# Optical Spectra of Nonmetallic Inorganic Transient Species in Aqueous Solution

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### Foreword

The National Standard Reference Data System provides access to the quantitative data of physical science, critically evaluated and compiled for convenience and readily accessible through a variety of distribution channels. The System was established in 1963 by action of the President's Office of Science and Technology and the Federal Council for Science and Technology, and responsibility to administer it was assigned to the National Bureau of Standards.

NSRDS receives advice and planning assistance from a Review Committee of the National Research Council of the National Academy of Sciences-National Academy of Engineering. A number of Advisory Panels, each concerned with a single technical area, meet regularly to examine major portions of the program, assign relative priorities, and identify specific key problems in need of further attention. For selected specific topics, the Advisory Panels sponsor subpanels which make detailed studies of users' needs, the present state of knowledge, and existing data resources as a basis for recommending one or more data compilation activities. This assembly of advisory services contributes greatly to the guidance of NSRDS activities.

The System now includes a complex of data centers and other activities in academic institutions and other laboratories. Components of the NSRDS produce compilations of critically evaluated data, reviews of the state of quantitative knowledge in specialized areas, and computations of useful functions derived from standard reference data. The centers and projects also establish criteria for evaluation and compilation of data and recommend improvements in experimental techniques. They are normally associated with research in the relevant field.

The technical scope of NSRDS is indicated by the categories of projects active or being planned: nuclear properties, atomic and molecular properties, solid state properties, thermodynamic and transport properties, chemical kinetics, and colloid and surface properties.

Reliable data on the properties of matter and materials are a major foundation of scientific and technical progress. Such important activities as basic scientific research, industrial quality control, development of new materials for building and other technologies, measuring and correcting environmental pollution depend on quality reference data. In NSRDS, the Bureau's responsibility to support American science, industry, and commerce is vitally fulfilled.

ERNEST AMBLER, Director

## **Preface**

This report is one of a series of data publications on radiation chemistry; the aim of the series is to compile, evaluate, and present the numerical results on processes occurring in systems which have been subjected to ionizing radiation. Various kinds of data are important in radiation chemistry. The quantities which were measured first were the observed radiation yields or G values (molecules formed or destroyed per 100 eV). Various indirect methods based on G values have been used to determine yields of transient species and relative rates of reactions. The spectral properties (optical, electron spin resonance) of transients have provided a direct method for their identification, and rates of the very fast reactions of transients which occur in irradiated systems have been measured directly by spectroscopic methods. Conductivity and luminescence methods have also provided a means of measuring properties of transients and their kinetics. Some reactions which occur in irradiated systems have also been studied by other methods, such as photochemistry, electric discharge, ultrasonics, chemical initiation, electron impact, etc. The emphasis in these publications is on the data of radiation chemistry, but where other pertinent data exist, they are included.

The data of radiation chemistry are voluminous; thousands of systems have been investigated. As a result there are certain collections, e.g. rate constants of particular types of reactions or certain properties of transients, for which tabulations of the data are considered essential, but for which critical assessment of each value is impossible. On the other hand, certain systems and properties have been studied so extensively that critical examination of these data is desirable and timely. Authors of this series of data publications have been asked to evaluate the extent to which the data can be critically assessed, to describe their criteria for evaluation, and to designate preferred values whenever possible.

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# Optical Spectra of Nonmetallic Inorganic Transient Species in Aqueous Solution

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Optical absorption spectra are presented for 77 nonmetallic, inorganic radicals and radical ions, produced by radiolysis or photolysis in aqueous solution. This atlas results from the compilation, evaluation, and replotting of spectra from the literature. Additional information is included such as extinction coefficients, initial chemical conditions, methods of transient generation and spectral acquisition, references, mechanisms of transient formation, and notes on dosimetry.

Key words: Absorption spectra; aqueous solution; extinction coefficients; inorganic; radical ions; radicals; transients.

### Introduction

Simple inorganic radicals and radical ions in solution have been one of the favorite topics for study in radiation chemistry since the introduction of pulse radiolysis in the early 60's. Some of these radicals had been studied earlier by flash photolysis, but the number of flash photolysis studies increased in parallel with the radiolysis studies. The peak of the research activity on the simple inorganic species came in the late 60's and early 70's. Although many interesting problems remain in this field, a consensus seems to have been reached concerning the identity of the chemical composition of many of these inorganic transients. Such transients are currently being used to form more complicated species in organic and biological systems through oxidation and reduction.

One of the most fruitful techniques that has been used to study the chemistry of the simple inorganic species has been optical absorption spectroscopy. Most of these species absorb ultraviolet and visible radiation. Although optical spectroscopy is not very helpful in determining the exact structure of transients, it is very valuable in following the progress of reactions taking place on the nanosecond or microsecond time scale. Because of this use as a tool to study the kinetics and the use of transient spectra to identify reaction intermediates and individual steps in reaction mechanisms, the optical spectra have become useful data to the experimenter. Since many spectra are now scattered through the literature and since many of these are well characterized, it now seems like an opportune

time to collect all of these spectra together in one volume.

It is the purpose of this volume to collect as many spectra of nonmetallic inorganic transients in aqueous solution as possible. The main emphasis has been to collect the most complete and well-characterized spectra that are available in the literature. In addition, special effort was made to obtain extinction coefficients for these spectra.

# Explanation of Display and Definition of Symbols and Units

The spectra are displayed on a uniform grid, with a standard 0 to 100 relative absorbance scale and with a standard 500 nm long wavelength scale (1000 nm scale is used for hydrated electron spectrum). The relative absorbance, A, is defined by

$$A = N \times \log_{10} (I_0/I) \tag{1}$$

In this equation  $I_0$  is the incident intensity of the monitoring light beam at the wavelength being monitored, I is the intensity of the monitoring light beam, at the same wavelength, that is transmitted through the sample, and N is a normalization factor chosen so that the largest absorbance is 100. The wavelengths are given in nanometers (nm) =  $10^{-9}$  meters. An alternative abscissa scale is used in these spectra on the top line of the plots in a scale that is

proportional to energy. The units used are cm<sup>-1</sup>. One eV is approximately 8068 cm<sup>-1</sup>. To the right of the spectral plots is a sliding extinction coefficient scale.

$$\epsilon = (L[X])^{-1} \log_{10} (I_0/I) \tag{2}$$

 $I_0$  and I are defined the same as above, L is the length of the optical path, and [X] is the concentration of the absorbing species, X.

In addition to the spectral display, additional information is given. Under the heading "Transient," the transient species is identified; other information which may be included under this heading is the wavelength of the maximum absorbance, an extinction coefficient, and a second-order decay constant, k, defined by

$$d[X]/dt = 2k[X]^2. (3)$$

The wavelength of the absorption maximum and the extinction coefficient are given under this heading only when explicitly quoted by the experimenters or when taken from another reference. If  $\lambda(\text{max})$  and  $\epsilon$  are quoted by the experimenters, then the smoothed curves are normalized (see below) to these numbers, both with respect to the ordinate and the abscissa. Extinction coefficients are given in L mol<sup>-1</sup> cm<sup>-1</sup> (dm³ mol<sup>-1</sup> cm<sup>-1</sup>); decay rate constants are given in L mol<sup>-1</sup> s<sup>-1</sup> (dm³ mol<sup>-1</sup> s<sup>-1</sup>). No systematic attempt was made to include the second-order decay rate constants.

The information following the heading "System" concerns the chemical composition of the starting system. Under the headings of "Transient generation" and "Spectral acquisition" the physical equipment used to create the transients and detect them are given. Of particular interest in these two sections is the nature of the dosimetry (measurement of the amount of radiation absorbed in the sample) and actinometry plus the precautions taken to eliminate artifacts, like scattered light. The heading "Reference" contains the literature source of the displayed spectrum.

The information contained under the heading "Transient formation" is of particular interest when transients are prepared in a pulse radiolysis experiment. As opposed to near-ultraviolet or visible photolysis of aqueous solutions where the solutes form the transients in a relatively direct fashion, radiolysis experiments involve the decomposition of the water as a first step. The three major radicals from this process are OH, e(aq), and H. The amount of H formed is five or six times less than that of either OH or e(aq). These three primary radicals can react with solutes or other

transients to form the transients to be studied. Various scavengers are added to prevent the formation of interfering species or to maximize the yield of the transient to be studied. The detailed mechanisms of the authors are given in this section whenever possible, but in many cases these equations are given only as plausible formation equations and should not be understood as precise mechanisms endorsed by the authors who measured the spectrum.

The "Footnotes" are designed mainly to give additional information on alternate extinction coefficients, assumptions on G values,

$$G(X)$$
 = number of X radicals formed for 100 eV  
of the radiation absorbed by the system (4)

and additional information on special purifications and other precautions.

#### Criterion for Inclusion

The following standards were set for the spectra to be included in this atlas. Attempts were made to include spectra that contain contributions from only one transient species, have chemical or physical proof of the chemical assignment, correct for instrumental effects like scattered light, give extinction coefficients, and contain a sufficient number of data points to indicate the precision of the smooth spectral curve. These criteria could not always be met. However, in cases where the transients were considered to be important, the spectra were included with some documentation indicating appropriate qualifications. One common problem, for example, is the failure to correct for spectral contributions of species resulting from H atom reactions in pulse radiolysis experiments in aqueous solutions. This is often neglected on the basis of the low yield of hydrogen atoms relative to hydroxyl radicals and hydrated electrons. However, this contribution is not always negligible, and a spectrum must be viewed with some caution when this type of omission occurs.

All of the spectra in this atlas are taken from the literature. Since the pioneering works on these inorganic transients are approximately 10 to 20 years old, newer spectra often exist in private collections and in the literature. These later spectra are often better characterized than the original works. This has been due in large part to refinements in techniques and to better understanding of the chemical processes. One improvement in equipment that has resulted in more reliable spectral shapes has been the interfacing of the experiment to computers. This has allowed the use of

convenient signal averaging techniques [1]<sup>1</sup>. In addition, the case of data handling with these systems has allowed the experimenter to explore a larger set of wavelengths [2]. Because of these general advances, spectra that have appeared later in the literature than the pioneering works are often chosen for this atlas over the original works.

Even though spectra obtained from private collections often have these same advantages, it was decided that they should be excluded from the atlas since so many of them were inaccessible to this Data Center. However, the Radiation Chemistry Data Center is collecting these newer spectra and is storing them in a computer-readable form. Spectra will be updated as the Center is made aware of new measurements. Some spectra from private collections are already on file.

## Difficulties with Yields and Extinction Coefficients

There is probably much less variability among the various reported spectra of a given transient than among the various reported extinction coefficients of that transient. Many factors could contribute to this difficulty. Among the possible causes are nonuniform irradiation of sample, poor wavelength calibration, faulty dosimetry, scattered light problems, and poor time resolution. Discussion of these and other experimental problems has been thoroughly covered elsewhere [3,4]. However, even when the normal experimental precautions are taken, there remain difficulties. The ultimate source of the problem with extinction coefficients lies in the difficulty of knowing the concentration of a short-lived transient (see eq (2)). Since most of the extinction coefficients in this work are taken from radiolysis works, this means that it is crucial to know the yields (or G values) of the primary radicals, the mechanism of production of the transients from these radicals, and the dose of radiation absorbed by the sample in order to calculate the concentration of the species of interest.

The problem of knowing the G value is particularly acute, for both the sample itself and for the dosimetry. Usually G(X), itself, is not measured directly. It is usually based on a yield of primary radicals that has been estimated from measured yields at infinite dilution of all scavengers. These yields represent the number of primary radicals that diffuse out of the tracks formed by the ionizing radiation and their secondary electrons. Immediately after the arrival of the high energy radiation, the tracks contain a high density of radicals

and other fragments, and many of the radicals combine with each other. Thus only a fraction of the original primary radicals escape into the bulk solution and are measured.

In a usual radiolysis experiment in which absorption spectra are to be measured, scavengers and/or solutes must be added at some finite concentration to react with the primary radicals. However, as the concentration of the scavenger is raised, the chance that the scavenger molecule will initially be present in the volume of a track increases. If this eventually takes place, the primary radicals which would normally combine with other primary radicals in the track will react with the scavenger instead. Thus at high scavenger concentration, scavenging in the track increases the effective yield of primary radicals. The increase in yield depends not only on the scavenger or solute concentration but on the rate constant of the reaction of the scavenger or solute with the primary radicals. This problem has been treated for hydrated electrons [5] and has recently been addressed for hydroxyl radicals [6].

Extinction coefficients can be corrected for scavenging of primary radicals in tracks if the scavenger concentration is known, if the rate constant of the scavenging reaction is known, and if a calibration curve is available. Some calibration curves do now exist [5], but not enough supplementary information is available to make corrections for many cases in the literature. For example, since dosimetry usually involves measuring some well-known transient, it is necessary to know concentrations in the dosimeter as well as in the sample under study. Since corrections cannot be made in many cases, none of the extinction coefficients reported within have these corrections. However, much of the supplementary information that would be needed to make these corrections is included.

# Digitization Methods and Their Sources of Errors

The transcription of the spectra from the literature to the standard format reported in this atlas has led to some unavoidable errors in the resulting data. In order to document the errors, a brief description of the transcription procedure and the possible errors in each step are presented for reference. The data were taken from the literature by first photocopying and then digitizing the spectra from the photocopy. If the drawing was too small for accurate digitization, the photocopy was enlarged before the spectrum was digitized. Where possible the spectra were digitized directly from reprints of journal articles. The digitizer consisted of a digital voltmeter connected to the slide wires of an X-Y

<sup>&</sup>lt;sup>1</sup> Figures in brackets indicate literature references at the end of the text.

recorder. A seven-power comparator was attached to the slide wires to improve both accuracy and precision of each reading.

For normal-sized, full-column graphs taken from an American Chemical Society journal with the ordinate ranging from 0 to 100 and the abscissa covering a 200 nm range, the reproducibility of the digitizer was  $\pm 0.5\%$  for the ordinate and  $\pm 1$  nm for the abscissa. Most of the curves reproduced will have similar uncertainties in the ordinate from the digitization process itself. However, the uncertainties in the abscissa will vary since the wavelength range of the original figures cover a wide range. When the original drawing has the full 500 nm range, the uncertainty in the abscissa might be as large as  $\pm 3$  nm.

In general, the ultimate accuracy is probably not limited by the reproducibility of the digitizer. Other sources of error enter between the data from the author and the final reproduction. One source of distortions is the optics of the xerography and enlarging processes. Some pincushion distortions were found, particularly in the xerography process. In some cases it was found that figures in the literature had their own inherent distortions that had entered in the processing of the initial raw data to the final journal spectrum. Thus a loss in accuracy due to these processes is very difficult to estimate. However, in the overall reproduction process from the literature, it is estimated that the data presented here are accurate to better than  $\pm 2\%$  of the maximum extinction coefficient. The spectral plots were made with a Versatec electrostatic plotter. The resolution of the plots is 200 dots per inch.

All of the continuous spectral curves, but not the data points themselves, that were digitized from the literature were put through a smoothing procedure involving a fast Fourier transform [7]. This procedure had the advantage of averaging the digitization of the spectral curves from the literature, which have the best likelihood of representing the shape of the spectra. Intermediate steps in the smoothing process were displayed on a video GT 62 terminal made by the Digital Equipment Corporation. The output from the smoothing routine is a set of extinction coefficients at regular wavelength intervals. This output was stored in permanent computer files and was plotted along with the individual, digitized data points to give the displayed spectra in this volume. When an  $\epsilon$ (max) was quoted by the authors, the ordinate and abscissa of the plots containing the digitized data were checked and adjusted to match the quoted values.

#### Acknowledgements

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I would like to thank my coworkers in the Radiation Chemistry Data Center, Drs. A. B. Ross and W. P. Helman, for much good advice, criticism, and encouragement. Also, I would like to thank Professor R. H. Schuler and Drs. C. L. Greenstock, P. Neta, and I. K. Patterson for many discussions of the science and technology in this volume. Finally, I would like to thank Dr. A. K. E. Hagopian, who set up this spectral project initially.

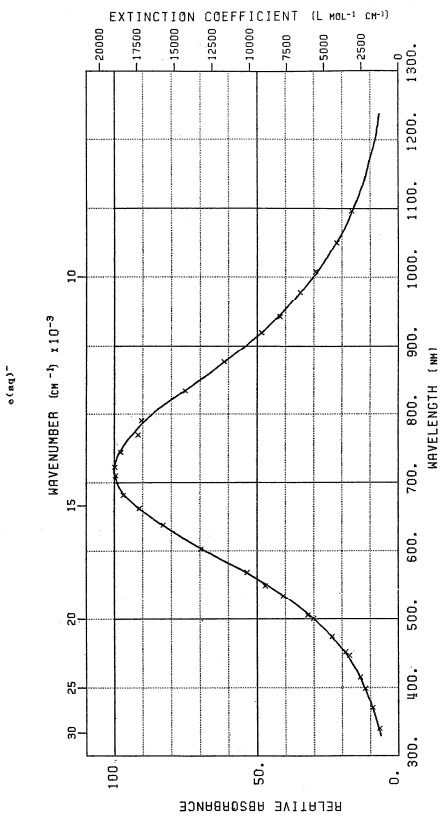
### Symbols and Abbreviations

- A absorbance (optical density)
- ε extinction coefficient (molar absorption coefficient)
- G radiation yield (molecules per 100 eV)
- k rate constant
- krad dose absorbed (6.24 $\times$ 10<sup>16</sup> eV per g) = 10 Gy
- K equilibrium constant
- λ wavelength (nm)
- $pK_a$  -log<sub>10</sub> dissociation constant
- PMT photomultiplier tube

# References

- [1] Patterson, L. K., and Lilie, J., Int. J. Radiat. Phys. Chem. 6(2), 129-41 (1974).
- [2] Gordon, S., Schmidt, K. H., and Martin, J. E., Rev. Sci. Instr. 45(4), 552-8 (1974).
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- [5] Balkas, T. I., Fendler, J. F., and Schuler, R. H., J. Phys. Chem. 74, 4497-505 (1970).
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- [7] Helman, W. P., unpublished work, 1979.

Spectra of Transients

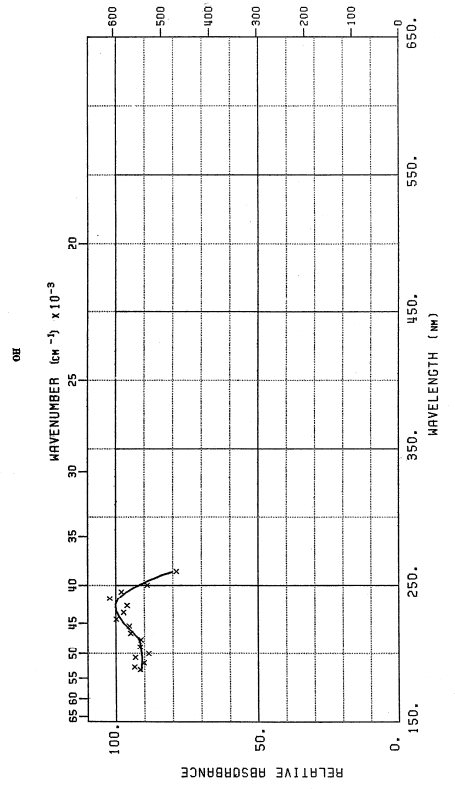


Transients: e(aq)", hydrated electron, \(\text{Max}\) = 720 mm, c720 = 1.9 x 104 L mol-1 cm-1 [11] System: water (triply distilled), 300 K, 1 bar

Reference: F.-Y. Jon and G.R. Freeman, J. Phys. Chem. 81(9): 909-15 (1977)

Transient generation: 2.1 MeV electrons; pulse width 1.0 µs; dosimetry KSCN [2]; steel pressure cell In UV range: Bausch & Lomb 33-86-07 grating in monochromator, band pass = 7 nm; RCA 1P28 PMT; Corning filter were used to eliminate higher order light interference. In visible and ir region [3] gratings were 33-86-02, -03, and -78; infrared photocell, Barnes Engineering A-10D/D108 InSb diode cooled to 77 K. Spectral acquisition:

[1] G.L. Bolton, K.N. Jha, and G.R. Freeman, Can. J. Chem., 54(10): 1497-506 (1976).
[2] Dosimetry oxygen saturated 5 x 10<sup>-3</sup> nol L<sup>-1</sup> KSCN using Ge<sub>478</sub> ((SCN)<sub>2</sub>) = 21000 (100 eV mol L<sup>-1</sup> cm)<sup>-1</sup>.
[100 eV mol L<sup>-1</sup> cm)<sup>-1</sup>.
[3] F.-Y. Jou and G.R. Freeman, Can. J. Chem. 54(23): 3693-704 (1976). Footnotes:



System: aqueous solution of 5.25 x 10-5 mol L-1 NaOH and 5.0 x 10-4 mol L-1 N2O, T = 302 K Transient [1]: OH, hydroxyl radical, c188 = 540 L mol-1 cm-1 [2]

Reference: S.O. Wielsen, B.D. Michael, and E.J. Hart, J. Phys. Chem. 80(22): 2482-8 (1976)

pulse radiolysis using 0.5 µs pulses of 7 MeV electrons; dose per pulse 6.6 krad [3]; dose variations monitored by charge collection method [4] Transient generation:

series [5]; bandwidth at half-max intensity z 4.8 nm; light path purged with  $N_2$  [6]; solar-blind PMT (Ri66, H.T.V. Go., Japan) [7]; XBO 450 W/4 xenon lamp (Osram) with Suprasil envelope operated with a transitorized pulser which intensifies lamp by 30-fold; time delay 0.8  $\mu$ s after pulse two Bausch & Lomb high-intensity grating monochromators connected in Spectral acquisition:

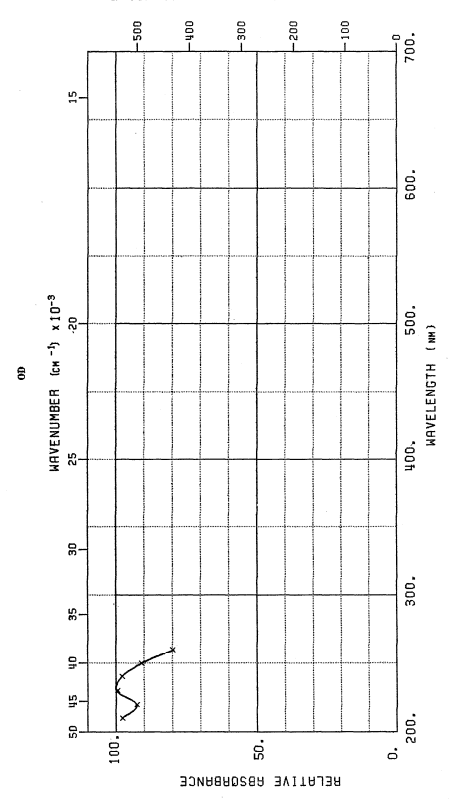
Transient formation:

OH formed in primary processes of radiolysis of H10 also

 $e(aq)^{-} + N_2O (+H_2O) \rightarrow OH + OH^{-} + N_2$ 

- There is a transition in H20 that is perturbed by the presence of an OH radical. [1] The long wavelength portion of the spectrum is thought to be a  $^2\Pi \rightarrow ^2\Sigma^+$ transition that is perturbed by H-bonding to H2O. At wavelengths shorter than 200 nm, the authors think that the intensity may be from an alternate explanation of the spectrum in the literature. Footnotes:
- Absorbances were corrected for  $0H^-$ ,  $H_2O_2$ , and  $N_2O$ . Reactions of these species and the transients were simulated by a computer program using available rate constants. Rate of  $2k(H+H) = 1.55 \times 10^{10} \text{ L mol}^{-1} \text{ s}^{-1}$  and  $G(e(aq)^-) = 2.80$ ;  $G(H^+) = 2.90$ ; G(OH) = 2.70;  $G(OH^-) = 0.10$ ; G(H) = 0.4B;  $G(H_2) = 0.45$ ;  $G(H_2) = 0.70$ ;  $G(H_2) = 0.02$ . 2
  - il Sample changed after each pulse.
- Total dose was obtained by matching the calculated e(aq) transient with that [4]
  - observed at 600 nm using alkaline solution. [5] Gratings blazed for maximum efficiency at 250 nm.
- [6] With this system transient spectra could be measured to  $182 \, \mathrm{nm}$ , but  $\mathrm{H}_2\mathrm{O}$  absorption interfered below  $188 \, \mathrm{nm}$ .
  - Amount of scattered light at the shortest wavelength was less than 1%.

# EXTINCTION COEFFICIENT (L MOL-1 CM-1)



System: D<sub>2</sub>0 solution of 3 x 10<sup>-5</sup> mol L<sup>-1</sup> DC104 and 2 x 10<sup>-3</sup> mol L<sup>-1</sup> N<sub>2</sub>0

Transient [1,2]: 0D

Reference: S.O. Nielsen, B.D. Michael, and E.J. Hart, J. Phys. Chem. 80(22): 2482-8 (1976)

pulse radiolysis; pulse energy 10-11 MeV; pulse length 1.0  $\mu s$ ; dosimetry K4Fe(CN)  $_6$  [4]; dose per pulse  $\approx$  5 krad Transient generation [3]:

resulting in an increase in brightness of 25-30 times the nonpulsed lamp; optical windows of cell were of Suprasil quartz; lenses were of Suprasil quartz; Zeiss MM 12 double quartz prism monochromator [5]; EMI 9558Q PMT [6]; time delay, extrapolated back to zero Osram xenon lamp XBO 450 with Suprasil quartz bulb; lamp pulsed Spectral acquisition [3]:

Transient formation:

OD from radiolysis of D2O

 $e(sol)^{-} + N_2O (+D_2O) \rightarrow OD + OD^{-} + N_2$ 

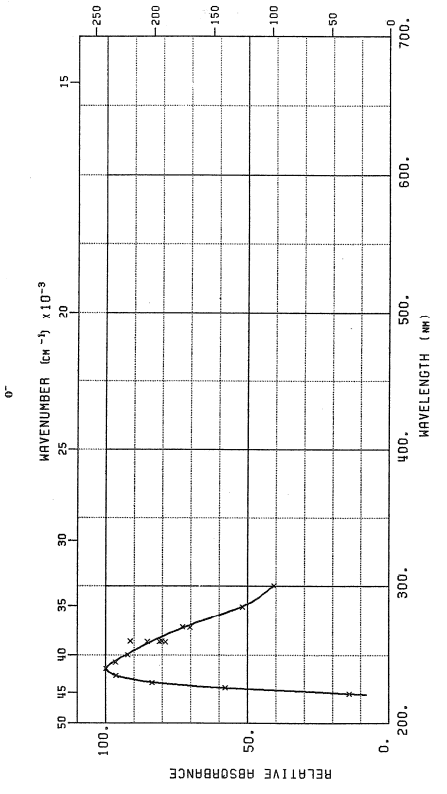
At wavelengths shorter than 200 nm, the authors think that the intensity may be from a transition in B20 that is perturbed by the presence of an OD radical. There is an [1] The long wavelength portion of the spectrum is thought to be a  $^2\Pi \to ^2\Sigma^+$  transition. alternate explanation of the corresponding OH spectrum in the literature. Footnotes:

Galculation of concentrations were made using G(D) = 0.44, G(e(sol)) = 2.99, G(OD) = 2.99, G(D<sub>2</sub>O<sub>2</sub>) = 0.62 with a small correction for radical recombination during pulse. P. Pagsberg, H. Christensen, J. Rabani, G. Nilsson, J. Fenger, and S.O. Nielsen, J. Phys. Chem. 73(4): 1029-38 (1969). [3]

Aqueous solution of  $10^{-3}$  mol L<sup>-1</sup> K<sub>4</sub>Fe(GN)<sub>6</sub> saturated with N<sub>2</sub>O; G(Fe(GN)<sub>6</sub><sup>3-</sup>) = G(OH) + G(e(aq)<sup>-</sup>) = 2.65 + 2.6 = 5.25 and c<sub>420</sub>(Fe(GN)<sub>6</sub><sup>3-</sup>) = 1000 L mol<sup>-1</sup> cm<sup>-1</sup>. No scattered light could be detected at 200 nm; at maximum slit width of 2.0 mm the [4]

bandwidth at 200 nm was 0.95 nm. [2]

[6] Monochromator and PMT set behind a 10 cm lead shield.



System: aqueous solution of 0.105 mol L.1 NaOH, pH = 13, 0.5 atmospheres N2O, carbonate free Transient: 0<sup>-</sup>, λ(max) = 240 nm, ε<sub>249</sub> = 240 L mol<sup>-1</sup> cm<sup>-1</sup> [1]

12

Reference: J. Rabani, Advan. Chem. Ser. 81: 131-52 (1968)

pulse radiolysis using syringe technique; electron beam monitored by split beam technique [2] Transient generation:

450 W Osram xenon monitoring lamp, corrections made for scattered light [3]; Bausch & Lomb monochrometors Spectral acquisition:

Transient formation:

 $H + OH^- + e(aq)^- (+H_2O)$ 

 $e(aq)^{-} + N_2O (+H_2O) \rightarrow N_2 + OH^{-} + OH$ 

 $e(aq)^{-} + N_20 \rightarrow N_2 + 0^{-}$ 

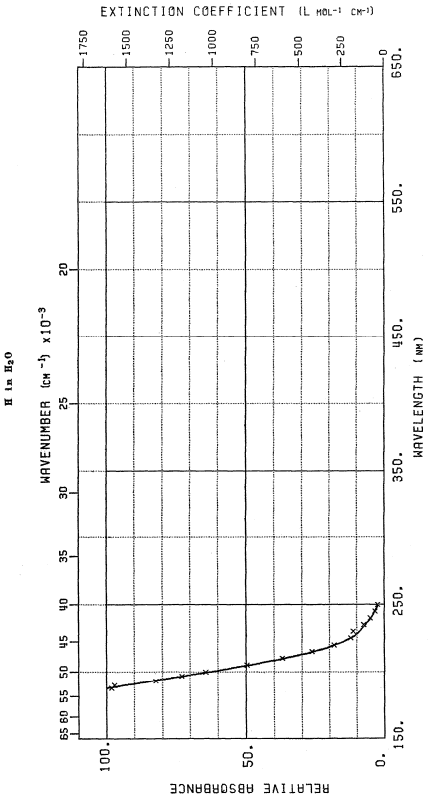
OH + OH" ← O" + H2O, pKa (OH) = 11.85

[1] Using G(OH) + G(O<sup>-</sup>) = 5.8 and G(H<sub>2</sub>O<sub>2</sub>) = 0.7. Corrections were made for Footnotes:

the absorption of OH and H202.

[2]

One beam was passed through a monochromator set at 260 nm to monitor the electron beam intensity. Absolute dosimetry used 5 x 10<sup>-4</sup> mol L<sup>-1</sup> K<sub>4</sub>Fe(CN)<sub>6</sub> under experimental conditions, taking c<sub>429</sub> = 1000 L mol<sup>-1</sup> cm<sup>-1</sup>. Depending on the optical alignment, the scattered light was 3-6% at 260 nm, 8% at 250 nm, 10-15% at 240 nm, 20-25% at 230 nm, and 50% at 222 nm. [3]



System: aqueous solution of 3.39 x 10<sup>-4</sup> mol L<sup>-1</sup> HG10<sub>4</sub>, 2.82 x 10<sup>-2</sup> mol L<sup>-1</sup> H<sub>2</sub> (30 atm), and 4.5 x 10<sup>-5</sup> mol L<sup>-1</sup> H<sub>2</sub>0<sub>2</sub> (steady state concentration from radiolysis), T = 307 K

Transient [1]: H atom in H2O, c188 = 1620 L mol-1 cm-1 [2]

Reference: S.O. Nielsen, B.D. Michael, and E.J. Hart, J. Phys. Chem. 80(22): 2482-8 (1976)

pulse radiolysis using 1.0  $\mu s$  pulses of 7 MeV electrons; dose per pulse 3.3 krad; dose variations monitored by charge collection method [3] Transient generation:

series [4]; bandwidth at half-max intensity = 4.8 nm; light path purged with N<sub>2</sub> [5]; solar-blind PMT (R166, H.T.V. Co., Jpn) [6]; XBO 450 W/4 xenon lamp (Osram) with Suprasil envelope operated with transistorized pulser which intensifies lamp by 30-fold; time delay 1.0  $\mu s$  after pulse two Bausch & Lomb high-intensity grating monochromators connected in Spectral acquisition:

Translent formation:

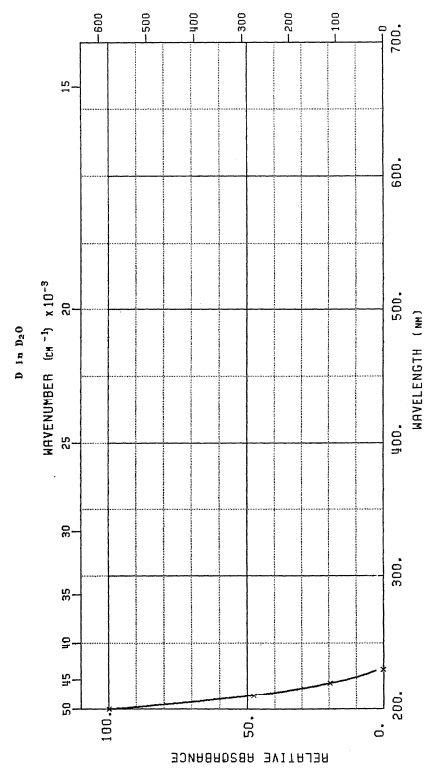
H from primary processes in radiolysis of H20

e(ad)" + H<sup>+</sup> → H

OH + H<sub>2</sub> + H<sub>2</sub>O + H

- The ground state was not perturbed by the H [1] Authors assign the spectrum to a transition that mainly involves the  ${\rm H}_2{\rm O}$  molecule. The excited state was thought to involve a partial negative charge There are alternate assignments of this spectrum in the literature. transfer to a neighboring H atom. Footnotes:
  - (2) Absorbances were corrected for presence of  $H_2O_2$ . This correction was made using the calculated concentrations for  $H_2O_2$  and transient species that were obtained by a computer simulation of the overall reaction using available rate constants. Yields used were  $G(e(aq)^{-}) = 2.89$ ;  $G(H^+) = 2.99$ ; G(0H) = 2.79;  $G(0H^-) = 0.19$ ; G(H) = 0.48;  $G(H_2) = 0.45$ ;  $G(H_2O_2) = 0.79$ ;  $G(HO_2) = 0.02$ . Correction
- was made for radiation-induced absorption of quartz cell used for high pressures. Total dose was obtained by matching the calculated e(aq) transient with that observed at 600 nm using alkaline solution.
  - [4] Gratings blazed for maximum efficiency at 250 nm. [5] With this system transient spectra could be measured to 182 nm, but  $\rm H_20$  absorption interfered below 188 nm.
    - Amount of scattered light at the shortest wavelength was less than 1%.

[9]



Transient [1,2]: D atom in  $D_2O$ System:  $D_2O$  solution of  $10^{-3}$  mol  $L^{-1}$   $DG10_4$ 

Reference: S.O. Nielsen, B.D. Michael, and E.J. Hart, J. Phys. Chem. 80(22): 2482-8 (1976)

pulse radiolysis; pulse energy 10-11 MeV; pulse length 1.0 µs; dosimetry K4Fe(GN)6 [4]; dose per pulse \* 5 krad Transient generation [3]:

resulting in an increase in brightness of 25-30 times the nonpulsed lamp; optical windows of cell were of Suprasil quartz; lenses were of Suprasil quartz; Zeiss MM 12 double quartz prism monochromator [5]; Osram xenon lamp XBO 450 with Suprasil quartz bulb; lamp pulsed EMI 95580 PMT [6]; time delay, extrapolated back to zero Spectral acquisition [3]:

Transfent formation:

D from radiolysis of D20

 $D^+ + e(sol)^- \rightarrow D$ 

D atom. There are alternate assignments of the corresponding H atom spectrum in the molecule. The excited state was thought to involve a partial negative charge transfer to a neighboring D atom. The ground state was not perturbed by the [1] Authors assign the spectrum to a transition that mainly involves the D20 literature. Footnotes:

Calculation of concentrations were made using G(D) = 0.44, G(e(sol) = 2.90, G(OD) = 2.90,  $G(D_2O_2)$  = 0.62 with a small correction for radical recombination during pulse. [2]

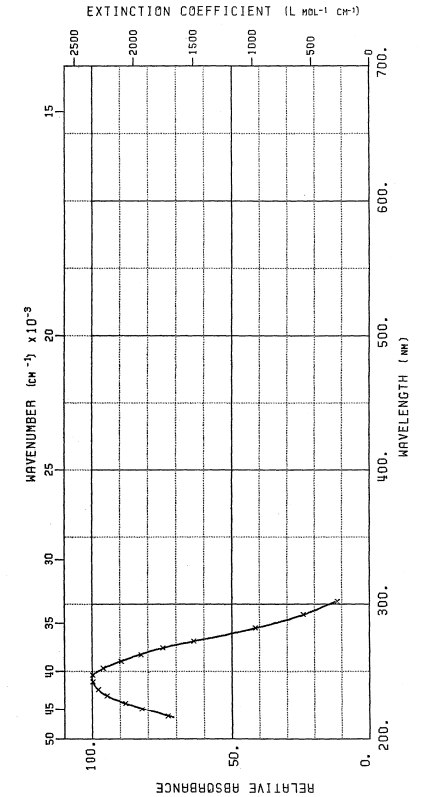
[3] P. Pagsberg, H. Christensen, J. Rabani, G. Nilsson, J. Fenger, and S.O. Nielsen, J. Phys. Chem. 73(4): 1629-38 (1969).

[4] Aqueous solution of 16<sup>-3</sup> mol L<sup>-1</sup> K<sub>4</sub>Fe(GN)<sub>6</sub> saturated with N<sub>2</sub>O; G(Fe(GN)<sub>6</sub><sup>3-</sup>) = G(OH) + G(e(aq)<sup>-</sup>) = 2.65 + 2.6 = 5.25 and e<sub>429</sub>(Fe(GN)<sub>6</sub><sup>3-</sup>) = 1000

[5] No scattered light could be detected at 200 nm; at maximum slit width of 2.0 mm, the bandwidth at 200 nm was 0.95 nm.

[6] Monochromator and PMT set behind a 10 cm lead shield.





Transient:  $0_2^-$ , superoxide radical ion,  $\lambda(\max) = 245$  nm,  $\epsilon_{245} = (2350 \pm 2)$ ) L mol<sup>-1</sup> cm<sup>-1</sup> [1,2], k( $0_2^- + 0_2^-$ ) less than 0.35 L mol<sup>-1</sup> s<sup>-1</sup>

System: air-saturated, aqueous [3] 10-2 mol L<sup>-1</sup> sodium formate solution, containing 10-4 mol L<sup>-1</sup> EDTA [4] and Na<sub>3</sub>PO<sub>4</sub> [5] added to adjust pH to 10.5

Reference: B.H.J. Bielski, Photochem. Photobiol. 28: 645-9 (1978)

Translent generation: pulse radiolysis, using stopped-flow technique [6]

Spectral acquisition:

Transient formation:

$$e(aq)^- + 0_2 \rightarrow 0_2^-$$

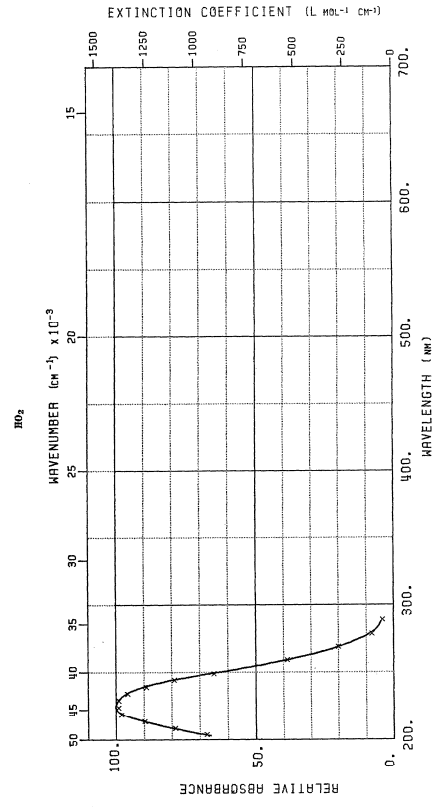
$$H + \theta_2 \rightarrow H\theta_2$$

OH + H0<sub>2</sub> 
$$\star$$
 0<sub>2</sub> + H<sub>2</sub>0, pK<sub>a</sub> (H0<sub>2</sub>) : 4.69  $\pm$  0.08

$$0H + HC00^{-} \rightarrow C0_{2}^{-} + H_{2}0$$
,  $pK_{a}(HC00H) = 3.77$ 

$$C0_2^- + 0_2 \rightarrow 0_2^- + C0_2$$

- Based (a) on extinction coefficient of nitroform radical anion, G(NO<sub>2</sub>)<sub>3</sub><sup>-</sup>, which was formed by the reaction of O<sub>2</sub><sup>-</sup> and tetranitromethane and (b) on difference in 550 nm extinction coefficients of ferrocytochrone c and ferricytochrone c. The ferrocytochrone c was formed by the reaction of O<sub>2</sub><sup>-</sup> with ferricytochrone c. No G(O<sub>2</sub><sup>-</sup>) was assumed with this method. [1] Footnotes:
  - Earlier values for c(max) are listed in the review by G. Czapski, Ann. Rev. Phys. Chem. 22: 171-208 (1971). [2]



Transient:  $HO_2$ , perhydroxyl radical,  $\lambda(\max) = 225$  nm,  $\epsilon_{225} = (1400 \pm 80)$  L mol<sup>-1</sup> cm<sup>-1</sup> [1,2], k(HO<sub>2</sub> + HO<sub>2</sub>) = (8.60 ± 0.62) x  $10^6$  L mol<sup>-1</sup> s<sup>-1</sup> System: air-saturated, aqueous [3] 10-2 mol L-1 sodium formate solution with pH adjusted to 1.5 using perchloric acid

Reference: B.H.J. Bielski, Photochem. Photobiol. 28: 645-9 (1978)

Transient generation: pulse radiolysis, using stopped-flow technique [4]

Spectral acquisition:

Transient formation:

e(aq) + H+ + H

 $H + HC00H \rightarrow CO_2H + H_2$ ,  $pK_a$  (HC00H) = 3.77

OH + HCOOH  $\rightarrow$  CO<sub>2</sub>H + H<sub>2</sub>O

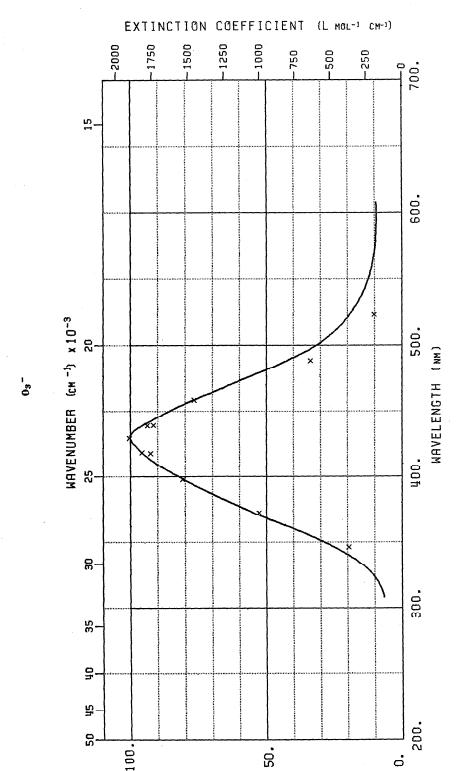
 $CO_2H + O_2 \rightarrow O_2^- + CO_2 + H^+$ 

 $0_2^- + H^+ + H0_2$ ,  $pK_n (H0_2) = 4.69 \pm 0.08$ 

Footnotes:

[1] Determined from calibrations of beam current based on 02 concentration, where 02 concentration was obtained without assuming a G(02).
[2] Earlier values for c(max) are given by G. Czapski in Ann. Rev. Phys. Chem. 22: 171-208 (1971).

[3] Water purified by a Milli-Q reagent grade water system.
[4] B.H.J. Bielski and H.W. Richter, J. Am. Chem. Soc. 99(9): 3019-23 (1977).



System: aqueous solution of  $0_2$  and  $N_20$  with  $[N_20]/[0_2] \approx 20$ , pH  $\approx 13.5$  [3] Translent [1]: 03, ozonide ion, £430 = (1900 ± 120) L mol-1 cm-1 [2]

RELATIVE ABSORBANCE

Reference: W.D. Felix, B.L. Gall, and L.M. Dorfman, J. Phys. Chem. 71(2): 384-92 (1967)

pulse radiolysis using a Varian V-7715A linear accelerator, syringe technique; dosimetry [4] Transient generation:

500 W Osram xenon lamp (Type XBO 450 W); 1P28 PMT; two Bausch & Lomb grating monochromators (Type 33-86-25, f/3.5) with light beam split by partially reflecting mirror; bandwidth: less than 1.8 nm Spectral acquisition:

Transient formation:

 $e(aq)^{-} + N_20 \rightarrow N_2 + 0^{-}$ 

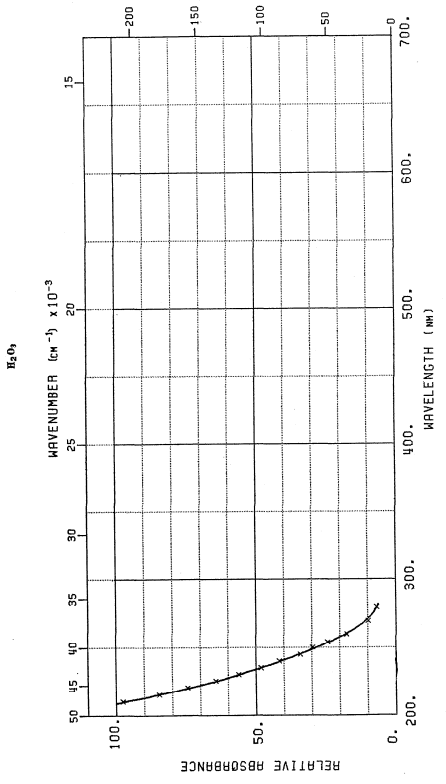
 $OH^{-} + OH + H_{2}O + O^{-}, pK_{a}(OH) = 11.85$ 

 $0^- + 0_2 \rightarrow 0_3$ 

ø [1] Solid curve is drawn from a composite of the data presented plus data from Footnotes:

solution free of  $\rm N_2O$ . Based on G(O<sup>-</sup>) = 3.3 at pH = 13.1 in O<sub>2</sub>-saturated solution. Under these conditions most e(aq)<sup>-</sup> are converted to O<sup>-</sup>.

[2] Based on G(0") = 3.3 at pu \_\_\_\_\_\_ are converted to U .
[3] Under these conditions most e(aq) are converted to U .
[4] Dosimeter was an O<sub>2</sub>-saturated sulfuric acid solution with no chloride ions and with 10<sup>-2</sup> mol L<sup>-1</sup> Fe<sup>2+</sup>. At high dose rates, G(Fe<sup>3+</sup>) = 15.6. At low dose, c<sub>304</sub>(Fe<sup>3+</sup>) = 2200 L mol<sup>-1</sup> cm <sup>-1</sup> was used to calculate dose. At high intensity, dose was calculated by monitoring 366 nm and by using c<sub>304</sub>/c<sub>366</sub> B.9 for Fe<sup>3+</sup>.



Transient:  $\rm H_2O_3$ , hydrogen sesquioxide [1] System: air-saturated, aqueous  $10^{-2}$  mol  $\rm L^{-1}$  HClO4, pH  $\approx 2$ 

Reference: B.H.J. Bielski and H.A. Schwarz, J. Phys. Chem. 72(11): 3836-41 (1968)

electrons from 2. MeV Van de Graaff generator; variable pulse length, a few hundred microseconds to several tenths of a second; ferrous dosimetry Transient generation:

deuterium analyzing lamp; optical path length = 6.1 cm; Bausch & Lomb f/3.5 monochromator; tandem nonochromators used below 240 nm [2]; dual Spectral acquisition:

beam oscilloscope

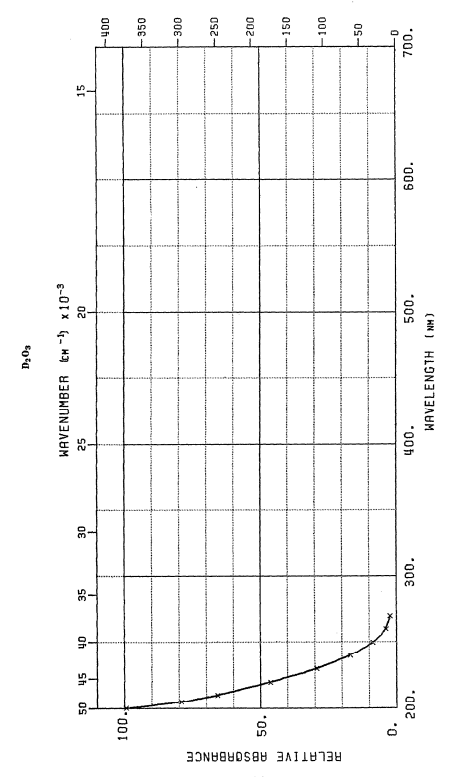
Transient formation:

 $0H + HO_2 \rightarrow H_2O_3$ ,  $pK_a(HO_2) = 4.69$  [3]

 $9 < pK_a(H_2O_3) < 10 [4]$ 

[1] Extinction coefficients were calculated using G(H2O3) = 1.7 from reference in footnote 4. Footnotes:

[2] Scattered light always kept below 2% of the analyzed light beam. [3] B.H.J. Bielski, Photochem. Photobiol. 28(4-5): 645-9 (1978). [4] G. Czapski and B.H.J. Bielski, J. Phys. Chem. 67(10): 2180-4 (1963).



Transient [1]:  $D_2\theta_3$ , deuterium sesquioxide System: oxygenated  $D_2\theta$  [1]

Reference: B.H.J. Bielski, J. Phys. Chem. 74(17): 3213-6 (1970)

Transient generation: pulse radiolysis using 1.95 MeV Van de Graaff generator; pulse length 0.1 s [2]; dosimetry-ferrous sulfate

deuterium monitoring lamp; 3-pass Suprasil quartz cell with optical path = 6.1 cm; two tandem coupled Bausch & Lomb f/3.5 monochromators [3]; 7200 RCA PMT Spectral acquisition:

Transient formation:

 $D + 0_2 \rightarrow D0_2$ 

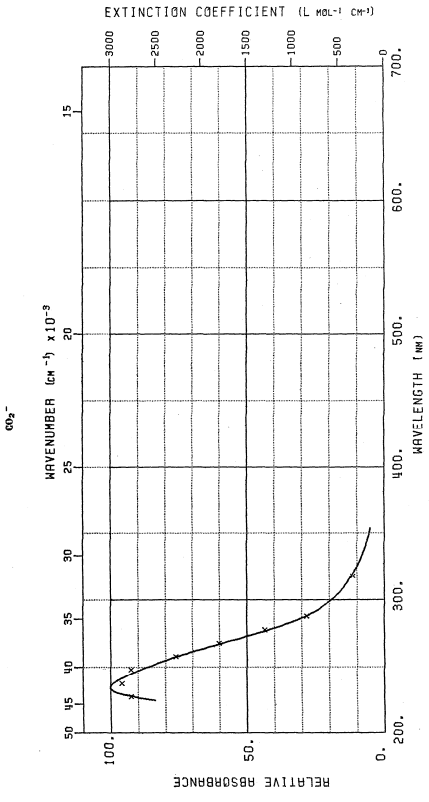
 $e(sol)^- + 0_2 \rightarrow 0_2^-$ 

 $D^+ + 0_2^- \rightarrow D0_2$ 

 $0D + D0_2 \rightarrow D_20_3$ 

[1] Extinction coefficient based on  $G(D_2O_3) = 1.86$  which was determined in a  $10^{-2}$  nol L<sup>-1</sup> Footnotes:

DC104 solution. [2] Spectrum corresponds to time at end of pulse. [3] Scattered light at 200 nm was < 0.3% of total light signal.



Transient [1]: CO2-, carboxyl radical, \(\text{(max)} = 235 nm, \(\epsilon\_{235} = 3000 L mol^{-1} cm^{-1} [2,3]\) System: aqueous solution of 3 x  $10^{-2}$  mol L<sup>-1</sup> HC00<sup>-</sup>, 1 atm N<sub>2</sub>0, pH 9

Reference: P. Neta, M. Simic, and E. Hayon, J. Phys. Chem. 73(12): 4207-13 (1969)

pulse radiolysis, using Febetron 705; pulse energy 2.3 MeV; pulse length  $\approx$  30 ns; dose per pulse 8 krad; dosimetry 0.1 mol L<sup>-1</sup> SCN<sup>-</sup> solution [4] Transient generation:

Osram XBO 450 W xenon lamp (pulsed); two high-intensity Bausch & Lomb monochromators, used in series to reduce scattered light [5]; two EMI 9558 QB PMT; dual beam Fairchild 777 oscilloscope Spectral acquisition:

Translent formation:

 $N_2O + e(aq)^- (+H_2O) \rightarrow N_2 + OH^- + OH$ 

 $HCO_2^- + OH \rightarrow CO_2^- + H_2O$ ,  $pK_a$  (HCOOH) = 3.77

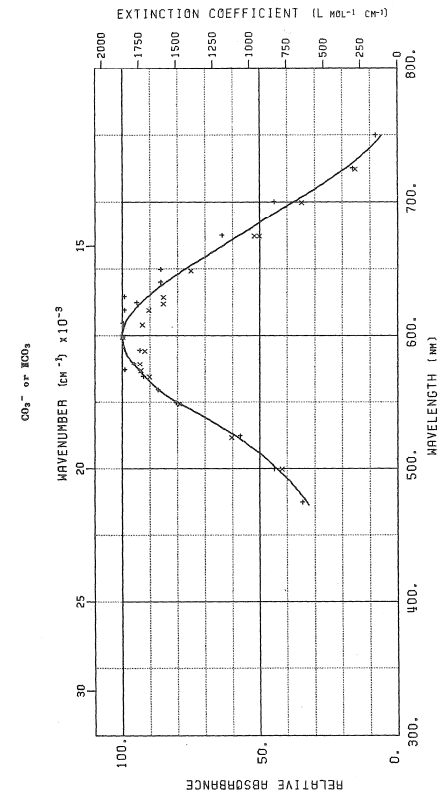
 $HCO_2$  +  $H \rightarrow CO_2$  +  $H_2$ 

[1] Smooth curve for spectrum is from a composite spectrum. Footnotes:

[2] Earlier results gave  $\lambda(\text{max}) = (250 \pm 5)$  nm and  $\epsilon_{250} = 2400$  L mol<sup>-1</sup> cm<sup>-1</sup>.

J.P. Keene, Y. Raef, and A.J. Swallow, Pulse Radiolysis, M. Ebert, J.P. Keene, A.J. Swallow, and J.H. Baxendale (eds.), Academic Press, London, 1965, pp.99-106.
[3] Using  $G(e(\text{aq})^-) = G(OH) = 2.8$ ; deviation  $\pm$  15%.
[4] Using  $\epsilon_{500} = 7600$  L mol<sup>-1</sup> cm<sup>-1</sup> for  $(SCN)_2^-$ .
[5] At 230 and 210 nm the amount of scattered light was 1% and 18%, respectively.

M. Simic, P. Neta, and E. Hayon, J. Phys. Chem. 73(11): 3794-8000 (1969).



the chemical system represented by the data points "x" was an aqueous solution of 1 mol L<sup>-1</sup> NaHCO<sub>3</sub>,  $10^{-3}$  mol L<sup>-1</sup> H<sub>2</sub>O<sub>2</sub>, N<sub>2</sub>-saturated, pH B; and the system represented by the data points "+" was an aqueous solution of 1 mol L<sup>-1</sup> Na<sub>2</sub>CO<sub>3</sub>, 2 x  $10^{-4}$  mol L<sup>-1</sup> H<sub>2</sub>O<sub>2</sub>, N<sub>2</sub>-saturated, pH 13 Transfent:  $CO_3^-$  or  $HCO_3$  [1],  $\lambda(\max) = 600$  nm,  $\epsilon_{600} = 1860$  L mol<sup>-1</sup> cm<sup>-1</sup> [2] System:

Reference: D. Behar, G. Czapski, and I. Duchovny, J. Phys. Chem. 74(19): 2206-10 (1970)

Transient generation [3]: flash photolysis, four  $0_2$ -filled (5 mm Hg) Spectrosil grade quartz lamps operated at 20 kV and 2  $\mu F$  fired through a triggered spark gap

Spectral acquisition [3]: monitoring light was 120 W tungsten lamp; two 9558B EMI PMT; split beam technique; two Bausch 8 Lomb monochromators types 567 AB and F.D. 999

Transient formation [4]:

 $H_2O_2 + h\nu \rightarrow 20H$ 

 $0H^{-} + 0H + 0^{-} + H_{2}0$ ,  $pK_{a}(0H) = 11.85$ 

OH + CO32 - - CO3 + OH-

 $0^{-} + C0_3^{2^{-}} (+H_20) \rightarrow C0_3^{-} + 20H^{-}$ 

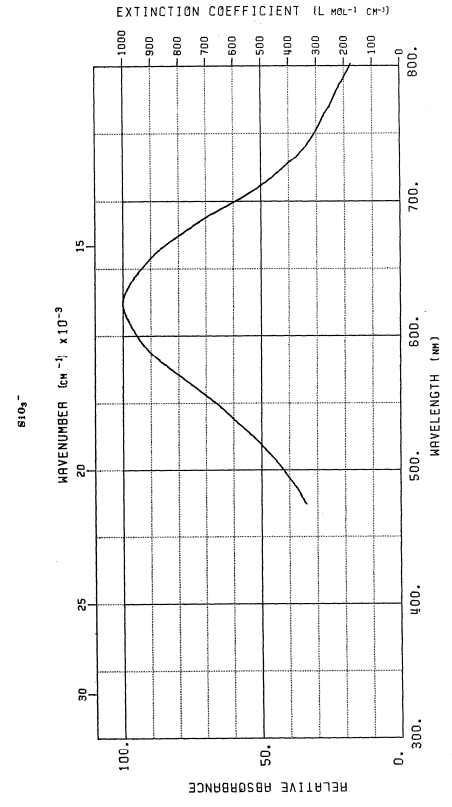
 $0H + HCO_3^- \rightarrow CO_3^- + H_2O$ 

Footnotes: [1] pKs.(HCO<sub>3</sub>) = 9.6 ± 0.3; S. Chen, V.W. Cope, and M.Z. Hoffman, J. Phys. Chem. 77(9): 1111-6 (1973). These authors reported no difference between the spectra of HCO<sub>3</sub> and CO<sub>3</sub>.

[2] J.L. Weeks and J. Rabani, J. Phys. Chem. 70(7): 2100-6 (1966).

[3] D. Behar and G. Czapski, Israel J. Chem. 6: 43-51 (1963).

[4] By measuring the rate constants in a pulse radiolysis experiment, it was found that reactions of OH with H<sub>2</sub>O<sub>2</sub> to form O<sub>2</sub> or HO<sub>2</sub> were negligible under the experimental conditions.



System: aqueous solution of 5 x 10<sup>-2</sup> mol L<sup>-1</sup> Na<sub>2</sub>SiO<sub>3</sub> and 1 x 10<sup>-2</sup> mol L<sup>-1</sup> Na<sub>2</sub>S<sub>2</sub>O<sub>8</sub> Transient: Si03, λ(max) = 625 nm, ε = 1.0 x 103 L mol-1 cm-1

Reference: V.A. Kuz'min and A.K. Chibisov, High Energy Cham. 4(2): 146-7 (1970)

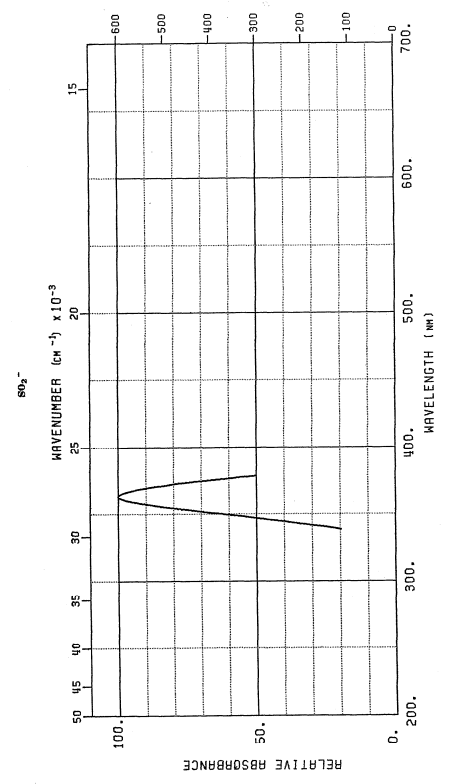
Transient generation: flash photolysis

Spectral acquisition:

Transient formation:

$$S_20_8^{2-} + h\nu \rightarrow 2S0_4^{-}$$
  
 $S0_4^{-} + S10_2^{2-} \rightarrow S0_4^{2-} + S$ 

$$S0_4^- + S10_3^{2-} \rightarrow S0_4^{2-} + S10_3^-$$



Transient [1]:  $S0_2^-$ ,  $\lambda(max) \approx 365$  nm,  $\epsilon_{365} \approx 600$  L mol<sup>-1</sup> cm<sup>-1</sup> [2,3],  $2k = 1.0 \times 10^{10}$  L mol<sup>-1</sup> s<sup>-1</sup>

System: aqueous solution of 6.0 x 10-2 HSO3-, N2 (1 atm), pH 4.0

Reference: E. Hayon, A. Treinin, and J. Wilf, J. Am. Chem. Soc. 94(1): 47-57 (1972)

Transient generation: flash photolysis; 1800 J; duration at "1/e" time of 5 μs; total pulse length 70 μs; 240 nm cutoff filter [4]

Spectral acquisition: Bausch & Lomb grating monochromator; PMT; spectrum was obtained by a point-by-point method from oscilloscope traces; each point was average of 3 or more experiments; time delay 80 µs after pulse

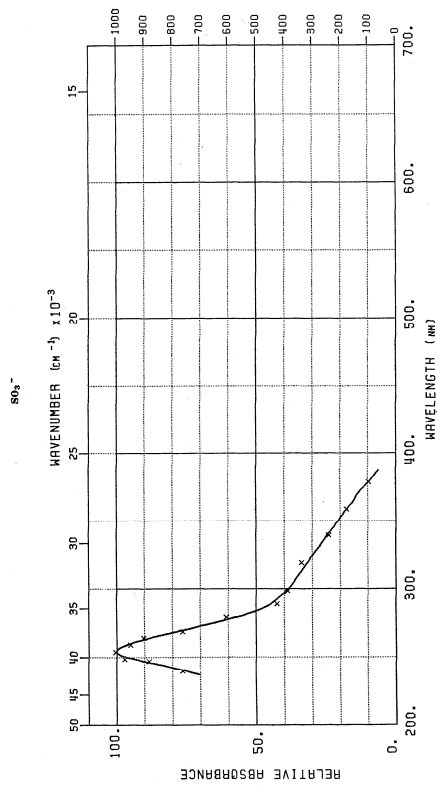
Transient formation [4]:

$$^{-}0_{3}S-S0_{2}^{-}$$
 + h $\nu$  +  $S0_{3}^{-}$  +  $S0_{2}^{-}$ 

[1] Spectrum obtained by subtracting contribution of  $S_20_4^{2-}$  formed from Footnotes:

$$2SO_2^- \rightarrow S_2O_4^{2-}$$
.

- [2] Based on  $\varepsilon$  (max) = 8300 L mol<sup>-1</sup> cm<sup>-1</sup> for  $S_2O_4^{2-}$ . [3] Displayed spectrum normalized to 600 L mol<sup>-1</sup> cm<sup>-1</sup> at 363 nm. [4] Cutoff filter permitted only  $S_2O_5^{2-}$  to be photolyzed.



Transient [1,2]: SO<sub>3</sub>, \(\)(max) = 255 nm, \(\epsilon\_{255} \pi 1600 L mol^1 cm^1, 2k = (1.1 \pi 0.2) x 100 L mol^1 s^1 System: aqueous solution of 2 x  $10^{-3}$  mol L<sup>-1</sup>  $SO_3^{2-}$ , N<sub>2</sub>0-saturated, pH 10.3

Reference: E. Hayon, A. Treinin, and J. Wilf, J. Am. Chem. Soc. 94(1): 47-57 (1972)

pulse radiolysis, using Febetron 705 System (Field Emission Corp.); pulse energy 2.3 MeV; pulse length  $\approx 30$ ns; dosimetry -  $\rm N_2O-saturated$ , aqueous solution of KSCN [3] Transient generation:

Osram XBO 450 W xenon lamp (pulsed); two high intensity Bausch 8 Lomb monochromators, used in series to reduce scattered light [4]; two EMI 9558 QB PMT; dual beam Fairchild 777 oscilloscope Spectral acquisition:

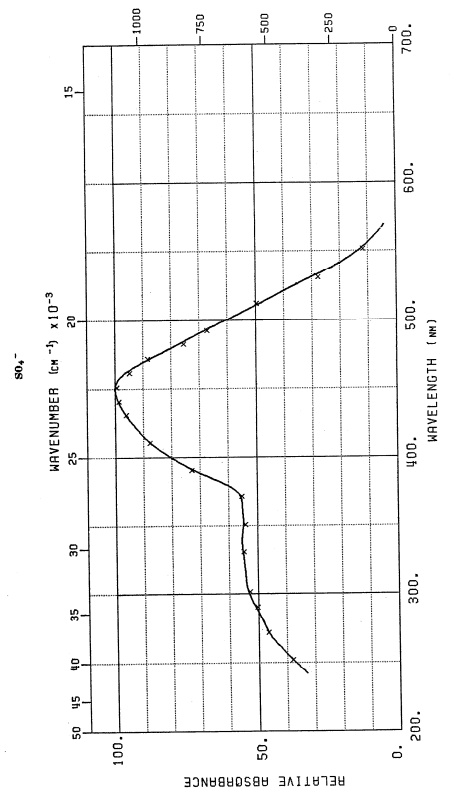
Translent formation:

 $N_2O + e(aq)^- (+H_2O) \rightarrow N_2 + OH + OH^-$ 

 $0H + SO_3^{2-} \rightarrow SO_3^{-} + 0H^{-}$ 

[1] Points in spectrum are averages of three or more experiments. Footnotes:

[2] Spectrum remains unchanged in pH range 4-10.
[3] Using e500 (SCN)<sub>2</sub> = 7600 L mol<sup>-1</sup> cm<sup>-1</sup> for  $10^{-1}$  mol L<sup>-1</sup> SCN<sup>-</sup>,
N<sub>2</sub>O-saturated solutions at pH 5.5 and assuming G = G(e(aq)<sup>-</sup>) + G(OH) = 5.5,
E. Hayon, J. Chem. Phys. 51(11): 4881-92 (1969).
[4] At 230 and 210 nm, the amount of scattered light was 1% and 18%, respectively.



Transient: SO4<sup>-</sup>, sulfate radical anion, λ(max) = 450 nm [1] , ε450 = 1100 L mol<sup>-1</sup> cm<sup>-1</sup> [1,2] System: aqueous solution of 2 x  $10^{-3}$  mol L<sup>-1</sup> S<sub>2</sub>08<sup>2-</sup>, Ar(1 atm), pH = 5.1

Reference: E. Hayon, A. Treinin, and J. Wilf, J. Am. Chen. Soc. 94(1): 47-57 (1972)

electrons from Febetron 705 System (Field Emission Corp.); pulse energy 2.3 MeV; pulse length  $\approx$  30 ns; dosimetry [3] N<sub>2</sub>O-saturated aqueous solution of 0.1 mol L<sup>-1</sup> SGN<sup>-</sup> at pH 5.5 Transient generation:

Osram XBO W xenon lamp; two high-intensity Bausch & Lomb monochromators used in series to reduce scattered light; two EMI 95580B PMT; dual beam Fairchild 777 oscilloscope; quartz cells with optical path lengths of 2 cm; spectrum is the average of 3 runs. Spectral acquisition:

Transient formation:

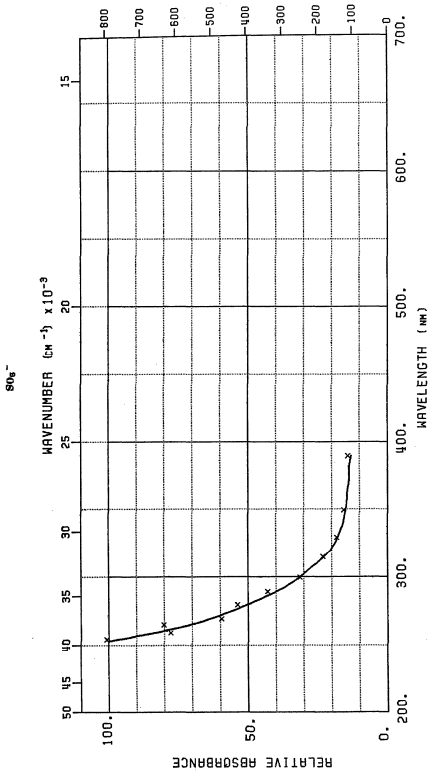
 $S_20_8^{2-} + e(aq)^- \rightarrow S0_4^{2-} + S0_4^-$ 

[1] With spectrum of SO4- displayed, Hayon, Treinin, and Wilf quoted (but apparently did not measure) a value of  $\lambda(\max) = 450$  nm and  $\epsilon_{480} = 1160$  L mol<sup>-1</sup> cm<sup>-1</sup> displayed spectrum is normalized to conform to these values. [2] Footnotes:

The extinction coefficient was measured by W. Roebke, M. Renz, and A. Henglein, Int. J. Radiat. Phys. Chem. 1(1): 39-44 (1969). They reported the maximum of the SO<sub>4</sub> spectrum to be at λ(max) = 460 nm. Earlier values of extinction coefficients were ε<sub>455</sub> = (450 ± 45) L mol<sup>-1</sup> cm<sup>-1</sup> from E. Hayon and J.J. McGarvey, J. Phys. Chem. 71(5): 1472-7 (1967) and ε<sub>456</sub> = (460 ± 25) L mol<sup>-1</sup> cm<sup>-1</sup> from L. Dogliotti and E. Hayon, J. Phys. Chem. 71(8): 2511-6 (1967).

Using  $\epsilon_{500}((SCN)_2^-) = 7600 \text{ L mol}^{-1} \text{ cm}^{-1}$  and assuming  $G = G(\epsilon(aq)^-) + G(OH) = 5.5$ , E. Hayon, J.Chem. Phys. 51(11): 4881-92 (1969). In these last two references, A(max) was quoted at 455 nm. [3]





System: aqueous solution of 2 x 10<sup>-3</sup> mol L<sup>-1</sup> HSO<sub>5</sub>, N<sub>2</sub>O-saturated, pH  $\approx$  6 Transient [1]: SO5, c265 \* 560 L mol" cm-1[2]

Reference: E. Hayon, A. Treinin, and J. Wilf, J. Am. Chem. Soc. 94(1): 47-57 (1972)

pulse radiolysis, using Febetron 705 System (Field Emission Corp.); pulse energy 2.3 MeV; pulse length \* 30 ns; dosimetry - N20-saturated, aqueous solution of KSCN [3] Transient generation:

monochromators, used in series to reduce scattered light [4]; two EMI 9558 QB PMT; dual beam Fairchild 777 oscilloscope; time delay 200  $\mu s$  after pulse Osram XBO 450 W xenon lamp (pulsed); two high intensity Bausch & Lomb Spectral acquisition:

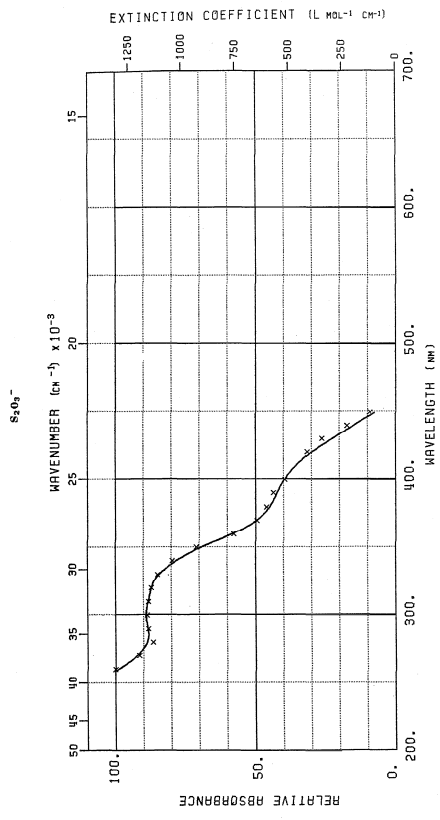
Transient formation:

 $e(aq)^- + N_2O (+H_2O) \rightarrow N_2 + OH^- + OH$ 

 $0H + HSO_5^{-} \rightarrow 0H^{-} + HSO_6 \text{ (or } SO_5^{-})$ 

[1] Points in spectrum are averages of three of more experiments. [2] Quoted extinction coefficient is a lower limit. Footnotes:

Impurities, sulfates,  $H_2O_2$ , and  $S_2O_6^{2-}$ , in the prepared salts KHSO<sub>5</sub> and NaHSO<sub>4</sub> made c uncertain. Earlier values of  $\varepsilon$  (SO<sub>5</sub>) and the shape of earlier spectra were somewhat different, W. Roekke, M. Renz, and A. Henglein, Int. J. Radiat. Phys. Chem. 1(1): 39-44 (1969). Using  $\varepsilon_{500}$  (SCN)<sub>2</sub> = 7600 L mol<sup>-1</sup> cm<sup>-1</sup> for  $[0^{-1}$  mol L<sup>-1</sup> SCN<sup>-</sup>,  $N_2O_{-8}$  saturated solutions at pH 5.5 and assuming  $G = G(\varepsilon(aq)^-) + G(OH) = 5.5$ , [3]



System: aqueous solution of  $10^{-4}$  mol L<sup>-1</sup>  $S_2 O_3^{2-}$ , saturated with  $N_2 O_4$  pH = 4.8

Translent [1]: S<sub>2</sub>O<sub>3</sub> [2]

Reference: M. Schoenseshoefer, Int. J. Radist. Phys. Chem. 5: 375-86 (1973)

pulse radiolysis [3] using Van de Graaff generator; energy 1.5 MeV; dose rate during the pulse was 700 rad per #s Transient generation:

Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 PMT; decay curves photographed from oscilloscope display; time delayimmediately after pulse Spectral acquisition:

Transient formation:

 $N_2O + e(aq)^- (+H_2O) \rightarrow N_2 + OH + OH^-$ 

 $OH + S_2 O_3^{2-} \rightarrow S_2 O_3^{-} + OH^{-}$ 

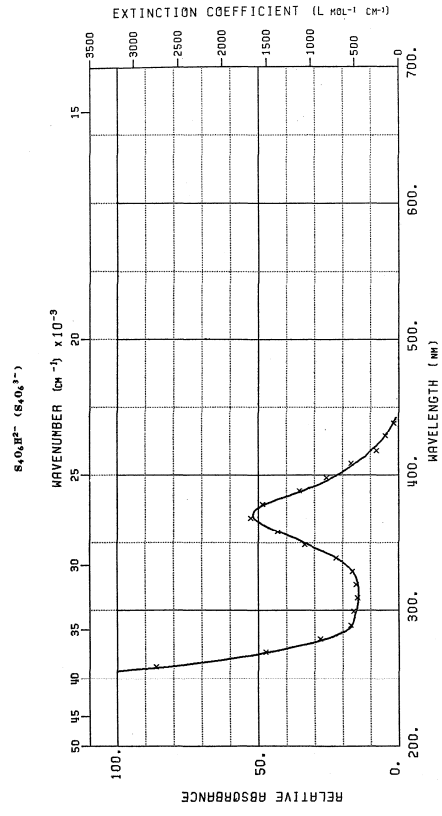
Footnotes:

[1] Spectrum corrected for  $S_203^- + S_203^{2-} + S_406^{3-}$  equilibrium.

Extinction coefficients were based on G(OH) = 6.

[2] Another spectrum has been assigned as the  $S_203^-$  spectrum: L. Dogliotti and E. Hayon, J. Phys. Chem. 72: 1890-7 (1968); G.E. Adams, J.W. Boag, and B.D. Michael, Proc. Chem. Soc. 411 (1964); G.E. Adams, J.W. Boag, and B.D. Michael, Trans. Faraday Soc. 61: 1674-80 (1965).

[3] A. Henglein, Allg. Prakt. Chem. 17(5): 295-391 (1966).



Transient:  $S_40_6H^{2-}$  [1]  $(S_40_6^{3-}$  [2]),  $\lambda(max) = 370$  nm,  $\epsilon_{370} = 1650$  L mol<sup>-1</sup> cm<sup>-1</sup> [3],  $2k = 1.1 \times 10^9$  L mol<sup>-1</sup> s<sup>-1</sup>

System: aqueous solution of 10-4 mol L-1 S2032", saturated with N20, pH = 4.8

## S406H2- (S4063-)

M. Schoeneshoefer, Int. J. Radiat. Phys. Chem. 5: 375-86 (1973)

pulse radiolysis [4] using Van de Graaff generator; energy 1.5 MeV; dose rate during the pulse was 700 rad per  $\mu$ s Transient generation:

Spectral acquisition: Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; ΕΜΙ 9558
PMΓ; decay curves photographed from oscilloscope display; time delay-spectrum measured after 25 μs and extrapolated to time zero

Transient formation [5]:

$$N_2O + e(aq)^- (+H_2O) \rightarrow N_2 + OH + OH^-$$

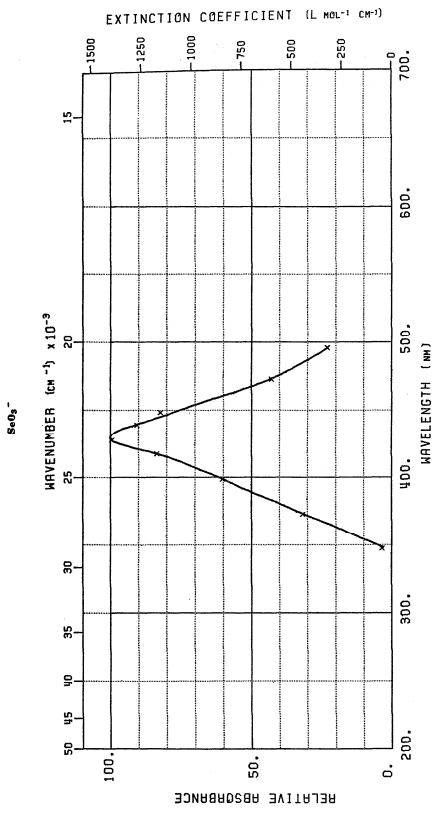
$$0H + S_2 0_3^2 \rightarrow S_2 0_3^2 + 0H^2$$

$$S_2O_3^- + S_2O_3^{2-} \rightarrow S_4O_6^{3-}$$

Similar spectra have been attributed to the S<sub>2</sub>0<sub>3</sub> radical: L. Dogliotti and E. Hayon, J. Phys. Chem. 72: 1800-7 (1968); G.E. Adams, J.W. Boag, and B.D. Michael, Proc. Chem. Soc. 411 (1964); G.E. Adams, J.W. Boag, and B.D. Michael, Trans. Faraday. Soc. 61: 1674-80 (1965). £13 Footnotes:

- Spectrum is almost independent of pH.
  - Using G(OH) = 6.
- A. Henglein, Allg. Prakt. Chem. 17(5): 295-301 (1966). Almost identical spectrum was obtained in an alkaline, aqueous solution of S406<sup>2</sup> containing tert-butanol. 2522

$$e(aq)^- + S_4O_6^{2-} \rightarrow S_4O_6^{3-}$$



System: aqueous solution of 10-2 mol L-1 Se032-, Ar-saturated, pH natural Transient: Se03, \(\)(max) = 430 nm, \(\)e430 = 1.4 x 103 L mol<sup>-1</sup> cm<sup>-1</sup> [1]

(1978) Reference: M.S. Subhani and T. Kausar, Rev. Roum. Chim. 23(11-12): 1627-

Transient generation: flash photolysis, using flash tubes [2]

monitoring lamp; monochromator; PMT; oscilloscope [3] Spectral acquisition:

Transient formation:

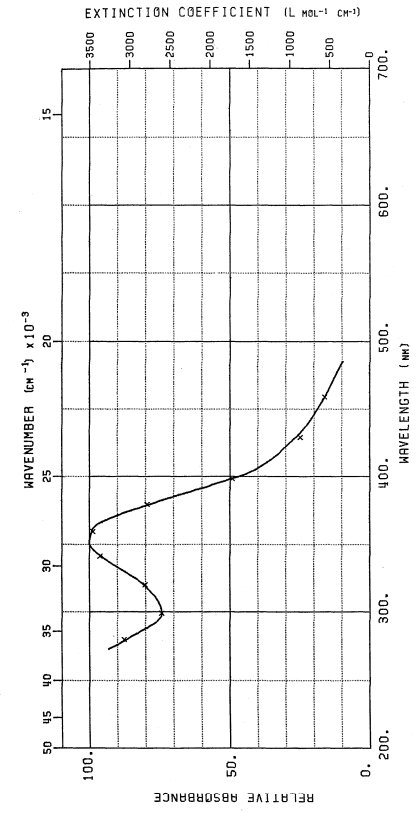
 $SeO_3^{2-}$  H<sub>2</sub>0 + h $\nu$  +  $SeO_3^{2-}$  H<sub>2</sub>0\*

 $SeO_3^2 - H_2O^* \rightarrow SeO_3^- + e(aq)^- (+H_2O)$ 

Footnotes: [1] Extinction coefficient obtained by assuming amount of 02, formed from e(aq) in 02-saturated solutions, is equal to that of Se0;.
[2] Apparatus similar to that in G.V. Buxton and M.S. Subhani, J. Ghem. Soc. Trans. 1 68(5): 958-69 (1972).
[3] Apparatus similar to that in K.W. Chambers, E. Collinson, F.S. Dainton, W.A. Seddon, and F. Wilkinson, Trans. Faraday Soc. 63: 1699-711 (1967).

raday





System: aqueous solution of 2 x 10-2 mol L-1 TeO32-, air-saturated, pH neutral Transient:  $TeO_3^-$ ,  $\lambda(max) = 350$  nm, c = 3500 L mol<sup>-1</sup> cm<sup>-1</sup>

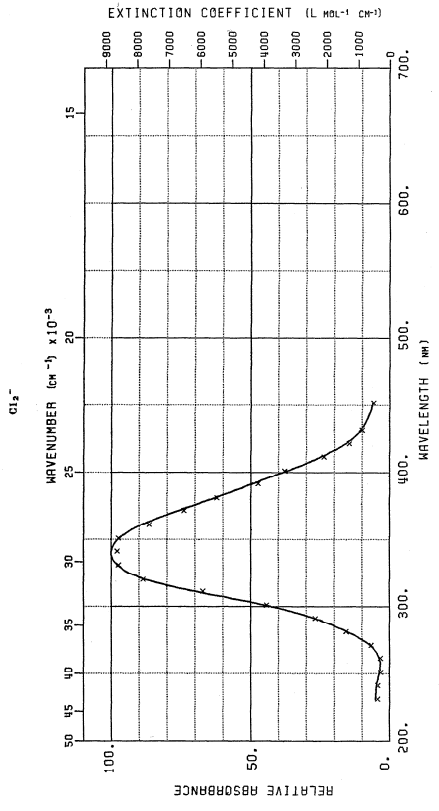
Reference: G.E. Adams, J.W. Boag, and B.D. Michael, Trans. Faraday Soc. 61: 1674-80 (1965)

Transient generation: pulse radiolysis, linear accelerator; energy 1.8 MeV; dosimetry, using ferrocyanide

450 W xenon monitoring lamp; Bausch & Lomb type 250 grating monochromator; EMI 9558 QC PMT; 545 Tektronix oscilloscope with 53/54L preamplifier Spectral acquisition:

Translent formation:

 $0H + Te0_3^{2-} \rightarrow Te0_3^{-} + 0H^{-}$ 



System: aqueous solution of 10<sup>-2</sup> mol L<sup>-1</sup> Gl<sup>-</sup>, Ar saturated, 10<sup>-1</sup> mol L<sup>-1</sup> perchloric acid, pH  $\approx$  1 Transient:  $Cl_2^-$ ,  $\lambda(max) = 340$  nm,  $\epsilon_{340} = (8800 \pm 500)$  L mol<sup>-1</sup> cm<sup>-1</sup> [1,2]

Reference: G.G. Jayson, B.J. Parsons, and A.J. Swallow, J. Chem. Soc., Faraday Trans. 1 69: 1597-607 (1973)

Transient generation: pulse radiolysis, electron linear accelerator; dose per pulse 1.58 x 103 rad; dosimetry-secondary emission chamber [3]

two Bausch & Lomb high-intensity monochromators used in series to reduce scattered light. Spectral acquisition:

Transient formation:

.HOID + CI + HO

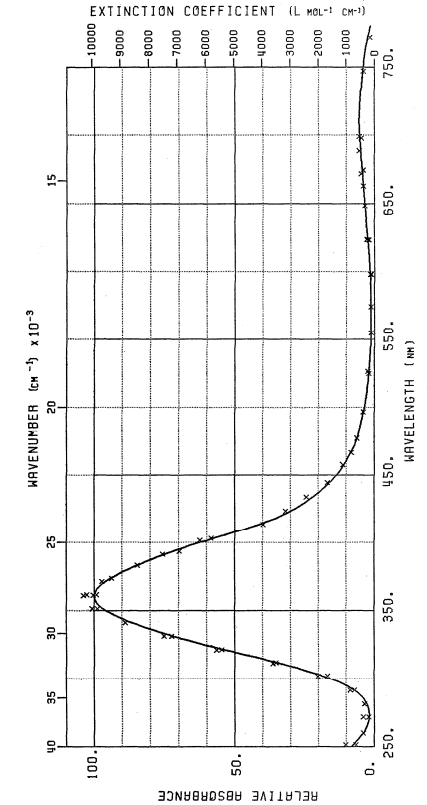
 $C10H^- + H^+ \rightarrow H_2O + C1$ 

CI + CI + CI2-

[1] An earlier value was c340 = 12,500 L mol<sup>-1</sup> cm<sup>-1</sup> from M. Anbar and J.K. Thomas, J. Phys. Chem. 68: 3829-35 (1964) and a later value was c = 12,400 L mol<sup>-1</sup> cm<sup>-1</sup> from R.K. Broszklewicz, Bull. Acad. Pol. Sci., Ser. Sci. Chim. 24(2): 123-31 (1976). [2] Assuming G(Gl<sub>2</sub>) = 2.9. Footnotes:

Calibrated by aqueous potassium thiocyanate taking  $G((SCII)_2^-) = 2.9$  and  $\epsilon_{500}((SCI)_2^-) = 7100$  L mol<sup>-1</sup>.





EXTINCTION COEFFICIENT

Transient:  $Br_2^-$ ,  $\lambda(max) = 360 \text{ nm}$ ,  $\epsilon_{360} = (9900 \pm 600) \text{ L mol}^{-1} \text{ cm}^{-1} [1,2]$ System: aqueous solution of 10-3 mol L-1 Br-, neutral, N20-saturated

Reference: D. Zehavi and J. Rabani, J. Phys. Chem. 76(3): 312-9 (1972)

pulse radiolysis; energy 5 MeV; pulse length 0.1 µs; doslmetry-induction coll [3] Transient generation:

split analyzing light beam: with two monochromators; extrapolated to zero time; PMT's type. R166 for 260-300 nm, 1P28 for 300-650 nm and R196 above 650 nm; dual-beam oscilloscope; filter used to prevent photolysis and second-order light; 150 W xenon lamp Spectral acquisition:

Transient formation:

 $N_2O + e(aq)^- (+H_2O) \rightarrow OH + OH^+ +$ 

OH + Br + BrOH"

BrOH - Br + OH

Br + Br + Br2

 $BrOH^- + Br^- \rightarrow Br_2^- + OH$ 

Footnotes:

other values for ease in the literature are lower. For example 9600 L mol<sup>-1</sup> cm<sup>-1</sup> [4], 8200 L mol<sup>-1</sup> cm<sup>-1</sup> [5], 8300 L mol<sup>-1</sup> cm<sup>-1</sup> [6], and 8560 L mol<sup>-1</sup> cm<sup>-1</sup> [7]. In addition ease = 7800 L mol<sup>-1</sup> cm<sup>-1</sup> [8]. D. Behar, private communication, 1980. Value quoted by authors was (12,000  $\pm$  1000) L mol  $^{-1}$  . However, 3

Internal desimpter using Br. absorption was also used.

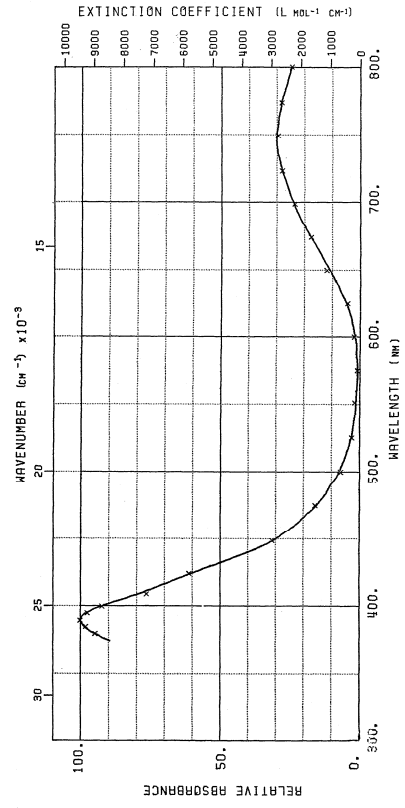
B. Gercek, M. Ebert, C.W. Gilbert, and A.J. Swallow, Pulse Radiolysis, eds. M. Ebert, J.P. Keene, A.J. Swallow, and J.H. Baxendale (Academic Press, 1965) p. 83. [3]

H.C. Sutton, G.E. Adams, J.W. Bong, and B.D. Michael, ibid., p. 61.

[2] [3] [4]

B. Cercek, Int. J. Radiat. Phys. Chem. 4(1): 25-30 (1972).
R.K. Broszkiewicz, Bull. Acad. Pol. Sci. Chim. 24(2): 123-31 (1976).
M.S. Matheson, W.A. Mulac, J.L. Weeks, J. Rabani, J. Phys. Chem. 70: 2092-9 (1966).





System: aqueous solution of 3.3 x 10"4 mol L-1 i, pH neutral, oxygen-saturated Transient [1]:  $I_2^-$ ,  $\epsilon$ (max)  $\approx 9400 \text{ L mol}^{-1} \text{ cm}^{-1} [2,3,4]$ 

Reference: R. Devonshire and J.J. Weiss, J. Phys. Chem. 72(11): 3815-20 (1968)

photolysis using the 185 nm mercury line from a low-pressure Spectrosil mercury lamp: photolyzing light intensity was modulated at 50 Hz by using a leakage reactance transformer giving 120 mA and 1650 V; pure Na was passed between the lamp and absorption cell Transient generation:

monitoring lamp was 150 W quartz tungsten-iodine lamp using a 24 V battery; absorption cell was a 50 cm Spectrosil tube of 0.5 cm inside diameter; was detected using phase sensitive technique using electronic components from Brookdeal Electronics Ltd. - amplifier (LA635), phase sensitive detector (PD629), phase shifter (PS946), and meter (MU947). heat filter was used in monitoring beam [5]; the output from PMT Spectral acquisition:

Translent formation:

I- + hv + (I-)\*

-e + I + \*(-I)

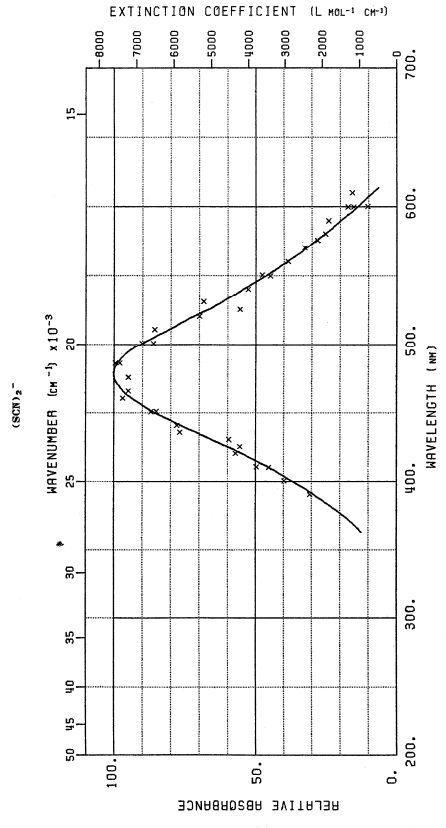
I + I + I2

[1] At wavelengths shorter than about 525 nm, the spectrum has been corrected for the build-up of 13 from the photolysis. Footnotes:

Listed extinction coefficient of 9400 L mol-1 cm 1 is from a spectrum where the maximum In contrast, the literature spectrum of  $1_2$ , after being digitized from the journal and smoothed, shows a maximum near 390 nm. For the purpose of this compilation, the literature spectrum is normalized to  $\epsilon(\max) = 9400$  L mol<sup>-1</sup> cm<sup>-1</sup>; but the original wavelength scale is used, so that the displayed spectrum has  $\lambda(\max) \approx 390$  nm. extinction was near 380 nm. Private communication, R.H. Schuler, 1979. [2]

3] In descrated aqueous solution L.I. Grossweiner and M.S. Matheson, J. Phys. Chem. 61: 1069-95 (1957), measured c<sub>395</sub> = (15,600 ± 3000) L mol<sup>-1</sup> cm<sup>-1</sup>. 4] Ratio of absorbance at 390 nm to absorbance at 750 nm = 3.5.

To further guard against large temperature fluctuations, the solutions were only irradiated for 30 s. [4]



System: aqueous 10-4 mol L-1 KSCN (freshly prepared), N2-saturated, pH 6.0 Transient:  $(SCN)_2^-$ ,  $\lambda(max) = 480$  nm [1],  $\epsilon_{475} = 7600$  L mol<sup>-1</sup> cm<sup>-1</sup> [2,3,4]

## (SCN)2

Reference: L. Dogliotti and E. Hayon, J. Phys. Chem. 72: 1800-1867 (1968)

Transient generation: flash photolysis; energy per pulse 1800 J; duration at "1/e time" of 5 µs; pulse length 70 µs Bausch 8 Lomb grating monochromator; EMI 95580B photomultiplier; spectra were obtained by a point-by-point method; time delay =  $80~\mu s$  after start of pulse Spectral acquisition:

Transfent formation:

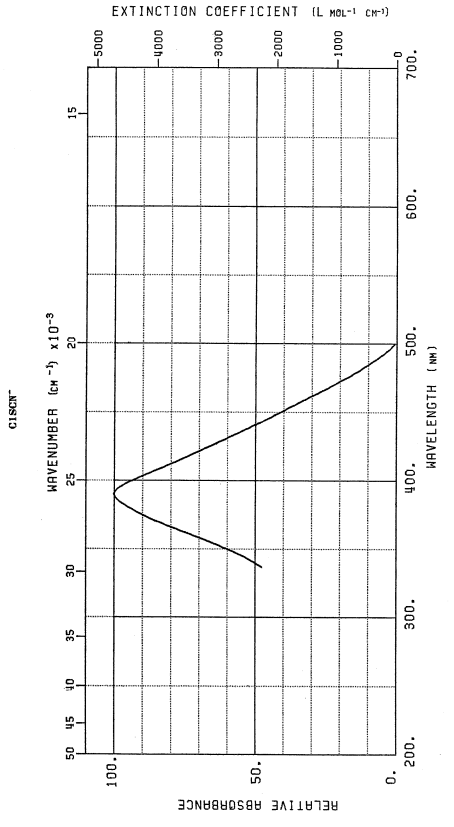
 $SGN^-H_2O + h\nu + SGN + e(aq)^- + H_2O$ 

SCN + SCN + (SCN)2

[1] A value of  $\lambda(\max) = 485$  nm was quoted by the authors. However, the digitization of spectrum showed that they apparently meant the maximum of the data points. The maximum of their smoothed curve was 478 nm. For the purpose of this compilation, 478 is rounded to 480 nm in the quoted maximum, but 478 nm is the maximum of the displayed spectrum. Footnotes:

[2] J.H. Baxendale, P.L.T. Bevan, and D.A. Stott, Trans. Faraday Soc. 64: 2389-97 (1968).

[3] The spectrum displayed has been normalized to correspond to  $\epsilon(\max) = 7600 \text{ L mol}^{-1}$  cm<sup>-1</sup>. [4] A recent spectrum of (SCN)<sub>2</sub> has been taken and shows  $\lambda(\max) = 472$  nm and  $\epsilon(\max) = 7580 \text{ L mol}^{-1}$  cm<sup>-1</sup>. The  $G((SCN)_2^-)$  was measured to be 6.13 in  $10^{-2}$  mol  $L^{-1}$  SCN<sup>-</sup> aqueous solutions, saturated with  $N_2O$ . R.H. Schuler, L.K. Patterson, and E. Janata, J. Phys. Chem. 84(16): 2088-9 (1980).



Transient [1]: CISCN,  $\lambda$ (max) = 390 nm,  $\epsilon_{390}$  = (4700 ± 900) L mol<sup>-1</sup> cm<sup>-1</sup> [2] System: aqueous solution of 3.8 mol L-1 Cl and 5 x 10-6 mol L-1 SCN-

Reference: M. Schoeneshoefer, Int. J. Radint. Phys. Chem. 1: 505-13 (1969)

pulse radiolysis [3] using Van de Grasff generator; energy 1.7 MeV Translent generation:

Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 PMT; decay curves photographed from oscilloscope display; time delay = 20 µs [4] Spectral acquisition:

Transient formation:

.HO I → ..IO + HO

C10H + C1 + 0H CI + CI + CI2  $C1_2$  + SCN  $\rightarrow$  2C1 + SCN

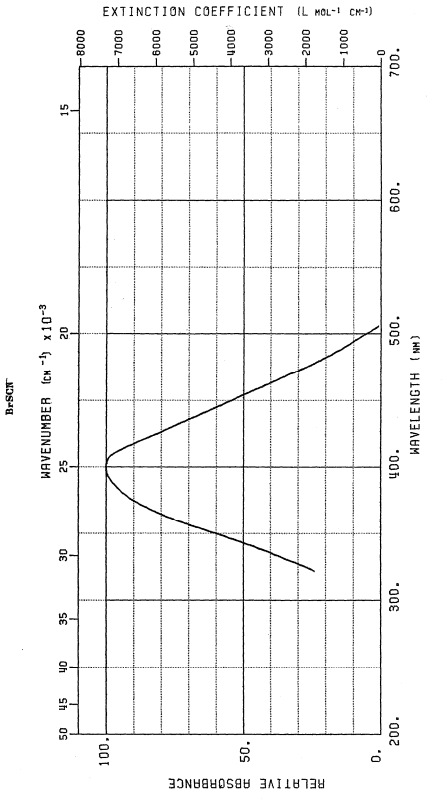
SCN + SCN - + (SCN)2-

 $(SCN)_2$  + Cl + Cl + Cl SCN + Cl SCN , K = 3.3 x 10-3

11] Obtained from a mixed spectrum due to (SGN)2" and CISCN". Since CISCN-absorbes very little at 520 nm, the contribution of (SGN)2" which does absorb at 520 nm could be determined.

[2] Based on \$\epsilon \text{520} = 5300 \text{ L mol}^{-1} \text{ cm}^{-1} \text{ for } (SCN)2^{-}.

[3] A. Henglein, Allg. Prakt. Chem. 17(5): 295-301 (1966). Footnotes:



System: aqueous solution of 2 x 10-2 mol L-1 Br and 1 x 10-4 mol L-1 SCN-Transient [1]: BrSCN<sup>-</sup>, λ(max) = 400 nm, ε<sub>400</sub> = 7300 L mol<sup>-1</sup> cm<sup>-1</sup> [2]

M. Schoeneshoefer and A. Henglein, Ber. Bunsenges. Phys. Chem. 73(3): 289-93 (1969) Reference:

Transient generation: pulse radiolysis [3] using Van de Graaff generator; energy 1.7 MeV

Osram XBO 450 monitoring lamp; Zeiss M4Q3 monochromator; EMI 9558 PMF; decay curves photographed from oscilloscope display; time delay greater than 10 µs [4] Spectral acquisition:

Transient formation:

Br + 0H + Br0H

BrOH" \* Br + OH" Br + Br & Br2

Br2 + SCN - 2Br + SCN

SCN + SCN + (SCN)2-

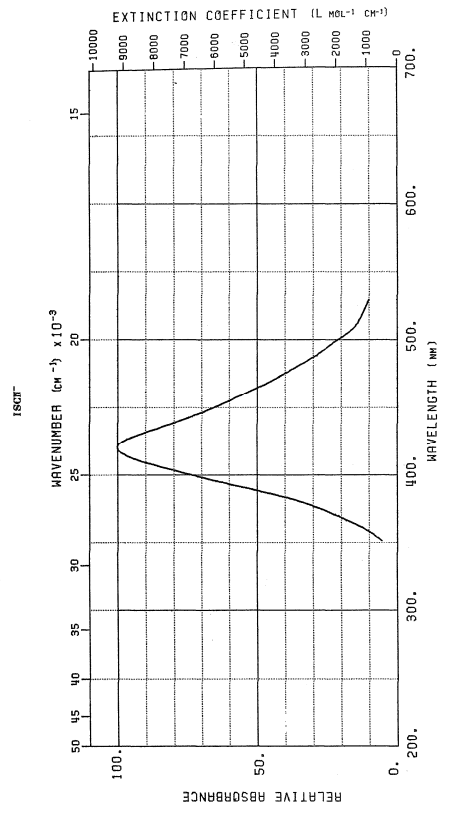
 $Br_2 + SCN + BrSCN + Br', K = 10^3$ 

SCN + Br + SCNBr

 $(SCN)_2$  + Br + SCN + BrSCN , K = 9.1 x 10-3

[1] Obtained from a mixed spectrum due to (SCN)2 and BrSCN. Since BrSCN absorbes very little at 520 nm, the contribution of (SCN)2 which does absorb at 520 nm could be determined. Footnotes:

Based on e529 = 5300 L mol-1 cm<sup>-1</sup> for (SCN)<sub>2</sub>. A. Henglein, Allg. Prakt. Chem. 17(5): 295-301 (1966). For ratios of [Br<sup>-</sup>]/[SCN<sup>-</sup>]  $\leq$  200, it was found that there was very little Br<sub>2</sub> present. 252



System: aqueous solution of 1 mol L-1 SCN- and 10-4 mol L-1 I-, saturated with N20 Transient [1,2]: ISCN,  $\lambda(\max) = 420$  nm,  $\epsilon_{429} = 9200$  L mol<sup>-1</sup> cm<sup>-1</sup>

Reference: M. Schoeneshoefer and A. Henglein, Ber. Bunsenges. Phys. Chem. 74(4): 393-8 (1970)

pulse radiolysis using Van de Graaff generator; energy 1.7 MeV Transient generation: Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 PMT; decay curves photographed from oscilloscope display; time delay 20 µs Spectral acquisition:

Transient formation:

 $e(aq)^- + N_2O (+H_2O) \rightarrow OH + OH^+ + N_2$ 

OH + SCN - HOSCN

HOSCN + OH + SCN

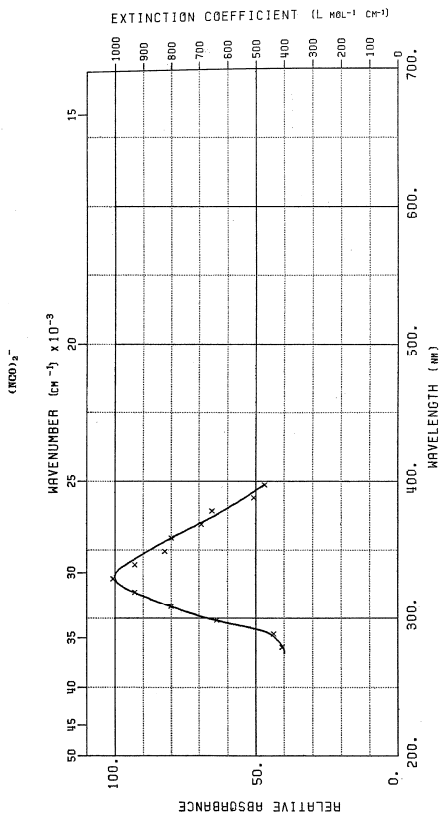
SCN + SCN + (SCN)2

 $(SCN)_2 + I^* + ISCN^+ + SCN^-, K = 400$ 

 $ISCN^{-} + I^{-} + I_{2}^{-} + SCN^{-}, K = 55$ 

does not contribute much because it is present in small quantities under the experimental conditions. Using the measured rate constants, [(SCN)2]: [ISCN]: [I2] = 25:1:0.0055.  $I_2$  and ISCN absorb very little at 550 mm, the contribution of (SCN)2 could be subtracted using the dose and yield of OH radicals of G=5.5[1] Obtained from a mixed spectrum due to (SCN)2, 12 and ISCN. Footnotes:

in this case much more absorption was evident near 350 nm. Since the ratio of  $[(SCN)_2^-]$ :  $[ISCN^-]$ :  $[I_2^-]$  = 2.5 : 1 : 0.055 in this latter experiment, the additional absorption at 350 nm may have been due to  $I_2^-$  instead of ISCN. Since the ratio of  $[(SGN)_2^{-1}]$ :  $[ISGN^{-1}] = 25$ , the shape of the spectrum is not very well established. Another spectrum was given in the reference for the initial conditions  $[SGN^{-1}] = 0.1 \text{ mol } L^{-1}$  and  $[I^{-1}] = 10^{-4} \text{ mol } L^{-1}$ . 2



System: aqueous solution of  $10^{-1}$  mol L<sup>-1</sup> NCO<sup>-</sup>, N<sub>2</sub>O-saturated, pH = 5.0  $\pm$  0.2 [2] Transient:  $(NCO)_2^-$ ,  $\lambda(max) = 330 \text{ nm}$ ,  $\epsilon_{330} = (1000 \pm 100) \text{ L mol}^{-1} \text{ cm}^{-1}$  [1]

Reference: M. Schoeneshoefer and A. Henglein, Ber. Bunsenges. Phys. Chem. 74(4): 393-8 (1970)

Transient generation: pulse radiolysis using Van de Graaff generator; energy 1.7 MeV

Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 PMT; decay curves photographed from oscilloscope display; time delay 20 µs Spectral acquisition:

Transient formation:

 $e(aq)^- + N_2O (+H_2O) \rightarrow OH + OH^- + N_2$ 

OH + SCN - HOSCN

HOSCN → OH + SCN

SCN + SCN + (SCN)2

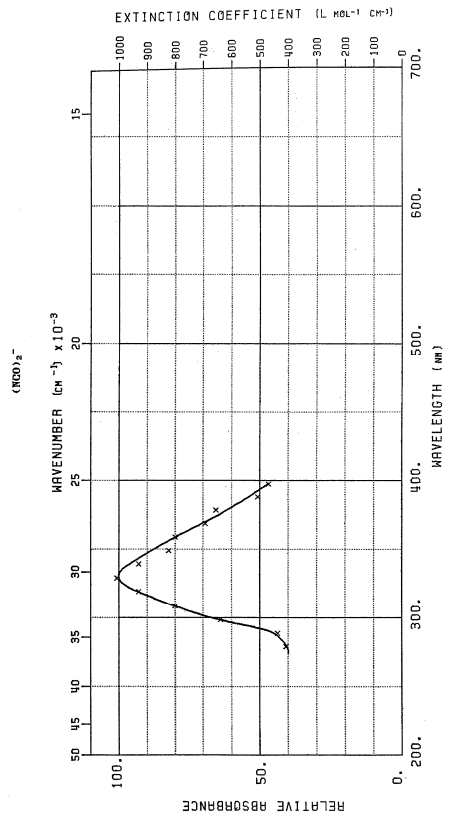
 $(SCN)_2$  + I + ISCN + SCN", K = 400

 $ISCN^{-} + I^{-} + I_{2}^{-} + SCN^{-}, K = 55$ 

does not contribute much because it is present in small quantities under the  $\rm I_2$  and ISCN absorb very little at 550 nm, the contribution of (SCN)<sub>2</sub> could be subtracted using the dose and yield of OH radicals of G = 5.5 [1] Obtained from a mixed spectrum due to (SCN)2, 12 and ISCN. Footnotes:

experimental conditions. Using the measured rate constants, [(SGN)<sub>2</sub>]: [ISCN]: [I<sub>2</sub>] = 25: 1: 0.0055.

Since the ratio of [(SGN)<sub>2</sub>]: [ISCN] = 25, the shape of the spectrum is not very well established. Another spectrum was given in the reference for the initial conditions [SGN] = 0.1 mol L<sup>-1</sup> and [I] = 10<sup>-4</sup> mol L<sup>-1</sup>. However, in this case much more absorption was evident near 350 nm. Since the ratio of [(SGN)<sub>2</sub>]: [ISCN] = 2.5: 1: 0.055 in this latter experiment, the additional absorption at 350 nm may have been due to [2] instead of ISCN. [2]



System: aqueous solution of  $10^{-1}$  mol L<sup>-1</sup> NCO<sup>-</sup>, N<sub>2</sub>O-saturated, pH = 5.0  $\pm$  0.2 [2] Transient:  $(NCO)_2^-$ ,  $\lambda(max) = 330$  nm,  $\epsilon_{330} = (1000 \pm 100)$  L mol<sup>-1</sup> cm<sup>-1</sup> [1]

Reference: J.G. Leopold and M. Faraggi, J. Phys. Chem. 81(8): 803-6 (1977)

Transient generation: pulse radiolysis [3]; energy 5 MeV; dosimetry-electron coil [4]

150 W xenon lamp; Bausch & Lomb high-intensity monochromator; 1P28A or R136 PMT; Type 556 Tektronix double beam oscilloscope with Polaroid camera Spectral acquisition:

Transient formation:

 $N_2O + e(sq)^- (+H_2O) \rightarrow N_2 + OH + OH^-$ 

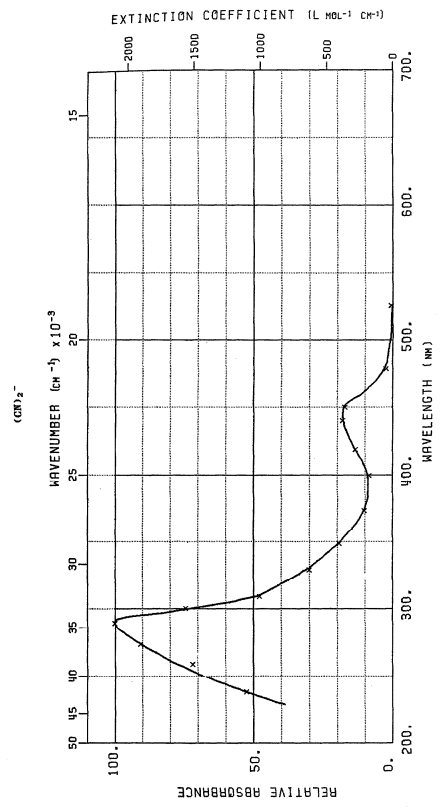
NCO + OH → HNCOO"

HNCO0- + H+ → HNCOOH

 $HNCOOH + NCO^- + (NCO)_2^- + H_2O$ 

Footnotes:

, using [1] Based on G(OH) = 5.6 for  $N_2D$ -saturated solutions. [2] Dose was such that [OH] = 9.7 x 10<sup>-6</sup> mol L<sup>-1</sup>. [3] M. Faraggi and Y. Tendler, J. Chem. Phys. 56(7): 3287-93 (1972). [4] Dosimetry used N<sub>2</sub>O saturated solutions of 10<sup>-2</sup> mol L<sup>-1</sup> Fe(GN)<sub>6</sub><sup>4-</sup>, G(Fe(GN)<sub>6</sub><sup>3-</sup>) = 6.1



Transient:  $(CN)_2$ , cyanegen radical ion,  $\lambda(\max) = 290$  nm,  $\epsilon_{290} = 2160$  L mol<sup>-1</sup> cm<sup>-1</sup>, and  $\lambda(\max) = 440$  nm,  $\epsilon_{440} = 360$  L mol<sup>-1</sup> cm<sup>-1</sup> [1]

System: aqueous golution of 3.7 x 10 tmol L-1 (CN)2, He-purged, at natural pH & 6

Reference: I.G. Draganic, Z.D. Draganic, and R.A. Holroyd, J. Phys. Chem. 75(5): 608-12 (1971)

pulse radiolysis using Febetron 705 system; energy 2 MeV; pulse length < 0.1  $\mu s$ ; dos- 4.72 krad/pulse; two dosimetry methods were used [2]; syringe technique Transient generation:

450 W xenon analyzing lamp with an arc booster [3]; path length 4.0 cm; high-intensity Bausch 8 Lomb grating monochromators; Amperex XP 1003 PMT; Tektronix 454 escilloscope Spectral acquisition:

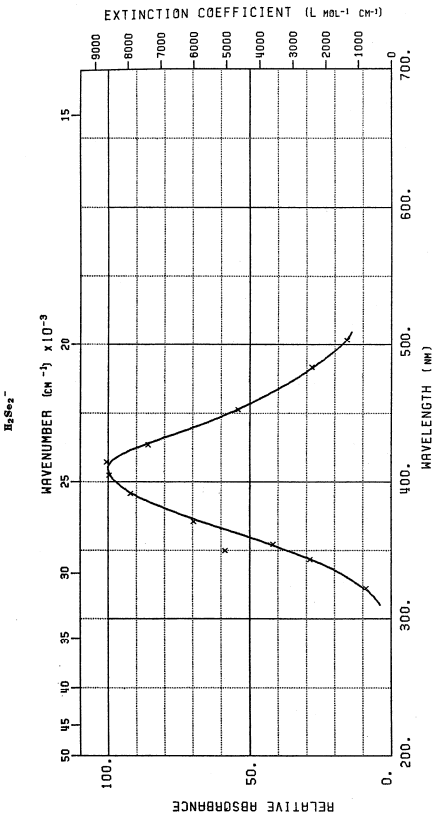
Transient formation [4]:

 $(CN)_2 + e(aq)^{\sim} \rightarrow (CN)_2^{\sim}$ 

[1] Corrected for OH absorption. [2] Footnotes:

The first method used N20-saturated aqueous solutions of 0.1 mol L<sup>-1</sup> thiocyanate using  $\varepsilon_{500}$  = 7600 L mol<sup>-1</sup> cm<sup>-1</sup> for (SCN)<sub>2</sub>. The second method used N<sub>2</sub>0-saturated aqueous solutions of  $10^{-3}$  mol L<sup>-1</sup> ferrocyanide containing a small amount of air. In this system  $\varepsilon_{420}$  = 1000 L mol<sup>-1</sup> cm<sup>-1</sup> for the ferricyanide formed. Both methods assumed G(product) = 5.4.

Increased lamp intensity in ultraviolet by 60-fold. The reaction of (GN)<sub>2</sub> with OH was too slow (k  $\leq$  10<sup>7</sup> L mol<sup>-1</sup> s<sup>-1</sup>) for the product to interfere with the spectrum. [3]



System: aqueous solution [2] of 10-3 mol L-1 H2Se, Ar-saturated, pH = 3.7 Transient:  $H_2Se_2^-$ ,  $\lambda(max) = 410$  nm,  $\epsilon_{410} = 8600$  L mol<sup>-1</sup> cm<sup>-1</sup> [1]

Reference: M. Schoeneshoefer, W. Karmann, and A. Henglein, Int. J. Radiat. Phys. Chem. 1: 407-23 (1969)

pulse radiolysis, Van de Graaff generator; energy 1.6 MeV; dose rate during the pulse was 700 rad per µs; dosinetry based on SCN<sup>-</sup> [3] Transient generation:

Spectral acquisition [4]: Osram XBO 450 monitoring lamp; Zeiss N403 monochromator; EMI 9558 PMT; decay curves photographed from oscilloscope display; time delay-immediately after pulse

Transient formation:

 $e(aq)^- + H_2Se \rightarrow HSe^- + H$ 

 $H + H_2Se \rightarrow H_2 + HSe$ 

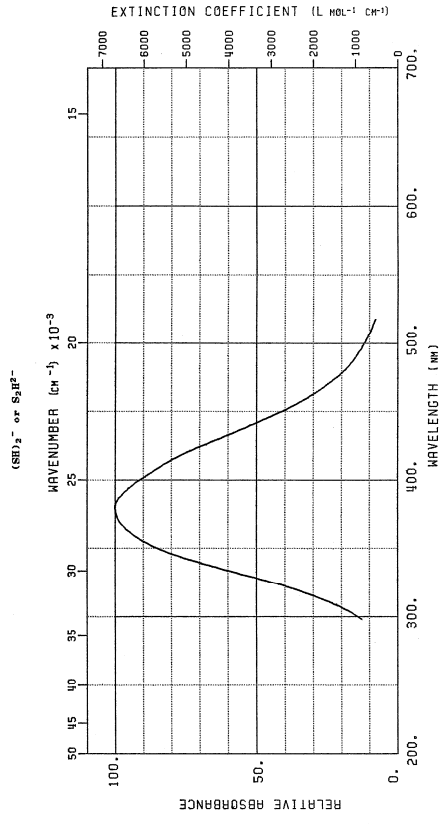
 $0H + HSe^- \rightarrow 0H^- + HSe$ ,  $pK_a(H_2Se) = 3.77$  $OH + H_2Se \rightarrow H_2O + HSe$ 

HSe + HSe + H2Se2

Footnotes:

[1] Based on G(H<sub>2</sub>Se<sub>2</sub><sup>-</sup>) = 6. [2] Oxygen was removed from water before H<sub>2</sub>Se was added to make solution. [3] Based on the measurement of SGN solutions under equivalent conditions, using egoe = 7100 L mol<sup>-1</sup> cm<sup>-1</sup>.

[4] A. Honglein, Ailg. Prakt. Chem. 17(5): 295-301 (1966).



Transient [1]: (SH)<sub>2</sub> or  $S_2H^2$ ,  $\lambda(max) = 380$  nm System: aqueous solution of  $10^{-3}$  mol L<sup>-1</sup> H<sub>2</sub>S, pH = 6.6

(SH)2 or S2H2-

Reference: W. Karmann, G. Meissner, and A. Henglein, Z. Naturforsch. Pt. B 22(3): 273-82 (1967)

Transient generation: pulse radiolysis [2] using Van de Graaff generator; energy 1.7 MeV

Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 PMT; decay ourves photographed from oscilloscope display Spectral acquisition:

Transient formation:

 $e(aq)^{-} + H_2S \rightarrow SH^{-} + H, pK_a(H_2S) = 7$ 

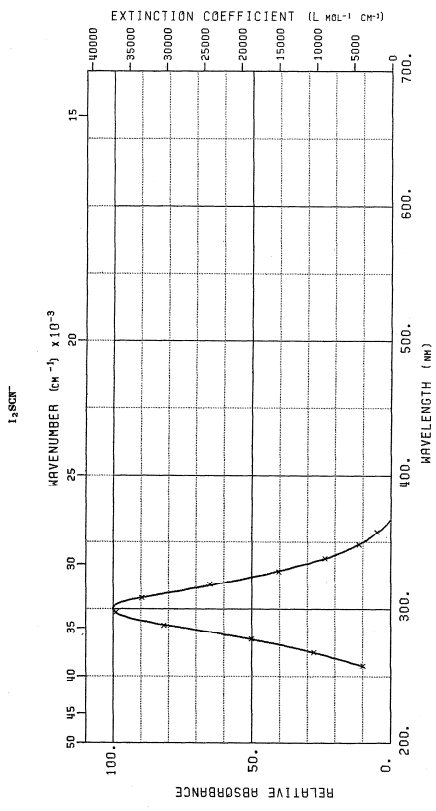
 $H + H_2S \rightarrow H_2 + HS$ 

 $OH + H_2S \rightarrow H_2O + HS$ 

 $HS + HS^- \rightarrow (SH)_2^-$ 

(SH)2 + S2H2 + H+

[1] Extinction coefficient based on G=6. [2] A. Henglein, Allg. Prakt. Chem. 17(5): 295-391 (1966). Footnotes:



System: aqueous solution of 10-1 mol L-1 SCN- and 2 x 10-4 mol L-1 I, saturated with N20 Transient [1]:  $I_2SCN^-$ ,  $\lambda(max) = 300$  nm,  $\epsilon_{300} = 37,000$  L mol<sup>-1</sup> cm<sup>-1</sup> [2]

Reference: M. Schoeneshoefer and A. Henglein, Ber. Bunsenges. Phys. Chem. 74(4): 393-8 (1970)

Transient generation: puise radiolysis using Van de Graaff generator; energy 1.7 MeV

Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 PMT; decay curves photographed from oscilloscope display; time delay 200 µs Spectral acquisition:

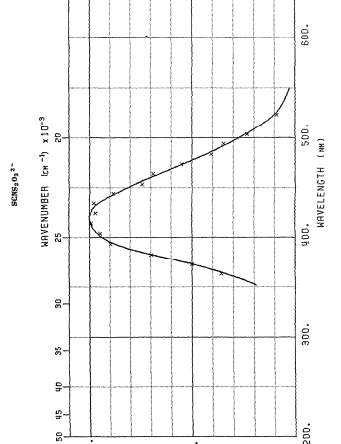
Transient formation:

 $I_2$  + (SCN)<sub>2</sub>  $\rightarrow$   $I_2$ SCN + SCN

 $I_2$  + ISCN  $\rightarrow$   $I_2SCN$  + I

ISCN + ISCN → I2SCN + SCN

[1] Compound is relatively stable in acid solutions.
[2] Assuming G = 2.75 and assuming G for the three complexes, I<sub>2</sub><sup>-</sup>, (SCN)<sub>2</sub><sup>-</sup>, and ISCN<sup>-</sup>, was collectively 5.5. Footnotes:



EXTINCTION COEFFICIENT (L MOL-1 CH-1)

0001

-3000

-2000

-1000

-5000

-6000

-7000

100.

15

Transient [1]:  $S(RS_2\theta_3^{2-}, \lambda(max) = 420 \text{ nm}$ ,  $\varepsilon_{420} = 7400 \text{ L mol}^{-1} \text{ cm}^{-1}$  [2] System: aqueous solution of 5 x 10<sup>-2</sup> mol L<sup>-1</sup> SCN<sup>-</sup> and 4 x 10<sup>-4</sup> mol L<sup>-1</sup> S<sub>2</sub>0<sub>3</sub><sup>2-</sup>, saturated with N<sub>2</sub>0 [3]

50,

RELATIVE ABSORBANCE

200.

0.

## SCNS2032-

M. Schoeneshoefer, Ber. Bunsenges. Phys. Chem. 77(4): 257-62 (1973) Reference:

pulse radiolysis [4] using Van de Graaff generator; energy 1.7 MeV Transient generation:

Spectral acquisition: Osram XBO 450 monitoring lamp; Zeiss M4Q3 monochromator; EMI 9558 PMI; decay curves photographed from oscilloscope display

Transient formation [5]:

 $N_2O + e(aq)^- (+H_2O) \rightarrow N_2 + OH + OH^-$ 

OH + SCN + HOSCN

HOSCN + OH-

SCN + SCN + (SCN)2"

 $(SCN)_2^- + S_2^- O_3^2^- + SCNS_2^- O_3^2^- + SCN^-, K = (1.6 \pm 0.4) \times 10^2$ 

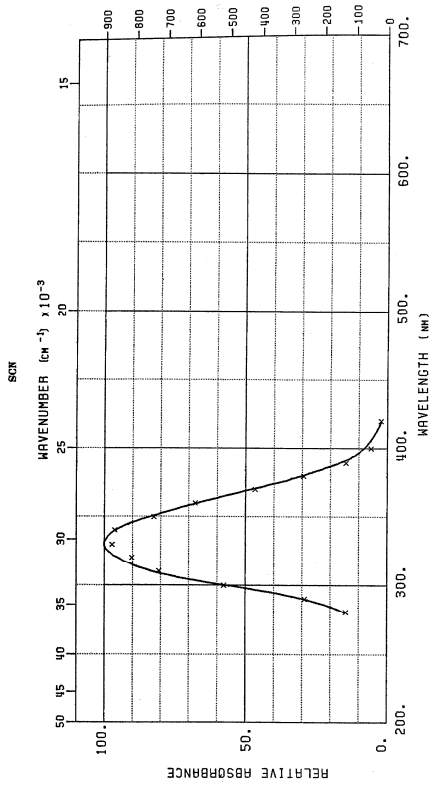
[1] Obtained from a mixed spectrum due to (SCN)<sub>2</sub><sup>-</sup>, S406<sup>3</sup><sup>-</sup>, and SCNS<sub>2</sub>09<sup>2</sup><sup>-</sup>. The SCNS<sub>2</sub>09<sup>2</sup><sup>-</sup> spectrum was obtained from the experimental spectrum using a kinetic analysis.

[2] Based on c = 1600 L mol<sup>-1</sup> cm<sup>-1</sup> for S<sub>4</sub>06<sup>3</sup><sup>-</sup> and 7100 L mol<sup>-1</sup> cm<sup>-1</sup> for the c of (SCN)<sub>2</sub><sup>-</sup>. Footnotes:

[3] G(0H) taken to be 6.

A. Henglein, Allg. Prakt. Chem. 17(5): 295-301 (1966). ScNS<sub>2</sub>03<sup>2-</sup> under Shows only the most prominent pathway for formation of SCNS<sub>2</sub>03<sup>2-</sup> under the experimental conditions. Several other equilibria complicate the analysis. In particular, there are two important mechanisms that lead to formation of S406<sup>3-</sup> starting from SCNS<sub>2</sub>03<sup>2-</sup>. [4]





System: aqueous solution of 10-4 mol L-1 SCN-, saturated with N2O, natural pH Transient [1]: SCN, λ(max) = 330 nm, ε<sub>330</sub> = 900 L mol<sup>-1</sup> cm<sup>-1</sup> [2]

Reference: D. Behar, P.L.T. Bevan, and G. Scholes, J. Phys. Chem. 76(11): 1537-42 (1972)

pulse radiolysis, linear accelerator; pulse length 0.1  $\mu s$ ; doses 0.1 to 1.0 krad Transient generation:

Spectral acquisition: kinetic spectroscopy; time delay 1 µs

Transient formation [3]:

 $e(aq)^{-} + N_2O (+H_2O) \rightarrow N_2 + OH + OH^{-}$ 

OH + SCN - HDSCN

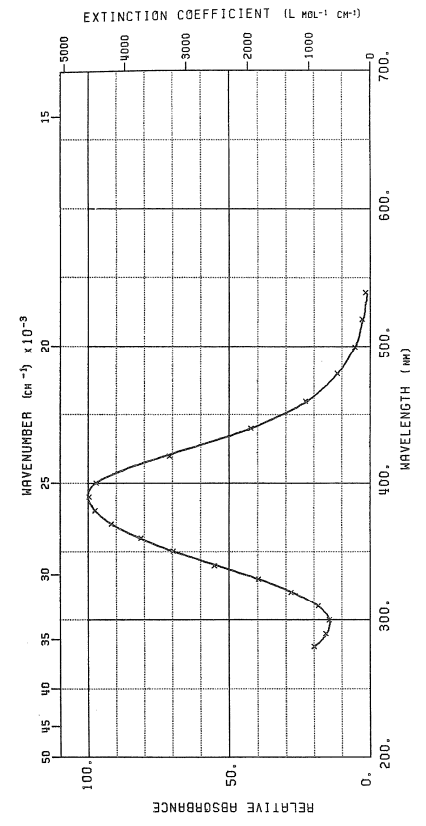
 $HOSCN^{-} + SCN + OH^{-}, K = 3.2 \times 10^{-2} \text{ nol } L^{-1}$ 

SCN + SCN + (SCN)2, K = 2 x 105 L mol-1

[1] Obtained from a composite spectrum by subtracting (SCN)<sub>2</sub> contribution. [2] Based on the kinetic analysis and the value 7600 L mol<sup>-1</sup> cm<sup>-1</sup> for c<sub>475</sub> of (SCN)<sub>2</sub>. Footnotes:

[3] Mechanism involving SCNO2 was eliminated by considering [OH] dependence.





System: aqueous solution of 10-4 mol L-1 SCN" and 10-1 mol L-1 OH-, saturated with N20 Transient [1]: HOSCN", \(\lambda\) = 390 nm, \(\epsilon\) = 4600 L mol^1 cm^1 [2]

## HOSCN-

Reference: D. Behar, P.L.T. Bevan, and G. Scholes, J. Phys. Chem. 76(11): 1537-42 (1972)

Transient generation: pulse radiolysis, linear accelerator; pulse length 0.1 µs; doses 0.1 to 1.0 krad

Spectral acquisition: kinetic spectroscopy; time delay 3.5 µs

Translent formation [3]:

 $e(aq)^- + N_2O (+H_2O) \rightarrow N_2 + OH + OH^-$ 

 $0H + 0H^- + 0^- + H_20$ 

OH + SCN - HOSCN

 $0^- + SCN^- \rightarrow OSCN^{2-}$ 

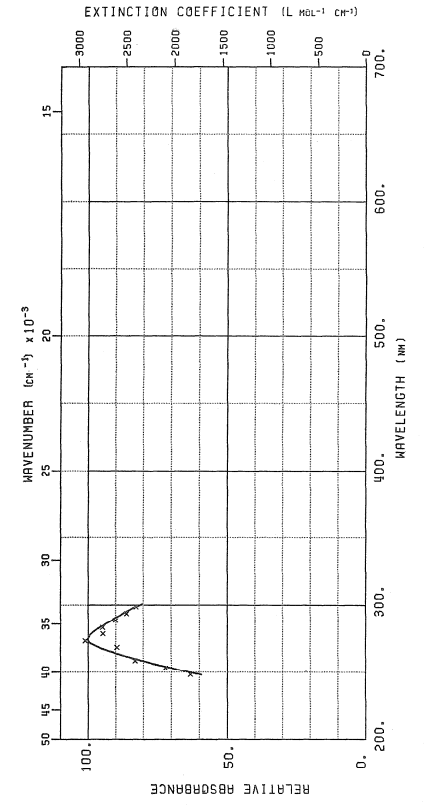
OSCN2 - + H2O + HOSCN + OH

HOSCN + SCN + (SCN)2 + OH

Footnotes:

[1] Obtained from a composite spectrum by subtracting (SGN)<sub>2</sub> contribution. [2] Based on extinction coefficients at 390 nm and 475 nm. [3] The transient OSGN<sup>2</sup> was shown to have a neglible concentration by a study of the dependence of the spectrum on OH concentration.





aqueous solution of 2.5 x 10<sup>-3</sup> mol L<sup>-1</sup> Br<sub>2</sub> and 1.4 mol L<sup>-1</sup> HClO<sub>4</sub>, argon saturated before Br<sub>2</sub> was added Transient [1]: Br [2], bromine atom,  $\lambda(\max) = 275 \text{ nm}$ ,  $\epsilon(\max) = 2900 \text{ L mol}^{-1} \text{ cm}^{-1}$  [3,4] System:

Reference: A. Treinin and E. Hayon, J. Am. Chem. Soc. 97(7): 1716-21 (1975)

Transient generation: laser flash photolysis, frequency-quadrupled neodynium laser, excitation wavelength 265 nm; pulse length  $\approx$  15 ns; energy per pulse  $\approx$  2 x 10<sup>-2</sup> J

Spectral acquisition: monitoring lamp was a pulsed 250 W Osram xenon lamp; monitoring beam formed a right angle to the laser beam; time delay 20 ns after pulse

Transient formation [5]:

Br<sub>2</sub> + hv → 2Br

Brg + hv + Br + Br2

[1] Spectrum corrected for depletion of Br2 and Br3 and for Footnotes:

formation of Bra

Spectrum is attributed to a complex with the transition being due to an electron transfer from H<sub>2</sub>O to Br. Extinction coefficient measured under conditions such that all Br is converted to Br<sub>2</sub>. 2

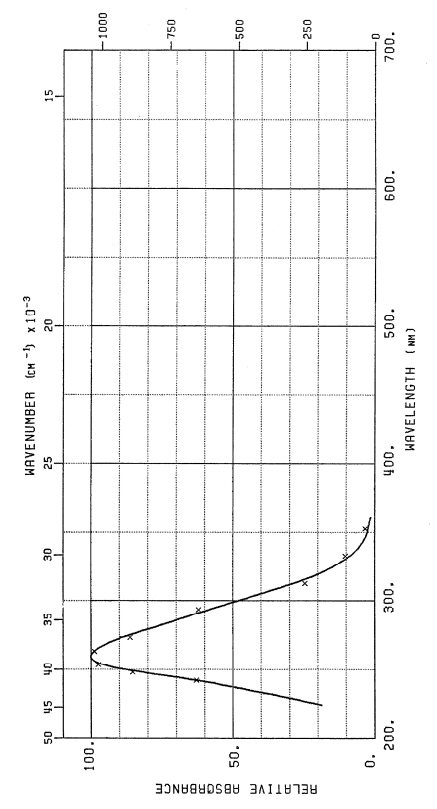
[3]

Extinction coefficient of Br2 is taken to be c360 = 9900 L mol-1 cm-1, [4]

D. Behar, private communication, 1980. [2]

Under conditions where no Br was added initially, authors calculated that their solutions contained 5 to 15 x 10<sup>-6</sup> mol L<sup>-1</sup> Br and 2 to 7 x 10<sup>-7</sup> mol L<sup>-1</sup> Brs. Under experimental conditions absorbance due to Brs was 0.1 to 0.2 at 265 nm.





Transfent: I, iodine aton,  $\lambda(max) = 260 \text{ nm}$ ,  $\epsilon(max) = 1040 \text{ L mol}^{-1} \text{ cm}^{-1}$  [1] System: aqueous solution of 3 x 10-4 mol L-1 I2

82

Reference: P. Fornier de Violet, R. Bonneau, and J. Joussot-Dubien, Chem. Phys. Lett. 19(2): 251-3 (1973)

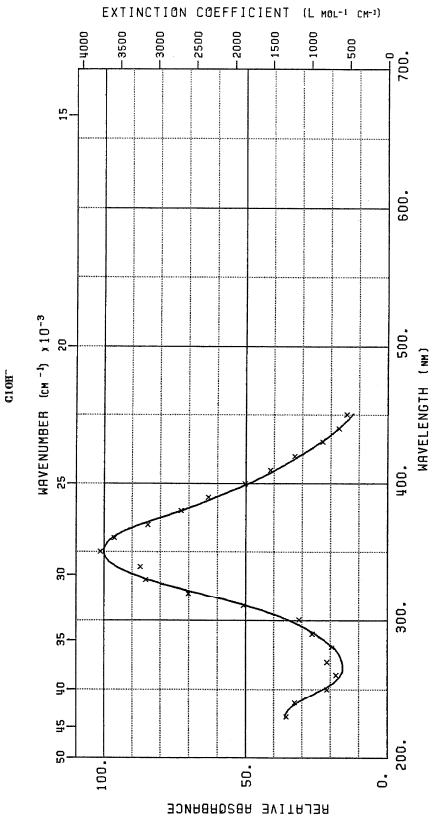
Transient generation: laser flash photolysis; excitation wavelength 529 nm; pulse energy 1 J

Spectral acquisition: pulsed xenon monitoring lamp; rise time of detection system < 10 ns; accuracy & 0.5 % on transmission measurements

Transient formation:

 $I_2 + h\nu + 2I$ 

Footnote: [1] Extinction coefficient attributed to these authors by A. Treinin and E. Hayon, J. Am. Chem. Soc. 97(7): 1716-21 (1975). Basis for extinction coefficient measurement is not known.



System: aquecus solution of 3 mol L<sup>-1</sup> NaCl, neutral pH, N<sub>2</sub>O or O<sub>2</sub> saturated [1] Transient [1]: C10H", \(\)(max) = 350 nm, \(\epsilon\_{350} = (3700 \pm 400) \) L mol \(^{-1}\) cm \(^{-1}\)

Reference: G.G. Jayson, B.J. Parsons, and A.J. Swallow, J. Chem. Soc., Faraday Trans. 1 69: 1597-607 (1973)

pulse radiolysis, electron linear accelerator; dose per pulse 1.63 x  $10^3$  rad; dosimetry-secondary emission chamber [4] Transient generation:

two Bansch & Lomb high-intensity monochromators used in series to reduce scattered light Spectral acquisition:

Transient formation:

OH + C1 → C1OH ; K = (0.70 ± 0.13) L mol-1

N20-saturated solutions had these reactions in addition to the extra C10H-Footnotes: [1] The resulting spectrum was obtained from the spectra obtained from N2O-saturated and from O2-saturated solutions. The absorption in O2-saturated solutions was thought to be due to both G10H and G12. The G12 was thought to be formed in the spurs by hole scavenging with G1 or by G10H + H \* H2O + G1 and G1 + G1 \* G12". The formed by OH formed from

 $N_2O + e(aq)^- (+H_2O) \rightarrow N_2 + OH^+ + OH$ 

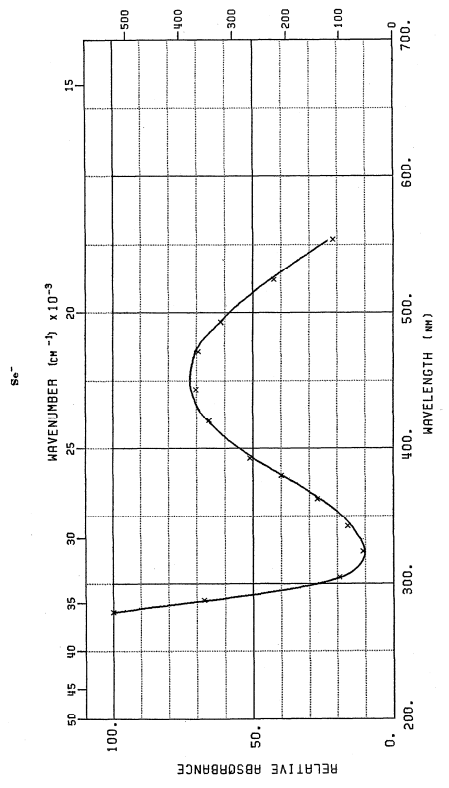
Spectra from 02-saturated solutions were corrected for 02 absorption, using  $\epsilon_{24.0} \approx 1720~L$  mol<sup>-1</sup> om<sup>-1</sup> for 0<sub>2</sub>.

Assumed excess yield of OH in N20 solutions was G(e(aq)") = 2.75 for OH formed from N20 reaction with hydrated electron. [2]

A more recent report gave a value for c(max) of C10H that was much lower, c(max) = 1200 L mol<sup>-1</sup> cm<sup>-1</sup>. [3]

R.W. Fessenden, private communication, 1980.

[41 Galibrated by aqueous potassium thiocyanate taking G((SCN)2") = 2.9 and c500((SCN)2") = 7100 L, mol<sup>-1</sup> cm<sup>-1</sup>.



Transient: Se<sup>-</sup>, \(\max\) = 450 nm, \(\epsilon\) = 380 L mol<sup>-1</sup> cm<sup>-1</sup> [1], 2k = (4.6 \tau 1.5) x 10<sup>9</sup> L mol<sup>-1</sup> s<sup>-1</sup> System: aqueous solution [2] of HSe-, N20-saturated, pH = 11

Reference: M. Schoeneshoefer, W. Karmann, and A. Henglein, Int. J. Radiat. Phys. Chem. 1: 407-23 (1969)

Transient generation: pulse radiolysis, Van de Graaff generator; energy 1.6 MeV; dose rate during the pulse was 700 rad per µs; dosimetry based on SCN-[3]

Spectral acquisition [4]: Osram XBO 450 monitoring lamp; Zeiss M4Q3 monochromator; EMI 9558 PMT; decay curves photographed from oscilloscope display; time delay 20 µB

Transient formation:

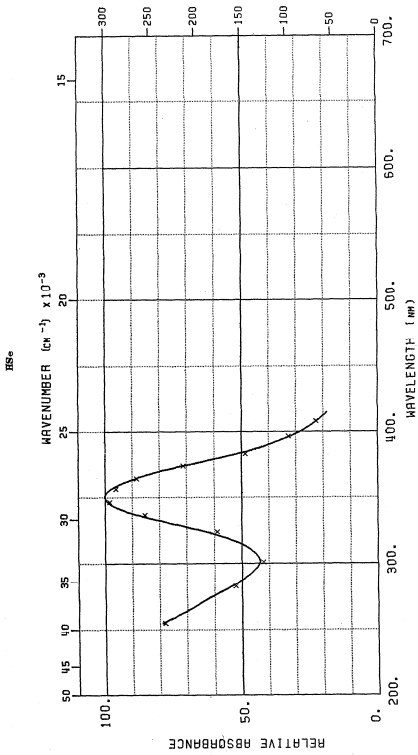
 $N_2O + e(aq)^- (+H_2O) \rightarrow OH + OH^- + N_2$ 

 $0H + HSe^- \rightarrow H_2O + Se^-$ ,  $pK_a(H_2Se) = 3.77$ 

 $H + HSe^- \rightarrow H_2 + Se^-$ 

Footnotes:

[1] Based on G(OH) = 6.
[2] Oxygen was removed from water before H<sub>2</sub>Se was added to make solution.
[3] Based on the measurement of SGN<sup>-</sup> solutions under equivalent conditions, using c<sub>500</sub> = 7100 L mol<sup>-1</sup>.
[4] A. Henglein, Allg. Prakt. Chem. 17(5): 295-301 (1966).



Translent: HSe, λ(max) = 350 nm, ε350 z = 300 L mol<sup>-1</sup> cm<sup>-1</sup> [1], 2k = (8.1 ± 2.0) x 10° L mol<sup>-1</sup> s<sup>-1</sup> System: aqueous solution [2] of H<sub>2</sub>Se, N<sub>2</sub>O-saturated, pH = 0.8

88

Reference: M. Schoeneshoefer, W. Karmann, and A. Henglein, Int. J. Radiat. Phys. Chem. 1: 407-23 (1969)

Transient generation: pulse radiolysis, Van de Graaff generator; energy 1.6 MeV; dose rate during the pulse was 700 rad per µs; dosimetry based on SGN [3]

Spectral acquisition [4]: Osram 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 PMT; decay curves photographed from oscilloscope display; time delay 5 µs after pulse

Transient formation:

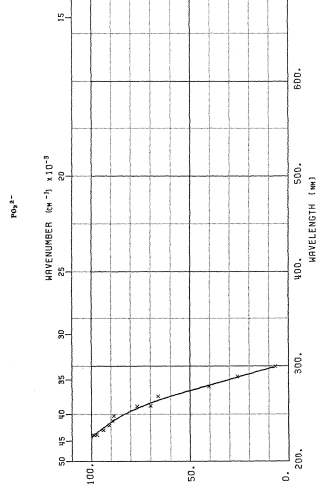
 $N_2O + e(aq)^- (+H_2O) \rightarrow OH + OH^- + N_2$ 

 $0H + H_2S_e \rightarrow HS_e + H_2O$ ,  $pK_a(H_2S_e) = 3.77$ 

 $H + H_2Se \rightarrow HSe + H_2$ 

Footnotes:

[1] Based on G(OH) = 6.
[2] Oxygen was removed from water before H<sub>2</sub>Se was added to make solution.
[3] Based on the measurement of SCN solutions under equivalent conditions, using coo = 7100 L mol<sup>-1</sup>.
[4] A. Henglein, Allg. Prakt. Ghem. 17(5): 295-301 (1966).



EXTINCTION COEFFICIENT (L MOL-1 CM-1)

-2000

1000

700.

-3000

0001

Transient:  $P0_3^{2-}$ , phosphite radical anion,  $\epsilon_2 \epsilon_0 = 3500$  L mol<sup>-1</sup> cm<sup>-1</sup> [1] System: aqueous solution of 2 x  $10^{-3}$  nol L<sup>-1</sup> phosphorous acid, saturated with N<sub>2</sub>O, pH 9

Reference: K. Schaefer and K.-D. Asmus, J. Phys. Chem. 84(17): 2156-60 (1980)

Transient generation: pulse radiolysis using Van de Graaff generator; energy i.6 MeV; dose rate during pulse was 700 rad per  $\mu s$ 

Spectral acquisition: Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 QB PMT

Transient formation:

 $e(aq)^- + N_2O (+H_2O) \rightarrow OH + OH^- + N_2$ 

 $0H + H_2P0_3^- \rightarrow HP0_3^- + H_20$ 

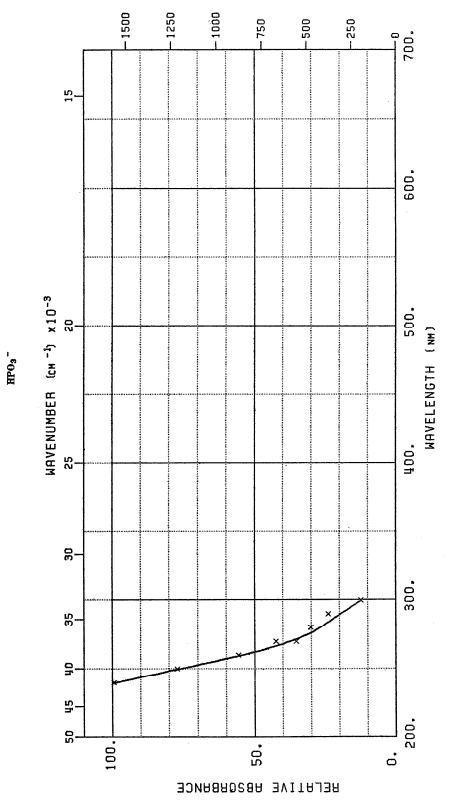
 $H + H_2PO_3^- \rightarrow HPO_3^- + H_2$ 

 $\mathrm{HPO_3}^- + \mathrm{OH}^- + \mathrm{H_2O} + \mathrm{PO_3}^{2-}, \ \mathrm{pK_a} (\mathrm{HPO_3}^-) = 5.75$ 

 ${\rm HPO_3}^{2^+} + {\rm H} \rightarrow {\rm H_2} + {\rm PO_3}^{2^-}$ 

 $\mathrm{HPO_3}^{2-} + \mathrm{OH} \rightarrow \mathrm{H_2O} + \mathrm{PO_3}^{2-}$ 

Footnote: [1] Assuming  $G(PO_3^{2-}) = G(OH) + G(H) + G(e(aq)^{-}) = 6.2$ 



EXTINCTION COEFFICIENT (L MOL-1 CM-1)

System: aqueous solution of 2 x 10-3 mol L-1 phosphorous acid, saturated with N2O, pH 4 Transient: HPO3, protonated phosphite radical anion, e250 : 1200 L mol-1 cm-1 [1]

Reference: K. Schaefer and K.-D. Asmus, J. Phys. Chem. 84(17): 2156-60 (1980)

Transient generation: pulse radiolysis using Van de Graaff generator; energy 1.6 MeV; dose rate during pulse was 700 rad per  $\mu s$ 

Spectral acquisition: Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 QB PMT

Translent formation:

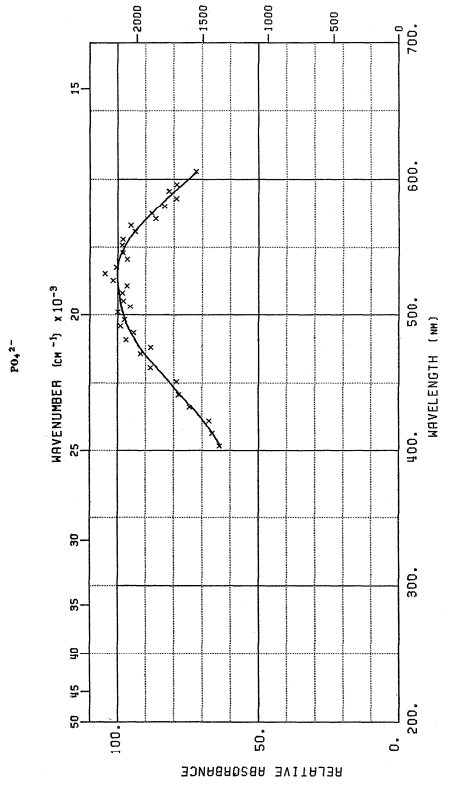
 $e(aq)^{-} + N_2O (+H_2O) \rightarrow OH + OH^{-} + N_2$ 

 $0H + H_2PO_3^- \rightarrow HPO_3^- + H_2O$ 

 $H + H_2PO_3^- \rightarrow HPO_3^- + H_2$ 

Footnote: [1] Assuming  $G(HPO_3^-) = G(OH) + G(H) + G(e(aq)^-) = 6.2$ 





Transient [1]:  $PO_4^{2-}$ ,  $\lambda(max) = 530$  nm,  $\epsilon_{530} = 2150$  L mol<sup>-1</sup> cm<sup>-1</sup>

System: aqueous solution of 2.5 x  $10^{-2}$  mol L<sup>-1</sup> peroxodiphosphate at pH 11

Reference: P. Maruthamuthu and P. Neta, J. Phys. Chem. 82(6): 710-13 (1978)

ARCO LP-7 linear accelerator; pulse length 5 ns; dosimetry thiocyanate Transient generation: Bausch 8 Lomb monochromator  $\Theta.5$  m; photomultiplier signals digitized by Biomation 8100 recorder; signal averaging with PDF-8 computer; time delay 2-10  $\mu s$  after pulse-formation was complete after this delay Spectral acquisition:

Transient formation:

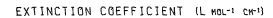
 $P_2O_6^{4-} + e(aq)^- \rightarrow PO_4^{3-} + PO_4^{2-}$ 

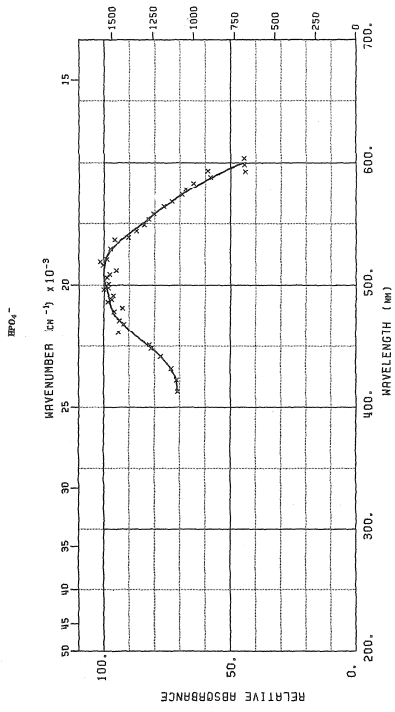
 $\mathrm{HPO_4}^- + \mathrm{OH}^- + \mathrm{PO_4}^2 + \mathrm{H_2O_4} \mathrm{pK_s} (\mathrm{HPO_4}^-) = 8.9$ 

[1] In other works where spectrs were taken at shorter wavelengths, another peak was observed in the 300-400 nm region. For examples see references in Footnotes:

footnotes 2 and 3.

[2] G. Levey and E.J. Hart, J. Phys. Chem. 79(16): 1642-6 (1975).
 [3] E.D. Black and E. Hayon, J. Phys. Chem. 74(17): 3199-203 (1970).





Transient [1]:  $\text{MPO}_4^-$ ,  $\lambda(\text{max}) = 510$  nm,  $\epsilon_{510} = 1550$  L  $\text{mol}^{-1}$  System: sque-us solution of 2.3 x  $10^{-2}$  mol L<sup>-1</sup>  $\text{HP}_20_5^{\,3}$ - at pH 7

Reference: P. Maruthamuthu and P. Neta, J. Phys. Chem. 82(6): 710-13 (1978)

Transient generation: ARCO LP-7 linear accelerator; pulse length 5 ns; dosimetry thiocyanate

Bausch & Lomb monochromator 0.5 m; photomultiplier signals digitized by Biomation 8100 recorder; signal averaging with PDF-8 computer; time delay 2-10 µs after pulse-formation was complete after this delay. Spectral acquisition:

Transient formation [3]:

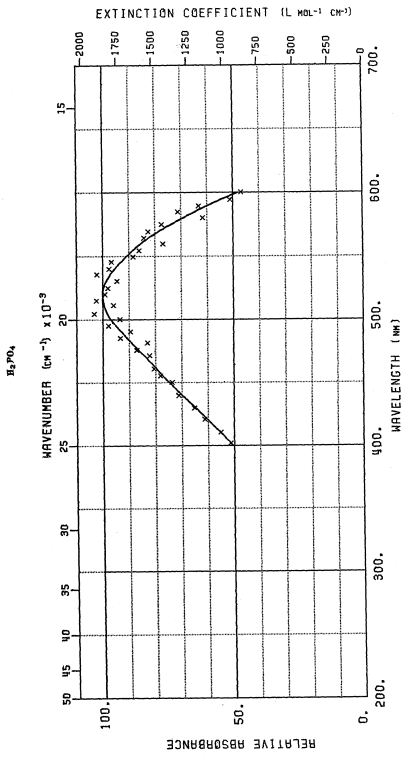
+ HPO42- + PO42-

 $PO_4^{2-} + H^+ + HPO_4^-, pK_n (HPO_4^-) = 8.9$ 

[1] In other works where spectra were taken at shorter wavelengths, another peak Footnotes:

[3]

was observed in the 300-400 nm region. For example see reference in footnote [2]. G. Levey and E.J. Hart, J. Phys. Chem. 79(16): 1642-6 (1975). At pH = 7, HP<sub>2</sub>06<sup>3-</sup> is in equilibrium with P<sub>2</sub>06<sup>4-</sup> which is about 18% of total peroxophosphate concentration, see p. 392, Topics in Phosphorus Chemistry, Vol. 7 edited by E.J. Griffith and M. Grayson. I.I. Greaser and J.O. Edwards, "Peroxophosphates".



Transient [1]:  $\rm H_2P0_4$ ,  $\lambda(\rm max)$  = 520 nm, c520 = 1850 L mol<sup>-1</sup> cm<sup>-1</sup> System: aqueous solution of 2.5 x 10<sup>-2</sup> nol L<sup>-1</sup> H<sub>2</sub>P<sub>2</sub>06<sup>2-</sup> ions at pH 4

Reference: P. Maruthamuthu and P. Neta, J. Phys. Chem. 82(6): 710-13 (1978)

Transient generation: ARCO LP-7 linear accelerator; pulse length 5 ns; dosimetry thiocyanate

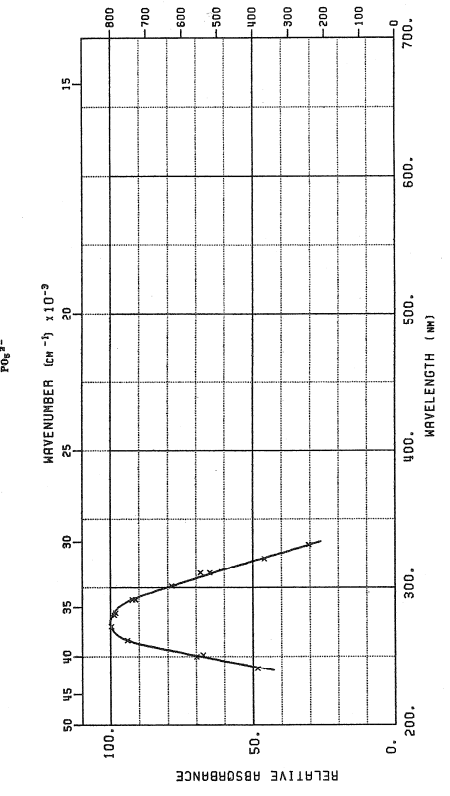
Bausch 3 Lomb monochromator 0.5 m; photomultiplier signals digitized by Bicmation 8100 recorder; signal averaging with PDP-8 computer; spectrum extrapolated to time zero. Spectral acquisition:

Transient formation:

 $e(aq)^- + H_2P_2O_8^{2-} \rightarrow HPO_4^- + HPO_4^{2-}$ 

 $\mathrm{HPO_4}^- + \mathrm{H}^+ + \mathrm{H}_2 \mathrm{PO_4}$ ,  $\mathrm{pK_a} (\mathrm{H_2PO_4}) = 5.7$ 

 [1] In other works where spectra were taken at shorter wavelengths, another peak
was observed in the 300-400 nm region. For example see reference in footnote [2].
 [2] G. Levey and E.J. Hart, J. Phys. Chem. 79(16): 1642-6 (1975). Footnotes:



Transfent:  $P0_5^2$ , peroxy phosphite radical anion,  $\lambda(\max) = 275$  nm,  $\epsilon_{275} = 800$  L mol<sup>-1</sup> cm<sup>-1</sup> System: aqueous solution of phosphite ions, containing oxygen, pH 9

Reference: K. Schnefer and K.-D. Asmus, J. Phys. Chem. 84(17): 2156-66 (1980)

Transient generation: pulse radiolysis using Van de Graaff generator; energy 1.6 MeV; dose rate during pulse was 700 rad per µs.

Spectral acquisition: Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 QB PMT

Transient formation:

 $e(aq)^{-} + N_2O (+H_2O) \rightarrow OH + OH^{-} + N_2$ 

 $OH + H_2PO_3^- \rightarrow HPO_3^- + H_2O_3^-$ 

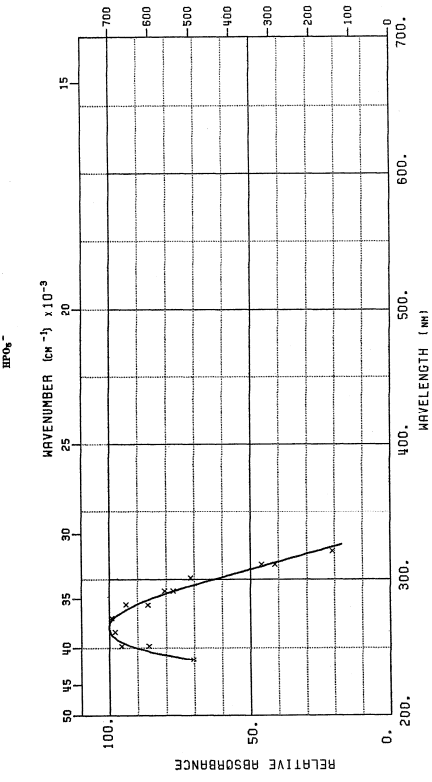
 $H + H_2PO_3^- \rightarrow HPO_3^- + H_2$ 

 $0H^- + HPO_3^- + PO_3^{2-} + H_2O, pK_a (HPO_3^-) = 5.75$ 

 $0H + HPO_3^{2-} \rightarrow H_2O + PO_3^{2-}$ 

H + HPO<sub>3</sub><sup>2</sup> - + H<sub>2</sub> + PO<sub>8</sub><sup>2</sup>

 $PO_3^{2-} + O_2 \rightarrow PO_5^{2-}$ 



Transient:  $\mathrm{HPO_5}^-$ , protonated peroxy phosphite radical anion,  $\lambda(\mathrm{max}) = 265~\mathrm{nm}$ ,  $\mathrm{c_{265}} = 700~\mathrm{L~mol}^{-1}~\mathrm{cm}^{-1}$ 

System: aqueous solution of phosphite ions, containing oxygen, pH 2.5

Reference: K. Schaefer and K.-D. Asmus, J. Phys. Chem. 84(17): 2156-60 (1980)

Transient generation: pulse radiolysis using Van de Granff generator; energy 1.6 MeV; dose rate during pulse was 700 rad per  $\mu s$ 

Spectral acquisition: Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 QB PMI

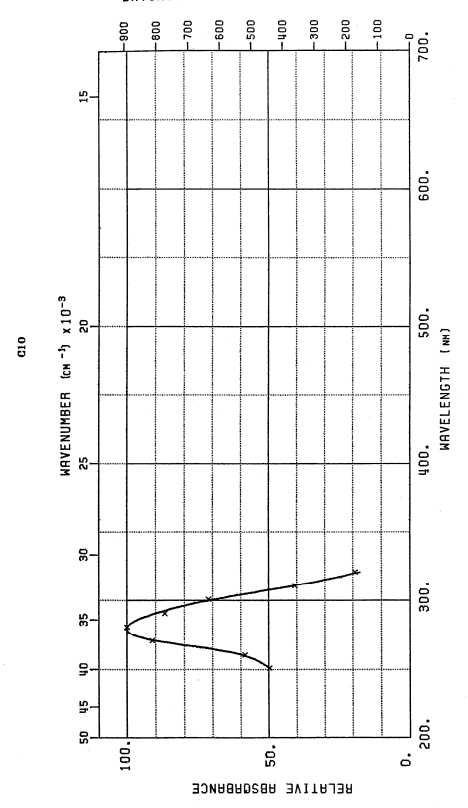
Transient fornation:

 $e(aq)^{-} + N_2O (+H_2O) \rightarrow OH + OH^{-} + N_2$ 

 $0H + H_2 PO_3^- \rightarrow HPO_3^- + H_2 O$ 

H + H<sub>2</sub>PO<sub>3</sub> → HPO<sub>3</sub> + H<sub>2</sub>

 $\mathrm{HPO_3}^- + \mathrm{O_2} \rightarrow \mathrm{HPO_5}^-$ 



Transient [11: C10, A(max) \* 280 nm, c280 \* 890 L mol-1 cm-1 [2,3] System: aqueous solution of 10-3 mol L-1 G10-, Ar-saturated, pH 11.4

Reference: G.V. Buxton and M.S. Subhani, J. Chem. Soc., Faraday 1 68: 947-57 (1972)

pulse radiolysis, Van de Graaff accelerator; dosimetry-secondary emission chamber [5] Transient generation [4]:

monitoring lamp - Philips 150 W GSM xenon lamp; Bausch & Lomb f/3.5 high intensity, grating monochromator; EMI 9558 BQ PMT; Tektronix 545 oscilloscope with type B input preamplifier; bandwidth 10 nm Spectral acquisition [4]:

Transient formation:

 $e(aq)^- + C10^- \rightarrow C1^- + 0^-$ 

 $0^- + C10^- + H_20 \rightarrow C10 + 20H^-$ 

 $e(aq)^- + C10^- + H_20 \rightarrow C1 + 20H^-$ 

C1 + C10" + C10 + C1"

H + C10 - C1 + OH

\_HO + 010 + 010 + 0H\_

Spectrum was corrected for depletion of C10 which absorbs in the region of [1] Footnotes:

reaction mechanism by using measured product yields in a steady state radiolysis Assuming G(G10) = G(e(aq)") + G(OH) + G(H). This value was calculated from the CIO absorption. 2

experiment.

890 L mol<sup>-1</sup> cm<sup>-1</sup> is the average of two extinction coefficients measured by the authors.

The average was rounded to the nearest 10 L mol<sup>-1</sup> cm<sup>-1</sup>.

The average was rounded to the nearest 10 L mol<sup>-1</sup> cm<sup>-1</sup>. [3]

K.W. Chambers, E. Collinson, F.S. Dainton, W.A. Seddon, and F. Wilkinson, Trans. Faraday Soc. 63: 1699-711 (1967). [4]

[5] Calibrated by ferrocyanide dosimeter, using G(Fe(GN), 3") = 3.2 and table (Fe(GN), 3") = 1000 L mol<sup>-1</sup> · cm<sup>-1</sup>.

Radical: G10<sub>2</sub>, λ(max) \* 360, ε<sub>360</sub> \* 1000 L mol<sup>-1</sup> cm<sup>-1</sup> [1] System: aqueous solution of 1.5 x 10<sup>-3</sup> mol L<sup>-1</sup> G10<sub>2</sub>

Reference: F. Stitt, S. Friedlander, H.J. Lewis, and F.E. Young, Anal. Chem. 26(9): 1478-84 (1954)

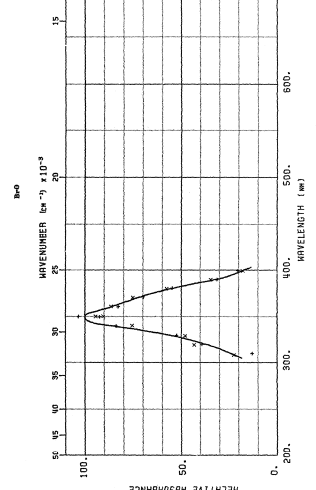
Transient generation: radical is stable

Spectral acquisition: Beckman Model DU spectrophotometer

Transient formation:

radical is stable

Footnote: [1] c = 495 L'mol - 1 cm 1 at \lambda = 404.7 nm.



EXTINCTION COEFFICIENT (L MOL-1 CM-1)

Transient: BrO, A(max) = 356 mm, case ~ 966 L mel-1 cm-1 [1]
System: aqueous 1.5 x 16-3 mel L-1 hypobromite [2]; data peints represented by "x" are for 0.1 mel L-2 MaOH, and data peints represented by "t" are for 16-3 mel L-1 NaOH.

50.

100.

Reference: G.V. Buxton, F.S. Bainton, and F. Wilkinson, Chem. Commun. (11): 320-321 (1966)

Transient generation: pulse radiolysis; pulse energy 3 MeV

Spectral acquisition:

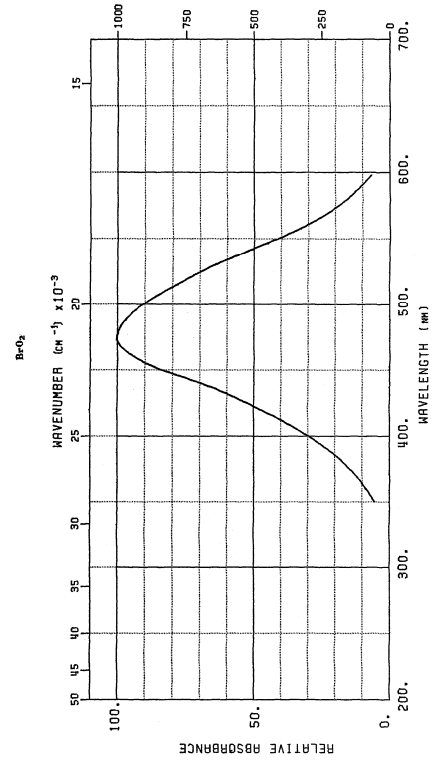
Transient formation [3]:

$$e(aq)^- + Br0^- \rightarrow Br^- + 0^-$$

$$0^- + H_20 + 0H + 0H^-$$
, pK<sub>a</sub> (0H) = 11.85

$$0H + Br0^- \rightarrow 0H^- + Br0$$

[1] Using G(e(aq)<sup>-</sup>) + G(OH) \* 6. [2] Br<sup>-</sup> was removed from the solution by adding Ag<sup>+</sup>. [3] Written for 10<sup>-3</sup> mol L<sup>-1</sup> NaOH system. Footnotes:



System: aqueous solution of BrO3, saturated with Ar, pH neutral or alkaline Transient: Br0<sub>2</sub>,  $\lambda$ (max) = 475 nm,  $\epsilon_{475}$  = (1000 ± 100) L mol<sup>-1</sup> cm<sup>-1</sup> [1]

 $Br0_2$ 

Reference: G.V. Buxton and F.S. Dainton, Proc. Roy. Soc. (London) Ser. A 304: 427-39 (1968) Transient generation: pulse radiolysis, Van de Graaff accelerator; energy 2.9 MeV; syringe technique; dosimetry-secondary emission chamber [2] Spectral acquisation [3]: monitoring lamp - Phillips 150 W GSX xenon lamp; Bausch & Lomb f/3.5 high intensity, grating monochromator; EMI 9558 BQ PMT; Tektronix 545 oscilloscope with type B input preamplifier; bandwidth 10 nm

Transient formation:

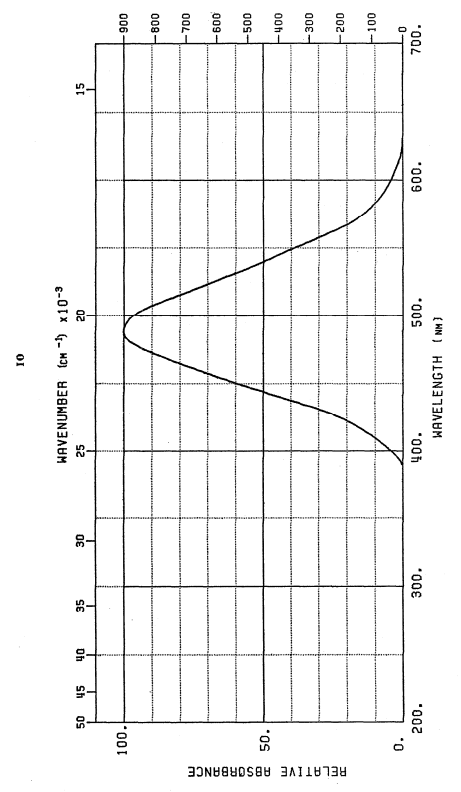
 $e(aq)^- + BrO_3^- (+H_20) \rightarrow (BrO_3^2^- + H_20) \rightarrow BrO_2 + 20H^-$ 

Based on assuming  $G(BrO_2) = G(e(aq)^-) + G(H) + G(0H)$  for alkaline solutions of  $BrO_2^-$  and  $BrO_3^-$ . In these systems  $OH + BrO_2^- + BrO_2^- + OH^-$  can occur in addition to the reduction of  $BrO_3^-$ . []] Footnotes:

Calibrated by ferrocyanide dosimeter, using G(Fe(CN)<sub>6</sub><sup>3-</sup>) = 3.2 and e<sub>420</sub> (Fe(CN)<sub>8</sub><sup>3-</sup>) = 1000 L mol<sup>-1</sup>.

K.W. Chambers, E. Collinson, F.S. Dainton, W.A. Seddon, and F. Wilkinson, Trans. [2]

Faraday Soc. 63: 1699-711 (1967). [3]



Transient: 10,  $\lambda(max) = 490$  nm, e499 = 960 ± 130 L mol<sup>-1</sup> cm<sup>-1</sup> [1] System: aqueous solution of 1.2 x 10<sup>-3</sup> mol L<sup>-1</sup> 10<sup>-</sup>, pH = 13.6

Reference: 0. Amichai and A. Treinin, J. Phys. Chem. 74(4): 830-5 (1970)

pulse radiolysis (Varian linear accelerator); energy 5 MeV; pulse length 1.5  $\mu s_1$  desimetry  $K_4 Fe(CN)_6$  [2] Transient generation:

two Bausch 8 Lomb monochromators, type 567 AB; R136 and 1P28 PMT; split monitoring beam; spectrum was recorded at 5 nm.intervals [3]; multiple reflections through cell, optical path 6 cm; time delay, 20  $\mu s$ Spectral acquisition:

Translent formation:

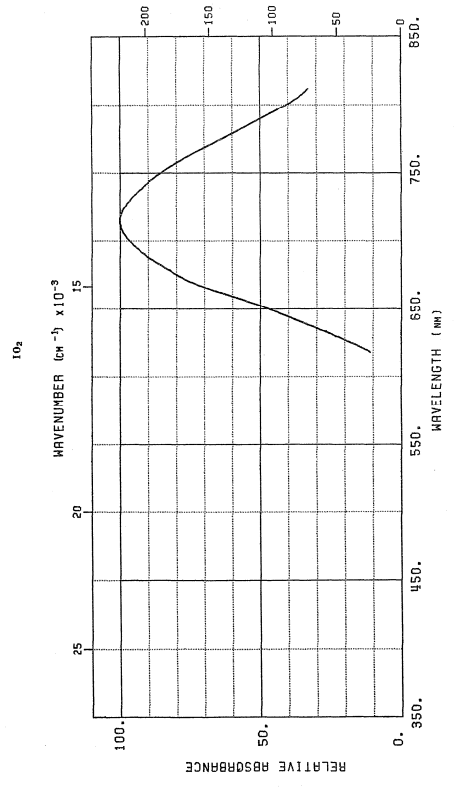
 $I0^- + e(aq)^- + I^- + 0^-$ 

 $0H^{-} + 0H \rightarrow 0^{-} + H_{2}0, pK_{a}(0H) = 11.85$ 

 $0^{-} + 10^{-} (+H_20) \rightarrow 10 + 20H^{-}$ 

Footnotes:

[1] Assuming G(10) = G(0H) + G(e(aq)<sup>-</sup>)
[2] Dosimeter consisted of air-saturated K<sub>4</sub>Fe(GN)<sub>6</sub> solution, taking e<sub>420</sub>(Fe(GN)<sub>6</sub><sup>3-</sup>) = 1000 L mol<sup>-1</sup> cm<sup>-1</sup> and G(Fe(GN)<sub>6</sub><sup>3-</sup>) = 2.7.
[3] Experimental points were within 5% of smoothed curve.



Transient [1]:  $10_2[2]$ ,  $\lambda(max) = 715$  nm,  $\epsilon_{715} = (220 \pm 30)$  L  $mol^{-1}$   $cm^{-1}$  [3] System: aqueous solution of 1.1 x 10-2 mol L-1 103-

Reference: 0. Amichai and A. Treinin, J. Phys. Chem. 74(4): 830-5 (1970)

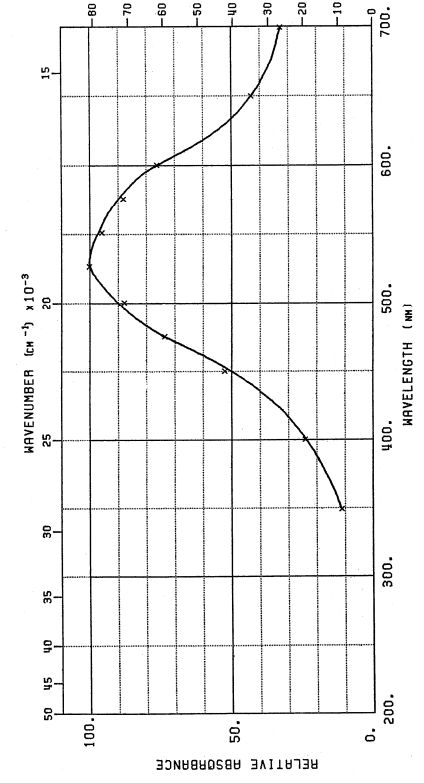
pulse radiolysis (Varian linear accelerator); energy 5 MeV; pulse length 1.5 µs; dosimetry K4Fe(CN)6 [4] Transient generation:

two Bausch 3 Lomb monochromators, type 567 AB; R136 and 1P28 PMT; split monitoring beam; spectrum was recorded at 5 nm intervals [5]; multiple reflections through cell, optical path 6 cm; time delay – 23  $\mu s$  after pulse Spectral acquisition:

Transient formation [6]:

 $e(aq)^- + 10_3^- (+H_20) \rightarrow 10_2 + 20H^-$ 

- [1] In the flash photolysis of aqueous solutions of KIO3, F. Barat, L. Gilles, B. Hickel, and J. Sutton, Chem. Commun. (24): 1485 (1969) assigned a spectrum having λ(max) = 480 nm to the IO<sub>2</sub> radical.
  [2] The spectrum was a composite spectrum due to IO<sub>3</sub>, IO, and IO<sub>2</sub>. The IO<sub>2</sub> component was resolved by Gaussian fit on a wavenumber scale. Footnotes:
- [3] From ratio of optical densities at zero tine, A(102)/A(103), the authors
- obtained  $e(\max)10_2/e(\max)10_3 = 0.55 \pm 0.08$  For  $10_3$ ,  $e(\max) = (400 \pm 60)$  L mol<sup>-1</sup>, assuming  $G(10_3) = G(0H)$ , see footnote 6. Dosimeter consisted of air-saturated  $K_4 Fe(GN)_6$  solution, taking  $e_{420}(Fe(GN)_6^{3-}) = 1000$  L mol<sup>-1</sup> cm<sup>-1</sup> and  $G(Fe(GN)_6^{3-}) = 2.7$ . <u>4</u>
  - Experimental points were within 5% of smoothed curve. [3]
    - Spectrum was complicated by absorption by 103 and 10 0H +  $10_3^-$  +  $10_3^+$  + 0H<sup>-</sup>  $10_2^-$  +  $10_3^-$  +  $10_4^-$



NH2

Transient [1]: NH2, amino radical, \(\text{Max}\) = 530 nm, c630 = 81 L mol^1 cm^1 [2] System: aqueous solution of ammonia, N20 saturated

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Reference: P.B. Pagsherg, RISO-256, Jan. 1972. p. 269-21 [Report, AEK-Riso, Roskilde, Denmark]

Transient generation: pulse radiolysis; dose rates during the pulse were 1-10 krad/ $\mu s$ ; pulse length 0.4-2  $\mu s$ ; dosimetry-ferrocyanide [3]

Spectral acquisition: optical path length of 8 cm [4]

Transient formation:

 $e(aq)^- + N_2O (+H_2O) \rightarrow OH + OH^- + N_2$ 

 $0H + NH_3 + NH_2 + H_20$ ,  $pK_a(NH_3^+) = 2.3$  [5]

[1] There was another band, that is not displayed, with a maximum at 215 nm. No extinction coefficient was given for this band, but it was assigned Footnotes:

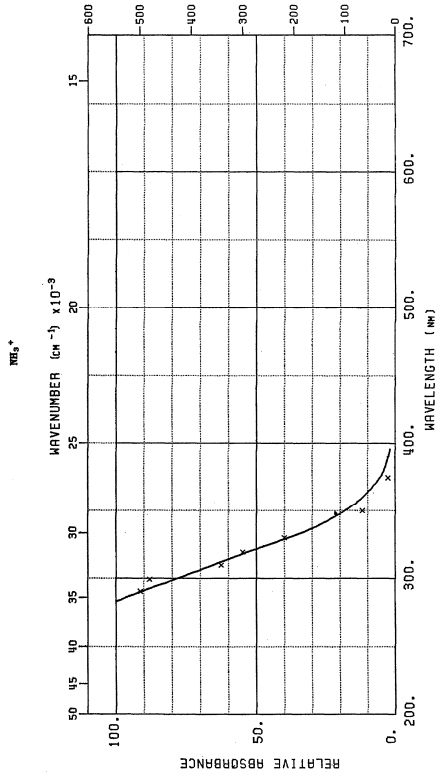
to the charge-transfer complexes (NHz, H<sub>2</sub>O) and (NHz, NH<sub>3</sub>).

[2] Based on the solution to the rate equation for the above formation equations, assuming a constant production of OH during the pulse.

[3] Dosimeter consisted of aqueous solution of 10<sup>-3</sup> mol L<sup>-1</sup> K<sub>4</sub>Fe(CN)<sub>6</sub>, 2.5 x 10<sup>-2</sup> mol L<sup>-1</sup> N<sub>2</sub>O, and about 10<sup>-4</sup> mol L<sup>-1</sup> O<sub>2</sub>. G(Fe(CN)<sub>6</sub><sup>3</sup>) assumed to be equal to G(e(aq)<sup>-</sup>) + G(OH) and c<sub>4.20</sub>(ferricyanide) = 1000 L mol<sup>-1</sup> cm<sup>-1</sup>.

[4] Optical densities were of the order of 0.03. The signal to noise ratio was 100.

[5] M.Z. Hoffman and K.R. Olson, J. Phys. Chem. B2(24): 2631-2 (1978).



Transient: NH3+, protonated amino radical, c290 \* 500 L mol-1 cm-1 [1], 2k \* 7 x 108 L mol-1 s-1 System: aqueous solution of 2 x 10-5 mol L-1 Co(NH3)63+, pH 1 [2], deserated

NH3+

Reference: M.Z. Hoffman and K.R. Olson, J. Phys. Chem. 82(24): 2631-2 (1978)

Transient generation: flash photolysis

Spectral acquisition: quartz-jacketed optical cell, path length 22 cm; time delay 50 µs after pulse

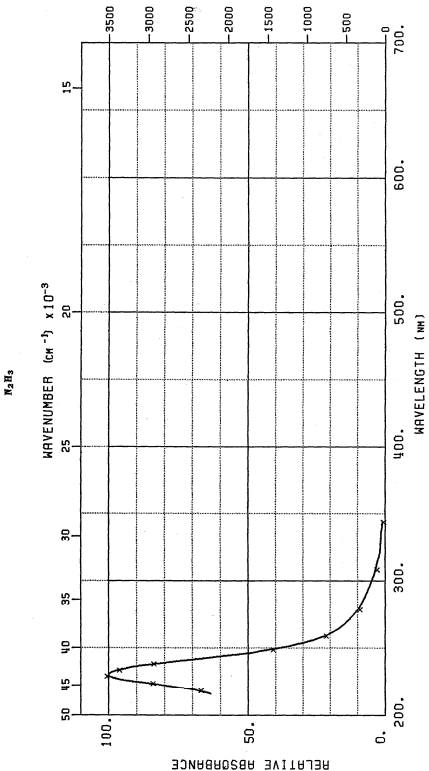
Transient formation [3]:

 $C_{0}(III)(NH_{3})_{6}^{3+} + h_{V} \rightarrow C_{0}(II)(NH_{3})_{5}(NH_{3}^{+})^{3+}$ 

 $Co(II)(NH_3)_5(NH_3^+)^{3+}$  (+5H<sup>+</sup>)  $\rightarrow Co(aq)^{2+} + 5NH_4^+ + NH_3^+$ 

 $NH_3^+ \leftrightarrow NH_2^- + H^+, pK_a(NH_3^+) = 2.3 [4]$ 

[1] Based on \* 4 x 10<sup>-6</sup> mol L<sup>-1</sup> Co(aq)<sup>2+</sup> formed as determined by the method of R.E. Kitson, Anal. Chem. 22: 664-7 (1950)
[2] Acidity adjusted with HG104.
[3] Quantum yield at 254 nm = 0.16, M.F. Manfrin, G. Varani, L. Moggi, and V. Balzani, Mol. Photochem. 1(4): 387-402 (1969).
[4] At pH 4, absorption spectrum could not be distinguished from scattered light. Footnotes:



EXTINCTION COEFFICIENT

(L MOL-1 CM-1)

Transient: N<sub>2</sub>H<sub>3</sub>, hydrazyl radical,  $\lambda(\max) = 230$  nm,  $\epsilon_{230} = 3500$  L mol<sup>-1</sup> cm<sup>-1</sup> [1],  $2k = 2.4 \times 10^9$  L mol<sup>-1</sup> s<sup>-1</sup> System: aqueous solution of  $2 \times 10^{-2} \text{ mol L}^{-1} \text{ N}_2\text{H}_4$ , N<sub>2</sub>O (1 atm), pH 9.2

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Reference: E. Hayon and M. Simic, J. Am. Chem. Soc. 94(1): 42-7 (1972)

Transient generation: pulse radiolysis, Febetron 705; pulse energy 2.3 MeV; pulse length: \* 30 ns; dose \* 8 krad per pulse; dosimetry KSCN

double monochromator; 450 W xenon lamp, pulsed to increase output by factor of  $\approx 20\text{--}25$ ; time delay  $\approx 0.1~\mu s$  after pulse Spectral acquisition:

Transient formation:

 $N_2O + e(aq)^- (+H_2O) \rightarrow N_2 + OH^- + OH$ 

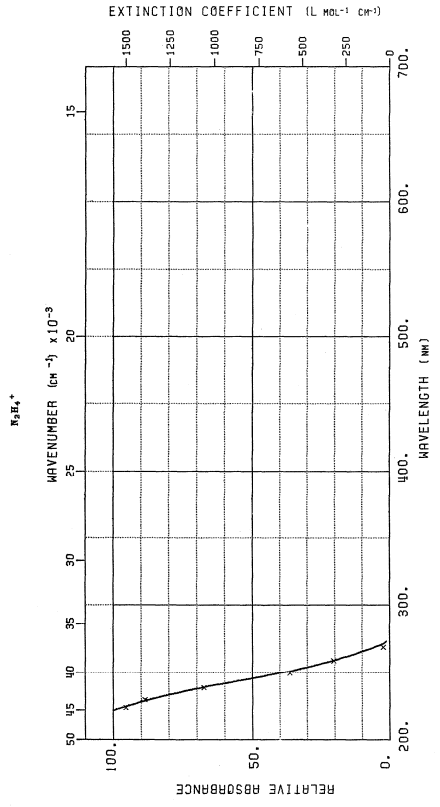
 $0H + N_2H_4 \rightarrow N_2H_4^{+} + 0H^{-}$ 

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either  $OH + N_2H_4 \rightarrow N_2H_3 + H_2O$ 

 $N_2H_4^+ + 0H^- + N_2H_8 + H_2O$ ,  $pK_a(N_2H_4^+) = 7.1 \pm 0.1$ 

Footnote: [1] Using C(e(aq)") = G(OH) = 2.8.



Transient: N2H4+, c225 = 1500 L mol-1 cm-1 [1], 2k = 6.0 x 108 L mol-1 s-1 System: aqueous solution of 2 x  $10^{-2}$  mol L<sup>-1</sup>  $N_2H_4$ ,  $N_2O$  (1 atm), pH 3.7

Reference: E. Hayon and M. Simic, J. Am. Chem. Soc. 94(1): 42-7 (1972)

Transient generation: pulse radiolysis, Febetron 705; pulse energy 2.3 MeV; pulse length & 30 ns; dose & 8 krad per pulse; dosimetry KSCN

double monochromator; 450 W xenon lamp, pulsed to increase output by a factor of  $\approx 20\text{--}25$ ; time delay  $\approx 0.1~\mu s$  after pulse Spectral acquisition:

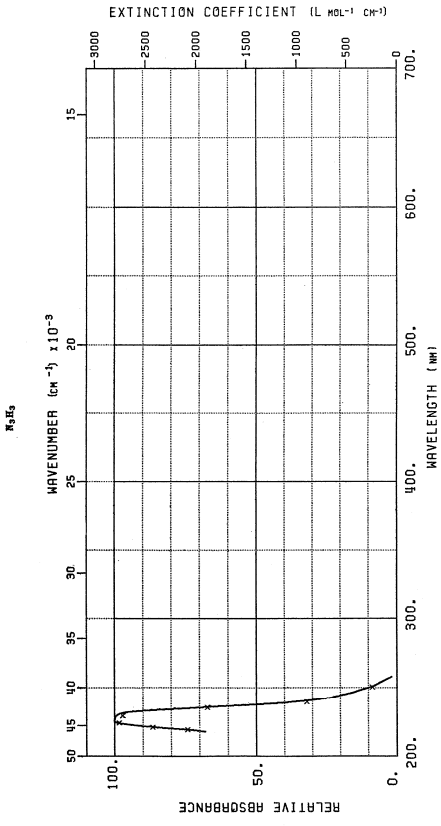
Transient formation:

 $N_2O + e(aq)^- (+H_2O) \rightarrow N_2 + OH + OH^-$ 

 $N_2H_5^+ + 0H \rightarrow N_2H_4^+ + H_2O$ ,  $pK_a(N_2H_5^+) = 8.1 \pm 0.1$ 

and  $pK_n(N_2H_4^+) = 7.1 \pm 0.1$ 

Footnote: [1] Using G(e(sq)") = G(OH) = 2.8.



System: aqueous solution of 2 x 10-2 mol L-1  $\rm N_2H_4$ ,  $\rm N_2O$  (1 atm), pH 9.2 Transient:  $N_3H_3$ ,  $\lambda(max) = 225$  nm,  $\epsilon_{225} = 2800$  L mol<sup>-1</sup> cm<sup>-1</sup>

Reference: E. Hayon and M. Simic, J. Am. Chem. Soc. 94(1): 42-7 (1972)

Transient generation: pulse radiolysis, Febetron 705; pulse energy 2.3 MeV; pulse length ~ 30 ns; dose ~ 8 krad per pulse; dosimetry KSCN

double monochromator; 450 W xenon lamp, pulsed to increase output by factor of  $\approx 20-25$ ; time delay 10 ms after pulse Spectral acquisition:

Transient formation:

$$N_20 + e(aq)^- (+H_20) \rightarrow N_2 + OH + OH^-$$

$$OH + N_2H_4 \rightarrow N_2H_3 + H_2O$$
 [1]

$$N_4H_8^{2+} + H_2O \rightarrow NH_3 + N_3H_4^+ + H_3O^+$$

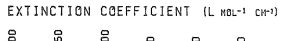
$$0H^{-} + N_{3}H_{4}^{+} + N_{3}H_{3} + H_{2}O, pK_{a}(N_{3}H_{4}^{+}) = 7.0 \pm 0.2$$

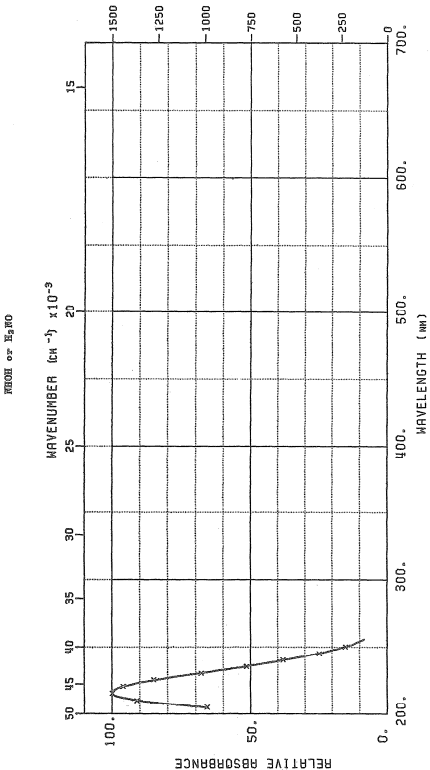
Footnotes: [1] Alternative formation of N2H3 radical is

$$0H^- + N_2H_4^+ + N_2H_3 + H_2O$$
,  $pK_n(N_2H_4^+) = 7.1 \pm 0.1$ .

[2] Disproportionation

was ruled out because  $\rm N_2H_2$  should give an equivalent amount of  $\rm N_2$  and  $\rm H_2$  , and this was not observed.





System: aqueous solution of 10-3 mol L-1 hydroxylamine, 2 x 10-2 mol L-1 N2O, pH 7.6 or 5.4 [6] Transient [11: NHOH[2] or H<sub>2</sub>NO[3],  $\lambda$ (max) = 216 nm,  $\epsilon_{216} \approx 1500$  L mol<sup>-1</sup> cm<sup>-1</sup> [4,5]

## NHOH or H2NO

Reference: D.Behar, D. Shapira, and A. Treinin, J. Phys. Chem. 76(2): 180-6 (1972)

Transient generation: pulse radiolysis, using linear accelerator; pulse energy 5 MeV; dosimetry modified Fricke [8]

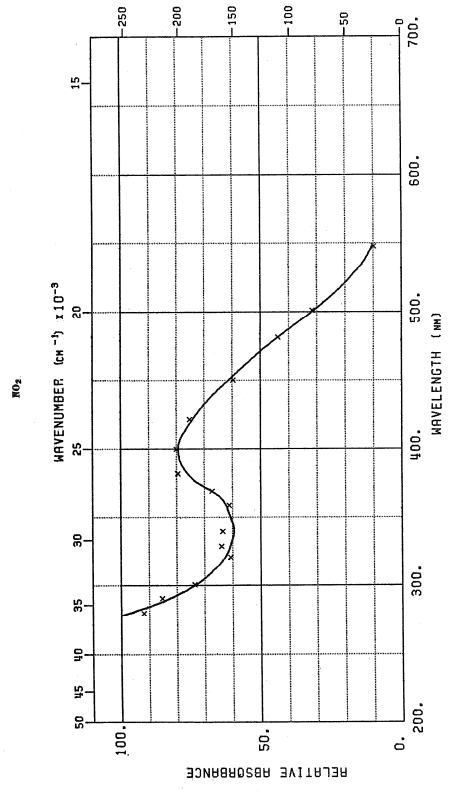
Spectral acquisition: 60 W deuterium monitoring lamp; "solar blind" R166 PMT [9]

Transient formation [10]:

 $e(aq)^- + N_2O (+H_2O) \rightarrow N_2 + OH + OH^-$ 

 $0H + NH_2OH \rightarrow NHOH (or H_2NO) + H_2O [111]$ 

- [1] Due to conflicts between information given in footnotes 2 and 3, the assignment of the spectrum is currently in doubt. P. Nets, private communication, 1978. Footnotes:
  - Simic and Hayon [7] assigned the spectrum to NHOH and not to  $\rm H_2NO$  because (a) the similarity of the spectra of the intermediates formed in the reaction of OH with NH<sub>2</sub>OH and NH<sub>2</sub>OCH<sub>3</sub> and (b) the large difference in Simic and Hayon used (a) and (b) to suggest that the radicals from NH2OH and the acid-base properties of these two radicals from NH2OH and NH2OCH3.
    - NH20CH3 are NHOH and NHOCH3, respectively. From esr data 0.P. Chawla and R.W. Fessenden, J. Phys. Chem. 79(24): 2693-700 (1975) assigned the radical, observed from the reaction of NH20H with [3]
      - SO4", as H2NO.
- Messurement at 230 nm gave  $\epsilon_{239}$  = (1020  $\pm$  30) L mol<sup>-1</sup> cm<sup>-1</sup> at pH 5.5. Another value was reported in the literature,  $\epsilon_{217}$  = 2500 L mol<sup>-1</sup> cm<sup>-1</sup> [7]. Each data point is the average of six measurements, three measurements at each pH. **4 5 9** 
  - The scatter of these points was 15%.
  - M. Simic and E. Hayon, J. Am. Chem. Soc. 93: 5982-6 (1971). M.S. Matheson and L.M. Dorfman, Pulse Radiolysis, MIT Press, Cambridge, 1969.
    - Corrections were made for stray light. Written for pH 7.6.
- pKa (NH3 OH+) = 6.0 ± 0.1 and pKa (NH2 OH+) = 4.2 ± 0.1.



Transient: NO2, \(\lambda\) (max) = 400 mm, \(\epsilon\) = 201 L mol^{-1} cm^{-1}, 2k = (9.0 \pm 2.0) \times 10^8 L mol^{-1} s^{-1} System: aqueous solution of NO2", saturated with N20

Reference: M. Graetzel, A. Henglein, J. Lilie, and G. Beck, Ber. Bunsenges. Physik. Chem. 73(7): 646-53 (1969)

Transient generation: pulse radiolysis [1] using Van de Graaff generator; energy 1.6 MeV

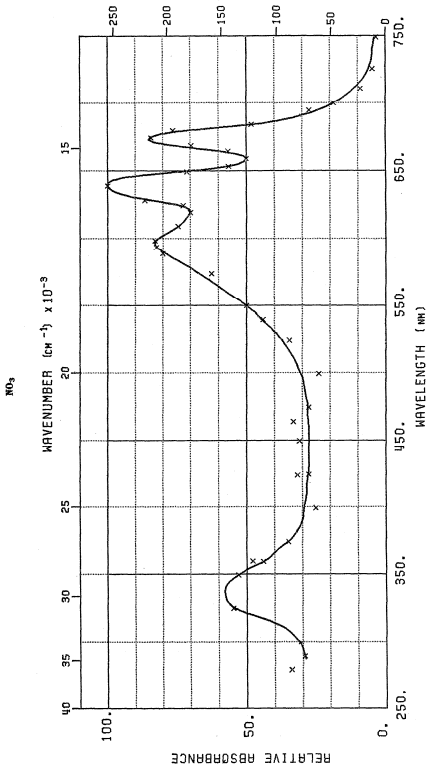
Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 PMT; decay curves photographed from oscilloscope display Spectral acquisition:

Translent formation:

 $N_2O + e(aq)^- (+H_2O) \rightarrow N_2 + OH + OH^-$ 

 $0H + NO_2^- \rightarrow 0H^- + NO_2$ 

Footnote: [1] A. Henglein, Alig. Prakt. Chem. 17(5): 295-301 (1966).



Transient [1]:  $NO_3$ ,  $\lambda(max) = 595$ , 640, 675 nm,  $c_{535} = (250 \pm 90)$  L  $mol^{-1}$  cm<sup>-1</sup> [2] System: aqueous solution of 0.1 mol L<sup>-1</sup>  $K_2Ce(NO_3)_6$ , pH 0.65 [3]

Reference: L. Dogliotti and E. Hayon, J. Phys. Chem. 71: 3802-8 (1967)

Transient generation [4]: flash photolysis; oxygen flash lamp with total energy 1800 J per flash; total flash duration 70  $\mu s$ , duration at "1/e time" 5  $\mu s$ 

monitoring lamp, attention given to minimize exposure of the solution to this lamp; EMI 9552 B PMT; Tektronix 535A oscilloscope, traces recorded on film; time delay 200 µs after start of flash Bausch 8 Lomb 250 nm grating monochromator; Osram 450 W xenon Spectral acquisition [4]:

Transient formation [5,6]:

 $Ce(IV) \cdot NO_3^- + h\nu + [Ce(III) \cdot \cdot \cdot NO_3]^*$ 

 $[Ce(III)\cdots NO_3]^* \rightarrow Ce(III) + NO_3$ 

Footnotes:

[1] Band at 340 mm was tentatively suggested to be due to N204, not N03.

R.K. Broszkiewicz, Bull. Acad. Pol. Sci., Ser. Sci. Chim. 24(3): 221-9 (1976).

[2] R.W. Glass and T.W. Martin, J. Am. Chem. Soc. 92: 5084-93 (1970).

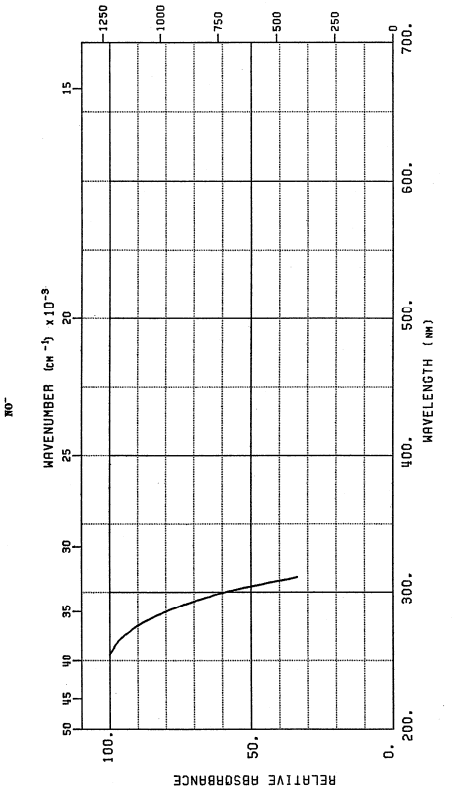
[3] Due to hydrolysis of the saft. Solutions were unbuffered.

[4] L. Dogliotti and E. Hayon, J. Phys. Chem. 71: 2511-6 (1967).

[5] Mechanism proposed by T.W. Martin and M.V. Stevens, The 12th Informal Conference on Photochemistry, June 28 - July 1, 1976, Natl. Bur. Stds., Gaithersburg, MD, Extended Abstracts, C2, p.1-5.

[6] Other mechanisms for formation have been proposed, i.e. see reference in footnote [4].





Transient: NO<sup>-</sup>,  $\lambda(max) \approx 260$  nm,  $c_{260} \approx 1290$  L mol<sup>-1</sup> cm<sup>-1</sup> [1,2] System: aqueous solution of  $\approx 10^{-4}$  mol L<sup>-1</sup> NO, pH 7.0 (buffered) [3]

W.A. Seddon, J.W. Fletcher, and F.C. Sopchyshyn, Can. J. Chem. 51(7): 1123-30 (1973) Reference:

pulse radiolysis, using Van de Grasif generator; energy 2.25 MeV; pulse length 1 µs; dose per pulse \* 3 krad; dosinetry [4] Transient generation:

Spectral acquisition: monitoring lamp [5]; Bansch & Lomb f/3.5 high intensity monochromator; RCA 1P28 or HTV R166 PMT [6]; Tektronix 556 oscilloscope; time delayimmediately after pulse

Translent formation [7]:

-0M + NO → MO-

H + NO + HNO

 $HNO + NO^{-} + H^{+}, pK_{n}(HNO) = 4.7$ 

[1] Extinction coefficient based on G(NO<sup>-</sup>) = G(e(aq)<sup>-</sup>) + G(H) = 3.2. [2] Earlier value - 1400 L mol<sup>-1</sup> cm<sup>-1</sup>, M. Graetzel, S. Taniguchi, and A. Henglein, Ber. Bunsenges Physik. Chem. 74(10): 1003-10 (1970). Footnotes:

[3] Fresh solutions were used for each pulse. [4] Using serated, 2 x 10-3 mol L-1 SCN solutions and taking Ge = 2.10 x 10- for (SCN)2

at 475 nm. Relative dose monitored by integrating current striking an Al block.

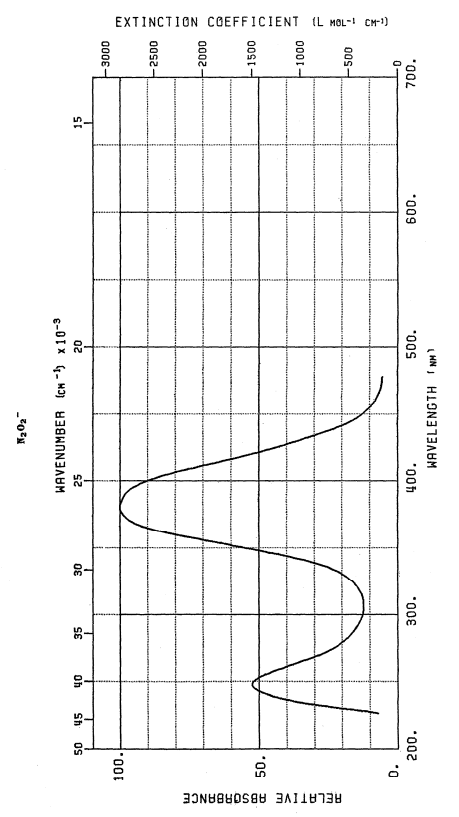
[5] Either a Hanovia 300 W high pressure xenon lamp operated in DC or pulsed mode, a CE 62 W quartz-lodine lamp, or a 30/50 W deuterium lamp was used.

[6] For wavelengths shorter than 300 nm the Ri66 PMT was used with narrow band pass

[7] Hydroxyl radical reactions were interference filters.

OH + NO → HNO<sub>2</sub>

 $HNO_2 + H^+ + NO_2^-$ ,  $pK_n (HNO_2) = 3.5$ 



Transient:  $N_2O_2^-$ ,  $\lambda(max) = 380$  nm,  $\epsilon_{380} = (2850 \pm 200)$  L mol<sup>-1</sup> cm<sup>-1</sup> [1,2] System: aqueous solution of  $\approx 2 \times 10^{-3} \text{ mol L}^{-1} \text{ NO, pH 7.0 (buffered) } [3]$ 

Reference: W.A. Seddon, J.W. Fletcher, and F.C Sopchyshyn, Can. J. Chem. 51(7): 1123-30 (1973)

pulse radiolysis, using Van de Graaff generator; energy 2.25 MeV; pulse length 1  $\mu s$ ; dose per pulse \* 3 krad; dosimetry [4] Transient generation:

monitoring lamp [5]; Bausch & Lomb f/3.5 high intensity monochromator; RCA 1P28 or HTV R166 PMT [6]; Tektronix 556 oscilloscope; time-delayimmediately after pulse Spectral acquisition:

Transient formation [7]:

e(aq) + NO + NO

H + NO + HNO

 $HNO + NO^- + H^+$ ,  $pK_n$  (HNO) = 4.7

 $NO + NO^- + N_2O_2^-$ ,  $pK_a(HN_2O_2) = 3.5$ 

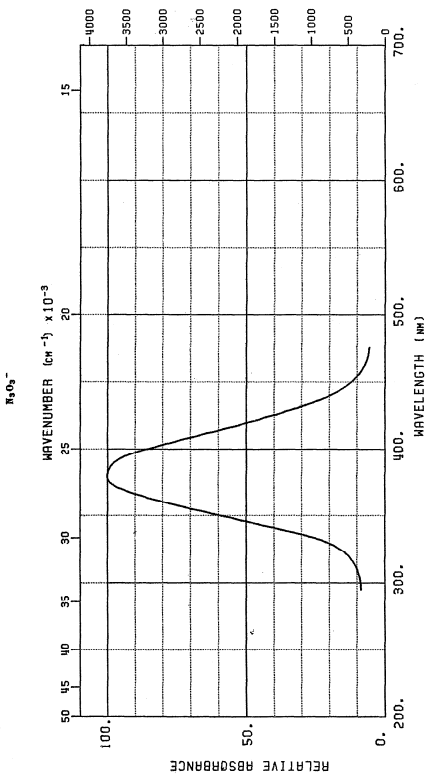
Extinction coefficient based on  $G(N_2O_2^-) = G(e(aq)^-) + G(H) = 3.2$ . Earlier value - 3300 L mol<sup>-1</sup> cm<sup>-1</sup>, M. Graetzel, S. Tanlguchi, and A. Henglein, Ber, Bunsenges Physik. Chem. 74(10): 1003-10 (1970). 52 Footnotes:

[3] Fresh solutions were used for each pulse.
[4] Using aerated, 2 x 10<sup>-3</sup> mol L<sup>-1</sup> SCN solutions and taking Ge = 2.10 x 10<sup>4</sup> for (SCN)<sub>2</sub> at 475 nm. Relative dose monitored by integrating current striking on Al block.
[5] Either a Hanovia 300 W high pressure xenon lamp operated in DC or pulsed mode, a GE 62 W quartz-iodine lamp, or a 30/50 W deuterium lamp was used.
[6] For wavelengths shorter than 300 nm the Ri66 PMT was used with narrow band pas-

[7] Hydroxyl radical reactions were interference filters.

OH + NO → HNO<sub>2</sub>

 $HNO_2 + H^+ + NO_2^-$ ,  $pK_a(HNO_2) = 3.5$ 



EXTINCTION COEFFICIENT (L mol-1 cm-1)

Transient: N<sub>3</sub>O<sub>3</sub>,  $\lambda$ (max) = 380 nm,  $\epsilon_{380}$  = (3750 ± 300) L mol<sup>-1</sup> cm<sup>-1</sup> [1,2] System: squeous solution of  $\approx 2 \times 10^{-3} \text{ mol L}^{-1} \text{ NO}$ , pH 7.0 (buffered) [3]

Reference: W.A. Seddon, J.W. Fletcher, and F.C. Sopchyshyn, Can. J. Chem. 51(7): 1123-30 (1973)

Transient generation: pulse radiolysis, using Van de Graaff generator; energy 2.25 MeV; pulse length 1 µs; dose per pulse \* 3 krad; dosimetry [4]

Spectral acquisition: monitoring lamp [5]; Bausch & Lomb f/3.5 high intensity monochromator; RCA 1P28 or HTV Ri66 PMT [6]; Tektronix 556 oscilloscope; time delayextrapolated back to end of pulse

Transient formation [7]:

$$e(aq)^- + NO \rightarrow NO^-$$

H + NO + HNO

 $HNO + NO^- + H^+$ ,  $pK_a$  (HNO) = 4.7

 $NO^- + NO + N_2O_2^-$ ,  $pK_a (HN_2O_2) = 3.5$ 

 $N_2O_2^- + NO + N_3O_3^-$ ,  $pK_a(HN_3O_3) = 3.1$ 

Footnotes:

[1] Extinction coefficient based on  $G(N_3O_3^-)$  =  $G(e(aq)^-)$  + G(H) = 3.2. [2] Earlier value - 4000 L mol<sup>-1</sup> cm<sup>-1</sup>, M. Graetzel, S. Taniguchi, and A. Henglein, Ber. Bunsenges Physik. Chem. 74(10): 1003-10 (1970).

Fresh solutions were used for each pulse. Using serated,  $2 \times 10^{-3} \text{ mol L}^{-1} \text{ SCN}^{-}$  solutions and taking  $\text{Gc} = 2.10 \times 10^{4} \text{ for (SCN)}_{2}^{-}$  at 475 nm. Relative dose monitored by integrating current striking an Al block. Either a Hanovia 300 W high pressure xenon lamp operated in DC or pulsed mode, [8]

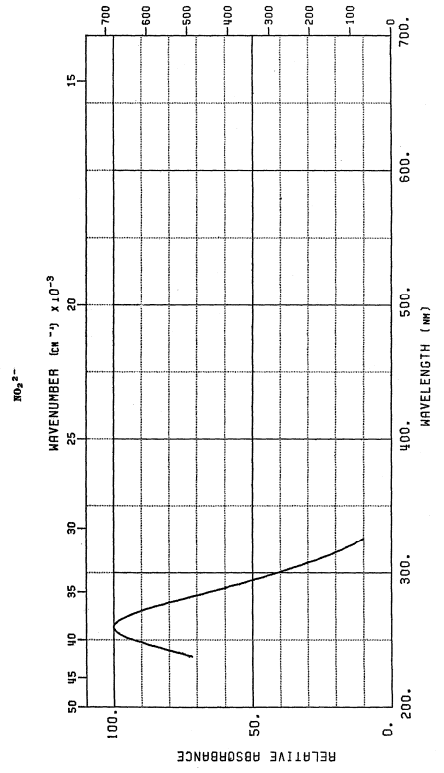
[2]

a GE 62 W quartz-lodine lamp, or a 30/50 W deuterium lamp was used. For wavelengths shorter than 300 nm the R166 PMT was used with narrow band pass [9]

[7] Hydroxyl radical reactions were interference filters.

 $OH + NO \rightarrow HNO_2$ 

 $HNO_2 \leftrightarrow H^+ + NO_2^-, pK_a (HNO_2) = 3.5$ 



System: aqueous solution of 2.5 x 10-4 mol L-1 NO2 and 0.25 mol L-1 tert-butanol, pH = 9, Ar-saturated Transient:  $NO_2^2$  [1],  $\lambda(max) = 260 \text{ nm}$ ,  $\epsilon_{260} = 680 \text{ L mol}^{-1} \text{ cm}^{-1}$ 

NO22-

Reference: M. Graetzel, A. Henglein, J. Lilie, and G. Beck, Ber. Hunsenges. Physik. Chem. 73(7): 646-53 (1969)

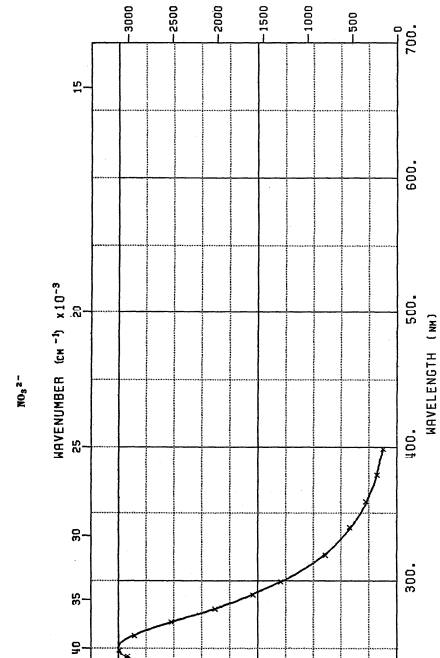
Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 PMT; decay curves photographed from oscilloscope display Transient generation: pulse radiolysis [2] using Van de Graaff generator; energy 1.6 MeV Spectral acquisition:

Transient formation:

 $0H + (CH_3)_3COH \rightarrow H_2O + CH_2C(CH_3)_2OH$ 

 $NO_2^- + e(aq)^- \rightarrow NO_2^{2-}$ 

Footnotes: [1] pKa(HND2-) = 7.7. [2] A. Henglein, Allg. Prakt. Chem. 17(5): 295-301 (1966).



EXTINCTION COEFFICIENT (L MOL-4 CM-3)

System: aqueous solution of 2 x 16-4 nol L-1 NO3 and 0.1 nol L-1 tert-butanol, pH 9.5 Transient:  $NO_3^{2-}$ ,  $\lambda(max) = 250 \text{ mm}$ ,  $\epsilon_{250} = 3100 \text{ L mol}^{-1} \text{ cm}^{-1}$  [11]

200.

0

RELATIVE ABSORBANCE

50.

£2-

20

Reference: M. Graetzel, A. Henglein, and S. Taniguchi, Ber. Bunsenges. Physik. Chem. 74(3): 292-8 (1970)

Transient generation: pulse radiolysis [2], using Van de Graaff generator

Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 PMT; decay curves photographed from oscilloscope display; time delay, immediately after pulse Spectral acquisition:

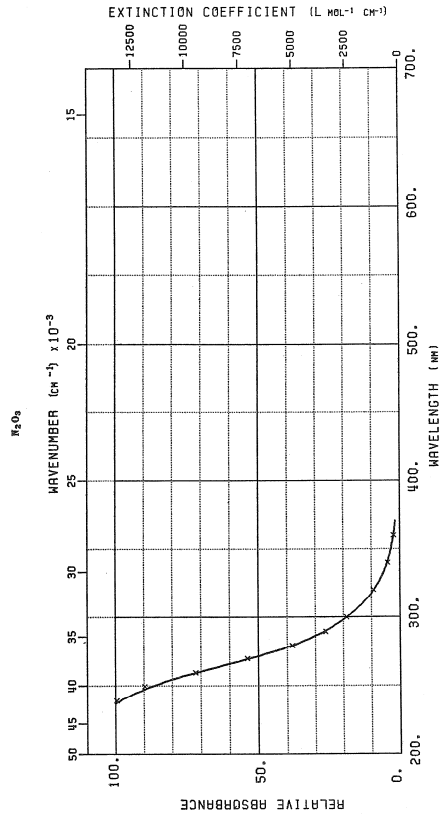
Transient formation:

OH + (CH<sub>3</sub>)<sub>3</sub>COH → H<sub>2</sub>O + CH<sub>2</sub>C(CH<sub>3</sub>)<sub>2</sub>OH

 $e(aq)^- + NO_3^- \rightarrow NO_3^{2-}$ ,  $pK_a(HNO_3^-) = 7.5$ 

[1] Based on G(e(aq)<sup>-</sup>) = 2.7. Effect of H atom was neglected because rate constant for reaction of H with NO<sub>3</sub><sup>-</sup> is small, k = 3.8 x 10<sup>6</sup> L mol<sup>-1</sup> s<sup>-1</sup> from G. Navon and G. Stein, J. Phys. Chem. 69: 1384-9 (1965).

[2] A. Henglein, Allg. Prakt. Chem. 17(5): 295-301 (1966). Footnotes:



System: aqueous solution of 2.6 x 16<sup>-5</sup> to 1.64 x 16<sup>-4</sup> mol L<sup>-1</sup> NO, 2 x 16<sup>-4</sup> to 3 x 16<sup>-4</sup> mol L<sup>-1</sup> NO<sub>2</sub><sup>-1</sup>, and 2.4 x 16<sup>-2</sup> mol L<sup>-1</sup> N<sub>2</sub>0 Transient: N203 [1,2]

Reference: M. Graetzel, S. Taniguchi, and A. Henglein, Ber. Bunsenges. Physik. Chem. 74(5): 488-92 (1970)

Transient generation: pulse radiolysis [3], using Van de Graaff generator; energy 1.5 MeV; dose rate during the pulse was 700 rad per  $\mu s$ 

Osram XBO 450 monitoring lamp; Zeiss M403 monochromator; EMI 9558 PMI; decay curves photographed from oscilloscope display; time delay 20 µs Spectral acquisttion:

Transient formation [4]:

$$N_2O + e(aq)^- (+H_2O) \rightarrow N_2 + OH + OH^-$$

$$NO_2$$
 + OH +  $NO_2$  + OH

$$NO_2 + NO + N_2O_3$$
,  $K = 1.4 \times 10^4 \text{ L mol}^{-1}$ 

[1] Extinction coefficient was obtained by assuming that the yield of  $NO_2 + N_2O_3$ Footnotes:

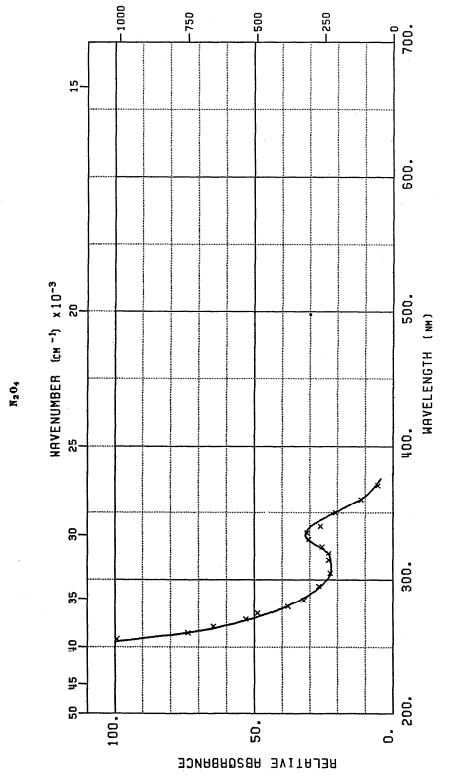
was equal to the initial yield of NO<sub>2</sub>.

Spectrum corrected for NO<sub>2</sub> absorption.

A. Henglein, Alig. Prakt. Chem. 17(5): 295-301 (1966).

Reaction of OH with NO does not involve complications since little reacts under the experimental conditions. Any NO that does react with OH only reforms the initial reactants

 $NO + OH \rightarrow NO_2 - + H^+$ 



aqueous solution of 9.5 x 10<sup>-4</sup> mol L<sup>-1</sup> NO<sub>2</sub>, saturated with N<sub>2</sub>O (\* 2.5 x  $10^{-2}$  mol L<sup>-1</sup>), pH 5.0 Transient [1]:  $N_2O_4$ ,  $\lambda(\max) \approx 340$  nm,  $\epsilon_{335} = (320 \pm 60)$  L mol<sup>-1</sup> cm<sup>-1</sup> [2] System:

Reference: A. Treinin and E. Hayon, J. Am. Chem. Soc. 92(20): 5821-8 (1970)

pulse radiolysis, using Febetron 705 System (Field Emission Corp.); pulse energy 2.3 MeV; pulse length  $\approx 30$  ns; desimetry using N<sub>2</sub>0saturated aqueous solution of 6.1 KSCN [4] Transient generation [3]:

Osram XBO 450 W xenon lamp (pulsed); two high-intensity Bausch & Lomb monochromators, used in series to reduce scattered light [5]; two EMI 9558 QB PMI; dual beam Fairchild 777 oscillosocope; time delay 160 µs after pulse Spectral acquisition [3]:

Transient formation:

 $N_2O + e(aq)^- (+H_2O) \rightarrow OH + OH^- + N_2$ 

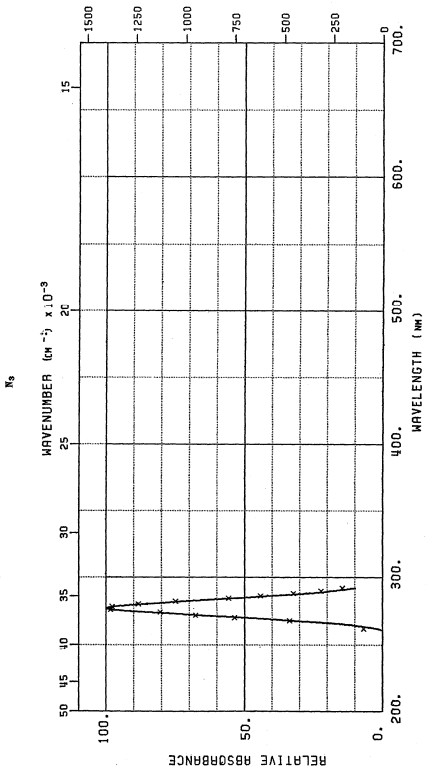
 $OH + NO_2^- \rightarrow NO_2 + OH^-$ 

 $2NO_2 + N_2O_4$ ,  $K^{-1} = (1.3 \pm 0.4) \times 10^{-6} \text{ mol L}^{-1}$ 

Footnotes: [1] Absorption at wavelength shorter than 300 nm may be partially due to  $\Pi_2 O_3$ which can be formed by

 $NO + NO_2 \rightarrow N_2O_3$ 

[2] Based on equilibrium equations for equilibrium of NO<sub>2</sub> and N<sub>2</sub>O<sub>4</sub> and taking  $\epsilon(NO_2) = 200 L \text{ mol}^{-1}$  at the maximum of the NO<sub>2</sub> spectrum.
[3] M. Simic, P. Neta, and E. Hayon, J. Phys. Chem. 73(11): 3794-800 (1969).
[4] Using  $\epsilon_5$ 00 (SCN)<sub>2</sub> = 7600 L mol<sup>-1</sup>.
[5] At 230 and 210 nm the amount of scattered light was 1% and 18%, respectively. where NO was produced by H atom reactions.



Transfent [1]: Ns. szide radical,  $\lambda(max) = 277$  nm,  $\epsilon_{275} = 1460$  L mol<sup>-1</sup>  $\epsilon m^{-1}$  [2,3],  $2k = (8.0 \pm 0.7) \times 10^{9}$  L mol<sup>-1</sup> s<sup>-1</sup> [2]

System: aqueous solution of 10-4 mol L.1 NaN3, pH ~ 9.4, N2-saturated

Reference: A. Treinin and E. Hayon, J. Chem. Phys. 50: 538-9 (1969)

Transient generation [4]: flash photolysis with four exygen flash lamps; total flash energy 1500 J; duration at "I/e" time 5 µs, total flash duration 70 µs

Spectral acquisition [4]: xenon Osram XBO 450 W monitoring lamp; Bausch 8 Lomb grating monochronator; PMT; Tektronix 535A oscilloscope; spectrum was obtained by a point-by-point method; time delay 120  $\mu s$  after start of flash

Translent formation:

Ng + hv > Ng + e(aq) -

Each data point was the average of six runs. Footnotes:

From G.V. Buxton and I. Janovsky, J. Chem. Soc., Faraday 1 72(8): 1884-6 (1976).

Other sets of c and 2k for N3 are c278 = 2300 L mol<sup>-1</sup> cm<sup>-1</sup> and

2k = (9 ± 1) x 10<sup>9</sup> L mol<sup>-1</sup> s<sup>-1</sup> from E. Hayon and M. Simic, J. Am. Chem. Soc.

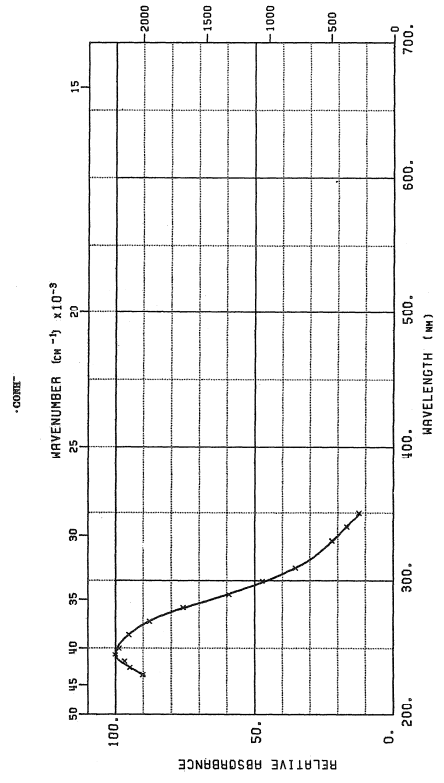
92(25): 7486-7 (1970); and c275 = 1300 L mol<sup>-1</sup> cm<sup>-1</sup> and 2k = 6 x 10<sup>9</sup> L mol<sup>-1</sup> s<sup>-1</sup>

from E.J. Land and W.A. Prutz, Int. J. Radiat. Biol. Relat. Stud. Phys., Chem. Med.
36(1): 75-83 (1979).

Although the displayed spectrum has λ(max) = 277 nm, the c(max) is shown as 1400 L mol<sup>-1</sup> cm<sup>-1</sup> on the assumption that all the authors in this case were making measurements at the maximum. [3]

L. Dogliotti and E. Hayon, J. Phys. Chem. 71(8): 2511-6 (1967). [4]





Transient: .CONHT, \(\)(max) = 245 nm, \(\epsilon\) = 2200 L mol^1 cm^1 [1], \(2k = 2.2 \times 10^9 \) L mol^1 s^1 System: aqueous solution of 10-3 nol L-1 CN- in 1 nol L-1 KOH, saturated with N20

Reference: D. Behar, J. Phys. Chem. 78(26): 2666-3 (1974)

pulse radiolysis, using electrons from Van de Grasif generator; energy 2.8 MeV; pulse length 0.5-2 µs; dose 1-5 krad [2] Translent generation:

Spectral acquisition [3]: single-pass cell, pulsed Osram 450 W XBO xenon monitoring lamp; Bausch 8 Lomb monochromator, focal length 500 cm, bandwidth 2 nm; EMI 9558 QC PMT; corrected for stray light below 240 nm

Translent formation [4]:

 $OH^- + OH \rightarrow O^- + H_2O$ ,  $pK_n(OH) = 11.85$ 

0\_ + CN\_ + \_0-C=N\_

 $-0-C=N^- + H_2O \rightarrow -0-C=NH + OH^-$ 

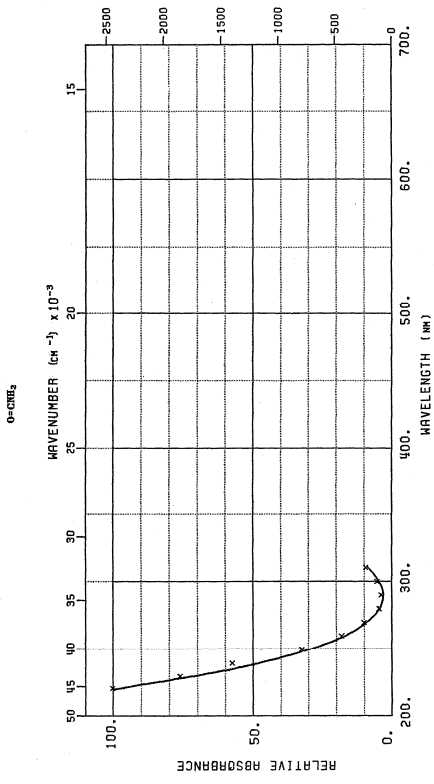
 $^{-}0-C=NH \rightarrow 0=C-NH^{-}, pK_a(NH_2C=0) = 12.25$ 

e(max) = 7600 L mol<sup>-1</sup> cm<sup>-1</sup> and by assuming G(radicals) = G(e(aq)<sup>-</sup>) + G(OH).

[2] Electron beam monitored by a secondary emission monitor.

[3] L.K. Patterson and K.M. Bansal, J. Phys. Chem. 76(17): 2392-9 (1972).

[4] H. Buechler, R.E. Buchler, and R. Cooper, J. Phys. Chem. 80(14): 1549-53 (1976). Footnotes:



EXTINCTION COEFFICIENT (L MOL-1 CM-1)

System: variety of aqueous solutions of either  $10^{-1}$  mol  $L^{-1}$  GN or  $10^{-2}$  mol  $L^{-1}$  GN, pH 9.85 to 11.5; all solutions saturated with  $N_20$ Transient: 0=CNH2 [1,2], 2k = 6.2 x 10° L mol-1 s-1

## 0=CNH2

Reference: H. Buechler, R.E. Buehler, and R. Cooper, J. Phys. Chem. 80(14): 1549-53 (1976)

pulse radiolysis, using Febetron 705 (Field Emission Corp.); energingly; dose 25.8 to 31.2 krad Transient generation [3]:

xenon XBO 450 monitoring lamp; Bausch & Lomb High Intensity monochromator; Philips XP 1003 PMT; Keithley pulse amplifier (Model 105); dual-beam oscilloscope (Tektronix 556) Spectral acquisition [3]:

Transient formation:

 $e(aq)^- + N_2O(+H_2O) \rightarrow N_2 + OH + OH^-$ 

OH + CN - + HOC=N-

 $H_2O + HOC=N^- + HOC=NH + OH^-, pK_a (HOC=NH) = 19.2$ 

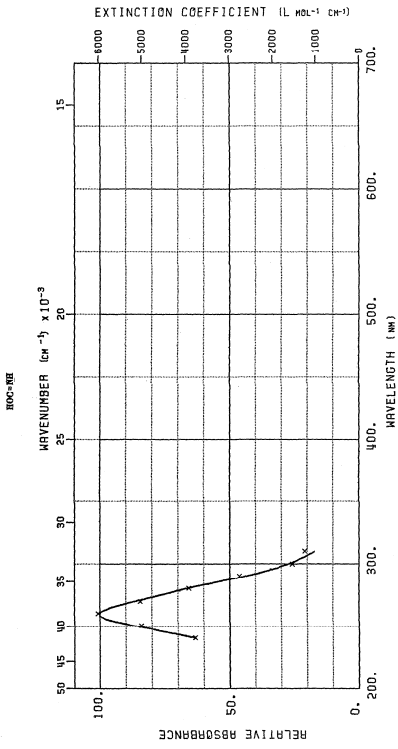
HOC=NH → 0=CNH2

[1] Spectrum corrected for spectra of the precursors HOC=NH and HOC=NT. Footnotes:

This was done by computer simulation of the kinetic scheme. Earlier spectrum by D. Behar, J. Phys. Chem. 78(26): 2660-3 (1974), shows a weak maximum in the region of 320 nm.

B. Hurni, U. Bruchiman, and R.E. Buchler, Int. J. Radiat. Phys. Chem. 7: 499-506 2

(1975). [3]



System [2]: aqueous solution of 10-2 mol L-1 GN-, pH 9.85, saturated with N20

Transient: HOC=NH [1]

## HOC=NH

Reference: H. Buechler, R.E. Buehler, and R. Cooper, J. Phys. Chem. 80(14): 1549-53 (1976)

pulse radiolysis, using Febetron 705 (Field Emission Corp.); energy 2 MeV; dosimetry using calorimetry; dose = 25.8 krad Transient generation [3]:

xenon XBO 450 monitoring lamp; Bausch & Lomb High Intensity monochromator; Philips XP 1003 PMT; Keithley pulse amplifier (Model 105); dual-beam oscilloscope (Tektronix 556); time delay variable [4] Spectral acquisition [3]:

Transient formation:

 $e(aq)^- + N_20(+H_20) \rightarrow N_2 + OH + OH^-$ 

OH + CN → HOC=N"

 $H_2O + HOC=N^- + HOC=NH + OH^-, pK_a (HOC=NH) = 10.2$ 

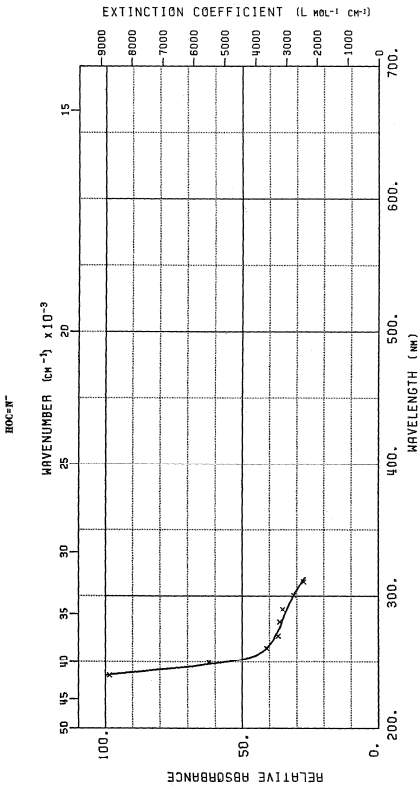
[1] Spectrum was obtained from a computer simulation using the above equations in addition to known rate constants and Footnotes:

HOC=NH → O=CNH2.

Two other chemical systems were used. One had  $[CN^-] = 10^{-2} \text{ mol L}^{-1}$ , pH 10.7, and received a dose = 25.8 krad. The other had  $[CN^-] = 10^{-1} \text{ mol L}^{-1}$ , pH = 11.4, and received a dose = 28.1 krad. Both of these solutions were saturated [5]

[3] B. Hurni, U. Bruehlman, and R.E. Buehler, Int. J. Radiat. Phys. Chem. 7: 499-506 with N20.

[4] Transient makes contribution to experimental spectrum only for time less than 1 µs.



System [2]: aqueous solution of 10-2 mol L-1 CK-, pH 19.7, saturated with N20

Transient: HOC=N- [1]

Reference: H. Buechler, R.E. Buchler, and R. Cooper, J. Phys. Chem. 80(14): 1549-53 (1976)

pulse radiolysis, using Febetron 705 (Field Emission Corp.); energy 2 MeV; dosimetry using calorimetry; dose = 25.8 krad Transient generation [3]:

xenon XBO 450 monitoring lamp; Bausch & Lomb High Intensity monochromator; Philips XP 1603 PMT; Keithley pulse amplifier (Model 165); dual-beam oscilloscope (Tektronix 556); time delay variable [4] Spectral acquisition [3]:

Transient formation:

 $e(aq)^{-} + N_2\theta (+H_2\theta) \rightarrow N_2 + 0H + 0H^{-}$ 

 $0H + CN^- \rightarrow HOC=N^-$ ,  $pK_a (HOC=NH) = 10.2$ 

[1] Spectrum was obtained from a computer simulation of the Footnotes:

OH + CN → HOC=N-

H<sub>2</sub>O + HOC=N → HOC=NH + OH

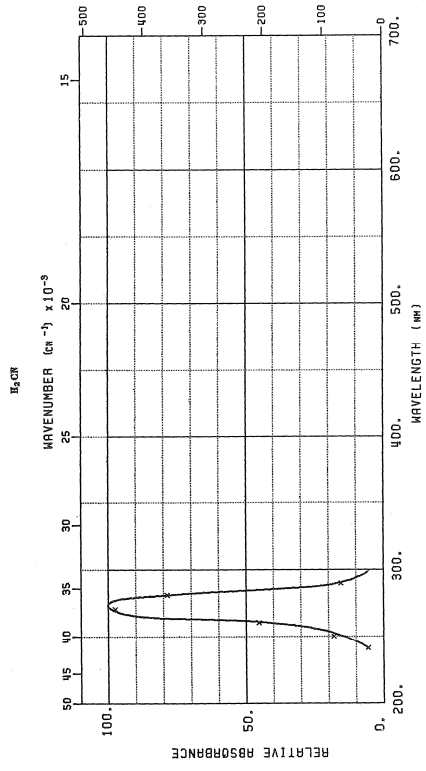
HOC=NH > 0=CNH2

system, using known rate constants.

Two other chemical systems were used at pH 9.85 and 11.4 [CN-], [N<sub>2</sub>0], and the dose were the same in the pH 9.85 system; but [CN-] = 10<sup>-1</sup> in the pH 11.4 system, and the dose was 28.1 krad in the pH 11.4 system.

B. Hurni, U. Bruehlman, and R.E. Buehler, Int. J. Radiat Phys. Chem. 7: 499-506 2

[4] Transient makes contribution to experimental spectrum only for time less than 1 µs. (1975). [3]



Transient:  $H_2GN$  [1],  $2k = (2.6 \pm 0.8) \times 10^9 L \text{ mol}^{-1} \text{ s}^{-1}$ System: agreeous solution of  $10^{-1} \text{ mol L}^{-1} \text{ HCN}$ , pH 1.9

Reference: H. Buechler, R.E. Buehler, and R. Cooper, J. Phys. Chem. 80(14): 1549-53 (1976)

pulse radiolysis, using Febetron 705 (Field Emission Corp.); energy 2 MeV; dosimetry using calorimetry; dose = 34.8 krad Transient generation [2]:

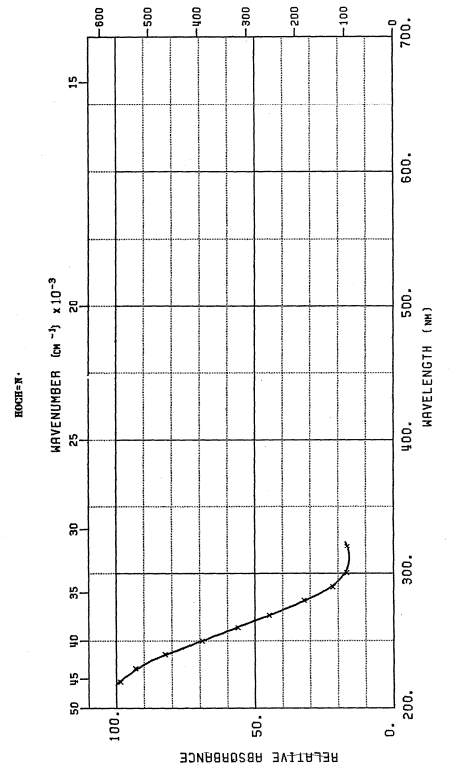
xenon XBO 450 monitoring lamp; Bausch 8 Lomb High Intensity monochromator; Philips XP 1003 PMT; Keithley pulse amplifier (Model 105); dual-beam oscilloscope (Tektronix 556) Spectral acquisition [2]:

Transient formation:

e(aq) - + H+ + H

H + HCN → H2CN

[1] Spectrum determined using computer optimization of some of the rate constants plus computer analysis of spectra from the pH 1.9 HGN system and other acidic HCN solutions which were saturated with N<sub>2</sub>O.
[2] B. Hurni, U. Bruchlman, and R.E. Buchler, Int. J. Radiat. Phys. Chem. 7: 499-506 (1975). Footnotes:



System: aqueous solution of 10-1 mol L-1 HCN, pH 2.85, saturated with N20 Transient:  $HOCH=N \cdot [1]$ ,  $2k = (2.8 \pm 0.5) \times 10^{9} \text{ L mol}^{-1} \text{ s}^{-1}$ 

## HOCH=N.

Reference: H. Buechler, R.E. Buehler, and R. Cooper, J. Phys. Chem. 80(14): 1549-53 (1976)

pulse radiolysis, using Febetron 705 (Field Emission Gorp.); energy 2 ReV; dosimetry using calorimetry; dose = 34.8 krad Transient generation [2]:

xenon XBO 450 monitoring lamp; Bausch 3 Lomb High Intensity monochromator; Philips XP 1003 PMT; Keithley pulse amplifier (Model 105); dual-beam oscilloscope (Tektronix 556) Spectral acquisition [2]:

Transient formation:

 $e(aq)^{-} + N_2O (+H_2O) \rightarrow N_2 + OH + OH^{-}$ 

OH + HCN - HOCH=N., pKa (HOCH=N.) = 5

 [11] Spectrum corrected for OH absorption and the H atom reaction product, H<sub>2</sub>CN.
 This was done using computer simulation.
 [21] B. Hurni, U. Bruchiman, and R.E. Buchler, Int. J. Radiat. Phys. Chem. 7: 499-506 Footnotes:

(1975).