

· · REPORT DOCUMENTATION	PAGE	READ INSTRUCTIONS
SCASH PB PO DECA	2. GOVT ACCESSION NO.	3. RECIPIENT'S CATALOG NUMBER
Arusk-1k- 82-0600	AD-A118578	
TITLE (and Sublishe)		S. TYPE OF REPORT & PERIOD COVE
		Final ••
LABORATORY STUDIES OF SOLVATED GAS	-PHASE AN IONS	6. PERFORMING ORG. REPORT NUMB
. AUTHOR(e)		8. CONTRACT OR GRANT NUMBER(+)
Pater M. Hierl		AFOSR-81-0105
DEDERMING CHEANIZATION NAME AND ADDRESS		10. PROGRAM ELEMENT, PROJECT, TA
The University of Kansas		61102F
Lawrence, Kansas 66045		2303/B1
. CONTROLLING OFFICE NAME AND ADDRESS		12. REPORT DATE
Dept. of the Air Force		Feb 198
Air Force Office of Scientific Res	earch/NC	13. NUMBER OF PÅGES
BOILING AIT FOTCE BABE, D.C. 2033 4. MONITORING AGENCY NAME & ADDRESS(11 dillorad	from Controlling Office)	15. SECURITY CLASS. (of this report)
		linglassifisi
		UNCLESSIFICATION/DOWNGRADI
		SCHEDULE
Approved for public release; distr: 7. DISTRIBUTION STATEMENT (of the abstract enfored in	ibution unlimite	d n Report)
Approved for public release; distr: 7. DISTRIBUTION STATEMENT (of the abeliact entered in	ibution unlimite	d n Report)
Approved for public release; distr: DISTRIBUTION STATEMENT (of the abstract entered in B. SUPPLEMENTARY NOTES	ibution unlimite	d n Report)
Approved for public release; distr: 7. DISTRIBUTION STATEMENT (of the abeliact entered in 8. SUPPLEMENTARY NOTES	ibution unlimite	d <b>Report)</b> <b>E</b> (100 0 1000
Approved for public release; distr: DISTRIBUTION STATEMENT (of the abstract entered in B. SUPPLEMENTARY NOTES	ibution unlimite	d Report) NUS 2 3 1982
Approved for public release; distr: 7. DISTRIBUTION STATEMENT (of the abeliact entered in 8. SUPPLEMENTARY NOTES	ibution unlimite	d n Report) NUS 2 3 1982
Approved for public release; distr: DISTRIBUTION STATEMENT (of the obstract entered in B. SUPPLEMENTARY NOTES D. KEY WORDS (Continue on reverse side if necessary and	ibution unlimite n Block 20, 11 different free (Identify by block number)	d Report) AUG 2 3 1982 E
Approved for public release; distr: Approved for public release; distr: DISTRIBUTION STATEMENT (of the obstract entered in B. SUPPLEMENTARY NOTES C. KEY WORDS (Continue on reverse side if necessary and Atmospheric chemistry Ion-Molecule Reactions Cluster Ions	ibution unlimite n Block 20, 11 different free (Identify by block number)	d Report) AUS 2 3 1982 E
Approved for public release; distr: Approved for public release; distr: DISTRIBUTION STATEMENT (of the obstract entered in B. SUPPLEMENTARY NOTES AUPPLEMENTARY NOTES AMOSPHERIC Continue on reverse side if necessary and Atmospheric chemistry Ion-Molecule Reactions Cluster Ions ABSTRACT (Continue on reverse side if necessary and Cross sections are reporte OHI (H2O) with CO2 and SO2 ov energy 0515-25.0 eV (LAB). R collisional dissociation. and	ibution unlimite a Block 20, 11 dilierent free identify by block number) identify by block number) d for the reacti er the range of eactions observe charge transfer	d AUG 2 3 1982 E E tons of the solvated anion reactant translational ed include solvant switching These reactions are
Approved for public release; distr: Approved for public release; distr: DISTRIBUTION STATEMENT (of the abstract entered in B. SUPPLEMENTARY NOTES C. KEY WORDS (Continue on reverse side if necessary and Atmospheric chemistry Ion-Molecule Reactions Cluster Ions ADSTRACT (Continue on reverse side if necessary and Cross sections are reporte OHT (H2O) with CO2 and SO2 ov energy 0.15-25.0 eV (LAB). R collisional dissociation, and particularly important to the below 80 km, where considerab observed.	ibution unlimite a Block 20, If different free (identify by block number) d for the reaction eactions observed charge transfer ion chemistry of le hydration of	d AUS 201982 E E E E E E E E E E E E E
Approved for public release; distr: Approved for public release; distr: DISTRIBUTION STATEMENT (of the abstract entered in B. SUPPLEMENTARY NOTES AUDITARY NOTES AUDITARY WORDS (Continue on reverse side if necessary and Atmospheric chemistry Ion-Molecule Reactions Cluster Ions ABSTRACT (Continue on reverse side if necessary and Cross sections are reporte OHI (HgO) with CO2 and SO2 ov energy 0915-25.0 eV (LAB). R collisional dissociation, and particularly important to the below 80 km, where considerab observed. D 1 JAN 75 1473 EDITION OF 1 NOV 65 IS OBSOLE	ibution unlimite n Block 20, If different breact I dentify by block number) d for the reacti er the range of eactions observe charge transfer ion chemistry c le hydration of ETE	d AUS 2 3 1982 E tons of the solvated anion reactant translational of include solvant switchi These reactions are of the earth's atmosphere negative ions has been Unclassified

. i

. 4 Unclassified 12 JUL 1882 SECURITY CLASSIFICATION OF THIS PAGE (When Date Entered) READ INSTRUCTIONS REPORT DOCUMENTATION PAGE BEFORE COMPLETING FORM 2. GOVT ACCESSION NO. 3. RECIPIENT'S CATALOG NUMBER AFOSK-TR. 82-0660 S. TYPEOF REPORT & PERIOD COVERED 4. TITLE (and Subling) Final . . LABORATORY STUDIES OF SOLVATED GAS-PHASE ANIONS 6. PERFORMING ORG. REPORT NUMBER CONTRACT OR GRANT NUMBER(+) 7. AUTHOR(#) ŝ. Peter M. Hierl AFOSR-81-0105 PROGRAM ELEMENT, PROJECT, TASK AREA & WORK UNIT NUMBERS DEPT OF CHEMISLY The University of Kadeas 61102F Lawrence, Kansas 66045 2303/B1 12. REPORT DATE 11. CONTROLLING OFFICE NAME AND ADDRESS 1982 Dept. of the Air Force 13. NUMBER OF PAGE Air Force Office of Scient fic Research/NC 8 Bolling Air Force Base, D. 0. 20332 14. MONITORING AGENCY NAME & ADDRESS (V dillerent from Controlling Office) 15. SECURITY CLASS. (of this report) Unclassified 15. DECLASSIFICATION/DOWNGRADING SCHEDULE 16. DISTRIBUTION STATEMENT (of this Report) Approved for public release; distribution unlimited 17. DISTRIBUTION STATEMENT (of the abstract entered in Block 20, If different from Report) 18. SUPPLEMENTARY NOTES 19. KEY WORDS (Continue on reverse side if necessary and identify by block number) Atmospheric chemistry Ion-Moleculy Reactions Cluster Ions 20. ABSTRACT (Continue on reverse side if necessary and identity by block number) Cross sections are reported for the reactions of the solvated anions OH- $(H_2O)$  with CO<sub>2</sub> and SO<sub>2</sub> over the range of reactant translational energy 0.15-25.0 eV (LAB). Reactions observed include solvent switching, collisional dissociation, and charge transfer. These reactions are particularly important to the ion chemistry of the earth's atmosphere below 80 km, where considerable hydration of negative ions has been observed. DD 1 JAN 73 1473 EDITION OF 1 NOV 65 15 OBSOLETE Unclassified SECURITY CLASSIFICATION OF THIS PAGE (When Date Entered) s . when we get a ser

.1

# AFOSR-TR- 82-0660

JUL 1002

# FINAL SCIENTIFIC REPORT

TITLE: Laboratory Studies of Solvated Gas-Phase Anions

PRINCIPAL INVESTIGATOR: Professor Peter M. Hierl Department of Chemistry University of Kansas Lawrence, Kansas 66045

INCLUSIVE DATES: 1 March 1981 - 28 February 1982

GRANT NUMBER: AFOSR-81-0105

COSTS AND FY SOURCE: \$9,944, FY 82 (MINI GRANT)





topressed for public releases distribution unlimited.

and a second of the second

### A. RESEARCH OBJECTIVES

The objective of this research project was the study of the gas-phase reactions of solvated anions such as  $OH \cdot nH_2O$  (where n ranges from 0 to 6) with neutral molecules of atmospheric interest such as  $CO_2$  or  $SO_2$ . These reactions, which include solvent switching, electron transfer, proton transfer, isotope exchange, and collision-induced dissociation, are important in predicting the composition of the Earth's atmosphere subsequent to natural and artificial perturbations such as aurora, nuclear detonation, and missle reentry. Improved knowledge of this subject is necessary for better detection and tracking of spacecraft and post boost vehicles.

More specifically, this project had the following objectives:

- 1 -

(1) The processing of kinetic data collected at the Air Force Geophysics Laboratory, where the Principal Investigator spent ten weeks during the summer of 1980 as an AFOSR Summer Faculty Research Program participant.

(?) The analysis and interpretation of the processed kinetic data. For the exothermic solvent switching reactions, this involves the comparison of the measured reaction rates with calculated rates derived from current theories of ion-molecule reaction and, when possible, with rates measured in other laboratories. For the endothermic collisional dissociation processes, this involves the analysis of the energy-dependence of the reaction cross section near threshold to obtain information on the bond strengths and the structures of the ions undergoing dissociation.

(3) The development of new ion sources capable of producing intense, monoenergetic beams of cluster ions of the type  $OH \cdot nH_2O$ , where n extends up to 5 or 6, and of mixed clusters such as  $HCO_3 \cdot nH_2O$ .

# B. STATUS OF THE RESEARCH

#### Processing of kinetic data

During the ten week period the Principal Investigator spent at the Air Force Geophysics Laboratory, kinetic data were collected for the following reactions: AIR FORCE OFFICE OF SCIENTIFIC RESEARCY (ATSO)

NOTICE OF TRANSFILTTAL TO DTIC This technic for out has been reviewed and is approved to mail to solve the AFR 190-12. Distribution to walimited. MATTHEM J. KENDER Chiei, Technical Information Division

OH +  $SO_2 \rightarrow SO_2$  + OH(1) $OH(H_2O) + Ar \rightarrow OH + H_2O + Ar$ (2) $+ SO_{2} \rightarrow OH^{-} + H_{2}O + SO_{2}$ (3a) $\rightarrow HSO_3 + H_2O$ (3b)  $OH^{-}(H_{2}O)_{2} + SO_{2} - OH^{-}(H_{2}O) + H_{2}O + SO_{2}$ (4a) $--HSO_3 + 2H_2O$ (4b)  $OH^{-}(H_{2}O)_{3} + SO_{2} \rightarrow HSO_{3}^{-} + 3H_{2}O_{3}^{-}$ (5a) $\rightarrow$  HSO<sub>3</sub> (H<sub>2</sub>O) + 2H<sub>2</sub>O (5b)  $OH^{-}(H_{2}O) + CO_{2} - HCO_{3} + H_{2}O$ (6)  $OH^{-}(H_{2}O)_{2} + CO_{2} \rightarrow HCO_{3}^{-} + 2H_{2}O$ (7a)  $- HCO_3 (H_2O) + H_2O$ (7b)  $OH^{-}(H_{2}O)_{3} + CO_{2} \rightarrow HCO_{3}^{-} + 3H_{2}O_{3}$ (8a) $\rightarrow HCO_{3}(H_{2}O) + 2H_{2}O$ (85) $\rightarrow \text{HCO}_3$  (H<sub>2</sub>O)<sub>2</sub> + H<sub>2</sub>O (8c)  $OH + CH_3Br \rightarrow CH_2Br + H_2O$ (9a)  $\rightarrow$  Br + CH<sub>3</sub>OH (9Ъ)  $OH^{-}(H_{2}O) + CH_{3}Br \rightarrow CH_{2}Br^{-} + 2H_{2}O$ (1.0a) $\rightarrow$  Br + CH<sub>3</sub>OH + H<sub>2</sub>O (10b)  $\rightarrow$  Br<sup>-</sup>(H<sub>2</sub>O) + CH<sub>3</sub>OH (10c) $\rightarrow$  OH<sup>-</sup> + H<sub>2</sub>O + CH<sub>3</sub>OH (104)  $OH^{-} + CH_3C1 \rightarrow CH_2C1^{-} + H_2O$ (11a)  $\rightarrow$  C1 + CH<sub>3</sub>OH (11b)  $OH^{-}(H_{2}O) + CH_{3}C1 \rightarrow CH_{2}C1^{-} + 2H_{2}O$ (12a) $\rightarrow$  C1 + CH<sub>2</sub>OH + H<sub>2</sub>O (12b)  $\rightarrow C1^{-}(H_{2}O) + CH_{3}OH$ (12c)  $\rightarrow$  OH + H<sub>2</sub>O + CH<sub>3</sub>C1 (12d)  $OH^{-}(H_{2}O)_{2} + CH_{3}C1 \rightarrow CH_{2}C1^{-} + 3H_{2}O$ (13a)→ C1 + CH<sub>3</sub>OH + 2H<sub>2</sub>O (13b) -1 C1 (H<sub>2</sub>O) + CH<sub>3</sub>OH + H<sub>2</sub>O (13c)  $\rightarrow$  OH (H<sub>2</sub>O) + H<sub>2</sub>O + CH<sub>3</sub>C1 (1 ....  $\rightarrow$  OH + 2H<sub>2</sub>O + CH<sub>3</sub>C1 (13e)

Typically, data for a particular reaction were obtained for 2 to 5 replicate runs in which the reactant and product ion intensities  $[I^{\circ}(A^{-})]$  and  $I(C^{-})$ , respectively] were measured at each of 30 to 50 selected col-

- FE WARDER

lision energies covering the range 0.15 to 25 eV (LAB).

Data processing, which was completed during the period covered by this report, consisted of the following steps:

(a) Correcting the intensities  $I^{\circ}(A^{-})$  and  $I(C^{-})$  measured in a given run for any possible differences in the transmission efficiency of the quadrupole mass spectrometer employed to detect the ions  $A^{-}$  and  $C^{-}$ .

(b) Correcting the observed intensities  $I^{\circ}(A^{\overline{}})$  for those ions arriving at the detector during the "dead time" of the pulse counting system.

(c) In the studies of collisional dissociation, correcting the observed rate for the effect of collisions with molecules of background gas.

(d) Averaging the results of the replicate runs for each particular reaction.

(e) Plotting the averaged cross sections for each reaction versus the relative translational energy of the reactants to produce plots of what are known as excitation functions for each reaction.

# (2) Data Analysis and Interpretation

(a) The dependence of the reaction cross section  $\sigma$  on the reactants' relative translational energy  $E_T$  was determined for the various exothermic reactions by plotting log  $\sigma$  <u>vs</u>. log  $E_T$ . Such plots usually showed a linear dependence at low collision energies ( $E_T$  less than 1 eV), a curved transition region at intermediate energies (1-2 eV), and a second linear region at higher energies. Thus, in the limit of low or high energy, the dependence of the reaction cross section on collision energy can be expressed as  $\sigma(E_T) \ll E_T^{-s}$ , where the parameter s, determined from the slopes of the linear portions of these log-log plots, usually has a value of about 0.5 at low energy and a value of 5-7 at high energy.

(b) The cross sections measured at a particular collision energy  $E_T$  were converted to monoenergetic rate coefficients  $k(E_T)$  by the relation  $k(E_T) = v_T \sigma(E_T)$ , where  $v_T$  is the relative velocity between the reactants corresponding to a relative translational energy  $E_T$ . Quasi thermal energy rate coefficients were then obtained by extrapolating the derived monoenergetic rate coefficients to thermal energy. The exothermic solvent switching reactions between  $OH^{-}(H_{2}O)_{n}$  and  $CO_{2}$  and  $SO_{2}$  were found to be very rapid; the derived thermal energy rate coefficients were on the order of  $1-3\times10^{-9}$  cm<sup>3</sup>/molecule-sec, suggesting that reaction occurs on every or nearly every collision between the reactants. Increasing the degree of solvation of the reactant  $OH^{-}(H_{2}O)_{n}$  ion was found to have no effect on the rate of the reaction with  $SO_{2}$ , and only slightly decreased the rate of the reaction with  $CO_{2}$ .

Theoretical rate coefficients for these solvent switching reactions were calculated from several current models for ion-molecule reactions. The AADO theory of Su, Su and Bowers [T. Su, E.C.F. Su, and M.T. Bowers, J. Chem. Phys. <u>69</u>, 2245 (1978)] predicted rate coefficients which were within a factor of two of those measured in the present study. Moreover, this model successfully predicted that SO<sub>2</sub> would react more rapidly with OH<sup>-</sup>(H<sub>2</sub>O)<sub>n</sub> than does CO<sub>2</sub> because the permanent dipole moment of SO<sub>2</sub> enhances the long-range attractive forces and thus the frequency of collisions (CO<sub>2</sub> is non-polar).

One interesting observation made in this study was the fact that the major products of the solvent switching reactions are the unsolvated ion  $HCO_3^-$  and  $HSO_3^-$ , even when the reactant ion was  $OH^-(H_2O)_3$ . This failure of the water molecules to solvate the ionic product (thus forming energetically more favorable products) we attribute to the high exothermicity of these reactions. Because the  $OH^--SO_2$  and  $OH^--CO_2$  bond energies are much greater than the  $OH^--H_2O$  bond energy, sufficient energy is released in these switching reactions to cause the decomposition of any solvated ionic products via "solvent boil-off".

In analyzing the data on the collisonal dissociation processes, we had to take into account the thermal motion of the target molecules because such motion tends to smear out the onset of dissociation at collision energies above threshold due to the Doppler effect. This was done by assuming a functional form for the true microscopic reaction cross section, convoluting this assumed function with the experimental conditions, and then comparing the calculated results with the phenomenological cross sections.

We assumed that the true cross sections  $\sigma(E_T)$  for these endothermic processes increase linearly with relative translational energy  $E_T$  above the

• 4 -

threshold value Eo according to the equation

$$\sigma(E_{T}) = \begin{cases} 0 & \text{if } E_{T} \leq E_{O} \\ Q(E_{T} - E_{O}) & \text{if } E_{T} \geq E_{O} \end{cases}$$

- 5 -

Assuming that a monoenergetic ion beam interacts with target gas molecules having an isotropic Maxwellian velocity distribution, we have convoluted the assumed cross section with the distribution in relative velocities according to the procedure described by Chantry [P.J. Chantry, J. Chem. Phys. <u>55</u>, 2746 (1971), Eq. (30)]. The results were then compared with the phenomenological cross sections, and the parameters Q and  $E_0$  were varied until satisfactory agreement was achived.

Agreement between the calculated excitation function and the measured cross sections was generally very good, indicating (1) a linear excitation function above threshold is adequate to describe the energy dependence of the cross section for the collision-induced processes, and (2) our reactant ions possessed little if any internal excitation which, if present, would have caused more tailing than observed in the measured cross sections at energies below threshold.

The threshold energies correspond to the dissociation energy of the bond being broken in the dissociation process. Values of  $1.12 \pm 0.10$ eV and  $0.80 \pm 0.10$ eV were obtained for the dissociation of OH -H<sub>2</sub>O and OH (H<sub>2</sub>O)-H<sub>2</sub>O, respectively, in very good agreement with values obtained in other laboratories employing different methods. Moreover, we obtained a value of  $3.8 \pm 0.25$  eV for the OH -CO<sub>2</sub> bond dissociation. Since such a high value indicates appreciable chemical bonding (as opposed to simple electrostatic attraction), we have chosen to write this ion as HCO<sub>3</sub> rather than as OH (CO<sub>2</sub>). Our observation that SO<sub>2</sub> rapidly reacts with HCO<sub>3</sub> to produce HSO<sub>3</sub> indicates that the reaction is exothermic and therefore that the bond dissociation energy of OH -SO<sub>2</sub> is greater than 3.8 eV.

Manuscripts describing these results in more detail have been prepared for publication, and copies will forewarded to the AFOSR in the near future.

#### (3) Ion Source Design

Attempts were initiated to design and contruct ion sources capable of

A Street Contraction

producing intense, monoenergetic beams of cluster ions such as  $OH^-(H_2O)_n$ , where n extends up to 5 or 6. Unfortunately, only preliminary results have been obtained during the period of this project. These preliminary results, however, have served to identify the problems that must be overcome in the design of such an ion source.

The kinetics of the three-body association reactions leading to cluster formation are sufficiently well known that appropriate source conditions can be calculated. The source must operate at gas pressures of up to 1 Torr, and the ions should have a residence time of at least several microseconds in the source. (Although it is possible to compensate for lower pressures with longer residence times, this trade-off is unfavorable, since a ten-fold decrease in pressure would require a hundred-fold increase in residence time.)

By using field-penetration from an extraction plate in front of the ion source rather than a repeller plate within the source, we have been able to form thermal-energy ions whose residence time in the source exceeds seve ral microseconds.

Operating an ion source at pressures of 1 torr, however, creates at least three problems. (1) Because of the great pressure differential between the source and the surrounding vacuum system, the source must be very "tight" and high speed vacuum pumps must be available. However, such tight sources are commonly used in chemical ionization mass spectrometry, so the problem here is one of optimizing the design of such ion sources. (2) Because the ionizing  $\phi_{i}$ electron beam is rapidly attentuated at high gas pressures, adequate penetration of the beam from an external filament into the source can present a problem. We are currently investigating the possibility of using a very high energy electron beam which, upon collisions with the source wall or the gas within the source, will eject secondary electrons that actually cause the desired ionization. (3) The large voltage difference between the ion source and the gas inlet system produces a discharge through the gas inlet line when the gas pressure in the line approaches 1 Torr. Again, there are known to be several solutions to this problem, and we are currently investigating them.

S. Contactor and

- 6 -

# C. PUBLICATIONS

- 1. John F. Paulson and Peter M. Hierl, "Translational Energy Dependence of Cross Sections for Reactions of  $OH^{-}(H_2O)_n$  with  $CO_2$  and  $SO_2$ ," manuscipt in final stages of preparation for submission to The Journal of Chemical Physics.
- 2. John F. Paulson, Michael J. Henchman, and Peter M. Hierl, "Effect of Reactant Ion Solvation on Gas-Phase  $S_N^2$  Reactions," manuscript in final stages of preparation for submission to The Journal of Chemical Physics.

# D. PROFESSIONAL PERSONNEL

Calvin Cole, Ph.D. Candidate, Department of Chemistry, University of Kansas, Lawrence, KS 66045

المراجع في المعالم المعالم الم

#### E. INTERACTIONS

1. Oral Presentations

P.M. Hierl, M.J. Henchman, and J: 1 Paulson, "Tandem Mass Spectrometric Studies of the Reactions of Solvated Anions," presented at the 29th Annual Confernce on Mass Spectrometry and Allied Topics, Minneapolis, Minnesota, May 24-29, 1981.

2. Consulting - none

#### F. NEW DISCOVERIES, INVENTIONS

none

