Luminescent Photoproducts in UV-Irradiated Ice

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ABSTRACT

This Account describes the near-UV and visible luminescences emitted from crystalline, polycrystalline, and amorphous ices as a result of excitation by UV light. Vibrationally resolved, short-lived luminescence around 340 nm arises from excited O₂ formed by the reaction of two O atoms. Long-lived luminescence around 420 nm is tentatively assigned to a spin-forbidden ${}^{4\Sigma^{-}} \rightarrow X^{2}\Pi$ transition of OH. This Account gives a history of the research into this little-known phenomenon, places it in the context of other spectroscopic studies of gaseous and solid water, and proposes future directions for the work.

I. Introduction

The photochemical and photophysical properties of water ice are of such importance and complexity that research interest in this area is increasing. The complexity of the photochemistry and photophysics of ice arises for two reasons. First, the aggregation of isolated water molecules into a solid matrix perturbs their spectroscopic characteristics significantly. Second, significant quantities of photoproducts produced by initial irradiation are trapped in the ice matrix and strongly alter the spectroscopic characteristics of the host, leading to additional absorption features and an even more diverse set of photoproducts. Consequently, the photolysis of ice into hydroxylic species is more probable than would at first be expected from a

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Terry Quickenden graduated with a Doctorate from the University of Canterbury in New Zealand in 1967 and has published extensively on the luminescence properties of gaseous, liquid, and solid water over the last 30 years. Another area of interest has been that of the ultraweak luminescences of biological systems. In addition, he has contributed significantly to the literature of electrochemistry—particularly in the areas of photoelectrochemistry and solar energy conversion. substance that is often (mistakenly) perceived as being completely transparent from 200 to 700 nm. Photoproducts identified in UV-irradiated ice include $O_{2,1}^{1}$ atomic H,² OH,^{2,3} and H₂O₂.²

Interest⁴ in the spectral properties of water ice has received a substantial impetus in recent years from its identification on outer solar system bodies (such as Jupiter's Galilean satellites, Europa, Ganymede, and Callisto) and in comets, which are largely comprised of "dirty" ice. The occurrence of ice in extraterrestrial environments is of particular importance because it could allow life to be supported at these remote locations on a permanent basis. Optical techniques for remote identification of ice currently use reflectance spectroscopy, but luminescence spectroscopy is also a possibility.⁴

This Account brings together the published work on the luminescence of UV-excited H_2O and D_2O ices and examines the implications that such work has for the identification and monitoring of photoproducts in an ice matrix. The Account does not cover the luminescences produced in ice excited by electrons and other high-energy radiations. It is believed⁵ that the emissions caused by such high-energy radiation result from different mechanisms than those produced by light of wavelength greater than 200 nm, which are the subject of the present Account.

Methods for the Detection of Photogenerated Species in Ice. Definitive assignment of features in the absorption, luminescence, or electron paramagnetic resonance (EPR) spectra of ice is often difficult for several reasons. First, the ice lattice often significantly perturbs photoproducts, such that their absorption and emission lines shift and broaden and their electronic *g* values change. Second, broadening in optical absorption and/or luminescence spectra often obscures fine structure.

Definitive identification of photoproducts may be assisted by the use of several techniques. This strategy was employed to assign the broad absorption band at 280 nm in electron-excited ice (Figure 1a) to the hydroxyl (OH) radical,⁶ because the absorption intensity decayed with temperature according to the known^{6,7} behavior of EPRmonitored OH. Unfortunately, similar correlations have not yet been made for UV-excited ice.

Ghormley and Hochanadel² flash-photolyzed ice at 263 K using a pulsed xenon lamp. Their transient absorption spectra (shown in Figure 1b) revealed H[•] and OH[•] photoproducts, the former only on the microsecond time scale. HO₂ was not observed. Soon after, Kuwata et al.⁸ used EPR to detect photoproducts formed after photolysis of ice by 121.6-nm UV light at 77 K. Their assignment⁸ of the EPR to HO₃ is perplexing, since the EPR of high-energy-irradiated ice has not revealed such species.^{9,10} Moreover, their EPR spectra of irradiated water vapor,⁸ where the formation of HO₃ would be unlikely, look remarkably similar to those of ice. Clarification of this work is necessary. Nevertheless, these reports act as useful point-

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FIGURE 1. (a) UV absorption spectrum of electron-irradiated ice at 175 K. Adapted from ref 6 with permission from Dr. I. A. Taub and the American Institute of Physics. (b) Transient absorption spectra of flash-photolyzed ice at 263 K, obtained 40 μ s and 8 ms after the flash. Reprinted with permission from ref 2. Copyright 1971 the American Chemical Society.

ers to the assignment of luminescent photoproducts, although care must be taken in relating these data to the luminophores. For example, species detected by EPR or absorption may not luminesce, or, alternatively, the luminophore may be EPR-inactive or may be below the detection limit of EPR and/or optical absorption spectroscopy. Hence, it is important to relate the UV-excited luminescence of ice to relevant data from a range of sources, including EPR spectroscopy and gas-phase and matrix-isolation optical spectroscopy. Such comparisons form part of this Account.

Our Account is arranged as follows. Section II gives an overview of the ice luminescence literature, dealing sequentially with the distinct luminescence features. New thoughts on the interpretation of the experimental data are offered in section III. In section IV, we summarize the current state of knowledge and suggest future directions for the work on the luminescence of UV-excited ice.

II. History of Studies of Luminescence from UV-Irradiated Ices

Prior to 1970, reports^{11–14} of luminescence from UVexcited ice arose from qualitative, visual observations, and neither spectroscopic nor kinetic measurements were made. The water purity was not specified in these early works, and thus it cannot be ascertained whether the luminescence originated from the ice or organic impurities therein.



FIGURE 2. Luminescence spectra of UV-irradiated crystalline H_2O ice at 88 K: (a) 220-nm and (b) 260-nm excitation. Accumulation of luminescence intensity is shown as a function of the irradiation time in hours. Reprinted with permission from ref 16. Copyright 1985 Elsevier Science.

In 1970, Vierke and Stauff¹⁵ obtained the first quantitative luminescence spectra of ice in experiments designed to detect long-lived (>5 ms) luminescence after 290-nm flash irradiation at 77 K. The luminescence bandwidth was broad and extended from 390 to 600 nm. Superimposed on it were two weak, high-energy peaks and two more intense, low-energy peaks. Vierke and Stauff used highly purified water in their experiments. Hence, their proposal that the observed luminescence was due not to ice, but to the presence of two unknown organic impurities, either airborne or residing on the walls of the sample cuvette, is rather surprising.

Since the work of Vierke and Stauff,¹⁵ UV-excited ice luminescence has been investigated primarily by the Quickenden research group at the University of Western Australia.^{1,3,16–19} Results from our first publication¹⁶ are shown in Figure 2. Broad excitation bands at about 220 and 260 nm in crystalline ice gave rise to broad emission bands at about 340 and 420 nm, respectively.¹⁶ These features have appeared consistently in later works, and we retain these wavelength values as subsection titles and band labels, even when their fine-structure components have been resolved or when the peak wavelength shifts slightly with different ice morphology.

Luminescent photoproducts are produced by photolysis of H_2O in the ice lattice and by subsequent photolysis or reactions of these photoproducts. Photolysis occurs at wavelengths greater than 200 nm, despite the fact that the peak absorption of the dissociative ${}^{1}A_{1}$ state of H_2O occurs below 150 nm,²⁰ because of a weak, broad absorption tail.^{21,22} A similar tail is observed for H₂O in rare-gas matrixes.²³ Thermodynamically,¹ dissociation of H₂O in ice can occur for wavelengths as long as ~270 nm, due to stable trapping of the OH photoproduct in the ice lattice. Hence, ice absorption increases with irradiation time since more strongly absorbing photoproducts, such as OH, accumulate in the ice lattice.

Quickenden and co-workers have given a detailed description of their triple distillation procedure for purifying water.²¹ The water has properties characteristic of ultrapure water: a high surface tension,²¹ a low level of particulates,²¹ and very low UV^{21,24} and visible²⁴ absorptivities (<1 ppb of UV-absorbing impurities). Great care was taken to avoid water contamination when preparing crystalline ice and depositing amorphous and polycrystalline ice under high-vacuum conditions.

Ice Morphology. Three ice morphologies are encountered here: crystalline hexagonal ice I_h , polycrystalline ice I_h , and amorphous (or vitreous) ice.²⁵ The latter is prepared only by very slow deposition of water vapor onto a surface held at <130 K and is thermally unstable. At higher deposition rates and/or warmer temperatures, polycrystalline ice is formed. Crystalline ice I_h is formed at atmospheric pressure at temperatures from 0 to 273 K. Luminescences from the different ice morphologies will be treated together, since the small variations do not require different interpretations. The primary changes³ are an increase in luminescence quantum efficiency from crystalline to polycrystalline to amorphous ice and a change in the relative intensities of the 340- and 420-nm emissions.

Small- to medium-sized water clusters of ice (or nanocrystals) show completely different luminescence behavior²⁶ compared to the bulk forms of ice. Irradiation of water clusters containing up to ~1500 molecules with UV photons from a synchrotron source²⁶ yielded fluorescence excitation spectra similar to those of the free water molecule, rather than bulk ice.^{1,16} The proposed mechanism²⁶ is given in eq 1. *n* is the original number of water

$$(H_2O)_n + h\nu \rightarrow (H_2O)_n^* \rightarrow (H_2O)_{n-1} + H_2O^* \rightarrow (H_2O)_{n-1} + OH(A^2\Sigma^+)^* + H$$
 (1)

molecules in the cluster. The excited OH then emits its characteristic luminescence.

The 340-nm Luminescence of Ice. The 340-nm luminescence of UV-excited ice is excited predominantly at 220 nm (Figure 2).¹⁶ Early work on crystalline H₂O ice by Litjens and Quickenden¹⁷ suggested that the luminescence arose from the $A^{2}\Sigma^{+} \rightarrow X^{2}\Pi$ transition of OH, red-shifted from the gas phase²⁷ because of the modified Franck– Condon factors in ice that concomitantly blue-shift the excitation maximum.⁶ Both the luminophore's accumulation with increased irradiation time¹⁶ and its thermal behavior¹⁷ were similar to those of EPR-detected OH in high-energy-irradiated ices.^{7,9} The luminescence lifetime was shorter than the 12- μ s lower limit of the instrumentation,¹⁸ which was consistent with an assignment to OH, since OH has a gas-phase lifetime of ~700 ns.²⁸ The



FIGURE 3. Better resolved luminescence spectra of crystalline H₂O and D₂O ice at 78 K, excited by 260-nm light. Features at wavelengths less than ~400 nm are attributed to the Herzberg I or III systems of O₂. Transitions from the upper to lower vibrational states for the Herzberg III system are indicated, as is the tentative assignment of the long-lived 420-nm luminescence to the OH ${}^{4}\Sigma^{-} \rightarrow X {}^{2}\Pi$ transition. Reprinted with permission from ref 1. Copyright 1993 the American Chemical Society.

observed linear dependence of luminescence intensity on sample thickness in amorphous and polycrystalline ice indicated a homogeneous distribution of luminophores through the sample.¹⁸

Figure 3 shows the later work of Matich et al.,¹ which resolved structure not previously observed^{16,17} in the UVexcited luminescence of crystalline ice. The broad 340nm band^{16,17} is comprised of four evenly spaced bands that are insensitive to deuterium substitution (Figure 3),¹ and hence the earlier ad hoc assignment^{17,29} to OH $A^{2\Sigma^{+}}$ $\rightarrow X^2 \Pi$ is invalidated. The structured luminescence was attributed to either the Herzberg I A ${}^{3}\Sigma_{u}^{+} \rightarrow X {}^{3}\Sigma_{g}^{-}$ or Herzberg III $C {}^{3}\Delta_{u} \rightarrow X {}^{3}\Sigma_{g}^{-}$ systems of O₂ because of the favorable vibrational spacings and their insensitivity to deuterium substitution. The $C {}^{3}\Delta_{u} \rightarrow X {}^{3}\Sigma$ transition is dominant in the near-UV luminescence spectra of matrixisolated O₂³⁰ and may be the more likely candidate. The ¹⁶O₂ luminescence lifetime in N₂ matrixes decreases from \sim 250 μ s at 4 K to \sim 25 μ s at 14 K.³⁰ Thus, at 77 K, the lifetime is likely to be less than 12 μ s, the upper limit obtained for the ice luminescence lifetime.¹⁸ In ice, the O_2 luminescence is a chemiluminescence of O_2 , the intensity growing as O2 forms. It arises neither from excitation of O₂ trapped in the ice lattice during deposition, nor from re-excitation of the O₂ photoproduct.¹ Two mechanisms for formation of O2* involve either an O atom and H₂O₂ or two O atoms.¹ The latter is favored due to higher exothermicity. A generalized scheme is shown in eq 2.¹ In this scheme, OH_T and O_m are, respectively,

$$H_2O + h\nu \rightarrow H + OH_T \rightarrow H + O_m \xrightarrow{X_T} O_2^* \rightarrow O_2 + h\nu'$$
(2)

trapped OH radicals and mobile O atoms. X_T is trapped H₂O₂ or O. Predissociation in the $A^2\Sigma^+$ state of OH can account for efficient formation of O atoms in UV-irradiated ice (section III).

			luminescence lifetime/s		
sample	temperature/K	band maximum/nm	component a	component b	ref
crystalline ice	88	420	1.7 ± 0.2		29
0	77	430	5		32
	77	417	1.9		1
amorphous ice	78	410, 475	1.6 ± 0.1	4.2 ± 0.3	3
polycrystalline ice	78	420	0.8 ± 0.1	2.5 ± 0.4	19
0.1 M NaOH/H ₂ O	77	400	0.9	2.9	33
	?	400	1.35		35
8 M NaOH/H ₂ O	77	400	1.1		34
0.1 M NaOH/MeOH	?	400			35
0.1 M NaOD/D ₂ O	?	390	1.55		35
8 M KOH/H ₂ O	77	400	0.9		34
KBr:OH ⁻	77	400	0.63	0.2	33
Ba(OH) ₂ ·2H ₂ O	77	400	2.0		34
10 M H ₂ SO ₄ /H ₂ O	?	410	1.2		35
$1 \text{ M H}_3\text{PO}_4/\text{H}_2\text{O}$?	410	1.2		35
10 M HF/H ₂ O	?	410	1.2		35

 Table 1. Characteristics of Long-Lived Luminescences from UV-Excited Pure and Doped Ices and Metal

 Hydroxide Crystals

The above evidence^{1,16–18,29} strongly supports assignment of the 340-nm luminescence to the Herzberg I or III systems of O_2 .¹ The temperature dependence^{17,29} of this band follows closely that of the OH radical in ice, because OH is the source of O atoms. However, it is unclear whether other broad luminescence bands having lifetimes and thermal behavior similar to those of O_2 might underlie the O_2 emission. The OH $A \, {}^2\Sigma^+ \rightarrow X \, {}^2\Pi$ luminescence is a possible candidate, although in this case isotopic effects would have been expected and should have been detected, unless the OH emission was only a small component of the total luminescence.

The 420-nm Luminescence of Ice. Quickenden and coworkers¹⁶ first observed the 420-nm luminescence of UVirradiated crystalline H_2O ice at 88 K and found that it was excited primarily at 260 nm (Figure 2). The band was blue-shifted by 1.4 nm in D_2O ice.¹ In amorphous ice, the band was resolved into two components centered at 410 and 475 nm.³

The luminescent photoproduct (or its precursor) that leads to the 420-nm emission was found to accumulate²⁹ in the ice lattice at liquid N_2 temperature, where it survived for at least several hours and was re-excited by further irradiation.

The 420-nm luminescence has a long lifetime of the order of 1 s but varies somewhat with ice morphology (Table 1). The lifetime is insensitive to deuterium substitution in crystalline ice.³¹ Early work^{1,29} fits the data to second-order kinetics, but the identical rise and fall of luminescence intensity³ is better represented by biexponential kinetics.³ The two components of the luminescence derived from the biexponential fit are tentatively attributed to the same luminophore occupying substitutional and interstitial sites in the ice lattice,³ with the former, shorter-lived component persisting at higher temperatures.¹⁹ Arrhenius treatment of the temperature dependence of the luminescence decay in polycrystalline ice¹⁹ showed that the decay had negligible activation energy, which is consistent with the rate-determining step being an electronic transition, rather than a chemical step.

Khusnatdinov and Petrenko³² have detected similar 420-nm luminescence from crystalline ice excited between

180 and 300 nm, but they have not reported the 340-nm emission. Their decay data were fitted with a single-exponential expression, yielding a lifetime of 5 s, which is somewhat longer than our data for crystalline ice (Table 1).^{1,29}

The 420-nm luminescence was originally tentatively assigned to the OH A ${}^{2}\Sigma^{+}$ (v = 0) $\rightarrow X {}^{2}\Pi$ (v = 2) transition.¹⁷ Later work showed that this assignment was unsatisfactory because the band energy was almost the same for H and D isotopes,1 contrary to expectations.27 Furthermore, there was no isotopic effect on luminescence lifetime.³¹ Khusnatdinov and Petrenko³² assigned the luminescence to decay from a low-lying triplet state of H₂O, but this assignment is very unlikely according to work by Quickenden and Irvin.²¹ A more recent and much more satisfactory assignment³ has been to the spinforbidden ${}^{4}\Sigma^{-} \rightarrow X^{2}\Pi$ transition of OH, which is likely to be long-lived. Maria and McGlynn³³ first made the ${}^{4}\Sigma^{-} \rightarrow$ $X^2\Pi$ assignment to the long-lived 400-nm luminescences from hydroxide-doped alkali halides and frozen aqueous metal hydroxide solutions excited by UV light. Their data are summarized in Table 1. Subsequent literature, while disagreeing with this assignment,^{34,35} has not provided a convincing alternative.^{29,31} Quickenden and co-workers^{3,19} have adopted the ${}^{4}\Sigma^{-} \rightarrow X^{2}\Pi$ assignment for the 420-nm luminescence of UV-excited ice because the negligible activation energy and long lifetime suggest that an electronic transition is rate-determining.

Although the ${}^{4\Sigma^{-}}$ state of OH is dissociative in the gas phase,^{36,37} Quickenden et al.^{3,19} propose that it becomes bound in the ice lattice. This proposal does not appear unreasonable because the ice lattice strongly perturbs OH (section III). A high-level calculation of the ${}^{4\Sigma^{-}}$ state's potential energy (PE) curve in ice and/or direct detection of the ${}^{4\Sigma^{-}}$ state by EPR are necessary before a definitive statement can be made regarding the assignment's validity.

III. New Perspectives

In this section, the ice luminescence results of section II are interpreted with the help of the well-known gas-phase



FIGURE 4. Potential energy curves for OH taken from ref 40, except for the dotted curve, which is a hypothetical curve for a bound ${}^{4}\Sigma^{-}$ state in ice. The region of the important crossings is enlarged for clarity.

predissociations of OH $A^{2}\Sigma^{+}$. This is then followed by an attempt to draw together the optical and EPR experimental results for OH in a variety of environments as we seek to understand the absence of the normally ubiquitous $A^{2}\Sigma^{+} \rightarrow X^{2}\Pi$ luminescence when OH is embedded in ice.

Predissociation of OH *A* ²Σ⁺. In the gas phase, repulsive ⁴Σ⁻, ²Σ⁻, and ⁴Π states of OH efficiently predissociate the *A* ²Σ⁺ state for vibrational levels *v* > 1 (Figure 4).³⁷⁻⁴⁰ The ⁴Σ⁻ state is responsible for the lowest-energy predissociation⁴⁰ of the *A* ²Σ⁺ state at ~38 000 cm⁻¹ (262 nm),²⁸ with a predissociation probability that is a few times greater than the radiative decay probability.³⁹ The higherlying ²Σ⁻ and ⁴Π states are ~900 cm⁻¹ apart³⁷ and predissociate the *A* ²Σ⁺ state at *v* ≈ 5, or ~46 000 cm⁻¹ (217 nm) above the ground state.^{37,38,40} The predissociation induced by the ⁴Π state is very strong,³⁸ reducing the lifetimes of rovibrational levels to ~10⁻¹¹−10⁻¹² s.^{38,41} Because of the small separation between the ²Σ⁻ and ⁴Π states and the broad OH absorption bands in ice, we do not consider the ²Σ⁻ predissociation here.

The situation for OH in ice may well be somewhat modified. If the ${}^{4}\Sigma^{-}$ state is made associative by the ice matrix, as proposed by Quickenden et al.,^{3,19} a reasonably efficient pathway for the preparation of OH(${}^{4}\Sigma^{-}$) should be accessible at ~260 nm, although the crossing energy may be shifted from its gas-phase value. Dissociation of OH is prevented if the crossing point of the $A {}^{2}\Sigma^{+}$ and ${}^{4}\Sigma^{-}$ states corresponds to a bound portion of the ${}^{4}\Sigma^{-}$ PE curve. With these constraints in mind, a possible form for a bound ${}^{4}\Sigma^{-}$ state is shown as the dotted curve in Figure 4. Figure 5 shows a schematic Jablonski diagram summarizing the proposed pathway to the luminescence, which terminates at $v \approx 4$ of the $X {}^{2}\Pi$ state.³³ A calculation of the PE curves of OH in ice would, indeed, be most helpful but is not available at present.



FIGURE 5. Schematic Jablonski diagram showing the proposed processes leading to OH ${}^{4}\Sigma^{-} \rightarrow X {}^{2}\Pi$ luminescence in ice and the absence of $A {}^{2}\Sigma^{+} \rightarrow X {}^{2}\Pi$ luminescence.

The strong predissociation of the $A^{2}\Sigma^{+}$ state by the ${}^{4}\Pi$ state ensures that a large number of O atoms are formed when OH is excited to $A^{2}\Sigma^{+}$ ($v \approx 5$). The yield will be moderated somewhat by small Franck–Condon overlap for OH $X^{2}\Pi$ (v = 0) $\rightarrow A^{2}\Sigma^{+}$ (v = 5) absorption, but the shift of the absorption maximum from 306 nm²⁷ in the gas phase to 280 nm⁶ in ice suggests that overlap is greater in ice. The ${}^{4}\Pi$ predissociation occurs at an energy very similar to that of the peak of the ice excitation spectrum when it is detected in the O₂ luminescence (340 nm; section II). Thus, the strong predissociation of OH $A^{2}\Sigma^{+}$ by ${}^{4}\Pi$ is very probably responsible for more efficient production of O atoms at ~220 nm than at longer wavelengths. A higher concentration of O atoms leads to more O₂* and hence to more intense 340-nm luminescence.

Predissociation of OH $A^{2}\Sigma^{+}$ by ${}^{4}\Sigma^{-}$ and ${}^{4}\Pi$ states supports the observed¹⁶ relationship of 220- and 260-nm excitation to 340- and 420-nm emissions, respectively. At shorter wavelengths there is a higher yield of O atoms that gives rise to the O₂ luminescence around 340 nm.¹ At longer excitation wavelengths, more OH crosses to the bound portion of the ${}^{4}\Sigma^{-}$ potential curve (Figure 4), giving greater intensity to the 420-nm band assigned to the OH ${}^{4}\Sigma^{-} \rightarrow X {}^{2}\Pi$ transition.^{3,19}

The Intriguing Absence of OH $A^{2}\Sigma^{+} \rightarrow X^{2}\Pi$ luminescence. If the recent assignments of the 340- and 420-nm luminescence bands to the respective Herzberg and $4\Sigma^{-1}$ transitions are indeed correct, this removes the wellknown and often dominant $A^{2}\Sigma^{+} \rightarrow X^{2}\Pi$ emission of OH from consideration in UV-excited ice (Figure 5). Furthermore, it is now known⁵ that above 77 K the emissions from electron-excited ice are dominated by the $C {}^{1}B_{1} \rightarrow A {}^{1}B_{1}$ excimer transition of H₂O and not the $A^{2}\Sigma^{+} \rightarrow X^{2}\Pi$ transition of OH. We note that the ${}^{4}\Sigma^{-}$ predissociation of the A ${}^{2}\Sigma^{+}$ state cannot totally quench this luminescence because, at its strongest, the predissociation probability is only a few times greater than the luminescence probability.³⁹ EPR and luminescence data reveal that OH trapped in ice is very different from OH in the gas phase, where the $A^{2}\Sigma^{+} \rightarrow X^{2}\Pi$ transition is so dominant.

Table 2. Optical Absorption and Luminescence Datafor the OH $A^{2}\Sigma^{+} - X^{2}\Pi$ Transition in Various Media

		(0,0) luminescence				
medium	(0,0) absorption/ nm	maximum/ nm	lifetime/ ns	quantum efficiency		
gas phase	306.4^{27}	306.427	693 ± 20^{28}	~ 1		
Ne matrix ⁴⁶	308.1	309.1	582 ± 10	>0.9		
Ar matrix ⁴⁷	310.8	340	475 ± 10	>0.9		
Kr matrix ⁴⁷	а	363	450 ± 20	>0.9		
Xe matrix ⁴⁷	а	440	100 ± 4	>0.9		
KCl ³³	b	300	С	С		
KBr ³³	b	311	С	С		
ice	280 ⁶	d	d	d		

^a Not stated. ^b Below detection limit. ^c Not measured. ^d Absent.

Consider first the EPR data. The EPR of gas-phase OH gives an orbital angular momentum g value (g_l) of unity for the electronic ground state of OH.⁴² Detection of OH in ice by EPR after β -,^{7.9} γ -,^{43,44} or X-irradiation¹⁰ is well documented. The measured g value is very close to that of a free electron, and thus the orbital angular momentum is almost completely quenched^{9,43} ($g_l \approx 0$). Quenching occurs because hydrogen bonding of the OH radical to H₂O molecules of the ice lattice lowers the degeneracy of the OH p_x and p_y orbitals.^{43,45}

Consider now the $A^{2}\Sigma^{+} \rightarrow X^{2}\Pi$ luminescence of OH, which might be expected from UV-irradiated ices because H and OH radicals are probably the only primary photoproducts formed at excitation wavelengths greater than 200 nm.¹ The abundance of rovibrational levels in $A^{2}\Sigma^{+}$ should mean that once formed, OH would be relatively easily excited into one of these levels and the characteristic luminescence observed. This is *not* the case in ice, as we have seen above.

Table 2 summarizes the optical luminescence data for OH $A^{2}\Sigma^{+} \rightarrow X^{2}\Pi$ in a variety of environments. The wellknown gas-phase $A^{2}\Sigma^{+} \rightarrow X^{2}\Pi$ emission of OH^{27,28} is observed when OH is embedded in rare-gas matrixes.^{46,47} The luminescence quantum yield remains high despite OH being increasingly perturbed by weak hydrogen bonding to a rare-gas atom⁴⁷ along the series Ne,⁴⁶ Ar,⁴⁷ Kr,⁴⁷ and Xe.⁴⁷ OH luminescence is also observed from UVirradiated alkali halide crystals containing a small amount of alkali hydroxide.³³ The (0,0) and (0,1) transitions are little shifted from their gas-phase values.

The absence of the OH $A^{2}\Sigma^{+} \rightarrow X^{2}\Pi$ emission in ice is not due to the crystal field imposed by the ice lattice, because then quenching would occur to some extent in alkali halides and rare-gas matrixes. OH in Ar matrixes exhibits partial quenching of orbital angular momentum in some sites,⁴⁸ due very likely to weak hydrogen bonding of OH to an Ar atom, which is insufficiently strong to quench luminescence.⁴⁷ OH $A^{2}\Sigma^{+} \rightarrow X^{2}\Pi$ emission is apparently quenched in ice by strong hydrogen bonding of OH to water molecules of the ice lattice. The quenchings of both luminescence and orbital angular momentum are evidently related.

Miyazaki et al.^{49,50} have observed OH $A {}^{2}\Sigma^{+} \rightarrow X {}^{2}\Pi$ emission from electron-irradiated ice at temperatures lower than 77 K. However, this emission arises from OH

 $A^{2}\Sigma^{+}$ formed by dissociative recombination of $H_{2}O^{+}$ and e^{-} , which is significant only below 77 K.⁵⁰ The emitting OH radicals are not the stably trapped OH observed by EPR, because they do not show the known persistence to temperatures above 100 K.^{6,7}

Finally, since the hydrogen bonding of OH to H₂O in ice has a profound effect on both the orbital angular momentum of OH ($X^{2}\Pi$) and the $A^{2}\Sigma^{+} \rightarrow X^{2}\Pi$ luminescence, it may indeed not be surprising that the repulsive ${}^{4}\Sigma^{-}$ state becomes bound.^{3,19}

IV. Summary and Outlook

We have reviewed the published studies of UV-excited ice luminescence. The assignment of the 340-nm luminescence band to the Herzberg I or III systems seems entirely reasonable. The resolved structure has spacings corresponding to O₂ vibrations, the spacings are not affected by deuterium substitution for hydrogen, and the lifetime is in the correct range. Measurement of the decay kinetics should reveal the luminescence lifetime and provide a new test of the mechanism for O2 production. Excitation at \sim 220 nm leads to efficient predissociation of OH, producing O atoms that react to give chemiluminescent O_2 . Hence, the temperature dependence of the O₂ luminescence is similar to that of the OH radical. It remains to be seen whether other luminescence bands underlie the O₂ structure. If they do, their temperature dependence is evidently similar to that of O_2 , and they, like O_2 , have a lifetime shorter than 12 μ s. Measurement of the temperature dependence of the luminescence decay may answer this question.

Assignment of the 420-nm luminescence remains less certain. Many aspects of its behavior could correspond to the spin-forbidden ${}^{4}\Sigma^{-} \rightarrow X {}^{2}\Pi$ transition of OH. The ${}^{4}\Sigma^{-}$ state is dissociative in the gas phase but is proposed to become associative in ice. The long luminescence lifetime and zero activation energy for the process provide support for this proposal. However, the temperature dependence of the band differs from that of OH by steadily decreasing with increasing temperature, whereas OH detected by EPR disappears very suddenly between 100 and 130 K. Early work proposed that a stable precursor to OH, for example H_2O_2 , could be responsible for such behavior. Soon, we intend to use EPR as a probe for OH and other EPR-active photoproducts. Support for a bound ${}^{4}\Sigma^{-}$ state of OH embedded in ice might also be provided by high-level theoretical calculations.

The absence of OH $A^{2}\Sigma^{+} \rightarrow X^{2}\Pi$ luminescence in ice is intriguing and warrants further study. Shortly, we will undertake matrix-isolation experiments to probe the OH luminescence behavior when it is embedded in water clusters of varying size. These experiments should provide valuable insight into the nature of the luminescence quenching and possibly show the ${}^{4}\Sigma^{-}$ emission in the hydrogen-bonded environment. Measurement of the UVexcited luminescence of water clusters embedded in solid, inert environments may provide data in closer agreement with those from bulk ice than were obtained by transient excitation of gas-phase clusters.

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