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\[ \Phi_{\text{H}_2\text{O}_2} \times 10^{-4} \]

Irradiation time (min)

\[ \text{H}_2\text{O}_2 \text{ (nM)} \]

41x37mm (300 x 300 DPI)
Photochemically produced hydrogen peroxide (H$_2$O$_2$) can negatively affect microbial activity and play an important role in many chemical reactions involving organic matter or metals in marine waters. To model production rates, wavelength and temperature dependent H$_2$O$_2$ production rates were determined as a function of absorbed irradiance (apparent quantum yields) for diverse marine environments from the Southern Ocean, Pacific Ocean, Atlantic Ocean and Gulf of Mexico. Apparent quantum yields were remarkably similar among these environments irrespective of expected differences in composition or concentrations of metals and organic matter or prior light-exposure history. The comprehensive data set for H$_2$O$_2$ apparent quantum yields reported here can be used to model H$_2$O$_2$ photochemical production rates globally based on remotely sensed optical and temperature data.

One Sentence Summary (20 word max)

Wavelength, temperature and light-dose dependent hydrogen peroxide photoproduction quantum yields were determined in subtropical, temperate and polar marine waters.
Wavelength and temperature-dependent apparent quantum yields for photochemical formation of hydrogen peroxide in seawater

David J. Kieber1*, Gary W. Miller1#, Patrick J. Neale2 and Kenneth Mopper3

1State University of New York, College of Environmental Science and Forestry, Department of Chemistry 1 Forestry Drive Syracuse, New York 13210
2Smithsonian Environmental Research Center Photobiology and Solar Radiation Laboratory, Edgewater, Maryland 21037
3Old Dominion University Department of Chemistry and Biochemistry Norfolk, Virginia 23529

*Corresponding author (djkieber@esf.edu)
#Current Address: Daiichi Sankyo Pharma Development, Edison, New Jersey 08837
Wavelength and temperature-dependent apparent quantum yields (AQYs) were determined for the photochemical production of hydrogen peroxide using seawater obtained from coastal and oligotrophic stations in Antarctica, the Pacific Ocean at Station ALOHA, the Gulf of Mexico, and at several sites along the East Coast of the United States. For all samples, AQYs decreased exponentially with increasing wavelength at 25 °C, ranging from $4.6 \times 10^{-4} - 10.4 \times 10^{-4}$ at 290 nm to $0.17 \times 10^{-4} - 0.97 \times 10^{-4}$ at 400 nm. AQYs for different seawater samples were remarkably similar irrespective of expected differences in the composition and concentrations of metals and dissolved organic matter (DOM) and in prior light exposure histories; wavelength-dependent AQYs for individual seawater samples differed by less than a factor of two relative to respective mean AQYs. Temperature-dependent AQYs increased between 0 and 35 °C on average by a factor of 1.8 per 10 °C, consistent with a thermal reaction (e.g., superoxide dismutation) controlling $\text{H}_2\text{O}_2$ photochemical production rates in seawater. Taken together, these results suggest that the observed poleward decrease in $\text{H}_2\text{O}_2$ photochemical production rates is mainly due to corresponding poleward decreases in irradiance and temperature and not spatial variations in the composition and concentrations of DOM or metals. Hydrogen peroxide photoproduction AQYs and production rates were not constant and independent of the photon exposure as has been implicitly assumed in many published studies. Therefore, care should be taken when comparing and interpreting AQY or photochemical production rate $\text{H}_2\text{O}_2$ results from published studies. Modeled depth-integrated $\text{H}_2\text{O}_2$ photochemical production rates were in excellent agreement with measured rates obtained from in situ free-floating drifter experiments conducted during a Gulf of Maine cruise, with differences (ca. 10%) well within measurement and modeling uncertainties. Results from this study provide a comprehensive data set of wavelength and temperature-
dependent AQYs to model and remotely sense hydrogen peroxide photochemical production rates globally.
1. INTRODUCTION

Hydrogen peroxide ($\text{H}_2\text{O}_2$) is an important species in natural waters that has been intensively studied ever since Van Baalen and Marler first quantified $\text{H}_2\text{O}_2$ concentrations in the Gulf of Mexico\textsuperscript{1}. Hydrogen peroxide concentrations have been determined in a wide range of coastal and open ocean waters\textsuperscript{2-26}, wherein $\text{H}_2\text{O}_2$ has been shown to (1) adversely affect microorganisms at ecologically relevant concentrations\textsuperscript{27,28}; (2) oxidize DOM through transformations involving the Fenton reaction\textsuperscript{29,30}; (3) affect the redox chemistry of trace metals such as iron, copper, chromium and manganese\textsuperscript{31-36}; and (4) serve as a water mass tracer for vertical advection\textsuperscript{11,37-39}.

Hydrogen peroxide concentrations vary spatiotemporally and diurnally in the oceans due to several competing processes that affect $\text{H}_2\text{O}_2$ concentrations in the water column. The primary removal pathway for $\text{H}_2\text{O}_2$ in seawater is through its biological decay\textsuperscript{5,21,40,41}. The half-life for the biological decay of $\text{H}_2\text{O}_2$ in unfiltered seawater generally ranges from a few hours in coastal waters to several days in the open ocean\textsuperscript{3,21,23,41}. Although not as important as its biological loss, photochemical decomposition of $\text{H}_2\text{O}_2$ has been shown to occur in sunlight surface waters at rates that were on average 5% of photochemical production rates\textsuperscript{42}.

Microorganisms remove $\text{H}_2\text{O}_2$ from the water column, but they are also responsible for its biological production, which is expected to occur throughout the water column\textsuperscript{43}. A wide variety of algae and bacteria produce hydrogen peroxide and its precursor superoxide in culture\textsuperscript{43,44}, and biological $\text{H}_2\text{O}_2$ production has been shown to occur in oligotrophic waters under nitrogen-limiting conditions\textsuperscript{18,45}. Biological production is the main source of $\text{H}_2\text{O}_2$ deeper in the water column, but in some cases may also be important in the photic zone\textsuperscript{3,46,47}. Rain inputs are important as well\textsuperscript{3,13,48}, but they are difficult to predict and quantify. When they occur they can significantly increase $\text{H}_2\text{O}_2$ concentrations by a factor of 3 to 4 in the upper 20 m\textsuperscript{8,14,49,50}. The
importance of rain as a source of H$_2$O$_2$ in surface waters is expected to vary with latitude, as the largest rain inputs occur in the subtropics and equatorial region$^{6,23}$. The primary pathway for the formation of hydrogen peroxide in sunlit surface waters is through DOM photoreactions involving ultraviolet (UV) and visible (vis) solar radiation$^{5,51}$. Midday photochemical production rates in marine environments are in the 1-10 nM h$^{-1}$ range, with the highest rates observed in DOM-rich coastal waters and the lowest rates observed in cooler, polar waters$^{3,10,11,16,22,23,52}$. Photoproduction of H$_2$O$_2$ is proposed to involve intramolecular electron transfer reactions by excited state DOM forming radical intermediates that subsequently react with dissolved molecular oxygen to form the superoxide anion, followed by its disproportionation to form H$_2$O$_2$.$^{53}$ Early work suggested that 51-76% of the superoxide disproportionated to form H$_2$O$_2$ with an uncatalyzed, second-order rate constant of 2.2 x 10$^4$ M$^{-1}$ s$^{-1}$ in oligotrophic seawater, pH 8.3.$^{54}$ However, recent studies indicate that superoxide decay is more complicated involving metal complexes and metal-catalyzed reactions, with a smaller percentage of superoxide forming H$_2$O$_2$ than previously indicated.$^{55-59}$ Wavelength-dependent apparent quantum yields (AQYs) for the photochemical formation of H$_2$O$_2$ have been determined in seawater$^{16,22,60,61}$ to assess the importance of UV and vis radiation in H$_2$O$_2$ photoproduction$^9$ and to model photoproduction rates$^{62}$. Wavelength-dependent AQYs are remarkably similar in diverse marine waters$^{16,22,60}$, decreasing exponentially from $\sim$5 x 10$^{-3}$ at 290 nm to $\sim$0.1 x 10$^{-4}$ at 400 nm, with corresponding sunlight-normalized H$_2$O$_2$ production in marine waters primarily in the UV-B (280-320 nm) and UV-A (320-400 nm).

Building on these prior results, we conducted an extensive study to determine and compare wavelength and temperature-dependent AQYs for the photochemical production of H$_2$O$_2$ in a wide range of marine environments that included coastal and oligotrophic sites along the East
Coast of the United States, Gulf of Mexico, Hawaii, and Antarctica. Apparent quantum yields were used to calculate depth-integrated photochemical production rates that were compared to measured depth-integrated production rates obtained from in situ drifter experiments deployed at sea.

2. MATERIALS AND METHODS

2.1 Chemicals

Quinine sulfate dihydrate (Ultrex) and 30% aqueous H₂O₂ (Baker Analyzed, ACS reagent) were obtained from J.T. Baker. Reagent grade glacial acetic acid, hydrofluoric acid and 1,10-phenanthroline were obtained from Fisher Scientific. Certified ACS grade potassium oxalate monohydrate was obtained from Sigma-Aldrich. Certified ACS grade sulfuric acid and ferric chloride were obtained from Mallinckrodt. Certified ACS grade hydrochloric acid was purchased from VWR. Catalase (from bovine liver; 58,000 units mg protein⁻¹), tris (hydroxymethyl) aminomethane (tris), p-hydroxyphenylacetic acid (POHPAA) and horseradish peroxidase (type VI, 290 purpurogallin units mg⁻¹) were obtained from Sigma-Aldrich. The POHPAA crystals were further purified by three successive re-crystallizations from Milli-Q water. High purity, HPLC-grade methanol (MeOH) and acetonitrile (ACN) were obtained from Burdick and Jackson. The ethylene glycol used in the water baths for AQY determinations was spectrophotometric grade (>99% purity) from Sigma-Aldrich. Laboratory water (Milli-Q water) was obtained from a Millipore system consisting of a RiO8 reverse osmosis system to remove particles and chlorine and a Milli-Q Gradient system to remove trace ions and organic compounds (Millipore).

A 20 mM aqueous potassium ferrioxalate actinometer solution was prepared by adding 5 mL of 1.2 M aqueous potassium oxalate and 5 mL of 0.4 M aqueous ferric chloride to 90 mL Milli-Q
water in an aluminum foil-wrapped Qorpak bottle and sealed with a Teflon-lined cap. The 0.6 M, pH 4.5 acetate buffer solution used for the ferrioxalate actinometer was prepared in 0.18 M aqueous sulfuric acid. A 100 mL solution of 0.2% w/w 1,10-phenanthroline in Milli-Q water was prepared in subdued lab lighting and stored in an aluminum foil-wrapped Qorpak bottle. All actinometer solutions were stored at 4 °C.

The fluorometric reagent for H₂O₂ quantification was prepared by adding 0.5 mL of 25 mM purified POHPAA and 3.1 mg horseradish peroxidase to 49.5 mL of 0.25 M aqueous tris buffer (pH 8.80) in an aluminum foil-wrapped Qorpak bottle. The tris buffer, POHPAA solutions and fluorometric reagent were stored at 4 °C, and the solid horseradish peroxidase was stored at –20 °C. The fluorometric reagent blank increased with time, therefore, a fresh solution was prepared every three days. The tris and POHPAA solutions used to make the fluorometric reagent were stable and prepared as needed.

Stock solutions of H₂O₂ were prepared by adding 20 μL of 30% aqueous H₂O₂ to 20 mL Milli-Q water in a scintillation vial. The absorbance of the stock solution was measured in a 1 cm quartz cell and referenced with Milli-Q water using a Hewlett Packard 8453 UV-vis photodiode array spectrophotometer (Agilent Technologies). The concentration of the stock solution (10.1 mM) was determined spectroscopically using a molar absorptivity of 38.1 ± 1.4 M⁻¹ cm⁻¹ at 240 nm.

### 2.2 Hydrogen Peroxide Quantification in Seawater

Hydrogen peroxide concentrations were determined from batch fluorescence measurements of the POHPAA dimer using the method outlined in Miller and Kester, as modified in Yocis et al., and a Hitachi F-1200 fluorometer, with the excitation and emission wavelengths set at
315±7.5 nm and 400±7.5 nm, respectively. A Rainin Rabbit Plus peristaltic pump was used to pull samples through a 12 μL quartz, fluorescence flow cell using Teflon tubing (1.59 mm o.d. x 0.82 mm i.d.) and a flow rate between 0.87 to 1.0 mL min\(^{-1}\). The fluorometer was periodically calibrated with 100 nM aqueous quinine sulfate in 0.05 M aqueous sulfuric acid. The F-1200 data were collected using E-Lab data acquisition software (OMS Tech).

To quantify H\(_2\)O\(_2\) in seawater, a 100 μL aliquot of the fluorometric reagent was added to 5 mL seawater and reacted for 30 min followed by its fluorometric analysis. Unless otherwise noted, samples and the three blanks were analyzed in triplicate. The mV fluorescence responses recorded for the reacted seawater samples were corrected by subtracting the total blank mV response (see below). Blank-corrected responses were then used to calculate H\(_2\)O\(_2\) concentrations by the method of standard additions. Hydrogen peroxide standards were prepared by adding μL aliquots of a 10.1 μM standard to 5.0 mL of seawater yielding concentrations between 5.0–150 nM. The detection limit of the method, 1.3 nM, was three times the standard deviation obtained from analysis of the H\(_2\)O\(_2\) content in seven aliquots of a 0.2 μm-filtered seawater sample; the average concentration of H\(_2\)O\(_2\) in this sample was 7.1 nM.

Three blanks were analyzed to quantify the fluorescence response of (1) seawater (NAT blank) (2) catalase (CAT blank), and (3) the fluorometric reagent (FL blank). The NAT blank was determined by measuring the fluorescence of the seawater sample without addition of the fluorometric reagent or catalase. The CAT blank was determined to allow for the determination of FL blank by reacting and removing H\(_2\)O\(_2\) from the seawater sample. The CAT blank was determined by adding 25 μL of aqueous catalase (1.02 \times 10^8 units catalase L\(^{-1}\)) to 5 mL of seawater in a 20 mL borosilicate vial followed by reaction for 8 min at room temperature (CAT blank). A 100 μL aliquot of fluorometric reagent was then added to the CAT blank sample and
allowed to react for an additional 30 min (FL blank). After fluorometric analysis of these three blanks, the total blank was calculated:

\[
\text{total blank} = \text{NAT blank} + (\text{FL blank} - \text{CAT blank})
\]

All fluorescence measurements were referenced against Milli-Q water.

2.3 Seawater Sample Collection

Seawater samples were collected from stations in the confluence of the Weddell and Scotia Seas (Sta. B), along the Antarctic Peninsula at Sta. N and Arthur Harbor (Fig. 1A), and along the East Coast of the United States (Fig. 1B). The Antarctic seawater samples were collected from October 7 to November 21, 1998 during a cruise aboard the R/V Lawrence M. Gould. Gulf of Maine and stations E and F samples were obtained during a cruise aboard the R/V Endeavor from July 9 to July 27, 1999. All cruise samples were collected at 5 to 10 m with 10 L Go-Flo bottles (General Oceanics) attached to a rosette sampler equipped with conductivity, temperature and depth sensors. Seawater samples were gravity filtered directly from the Go-Flo bottles using a Whatman POLYCAP 75 AS 0.2 μm filter and stored in 4 L Qorpak bottles (capped with PTFE-lined Thermoset caps) at 4 °C until analyzed back in the home laboratory in Syracuse, NY. Prior to the cruises, the Qorpak bottles were cleaned by several rinses with MeOH and Milli-Q water followed by baking at 550 °C for 8 h. POLYCAP filters were copiously rinsed alternately with ACN and Milli-Q water until the absorbance at 220 nm and fluorescence were lowered to background levels in the Milli-Q water64.

Seawater was also collected with a Go-Flo bottle from 5 m in the Gulf of Mexico during the summer 1998 (24° 57.8’ N, 85° 53.9’ W). A surface estuarine water sample was collected using an all-polypropylene bucket in the Rhode River Estuary in March 1999. These samples were
subsequently gravity filtered through a 0.2 \( \mu \text{m} \) POLYCAP filter into pre-cleaned 4 L Qorpak bottles. Unfiltered seawater samples were collected in 4 L Qorpak bottles from Banks Channel, North Carolina (Fig. 1B) and Station ALOHA, Hawaii (22\(^\circ\) 45.0′ N, 157\(^\circ\) 58.5′ W) during the summer 1999. These samples were filtered within two days of collection through a 47 mm diameter, 0.2 \( \mu \text{m} \) Nylon filter. All samples were stored in the dark at 4 °C until they were used in irradiations experiments, which were performed from less than a day to a few months after a sample was collected; most samples were analyzed within a week of sample collection.

2.4 Apparent Quantum Yields

2.4.1 Narrow-Bandwidth AQY Determinations: To determine an AQY, a 15 mL aliquot of 0.2 \( \mu \text{m} \)-filtered, air-saturated seawater was placed in a 5 cm pathlength, rectangular quartz spectrophotometer cell and sealed with a Teflon-lined screw cap (Spectrocell). The quartz cell containing the seawater sample was placed into an enclosed temperature-controlled sample holder equipped with a stirrer. Prior to irradiation, the sample was equilibrated for 5 min to the sample holder temperature that was regulated with a re-circulating water-glycol bath. Since the water-glycol coolant was contained within the cell holder and did not bathe the quartz cell, the temperature inside the quartz cell was periodically verified with a thermistor. Likewise, the water-glycol coolant re-circulated through the cell holder and did not affect the narrow bandwidth radiation impinging on or passing through the 5 cm quartz cell. Sample irradiation times varied from 10 to 240 min, depending on the absorbance of the seawater, and the wavelength and temperature selected. Two 5 mL aliquots were removed from each 15 mL irradiated sample and equilibrated to room temperature before analysis. For each irradiated sample, a corresponding
dark control was also examined in the same apparatus by blocking the incoming light.

Wavelength-dependent \( \text{H}_2\text{O}_2 \) production rates were converted to AQYs:

\[
\Phi_\lambda = \frac{d[H_2O_2]}{dt} \frac{V}{P_\lambda (1 - 10^{-A_\lambda})}
\]  

(2)

where \( \Phi_\lambda \) is the wavelength-dependent AQY for \( \text{H}_2\text{O}_2 \) formation (mol (mol quanta\(^{-1}\)),

\( d[H_2O_2]/dt \) is the measured rate of \( \text{H}_2\text{O}_2 \) photoproduction (mol L\(^{-1}\) min\(^{-1}\)), \( V \) is volume of the

irradiated seawater sample, \( P_\lambda \) is the spectral radiant flux (mol quanta min\(^{-1}\)) determined by

ferrioxalate actinometry, and \( A_\lambda \) and \((1-10^{-A_\lambda})\) are the wavelength-dependent absorbance and

fraction of radiation absorbed by 0.2 µm-filtered seawater in a 5 cm pathlength quartz cell, respectively. Absorbance spectra were determined from 200 to 800 nm using a 5 or 10 cm

pathlength quartz cell and a Hewlett Packard 8453 UV-vis photodiode array spectrophotometer;

0.2 µm-filtered seawater sample spectra were referenced against Milli-Q water and corrected for

scattering and refractive-index baseline offsets\(^{65}\).

The apparatus used to irradiate seawater samples consisted of a 1 kW xenon arc lamp and

LPS255HR power supply (Spectral Energy). Wavelengths were selected from 290 to 400 nm

using a Spectral Energy GM 252 high intensity quarter meter grating monochromator. The lamp,

monochromator and cell holder were held fixed along an optical rail so that the beam of narrow-

bandwidth radiation leaving the monochromator was perpendicular to (and smaller in diameter

than) the front window of the 5 cm quartz cell; the radiation passed through the front window and

exited out the rear window. Unless otherwise noted, for all AQY determinations the bandwidth

was set at 9.9 nm for wavelengths < 320 nm and 19.5 nm for wavelengths from 340 nm to 400

nm. For irradiations at wavelengths \( \geq 360 \) nm, a long-pass filter with a 314 nm cut-off was

placed between the monochromator and the sample holder to filter out shorter wavelengths due to
frequency doubling. The transmission spectrum for the long-pass filter is given in Miller\textsuperscript{66}. The spectral output of the irradiation system was periodically checked using an OL754 spectroradiometer (Optronics Laboratories) that was calibrated with a 1kW NIST-traceable OL 200EA quartz-halogen lamp\textsuperscript{66}.

Temperature-dependent AQYs were determined for selected seawater samples at 290, 300, 320, 360 and 400 nm. Four temperatures (0, 15, 25 and 35 °C) were examined at each wavelength. The activation energy was calculated from linear regression analysis of the equation:

\[
\ln \Phi_\lambda = \ln A - \frac{E_a}{RT}
\]  

(3)

where $\Phi_\lambda$ is the wavelength-dependent AQY, $A$ is the pre-exponential factor, $E_a$ is the activation energy (kJ mol\textsuperscript{-1}) for the formation of hydrogen peroxide, $R$ is the universal gas constant ($8.315 \times 10^{-3}$ kJ mol\textsuperscript{-1} K\textsuperscript{-1}), and $T$ is temperature (K).

For all laboratory AQY studies, spectral radiant fluxes were determined using the potassium ferrioxalate chemical actinometer\textsuperscript{67,68} and the procedure outlined in White et al.\textsuperscript{69}. Absorption values for the irradiated actinometer samples were referenced against non-irradiated potassium ferrioxalate blanks. From known ferrous production quantum yield values\textsuperscript{68}, spectral radiant fluxes ($P_\lambda$) were calculated from:

\[
P_\lambda = \frac{AV_1V_310^{-3}}{\varepsilon \Phi_\lambda tV_2L}
\]  

(4)

where $A$ is the absorbance at 510 nm of the blank-corrected irradiated actinometer solution, $V_1$ is the volume of irradiated actinometer solution (15 mL), $V_2$ is the volume of the irradiated solution taken for spectrophotometric analysis (0.5 mL), $V_3$ is the final volume of the actinometer prepared for spectrophotometric analysis (10 mL), $10^{-3}$ is a volume conversion factor, $\varepsilon$ is the molar absorption coefficient for ferrous phenanthroline at 510 nm ($1.11 \times 10^4$ M\textsuperscript{-1} cm\textsuperscript{-1}), $\Phi_\lambda$ is
the wavelength-dependent quantum yield for ferrous iron production at wavelength $\lambda$, $t$ is the irradiation time (min) and $L$ is the pathlength (1 cm) of the quartz cell used to measure the absorbance of the phenanthroline complex at 510 nm.

2.4.2. Polychromatic AQY Spectrum Determination: In addition to wavelength-dependent AQY spectra determined with stored seawater in the laboratory in Syracuse, NY employing a narrow-bandwidth irradiation system, a wavelength-dependent AQY spectrum was also determined at the Smithsonian Environmental Research Center with freshly collected seawater using a polychromatic irradiation system described by Cullen et al.\textsuperscript{70} and Neale and Fritz\textsuperscript{71}. For the present study, the polychromatic irradiation system was modified to hold capped quartz vials and there were only five slots per filter treatment instead of ten as shown in Neale and Fritz\textsuperscript{71}. Detailed diagrams and associated text for the polychromatic irradiation apparatus are presented in Neale and Fritz\textsuperscript{71}. Briefly, the irradiation system consisted of a 2500 W Xe arc lamp with its output focused onto a mirror that redirected the polychromatic irradiation vertically 90° where it passed through a panel of eight long-pass filters with cut-offs at 280, 295, 305, 320, 335, 350, 370, or 395 nm. The sample temperature was regulated to 9 °C with a re-circulating water-glycol bath; the sample temperature in each quartz cell was determined with a thermistor. A black anodized aluminum sample rack with machined cylindrical slots was placed in the water-glycol bath. Quartz vials were filled with 20 mL of 0.2 μm-filtered, air-saturated seawater, and placed into hollow black cylindrical vial holders that were threaded at the top. Each vial holder had an o-ring inserted into a groove around the inner bottom opening to hold the vial in place and seal it from the water-glycol coolant. After each quartz vial was filled with the filtered seawater sample and placed into the holder, a Teflon-lined plastic cap was screwed onto the top of the vial holder.
to secure and seal the quartz vial in the vial holder. Each vial holder was then placed into a cylindrical slot in the sample rack. There were five slots above each of the eight long-pass filters. In this study, two samples were irradiated per long-pass filter treatment. After samples were placed into the aluminum rack, they were equilibrated to 9 °C prior to an irradiation. Irradiation times ranged from 15 to 360 min, depending on the spectral treatment and sample photon exposure (time-integrated irradiance\textsuperscript{72}). Dark controls were prepared by wrapping quartz vials with black electrical tape. These controls were placed in the rack for the longest irradiation period (360 min). The spectral irradiance (mW m\textsuperscript{-2} nm\textsuperscript{-1}) was measured inside each irradiation cell (and therefore included transmission through the water-glycol bath) with a custom built fiber-optic spectroradiometer as described by Neale and Fritz\textsuperscript{71}. Since incoming light entered the capped quartz cells vertically from the base, it was necessary to correct irradiances for scattering and multiple reflections employing nitrite chemical actinometry\textsuperscript{73}. If this correction was not included, AQYs would be overestimated by 10-30%.

Measured H\textsubscript{2}O\textsubscript{2} production rates ($R_{H2O2}$) were used along with irradiance data ($E_\lambda$) and wavelength-dependent sample absorbance coefficient measurements ($a_\lambda$) to determine an AQY spectrum. Measured H\textsubscript{2}O\textsubscript{2} production rates were fit to:

\begin{equation}
R_{H2O2} = \int \Phi_\lambda \ a_\lambda \ E_\lambda d\lambda
\end{equation}

where $\Phi_\lambda$ is the wavelength-dependent AQY that was determined from the AQY spectrum which was assumed to follow an exponential function:

\begin{equation}
\Phi_\lambda = \Phi_{300nm} \ 10^{m_1 (\lambda - 300)}
\end{equation}

$m_1$ is the slope and $\Phi_{300nm}$ is the AQY for H\textsubscript{2}O\textsubscript{2} formation at 300 nm. Nonlinear regression (Marquardt algorithm) was used to estimate the $m_1$ and $\Phi_{300nm}$ that maximized $r^2$ and minimized the mean-square error between measured H\textsubscript{2}O\textsubscript{2} production rates and those predicted using
To perform this analysis, initial boundary conditions for $\Phi_{300nm}$ and $m_1$ were set using the AQY and $m$ determined from narrow bandwidth experiments.

### 2.5 Shipboard Photochemical Production Experiments

Photochemical production rates for H$_2$O$_2$ were determined during field studies in the 1998 austral spring aboard the R/V *Lawrence M. Gould* at several hydrographic stations in the confluence of the Weddell and Scotia Seas as well as along the Antarctic Peninsula. Production rates were also determined at stations in the Gulf of Maine and North Atlantic Ocean during the 1999 summer aboard the R/V *Endeavor*. Production rates were determined in the field with freshly collected seawater samples in order to compare to modeled rates that were determined using AQY results obtained with stored seawater in the home laboratory. Hydrogen peroxide production rates were determined at sea using the procedures described in Yocis et al. and Teflon-sealed quartz irradiation vessels filled with 0.2 μm-filtered, air-saturated seawater.

#### 2.5.1. Wavelength-Dependent H$_2$O$_2$ Production Rates:

Shipboard experiments were conducted to determine the photochemical production of H$_2$O$_2$ in the UV-B (290-320 nm), UV-A (320-400 nm), and visible region of the solar spectrum. Duplicate quartz tubes were placed in a flow-through surface seawater bath and exposed to sunlight on the ship’s aft deck between 10:00 and 16:00 local time. Details of the quartz tube design are given in Kieber et al. Production rates for H$_2$O$_2$ in these quartz tubes were compared to rates determined in quartz tubes that were wrapped in Mylar D polyester film or placed in a UF3 Plexiglas box. Mylar D polyester film and UF3 Plexiglas cut-off wavelengths were 313 and 400 nm, respectively (Fig. 2).
2.5.2. In Situ $\text{H}_2\text{O}_2$ Production Rates: Depth-dependent, daytime $\text{H}_2\text{O}_2$ production rates were determined by placing duplicate quartz tubes at six depths from 2 to 20 m using a free-floating drifter described by Kieber et al. The daytime production rate was also determined at the sea surface in a flow-through seawater bath on the aft deck of the R/V *Endeavor* in a location with minimal shading. The free-floating drifter and surface samples were deployed prior to sunrise (ca. 05:00) and retrieved after sundown (ca. 20:00) to obtain average daily photochemical production rates for hydrogen peroxide as a function of water-column depth. Controls were examined to determine hydrogen peroxide production during the deployment and retrieval of the drifter. Experimentally determined in situ production rates were compared to calculated depth-integrated, daily production rates:

$$ \Phi_{T,\lambda} \int \int \int \alpha_{z,\lambda} \frac{E_{z,\lambda,t}}{d z d \lambda d t} $$

where $R_{\text{H}_2\text{O}_2}$ is the depth- and wavelength-integrated, daytime production rate, $\Phi_{T,\lambda}$ is the temperature- and wavelength-dependent AQY, $\alpha_{z,\lambda}$ is the wavelength- and depth-dependent CDOM absorption coefficient, and $E_{z,\lambda,t}$ is the wavelength-, depth- and time-dependent spectral irradiance, and $d \lambda$ is the wavelength bandwidth (1 nm). $R_{\text{H}_2\text{O}_2}$ was calculated every 0.5 m ($dz$) from 0 to 20 m. The spectral irradiance ($E_{z,\lambda}$) at depth $z$ was determined from:

$$ E_{z,\lambda} = E_{0,\lambda} e^{K_{z,\lambda} z} $$

where $E_{0,\lambda}$ is the spectral irradiance at the sea surface and $K_{z,\lambda}$ is the wavelength-dependent downwelling attenuation coefficient. $K_{z,\lambda}$ was determined from linear regression analysis of $\ln \left( E_{z,\lambda} / E_{0,\lambda} \right)$ versus depth. Underwater irradiance measurements were made to determine $K_{z,\lambda}$ using a NIST-calibrated free-falling Profiling Multi-Channel Radiometer (Satlantic, Inc., Halifax, Nova Scotia) that measured the irradiance at thirteen spectral channels: 304, 323, 338, 380, 411, 442, 489, 510, 523, 555, 670, 683, and 700 nm. The bandwidth for each channel was 10 nm.
Satlantic Radiometer was deployed from the aft winch and allowed to free fall after it was situated approximately 30-60 m from the stern of the ship; several down casts were averaged to determine the spectral irradiance as a function of depth. Surface irradiance measurements were made every 15 min from 290 to 600 nm at 1 nm intervals with an NIST-calibrated Optronics Laboratories OL 754 spectroradiometer. Integrating surface irradiance measurements with time yielded $E_{0,\lambda,t}$; $E_{z,\lambda,t}$ were calculated using $E_{0,\lambda,t}$ and $K_{z,\lambda}$.

3. RESULTS AND DISCUSSION

3.1 Reciprocity for H$_2$O$_2$ Photoproduction

Experiments were conducted employing the narrow bandwidth irradiation system to determine if hydrogen peroxide production was a linear function of the photon exposure. For all wavelengths examined, H$_2$O$_2$ formation was initially a linear function of photon exposure. As an example, H$_2$O$_2$ production was linear from 0 to 20 min at 300 nm for a coastal seawater sample collected from the Gulf of Maine at Ammen Rock (Fig. 3). The photoproduction of H$_2$O$_2$ was non-linear for irradiations longer than approximately 25 min at 300 nm (Fig. 3) even though there was very little change in CDOM absorbance (< 5%). Several factors can result in this non-linear behavior, but the fact that the CDOM absorbance changed very little in these experiments indicated that the non-linearity was likely due to the loss of specific precursors leading to the formation of H$_2$O$_2$ or the loss of H$_2$O$_2$ (at the shorter wavelengths) through its direct or indirect photolysis. This non-linearity resulted in an approximately 15% decrease in the AQY between 25 and 60 min. A lack of reciprocity over short time scales was unexpected and may partly explain the subsurface maximum in average, net daytime hydrogen peroxide photoproduction rates that were observed in two in situ irradiation experiments conducted in the confluence of the
Weddell and Scotia Seas$^{22}$. These subsurface production rate maxima may have resulted from greater photon exposures in the near surface samples, especially in the UV, that reduced production rates at these depths due to non-reciprocity, whereas reciprocity may have been met for samples incubated deeper in the water column giving rise to slightly higher production rates at intermediate depths.

Lack of reciprocity was not only observed at 300 nm and was not limited to the Ammem Rock sample. Similar results were obtained for other seawater samples and at other wavelengths, although for longer wavelengths reciprocity occurred over much longer time frames (e.g., 6 h at 400 nm). All AQYs were determined in this study within the linear portion of the production plot where reciprocity was observed. Under these conditions, AQYs should be constant and not a function of photon exposure. Except for a few cases$^{78,79}$, reciprocity is rarely tested in natural-water photochemical studies when AQYs are determined. Nonetheless, in nearly all cases where H$_2$O$_2$ concentrations have been determined as a function of photon exposure using sunlight or a Xe-lamp (or Hg-lamp) based irradiation system, it has been shown that H$_2$O$_2$ accumulation is non-linear after exposures between 20 to 120 min (and in some cases even as short as 5 min) in a range of freshwater and marine samples$^{78,81-84}$ and aqueous solutions of organic matter isolates$^{17,53,85}$. Non-linear H$_2$O$_2$ accumulation rates imply that corresponding AQYs would decrease as well if CDOM absorbance coefficients photobleached more slowly than H$_2$O$_2$ accumulation rates decreased. Therefore, care should be taken in reporting, comparing and interpreting AQYs, and photochemical production or loss rates for indirect photochemical processes in seawater, since rates will not be a linear function of photon exposure upon long-term irradiation as has been implicitly assumed in many published studies.
3.2 Comparison of Polychromatic and Narrow Bandwidth AQY Spectra

An experiment was conducted with 0.2 μm-filtered Rhode River estuary water (salinity 11.1 ppt) to compare the wavelength-dependent AQY spectrum determined using the polychromatic irradiation system to that obtained with the narrow bandwidth irradiation system. Apparent quantum yields that were determined with the polychromatic and narrow bandwidth irradiation systems were in good agreement at all wavelengths that were examined (Fig. 4). For both approaches, AQY spectra decreased exponentially with increasing wavelength, and no localized maxima or minima were noted in the narrow-bandwidth generated AQY spectrum. When the polychromatic and narrow bandwidth spectra were compared to each other, the largest differences in AQYs (≤ 20%) were observed at wavelengths between 320 and 360 nm. For example, at 320 nm the average AQY determined by narrow bandwidth radiation was 2.0 x 10⁻⁴ compared to 2.4 x 10⁻⁴ determined with polychromatic radiation; this difference was not significant at the 95% confidence interval as was the case for comparisons at other wavelengths. The close agreement between polychromatic and narrow-bandwidth based AQY spectra indicated that there were no significant polychromatic wavelength interactions that affected AQYs. In other words, an AQY determined at one wavelength (e.g., 350 nm) by either narrow-bandwidth or polychromatic irradiations was not affected by irradiations at shorter (e.g., <340 nm) or longer (e.g., >360 nm) wavelengths. These results confirm the suitability of applying an exponential function to phenomenologically describe H₂O₂ AQY spectra in marine waters. Given the good agreement between these two approaches in determining AQYs for H₂O₂ photoproduction, it would be advantageous to determine AQYs using the polychromatic irradiation system because AQY spectra can be obtained in a fraction of time (e.g., day) required to determine AQY spectra using a narrow-bandwidth irradiation system (e.g., week). Furthermore, our results validate the use of...
narrow-bandwidth determined H₂O₂ photoproduction AQY to model H₂O₂ photochemical production rates in natural waters that are exposed to polychromatic solar radiation.

In a separate experiment, the 0.2 μm-filtered Rhode River estuary water was stored for 2.5 months at 4 °C, and wavelength-dependent AQYs were determined using the narrow bandwidth irradiation system to determine the effect of sample storage on AQYs. Apparent quantum yields did not change compared to initial wavelength-dependent AQY values determined with the freshly collected water (no significant difference noted at the 95% confidence level employing a two-tailed t test) demonstrating that there were no significant changes in AQYs due to long-term, cold storage in the dark (Fig. 4). The same result was seen in CDOM spectra (i.e., no changes were seen after 2.5 months, data not shown).

3.3 Wavelength-Dependent AQYs

Wavelength-dependent AQYs for the photochemical formation of hydrogen peroxide were determined at 25 °C in 0.2 μm-filtered seawater samples collected from Booth Bay Harbor, Ammen Rock, Rhode River estuary, Banks Channel, Gulf of Mexico, Station ALOHA, and Antarctic stations B, N and Arthur Harbor. For all seawater samples, AQYs decreased exponentially with increasing wavelength, ranging from 3.6 x 10⁻⁴ - 10.4 x 10⁻⁴ at 290 nm to 0.17 x 10⁻⁴ - 0.97 x 10⁻⁴ at 400 nm (Fig. 5). A tabular listing of all AQYs determined at 25 °C is given in Miller⁶⁶. Surprisingly, wavelength-dependent AQYs for individual seawater samples differed by less than a factor of two relative to mean AQYs determined by non-linear regression analysis of all the AQY data for all seawater samples that were examined in this study (Fig. 5). In addition, comparison of AQYs from different seawater samples shown in Fig. 5 indicated that there was no clear trend that would suggest that AQYs were higher or lower for coastal stations.
compared to oligotrophic stations. It is remarkable how similar wavelength-dependent AQYs are among the different water samples even though the concentrations and speciation of metals and sources and concentrations of DOM giving rise to H₂O₂ photoformation are presumably different (e.g., terrestrial vs. marine), and the samples will have had different light-exposure histories. Given the similarity in wavelength-dependent AQYs among the seawater samples that were examined in this study, data were combined to yield an AQY spectrum (see best-fit line in Fig. 5, equation 9) that was derived from non-linear regression analysis of the composite AQY data set; this equation can be used to estimate wavelength-dependent AQYs for H₂O₂ photoproduction at 25 °C:

\[
\Phi_\lambda = 1.70 e^{0.0272\lambda} \quad (9)
\]

The slope (±95% confidence interval) of this line, -0.0272 ± 0.0014 nm⁻¹, is not statistically different from the slope (-0.0267 ± 0.0028 nm⁻¹) reported by Yocis et al.²² using pooled AQY data from the Caribbean Sea, Orinoco River outflow (Venezuela), Suwanee River (GA), Vineyard Sound (MA), Shark River outflow (FL), and the Antarctic. This provides further evidence that AQYs are similar in all marine waters tested to date irrespective of expected differences in local DOM composition and light history. However, this finding may not extend to some terrestrially-dominated freshwater or ground water systems where a few relatively high AQYs have been reported⁴,⁶¹,⁸¹.

### 3.4 Temperature-Dependent AQYs

Temperature dependence studies were conducted with seawater from Booth Bay Harbor, Rhode River estuary, Station ALOHA, and Antarctic stations B, N and Arthur Harbor. As evident from the example Arrhenius plots shown in Fig. 6, AQYs decreased with decreasing
temperature for all seawater samples and all wavelengths that were examined. On average, AQYs decreased by a factor 1.8 per 10 °C. A tabular listing of temperature-dependent AQYs is given in Miller.

Activation energies for the photochemical production of H\(_2\)O\(_2\) in seawater determined from linear regression analysis of the Arrhenius plots ranged from 8.3 to 52.7 kJ mol\(^{-1}\) (Table 1). For all water samples tested, \(E_a\) increased with increasing wavelength (Table 1, Fig. 7). The largest difference in the \(E_a\) was seen for the Antarctic seawater sample collected from Antarctic Station B, which had a lower \(E_a\) at 290, 300 and 320 nm compared to the other samples including two other Antarctic waters (Sta. N and Arthur Harbor). Likewise, for all samples, the largest difference was observed when comparing \(E_a\) at 290 to 400 nm, where the average value for \(E_a\) increased from 16.6 to 31.9 kJ mol\(^{-1}\). Our average \(E_a\) determined at 400 nm, 31.9 ± 12 kJ mol\(^{-1}\) (Fig. 7) agreed well with the ≥ 365 nm \(E_a\) value of 37.4 kJ mol\(^{-1}\) determined by Szymczak and Waite. In their study, they used a 365 nm band-pass filter and a Hg lamp to determine \(E_a\) for the photochemical formation of H\(_2\)O\(_2\) in an estuarine sample near the mouth of Port Hacking River Estuary. This \(E_a\) included the temperature dependence for H\(_2\)O\(_2\) photoformation not only for the 365 nm Hg emission line but also from several strong emission lines greater than 400 nm including ones at 405 and 436 nm.

Although differences were observed in our study in \(E_a\) among samples and as a function of wavelength, overall differences were not large except in a few cases. Therefore, for the purpose of aiding modeling efforts, we determined the average \(E_a\) between 300 and 370 nm where \(E_a\) differences among samples and wavelengths were relatively small, and this wavelength range encompasses the main spectral H\(_2\)O\(_2\) photoproduction bandwidth observed in coastal and oligotrophic seawater based on published spectra and results presented here. Within this UV
bandwidth, $E_a$ for H$_2$O$_2$ photoproduction were the same for all samples tested within the 95% CI (except for Arthur Harbor, Antarctica), with a mean value of 21.8 kJ mol$^{-1}$. Using this mean activation energy, the temperature dependence for $\Phi_{\lambda,T}$ was determined from:

$$ln \frac{\Phi_{\lambda,T}}{\Phi_{\lambda,298}} = \frac{21.8}{R} \left( \frac{1}{298} - \frac{1}{T} \right)$$

where $\Phi_{\lambda,298}$ is the wavelength-dependent AQY at 298 K calculated from equation 9.

The increase in activation energies between 290 and 400 nm (Table 1) and the factor of 1.8 increase in the average AQY per 10 °C increase in temperature are consistent with a thermal process controlling the formation of hydrogen in seawater. Likely rate-limiting reactions controlling production are the uncatalyzed$^{54}$ and catalyzed$^{55-59,86-89}$ thermal disproportionation reactions to form H$_2$O$_2$. These reactions are complex and not fully resolved$^{87}$. Our results suggest that the relative importance of the different disproportionation reactions may vary with wavelength, due to changes in associated light-driven reactions, which in turn may lead to wavelength-dependent changes in $E_a$. It is also possible that the branching ratio for catalyzed O$_2^-$ dismutation versus non-dismutation decay pathways for O$_2^-$ involving metals or DOM$^{58,78,87,91}$ may vary with wavelength, thereby affecting observed $E_a$ values. Despite uncertainties regarding why $E_a$ increases with increasing wavelength of irradiation, the fact that $E_a$ does change with wavelength and between water samples indicates that there are some very interesting wavelength-dependent changes in the main pathways leading to H$_2$O$_2$ photoformation in natural waters that warrant further research.

### 3.5 Sunlight-Normalized H$_2$O$_2$ Production Rates

Sunlight normalized production rates for hydrogen peroxide at the sea surface are shown in Fig. 8 for two coastal and two oligotrophic seawater stations. Spectrally-resolved, sunlight-
normalized rates (i.e., the product of $E_i a_i \Phi_i$) were calculated at 1 nm intervals from 290 to 500 nm. Spectral irradiance data were obtained from Optronics spectroradiometer measurements made every 15 min at the Damariscotta River outflow station on a cloudless day, July 20, 1999.

The mean response wavelength for hydrogen peroxide photoproduction at the sea surface was determined by integrating the area under each curve in Fig. 8 between 290 and 500 nm and calculating the wavelength corresponding to half the area. Based on this calculation, the mean response wavelengths for the four stations were 330 nm for Ammen Rock, 322 nm for the Rhode River estuary, 318 nm for Station ALOHA, and 322 nm for Antarctic Station N (Fig. 8). In addition to the mean response wavelength, the response bandwidth was also determined. The response bandwidth, defined as the width at half-height of the response curve, varied from 302-335 nm for the Rhode River estuary, 301-333 nm for Station ALOHA, 302-335 nm for Antarctic Station N, to 303-346 nm for the Gulf of Maine, Ammen Rock Station (Fig. 8). Other than a slight red shift in the sunlight-normalized spectrum at Ammen Rock, these dissimilar samples had nearly the same spectral shape and peak response wavelength. Peak responses and spectral shapes were also quite similar to that observed for several other species that are photochemically formed or removed from seawater including dimethylsulfide$^{64,92,93}$, carbonyl sulfide$^{94}$ and ammonia$^{95}$, even though the processes leading to the production or loss of these compounds are undoubtedly quite different. However, not all compounds show the same spectral shape and peak response including (1) dissolved molecular oxygen whose spectrally-dependent photochemical loss is quite broad spanning the UV and extending into the blue portion of the solar spectrum$^{79}$, (2) formaldehyde photoproduction that is initiated by UV-B solar radiation$^{96}$ and (3) DIC and CO photoformation rates that are red shifted with a significant contribution at wavelengths greater than 400 nm$^{69,97}$. None of these differences are surprising, as they reflect fundamentally different
reactants, photosensitizers and pathways leading to the formation or loss of these different compounds.

The importance of UV in controlling H$_2$O$_2$ photoproduction was also evident from integrating the area under the curve for each seawater sample in Fig. 8; 29-51% of the total photoproduction was in the UV-B, 48-63% was in the UV-A and only a small fraction, 1-9%, was greater than 400 nm. This finding is in good agreement with the Gulf of Maine and Palmer Station field studies where H$_2$O$_2$ production rates determined in quartz tubes were compared to production rates in Mylar-wrapped quartz tubes and quartz tubes enclosed in UF3 Plexiglas (Fig. 9). These plastic coverings approximated UV-B and total UV (< 400 nm) exclusion filters, respectively. Based on differences in H$_2$O$_2$ photoproduction between the different treatments, 38-52, 45-47 and 4-16% of the total H$_2$O$_2$ production was observed in the UV-B, UV-A and > 400 nm, respectively, in the Gulf of Maine samples; and 32, 64 and 3% was observed in the UV-B, UV-A and > 400 nm, respectively, in seawater collected from Arthur Harbor, Palmer Station, Antarctica. These field results obtained with freshly collected and 0.2 $\mu$m-filtered samples agreed well with the relative trends observed in our lab-based results with stored 0.2 $\mu$m-filtered seawater samples indicating that sample storage did not significantly affect samples with respect to H$_2$O$_2$ photoproduction.

3.6 In Situ and Modeled H$_2$O$_2$ Photoproduction Rates

During the 1999 Gulf of Maine cruise, a free-floating drifter$^{76}$ was deployed to measure total daytime photochemical production rates for hydrogen peroxide as a function of depth at the coastal stations Ammen Rock and the Damariscotta River outflow, and at the oligotrophic station E in the northwest Atlantic Ocean (Fig. 1B). For all free-floating drifter studies, total daytime photochemical production rates for H$_2$O$_2$ were highest at the sea surface and decreased
exponentially with increasing depth (Fig. 10). Mean surface photochemical production rates
determined at the two coastal stations, 207 and 277 nM d\(^{-1}\), were approximately three times
greater than at the oligotrophic station E (72 nM d\(^{-1}\)). Production rates at the two coastal stations
decreased rapidly below the sea surface to 2.7 and 9.9 nM d\(^{-1}\), respectively, at 15 m, whereas rates
at Station E in the northwest Atlantic Ocean (Fig. 1B) decreased more slowly with depth to 24
nM d\(^{-1}\) at 15 m. Differences in the vertical attenuation of production rates in the water column
observed between the coastal and oligotrophic stations were consistent with differences in
downwelling attenuation coefficients, \(K_d\), among these stations. In particular, solar radiation was
attenuated in the water column much faster at the coastal stations compared to the oligotrophic
station E, especially at the shorter wavelengths in the UV. For the Damariscotta River outflow
and Ammen Rock stations, \(K_d\) at 323, 338 and 411 nm were 1.5, 1.0, and 0.43 m\(^{-1}\), and 0.70, 0.56
and 0.21 m\(^{-1}\), respectively, whereas \(K_d\) values at Station E were considerably smaller at 0.19, 0.09
and 0.04 m\(^{-1}\), respectively. Likewise, vertical attenuation of H\(_2\)O\(_2\) production rates in the water
column (see Yocis et al.\(^2\)) for details regarding this calculation) were 0.32, 0.20, and 0.07 m\(^{-1}\) for
the Damariscotta River outflow, Ammen Rock and Station E, respectively, paralleling differences
in \(K_d\) among these stations (i.e., lower \(K_d\) values corresponded to lower photochemical attenuation
rates).

Since reciprocity was not met for fairly short irradiations (ca. 20-60 min) of Ammen Rock
seawater (Fig. 3) and given the length of the drifter studies (ca. 05:00-20:00 local time), it is not
surprising that day-long exposure of Ammen Rock seawater to solar radiation in the drifter study
(Fig. 10A) led to 41% lower observed daytime rates in the near-surface exposed samples
compared to modeled rates determined from equation 7. This near-surface difference was not
observed in the Damariscotta River outflow drifter study (Fig. 10B), perhaps due to the faster
attenuation of UV in the water column at this station (as evidenced by larger $K_d$ values) compared
to the Ammen Rock station. Lower than expected in daily rates observed at Ammen Rock and in
the Antarctic\textsuperscript{22} in near surface waters may represent the norm, as there is no reason to expect that
production rates should be constant with increasing light dose in a natural setting, especially near
the sea surface. This lack of reciprocity, as evidenced by lower rates, will be an issue when
comparing results from different studies when solar (or solar simulator) exposure times are
significantly different or when applying AQYs to model daily rates.

Deeper in the water column ($\geq 4$ m), modeled rates underestimated in situ rates from 30 to
90%, but these differences are more likely due to uncertainties in the parameters ($\Phi_\lambda$, $a_\lambda$, $K_{d,\lambda}$, $E_{z,\lambda}$)
used in the model\textsuperscript{98} and variability in observed rates as opposed to “real” differences between the
model and observations. For example, uncertainties for modeled wavelength-dependent AQYs
ranged from 27% at 290 nm to 12% at 400 nm. In addition, absorption coefficients, irradiances
and downwelling attenuation coefficients had errors ranging from $< 1$ to $> 100\%$, depending on
the wavelength and depth in the water column. Using estimates for wavelength-dependent errors
for each of these parameters, propagation of error analysis was conducted to determine the
coefficient of variation of wavelength-dependent production rates for each depth. The coefficient
of variation of modeled production rates for the two drifter deployments ranged from 4-5% at the
surface, 8-13% at 2 m, 14-25% at 4 m, 22-44% at 6 m, 31-64% at 8 m, 39-103% at 10 m and 60-
450% at 15 m. This error analysis did not take into account errors associated with surface
seawater reflection or changes in CDOM absorption (we assumed 100\% radiative transfer and
constant CDOM absorbance—i.e., no photobleaching), which would further increase modeling
uncertainties.
Despite uncertainties in our photochemical model, modeled depth-integrated production rates between 0 and 20 m at Ammen Rock and the Damariscotta River outflow differed from measured depth-integrated rates over the same depth interval by only 9 and 14%, respectively (Table 2), and any differences noted at a given depth (as previously discussed) cancelled out when rates were integrated vertically in the upper 20 m. In addition, although surface production rates at the oligotrophic Station E in the northwest Atlantic Ocean were much lower than at the coastal stations in the Gulf of Maine, depth-integrated production rates were nearly the same at all three stations (Table 2) because H$_2$O$_2$ was produced at greater depths at Station E relative to the coastal stations (Figure 10). Yocis et al.\textsuperscript{22} observed that the depth-integrated production rate at an oligotrophic station, Crystal Sound, Antarctic was almost twice the rate at a coastal Antarctic seawater station undergoing a \textit{Cryptomonas} bloom. These comparisons demonstrate the importance of oligotrophic waters in H$_2$O$_2$ photoproduction, and more broadly illustrate the importance of considering the entire photochemically-active water column in assessing the importance of a photoprocess in marine waters.

\section*{4. CONCLUSIONS}

The main finding of this study is that, although differences are observed, wavelength-dependent AQYs for the photochemical formation of H$_2$O$_2$ in seawater are remarkably similar among diverse marine environments that presumably contain varying composition and concentrations of metals and DOM from a range of terrestrial and marine sources and that have different light-exposure histories. Thus, the question arises: Why are wavelength-dependent AQYs for H$_2$O$_2$ production in Antarctic waters so close in value to wavelength-dependent AQYs determined in estuarine water from Banks Channel, NC? Perhaps AQY are similar because
marine DOM, despite expected source differences, is reworked by marine microorganisms yielding a similar reservoir of DOM or metal-DOM complexes that serve as photochemical precursors for superoxide formation or as catalysts for \( \text{O}_2 \) dismutation and \( \text{H}_2\text{O}_2 \) formation.

Results presented in this study provide the data needed to model \( \text{H}_2\text{O}_2 \) production rates in seawater on a global scale using remotely sensed CDOM absorbance, sea-surface spectral irradiance and sea-surface temperature data, as was done in a companion paper\(^5\). For future studies it will be important to investigate the assumption that AQYs are constant as a function of photon exposure not only for \( \text{H}_2\text{O}_2 \) production but for other compounds as well (e.g., DIC, CO, DMS, COS, acetaldehyde). If AQY vary with photon exposure as suggested by our results, then it will be important to assess the ramifications of these findings to remotely sensed production or loss rate estimates or for conclusions regarding carbon cycling in marine waters based on relatively short photochemical exposure experiments\(^9\).

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polychromatic irradiance data. Thanks are also extended to two anonymous reviewers for their thorough and thoughtful comments that improved the clarity and quality of this manuscript.
REFERENCES


Table 1. Activation energy for the photochemical production of hydrogen peroxide in seawater at five wavelengths; the irradiation bandwidth was set at ± 5 nm for 290, 300 and 320 nm and ± 9.8 nm for 360 and 400 nm. The error for $E_a$ denotes a 95% confidence interval.

<table>
<thead>
<tr>
<th>Sample</th>
<th>290 nm</th>
<th>300 nm</th>
<th>320 nm</th>
<th>360 nm</th>
<th>400 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Booth Bay Harbor</td>
<td>12.4 ± 4.0</td>
<td>12.3 ± 2.2</td>
<td>15.6 ± 2.7</td>
<td>18.0 ± 3.8</td>
<td>24.0 ± 5.7</td>
</tr>
<tr>
<td>Rhode River estuary</td>
<td>19.2 ± 2.2</td>
<td>15.9 ± 2.0</td>
<td>17.7 ± 2.8</td>
<td>21.7 ± 5.4</td>
<td>26.2 ± 8.1</td>
</tr>
<tr>
<td>Hawaii Station ALOHA</td>
<td>21.7 ± 3.3</td>
<td>20.4 ± 3.1</td>
<td>23.3 ± 3.0</td>
<td>24.9 ± 6.4</td>
<td>34.1 ± 12.2</td>
</tr>
<tr>
<td>Antarctic Station B</td>
<td>9.6 ± 2.3</td>
<td>8.3 ± 2.6</td>
<td>13.6 ± 3.6</td>
<td>21.7 ± 26.0</td>
<td>20.8 ± 48.8</td>
</tr>
<tr>
<td>Antarctic Station N</td>
<td>16.9 ± 3.7</td>
<td>21.1 ± 3.6</td>
<td>22.5 ± 6.3</td>
<td>28.9 ± 13.4</td>
<td>33.5 ± 9.4</td>
</tr>
<tr>
<td>Arthur Harbor, Palmer Station</td>
<td>19.7 ± 2.2</td>
<td>28.7 ± 3.5</td>
<td>35.2 ± 12.8</td>
<td>42.1 ± 15.3</td>
<td>52.7 ± 9.7</td>
</tr>
</tbody>
</table>
Table 2. Total daytime, depth-integrated photochemical production rate for H$_2$O$_2$ in seawater determined from free-floating drifter studies conducted at Ammen Rock and the Damariscotta River outflow in the Gulf of Maine, and at Station E in the northwest Atlantic Ocean. The reported error is the standard deviation for measured and modeled rates. Modeled rate could not be determined (ND) at Station E because measured irradiance data were not available at this station.

<table>
<thead>
<tr>
<th>Station</th>
<th>Sampling Depth (m)</th>
<th>Measured</th>
<th>Modeled</th>
<th>% Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ammen Rock</td>
<td>15</td>
<td>1.10 ± 0.025</td>
<td>0.96 ± 0.028</td>
<td>9.4</td>
</tr>
<tr>
<td>Damariscotta River outflow</td>
<td>15</td>
<td>0.68 ± 0.016</td>
<td>0.58 ± 0.020</td>
<td>14.0</td>
</tr>
<tr>
<td>Station E (NW Atlantic Ocean)</td>
<td>20</td>
<td>0.77 ± 0.012</td>
<td>ND</td>
<td>ND</td>
</tr>
</tbody>
</table>
**FIGURES**

**Figure 1.** Location of hydrographic stations in (A) the confluence of the Weddell and Scotia Seas, and the Antarctic Peninsula; and (B) along the East Coast of the United States. Not shown are Hawaii Station ALOHA and the Gulf of Mexico station.

**Figure 2.** Transmission spectra for Mylar D polyester film and UF3 Plexiglas.
Figure 3. Photochemical formation of hydrogen peroxide in 0.2 μm-filtered Ammen Rock, Gulf of Maine seawater and the apparent quantum yield as a function of irradiation time at 300 ± 5 nm. Apparent quantum yields were corrected to account for losses in absorbance observed for longer irradiations. The solid line is the best-fit line of the production data as determined from linear regression analysis, excluding data from 25-60 min. The dashed line is the best-fit line determined from non-linear regression analysis.

Figure 4. Comparison of wavelength-dependent apparent quantum yields in freshly collected Rhode River estuary water determined with narrow bandwidth (○) and polychromatic irradiation systems (solid line). The dashed lines denote the upper and lower 95% confidence interval for the polychromatic study. The narrow bandwidth experiment was repeated 2.5 months later with stored (4 °C) 0.2 μm-filtered Rhode River estuary water (●). All apparent quantum yields were determined at 9 °C.
Figure 5. (A) Wavelength-dependent apparent quantum yields for the photochemical formation of hydrogen peroxide in nine marine waters at 25 °C with low (< 0.5 m⁻¹, ▲) or high (> 0.7 m⁻¹, ○) DOM absorption coefficients at 300 nm. The best-fit line was determined from non-linear regression analysis. Data shown in this figure are tabulated in Miller (2000). The value of n above each data set represents the total number of samples analyzed at that wavelength.

Figure 6. Natural logarithm of the apparent quantum yield plotted as a function of inverse temperature for 0.2 μm-filtered (A) Rhode River estuary water and (B) Hawaii Station ALOHA seawater. The activation energy was determined from the slope of the best-fit line determined from linear regression analysis of the data at (●) 290, (▼) 300, (■) 320, (♦) 360 and (▲) 400 nm. Three irradiation temperatures are listed in Panel B above the 290 nm data set.
Figure 7. Average activation energy for the data shown in Table 1 plotted as a function of wavelength. Error bars denote the 95% confidence interval.

Figure 8. Sunlight normalized production rates for hydrogen peroxide at the sea surface in (A) coastal seawater and (B) oligotrophic seawater. Note the y-axis scale change in panels A and B.
Figure 9. Hydrogen peroxide photochemical production rates in 0.2 μm-filtered seawater samples, collected from the northwest Atlantic Ocean stations E and F, and Arthur Harbor, Palmer Station, Antarctica, that were exposed to sunlight in quartz tubes or quartz tubes enclosed with long-pass filters, Mylar D or UF3 Plexiglas. Error bars denote the standard deviation of four replicates.
Figure 10. Hydrogen peroxide photochemical production rates as a function of depth determined from a free-floating drifter study (●) and predicted from a photochemical model (−) for (A) Ammen Rock, Gulf of Maine and (B) the Damariscotta River outflow, Gulf of Maine. (C) Hydrogen peroxide photochemical production rates as a function of depth determined from a floating drifter study in the northwest Atlantic Ocean, Station E. Production rates were not modeled at Station E, since measured spectral photon exposure data were not available for this station.