessary to titrate the O-atoms in the flow tube and is equal to the flow of atomic oxygen into the SFR; V is the SFR volume and t is the residence time. Rate constants  $k_6$  and  $k_7$  were taken from the Hampson and Garvin,<sup>\$0</sup> and Baulch et al.,<sup>11</sup> respectively.

The solution of equation (11) requires rate constant  $k_2$  and must be performed iteratively as described below. Since secondary reaction (9) affects [PNA<sup>7</sup>, equation (4) also must be modified. A steady state analysis gives

$$\frac{[PNA^{\gamma}o]}{[PNA^{\gamma}]} = 1 + k_{2}t[O^{\gamma}\left\{\frac{t^{-1} + 2 k_{9}[PNA^{\gamma}]}{t^{-1} + k_{9}[PNA^{\gamma}]}\right\}$$
(12)

This expression can be rearranged to give

$$S = k_{2}t[0] = \left(\frac{[PNA]_{0}}{[PNA]} - 1\right)\left\{\frac{t^{-1} + k_{9}[PNA]}{t^{-1} + 2k_{9}[PNA]}\right\}$$
(13)

Since  $k_9$ , [PNA], [PNA], and t are known or are measured, S is directly measured in each experimental run, giving an experimental value for  $k_2$ (run), if [O] is known, for each experimental run.

The effects of secondary reaction are explicitly taken into account in equations (11) and (13). The objective in the data analysis is to solve equations (11) and (13) in a self-consistent manner in order to devise [0] for each individual run and  $k_2(T)$  for each set of runs at a given temperature. This goal was expedited since S does not depend on  $k_2$  and [0] depends on  $k_2$  only weakly. The analysis procedure was begun by taking the data for each individual experimental run and solving equations (11) and (13) iteratively for [0] and for a value of  $k_2$ :  $k_2$ (run). Convergence was rapid and was complete to within 0.1% after the five iterations used for each calculation. The second step of the analysis was to consider each set of several runs corresponding to the same temperature and average the derived values for  $k_2(run)$  to give  $\langle k_2 \rangle$ . This approximate value for  $k_2$  was then used to recalculate [0] for each run in the series so that the values of S and the recalculated [0] values could be used to derive a final self-consistent value for  $k_2(T)$ . In principle, this procedure could be extended further by using  $k_2(T)$  in

place of  $\langle k_2 \rangle$  to produce a still more self-consistent value for  $k_2(T)$ , but this extension was not necessary, because of the weak dependence of [Olon  $k_2$ , according to equation (11). The final value of  $k_2(T)$  was derived from equation (13) according to either of two least-squares procedure, depending on the number of data points at a given temperature. If more than four data points were available, an unconstrained least-squares analysis was used, but the intercept was constrained to the origin if there were four or fewer data points. Values of  $k_2(T)$  for a given data set never differed from  $\langle k_2 \rangle$  for that data set by more than 25%. A 25% variation in  $\langle k_2 \rangle$  was found to give a variation in  $k_2(T)$  of  $\sim 2\%$ , far less than the other errors involved.

The data are presented in Table 1 along with the derived values for  $[O^{7}, S, and k_{2}(T)]$ . The values of  $k_{2}(T)$  are summarized in Table 2. To determine the Arrhenius parameters for rate constant  $k_{2}$ , a weighted least-squares analysis was performed using the Taylor series method described by Cvetanovic and coworkers.<sup>12</sup> The uncertainties associated with  $k_{2}$  at each temperature were adjusted<sup>12b</sup> by Student's t-factor according to the number of data points and the resulting "adjusted" standard deviations were used in the least-square analysis. The results were  $(k_{2} \pm \sigma) = (7.0 \pm 12.2) \times 10^{-11} \exp(-3369 \pm 489/T) \text{ cm}^{3} \text{ s}^{-1}$ ; covariance = 5.97 x  $10^{-8}$ . This analysis does not take potential systematic errors into consideration, however, and such errors might change these results by an estimated factor of two, to be conservative.

Table 1 DATA SUMMARY

P		~ <b>*</b>			[ DNA ]		[ 0 ]
'T torr)	(s)	NO <sub>2</sub>	°2,	PNA Jo	[PNA]	S	[0]
97 K							
7.70	0.851	9.72	5.40	1.49	1.29	0 210	<b>)</b> 0
2.70	0.957	9 04	5 40	1 70	1 77	0.207	217
2.20	0.053	0 04	5 57	1.30	1 . 37	0.247	3.0
2.70	0.053	0 0 0	1.13	1.63	1.33	0.24/	3.0
2.20	0.003	0.02	4.70	2+80	1.25	0.1/3	2.0
2.20	0.853	6.67	3.52	2.89	1.20	0.138	2.0
2.72	0.914	/.05	4.10	1.92	1.22	0.161	2.1
(./2	0.914	8.//	5+16	7.79	1+34	0.198	2.6
2.72	0.914	7.48	4.40	6.12	1.22	0.133	2.2
2.72	0.914	1.21	0.71	2.91	1.01	0.007	0.3
2.50	1.390	1.07	0.63	20.60	1.15	0.079	0.3
2.50	1.390	6.58	4.70	10.20	1.52	0.281	2.0
2.50	1.390	6.19	5.16	7.24	1.56	0.310	1.8
2.60	0.874	10.90	5.43	2.22	1.33	0.234	3.3
+60	0.874	9.40	4.70	2.22	1.34	0.241	2.8
			(k. ±	$1\sigma$ ) = (9.3	9 + 1.30	x 10 <sup>-16</sup> c	n <sup>3</sup> s <sup>-1</sup>
8 <u>2 K</u>			2 -				
1.42	1.330	10.70	4.70	14 10	• •	A 100	
1.42	1.770	20.20	7.77	14.10	1.30	0.192	4.4
2.05	1.044	14 40	3./3	12.20	1.45	0.241	8.3
2.05	1 044	14.70	2+19	13.90	1.56	0.292	6.0
2103	1.744	8.48	4.84	14.80	1.54	0,281	3.57
			(k <sub>2</sub> ±	$1\sigma$ ) = (2,9	8 ± 0.83)	x 10 <sup>-16</sup> cr	a <sup>3</sup> s <sup>-1</sup>
68 K							
2.05	1 540	0 04	<b>F</b> 20	E 00			
	4.007	0.04	5.20	5.80	1+45	0.253	3.5
1 10	0.9//	8.40	5.20	20.00	1.28	0.150	3.5
1.40	1.229	16.50	5.20	21.40	1.31	0.163	6.5
1.40	1.229	16.00	5.40	11.30	1.34	0.185	6.2
1.40	1,229	16.00	5,40	11.30	1.31	0.169	6.2
1.40	1.229	14.60	5.20	10.90	1.35	0.191	5.7
2.42	2.090	16.00	5.10	10.80	1.64	0.335	6.2
2.42	2.090	13.70	5.20	9.14	1.56	0.296	5.3
3.47	3.003	13.70	5.20	13.10	1.47	0.242	5 4/
3.47	3.003	18.80	5.20	3.20	2.23	0.664	7 7
3.47	3.003	16.00	5.20	2.64	2.00	0.552	6.2
063 V			$(k_2 \pm$	$1\sigma$ ) = (2.	53 ± 0.45)	x 10 <sup>-16</sup> c	:m <sup>3</sup> s <sup>-1</sup>
200 K							
2.05	1.610	4.60	5.00	2.56	1.20	0.127	1.70
2.05	1.610	4.60	5.00	2.77	1.25	0.154	1.70
1.60	1.739	16.60	6.07	2.58	1.16	0.101	5.74
• 60	1.739	16.60	6.07	2.58	1.25	0.156	5.74
							5.7.0
			(k <sub>2</sub> ± )	207) = (3.2	3 ± 2.30)	x 10 <sup>-16</sup> cm	<sup>3</sup> s <sup>-1</sup>
644 K							
4.60	1.860	16.60	6.07	3.40	1.14	0.004	
1.45	1.140	10.70	5.70	7 70	1+14	0.04	6.92
	<b>t</b>		2.20	3.32	1.0/	0.046	4.48
			(k2±)	$l\sigma$ ) = (0.78	8 ± 0.17)	x 10 <sup>-17</sup> cm	3 5-1
228 K							
<u> </u>							
1.57	2.260	11.80	2.87	5.03	1.04	0.034	4.04
	1.850	9.40	4.90	5.51	1.08	0.046	3.44
1.32							3,40
.32							

dunits: 10<sup>14</sup> molecules cm<sup>-3</sup>.

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т	Data Points	$(k_2 \pm 1\sigma_c)^a$
297	14	$9.39 \pm 1.35^{b}$
282	4	$2.98 \pm 1.00$
268	11	2.53 <u>+</u> 0.48 <sup>c</sup>
263	4	3.23 <u>+</u> 2.76
244	2	0.78 <u>+</u> 0.31
228	2	0.54 <u>+</u> 0.44

Table 2  $O + HOONO_2 \rightarrow Products$ 

<sup>a</sup>Units are  $10^{-16}$  cm<sup>3</sup> s<sup>-1</sup>; uncertainties are one standard deviation, corrected by student's t factor.

b Linear least-squares result; (intercept  $\pm 1\sigma$ ) =  $-0.01 \pm 0.03$ .

<sup>c</sup> Linear least-squares result; (intercept  $\pm 1\sigma$ ) = 0.01  $\pm 0.06$ . Although the slow rate constant makes reaction (2) unimportant in most of the atmosphere, it is interesting to compare the Arrhenius parameters obtained for reaction of 0 + PNA to those obtained for other oxygen-atom reactions:

	$A \times 10^{12}, \text{cm}^3 \text{ s}^{-1}$	E/R	Ref
$0 + H_2O_2$	$2.75 \pm 0.41$	$2125 \pm 261$	13
$0 + HONO_2$	$(k < 3 \times 10^{-17})$	at 300 K)	14
$0 + HOONO_2$	70 ± 122	<b>33</b> 69 ± 489	This work

Since A-factors for O-atom abstraction reaction are expected to be  $\sim 1-5 \times 10^{-11} \text{ cm}^3 \text{ s}^{-1}$ , the A-factor determined in the present work is not unreasonable, although highly uncertain. That for  $0 + \text{H}_2\text{O}_2$  is unexpectedly low, but the activation energy is also low, leading to absolute rate constants within about a factor of 2-5 of those determined in the present study. Thus, the chemistry of PNA appears to be more similar to that of  $\text{H}_2\text{O}_2$  than to that of nitric acid, which is quite unreactive.

### Acknowledgement

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This paper is dedicated to the late W. H. Duewer, who was active in investigating the role of PNA in atmospheric chemistry.

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## FIGURES

- 1 Low-Pressure Stirred-Flow Reactor Note that the reactor is coupled to a discharge flow tube (for preparation of reactive species) and to the modulated-beam mass spectrometer.
- 2 Stirred-Flow Reactor Results for 0 + PNA → Products Plotted as S versus t [0] for (a) 297 K and (b) 268 K.
- 3 Arrhenius Plot for the Rate Constant for 0 + PNA  $\rightarrow$  Products. Number associated with each data point shows the number of kinetic runs at each temperature. Confidence limits were derived from Propagation of Errors, including the covariance  $\sigma_{AB} = 5.97 \times 10^{-8}$ .



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# Appendix C

H + HOONO<sub>2</sub>  $\rightarrow$  PRODUCTS: TEMPERATURE-DEPENDENT RATE CONSTANT (Paula L. Trevor and John R. Barker)

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## A. Introduction

Pernitric acid (PNA) is the product of the combination reaction of  $HO_2$  radicals and  $NO_2$ :

$$HO_2 + NO_2 \rightarrow HOONO_2$$
 (1)

PNA has been the subject of considerable interest<sup>1-6</sup> since it represents a direct coupling pathway between the HO<sub>X</sub> and NO<sub>X</sub> catalytic cycles in the earth's atmosphere. In our laboratory, we have studied the reactions of OH, <sup>7</sup> O<sub>3</sub>, <sup>8</sup> O atom, <sup>9</sup> and H atom with PNA in order to determine the chemical reaction rate constants and assess the relative importance of the various processes that affect PNA in the atmosphere. Recent measurements of the PNA absorption cross sections in the ultraviolet <sup>10-12,21</sup> indicate that photolysis may be less important than chemical reaction. Since atomic hydrogen is present in the upper stratosphere, reaction (2) is an appropriate one for study.

$$H + HOONO_2 \longrightarrow H_2 + O_2NO_2 \qquad (2a)$$

$$\downarrow \\ O_2 + NO_2$$

$$\rightarrow$$
 H<sub>2</sub>0 + NO<sub>3</sub> (2b)

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$$\rightarrow$$
 H<sub>2</sub>O<sub>2</sub>+ NO<sub>2</sub> (2c)

$$\rightarrow$$
 HO + HONO<sub>2</sub> (2d)

Although its possible importance in the atmosphere was the primary reason for studying reactions (2), PNA is an interesting chemical species that represents several homologous series, including the nitrogen acids, the peroxides and the hydroperoxides. We are not yet able to identify the reaction products (although we can rule out  $HONO_2$ ) and so it was hoped that the reaction rate constant could provide clues to the dynamical pathways involved, giving us some insight into the chemistry of the various homologous series of which PNA is a member. As described below, the measured rate constants exhibit such unusual behaviour that interpretation is tenuous and uncertain, but it is also clear that the reaction is too slow to be important in the stratosphere.

## B. Experimental

The reaction of hydrogen atoms with PNA was studied under hydrogenatom-rich conditions due to the ubiquitous presence of NO<sub>2</sub> as an impurity in the PNA. The PNA was prepared as described in detail elsewhere <sup>13</sup> by slowly and carefully dissolving 0.6-g NO<sub>2</sub>BF<sub>4</sub> in 2.0-g 90% H<sub>2</sub>O<sub>2</sub> that was stirred vigorously at 0°C. The solution temperature was maintained at 0°C or slightly above in order to obtain the desired rate of evolution of PNA, and the vapor that evolved was introduced into the reactor via a short capillary tube. This preparative technique gives gas-phase PNA that contains NO<sub>2</sub> and H<sub>2</sub>O<sub>2</sub>, as well as O<sub>2</sub> and H<sub>2</sub>O, which were not measured.

Because of the  $NO_2$  impurity and because  $NO_2$  is one of the likely products of the reaction of H-atoms with PNA, the kinetics were studied under atom-rich conditions, where the atom concentration is not significantly affected reactions with the  $NO_2$  impurity. The kinetics were studied in a stirred-flow reactor (SFR), as depicted in Figure 2. A very similar reactor was used previously for the study of the oxygen atom reaction<sup>9</sup> with PNA, and all of the experimental details are quite similar in both studies.

Briefly, the SFR is coupled to a modulated molecular-beam mass spectrometer (MMBMS) that has been described elsewhere.<sup>14</sup>,<sup>15</sup> Hydrogen atoms were generated by a microwave discharge and swept by the helium carrier gas into the SFR of volume 2106 cm<sup>3</sup>. Wall reactions were inhibited by a halocarbon wax coating (Halocarbon Products Corp.). The helium flow rate was measured with a calibrated rotameter, and the flow rates of minor species were measured by the rate of pressure increases in calibrated volumes. The total pressure in the SFR was measured with a calibrated Wallace and Tiernan gauge.

Pernitric acid concentrations were monitored with the MMBMS at m/e = 46 and the absolute concentration was determined relative to the NO<sub>2</sub> sensitivity, as described in detail previously.<sup>7</sup> The hydrogen-atom flux into the SFR was measured by adding ~ 2-fold excess of C1NO and observing the resulting concentration of HCl product:

$$H + C_NO \longrightarrow HC1 + NO$$
(3)

A weakness of this method is that the H-atom reaction with ClNO is so fast<sup>16</sup> that potential wall reactions of H-atom cannot compete; thus, this method gives only a measure of the flux of H-atom into the reactor. not the true steady-state concentration encountered by the PNA. This may not be a severe weakness, however, since wall reactions of H-atom are usually relatively slow as measured in flow-tube reactors, and the SFR has a much lower surface/volume ratio than do flow-tube reactors. For typical flow-tube reactors (2.5 cm i.d.), the first-order wall-loss rate constants usually range from  $0-5 \text{ sec}^{-1}$ ; for the flow tube coupled to the SFR in the present experiments, no wall loss could be observed. implying  $k_{w} \lesssim 1 \text{ sec}^{-1}$ . (It should be pointed out, however, that the halocarbon wax coating on the flow tube has a melting point of 150°C, and the wax coating on the SFR has a melting point of 80°C. Thus, although the waxes are chemically similar, they have different average molecular weight and are from different batches.) For the flow tube,  $S/V = 1.6 \text{ cm}^{-1}$ , whereas that for the SFR is ~ 0.38 cm<sup>-1</sup>; thus, if the wall-reaction rate constant for the flow tube is conservatively estimated to be  $k_w = 2 \pm 2 \text{ sec}^{-1}$ , that for the SFR will be  $k'_w = 0.5 \pm 0.5 \text{ sec}^{-1}$ . Since the residence time in the SFR was sometimes as short as one second, the higher value of  $k_w^\prime$  could cause a factor of 2 error in the H-atom concentration, but the error is expected to be much less. Oxygen atoms

also do not combine rapidly on walls treated with halocarbon wax. In our study<sup>9</sup> of the O + PNA reaction,  $k_w$  for O-atoms was too slow to be detected in the SFR. If H-atoms are similar,  $k_w$  for H-atoms is negligible. After the data for this study was collected, another inlet was attached to the system immediately before the throttling stopcock and after the SFR. A photomultiplier tube (RCA 1P28) was nounted so as to view the region just beyond this additional inlet. Nitric oxide was introduced into the system (in excess of the H-atom concentration) at three different points, namely, the flow tube, the SFR, and the additional inlet while the intensity of the red HNO<sup>\*</sup> emission was recorded. No difference in the HNO<sup>\*</sup> emission was detected ( $\pm$  20%) regard as of the titration point. Also, no change was observed whether or not PNA was being added to the system. These data further evidence the fact that wall loss of H-atoms probably is not a problem in the kinetic study.

Because  $NO_2$  is present as an impurity and is also a possible product of the reaction between H-atom and PNA, we added carbon monoxide to the gas stream to act as an OH-radical scavenger that converts OH back to H:

 $H + NO_2 \longrightarrow OH + NO$  (4)

$$OH + CO \longrightarrow H + CO_2$$
 (5)

Typically,  $\sim 1-2 \times 10^{15}$  molecules/cm<sup>3</sup> of CO were added, leading to a rate for reaction (5) that is far greater than other potential losses of OH in this system.<sup>17</sup> Thus, the observed extent of reaction should not reflect contributions due to OH attack on PNA.

Typically, experiments were performed at 1-3 torr total pressure, and residence times in the reactor were 1-2 seconds. For temperature control, the reactor was covered with heavy aluminum foil and was then closely wrapped with a coil of Tygon tubing through which a cooling fluid was circulated from a constant temperature bath (Neslab, Model LT-50). A closed-cell flexible foam sheet was then used as insulation. The temperature was monitored by seven chromel-constantan thermocouples attached to the surface of the bulb; they showed excellent temperature uniformity:  $\pm 1$  K at 230 K.

# C. Results and Discussion

The data analysis for the SFR was performed as described earlier.<sup>9</sup> Briefly, the residence time t is related to the total gas flow rate  $F_T$ , the total concentration  $C_T$ , and the reactor volume V by:

$$t = \frac{\frac{C_T V}{T}}{\frac{F_T}{T}} \quad . \tag{6}$$

The concentration of the i<sup>th</sup> component of the mixture is given by:

$$C_{i} = \frac{F_{i}}{F_{T}} C_{T}$$
(7)

where  $F_i$  is the flow rate of the i<sup>th</sup> component, and  $C_T$  is the total gas concentration as related to  $P_T$  by the ideal gas law.

The input flux of H-atoms was measured by ClNO titration, as discussed earlier. Using this technique, the H-atom concentration is given by:

$$\left[H^{\intercal} = \left[HC1\right] \left(\frac{1}{1 + tk_{w}}\right)$$
(8)

where [HCl<sup>¬</sup> is the concentration of HCl as measured with the calibrated MMBMS at m/e = 36, and  $k'_w$  is the first-order wall reaction rate for loss of H-atoms in the SFR. As mentioned earlier, an estimated upper limit to  $k'_w$  is ~ 1 sec<sup>-1</sup>, but it is probably much lower. We will assume  $k'_w \approx 0$  and [H<sup>¬</sup> = [HCl<sup>¬</sup> as determined in the ClNO titration.

For bimolecular reaction (1) where [H7 is maintained in great excess so that pseudo-first-order conditions prevail, a simple steady-state analysis gives:

$$\frac{I_0}{I} = \frac{[PNA]_0}{[PNA]} = 1 + tk_2[H^{\gamma}]$$
(9)

where  $I_0$  and I are mass spectral intensities at m/e = 46 and [PNA]<sub>0</sub> and [PNA] are the concentrations of PNA in the absence and in the presence of [H], respectively; t is the residence time in the SFR, and  $k_2$  is the bimolecular reaction rate constant. Data for [PNA]<sub>0</sub>/[PNA] plotted versus t[H] should give a straight line of slope  $k_2$  and unity intercept. A sufficient excess of [H] was used so that even if the primary products reacted further with H-atoms, there was no significant variation in [H]. Moreover, if OH is a reaction product, it is converted back to H atoms by reaction with the added CO. The data obtained in the present study are presented in Table 1 and are plotted in Figure 1, where the lines shown were computed by least squares analysis. Slopes and intercepts obtained from such data for all of the temperatures studied are summarized in Table 2. The slopes may be identified with the overall rate constant for reaction (2). The calculated intercepts are unity (within one standard deviation) for all except the three lowest temperatures. At these temperatures, long periods of time were necessary for stabilization of the PNA intensity at m/e = 46 and there appeared to be a systematic variation toward higher extents of reaction (for the same [H])over a period of time; this pathological behavior may be symptomatic of heterogeneous reactions, and thus the data for the lowest three temperatures are somewhat suspect. At 226 K, all of the PNA reacted, and the rate constant could not be determined. On the other hand, the data for the higher temperatures were found to be reproducible.

Aside from the lower temperature data points, the apparent bimolecular rate constant shows a remarkable constancy with temperature. Considering the rather low average value for  $k_2$ , the constancy with temperature is difficult to explain. This is because the constancy with temperature implies that the measured rate constant corresponds to the A-factor, and the activation energy is zero. Typical A-factors for H-atom reactions are usually  $\geq 10^{-11}$  cm<sup>3</sup> sec<sup>-1</sup>, nearly three orders of magnitude larger than the observed rate constant. Thus, a puzzle exists.

		STIRRED-FD.	W REACTOR EXPERIMENT.	AL DATA	
т (к)	P <sub>T</sub> (torr)	t (sec)	$[HOONO_2]_0$ (molecule.cm <sup>-3</sup> )	[H] <sub>o</sub> (molecule·cm <sup>-3</sup> )	I o I
315	2.00	1.00	$3.2 \times 10^{12}$	$3.51 \times 10^{13}$	1.84
	2.00	1.00	$3.2 \times 10^{12}$	$4.62 \times 10^{13}$	2.15
	2.00	1.00	$3.2 \times 10^{12}$	$6.35 \times 10^{13}$	2.48
	2.00	1,00	$3.2 \times 10^{12}$	$7.83 \times 10^{13}$	2.74
	2.00	1.00	$3.2 \times 10^{12}$	$9.35 \times 10^{13}$	3.17
	2.00	1.00	$3.2 \times 10^{12}$	$1.54 \times 10^{14}$	4.23
	2.00	1.00	$3.2 \times 10^{12}$	$1.91 \times 10^{14}$	5.81
				$k_2 = (2.38 \pm 0.1)$	6) x $10^{-14}$
			10		
297	2.00	1.06	$2.0 \times 10^{12}$	$2.38 \times 10^{13}$	1.71
	2.00	1.06	$2.0 \times 10^{-2}$	$3.42 \times 10^{-3}$	1,80
	2.00	1.06	$2.0 \times 10^{12}$	$4.04 \times 10^{13}$	2.53
	2.00	1.06	$2.0 \times 10^{12}$	6.92 x 10 <sup>+3</sup>	2,72
	2.00	1.06	$2.0 \times 10^{12}$	$4.50 \times 10^{13}$	3.14
	2.00	1.06	$2.0 \times 10^{12}$	$4.50 \times 10^{13}$	2.53
	2.05	1.24	$1.0 \times 10^{12}$	$9.20 \times 10^{13}$	4.41
	2.05	1.24	$1.4 \times 10^{42}$	$4.40 \times 10^{-3}$	2.27
	2.05	1.24	$1.4 \times 10^{12}$	$2.30 \times 10^{13}$	1.68
				$k_2 = (2.87 \pm 0.5)$	i0) x $10^{-14}$
274	2.00	1.15	$2.5 \times 10^{12}$	$2.16 \times 10^{13}$	1.72
	2.00	1.15	$2.5 \times 10^{12}$	$2.79 \times 10^{13}$	1.82
	2,00	1.15	$2.5 \times 10^{12}$	7.81 x 10 <sup>13</sup>	2.50
	2.00	1.15	$2.5 \times 10^{12}$	$1.14 \times 10^{14}$	4.95
	2,00	1.15	$2.5 \times 10^{12}$	9.86 x 10 <sup>13</sup>	4.90
	2.00	1.15	$2.5 \times 10^{12}$	$4.63 \times 10^{13}$	2.27
	2,00	1.15	$2.5 \times 10^{12}$	5.41 x $10^{13}$	2.92
	2.00	1.15	$2.5 \times 10^{12}$	$6.61 \times 10^{13}$	3.60
	2.20	1.15	$2.5 \times 10^{12}$	8.71 x 10 <sup>13</sup>	4.44
				$k_2 = (3.28 \pm 0.5)$	6) x 10 <sup>-14</sup>
258	2.00	1.22	$2.6 \times 10^{12}$	$1.41 \times 10^{14}$	4.48
	2.00	1.22	$2.6 \times 10^{12}$	$1.03 \times 10^{14}$	4.30
	2.00	1.22	$2.6 \times 10^{12}$	$4.40 \times 10^{13}$	2.05
	2.00	1.22	$2.6 \times 10^{12}$	$5.98 \times 10^{13}$	3.09
	2.00	1.22	$2.0 \times 10^{12}$	$1.40 \times 10^{14}$	5.00
	2.00	1.22	$2.0 \times 10^{12}$	$7.62 \times 10^{13}$	1,88
	2,00	1.22	$2.0 \times 10^{12}$	5.61 x $10^{13}$	1.84
	2.00	1.22	$2.0 \times 10^{12}$	$3.77 \times 10^{13}$	1.49
	2.00	1.22	$2.0 \times 10^{12}$	$1.76 \times 10^{13}$	1.36
	2.00	1.22	$2.0 \times 10^{12}$	$4.01 \times 10^{13}$	1.85
	2.00	1.22	$2.0 \times 10^{12}$	5.61 x $10^{13}$	2.62
	2.00	1.22	$2.0 \times 10^{12}$	4.46 x 10 <sup>13</sup>	2.46
				$k_2 = (2.35 \pm 0.3)$	2) x $10^{-14}$

Table 1 HOONO<sub>2</sub> + H → PRODUCTS; IREED-FLOW REACTOR EXPERIMENTAL DATA

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т (к)	P <sub>T</sub> (torr)	t (sec)	$[HOONO_2]_0$ (molecule·cm <sup>-3</sup> )	[H] <sub>o</sub> (molecule·cm <sup>-3</sup> )	I º /I
248	2.00	1.27	$3.0 \times 10^{12}$	$4.04 \times 10^{13}$	1.92
	2.00	1.27	$3.0 \times 10^{12}$	$2.14 \times 10^{13}$	1.47
	2.00	1.27	$3.0 \times 10^{12}$	$4.34 \times 10^{13}$	2.36
	2.00	1.27	$3.0 \times 10^{12}$	$1.06 \times 10^{14}$	7.11
	2.00	1.27	$3.0 \times 10^{12}$	$3.56 \times 10^{14}$	4.94
				$k_2 = (3.84 \pm 2.94)$	i) x 10 <sup>-14</sup>
238	2.00	1.32	$2.5 \times 10^{12}$	$4.30 \times 10^{13}$	5,25
	2.00	1.32	$2.5 \times 10^{12}$	$2.46 \times 10^{13}$	2.57
	2.00	1.32	$2.5 \times 10^{12}$	$3.38 \times 10^{13}$	4.09
				$k_2 = (6.41 \pm 1.40)$	) x $10^{-14}$
226	2.00	1.39	$2.7 \times 10^{12}$	$1.69 \times 10^{14}$	LARGE
	2.00	1.39	$2.7 \times 10^{12}$	9.80 x 10 <sup>13</sup>	LARGE
	2.00	1.39	$2.7 \times 10^{12}$	$2.35 \times 10^{13}$	LARGE
				k <sub>2</sub> = FAST	

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Table 1 (concluded)

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т	k <sub>2</sub> ± 1σ	Intercept	Data
(K)	$(cm^3 \cdot molecule^{-1} \cdot sec^{-1})$	± 1σ	Points
315	$(2.38 \pm 0.16) \times 10^{-14}$	$0.95 \pm 0.18$	7
207	$(2.87 \pm 0.50) \times 10^{-14}$	1 02 6 90	٥
451	$(2.87 \pm 0.50) \times 10$	1.03 ± 0.29	5
274	$(3.28 \pm 0.56) \times 10^{-14}$	$0.77 \pm 0.46$	9
258	$(2.35 \pm 0.32) \times 10^{-14}$	$0.75 \pm 0.30$	12
200		0.10 2 0.00	
248	$(3.84 \pm 2.94) \times 10^{-14}$	0.55 ± 1.33	5
238	$(6.41 + 1.40) \times 10^{-14}$		3
	(0)		Ū
226	(too large for measurem	ent)	3

Table 2

HOONO<sub>2</sub> + H → PRODUCTS; SUMMARY OF APPARENT BIMOLECULAR RATE CONSTANTS

Weighted average from 248 K to 315 K:

 $(k_2 \pm 2\sigma) = (2.46 \pm 0.35) \times 10^{-14} \text{ cm}^3 \text{ s}^{-1}$ .

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Based upon analogy with  $H_2O_2 + H$  and  $CH_3OOH + H$ ,<sup>17</sup> several reaction paths are possible for reaction (2). The simple abstraction reaction (a) is expected to have an A-factor near  $10^{-11}$  cm<sup>3</sup> sec<sup>-1</sup> and an activation energy. The three "addition" or "displacement" paths will give the products shown. The Arrhenius parameters for such addition reactions are difficult to estimate, since few examples are known for this class of compounds. However, if res s for H +  $H_2O_2$  and H +  $CH_3OOH$  are taken at face value, the addition reactions can contribute from 10% to 70% of the total reaction.

For the case H + HOONO<sub>2</sub>, reaction channel (d) can be immediately ruled out because we observe no evidence for nitric acid formation. Because of the slow reaction between H and HNO<sub>3</sub>, nitric acid should survive under our conditions and, if present, it would be detected at m/e = 46. At this cracking peak, HNO<sub>3</sub> gives a signal ~ 90% as intense as that of HOONO<sub>2</sub>, for the same concentration. If nitric acid were formed the plots of I<sub>0</sub>/I versus [H<sup>¬</sup>t would show nonlinearity, tending to level off at high [H<sup>¬</sup>t. Inspection of the data shown in Figures 1(a) and 1(b) shows no tendency toward nonlinear behavior, and thus reaction channel (d) must contribute less than 10% of the total reaction. Under our conditions, it was not possible to detect the products of the other channels due to their fast reaction with the excess [H]. (Thus, there is also no interference at m/e = 46 due to product NO<sub>2</sub>, for example.)

In the absence of the identification of reaction products, the observed rate constant does not provide sufficient information for an unambiguous interpretation of the reaction dynamics. Nevertheless, several reasonable conclusions can be reached although they cannot be proved.

First of all, the reaction is probably <u>not</u> a simple H-atom abstraction reaction since such reactions usually have A-factors of about  $1-5 \times 10^{-11} \text{ cm}^3 \text{s}^{-1}$  and activition energies of ~ 5-10 kcal/mole. An abstraction reaction with these Arrhenius parameters might make itself evident at a temperature of ~ 400 K, but at the temperatures of the present study it would be too slow to be observed. These considerations suggest, however, that if additional studies were performed at higher temperatures, the observed reaction rates might reflect the presence of the abstraction channel (2a).

The "addition" or " isplacement" reactions can be written

$$H + PNA \xleftarrow{a}_{-a} [H_2 NO_4]^* \xrightarrow{b} Products (10)$$

where the  $H_2NO_4^*$  complex is a vibrationaly excited species formed by chemical activation. For this reaction scheme, the observed rate constant will be<sup>18</sup>

$$k_{obs} = \frac{k_{a}k_{b}}{k_{-a}+k_{b}} = \epsilon k_{a}, \qquad (11)$$

where  $\epsilon$  in an "efficiency factor".

Since  $k_a$  is expected to be ~ 1 x 10<sup>-11</sup> cm<sup>3</sup> s<sup>-1</sup>, similar to other H-atom addition reactions, the efficiency factor at room temperature is  $\epsilon(300) \approx 2 \times 10^{-3}$  and we may conclude that  $\epsilon \approx k_b/k_{-a}$ . Since reaction (-a) and (b) are unimolecular reactions almost certainly at the low pressure limit, we can use classical RRK theory and write

where E is the internal energy,  $E_b$  and  $E_{-a}$  are the respective critical energies for the two reaction channels, and s is the effective number of oscillators, which is usually about 2/3 of the actual number of oscillators in the molecule. The energy E is given by  $E \approx E_{-a} + skT$ ; therefore

$$\epsilon \approx \frac{A}{A_{-a}} \left[ 1 - \frac{\Delta E}{skT} \right]^{S-1}$$
 (13)

where  $\Delta E = E_{-a} - E_{b}$ , the difference in reaction critical energies.

To evaluate the A-factors,<sup>19</sup> we must estimate the entropy of activation. Forreaction (-a), the A-factor is expected to be similar to those for other H-atom bond fissions and we can estimate  $\log A_{-a} \approx 13.5$ . For reaction (-b), we can invoke a cyclic transition state:



If the three internal rotors in  $H_2NO_4$  are free rotors with no barriers, the major contribution to  $\Delta S^{\pm}$  will be the loss of entropy due to tying them up. This represents a limiting case (i.e. lowest estimate for  $k_b$ ) since significant barriers to internal rotation in  $H_2NO_4$  will reduce the amount of entropy lost in going to the cyclic transition state. For the three rotors,  $S_{int}^{max} \cong 10$  Gibbs/mole, giving log  $A_b \gtrsim 11$ . Thus  $A_b/A_{-a} \gtrsim 3 \times 10^{-3}$  and equation (13) becomes

$$\varepsilon \gtrsim 3 \times 10^{-3} \left[ \frac{\text{skT} - \Delta E}{\text{skT}} \right]^{S-1}$$
 (15)

Since the observed efficiency factors appear to be virtually independent of temperature for the more reliable higher temperature data, this analysis indicates that  $\Delta E \approx 0$ . A similar analysis using quantum RKK theory leads to the same general result.

The magnitude of  $\varepsilon$  predicted by equation (15) is within a factor of two of that deduced from the experimental data, indicating that this picture of the reaction dynamics is not inconsistent with experiment. On the other hand, the uncertainties in this analysis are difficult to estimate and are probably large. The idea that H-atoms can attach to the O-atom of the NO<sub>2</sub> group does not seem outlandish, but there is no clear precedent for this behavior.

Other dynamical reaction pathways are also conceivable, but in invoking them, it is more difficult to reproduce the value of  $\epsilon$  determined from experiment. For example, there exists some evidence that H-atoms attack the O-atom in H<sub>2</sub>O<sub>2</sub> and in HOOCH<sub>3</sub>.<sup>17</sup> If this were the case for the reaction

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of H + PNA, one could write a transition state involving a 3 or 4 member ring. Such structures do not tie up as many free-rotors and thus the  $\Delta S^{\pm}$  values would lead to larger values for  $A_b$ , requiring  $\Delta E > 0$  to reproduce the value of  $\varepsilon$  at room temperature. This leads to an inconsistency, however, since it would also predict that  $\varepsilon$  increases with temperature, contrary to experiment.

In conclusion, the interpretation of the experimental results is not straight forward, although addition of H to PNA followed by a cyclic elimination reactors appear to be consistent with the data. More extensive data over a wider temperature range may help to elucidate the reaction mechanism and, specifically, the simple abstraction reaction should become apparent at higher temperatures. A careful reaction product study might also clear up this issue.

Considering the concentration of H-atoms at various altitudes in the atmosphere, it is estimated that at 40 km,  $[H] \approx 10^5$  cm<sup>-3</sup> <sup>20</sup>; below 40 km, [H] is insignificant. Taking the measured rate constant from this study, the estimated lifetime of PNA with respect to reaction with H-atoms is  $_{T} \gtrsim 13$  years. Thus, this reaction is not expected to have any significant effect on the PNA concentration in the upper atmosphere. It is also worth noting that even if systematic errors have affected our rate constant measurements, the measured rate constants must be too slow by a factor of 10<sup>3</sup> or more for the reaction of H-atoms to affect the atmospheric PNA concentration.

# Acknowledgements

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# Appendix D

OH + HOONO<sub>2</sub>  $\rightarrow$  PRODUCTS: TEMPERATURE-DEPENDENT RATE CONSTANT

(Paula L. Trevor, Graham Black, and John R. Barker)

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#### A. Introduction

Both  $HO_2$  and  $NO_2$  play crucial roles in the chemistry of the upper atmosphere. For several years, these two species were thought to react via a radical disproportionation reaction, although it was suggested<sup>1</sup> that a longer-lived complex could be formed according to reaction (1):

$$HO_{2} + NO_{2} + M \approx HOONO_{2} + M$$
 (1)

The importance of reaction (1) has been verified by direct observation of  $HO_2NO_2$  using Fourier transform infrared spectroscopy.<sup>2</sup> Moreover, the rate of reaction (1) has been measured,<sup>3</sup> and its reverse reaction (-1) has been studied.<sup>4</sup> Also, RRKM theory has been applied to these data for reaction (-1) so that its rate can be reliably estimated as a function of both temperature and pressure.<sup>5</sup> In another publication, the effect of  $HO_2NO_2$  on tropospheric photochemical smog chemistry<sup>6</sup> was discussed.

A concerted effort is under way to investigate pernitric acid (PNA) and determine the rates of various loss mechanisms that affect its residence time in the stratosphere. In addition to the experimental studies reported on its thermal decomposition rate, results of investigations on optical absorption cross sections have been reported,<sup>7-9</sup> permitting calculation of the photolysis rates appropriate for the atmosphere.

In our laboratory, reactions of PNA with several atmospheric species have been studied to determine bimolecular rate constants for the destruction of PNA, including reactions with O-atoms,<sup>10</sup> H-atoms,<sup>11</sup> and  $O_3$ .<sup>12</sup> Each of these three reactions is too slow to be important for atmospheric chemistry.

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Of the atmospheric species, the OH radical is probably the most reactive, and its reaction with PNA can have a profound effect on the chemistry of the stratosphere. The reaction products are not known, but several set are thermochemically possible:

$$\rightarrow$$
 HO<sub>2</sub> + HNO<sub>2</sub> (2c)

If the reaction products are given by equation (2a), model calculations<sup>13</sup> show the net result of reactions (1) and (2) is the  $M_2$ -catalyzed disproportionation of OH and  $HO_2$  radicals. The effect of this reaction is to reduce the calculated depletion of the stratospheric ozone column by about a factor of two.<sup>13</sup> Because of the sensitivity of stratospheric chemistry calculations to the rate and products of reaction (2), we have made a great effort to obtain a reliable experimental measurement of the reaction rate constant.<sup>14</sup> Unfortunately, we are not yet able to identify the reaction products.

Although the primary reason for studying PNA reactions is their potential importance in the atmosphere, PNA also is an interesting chemical that is part of the fabric of H-N-O species. In this sense, it is a member of the homologous series of nitrogen acids (e.g., HONO,  $HONO_2$ ,  $HOONO_2$ ), as well as being the first member of a series of pernitrates (e.g.,  $HOONO_2$ ,  $CH_3OONO_2$ ,  $CH_3COONO_2$ ); in addition, it is a hydroperoxide (e.g., HOOH, HOOR,  $HOONO_2$ ). All these species are important for various reasons, and data are needed on their reactions. Such data can then be used as a basis for predicting reaction rate constants for species that have not, as yet, been studied experimentally.

### B. Experimental

The experiments were performed in four series, each differing from the other in various ways, mostly due to improvements in technique. The Series A was performed using a flow-tube apparatus and was essentially unsuccessful, but Series B, C, and D were performed using variations on the flash-photolysis resonance-fluorescence (FPRF) technique coupled with mass spectrometric measurement of (PNA), and they were fully successful. Although the flow-tube experiments were relatively unsuccessful, much of the apparatus is common to the other series.

## Series A: Flow-Tube Apparatus<sup>14</sup>

The apparatus<sup>15</sup> consists of two main parts, the flow system and the mass spectrometric detection system. The flow system consists of a flow tube, a gas-metering system, a mass spectrometric-sampling device, and a high-capacity pump. As shown in Figure 1, the main flow tube is a Pyrex tube, 1-meter long and 2.5-cm i.d., equipped with a manifold to prepare the main flow mixture before it is introduced into the flow tube. Typically, helium mixed with a small amount of hydrogen constitutes the main flow which is then passed through a microwave discharge (2450 MHz) before being introduced into the flow tube. A Wood's horn is used to prevent light generated by the microwave discharge from entering the main flow tube. The He flow is controlled with a needle valve and is measured by a Fisher-Porter rotameter, and reactant flows are measured by the pressure change in a calibrated volume fitted with a Validyne pressure transducer ( $\pm$  0.1 psi).

It is important to maintain the purity of the helium carrier gas because minor impurities easily produce hydrogen-atom concentrations in excess of  $10^{12}$  cm<sup>-3</sup>. For this purpose, ultra-high purity helium (99.999%, Matheson Gas Products, Inc.) was used as carrier gas. Before the helium was admitted to the flow tube, hydrogen-containing impurities were removed by passing the gas over a catalyst (No. R3-11, Chemical Dynamics Corp.) at 200°C followed by three liquid nitrogen-cooled molecular sieve (No. 5A) traps to remove water,  $CO_2$ , or other byproducts. The resulting purity was such that hydrogen-atom concentrations less than  $10^{10}$  cm<sup>-3</sup> were obtained from discharging the carrier gas alone.

A coaxial sliding injector inside the flow tube is used to add the reactant to the main flow. The injector is composed of a 0.3-cm o.d. Pyrex tube, O-ring-sealed to the main flow tube at the upstream end. A stainlesssteel valve and a length of 0.25-inch Teflon tubing connects the injector to the PNA generator. At the end of the Pyrex injector, a small Teflon tip (cap) that has radial pinholes is used to provide mixing of the reactant into the main flow. Both the inside surface of the main flow tube and the outside surface of the movable injector were coated with halocarbon wax (Halocarbon Products Corp., Type No. 15-00) to minimize wall reactions of OH radicals.

To determine the wall-loss rate of the OH, the radicals were generated at different axial positions along the flow tube. In this experiment, helium carrying a small amount of hydrogen was discharged to generate H-atoms at a concentration of  $\sim 2 \times 10^{11}$  cm<sup>-3</sup>. A fixed amount of NO<sub>2</sub>, usually on the order of  $10^{12}$  cm<sup>-3</sup>, is then added through the movable injector to generate known amounts of OH radicals through the reaction H + NO<sub>2</sub>  $\rightarrow$  OH + NO at different positions along the length of the flow tube. The first-order wall-loss rate constants obtained in this manner are usually less than 15 s<sup>-1</sup>.

The pressure in the main flow tube is monitored by a Wallace-Tiernan gauge or by a pressure transducer (Validyne,  $\pm 1$  psi) connected to upstream and/or downstream pressure ports.

The temperature of the reaction region is controlled by the use of heavy aluminum foil and a coil of tubing that circulates fluid from a constant temperature bath. The temperature is monitored by three chromelconstantan thermocouples attached to the outside surface of the main flow tube. The temperatures were constant within  $\pm 2^{\circ}C$  during the experiment.

Sampling of the reactants and the products from the flow tube to the mass spectrometer is accomplished by a Teflon cap that is situated at the downstream end of the flow tube. In the center of the Teflon cap, a 0.010-inch-diameter pinhole, situated close to the center of the flow tube, serves as a molecular beam source; the beam is modulated by a mechanical chopper and detected by the mass spectrometer. The pressure in the first pumping chamber after the pinhole is typically  $\sim 1-3 \times 10^{-5}$  torr, while the pressure in the second chamber (containing the Finnigan quadrupole) is  $\sim 10^{-6}$  torr. The same modulated-beam mass spectrometer has been used for detection in various other studies, such as very low-pressure pyrolysis and very low-pressure photolysis, and a description can be found elsewhere.<sup>16</sup>

Pumping of the main flow is achieved by a mechanical pump (Alcatel 1P4A or Welch 1397) connected to the system through a stopcock and ballast bulb to damp-out any pressure fluctuations caused by the pump. Molecular sieve (Union Carbide 4X) is placed in the bulb to prevent pump-oil vapor from diffusing back into the flow system. Velocities of 500 to 1000 cm/s are achieved by adjusting both the flow rate of gases and the stopcock openings.

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Although not shown in Figure 1, the flow tube is also equipped with a resonance-fluorescence detection system for OH radicals. The resonance lamp is fitted with a quartz window and is powered by a microwave generator (Opthos Instrument Company) that is operated in a chopped mode. Helium at ~ 2.5 torr is passed over water prior to flowing into the lamp. The OH radical resonance-fluorescence is detected through a window in the flow tube situated at 90° to the orientation of the lamp. An interference filter centered at 3100 Å (100 Å FWHM; Corion Corporation) helps to eliminate scattered light and the fluorescence light is detected by a photomultiplier tube (EMI Model 6255B) whose output is amplified by a lock-in amplifier. For a lock-in amplifier averaging time constant of 10 seconds, the detection limit for OH is ~ 2 x  $10^{10}$  particle/cm<sup>3</sup>. The resonancefluorescence cell is situated about 2-3 cm upstream from the mass spectrometer sampling point.

Pernitric acid was prepared<sup>17</sup> by slowly and carefully dissolving 0.6 g  $NO_2BF_4$  in 2.0 g 90%  $H_2O_2$  that was stirred vigorously and maintained in an ice bath. The solution temperature was then adjusted, and the PNA vapor that evolved was introduced into the flow tube through the movable glass injector. This preparative technique will be described below in more detail. The PNA concentration was determined by monitoring the intensity of m/e = 46 and comparing it to the intensity of a known concentration of  $NO_2$ . The relative sensitivities of PNA and  $NO_2$  were determined by diverting the flow of PNA through a heated inlet and measuring the  $NO_2$  produced in the thermal decomposition of PNA, as described below.

#### Series B: 248-nm Flash-Photolysis Resonance-Fluorescence (FPRF)

The apparatus used in experiment Series B, C, and D is shown schematically in Figure 2. For this setup, the flow tube described above was coupled to a resonance-fluorescence cell as shown in the figure. The cell is not the same one used for Series A and is 70 cm on the long axis to reduce scattered light from the  $CaF_2$  windows that transmit the laser beam. The top and side windows are quartz to transmit the OH radical resonance-fluorescence light centered near 3100 Å. Light from the OH resonance lamp (2450 MHz; 9 torr He; H<sub>2</sub>O) is collimated or weakly focussed with a quartz lens and directed through the side window of the cell. The flow tube is coupled to the cell directly opposite the side window. Since the long axis of the cell is oriented at ~35° from the axis of the flow tube, the light from the resonance lamp does not pass down the length of the tube, but is stopped by the bend of the coupling tubulation. This arrangement acts as a crude Wood's horn to reduce scattered light from the lamp.

The photomultiplier tube was rewired for single photon counting and viewed the OH fluorescence through the top window of the cell. The 3100 Å interference filter was used to reduce scattered light. Output from the photomultiplier was amplified and an amplifier/discriminator was used to detect single-photon pulses. The output of the discriminator was stored in a multichannel analyzer (Nuclear Data, Model ND100) that was operated in the multi-channel scaling mode. After several thousand laser shots, the memory contents were recorded on an x-y chart recorder and were then analyzed to determine the fluorescence decay rate.

The gas flow entered the cell at the port opposite the side window and then turned  $90^{\circ}$  downward through the exit tube that is opposite the top window. About 8 cm from the optical center, the gas stream was sampled through a Teflon cap, perforated with a hole ~ 0.008-inch in

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diameter that acted as the originating orifice of the modulated molecularbeam mass spectrometer described above. At the exit port of the cell (see Figure 2), a 10-mm glass or Teflon-bore stopcock fitted with viton O-rings was used as a throttle valve to control gas flow velocity and total pressure. From this valve, the gas flow was conducted directly to the main flow-tube pump.

In these experiments, the flow tube was used primarily as a gas-mixing manifold; linear flow velocities were usually of the order of 0.5-3 m/s. but faster velocities were used occasionally. Concentration calibrations for  $NO_2$ ,  $H_2$ , and  $HNO_3$  were performed by measuring the rate of pressure increase in calibrated volumes, as described previously. Dry  $HNO_3$  was prepared by mixing  $NaNO_3$  with concentrated  $H_2SO_4$  in an evacuated bulb. Ozone concentration calibrations were performed daily by measuring the attenuation<sup>18</sup> of light from a Hg 2537 Å penlight source viewed through an interference filter by a 1P28 photomultiplier down the long axis of the photolysis cell.

Ozone was prepared in a separate vacuum line by an electric discharge in  $O_2$  and was trapped and stored in a silica gel-packed trap maintained at  $-78^{\circ}$ C. The ozone concentration was controlled by sweeping it out of the trap with a stream of helium diverted from the main flow. Mass spectrometric analysis showed that  $H_2O$  vapor was released from the trap along with  $O_2$  and  $O_3$ .

Pulses of OH radicals were generated according to the following scheme:

$$O_3 + h_V(249 \text{ nm}) \rightarrow O(^1\text{D}) + O_2$$
 (3)

$$O(^{1}D) + H_{2} \rightarrow OH + H$$
(4)

$$O(^{1}D) + PNA \xrightarrow{i} OH + O_{2} + NO_{2}$$
 (5)

$$H + O_3 \rightarrow OH(vib) + O_2 \tag{6}$$

The 248-nm light was generated by a Lambda-Physik excimer laser (Model EMG 101) operated with a mixture of krypton, fluorine, and helium. Output energies were measured with a Scientech calorimeter-type power meter (Model 360203) and averaged ~ 80 mJ/pulse, of which energy about 25% actually entered the cell after collimation. Typically, pulse repetition frequencies used were ~ 5-10 Hz. Since gas residence times in the cell body were ~ 1/2 sec, the  $O_3$  was significantly depleted by the laser beam, but the PNA concentration showed less than ~ 5% depletion, demonstrating the validity of pseudo-first-order conditions. It was found by experimentation that the observed OH fluorescence decay rates were independent of  $O_3$  pressures (at this laser pulse energy) as long as they were less than ~ 1 mTorr, corresponding to an initial OH concentration of  $\leq 2 \times 10^{13}$  radicals/cm<sup>3</sup>. Hydrogen concentrations were high enough (2 x 10<sup>15</sup> atoms/cm<sup>3</sup>) so that all of the O(<sup>1</sup>D) reacted by reaction (4) within ~ 10 µsec after the laser pulse.<sup>19</sup>

The troublesome probability existed that vibrationally excited OH was present initially, and an observed spectrum showed our resonance lamp could excite OH(v=1), if present. Thus, the fluorescence data for times shorter than 2.5 msec after the laser pulse were ignored to allow some time for vibrational relaxation of OH(vib). Several spikes due to electronic pickup from the laser appeared on each decay curve, providing a convenient time marker for the 2.5 msec delay.

The relative mass spectrometric sensitivities of PNA and  $NO_2$  at m/e = 46 were measured in a series of experiments described below, and these data were used with the daily measurements of absolute  $NO_2$  sensitivity to determine the absolute PNA concentrations.

In this series of experiments, it was found that PNA decomposition on the pyrex surface could be somewhat inhibited by application of a thin coating of halocarbon wax. This treatment permitted the incorporation of

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a cold-trap in the PNA preparation train that helped to reduce unwanted  $H_2O_2$ . Moreover, higher total pressures were obtainable without excessive decomposition of the PNA. The primary high pressure limitation, however, was the residual gas pressure tolerable for the mass spectrometer.

# Series C: 266-nm FPRF<sup>20</sup>

This series of experiments was performed in a manner similar to Series B, with a few differences. The light source for this series was a Quanta Ray Nd:YAG laser whose output was quadrupled in frequency to give 266-nm light (~ 20 mJ/pulse, 90% of which entered the cell). The source of OH was:

$$O_3 + h\nu(266 \text{ nm}) \rightarrow O(^1\text{D}) + O_2$$
 (3)

$$O(^{1}D) + H_{2}O \rightarrow 2OH$$
 (7)

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This source of OH was "cleaner" than in Series B, because the initial vibrational excitation in OH produced by reaction (7) is quite low,<sup>21</sup> and there is no subsequent reaction of H +  $O_3$  to produce highly vibrationally-excited radicals. Due to the absence of vibrationally-excited radicals, the data analysis could be carried out for all data from t = 0. The initial OH concentration is estimated to be  $\leq 2 \times 10^{13}$  radicals/cm<sup>3</sup> for 1 mTorr  $O_3$ .

PNA was prepared and its concentration was determined as described previously, except that a new quadrupole mass spectrometer (Balzers Models 311,140) was substituted for the Finnigan instrument. The new equipment performed very well, and a new series of relative mass spectrometer sensitivity measurements was performed.

One additional change was made in the data analysis procedure. The trigger sequence was arranged so that background fluorescence was recorded for several tenths of a millisecond before the laser was pulsed, enabling us to digitally subtract the background light intensity. After subtraction of background, the memory contents of the MCA were passed through a logarithmic amplifier prior to the x-y recorder. These steps considerably reduced the time consumed in the data analysis process.

#### Series D: 266-nm FPRF

This series of experiments was performed in just the same way as Series C, except that the lens elements of the mass spectrometer were cleaned and the electron multiplier was rejuvenated (because of an accidental vacuum failure), necessitating a new series of relative sensitivity measurements.

### PNA Preparation, Purity, and Concentration Determination

The preparation, purity, and concentration determination of PNA require a separate section because this aspect of the experiments is the single largest source of error and uncertainty.

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Our experience is that PNA tends to decompose relatively rapidly (seconds) on pyrex and metal surfaces. Halocarbon wax and teflon tend to be more inert, but some decomposition still takes place. Due to this behavior by PNA, we did not use static mixtures of PNA in storage bulbs, but generated the PNA batch-wise and mixed it with the gas stream in the flow tube. Small preparative batches of PNA are advised because, during this study (CAUTION!!) two minor explosions took place, one of which destroyed a heated glass tube, even though the partial pressure of PNA was probably less than 5 Torr, and the total pressure was less than 20 Torr.

As described eisewhere, PNA was prepared<sup>17</sup> by cautiously and slowly adding 0.6 g  $NO_2BF_4$  (Alfa-Ventron) to 90%  $H_2O_2$  maintained at  $O^0C$  and rapidly stirred. This preparative solution was kept at  $O^0C$  until the

small vessel (~ 25 cm<sup>3</sup>) was O-ring sealed to the preparative train and pumped down to the flow tube pressure. PNA was carried out of the preparative vessel by a stream of helium (diverted from the main flow) that was passed over the liquid surface. Bubbling the carrier gas through the solution did not significantly improve the PNA yield. The flow-tube PNA concentration was controlled by varying the helium flow rate and by varying the bath temperature of the PNA preparative solution up to ~  $15^{\circ}$ C; it was deemed too dangerous to allow the temperature to go higher.

In experimental Series B, C, and D, a U-trap followed the preparation vessel, and when held at a temperature of  $\sim -15^{\circ}$ C, it seemed to reduce the relative amount of impurity  $H_2O_2$  that was released.

Potential impurities, such as  $NO_2$ ,  $H_2O_2$ , and  $HNO_3$ , were specifically sought using the mass spectrometer, and it was found that  $HNO_3$  either was not present or its level was < 5% of the PNA (see below). Hydrogen peroxide appeared as a highly variable impurity, ranging from undetectable levels (m/e = 34) up to concentrations several times as large as the PNA concentration.  $NO_2$  was often present as an impurity, but its level was usually  $\leq 10\%$ , and never more than ~ 50% of the PNA concentration. As discussed later, the reaction rate of OH with  $NO_2$  at the total pressures of He used here are slow, and thus the effect of  $NO_2$  is small.  $H_2O$  and  $O_2$  were also impurities, but were not monitored.

The basic idea behind the relative mass spectral sensitivity measurements for PNA and  $NO_2$  is that the thermal decomposition of PNA quantitatively yields  $NO_2$ :

$$HOONO_2 \xrightarrow{\text{heat}} HOO + NO_2$$
 (8)

$$2 HO_{2} - H_{2}O_{2} + O_{2}$$
(9)  
$$H_{2}O + 1/2 O_{2}$$

The experimental procedure was as follows:

For a constant flow of PNA,

- (1) Measure the mass spectral intensity at m/e = 46: I,
- (2) Using the flow-tube microwave-discharge, add O-atoms and titrate away any  $NO_2$  impurity using the  $NO_2^*$  afterglow technique; measure the mass spectral intensity:  $I_2$
- (3) With the O-atom concentration from step (2), divert the PNA flow through the heated bypass [a 1/4-inch x 50-cm stainless steel (because of explosion hazard) U-tube maintained at  $300^{\circ}$ C], and measure the mass spectral intensity: I<sub>3</sub>
- (4) Add more O-atoms to titrate any  $NO_2$  produced by the thermal decomposition of PNA, and measure the mass spectral intensity:  $I_4$  (Titration of  $NO_2$  by O-atoms is an effective technique, because the rate constant  $O + NO_2$  is about three orders of magnitude faster than that of the reaction O + PNA. As long as a modicum of care is taken, PNA is not depleted by reaction with O-atoms under these conditions.)

The difference  $(I_3 - I_4)$  is proportional to the amount of NO<sub>2</sub> produced by the thermal decomposition of PNA. The difference  $(I_2 - I_4)$  is proportional to the amount of PNA decomposed. The ratio of these quantities is related to the mass spectrometer sensitivities for the two compounds:

$$\frac{I_3 - I_4}{I_2 - I_4} = S^{-1} \tag{8}$$

**a** nd,  $[PNA] = S^{-1} \alpha_{NO_2} \cdot I_2$ , where  $\alpha_{NO_2}$  is the measured NO<sub>2</sub> sensitivity (molecules cm<sup>-3</sup>/millivolt).

To determine whether  $\text{HNO}_3$  contributes to the observed mass spectral intensity of m/e = 46, the PNA is diverted through the heated bypass and a great excess of O-atoms (i.e., >  $10^{14}$  atom/cm<sup>3</sup>) is added. Under these conditions (reaction time ~ 1 sec), it is observed that virtually all of the intensity at m/e = 46 disappears, indicating the absence of  $\text{HNO}_3$  since

 $HNO_3$  does not thermally decompose under these conditions, and it reacts only very slowly with O-atoms. On the other hand, PNA both thermally decomposes and reacts with O-atoms at a moderate rate ( $k \sim 10^{-15}$  cm<sup>3</sup> sec<sup>-1</sup> at 300 K)<sup>10</sup> All experiments of this type that were performed showed negligible amounts of  $HNO_3$ .

It was concluded that the thermal decomposition of PNA under our conditions quantitatively gives NO, for the following reasons:

- (1)  $HNO_3$  was shown not to be a significant product.
- (2) The ratio of intensities at m/e = 46 and at m/e = 30 were consistent with product  $NO_2$ , although the experimental scatter was large due to PNA interference and due to taking the differences of relatively noisy values.
- (3) Experiments in which mixtures of  $NO_2$  and  $H_2O_2$ were passed through the heated bypass showed that the  $NO_2$  was unaffected by the treatment.
- (4) In the highly oxidizing environment containing  $HOONO_2$  and  $H_2O_2$ , it seems unlikely that the PNA decomposition product would be HONO or other reduced forms of nitrogen.

The results obtained for the relative sensitivities of  $NO_2$  and PNA in experiment Series B, C, and D, are presented in Table 1.

Experimental	S ± 1σ	Data Points	Mass Spectrometer
B	7,48 ± 1.05	21	Finnigan
С	4.44 ± 0.74	9	Balzers
D	5.07 ± 0.34	14	Balzers

Table 1 RELATIVE MASS SPECTROMETER SENSITIVITIES FOR PNA AND NO2

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### C. Results and Discussion

To attain pseudo-first-order conditions, PNA was used in excess in all the experiments. Under these conditions, the reaction can be written:

$$\begin{array}{c} k_2 \\ \text{OH + PNA} \xrightarrow{} \text{Products} \end{array} (2)$$

where  $k_2$  is the bimolecular rate constant for the reaction. The rate expression for the removal of OH can be written:

$$\frac{d[OH]}{dt} = -k_2[OH][PNA] = -k^{I}[OH]$$
(10)

where  $k^{I}$  is the pseudo-first-order constant and equal to  $k_{2} \times [PNA]$ . For the flow-tube experiment Series A, the plug flow approximation is valid, and time resolution is achieved by varying the mixing distance of the two reacting species. Therefore,  $k^{I}$  is expressed as:

$$k^{I} = -\frac{d\ln[OH]}{dt} = -\frac{v}{v} \frac{d\ln[OH]}{dx}$$
(11)

where the average flow velocity,  $\overline{v}$ , depends on temperature, pressure, total flow rate, and the cross section of the flow tube. Therefore,  $k^{I}$  is the product of the average flow velocity and the slope of the semi-log plot of disappearance of [OH] versus mixing distance. The bimolecular rate constant,  $k_2$ , is then obtained from the slope of the plot of  $k^{I}$  versus [PNA]. As an alternative method to varying the reaction time for fixed concentrations of PNA, the time may be held fixed while the concentration of PNA is varied:

$$k_{2} \frac{x}{\overline{v}} = -\frac{\ln([OH]/[OH]_{0})}{[PNA]} = k_{2}t = z \qquad (12)$$

Therefore, the slope of the plot of ln[OH] versus [PNA] for fixed time gives the product  $k_2t$ . The bimolecular rate constant is then obtained from the slope of a linear plot of z versus t. The latter method was used in the present study, because it was found that an inordinate amount of time was necessary for the signal to stabilize after moving the probe position. This behavior is symptomatic of wall reactions involving OH, PNA, or both species. Moreover, the use of the second method was necessitated because during the long periods of time necessary for a complete experiment, the PNA concentration often varied significantly, necessitating corrections for such variations.

It is noteworthy that although the first-order plots of [OH] decay in the flow tube were very well behaved, the second-order plots show considerable scatter. Indeed, the second-order plot for 275 K showed two distinct groups of data and, for any given day, the data seemed to fall in one or the other of them at random.

Considerable effort was spent in attempting to isolate the cause of the data scatter or to eliminate it. In most of our other work with PNA, the major source of error and experimental scatter is in measurements of the PNA concentration. If this was the cause of the scatter in the present work, the first-order plots according to Method II should show such scatter, since the PNA concentration enters directly. But for these experiments, there is not an inordinate amount of scatter, and the plots give good straight lines. Thus, uncertainties in PNA concentration are probably not at fault.

Another potential problem is the possible production of major quantities of impurities in some batches of PNA. This possibility can be eliminated, however, since two or more batches of PNA were used on each day of experiments, and all of the experiments for any single day were self-consistent.

In an effort to test our general technique, we measured the reaction rate constant for the reaction OH + isobutane  $\rightarrow$  products. Good linear first-order plots and second-order plots were obtained that gave  $(k \pm 1\sigma) =$  $(2.2 \pm 0.3) \times 10^{-12} \text{ cm}^3 \text{ sec}^{-1}$ , in good agreement with literature values.<sup>22</sup> The agreement indicates that our experimental techniques are satisfactory and that there are no unrecognized errors of technique that are affecting the rate constants obtained when PNA is a reactant.

After many efforts to eliminate the anomalous scatter in the data, we have concluded that this behavior may be due to a wall reaction of OH and PNA that is dependent on the PNA concentration and/or  $H_2O_2$  concentration and the history of the flow tube. Although we cleaned the flow tube and recoated it with a fresh layer of halocarbon wax several times, the problem persisted. We feel that of the various wall coatings available, halocarbon wax is among the least likely to cause adsorption of PNA, and thus is one of the best for inhibiting wall reactions. Since it has turned out that this coating is not effective, we concluded that further experiments using the flow-tube technique on this reaction would not be fruitful, and we turned to the FPRF approach. In anticipation of the FPRF results, it is of interest to note that the flow-tube results give a bimolecular rate constant  $(k_2 \pm 1\sigma) = (3.3 \pm 0.9) \times 10^{-12}$  cm<sup>3</sup> sec<sup>-1</sup> at room temperature.

For the FPRF technique, the OH concentration change with time is conventionally written: $^{23}$ 

$$\frac{d[OH]}{dt} = -(k_2[PNA] + k_p)[OH]$$
(13)

and the inverse decay lifetime of the fluorescence intensity  $I_f$  is:

$$\tau^{-1} = k_2 [PNA] + k_D \tag{14}$$

where  $k_2$  is the bimolecular reaction rate constant, and  $k_p$  is a parameter that describes the effective "rate constant" for loss of OH due to physical processes. For the particular conditions in the present experiments, the physical processes include the translational motion of gas parcels as they pass into and out of the intersection of the detector field of view and the region illuminated by the resonance lamp. Such motions are due to the gas-stream velocity as it is pumped through the cell and to mass diffusion of [OH]. Neither of these processes is exactly represented by the first-order "rate constant"  $k_p$  since the pumping velocity is clearly independent of the OH concentration and spherical diffusion subsequent to excitation in a cylindrical profile coupled with crossed-optical-beam detection is not expected to be so simple. Nevertheless, equation (14) is a good, pragmatic representation as long as the expression is employed for data taken at the same average elapsed time following the laser pulse.

Under pseudo-first-order conditions, a plot of log  $I_f$  versus time will give a nearly straight line, if  $k_p$  is only slightly dependent on time. Two such plots are shown in Figure 3 for experiment Series C. For Series C and D, the protocol was established that the initial slopes on first-order plots be taken; this protocol is supported because secondorder plots of  $\tau^{-1}$  versus reactant concentrations give good straight lines. Two such examples are presented in Figures 4 and 5 for the reaction:

$$OH + H_2O_2 \rightarrow H_2O + HO_2$$
(15)

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and for

$$OH + NO_{2} + He \rightarrow HONO_{2} + He$$
 (16)

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Both second-order plots show very good straight-line behavior. The absolute rate constant for reaction (15) was not determined only because the mass spectrometer could not easily be calibrated for sensitivity to  $H_2O_2$ . The absolute rate constant for reaction (16), determined from the data in Figure 5, is 1.6 x  $10^{-30}$  cm<sup>6</sup> molecule<sup>-2</sup> s<sup>-1</sup>, in excellent agreement with literature data.<sup>24</sup> Similar tests were performed using isobutane as reactant, also leading to rate constants in excellent agreement with literature values.<sup>22</sup> The test reaction rates determined are summarized in Table 2.

		Results of 1	Present Study
Reaction	T(K)	P(torr)	k <sup>II</sup> (cm <sup>3</sup> s <sup>-1</sup> )
$OH + iC_4H_{10}$	298	9.2	$3.6 \times 10^{-12}$
	324	10.1	$(3.62 \pm 0.40) \times 10^{-12}$
	267	10.1	$(2.70 \pm 0.20) \times 10^{-12}$
OH + NO <sub>2</sub> + He	246	15.1	$(9.23 \pm 0.22) \times 10^{-13}$

Table 2 TEST REACTIONS

For experiment Series B, the protocol for data treatment was somewhat different for two reasons. First, because it was felt that OH(vib) could cause problems, the first 2.5 msec of data from each run were ignored, and the first-order plots were constructed from data beginning at 2.5 msec after the laser pulse. The second reason for this procedure was that two or three "spikes" due to electronic pickup from the excimer laser appeared in each run, causing some interference. The last spike occurred at 2.5 msec after the laser pulse, providing a convenient time marker.

Because the data for Series B corresponded to lower OH concentrations (due to decay after the pulse), the fluorescence data were significantly

noisier than those of Series C and D, but they, again, gave good straight second-order plots and absolute rate constants, in good agreement with the literature. This finding tends to support the use of the approximately accurate equation (14) for the second-order plots.

In all these series--B, C, D--the intercept of the second-order plots can be identified with  $k_p$ . It was observed that  $k_p$  is larger at lower pressures, as expected for diffusive loss, and it seemed to depend upon day-to-day variation in optical alignment and linear flow velocity through the optical intersection region. N further significance is placed on  $k_p$  in the present work.

When  $H_2O_2$  and  $NO_2$  are present as impurities, equation (14) must be generalized to:

$$\tau^{-1} = k_2 [PNA] + k_{16} [He] [NO_2] + k_{15} [H_2O_2] + k_P$$
(17)

As mentioned earlier, the  $NO_2$  impurity is relatively small, and at the pressures and temperatures of the present study, its effect on the observed  $\tau$  is also small and can be neglected (see below). Since  $k_p$  varies on a day-to-day basis, the best way of combining all of the data for a given temperature and pressure is to consider only the net first-order rate constant resulting from the following expression:

$$k_{\text{net}}^{I} = k_{2}[PNA] = \tau^{-1} - k_{15}[H_{2}O_{2}] - k_{p}$$
 (16)

For experimental runs performed using just  $H_2O_2$ , we have:

$$k_{H_2O_2}^{I} = k_{15} [H_2O_2] + k_p$$
 (17)

For each set of conditions, runs were cauried out using only  $H_2O_2$  as reactant, and a linear least-squares analysis gave values for  $k_{15}$  and  $k_p$ . Since  $H_2O_2$  concentration was measured in arbitrary units,  $k_{15}$  has units

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of (arbitrary unit)<sup>-1</sup> s<sup>-1</sup>;  $k_p$  is obtained in units of s<sup>-1</sup>. Once  $k_p$  and  $k_{15}$  are known,  $H_2O_2$  and PNA concentrations are measured mass spectrometrically and recorded along with  $\tau^{-1}$ . From these quantities,  $k_{net}^{I}$  is determined, and second-order plots of  $k_{net}^{I}$  versus [PNA] can be constructed as shown in Figure 6.

All of the data obtained in experiment Series B, C, and D are tabulated in the Appendix.

The data for 298 K presented in Figure 6 were obtained on several different days for three different total pressures and several different linear flow velocities. Clearly, the data are quite consistent with one another, showing no significant dependence on total pressure. For the purpose of illustration, the data for 246 K are presented in Figure 7, and again, there is no significant dependence on total pressure.

From the  $O_3$  concentration, laser energy, and ozone absorption coefficient,<sup>18</sup> we can estimate that the initial OH concentration does not exceed ~ 2 x  $10^{13}$  radicals/cm<sup>3</sup>. Several tests were performed to determine whether the observed fluorescence decay rates depended upon either the source of OH radicals or on their initial concentration. In experiment Series B, the  $O_3$  partial pressure was varied from 0.23 to 0.8 mtorr with no perceptable variation in either decay rate or in the shape of the fluorescence decay curve, although the absolute fluorescence intensity varied. When the  $O_3$  partial pressure was raised to 3 mTorr, a faster decay rate occurred at early times, but by the 2.5 msec time marker, there was little effect.

In experiments Series C, more extensive tests were performed. With 80 mTorr  $H_2O$ , the  $O_3$  partial pressure was varied from 0.4 to 4.0 mTorr, resulting in a variation of ~ 10% in the OH +  $H_2O_2$  bimolecular rate constant. The 4 mTorr  $O_3$  data for OH + PNA may have been ~ 15% lower than the usual, but this variation is not significant and the data were included in the overall data set.

In another series of test runs in Series C,  $H_2$  was substituted for  $H_2O$ , to reproduce the conditions of experiment Series B. Substitution of 85 mTorr  $H_2$  for the usual  $H_2O$  gave no significant difference in observed rate constant. Indeed, runs with <u>no</u>  $H_2$  or  $H_2O$  added to the gas stream also agreed with the rest of the data. This result can be understood, since reaction of  $O(^{1}D)$  with PNA may give OH and since  $H_2O$  was observed evolving both from the PNA preparation train and from the  $O_3$  trap. Thus, it appears that  $H_2O$  was always present in significant quantities. Since  $H_2O + O(^{1}D)$  gives relatively "cool" OH, and because  $H_2O$  is an efficient quencher for vibrationally excited OH, it appears that if any residual vibrational excitation remained, it played no significant role in the observed rate constants.

In addition to varying the  $O_3$ ,  $H_2O$ , and  $H_2$  partial pressures, the laser power was also varied in experiment Series C. When the laser pulse energy was reduced by about a factor of 10 from the usual 20 mJ/pulse, there was a reduction in fluorescence intensity, but no significant variation in observed rate constant. Thus, we may conclude that (1) secondary reactions of OH and reaction products are not significant, (2) depletion of PNA concentration is not significant, and (3) OH disproportionation is not significant. Thus, the system appears to be kinetically well behaved.

The measured bimolecular rate constants for each temperature are summarized in Table 3 and Figure 8, along with least-squares and propagation of errors estimates of the associated experimental uncertainties. The first uncertainty listed for each rate constant is just the statistical uncertainty associated with the second-order plot of  $k_{net}^{I}$  versus nominal [PNA]. The second uncertainty listed is that associated with the PNA/NO<sub>2</sub> relative mass spectrometric sensitivity. The third uncertainty listed is the propagation of errors combination of these two sources of error.

T(K)	Data Points	$10^{12}$ k cm <sup>3</sup> /sec	10 <sup>12</sup> FPRF	10 <sup>12</sup> NO <sub>2</sub> /PNA	10 <sup>1 2</sup> J net
Series B:	248 nm FPRF				
251	11	3.84	0.28	0.54	0.61
273	17	4.60	0.41	0.65	0.77
298	23	6.05	0,52	0.84	0.99
316	12	4.95	0.31	0.69	0.76
Series C:	266 nm FPRF				
246	42	4.80	0.46	0.80	0.92
267	22	4.83	0.21	0.81	0.84
298	58	3.61	0.16	0.60	0.62
324	20	3.79	0.21	0.63	0.66
Series D:	266 nm FPRF				
Serres D.					

Table 3 OH + PNA → PRODUCTS: RATE CONSTANTS

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The effect of any NO<sub>2</sub> impurity can be shown to be small. In the worst <u>case</u>, for our lowest temperature (246 K) and highest pressure (15.1 torr), the effective bimolecular rate constant for reaction of NO<sub>2</sub> with OH is  $k_{16}$  [He] = (9.23 ± 0.22) x 10<sup>-13</sup> cm<sup>3</sup> sec<sup>-1</sup> (Figure 4). If, for our worst case estimate, the NO<sub>2</sub> impurity is 50% of the PNA concentration, the relative contribution of NO<sub>2</sub> to the observed rate constant is less than 10%. Since this worst case scenario is intended to be pessimistic and the effect is smaller than the other sources of error, effects due to the small NO<sub>2</sub> impurity have been neglected.

The data from Table 3 show very little variation with temperature, but a weighted least-squares calculation<sup>25</sup> of the Arrhenius parameters was performed, giving  $(k \pm 1\sigma) = (8.05 \pm 5.69) \times 10^{-12} \exp(-193 \pm 194/T)$ cm<sup>3</sup> s<sup>-1</sup>, covariance = 1.098 x 10<sup>-9</sup>; at the center of our temperature range, these parameters give a one standard deviation uncertainty of about 8%.

Because the temperature dependence of the rate constant is so small, a simple weighted average of the nine rate constants in Table 3 may be of equal usefulness. Such an average gives  $(k \pm 1\sigma) = (3.97 \pm 0.27) \times 10^{-12}$ cm<sup>3</sup> s<sup>-1</sup>. Since the uncertainties quoted refer only to statistical fluctuation and do not include systematic errors, our <u>recommended</u> rate constant and 95% limit uncertainty is:

 $(k \pm 2\sigma) = (4.0 \pm 1.0) \times 10^{-12} \text{ cm}^3 \text{ s}^{-1}$ 

where the uncertainty is assigned by a subjective estimate of systematic error.

Our recommended value of  $(4.0 \pm 1.0) \times 10^{-12}$  can be compared to the Series A<sup>14</sup>result of  $(k \pm 10) = (3.3 \pm 0.9) \times 10^{-12}$  that was rejected due to our suspicions that wall reactions played a role in those experiments.

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Similarily, it can be compared to the "upper limit" rate constant of  $\sim 3 \times 10^{-12}$  estimated by Graham, Winer, and Pitts<sup>4</sup> on the basis of a competitive rate measurement in a smog chamber. More recently, an excellent molecular-modulation spectroscopy study was reported by Littlejohn and Johnston<sup>26</sup> that gave a rate constant  $k_2 = (2 \frac{+2}{-1}) \times 10^{-12}$ . This last result is noteworthy since it depends upon the accuracy and completeness of a reaction mechanism consisting of more than a dozen reactions, and yet the result is in good agreement with the present study. Moreover, the PNA in that study was prepared by an entirely different technique and the reaction mixture was analyzed spectroscopically, rather than by mass spectrometry. Thus, the agreement of these two very different studies is quite satisfactory. Further experiments using the molecular modulation technique may give information on the reaction products, as well.

Reaction	Rate Constant (cm <sup>3</sup> s <sup>-1</sup> )	Reference
ОН + Н О	$(2.51 \pm 0.6) \times 10^{-12} \exp(-126 \pm 76/T)$	29
	$(2.96 \pm 0.50) \times 10^{-12} \exp(-164 \pm 52/T)$	30
OH + HOONO <sub>2</sub>	$ \lesssim 3 \times 10^{-12} $ (4.0 ± 1.0) × 10 <sup>-12</sup> independent of T (2 $^{+2}_{-1}$ ) × 10 <sup>-12</sup>	4 present work 26
OH + HONO <sub>2</sub>	$(1.52 \pm 0.38) \times 10^{-14} \exp(649 \pm 69/T)$	27
OH + HONO	$7 \times 10^{-12}$	31

### Table 4 RATE CONSTANT COMPARISONS

In Table 4 the present results are compared to rate constants measured for  $H_2O_2$ , HONO, and HONO<sub>2</sub>. The most perplexing of these rate constants is the recent result<sup>27</sup> for OH + HNO<sub>3</sub>, since it exhibits a "negative activation energy." (The Arrhenius form of data expression is probably not appropriate for such reaction rate constants, but it is convenient in the present context.) Although the reaction products are not beyond question, indications are that it is a simple abstraction reaction<sup>28</sup> making interpretation of the reaction dynamics even more obscure (a displacement reaction could be explained on the basis of long-range chemical attraction). The OH + PNA reaction rate constant seems to be unexceptional in this group, since no clear trends or interpretations are apparent. The A-factor is appropriate either for abstraction or for displacement or complex formation. Further work should be done on this series of reactions to elucidate the dynamical pathways involved.

### Acknowledgements

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### D. Data Tables

Each of the following tables summarizes the experimental data obtained on a given day and set of conditions. Within each set of data are experiments run with [PNA] = 0, and these were analyzed by the method of least-squares to obtain the slope and intercept for the equation:

$$\tau^{-1} = k_{15} [H_2 O_2] + k_p$$
,

where  $\tau^{-1}$  is the observed first-order rate,  $k_{15}$  is the slope, and  $k_p$  is the intercept, as recorded in the table.

The absolute mass spectrometric sensitivity for  $NO_2$  is determined daily and used along with the  $NO_2/PNA$  sensitivity ratio to give absolute [PNA], as described in the text. The approximate  $NO_2/H_2O_2$  sensitivity ratio was determined using the phenomenological  $H_2O_2$  rate constant, the assumed absolute rate constant 1.6 x  $10^{-12}$  cm<sup>3</sup>/sec, and the measured  $NO_2$ sensitivity. The "net rate" is given by:

$$k_{net}^{I} = \tau^{-1} - k_{15} [H_2 O_2] - k_p$$

All of the data were used in the evaluation of the rate constants unless otherwise indicated. Specifically, the "test data" of 18 November 1980 were not used, although they are consistent with the rest of the data. Also, data were arbitrarily rejected if  $k_{net}^{I} \leq 0.5 k_{15}[H_2O_2]$  for any given run.

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CH + PERNIIRIC ACID DAIA--Beries B

TEMP = 298, KELVIN KELUN TOTAL PRESSURE = 3.40 TORR ND2 SENSITIVITY = 2.894E+11 NOLECULES/CE/NV ND2/PNA SENSITIVITY RATIO = 7.48

MPROXIMATE NO2/H2O2 SEMSITIVITY RATIO =

0.4466

NCOZ PNEMDNEMOLDEICAL RATE LEAST-SQUANES; SLOPE = 6.4800 Intercept = 295.0 Sec-1

OH + PERNITRIC ACID DATA -- Series B

TEMP = 273.KELVIN MELLUN TOTAL PRESSURE = 3.80 TORR MELLUN TOTAL PRESSURE = 3.306F411 MOLECULES/CC/MV MOZ/PMA SENSITIVITY RATIO = 7.48

0.5800 APPROXIMATE NO2/H2O2 SENSITIVITY RATIO =

H2O2 PHENNONENOLDGICAL RATE LEAST-SQUARES: SLOPE = 5.7000 INTERCEPT = 307.0 SEC-1

OX MTORR) H2X E-15 H202(ARB, UNITS) PNA(MOLEC./CC) DBS. RATE(SEC-1) MET RATE(SEC-1)

TN XCO	(JRR )	H2X E-13	5 H202	(ARB. UNITS)	PNAK MOLEC . / CC )	ODS. RATE(SEC-1)	NET RATE(SEC-1)	03K HTORR ) H2	X E-15 H2	02(ARB. UNITS)	PNAK MOLEC . /CC )	OBS. RATE(SEC-1)	NET RATE(S
	0.2	7.2.	.10	0.00	0.0005-01	226.	-59.	0.23	3.40	20.01	0.000E-01	446.	25. •
	2.3	°.	.10	8.0	0.000E-01	363.	78.	0.23	3.40	31.78	0.000E-01	478.	-10.
	1.9	0 2.	<u>9</u>	3,95	3.332E+13	423.	112.	0.23	3.40	62.62	0.000E-01	607.	-57. 0
		7 2.	.10	0.00	0.000E-01	293.	.8	0.23	3.40	86.16	0.000E-01	710.	-88.
		7 2.	.10	4.41	5.JB1E+13	570.	222	0.23	3.40	3.06	5.303E+13	369.	<b>4</b> 5.
	0.8	7 2.	.10	5.13	9.44 E+13	668.	366.	0.23	3.40	1.98	6.807E+13	492.	174.
	.8.0	7 2.	97.	4.14	7.5056413	637.	325.	0.23	3.40	1.41	3.959E413	467.	152.
	8.0	7 2.	.10	4.24	3.8266413	508.	196.	0.23	3.40	0.94	1.550E+13	260.	-52.
		7 2.	9.	4.24	2.536E+13	329.	17.	0.23	3.40	3.77	4.291E+13	562.	234.
	0.8	7 2.	.10	7.06	3.300E+13	597.	266.	0.23	3.40	12.19	1.387E+14	929.	553.
	0.8	7 2.	.10	7.72	4.717EH13	680.	345.	0.23	3.40	0.0	0.000E-01	237.	-70.
	6.0	7 2.	10	4.05	4.838E+13	574.	263.	0.70	3.40	0.0	0.000E-01	297.	-10.
1	0.8	7 2.	•10	1.51	7.9556412	311.	16.	0.70	3.40	0.00	0.000E-01	322.	15.
0		7 2.	9.	ft.4	3.294E+13	490.	177.	0.70	3.40	9.65	4.640E413	412.	50.
0	0.8	7 2.	.10	6.40	3.1336413	532.	206.	0.70	0+ · E	10.59	4.595E413	438.	71.
	.8.0	~ ~	9.	47.08	5.727E+13	784.	194.	0.70	3.40	35.31	1.5916414	1274.	766.
	8.0	7 2.	9.	37.19	5.301E+13	910.	384.	0.70	3.40	12.95	6.1896413	576.	195.
		7 2.	-10	36.49	4.492E+13	920.	399.	0.70	3.40	8.94	5.1246413	469.	111.
	8.0	7 2.	-10	48.96	1.029E+14	1207.	605.	0.70	3.40	3.77	2.212E+13	317.	-11.
	9.0	2.	.10	0.0	0.000E-01	274.	-11.	0.70	3.40	3.30	1.460E413	297.	-29.
	.0	7 2.	•10	0.0	0.000E-01	33.	10.	0.70	3.40	44.26	0.000E-01	660.	101. 0
	0.8	7 2.	.10	12.48	2.246E+13	412.	46.	0.60	0 <b>1</b> .40	40.02	0.000E-01	544.	9. 8
	0.8	7 2.	.10	69.68	1.4516+14	1752.	1015.	0.60	3.40	74.39	0.000E-01	782.	51. •
	8.0	7 2.	9	6.78	9.4526412	310.	-19. 8	0.60	3.40	77.21	0.000E-01	776.	29. 4
	0.8	7 2.	-10	240.11	6.111E+13	2148.	307. 8	0.60	3.40	0.00	0.000E-01	310.	'n.
	8.0	7 2.	<u>9</u>	100.75	0.000E-01	978.	40. 0						
	0.6	7 2.	2	16.79	0.000E-01	849.	-21. 0						
	8.0	7 2.	.10	43.78	0.000E-01	523.	-46. 8	<b>B MET RATE LES</b>	S THAN 50	PERCENT OF H202	RATE		
								# PURE H202 RU	INS FOR THI	S DATE:			
								18-JULY-80					

0 NET MATE LESS THAN SO PERCENT OF M202 RATE 9 Pure M202 Runs for This Date: 17-Jalt-90

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ON + PERNITRIC ACID DATA--Series B

TEMP = 316.KELVIN MELLUNTY = 3.78TOMR MELLUNTY = 2.302F11.MOLECULES/CC/NV MEZ/PMA SENSITIVITY ATTO = 7.46 #PROXIMATE M02/H202 SEMSITIVITY RATIO = 0.4203

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KZO2 PHEMONEMOLDGICAL RATE LEAST-SQUARES: SLOPE = 3,4000 INTERCEPT = 289-0 SEC-1

INTERCEPT = 289.0 SEC-1

03(MTORR) H2X E-15 H202(ARB, UNITS) PHAKMOLEC./CC) OBS. RATE(SEC-1) NET RATE(SEC-1)

- MET 8415							
	481.	1229.	9.357E+13	83.80	3.20	0.75	
0.6	341.	867.	6.464E113	43.31	3.20	0.75	
0.8	311.	.661	5.2396413	23.42	3.20	5/*0	
9.0	266.	645.	4.003E113	14.48	3.20	0.75	
0.9	75.	413.	1.6586413	8.94	3.20	6.75	l
	69.	422·	2.017EH13	12.24	3.20	0.75	01
0.9	<b>6</b> .	33.	0.000E-01	0.00	3.20	0.75	1
8.0	20. 8	383.	1.943E413	13.42	3.20	0.75	
8.0	373.	805.	5.851E413	<b>12.</b> 11	3.20	0.75	
0.0	358.	870.	8.9246413	44.26	3.20	0.75	
0.8	786.	1374.	1.5246+14	54.61	3.20	0.75	
0.8	-10.	279.	0.0005-01	0.00	3.20	0.75	
0.9	3.	. <b>11</b>	0.0005-01	0.00	3.20	0.75	
0.8	-23. •	-98-	0.0005-01	40.49	3.20	0.75	
0.0	-23. +	674.	0.0005-01	74.39	3.20	0.75	
0.9	-36. •	53.	0.0005-01	73.44	3.20	0.73	
8.0	52. 0	1063.	0.000E-01	131.82	3.20	5.5	
0.0	1. •	336. 2	0.000E-01	48.49	3.20	50	

A MET RATE LESS THAM SO PERCENT OF H2O2 RATE 9 PUME H2O2 RUNS FON THIS DATE: 19-JULY-80

OH + PERNITRIC ACT" DATA -- Series B

TEMP = 251.KELVIN KELIUM TOTAL PRESSURE = 3.94 TORR NO2 SENSITIVITY = 2.560E+11 MOLECULES/CC/NV NO2/PMA SENSITIVITY RATIO = 7.48 MPROXEMATE NO2/H202 SENSITIVITY RATIO = 0.5378

H2D2 PMEMOMENOLOGICAL RATE LEAST-SQUARES; 5.OPE = 4.7600 Intercept = 290.0 Sec-1 OXIMTORRI) H2X E-15 H2O2(ARB. UNITS) PNA/HOLEC./CC) OBS. RATE(SEC-1) NET RATE(SEC-1)

-37.	6.	-65.	-37. 4	ę. ę	-30. 0	108. 0	-59. 4	18.	17.	564.	146.	164.	252.	222.	B0.	л.
253.	296.	309.	./86	637.	495.	828.	760.	308.	307.	1163.	486.	523.	665.	765.	448.	361.
0.000E-01	0,000E-01	0.000E-01	0.000E-01	0.000E-01	0.000E-01	0.000E-01	0,000E-01	0.000E-01	0.000E-01	1.369E+14	3.391E+13	4.380€+13	5.8176+13	7.255€+13	2.684E+13	0.000E-01
0.00	0.00	17.66	28.25	71.56	64.43	90.39	111.11	0.00	0.00	64.97	10.59	14.59	25.89	53.20	16.4B	0.00
2.11	2.11	2.11	2.11	2.11	2.11	2.11	2.11	2.11	2.11	2.11	2.11	2.11	2.11	2.11	2.11	2.11
0.00	0.90	0.90	0.90	0.80	0.90	0.90	0.80	0.90	0.80	0.00	0.90	0.90	0.90	0.0	0.90	0.00

6 NET RATE LESS THAM SO PERCENT OF H202 RATE 0 Fure H202 Runs for this "Ate: 20-Jul-90

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	5	PERMITRIC	C ACIT DATA	•				5	+ PEKNUTULU	. AULU UNIA 8	eries C		
MEL I MEL I MEZ I	= 298, k un tota, p sensitivit pna sensit	LELVIN PRESSURE = 17 = 7.03 12 TY RATI	9.80 TC 13E+13 MOLE 10 = 4	BRR SCULES/CC 1.44	5. <b>æ</b>			TENP = 298. Helium Total ND2 Sensitivi ND2/Pha Sensi	KELVIN PRESSURE = 117 = 9.3d 111VITY RATI	5,06 Tarr 576413 Nolecul 10 = 4,44	S/CC/W		
A PR	OXIMATE NO	12/H202 SEN	1 1111111	*ATIO =	1.4427		·	APPROXIMATE N	KO2/H2O2 9EN	SITIVITY RATI	1 = 2.0587		
NC:02	PNENDNEND SLOPE = INTERCEPT	N.DG.ICAL RA 78.0000 1 = 505	NTE LEAST-: 1.0 sec-1	SQUARES:				N202 PMENOMEN SLOFE = Intercer	40L0CICAL RA 72.8000 PT = 807	ate least-soum 7.4 sec-1	fESt		
N 160	TORR > N2	OK MTORR )	H2D21 ARD.	UNITS)	PMM NOLEC./CC)	OBS. RATE(SEC-1)	NET RATE(SEC-1)	03( MTORR )	H20( MT0RR )	H202( ARD . UNI	IS) PNAK NOLEC./CC)	ODS. RATE(SEC-1)	NET RATE(SEC-1
	0.30	80.00	7.00	Ő	.000E-01	1060.		0.31	80.00	9.30	0.0006-01	1504.	20. 0
	89	8.8	8.7	<b>6</b> 6	.000E-01 .000E-01	780. 684.	-21. •	0.31	80.08 80.08	12.00 18.40	0.000E-01 0.000E-01	1729.	-17
	0.10	B0.00	0.0	Ó	0006-01	576	-1.	IE O	80.08	1.53	0.0006-01	1988.	- 11-
	0.30	80.00	0.0	Ő	.0006-01	510.	ŝ	0.31	80.00	4.30	0.000 -01	1123.	•
	0.30	80.00	0.69	m	.3266113	689.	130.	0.31	80.00	0.08	8.841E+12	795.	-19.
1	9.9	80.00	12.30	CÍ I	+143124	2450.	986. 201	16.0	80.00	0.04	2.004E+13	782.	-28.
10:	9,9	88	14.10	~ ~	.2046414 .1206413	1,760.	142. 1	17-0 17-0	00.08	0.01	3,3346412	/82. 1405.	-26.
2	2.2	80.00	J. 20	, ny	.069E+13	6664	.151	IE.0	80.00	05.4	1.7936414	1799.	533.
	00	80.00	15.00	-	.B06E+14	2163.	400. 1	15.0	8.0	5.70	2.5326414	2149.	927.
								15.0	88	• • •	1.110E414 · ADTF::/	-2/21	4.30.
×.	T RATE LES	S THAN 50	PERCENT OF	5 H202 RI	ATE			0.31	8.8	10.50	1.07/E114 1.899E414	2302.	301. 730.
2	RE N202 RU	MS FOR THE	IS DATE:					0.31	80.08	10.50	1.7306+14	2124.	552.
12-W	08-00							0.31	80.00	1.20	4.747E+13	1027.	132.
								15.0	8.8	0.60	3.9246413	1062.	211.
								15.0	00.08	0, 10	2.8486413 	628.	. ZA.
								12.0	8.8	0.10	7.384€+12	822.	
								11.0	8.8	6, 30	2.5956414	1016. 2160.	50/. 1994.
								0.31	80.00	4.60	2.0896+14	1837.	495.
								11.0	8.8	2.61	1.076E+14	1243.	246.
								17:0	80.00 1	1.20	4.8736413		<b>6</b> 2.
								12.9	M0.00	c/*0	3.000113	.653.	-9. 8

A NET RATE LESS THAN SO PERCENT OF H202 RATE 9 Pure H202 Runs for This date: 13-MDV-80

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OH + PERNITRIC ACID DATA--Series C

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TEMP = 290. KELVIN MELIUN TOTAL PRESSURE = 10.15 TORR MEZ SENSITVITY = 7.280E413 MCLECULES/CC/NV MG2/PMN SENSITVITY RAITO = 4.44

1.4726 #PROXIMATE NO2/H2O2 SENSITIVITY RATIO =

N202 PHEMONEMOLDGICAL RATE LEAST-SOUMMES: 54.0PE = 79.1000 Intercept = 447.0 Sec-1

TEMP = 299. KELVIN HELIUN TOTAL PRESSUME = 10.10 TORR MOZ SEMSTITUTY = 4.330E+13 MOLECULES/CC/WV MOZ/PHAN SENSTITUTY RAITO = 4.44 APPROXIMATE NO2/H202 SENSITIVITY RATIO =

OH + PERNITRIC ACID DATA--Beries C

1.3817

NZO2 PHEMONEMOLOGICAL RATE LEAST-SQUARES: SLOPE = 73.3000 INTERCEPT = 464.0 SEC-1

O34 MTORR) H204 MTORR) H204 MR. D. UMITS) PMM MOLEC./CC) OBS. RATE(SEC-1) NET RATE(SEC-1) O34 MTORR) H204 MTORR) H2024 ARB. UMITS) PMM MOLEC./CC) OBS. RATE(SEC-1) NET RATE(SEC-1)

	0.30	80.00	<b>4</b> .50	0.000E-01	629.	36. 0	0**0	80.00	5.20	0.000E-01	894.	46. 8	
	0.30	80.00	8.10	0-0006-01	1238.	130. 0	0.40	80.00	10.80	0.000E-01	1393.	137. 0	
	030	B0.00	15.90	0.000E-01	1782.	57. 4	0**0	80.00	22.80	0.000E-01	2548.	413. 0	
	0.30	80.00	19.20	0.000E-01	1946.	-40. #	0.40	80.00	33.00	0.0006-01	3003.	120. 0	
	06.0	80.00	12.00	0.000E-01	1315.	-101. •	0.40	80.00	4.50	0.000E-01	742.	-52.	
	0.30	80.00	1.71	0.000E-01	520.	-82. 0	0.40	80.00	1.32	0.000E-01	577.	16. 4	
	0.30	80.00	0.22	2.263E+13	595.	111.	4.0	80.00	7.30	0.0006-01	960.	- 39. 4	
	030	80.00	0.54	6.5596413	735.	225.	4.8	80.00	9.90	0.000E-01	1196.	ę. ę	
	0.30	80.00	16.20	1.7226+14	2317.	569. #	4.00	80.00	63.00	0.000E-01	5076.	<b>•</b> •	
	0.30	90.00	15.90	1.541E+14	2200.	475. 8	4.00	80.00	35.00	0.000E-01	2996.	- 34 . 4	
	0.30	B0.00	0.20	2.607E+13	510.	27.	4.0	80.00	25.80	0.000E-01	2432.	77. 4	
	0.30	80.00	0.15	5.411E+13	622.	143.	4.00	80.00	1.35	1.668E+14	1267.	704.	
10	0.30	80.00	0.12	1.082E+14	<b>5</b> 33.	477.	4.00	80.00	13.80	2.139E+14	2070.	594.	
)3	0.30	80.00	0.40	5.739E+13	712.	198.	4.8	B0.00	1.80	1.1556+14	1084.	488.	
	0.30	80.00	3.60	1.6726414	1406.	654.	4.00	80.00	1.14	6.984E+13	849.	301.	
	0.30	80.00	2.10	1.3126+14	1091.	458.	4.00	80.00	0.75	5.132E+13	769.	250.	
	0F.0	80.00	0.90	7.0506413	179.	241.	4.00	80.00	0.60	3.6786413	759.	251.	
	0.30	27.00	2.50	7.378€+13	884.	221.	4.00	80.00	18.90	1.9256414	2538.	689. 1	
	0.30	27.00	7.20	1.2796414	1401.	364.	4.00	80.00	4.40	1.1416+14	1243.	456.	
	0.30	27.00	0.90	5.7396413	704.	166.	4.00	80.00	5.00	1.255E+14	1214.	383.	
	0.30	27.00	2.40	1.164E+14	1084.	427.	<b>*</b> .00	80.00	3.00	8.411E+13	1029.	345.	
	0.30	27.00	3.40	1.4266414	1216.	480.	4.00	80.00	1.80	4.990E+13	851.	255.	
	0.30	27.00	6.60	1.623E+14	1495.	506.	4.00	80.00	0.60	2.695E+13	. 169	183.	
							4.00	80.00	6.00	1.7116+14	1757.	853.	
							4.00	80.00	5.40	1.411E+14	1500.	640.	
# #ET #	ATE LESS	THAN 50 P	ERCENT OF H	4202 RATE			4.00	80.00	5.40	1.411E+14	1505.	645.	

v rume h2d2 rums for this date: 14-mov-80

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0 MET RATE LESS THAN 50 PERCENT OF H202 RATE 0 Pure H202 Runs For This Date: 17-Nov-80

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ON + PERMITRIC ACID INIA--Beries C

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ПЕМР = 298. КЕLVIM MELINIOIAL PRESSURE = 15.10 ПОRR MEZ SENSITIVIY = 7.220€+13 MOLECULES/CC/MV MEZZ/MA SENSITIVIY #AID = 4.44 APPROXIMATE MO2/H202 SENSITIVITY RATIO = 1.6143

NZOZ PNEMDNEMOLOGICAL RATE LEAST-SOUMRES; SLOPE = 78.5000 INTERCEFT = 427.0 SEC-1 DX MTORR) H2OK MTORR) H2O2(ARB, UMITS) PMA MOLEC./CC) OBS, RATE(SEC-1) MET RATE(SEC-1) OX MTORR) H2 (HTORR) H2O2(ARB, UMITS) PMA MOLEC./CC) OBS, RATE(SEC-1) MET RATE(SEC-1)

				!		
	0.00	0.000E-01	342.	-95.	0.30	88
	17.40	0,000E-01	1900.	107. 0	00	8
	10.80	0.000E-01	1336.	61. 8	0.30	8.
	61.00	0.000E-01	5187.	-29. 0	00	88
	40.00	0.000E-01	3558.	-9. 0	0.30	86.
	24.90	0.000E-01	2387.	• •	0.30	\$
	13.50	0.000E-01	1475.	-12. 0	0.30	8.
	4.80	0.000E-01	771.	-33. 0	0.30	ġ
	26.70	2.8365+14	3515.	\$ . 24	0.10	8
	5.60	2.248E+14	1580.	713.	00	Ó
_	2.10	1.7126+14	1133.	541.	0.30	ŏ
	. +1 • 1	1.195E+14	846.	330.	0.30	Ó
_	0.72	8.741E+13	. 149	207.		
	0.36	6.778E+13	613.	158.		
					A NET RATE LESS	Ĩ
					A PURE H202 RUN	SFO
	PERCENT OF 5 DATE:	H202 RATE			18-MOV-80 H2	DATA
					T Test data: no	

H202 PHENDHENOLOGICAL RATE LEAST-SOUARES: SLOPE = 78.5000 Intercept = 427.0 Sec-1

1.8507

APPROXIMATE M02/H202 SENSITIVITY RATIO =

UH + FERNITRIC ACID DATA<sup>†</sup> --Bertes C

TEMP = 298. KELVIN MELTUM TOTAL PRESSURE = 15.10 TORR MELTUMTYT = 9.080£43 MQLECULES/CC/NV MD2/PMM SENSITIVITY RATIO = 4.44

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8.	86.00	0.90	1.452E+14	745.	247.
.30	86.00	2.40	2.331E+14	1095.	480.
05.0	86.00	22.80	3.374E+14	2547.	330. 1
95.0	86.00	2.70	1.902E+14	972.	333.
0.30	86.00	1.65	1.432E+14	810.	253.
0.30	86.00	26.70	3.006E+1+	2725.	202.1
0.30	86.00	7.50	2.270E+14	1434.	418.
01.0	86.00	7.50	2,270E+14	1361.	345.
90	0.00	7.50	2.270E+14	1457.	441.
00	8.0	1.65	1.063E+14	721.	164.
8.0	0.0	1.10	7.567E+13	559.	46.

# NET RATE LESS THAN SO PERCENT OF H202 RATE # PURE H202 RUNS FOR THIS DATE: 18-MOV-80 H2 DATA t Test data; not used in evaluation of rate constant.

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ON + PERNITRIC ACID DATA--Bertes C

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TEMP = 267, KELVIM MELIUM TOTAL PRESSURE = 10.10 TOMR ND2 SEMSITIVITY = 0.040E413 MGLEGUES/CC/WW MD2/PMM SEMSITIVITY RAITO = 4.44 APPROXIMATE NO2/H202 SENSITIVITY RATIO = 1.2187

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NCO2 PHENDNENOLOGICAL RATE LEAST-SQUARES: SLOPE = 79.3000 IMTERCEPT = 448.0 SEC-1

DH + PERMITRIC ACID DATA--Beries C

TEMP = 324. KELVIN MELIUM TOTAL PRESSURE = 10.10 TORR MO2 SENSITIVITY = 6.290E+13 MOLECULES/CC/NV MO2/PMA SENSITIVITY RATIO = 4.44 APPROXIMATE W02/H202 SENSITIVITY RATIO = 1.4649

HZO2 PHENDMENDLDGICAL RATE LEAST-SGUARES; SLOPE = 68.7000 IMTERCEPT = 588.0 SEC-1 03(HIDRR) N2OKHTORR) N2O2(ARD, UNITS) PMM NOLEC./CC) ODS. RATE(SEC-1) NET RATE(SEC-1) 03(HIORR) H2O(HIORR) H2O2(ARD, UNITS) PMM NOLEC./CC) ODS. RATE(SEC-1) NET RATE(SEC-1)

0.30 0.30 0.30 0.30 0.30 0.30 0.30 0.30	24-MDV-80
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\$\$\$\$\$ <b>\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$</b> \$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$	

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ON + PERNITRIC ACID DATA--Beries C

TEMP = 244. KELVIM MELINI DTAL PRESUME = 10.10 TORR M22 SENSTIVITY = 4.950E+13 MQLECULES/CC/MV M22 PMM SENSTIVITY RATIO = 4.44

1.4148 APPROXIMATE NO2/H202 SENSITIVITY RATIO =

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NCO2 PHENOMENOLOGICAL RATE LEAST-SOUMRES: 9.0FE = 78.6000 INTERCEPT = 513.0 SEC-1

OH + PERNITRIC ACID DATA -- Series C

TEMP = 246.KELVIM HELIUM TOTAL PRESSUME = 15.10 TORR MOZ SEMSITIVITY = 5.740E413 MOLECULES/CC/MV MOZ/PMA SEMSITIVITY RATID = 4.44

1.4330 APPROXIMATE MO2/H202 SEMSITIVITY RATIO =

H2D2 PHEMOMEMOLDEICAL RATE LEAST-SQUARES: 51.0PE = 44.0900 IMTERCEPT = 532.0 SEC-1

034 MTORR) H2OK MTORR) H2O2(ARB, UNITS) PMA MOLEC./CC) ORS. RATE(SEC-1) MET RATE(SEC-1) OJIMITORR) M2OCMTORR) M2O2ARB, UMITS) PMM MOLEC./CC) OBS, RATE(SEC-1) MET RATE(SEC-1)

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135. 0 135. 0 135. 0 135. 0 135. 0 135. 0 136. 0 136. 0 136. 0 136. 0 136. 0 136. 0 136. 0 136. 0 136. 0 137. 0 147. 0 14	334. 374. 375. 375. 375. 375. 375.
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A STATEMENT

ON + PERNITRIC ACID DATA--Bertes D

TEMP = 245. RELVIN MELINIPOTAL PRESSURE = 15.10 TOAR DOS SEMILIVITY = 9.400E+12 MOLECULES/CC/MU MOZ/PMA SEMILIVITY RAID = 5.07

0.7797 APPROXIMATE W02/H202 SENSITIVITY RATIO =

NCD2 FHENDMENDLDETCAL RATE LEAST-SOUMRES: SLOPE = 19.7000 INTERCEPT = 303.0 SEC-1 303.0 SEC-1 D34 MTORK) H20K MTORR) H202( ARB. UMITS) PNM MOLEC./CC) ODS. RATE(SEC-1) NET RATE(SEC-1)

OH + PERNITRIC ACID DATA --- Series D

TEMP = 254, KELVIN MELIUM TOTAL PRESSURE = 15.10 TORR MO2 SEMSITIVITY = 2.020E413 MOLECULES/CC/NV MO2/PMA SENSITIVITY RATIO = 5.07

0.7340 APPROXIMATE W02/H202 SENSITIVITY RATIO =

H202 PHENDHENOLOGICAL RATE LEAST-SOUARES: SLOPE = 44.0300 INTERCEPT = 175.0 SEC-1

D3(MTOKR) M20KMTORR) N202KARB. UNITS) PNAKMOLEC./CC) OBS. RATE(SEC-1) MET RATE(SEC-1)

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# MET RATE LESS THAN 50 FERCENT OF H202 RATE 9 PURE H202 RUNS FOR THIS DATE: 22-DEC-80

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#### CAPTIONS

- 1 Schematic Diagram of Fast-Flow-Discharge-Tube Apparatus.
- 2 Schematic Diagram of Laser Flash Photolysis Resonance-Fluorescence Experimental Apparatus. For experiment Series B, a KrF excimer laser (248 nm) was used, and for experiment Series C and D, the quadrupled output (266 nm) of a Nd:YAG laser was used.
- 3 First-Order Plot of Log I<sub>f</sub> versus Time. Rate data giving  $k^{I} = 1155 \text{ s}^{-1}$  are for [PNA] = 1.06 x 10<sup>14</sup> molecules/cm<sup>2</sup> and [H<sub>2</sub>O<sub>2</sub>] = 1.05 arbitrary units. Data giving  $k^{I} = 2531 \text{ s}^{-1}$  are for [PNA] = 1.51 x 10<sup>14</sup> molecules/cm<sup>3</sup> and [H<sub>2</sub>O<sub>2</sub>] = 21.9 arbitrary units (too high for use in second-order analysis). Both sets of data are from experiment Series C, T = 324 K.
- 4 Second-Order Plot of  $k^{I}$  versus  $[H_2O_2]$  from Experiment Series C, 324 K (uncertainties shown are  $1\sigma$ ).
- 5 Second-Order Plot of  $k^{I}$  versus [NO<sub>2</sub>]. The effective bimolecular rate constant is  $k^{II} = (9.23 \pm 0.22) \times 10^{-13} \text{ cm}^3 \text{ s}^{-1}$  (uncertainties shown are 1 $\sigma$ ).

6 Second-Order Plot of  $k^{I}$  versus [PNA] for Experiment Series C, T = 298 K.

- 7 Second-Order Plot of  $k^{I}$  versus [PNA] for Experiment Series C, T = 246 K.
- 8 Arrhenius Plot of Log k versus 1000/T
  - $\Delta$  Experiment Series B
  - - Experiment Series C
  - o Experiment Series D

The solid error-bars are the 25 statistical uncertainties not including calibration errors; the dashed error-bars are the total 25 statistical uncertainties from all sources.



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# Appendix E

 $O_3$  + HOONO<sub>2</sub>  $\rightarrow$  PRODUCTS. AN UPPER LIMIT FOR THE RATE CONSTANT (Paula L. Trevor and John E. Davenport)

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#### A. Introduction

The importance of reactions of pernitric acid  $(HOONO_2)$  for understanding atmospheric chemistry has been recognized since its formation by reaction (1) was suggested [Simonaitis and Heicklen, 1976]:

$$HO_2 + NO_2 + M \neq HOONO_2 + M$$
 (1)

After  $HOONO_2$  was directly observed in the infrared [Niki et al., 1977], a concerted effort was initiated to investigate  $HOONO_2$  reactions [Howard, 1977; Graham, Winer, and Pitts, 1977 and 1978] and to determine the various loss mechanisms that affect its residence time in the stratosphere [Baldwin and Golden, 1978; Graham et al., 1978; Cox and Patrick, 1979; Molina and Molina, 1980]. Because ozone is one of the most important minor species in the stratosphere, and because of the role  $HOONO_2$  could play in decreasing the concentration of catalytic destroyers of ozone, it is of interest to examine potential reactions between ozone and  $HOONO_2$ . In this letter, we describe our determination of the upper limit to the rate constant for the reaction of  $HOONO_2$  with ozone and discuss the significance of this limit to our understanding of atmospheric chemistry.

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#### B. Experimental

A static system was used to investigate the reaction kinetics since a slow reaction rate was anticipated. The apparatus consisted of a 20-m variable pathlength Monel cell (Wilks) fitted with a sample inlet and a rotary pump for evacuation; an additional inlet was added to facilitate expansion of  $O_3$  into the cell. A constant temperature circulating bath was attached to cooling coils around the cell to maintain a temperature of  $5^{\circ}C$ , as measured by an iron-constantan thermocouple on the outer cell surface underneath the outer fiberglass insulating layer. A Perkin-Elmer 467 scanning IR spectrophotometer was used to record spectra of species in the cell.

Ozone was prepared by tesla discharge in  $O_2$  and was stored and purified on silica gel. The infrared absorption at  $100^{\circ}$  cm<sup>-1</sup> showed that the  $O_3$  was  $\geq 95\%$  pure ( $O_2$  was probably the major impurity). Its concentration in the cell was determined by expansion of a known pressure of the gas from a calibrated volume ( $V_{vol}/V_{cell} = 4.5 \times 10^{-3}$ ), assuming that the  $O_3$  was pure. It was also assumed that no significant change in the  $O_3$  concentration occurred during the experiments. The  $O_3$  concentration was always at least 10 times greater than the HOONO<sub>2</sub> concentration at the time of the  $O_3$  addition. Because of the small absorption coefficient [Herzberg, 1945], (506 l·mole<sup>-1</sup>·m<sup>-1</sup>) at 1000 cm<sup>-1</sup> for  $O_3$  and interference from NaNO<sub>3</sub> (NaCl windows were used for

several experiments) at ~ 1000 cm<sup>-1</sup>,  $O_3$  concentration measurements by IR were not used in the data analysis. However, the  $O_3$  concentration obtained by the IR absorption at 1000 cm<sup>-1</sup> agreed within  $\pm$  30% of the value calculated by expansion of a pure sample.

 $HOONO_2$  was prepared by the addition of 0.6 g (4.5 x 10<sup>-3</sup> mol) of solid NO<sub>2</sub>BF<sub>4</sub> (Alfa-Ventron) in approximately 5-mg portions with vigorous stirring 90%  $H_2O_2$  (FMC) at  $0^{\circ}C$ . This reaction mixture to 2 g (5.3 x  $10^{-2}$  mol) was pipeted into a 10-ml bulb connected as close as possible to the inlet of the IR cell. The mixture was allowed to warm to ambient room temperature. Argon (~ 50  $cm^3/min$ ) was then bubbled through the reaction mixture, and the effluent was passed into the previously evacuated 20-m cell until the desired HOONO<sub>2</sub> concentration was obtained. More Ar was then added from a separate inlet to raise the cell pressure to the desired value. An IR spectrum was obtained using the preparative method described above and a flow-through 10-cm pathlength cell; the spectrum is shown in Figure 1. This spectrum indicates a low level of impurity from  $HNO_3$  (840 cm<sup>-1</sup>) and  $NO_2$  (1620 cm<sup>-1</sup>). Because  $HOONO_2$  and  $HNO_3$  have strong absorptions at 1720 cm<sup>-1</sup>, the  $HOONO_2$  absorption at 803 cm<sup>-1</sup>,  $\varepsilon = (1.26 \times 10^5 \text{ } 1 \cdot \text{mole}^{-1} \cdot \text{m}^{-1})$  [Graham et al., 1977 and 1978] was monitored for the experimental measurements. It was observed that HNO<sub>3</sub> increased after HOONO, contacted the Monel cell walls.

Throughout the study, it was clear that even after passivation of the cell walls, the  $HOONO_2$  reacted heterogeneously. Not only was a green substance

 $(Ni(NO_3)_2 \cdot x H_2 O)$  deposited on the walls, but the mirror surfaces were attacked, apparently through disintegration of the epoxy that bonded the gold mirror coating to the glass substrate.

Because of the nonreproducible behavior of the heterogeneous reaction, we performed experiments with ozone according to the following sequence. As previously described, the HOONO<sub>2</sub> was swept into the cell and then buffered to a total pressure of ~ 13 torr with Ar ( $P_T^0$ ); the IR spectrum was quickly scanned from 1800 cm<sup>-1</sup> - 700 cm<sup>-1</sup>. If the HOONO<sub>2</sub> proved to be relatively pure, the spectrometer was set at 803 cm<sup>-1</sup> and the absorbance was recorded as a function of time with a strip-chart recorder. When sufficient time had elapsed to determine the HOONO<sub>2</sub> half-life in the cell ( $k_{Obs}^0$ ), the contents of a 25-ml bulb of O<sub>3</sub> gas ( $\geq 95\%$ ) attached to the inlet system were flushed into the cell with Ar to a total pressure of ~ 15 torr ( $P_T$ ). The HOONO<sub>2</sub> decay continued to be monitored until the new HOONO<sub>2</sub> half-life was established ( $k_{Obs}$ ) and no more significant change was observed.

#### C. Results and Discussion

Experiments with only  $HOONO_2$  and Ar were performed according to the procedure described in the preceding section; a typical result is shown in Figure 2. Under these conditions, we assumed the following reactions were taking place in the cell:

$$HOONO_2 + M \rightarrow HO_2 + NO_2 + M$$
 (2)

$$HO_2 + NO_2 + M \rightarrow HOONO_2 + M$$
 (3)

$$HOONO_2 \xrightarrow{\text{wall}} Loss$$
 (4)

This mechanism implies that the observed first-order decay rate constant of  $HOONO_2$ ,  $k_{Obs}^0$ , is a direct measure of  $k_4$ . Wall loss of  $HO_2$  was considered unimportant since experimental evidence indicated that the first-order loss rate of  $HOONO_2$  was not affected by the addition of up to 1 torr of  $NO_2$ . This is also consistent with the fact that with the estimated impurity level of  $NO_2$  ( $[NO_2] \cong 0.1$  [ $HOONO_2$ ]), reaction (3) occurs faster than diffusion to the walls ( $t_{diff} \cong 5$  sec). By analogy with other  $HO_2$  reactions [Lloyd, 1974], reactions of  $HO_2$  and  $HOONO_2$  were judged to be too slow for inclusion in the above scheme.

When ozone was added to the cell, according to the procedure given in the previous section, the first-order loss rate of  $HOONO_2$  was increased. A typical experimental run demonstrating this effect is shown in Figure 3; a summary of the experimental data is given in Table 1. Besides reactions (2), (3), and

Table 1

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HOONO<sub>2</sub> + O<sub>3</sub> → PRODUCTS; SUMMARY OF EXPERIMENTAL DATA AT 278 K (see text for details

	[HO <sub>2</sub> NO <sub>2</sub> ]			[0 <sub>3</sub> ]0			[NO <sub>2</sub> ]
Experiment	(molecule cm <sup>-3</sup> )	PT (torr)	k <sup>0</sup> obs (sec <sup>-1</sup> )	(molecule cm <sup>-3</sup> )	P <sub>T</sub> (torr)	k obs (sec <sup>-1</sup> )	(molecule cm <sup>-3</sup>
* =	$2.5 \times 10^{15}$	11.8	$3.40 \times 10^{-4}$	5.8 x 10 <sup>16</sup>	13.3	8.43 x 10 <sup>-4</sup>	$2.5 \times 10^{16}$
7	$2.0 \times 10^{15}$	13.1	5.78 x 10 <sup>-4</sup>	$4.9 \times 10^{16}$	11.1	$5.54 \times 10^{-4}$	$2.0 \times 10^{16}$
e	$1.8 \times 10^{15}$	13.6	$8.75 \times 10^{-4}$	$9.1 \times 10^{16}$	16.5	1.99 x 10 <sup>-3</sup>	1.8 x 10 <sup>16</sup>
4	$1.1 \times 10^{15}$	13.4	6.79 x 10 <sup>-4</sup>	9.8 x 10 <sup>16</sup>	16.5	6.08 x 10 <sup>-4</sup>	$1.1 \times 10^{15}$
ß	$2.4 \times 10^{15}$	12.8	$7.70 \times 10^{-4}$	$3.3 \times 10^{16}$	14.5	7.22 x 10 <sup>-4</sup>	2.4 x 10 <sup>16</sup>
* 0	1.9 x 10 <sup>15</sup>	13.8	3.73 x 10 <sup>-4</sup>	$1.6 \times 10^{16}$	14.8	5.50 x 10 <sup>-4</sup>	1.9 x 10 <sup>16</sup>
* ~	2.1 x 10 <sup>15</sup>	11.8	$4.81 \times 10^{-4}$	$1.3 \times 10^{17}$	16,1	2.89 x 10 <sup>-3</sup>	$2.1 \times 10^{16}$
80	$1.7 \times 10^{15}$	13.4	5.02 × 10 <sup>-4</sup>	8.3 x 10 <sup>16</sup>	16.5	$1.54 \times 10^{-3}$	$1.7 \times 10^{16}$
5	3.4 x 10 <sup>15</sup>	13.1	$1.16 \times 10^{-3}$	$6.6 \times 10^{16}$	15.5	$1.75 \times 10^{-3}$	3.4 x 10 <sup>16</sup>
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\* Denotes simulated experimental run. ł

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م در از به مراجع از به (4), we included the following reactions in the mechanism:

$$HO_2 + O_3 \rightarrow OH + 2O_2 \tag{5}$$

$$HOONO_2 + O_3 \stackrel{?}{\rightarrow} OH + 2O_2 + NO_2$$
(6)

$$OH + HOONO_2 \stackrel{?}{\to} H_2O + NO_2 + O_2$$
 (7)

$$OH + NO_2 + M \rightarrow HNO_3 + M$$
 (8)

$$OH + O_3 \rightarrow HO_2 + O_2 \tag{9}$$

Wall loss and disproportionation of OH were ignored in view of the fast homogeneous reactions involving OH and the large concentrations of both  $O_3$ and HOONO<sub>2</sub> in the cell. Although the products of reaction (7) have not been measured directly, no experimental evidence indicates otherwise [Trevor et al., 1981; Littlejohn and Johnston, 1980]. Products were not measured in this kinetic study; however, those listed for reaction (6) seemed most likely. Since  $k_{Obs}^0$  indicated that the wall loss of HOONO<sub>2</sub> ( $t_{wall} \ge 150$  sec) occurred more slowly than diffusion to the walls and thus was not diffusion controlled, we assumed that the pressure increase due to the introduction of ozone into the system had no measurable effect on  $k_4$ .

The three experimental runs that best represented the range of ozone concentrations were simulated by means of a computer program employing Gear's method [Gear, 1971] to integrate the set of differential equations corresponding to reactions (2) through (9). The input parameters consisted of the rate constants for the reactions in the kinetic scheme given above and all initial non-zero reactant concentrations. Literature values were used for  $k_2$ ,  $k_3$ ,  $k_5$ ,  $k_7$ ,  $k_8$ ,

and  $k_9$  (see Table 2);  $k_{0DS}^0$  was substituted for  $k_4$  for each of the simulation runs. The measured initial concentrations of ozone and HOONO ied are listed in Table 1. Because the NO<sub>2</sub> concentration was not directly monitored, three different concentration values were used for each set of experimental conditions, while all other input parameters remained the same. The three NO<sub>2</sub> concentrations chosen were: (1) the most probable value; (2) zero; and, (3) the maximum possible value. The most probable NO<sub>2</sub> concentration was determined by taking the difference between the HOONO<sub>2</sub> initially present and the HOONO<sub>2</sub> remaining after  $k_{ObS}^0$  had been determined, just prior to the addition of ozone. The zero concentration was chosen simply to bracket the extreme cases. The maximum possible NO<sub>2</sub> concentration was chosen by taking ten times the total HOONO<sub>2</sub> concentration. At these high values, the computer simulations indicated deviations from first-order decay rates at low HOONO<sub>2</sub> which were not observed in any of the experimental data. Thus, these values do indeed represent the limiting cases for the initial NO<sub>2</sub> concentration.

The rate constant of interest,  $k_6$ , was allowed to assume values of zero,  $1 \times 10^{-20} \text{ cm}^3 \text{ s}^{-1}$ ,  $2 \times 10^{-20} \text{ cm}^3 \text{ s}^{-1}$ , and  $5 \times 10^{-20} \text{ cm}^3 \text{ s}^{-1}$ . During the actual simulation, the HOONO<sub>2</sub> concentration was monitored as a function of time and a first-order decay rate constant was calculated for all of the situations previously discussed. These results are listed in Table 3. For the three experiments studied, it was observed that if  $k_6$  was as large as  $2 \times 10^{-20} \text{ cm}^3 \text{ s}^{-1}$ ,  $k_{calc}$ was larger than  $k_{obs}$  for all cases investigated. In order to allow for random

Reaction	Reference
(2) HOONO <sub>2</sub> $\rightarrow$ HO <sub>2</sub> + NO <sub>2</sub> k <sub>2</sub> = 4.40 $\times 10^{-4}$ s <sup>-1</sup>	Graham, et al., 1977 Graham, et al., 1978
(3) $HO_2 + NO_2 + M \rightarrow HOONO_2 + M$ $k_3 = 3.07 \times 10^{-31} \text{ cm}^6 \text{ molecule}^{-2} \text{ s}^{-1}$	Graham, et al., 1977 Graham, et al., 1978 Baldwin and Golden, 1978
(5) $HO_2 + O_3 \rightarrow OH + 2O_2$ $k_5 = 1.74 \times 10^{-15} \text{ cm}^3 \text{ s}^{-1}$	Zahniser and Howard, 1980
(7) OH + HOONO <sub>2</sub> $\rightarrow$ H <sub>2</sub> O + NO <sub>2</sub> + O <sub>2</sub> k <sub>7</sub> = 4.0 x 10 <sup>-12</sup> cm <sup>3</sup> s <sup>-1</sup>	Barker et al., 1980 Trevor et al., 1981
(8) OH + NO <sub>2</sub> + M $\rightarrow$ HNO <sub>3</sub> + M k <sub>8</sub> = 3.24 x 10 <sup>-30</sup> cm <sup>6</sup> molecule <sup>-2</sup> s <sup>-1</sup>	Atkinson, et al., 1976 Anastasi and Smith, 1976 Wine, et al., 1979
(9) $OH + O_3 \rightarrow HO_2 + O_2$ $k_g = 5.44 \times 10^{-14} \text{ cm}^3 \text{ s}^{-1}$	Anderson and Kaufman, 1973 Ravishankara, et al., 1979

Table 2REFERENCES FOR REACTION SCHEME

## Table 3

SUMMARY OF GEAR ROUTINE RESULTS  $(k_{calc}^{\dagger})$  $k_{calc}$  versus NO<sub>2</sub>,  $k_6$  (see text for details)

Experiment 1  $k_{obs} = 8.43 \times 10^{-4} s^{-1}$ 

**.	[ NO		
к <sub>6</sub>	0	$7.6 \times 10^{14}$	2.5 x 10 <sup>16</sup>
0	7.31 x 10 <sup>-4</sup>	$5.82 \times 10^{-4}$	$3.51 \times 10^{-4}$
$1.0 \times 10^{-20}$	$1.28 \times 10^{-3}$	1.23 x 10 <sup>-3</sup>	9.41 x $10^{-4}$
$2.0 \times 10^{-20}$	$1.74 \times 10^{-3}$	1.75 x 10 <sup>-3</sup>	$1.49 \times 10^{-3}$
5.0 x $10^{-20}$	$3.44 \times 10^{-3}$	$3.79 \times 10^{-3}$	$3.12 \times 10^{-3}$

Experiment 6  $k_{obs} = 5.50 \times 10^{-4} \text{ s}^{-1}$ 

<sup>لا</sup> **	0	8.0 x $10^{14}$	1.9 x 10 <sup>16</sup>
0	$6.28 \times 10^{-4}$	4.83 x $10^{-4}$	$3.78 \times 10^{-4}$
$1.0 \times 10^{-20}$	8.09 x 10 <sup>-4</sup>	$6.94 \times 10^{-4}$	$5.52 \times 10^{-4}$
$2.0 \times 10^{-20}$	9.88 x 10 <sup>-4</sup>	$8.88 \times 10^{-4}$	$7.18 \times 10^{-4}$
5.0 x $10^{-20}$	$1.46 \times 10^{-3}$	$1.39 \times 10^{-3}$	1.19 x 10 <sup>-3</sup>

Experiment 7  $k_{obs} = 2.89 \times 10^{-3} s^{-1}$ 

** k <sub>6</sub>	0	$1.4 \times 10^{15}$	$2.1 \times 10^{16}$
0	9.06 x 10 <sup>-4</sup>	$6.92 \times 10^{-4}$	$5.10 \times 10^{-4}$
$1.0 \times 10^{-20}$	$2.14 \times 10^{-3}$	$1.74 \times 10^{-3}$	$1.70 \times 10^{-3}$
$2.0 \times 10^{-20}$	$2.98 \times 10^{-3}$	3.03 x 10 <sup>-3</sup>	$2.94 \times 10^{-3}$
5.0 x $10^{-20}$	5.95 x 10 <sup>-3</sup>	5.75 x $10^{-3}$	$6.17 \times 10^{-3}$

\* Units are molecules cm<sup>-3</sup> \*\* Units are cm<sup>3</sup> s<sup>-1</sup> ‡ Units are s<sup>-1</sup>. experimental fluctuations and the influence of systematic uncertainties, we estimated that 5 x  $10^{-20}$  cm<sup>3</sup> s<sup>-1</sup> is a conservative upper limit for the rate constant k<sub>6</sub> for the reaction of ozone with HOONO<sub>2</sub>.

### D. Conclusions

The potential importance of  $HOONO_2$  is due to its ability to act as a reservoir for both  $HO_2$  and  $NO_2$ . This unique role underlines its potential importance and makes it difficult to estimate its effects without detailed modeling calculations. Although  $HOONO_2$  can be shown not to influence tropospheric photochemical smog [Baldwin et al., 1977], it potentially has a large effect in the stratosphere because the lower temperatures and pressures combine to give it a much longer lifetime with respect to unimolecular decomposition. Table 4 briefly summarizes several of the effects of various processes on the first-order loss rate of  $HOONO_2$  in the atmosphere. The photolysis rates  $(J_{hv})$  are based on those for  $H_2O_2$ , multiplied by 1.5 to approximate the absorption cross sections of  $HOONO_2$  recently measured by Molina [1980]. If the correct cross sections are the larger ones measured by Graham et al. [1978], the photolysis rate of  $H_2O_2$  should be multiplied by ~ 6.

From inspection of Table 4, it is clear from the derived upper limit to the rate constant for reaction of ozone with  $HOONO_2$  that this reaction should have no direct effect of atmospheric importance.

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## Table 4

ATMOSPHERIC HOONO<sub>2</sub> FIRST-ORDER LOSS RATES ( $s^{-1}$ )

Altitude (km)	Thermal <sup>a</sup>	J <sub>hv</sub> b	<sub>k<sub>6</sub>(0<sub>3</sub>) с</sub>	Total	Lifetime (hr)
0	2.8(-2)	9.3(~6)	<u>≤</u> 5 (8)	2.8(-2)	0.01
20	2.6(-8)	5.2(~6)	<b>≤</b> 3 (-7)	≤5.5(-6)	51
30	2.4(-7)	1.1(~5)	<b>≤</b> 2 (−7)	1.1(-5)	2,5
40	1.3(-5)	5.2(~5)	≤ 4.5(-8)	6.5(-5)	4.3
L	L	L	l	L	

<sup>a</sup>[Graham et al., 1977 and 1978; Baldwin and Golden, 1978].

<sup>b</sup>Based on  $J(H_2O_2) \propto 1.5 [R. D. Hudson, 1977].$ 

 ${}^{c}O_{3}$  concentration from [Luther et al., 1979, as quoted from U.S. Standard Atmosphere, 1976], rate constant limit from present work.

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#### E. References (Appendix E)

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FIGURE 1 INFRARED SPECTRUM OF HOONO2 PREPARED AS DESCRIBED IN TEXT

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FIGURE 2 SEMI-LOGARITHMIC PLOT OF HOONO<sub>2</sub> DECAY Typical result for Argon buffer gas.



