Updated and validated solar irradiance reference spectra for estimating environmental photodegradation rates†

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Irradiance reference spectra are used to calculate environmentally relevant photodegradation half-lives, but the currently used spectra were originally published in the 1980s with limited validation. The goal of this work is to provide updated irradiance reference spectra using the Simple Model of the Atmospheric Radiative Transfer of Sunshine (SMARTS). The SMARTS irradiance spectra were validated against measurements from several high-resolution spectroradiometers, and the updated irradiance reference spectra use current measurements for atmospheric species that can affect the irradiance that reaches the Earth’s surface. These updated irradiance spectra are provided in 1 nm increments from 280 to 800 nm for 0° to 70° latitude at 10° increments in both the northern and southern hemisphere. Lastly, the influence of the input parameters on the modeled irradiance spectra was investigated. This work will allow users to calculate more accurate photodegradation half-lives using the updated irradiance reference spectra, and it also provides insight for users to calculate their own location- and time-specific irradiance spectra using SMARTS.

Environmental significance

Solar irradiance spectra are necessary for calculating environmentally relevant photodegradation half-lives of chemicals. This manuscript provides updated solar irradiance spectra, which were validated against high-resolution spectroradiometer measurements. These updated irradiances should be used to calculate more accurate photodegradation half-lives in the natural environment.

1 Introduction

Pollutants can be degraded or transformed by photochemical reactions in the aquatic environment. These reactions can either proceed through direct photodegradation, where the pollutant absorbs light and undergoes reaction, or through indirect photoreaction, where other chemical species absorb light and subsequently undergo reaction with the pollutant.

In both cases, the rate of pollutant degradation or transformation is dependent on the flux of photons that reach Earth’s surface, which is called the solar irradiance or the photon flux rate.

In laboratory studies, the photodegradation of chemicals is usually investigated using a well-characterized light source. The irradiance of a light source can be measured using a spectroradiometer, which has been calibrated beforehand with an independent light source of known spectral irradiance. Otherwise, the relative irradiance can be measured using an uncalibrated spectroradiometer or obtained from the literature, and a chemical actinometer can be used to determine the photon flux into the test solution. In contrast to laboratory conditions, the irradiance that reaches the surface of the Earth can vary substantially based on the geographic location, altitude, time of day, time of year, and the concentration of atmospheric gases and particles.

Because of this variability in irradiance in the environment, reference solar irradiance spectra are typically used to calculate environmentally relevant photochemical half-lives for chemicals in surface waters. Both the EPA guideline (OPPTS 835.2210) and the OECD guideline (316) recommend using the day-averaged solar irradiance values for a clear sky (i.e., cloudless) day originally published in The Kinetics of Environmental Aquatic Photochemistry by Leifer (1988). These values, denoted as $L_i$, represent the irradiance averaged over a 24 hour day and were an extension of the irradiance values previously published by Zepp and Cline (1977), which were for solar noon and denoted as either $W_s$ or $Z_s$.

The previously reported $L_i$, $W_s$, and $Z_s$ values have several shortcomings that make the calculation of new irradiance values desirable. First, the irradiances are reported in increments of 2.5–50 nm from 296.2 nm to 800 nm. These
increments are often inconvenient to work with. More importantly, this discretization can lead to errors in photodegradation calculations. In the ultraviolet-B (UVB) range (280–315 nm), the irradiance substantially increases as the wavelength increases, and for many chemicals of interest the molar absorptivity also substantially decreases in this wavelength range. Therefore, their product \((e_\lambda I_0)\) is sensitive to averaging and truncating in this region, which can hinder the calculation of the quantum yield \(\text{[e.g., clofibric acid]}^1\). Second, the documentation is not thorough enough to replicate the irradiance values or is hard to obtain. For example, the atmospheric ozone concentrations to compute values of \(L_1\) are not given, so it is difficult to say how accurate these irradiances are in the UVB region. Additionally, ref. 11, where the \(L_1\) values were published, is currently out of print. Lastly, Zepp and Cline (1977) originally validated the \(Z_0\) values by measuring the degradation of chemicals with relatively short photodegradation half-lives.\(^1\) This, however, only considers ranges of irradiance values as a whole instead of individual wavelengths, and a validation for the \(L_1\) values could not be found in the literature.

The purpose of this article is to provide updated solar irradiance reference spectra at solar noon as well as day-averaged values. These irradiance values were calculated using the Simple Model of the Atmospheric Radiative Transfer of Sunshine (SMARTS)\(^14\) at 1 nm increments from 280 to 800 nm, which fully encompasses the 290 to 800 nm wavelength range suggested in the OECD guidelines.\(^10\) The modeled irradiances from SMARTS were then compared to measurements from several high resolution spectroradiometers to validate the reference spectra. Finally, a sensitivity analysis on the input parameters for SMARTS was performed to determine (a) which parameters had the largest influence and (b) the variability of solar irradiance values that could be expected in the environment depending on local atmospheric conditions.

1.1 Theory and history

The general form of the equation typically used to describe the pseudo first order photoreaction, using a monochromatic light source of a given wavelength \((\lambda)\), is:\(^11\)

\[
-\frac{dC}{dt} = \Phi \frac{I_0}{z} (1 - 10^{-(\alpha_\lambda + \epsilon_\lambda \ell)}) \left( \frac{\epsilon_\lambda \ell C}{\alpha_\lambda + \epsilon_\lambda C} \right) \Delta \lambda
\]  

where \(C\) is the concentration of the chemical of interest \([\text{mol L}^{-1}]\), \(t\) is time [s], \(\Phi\) is the quantum yield \([\text{mol chemical/mol photon}]\), \(I_0\) is the solar photon irradiance on a molar basis \([\text{mmol photons cm}^{-2} \text{ s}^{-1} \text{ nm}^{-1}]\) for a given wavelength, \(z\) is the depth of the water [cm], \(\alpha_\lambda\) is the decadic absorbance or attenuation coefficient of the solution (excluding the chemical of interest) divided by the optical pathlength \([\text{cm}^{-1}]\), \(\epsilon_\lambda\) is the decadic molar absorptivity of the chemical \([\text{mol}^{-1} \text{ L cm}^{-1}]\), \(\ell\) is the pathlength of the light in the water [cm], and \(\Delta \lambda\) is the wavelength interval of the light [nm]. These units are consistent because of conversions from mmol to mol and cm\(^3\) to L that cancel. The first term of the equation, \(\Phi\), is the fraction of molecules that undergo reaction of those that absorb a photon. The second term, which is the irradiance divided by the pathlength, gives the incident light intensity per unit volume. The third term describes the fraction of photons at a given wavelength that is absorbed by the system over the pathlength \(\ell\), which is from the Beer–Lambert Law. Finally, the last term is the fraction of photons absorbed by the solution that were absorbed by the chemical of interest.

There are already several assumptions present in eqn (1) that should be acknowledged. First, the \(\Phi\) is typically assumed to be independent of wavelength. Second, the concentration of chemical species and the light intensity are assumed to be homogenous over the pathlength \(\ell\).\(^15\) In addition, in order to use the published \(L_1\) values, the absorbance over the pathlength needs to be <0.02 \(\text{[i.e., (}\alpha_\lambda + \epsilon_\lambda \ell)\text{]} < 0.02\) according to OECD guidelines or <0.05 according to EPA guidelines. When this is true, the values \((1 - 10^{-(\alpha_\lambda + \epsilon_\lambda \ell)})\approx 2.303(\alpha_\lambda + \epsilon_\lambda \ell)\). When \((\alpha_\lambda + \epsilon_\lambda \ell) = 0.02\), this simplification overestimates the photons absorbed by the system by 2%, and when \((\alpha_\lambda + \epsilon_\lambda \ell) = 0.05\), the overestimation is 6%. Yet, this simplification allows us to rewrite eqn (1) into a form that can be easily integrated (eqn (2)). Finally, if a polychromatic light source is used, the rate at each wavelength \((k_\lambda)\) must be summed to obtain the observed overall rate constant \((k)\) (eqn (3)).

\[
\frac{dC}{dt} = -2.303\Phi \int_{z_0}^{z} \Delta \lambda C = -k_\lambda C
\]

\[
\ln \left( \frac{C}{C_{i=0}} \right) = -2.303\Phi \int_{z}^{z_0} \left( I_0(\alpha_\lambda + \epsilon_\lambda \ell) \right) t = -kt
\]

In the above equations, \(\ell/z\) is the ratio of the pathlength of a photon \((\ell)\) and the vertical depth of the water \((z)\). In a laboratory experimental setup, \(\ell/z\) are assumed to be equal. In the environment, \(\ell\) can be longer than \(z\) because (a) the incident light usually does not enter the water surface at a perpendicular angle \(\text{i.e., zenith angle of 0}\), and therefore photons do not travel vertically in the water column, and (b) the presence of particles can scatter photons making their pathlengths longer. In non-turbid water, which is assumed in the derivation of eqn (2), \(\ell/z\) is typically between 1.05 and 1.3.\(^16\)

In eqn (2), the solar irradiance, denoted as \(I_{0,\lambda}\), is the global horizontal irradiance. The global horizontal irradiance is the sum of direct beam and diffuse (also known as sky) irradiance that intercept a plane that is horizontal to the Earth’s surface. However, both Zepp and Cline (1977) and Leifer (1988) adjusted the global horizontal irradiance for reflectance off the water surface and the increased pathlength within the water column (Fig. 1). This was done by correcting the direct beam and diffuse sunlight separately.

The reflected fraction for both was calculated using the Fresnel equations and assuming a refractivity index of water of 1.34. For diffuse sunlight, the reflected fraction was calculated by Zepp and Cline to be 7% on average assuming a uniformly bright sky \(\text{i.e., isotropic light scattering without circumsolar or horizon brightening scattered components}\).\(^12\) For direct beam sunlight, the reflected fraction is <5% up to a solar zenith angle (SZA) of 57° but increases quickly at larger values of SZA. For example, 10% of sunlight is reflected at SZA = 67°, 20% is...
reflected at SZA = 74°, 40% is reflected at SZA = 81°, and 60% is reflected at SZA = 85°. Zepp and Cline (1977) subtracted the reflected fraction of light to get the solar irradiance values they denote as $W_\lambda$.\textsuperscript{12}

The increased pathlength in the water column was calculated using Snell’s law to determine the angle of refraction. Again assuming a refractive index of 1.34 for water, Zepp and Cline calculated that the pathlength for diffuse sunlight would be 1.2 times longer on average at mid-latitudes.\textsuperscript{14} For direct sunlight, the increased pathlength is calculated using $z\sec(\theta)$ where $\theta$ is the angle of refraction. The pathlength will be longer for higher values of the SZA. For a SZA = 85°, $\theta = 48°$ and the pathlength increases by 50%.

When including the adjustments for the reflected fraction and the increased pathlength, the calculated solar irradiance is now denoted as $Z_\lambda$ for solar noon values, which is the time of day when the SZA is at its minimum (eqn (4)).\textsuperscript{12} Since the increased pathlength is incorporated into the solar irradiance values, then $\Delta z$ should be set to one in eqn (3). For 40°N latitude during midseason summer and at solar noon, the SZA = 20°. Therefore, $\approx 2.1\%$ of the direct beam sunlight is reflected and the pathlength is increased by $\approx 3.4\%$. On a clear sky day, approximately 90% of the global horizontal irradiance is direct beam sunlight at solar noon (40°N); therefore, there should be only a minor adjustment between $I_{0\lambda}$ and $Z_\lambda$ values ($\approx 2\%$ increase) and between $W_\lambda$ and $Z_\lambda$ values ($\approx 5\%$ increase).

$$Z_\lambda = I_{0,\text{direct}}(1 - \text{fraction reflected})\sec \theta + 1.2I_{0,\text{diffuse}}(1 - 0.07)(4)$$

Lastly, there are two important differences between the $Z_\lambda$ reported by Zepp and Cline and the $L_\lambda$ reported by Leifer that are used to calculate environmental photochemical half-lives. The first is that $Z_\lambda$ values are for solar noon while $L_\lambda$ values are day-averaged; therefore, the $L_\lambda$ values are more appropriate to use when half-lives are $>1$ day. The second important difference is that $L_\lambda$ values have the factor of 2.303 incorporated into the reported solar irradiance (eqn (5a)–(5c)). The actual calculated solar irradiance values are a factor of 2.303 lower, so $L_\lambda$ values must be used carefully.

$$k = 2.303\phi\sum_{\lambda}^\ell (e_\lambda I_{\lambda0}\Delta \lambda) \text{ or } (5a)$$

$$k = 2.303\phi\sum_{\lambda}^\ell (e_\lambda Z_{\lambda}(noon)\Delta \lambda) \text{ or } (5b)$$

$$k = \phi\sum_{\lambda}^\ell (e_\lambda L_{\lambda}(day-averaged)\Delta \lambda) \text{ or } (5c)$$

2 Methods

2.1 Data sources

National Renewable Energy Laboratory data (NREL; Golden, CO). Global horizontal spectral irradiance data were downloaded from the NREL website for 2014, which was the last available full year for this data type.\textsuperscript{17,18} The spectroradiometer was a LICOR LI-1800 with a nominal bandpass of 6 nm but collected data every 2 nm from 300–1100 nm. This instrument is not temperature-controlled, but the error is expected to be minor with a change in measurement of 0.1% per °C at 350 nm.\textsuperscript{18} A previous intercomparison study, which included this spectroradiometer, showed up to a 10% deviation relative to the average measurement in the study with the largest deviations generally being observed in the ultraviolet region.\textsuperscript{19} Spectra were recorded every 5 minutes and averaged over the 24 hour day to obtain an average daily irradiance. The solar calendar, also available on the NREL website, was used to choose clear sky days (Table S1). The total column ozone measurements for nearby Boulder, CO were available from NOAA’s Global Monitoring Division website and were based on remote sensing data.\textsuperscript{20} The aerosol optical depths at 500 nm from the nearby Table Mountain, CO station were downloaded from NASA’s Aerosol Robotic Network (AERONET), which is a ground-based remote sensing network with a reported uncertainty of up to 0.02 for AOD measurements.\textsuperscript{21}

National Oceanic and Atmospheric Administration data (NOAA; Boulder, CO). Global horizontal spectral irradiance data for 2016 were provided by Patrick Disterhoft (NOAA). This spectroradiometer had a nominal bandpass of 0.75 nm but collected data every 0.2 nm from 285–450 nm. The instrument is temperature-controlled at 30 °C.\textsuperscript{22} The data were linearly interpolated to 1 nm intervals. Spectra were recorded at irregular time intervals, so measured spectra were compared to the SMARTS output for solar noon. The solar calendar available on the NREL website was used to choose clear sky days (Table S2). The same data sources for ozone and aerosol optical depth were used as described for the NREL data.

National Science Foundation polar programs UV monitoring data (NSF; Barrow, AK & Ushuaia, Argentina). Global horizontal spectral irradiance data for the last available summertime period were provided by Dr Germar Bernhard (Biospherical Instruments). For Barrow, AK, which is now known as Utqiaġvik and is located at 71.3°N, data for June 2016 were used. For Ushuaia, which is located at 54.8°S, data from December 2007...
were used. Both stations had an SUV-100 spectroradiometer with a nominal bandpass of 1.0 nm but recorded in 0.2 nm (280–340 nm), 0.5 nm (340–400 nm), or 1 nm increments (400–600 nm). This instrument is temperature-stabilized. Spectra were recorded every 15 minutes, but there were insufficient data to confirm the presence of full clear sky days. In fact, cloudy skies are typical at both sites during summer. Therefore, clear sky days were chosen based on the days with the highest UV-B irradiation, and the times with the highest UV-B radiation (between 13:00 and 13:30 local standard time) were ultimately used for comparison to the SMARTS model. The detection limit of one, is 5 × 10⁻⁶ W m⁻² for SZAs above 70° and 10⁻⁵ W m⁻² for SZAs below 70°. This typically corresponds to a cutoff wavelength between 290 and 305 nm depending on the time of year, SZA, and cloud cover. Total column ozone came from the OMI satellite and was downloaded from the NSF’s website.²³ AERONET sites were at or near the spectroradiometer station (site names Barrow and CEILAP-RG), but data were not available for the clear sky day at the Barrow station; however, both sites had consistently low aerosol optical depths and a value of 0.03 was used when measurements were not available.

2.2 Simple model of the atmospheric radiative transfer of sunshine (SMARTS) and tropospheric ultraviolet and visible (TUV) radiation model

SMARTS was developed by Dr Christian Gueymard at NREL for use in the U.S. Department of Energy’s Solar Energy Technologies Program.¹⁴ This radiative transfer model can calculate clear sky spectral irradiances in the ultraviolet and visible range with a resolution of 0.5–1 nm. This work used version 2.9.5 installed on a PC with Windows 10 operating system. SMARTS is available for download free of charge from the NREL website and has an Excel-based user interface.

Calculated irradiances depend on ten parameters that are specified in the SMARTS program, and the values used for data comparison are in Table 1. Specific values for each spectroradiometer site are in Tables S1–S3. Water vapor, specified as precipitable water, was available in the AERONET data alongside the aerosol optical depth (at 500 nm), and daily total column ozone data were available from remote sensing data; therefore, site and day-specific data were used for each of these parameters in the SMARTS model. A CO₂ concentration of 407 ppm was chosen based on recent measurements by NOAA at the Mauna Loa station.²⁴ For the gaseous absorption and aerosol model, the ‘light pollution’ and the ‘Shettle & Fenn rural’ options were chosen based on the example configurations provided with SMARTS and previous literature of model and data comparisons.²⁵ The average total solar irradiance was set to 1361 W m⁻², and the most recent extraterrestrial spectrum (Gueymard 2004) was chosen.²⁶–²⁸ The albedo can be specified in SMARTS, but it was set to zero (no albedo) for the data comparison and reference spectra.

The TUV model is available through the National Center for Atmospheric Research (NCAR) website.²⁹ The web interface allows for the calculation of spectral irradiance based on the location and date or the solar zenith angle. The altitude, ozone concentration, and aerosol properties can also be specified. In addition, a single cloud layer can be accounted for in the web interface. For the comparison to the SMARTS model, the noontime spectral irradiances were compared for the two NSF sites and for the NREL and NOAA data from March, June, September, and December.

2.3 Reference spectral irradiance

Updated reference spectra were also calculated using the values in Table 1. For aerosol optical depth and ozone, average values were used, which were 0.1 for aerosol optical depth³⁰ and latitude-specific averages for ozone (Table S4 and Fig. S1–S8†). Water vapor pressures were calculated from the chosen reference atmosphere.

Global horizontal spectral irradiance was calculated for each solar hour during daylight. The irradiances were summed and divided by 24 to obtain a daily average solar irradiance. Daily average and solar noon values were calculated at 0°, 10°, 20°, 30°, 40°, 50°, 60°, and 70° latitude in both the northern and southern hemisphere.

2.4 SMARTS sensitivity analysis

Although reference spectra are sufficient for many applications, it may be desirable to calculate spectral irradiance values specific to location and time as well as to understand the error that can be introduced by using a reference spectrum. To ascertain which input parameters were the most influential on the calculations, SMARTS sensitivity analysis was performed. Table 1 lists the parameters and calculation methods used for comparison against data and calculation of reference spectra.

### Table 1

| Parameter                | Value                                                                 |
|--------------------------|----------------------------------------------------------------------|
| Site pressure            | Calculated based on latitude and altitude                            |
| Atmosphere               | Appropriate selection of reference atmosphere based on site latitude and season |
| Water vapor              | Variable (in cm, from AERONET data) or calculated from reference atmosphere |
| Ozone                    | Variable (from AERONET data) or 0.1                                  |
| Gaseous absorption       | Light pollution                                                       |
| Carbon dioxide           | 407 ppm                                                               |
| Extraterrestrial spectrum| 1361 W m⁻² Gueymard 2004                                              |
| Aerosol model            | Shettle & Fenn rural model⁰                                           |
| Turbidity                | Variable (from AERONET data) or 0.1                                  |
| Albedo                   | None                                                                 |
the spectral irradiance output, a sensitivity analysis was performed.

Water vapor was adjusted from 0.5 to 4 cm based on the range observed in the AERONET dataset. Total column ozone varies substantially depending on location and time of year with values typically in the range of 250–450 Dobson units (DU, equivalent to 0.01 mm pathlength in ozone). From the AERONET data, the aerosol optical depth varied from approximately 0.005 to 0.5 (measured at 500 nm). Carbon dioxide absorbs infrared radiation, but does not absorb radiation below 2700 nm, so it has no effect on these spectra. Other gaseous species that absorb ultraviolet and visible light can be designated in the gaseous absorption menu. The impact of four trace gases (SO$_2$, NO$_2$, O$_3$, H$_2$O) on the spectral output was determined by changing the air pollution input between pristine, light, moderate, or severe.

3 Results and discussion

3.1 Comparison of current reference spectra ($L_i$) to SMARTS outputs

The modeled SMARTS output values were first compared to the reported $L_i$ values at 0°N, 40°N, and 70°N. These latitudes were chosen because ozone concentrations were given in Zepp and Cline (Fig. 8 caption in ref. 12), presumably for summer conditions, and were assumed to also be used to calculate $L_i$ values. To put the $L_i$ values on the same basis for comparison as the SMARTS output values, the wavelength range of each value was set to 1 nm using interpolation and the factor of 2.303 was removed by dividing by this value. The corrected $L_i$ values were generally higher than the $I_{0i}$ values calculated by SMARTS (Fig. 2 and 3).

To compare the total irradiance in the UVB (280–315 nm), UVA (315–400 nm), and photosynthetically active region (PAR, 400–700 nm), irradiances were converted to units of W m$^{-2}$ and summed across the designated wavelengths. The total irradiance values derived from $L_i$ values were 17–23% higher in the UVB region, 22–29% higher in the UVA region, and 7–12% higher in the PAR region compared to the SMARTS irradiance values. Some of this difference may be due to accounting for the increased pathlength in water that is done with $L_i$ values, but this should be at least partially offset by accounting for the fraction of light that is reflected off the water surface. Another potential reason for these differences is that SMARTS also accounts for more gaseous species that can absorb light before it reaches the Earth’s surface (e.g., SO$_2$ and NO$_2$).

3.2 Validation of SMARTS solar irradiance spectra

To assess the accuracy of SMARTS, the modeled irradiances were compared to data from four spectroradiometers maintained by three U.S. government agencies (Table 2). These spectroradiometers measured global horizontal irradiances over differing wavelength ranges and resolutions and were located at four separate monitoring facilities. It was hoped that by comparing measured and modeled irradiances across all of these instruments and measurement sites, we could assess the accuracy of the calculated day-averaged and solar noon irradiances.

It should also be noted that the resolution of the SMARTS model and the spectroradiometers differ. The SMARTS program has an option to smooth the irradiance signal, which is typically done when comparing measurements to the model, but this was not used in the comparisons presented. Therefore, the
ratio of modeled and measured irradiances was expected to vary around the value of one, even in the case of perfect agreement.

**NREL comparison.** Solar irradiance comparisons with the NREL spectroradiometer were possible from 300 nm up to the desired upper limit of 800 nm and, since irradiance was measured at regular time intervals, day-averaged irradiances could be compared. Comparison of spectral irradiances for four clear sky days in March, June, September, and December showed overall good agreement, but there were several features to note (Fig. 4A).

There is a sharp decrease and recovery of the modeled-to-measured ratio at 760 nm due to light absorption by oxygen. This is because (1) SMARTS restricts the light absorption of oxygen to 10 nm band, but the spectroradiometer measurements are over affected over ≈16 nm and (2) the decrease in irradiance due to O₂ is higher in SMARTS than in the measurements (90% vs. 40% decrease, respectively). Between 715 and 735 nm (and 685–705 nm to a lesser extent), inaccurate characterization of water vapor leads to the deviations observed. Lastly, at the lower end of the spectrum (300–306 nm), the measured irradiances exceeded the SMARTS modeled irradiances. However, this is likely due to the lower resolution of this spectroradiometer (6 nm bandwidth) and the sharp decline in irradiance in this range, so even a small percentage of longer wavelength photons (>306 nm) being attributed to shorter wavelengths (<306 nm) can have a significant effect.

Broadband measurements (i.e., UVB, UVA, and PAR) were also compared for 10 days during 2014 (Fig. 4B–D). SMARTS modeled irradiances over-predicted the measured irradiances by 9–29% in the UVB region (300–315 nm), 7–15% in the UVA region (315–400 nm), and 2–9% in the PAR region (400–700 nm). UVB and UVA modeled irradiances showed the greatest deviations from October to February, but there was no apparent pattern with season for the PAR irradiance.

**NOAA comparison.** The NOAA spectroradiometer is able to measure irradiance values from 280 to 450 nm, but the irradiance at lower wavelengths (<295 nm) typically does not exceed the detection limit of the spectroradiometer. Additionally, irradiance measurements were taken at irregular time intervals, so only noontime irradiances were used. Comparison of the spectral irradiance for four clear sky days in March, June, September, and December again showed relatively good agreement (Fig. 5A). Unlike the comparison with NREL data, SMARTS modeled irradiances exceeded the measurements for wavelengths < 310 nm. These deviations match the absorption band of ozone, which suggests the measured ozone concentrations were biased low for these four days at solar noon. In order to match the measured irradiances values, ozone concentrations

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**Table 2** Locations and altitude of the four spectroradiometers used to verify the solar irradiances calculated by SMARTS

| Site location     | Latitude | Longitude | Elevation (m) |
|-------------------|----------|-----------|---------------|
| Golden, CO, USA (NREL) | 39.742 N | 105.18 W  | 1829          |
| Boulder, CO, USA (NOAA)  | 39.99 N  | 105.26 W  | 1628          |
| Barrow, AK, USA (NSF)   | 71.3167 N | 156.6833 W | 8             |
| Ushuaia, Argentina (NSF) | 54.8167 S | 68.3167 W | 18            |

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**Fig. 4** Comparison of day-averaged (24 h) solar irradiance data measured by the NREL facility in 2014. In panel (A), the data are compared as a function of wavelength to SMARTS model outputs on the clear sky days of March 23 (green), June 19 (red), September 24 (purple), and December 19 (blue). In panels (B–D), the total irradiances in the UVB (300–315 nm), UVA (315–400 nm), and PAR (400–700 nm) regions are compared for the NREL data and SMARTS model for 10 days in 2014 (data were not available for clear sky days in July and November). The black line is the 1 : 1 lines and the grey shaded area represents a 10% deviation from the 1 : 1.
would need to be ca. 50 Dobson units (DU) higher (e.g., from 273 to 325 DU for June 18th). It is possible for the reported ozone concentrations to be inaccurate since the OMI satellite only takes daily measurements, and day-to-day variations of this magnitude were seen within the OMI data set. Additionally, higher temporal resolution measurements have shown large variations in ozone concentrations throughout the day. 

Because of this bias at wavelengths <310 nm, the comparison of UVB broadband irradiances were worse for the NOAA data than the NREL data with modeled irradiances being 0–35% higher than the measurements (Fig. 5B). This deviation may be smaller than expected based on Fig. 5B, but that is because there are 300 times fewer photons at 295 nm than at 310 nm, so the largest deviations have a relatively small contribution to the broadband UVB measurement. In contrast, the comparison of measured and modeled UVA irradiance values were scattered around the 1:1 line with modeled irradiances being 75–112% of the measurements (Fig. 5C). Additionally, the largest deviation, which was from November 18, 2016, appears to be a measurement error. The irradiance measured on this day (38 W m⁻²) was higher than any other clear sky day in November or October (Fig. S9†). On another clear sky day (November 12th), the UVA irradiance was measured to be 28 W m⁻², which is in better agreement to the SMARTS modeled value of 29 W m⁻².

NSF comparison. The last two spectroradiometers were chosen for comparison because of their locations closer to the poles where atmospheric influences on the surface solar radiation is most pronounced. For these two sites, the irradiance values were compared during summer because (a) more photons at lower wavelengths would be detectable and (b) this is when photodegradation would be most important in the arctic and subarctic regions. Similar to the NOAA data, only nonnoontime irradiances were compared and there was a bias at wavelengths <310 nm (Fig. 6). This time, however, the measured ozone concentrations are seemingly too high by 10–15 DU. Adjusting the ozone concentrations from 293 to 278 for the Ushuaia site and from 342 to 332 for the Barrow site brings the measured and modeled irradiances into very good agreement for the entire range (Fig. S10†). Despite this deviation, the broadband irradiance comparison for these two sites actually showed the best agreement with differences <2% for both UVB and UVA.

Fig. 5 Comparison of solar noon solar irradiance data measured by the NOAA facility in 2016. In panel (A), the data are compared as a function of wavelength to SMARTS data on the clear sky days of March 20 (green), June 18 (red), September 25 (purple), and December 10 (blue). In panels (B) and (C), the total irradiances in the UVA and UVB regions are compared for the NOAA data and SMARTS model for 13 days in 2016 (two clear sky days in June were included). The black line is the 1:1 line and the grey shaded area represents a 10% deviation from the 1:1.

Fig. 6 Comparison of clear sky summertime solar irradiance data measured by the NSF facilities in Barrow (Utqiaġvik), AK, USA (green, 71.3°N, June 2016) and Ushuaia, Argentina (purple, 54.8°S, December 2007).
3.3 Comparison of SMARTS and the TUV calculator

The spectral irradiances modeled by SMARTS were compared to those using the TUV calculator, which is typically used in the atmospheric research community. Overall, there was very good agreement between the two models with a -6.1% difference in the UVB, UVA, and PAR regions (Fig. S11†). TUV modeled irradiances were mostly higher than SMARTS for two apparent reasons (Table S5†). In SMARTS, the selection of “light pollution” adds 23 ppbv ground-level O3, which lowers the irradiances in the UVB region slightly. It also appears that the TUV calculator does not account for light absorption by water vapor in the same manner as SMARTS, which resulted in higher PAR irradiances when using the TUV calculator. The use of either model should give similar photodegradation half-lives; however, since SMARTS already tends to overestimate the irradiances measured by spectroradiometers and it provides the flexibility to specifying trace gas concentrations, the SMARTS irradiances are expected to be more accurate than the TUV irradiances.

3.4 Updated reference spectra

Reference irradiance spectra are provided as ESI† for solar noon and as day-averaged (24 h) values for 0° to 70° in 10° increments in both the northern and southern hemisphere and at four times of year when the equinoxes (solar declination = 0°, approximately June 21st and December 22nd) and solstices (solar declination = ±23.4°, approximately March 20th and September 22nd) occur. The irradiance values for the northern and southern hemisphere are not equal or mirrored because (a) the Earth’s distance to the sun varies throughout the year with the sun being ≈5 million km closer in early January compared to early July and (b) ozone concentrations differ in the northern and southern hemisphere (Table S2†).

Reference spectra were calculated using the SMARTS inputs outlined in Table 1. For the aerosol optical depth, a typical value of 0.1 (measured at 500 nm) was used.11 Water vapor pressures (in cm of water) were calculated from the reference atmosphere chosen. Since the irradiances in the UVB region were sensitive to the ozone concentration chosen, data from the OMI satellite, averaged across all longitudes, were retrieved for the last eight years from the Giovanni website (giovanni.gsfc.nasa.gov) and median concentrations were used for calculation of the reference spectra (Table S4 and Fig. S1–S8†).

These reference spectra are useful for calculating typical photodegradation half-lives (e.g., for pesticide registration applications) and for reporting environmentally relevant half-lives in the literature. It should be emphasized that these irradiance values are representative of the sunlight that reaches the surface of the Earth at sea level. They have not been adjusted for reflection off the water’s surface or increased pathlength within the water column like Z0 and L0 values. The decision to not adjust the irradiance values was made because these adjustments tend to increase the magnitude of the irradiance, and several other assumptions made (e.g., no competing absorbance from organic matter, no decrease in irradiance through the water depth, no clouds, and no shading) already make the unadjusted irradiance values a “best-case scenario” when determining the rate of photodegradation. SMARTS, however, does provide the flexibility to output direct and diffuse horizontal irradiances separately that can then be adjusted as previously described.12

3.5 Influence of input parameters on the solar irradiance

While the reference spectra are useful in many cases, it may be desirable to determine spatially and temporally specific irradiances when conducting field studies in which case SMARTS can be used to calculate new irradiance spectra. In addition to location and time, SMARTS also allows users to input values for several other parameters (see Table 1). Therefore, the influence of the input parameters was investigated to determine which parameters have the largest effect and in which region(s) of the irradiance spectrum. With this information, users can decide which input parameters should be prioritized for obtaining site-specific information as well as what magnitude of variability should be expected when choosing a generic value.

Aerosol properties. There are two inputs in SMARTS that pertain to aerosols. One is the selection of an aerosol model from the list of 11 pre-loaded models or to input user-defined aerosol properties if that information is available. The other input is the aerosol optical depth (AOD), which is a measure of light extinction due to aerosol scattering and absorption. There are several measures used to represent the AOD that can be selected in SMARTS. In this work, the AOD at 500 nm was used, but the AOD at 550 nm, Ångström’s or Schüepp’s turbidity coefficient, or the meteorological visibility can also be used. Measures of AOD are available from the AERONET ground stations (aeronet.gsfc.nasa.gov) and from several satellites (e.g., MODIS, MISR, and OMI; data available at giovanni.gsfc.nasa.gov).15

Aerosol properties had relatively minor effect on irradiance in most conditions investigated. The Shettle and Fenn rural aerosol model at an AOD of 0.1 was used to calculate the reference spectrum. If one of the urban aerosol models was selected instead, then the irradiance would decrease by 1–9% depending on the wavelength, and if the maritime aerosol model was selected the irradiance would increase by up to 2% (Fig. S12†). Changing the AOD also affects the irradiance at all wavelengths. With the baseline Shettle and Fenn rural aerosol model, going from an AOD of 0.005 to 1.0 decreases irradiance by 50% at 300 nm but only 20–30% in the visible light range (Fig. S12†).15 However, an AOD of 1.0 represents a high aerosol load (i.e., a severe pollution event or dust storm). At a more common AOD of 0.5 the irradiance would decrease <20%, and at typical value of 0.1 irradiance decreases by <10% compared to an AOD of 0.005.

Gaseous absorption. Several gaseous species can be inputted into SMARTS (Table 1 and S6†). The carbon dioxide concentration does not have an effect since its first absorption band is in the infrared region. However, some of the more trace gas species can have a large influence on certain regions of the spectrum. These inputs can be changed in the water vapor, ozone, and gaseous absorption menus.

Ozone has a strong absorption band in the UVB region and a minor absorption band in the 500–700 nm range, and
satellite-based ozone measurements are available (giovanni.gsfc.nasa.gov). The global average for ozone concentration is 300 DU, but it can vary by as much as 200 DU depending on the location and time of year.32,36 These ozone concentration variations can lead to order of magnitude differences in irradiance depending on the wavelength (Fig. 7A and S13†). For the total UVB irradiance (280–315 nm), a decrease of 50 DU increases the irradiance by 16% and an increase in 100 DU decreases the irradiance by 24%; however, the larger changes at the lower wavelengths can have a significant effect on the photodegradation of organic pollutants that absorb light only in this range (<300 nm). For the minor absorption band, irradiance changes <5% going from 300 to 450 DU.

In the gaseous absorption menu, 4 pre-loaded pollution scenarios are available. Light pollution was chosen for the reference spectra, but the selection of a pristine or severe pollution atmosphere can have a large influence on irradiance (Fig. 7A and S13†). This selection actually changes the concentrations of 10 atmospheric gases, but only three were found to be responsible for the changes in irradiance (Fig. 7B and S13†).

As seen previously, ozone concentrations had a large influence in the UVB region but so did sulfur dioxide (SO2) concentrations (Fig. S12†). Nitrogen dioxide (NO2) also had a large influence in the UVB, UVA, and visible regions (Fig. 7B and S12†). Concentrations of ozone, SO2, and NO2 concentrations can all be obtained from satellite-based measurements (giovanni.gsfc.nasa.gov).31

Water vapor can be calculated from the chosen reference atmosphere or a measurement can be inputted by the user. There are several absorption bands for water vapor starting at 500 nm, but the largest bands are 690–750 nm. Changing the precipitable water from 0.5 cm to 4 cm (tropical climate) can decrease irradiance by up to 50% depending on the wavelength (Fig. S14†). However, since all of water vapor’s absorption bands are in the visible region, it is not expected that changing water vapor values will have a significant impact on the photodegradation of chemicals in aquatic systems, which typically absorb light in the ultraviolet region.

Altitude and albedo. Changing the altitude in SMARTS does have an influence on irradiance but only because there are fewer particles and gases that can scatter and absorb the light at higher elevations. For every kilometer above sea level, there is about a 5%, 3%, and 1% increase in UVB, UVA, and PAR irradiance, respectively (Fig. S14†).

SMARTS has many choices for albedo, which is the proportion of photons that are reflected off a surface. However, since SMARTS was originally designed for use in the solar power industry, adding an albedo selection only increases the calculated irradiance. The albedo was set to zero for the calculation of the reference spectra. The selection of an albedo, however, will only have a minor effect in most cases. For example, the calm ocean, coastal ocean, or grazing field selection will increase the irradiance by <3% at any given wavelength. In contrast, the selection of a snow covering does have a significant effect, especially in the ultraviolet region, with increases up to 70% for wavelengths around 320 nm (Fig. S14†). Therefore, on clear sky days in waterbodies located next to snow cover, photodegradation rates will be faster than those predicted by using the reference irradiance values.

4 Conclusions

In this work, the solar irradiance spectra currently used for calculating photodegradation half-lives, as specified in the EPA and OECD guidelines (originally from Leifer 1988), were compared to the irradiance calculated by the SMARTS and TUV models. The currently used irradiances were consistently higher than SMARTS and TUV values; therefore, irradiances from all three models were compared to measurements from high-resolution spectroradiometers. The best agreement with measured data was observed for SMARTS irradiances. Yet, SMARTS-modeled irradiances still tended to be biased high, particularly in the UVB region where O3 and SO2 absorb, showing the importance of using accurate concentrations of light-absorbing species. Updated solar irradiance reference spectra were calculated using current values of light-absorbing gaseous species and aerosols for both the northern and
southern hemispheres. It is recommended that these updated and validated reference spectra should be used in the calculation of photodegradation half-lives. This will result in longer calculated half-lives, but the relative increase in the calculated half-life will depend on the absorption spectra of the chemical of interest.

**Conflicts of interest**

There are no conflicts to declare.

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**References**

1 D. E. Latch, J. L. Packer, W. A. Arnold and K. McNeill, *Photochem. Photobiol.*, A, 2003, 158, 63–66.
2 K. M. Parker and W. A. Mitch, *Proc. Natl. Acad. Sci. U. S. A.*, 2016, 113, 5868–5873.
3 K. P. Mangalgiri and L. Blaney, *Environ. Sci. Technol.*, 2017, 51, 12310–12320.
4 N. C. Pflug, M. K. Hankard, S. M. Berg, M. O’Connor, J. B. Golier, E. P. Kolodzie, D. M. Cwiertny and K. H. Wammer, *Environ. Sci.: Processes Impacts*, 2017, 19, 1414–1426.
5 J. K. Challis, M. L. Hanson, K. J. Friesen and C. S. Wong, *Environ. Sci.: Processes Impacts*, 2014, 16, 672–696.
6 K. McNeill and S. Canonica, *Environ. Sci.: Processes Impacts*, 2016, 18, 1381–1399.
7 C. K. Remucal, *Environ. Sci.: Processes Impacts*, 2014, 16, 628–653.
8 C. M. Glover, S. P. Mezyk, K. G. Linden and F. L. Rosario-Ortiz, *Chemosphere*, 2014, 111, 596–602.
9 U.S. EPA, *Fate Transport, and Transformation Test Guidelines: 835.2210 Direct Photolysis Rate in Water by Sunlight*, Report EPA 712-C-98-060, 1998.
10 OECD, *Test No. 316, Phototransformation of Chemicals in Water – Direct Photolysis*, 2008.
11 A. Leifer, *The kinetics of environmental aquatic photochemistry: theory and practice*, American Chemical Society, 1988.
12 R. G. Zepp and D. M. Cline, *Environ. Sci. Technol.*, 1977, 11, 359–366.
13 J. L. Packer, J. J. Werner, D. E. Latch, K. McNeill and W. A. Arnold, *Aquat. Sci.*, 2003, 65, 342–351.
14 NREL, SMARTS: Simple Model of the Atmospheric Radiative Transfer of Sunshine, https://www.nrel.gov/grid/solar-resource/smarts.html, accessed November 29, 2018.
15 J. M. Parnis and K. B. Oldham, *J. Photochem. Photobiol., A*, 2013, 267, 6–10.
16 R. G. Zepp, in *Dynamics, exposure and hazard assessment of toxic chemicals*, ed. R. Haque, Ann Arbor Science, Ann Arbor, 1980, ch. 9, pp. 69–110.
17 A. Andreas and T. Stoffel, NREL Solar Radiation Research Laboratory (SRRL): Baseline Measurement System (BMS); Golden, Colorado (Data), Report DA-5500-56488, National Renewable Energy Lab.(NREL), Golden, CO (United States), 1981.
18 NREL, Solar Radiation Research Laboratory Baseline Measurement System, http://midcdmz.nrel.gov/srrl_bms/, accessed November 29, 2018, DOI: 10.5439/1052221.
19 A. Habte, A. Andreas, L. Ottoson, C. Gueymard, G. Fedor, S. Fowler, J. Peterson, E. Naranen, T. Kobashi, A. Akiyama and S. Takagi, *Indoor and Outdoor Spectroradiometer Intercomparison for Spectral Irradiance Measurement, Report NREL/TP-5D00-61476*, 2014.
20 NOAA Earth System Reasearch Laboratory Global Monitoring Division, ESRL/GMD FTP Data Finder, https://www.esrl.noaa.gov/gmd/dv/data/, accessed November 29, 2018.
21 Goddard Space Flight Center, AERONET Aerosol Robotic Network, aeronet.gsfc.nasa.gov/, accessed November 29, 2018.
22 S. Wuttke, G. Seckmeyer, G. Bernhard, J. Ehramjian, R. McKenzie, P. Johnston and M. O'Neill, *J. Atmospheric Ocean. Technol.*, 2006, 23, 241–251.
23 P. F. Levelt, J. Joiner, J. Tamminen, J. P. Veefkind, P. K. Bhartia, D. C. Stein Zweers, B. N. Duncan, D. G. Streets, H. Eskes, R. van der A, C. McLinden, V. Fioletov, S. Carn, J. de Laat, M. DeLand, S. Marchenko, R. McPeters, J. Ziemke, D. Fu, X. Liu, K. Pickering, A. Apituley, G. González Abad, A. Arola, F. Boersma, C. Chan Miller, K. Chance, M. de Graaf, J. Hakkarainen, S. Hassinen, I. Ialongo, Q. Kleipool, N. Krotkov, C. Li, L. Lamsal, P. Newman, C. Nowlan, R. Suleiman, L. G. Tilstra, O. Torres, H. Wang and K. Wargan, *Atmos. Chem. Phys.*, 2018, 18, 5699–5745.
24 NOAA Global Greenhouse Gas Reference Network, Trends in Atmospheric Carbon Dioxide, https://www.esrl.noaa.gov/gmd/ccgg/trends/weekly.html, accessed May 30, 2018.
25 C. A. Gueymard, *Sol. Energy*, 2008, 82, 260–271.
26 NASA Goddard Media Studios, Solar Variability and Total Solar Irradiance (TSI), https://svs.gsfc.nasa.gov/10396, accessed May 30, 2018.
27 O. Coddington, J. L. Lean, P. Pilawski, M. Snow and D. Lindholm, *Bull. Am. Meteorol. Soc.*, 2016, 97, 1265–1282.
28 C. A. Gueymard, *Sol. Energy*, 2004, 76, 423–453.
29 National Center for Atmospheric Research, Quick TUV Calculator, http://crpm.acom.ucar.edu/Models/TUV/Interactive_TUV/, accessed November 29, 2018.
30 E. P. Shettle and R. W. Fenn, *Models for the aerosols of the lower atmosphere and the effects of humidity variations on..."
their optical properties, Air Force Geophysics Lab Hanscom
Alb Ma, 1979.
31 NASA Goddard Media Studios, Aerosol Optical Thickness,
MODIS, 2000–2016, https://svs.gsfc.nasa.gov/12302,
accessed July 14, 2018.
32 M. Kroon, J. P. Veenkind, M. Sneep, R. D. McPeters,
P. K. Bhartia and P. F. Levelt, J. Geophys. Res.: Atmos., 2008,
113, D16S28.
33 C. Hill and R. L. Jones, J. Geophys. Res.: Atmos., 2000, 105,
9421–9428.
34 J. Kim, J. Kim, H. K. Cho, J. Herman, S. S. Park, H. K. Lim,
J. H. Kim, K. Miyagawa and Y. G. Lee, Atmos. Meas. Tech.,
2017, 10, 3661–3676.
35 H. Bibi, K. Alam, F. Chishtie, S. Bibi, I. Shahid and
T. Blaschke, Atmos. Environ., 2015, 111, 113–126.
36 NASA Gooddard Space Flight Center, NASA Ozone Watch,
https://ozonewatch.gsfc.nasa.gov/facts/dobson.html,
accessed May 30, 2018.