

Accepted Manuscript

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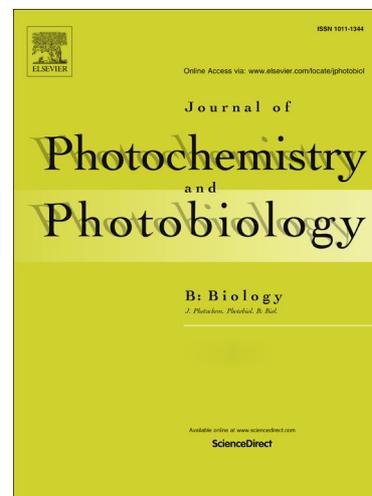
PII: S1011-1344(14)00042-6
DOI: <http://dx.doi.org/10.1016/j.jphotobiol.2014.02.005>
Reference: JPB 9668

To appear in: *Journal of Photochemistry and Photobiology B: Biology*

Received Date: 13 November 2013
Revised Date: 22 January 2014
Accepted Date: 6 February 2014

Please cite this article as: E. Navarro, S. Muñiz, M. Korkaric, B. Wagner, M.d. Cáceres, R. Behra, Ultraviolet radiation dose calculation for algal suspensions using UVA and UVB extinction coefficients, *Journal of Photochemistry and Photobiology B: Biology* (2014), doi: <http://dx.doi.org/10.1016/j.jphotobiol.2014.02.005>

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1 **Ultraviolet radiation dose calculation for algal suspensions using UVA and UVB**
2 **extinction coefficients**

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11

12 **Highlights**

13

14 • Extinction coefficients (k) for UVA and UVB were calculated using algal suspensions.

15 • Under controlled growth conditions, k may be estimated using OD₇₅₀.

16 • Beer-Lambert's law definite integral was used to calculate average UVR intensity.

17 • UVR dose calculations were 2-40 times more accurate than values obtained with previous
18 approaches.

19

20

21

22

23

24 **Abstract**

25

26 Although the biological importance of ultraviolet light (UVR) attenuation has been recognised in
27 marine and freshwater environments, it is not generally considered in in-vitro ecotoxicological studies
28 using algal cell suspensions. In this study, UVA and UVB extinction were determined for cultures of
29 algae with varying cell densities, and the data were used to calculate the corresponding extinction
30 coefficients for both UVA and UVB wavelength ranges. Integrating the Beer-Lambert equation to
31 account for changes in the radiation intensity reaching each depth, from the surface until the bottom of
32 the experimental vessel, we obtained the average UVA and UVB intensity to which the cultured algal
33 cells were exposed. We found that UVR intensity measured at the surface of *Chlamydomonas*
34 *reinhardtii* cultures lead to a overestimation of the UVR dose received by the algae by 2 to 40 times.
35 The approach used in this study allowed for a more accurate estimation of UVA and UVB doses.

36

37 **Keywords (up to 6)**

38 Algal cultures, UVR extinction, *Chlamydomonas reinhardtii*, *Synechocystis*, spectroradiometer,
39 radiometer

40

41 **1. Introduction**

42

43 **1.1. UVR measurement in ecotoxicological studies.** Ozone decline during the 1980s and 1990s
44 increased the amount of ultraviolet radiation (UVR) reaching the Earth's surface [1], raising concern
45 about the impacts of UVR on biological systems [2]. Since then, a huge research effort has focused on
46 assessing and predicting UVR impacts on human and natural ecosystems at different biological scales
47 [2].

48

49 Because of the importance of algae and macrophyta as primary producers in aquatic environments, the
50 effects of UVR effects on these organisms has been studied extensively [3, 4]. In aquatic environments,
51 numerous materials absorb and scatter light, contributing to vertical light attenuation. These include
52 dissolved substances [5, 6], particles [7, 8] and organisms [9]. Vertical light attenuation processes
53 result in a decrease of the intensity and changes in the light spectrum. While light attenuation is
54 routinely considered in ecological studies of UVR [10], it is not often considered in ecotoxicity studies
55 using algae or other aquatic organisms [11, 12]. In such studies, the responses of organisms are
56 generally related to doses of UV that are calculated using measurements of UV intensity at the surface
57 of experimental vessels or liquid media. While UVR extinction may be irrelevant under certain
58 conditions (i.e., low cell densities, low depth of experimental vessels and high media transparency),
59 the cell concentrations required for laboratory studies are expected to significantly attenuate UVR
60 intensity [13, 14]. Clearly, in that way, doses received by algae cells during UVR exposure can be
61 overestimated, leading researchers to conclude that the observed effects occur at lower doses than the
62 real ones [15-19].

63

64 **1.2. Integrative approaches for PAR modelling and UVR extinction assessment.** Standardised
65 procedures have been proposed for calculating the vertical attenuation coefficients for photo-
66 synthetically active radiation (PAR) models based on phytoplankton suspensions [13, 20]. In the case
67 of UVR, recent studies have demonstrated that simple laboratory measurements allow for establishing
68 reliable relationships between concentrations of optically active substances (such as chlorophyll,
69 dissolved organic matter and total suspended solids) and the underwater UVR light climate in natural
70 systems [10].

71

72 These approaches are based on the theoretical assumption that light passing through a dilute
73 suspension of cells should obey Beer-Lambert's law [10]. Accordingly, optical density is proportionate
74 to the number of cells [21, 22]. Nonetheless, other factors, such as cell size, cell shape and intracellular
75 pigment concentration, can modify light attenuation in a cell suspension [13, 20]. Even if the major
76 factors that determine light extinction characteristics in cell suspensions are identified, the specific
77 absorption of light is not a linear function of either pigment concentration or cell size [23]. The non-
78 linearity of light absorption results from the "package effect", which represents the decreased light
79 absorption of pigments contained in particles relative to the absorption of the same pigments in
80 solution [13]. Nevertheless, studies of various phytoplankton species have shown significant
81 relationships between cell size and volume and light absorption, indicating that morphological and
82 population parameters may be useful for developing improved models that link biological and optical
83 properties [14]. At present, there is a lack of methodological approaches for laboratory studies that
84 consider UVR extinction.

85

86 **1.3. Objectives of the study.** The goal of this study was to improve the measurement of UVR intensity
87 in algal cultures under agitation (assuming that the cells would be moving throughout the entire
88 medium in the vessel) by calculating UVR extinction coefficients. A spectroradiometer that provides a
89 spectral power distribution (power per unit area per unit wavelength) was used. We also tested a
90 radiometer equipped with simpler UVA and UVB integrative sensors. UVA and UVB extinction were
91 measured using different algal densities of two strains of *Chlamydomonas reinhardtii* and
92 *Synechocystis sp.* to test the robustness of our proposed approach. The extinction coefficients (k) for
93 UVA and UVB were calculated and modelled as a function of OD (the optical density, a proxy for
94 number of cells). We demonstrated that our approach was a rapid method for using light absorption to
95 estimate k_{uva} and k_{uvb} with algal cell suspensions.

96

97 **2. Materials and methods**

98

99 **2.1. Algal cultures.** The experiments were performed in various labs over various time periods, using
100 the cultures available at each lab: *Chlamydomonas reinhardtii*, CC-125 (Chlamydomonas Resource
101 Center, Univ. Minnesota, MN 55108, USA), and 137C+ 83.81 (Institute for Plant Physiology
102 (University of Göttingen, Germany). The two strains of this eukaryote algae showed differences in a

103 few genome sequences [24]. The culture medium was prepared according to the procedures described
104 by Le Faucheur et al. [25]. The algae were grown in a HT Multitron (Infors, Bottmingen, Switzerland)
105 at 25°C with continuous illumination of 120 $\mu\text{Mol photons sec}^{-1} \text{ cm}^{-1}$ (Philips Coolwhite TLD 15 W
106 fluorescent lamps) and shaken at 90 rpm. For cell number and cell volume measurements, a 200 μL
107 sample of the cell suspension was added to a final volume of 4 mL Isoton II diluent solution
108 (Beckman Coulter) and then counted using a Z2 Coulter Particle Counter (Beckman Coulter, Nyon,
109 Switzerland) within the 2.7-10.64- μm size range.

110
111 *Synechocystis* PCC 6803 wild type (Cyanophyta) was obtained from the Pasteur culture collection
112 (Institute Pasteur, France). The culturing procedure was similar to that described for *C. reinhardtii*, but
113 a culture medium (BG-11) was used [26] and the illumination was reduced to approximately 40 μMol
114 photons $\text{sec}^{-1} \text{ cm}^{-1}$. This photosynthetic prokaryote was selected to test the reliability of the
115 experimental approach with smaller sized algae. The word algae will be used through text indistinctly
116 for both the prokaryote and the eukaryote specie.

117
118 Experimental batch cultures were prepared by transferring an inoculum of algae in the exponential
119 growth phase at a starting density of approximately 6×10^5 cells mL^{-1} . Cell densities (see details in
120 Table 1) were obtained by centrifuging (3000 rpm, 10 min) the experimental batch cultures and then
121 resuspending the cell pellet until the desired density was reached.

122
123 For the estimation of the chlorophyll *a*, algal cells from sampled aliquots were centrifuged at
124 acceleration $16,000 \times g$, resuspended in 80% cold acetone and incubated for 5 min on ice in the dark.
125 Following another centrifugation step, the supernatants were transferred into a transparent 96-well
126 plate and absorption was measured at wavelengths 750, 663, 647, and 470 nm for calculating
127 concentrations according to a previous development [27].

128
129 **2.2. UVR intensity measurements.** Two different systems were used. The first system consisted of a
130 borosilicate beaker placed under a fluorescent lamp (Arimed B, 40 W; Cosmedico Light, Germany)
131 with an emission maximum between 330 and 350 nm (Fig. 1). Because borosilicate is transparent to
132 UVR radiation, the beaker walls were wrapped with black tape to prevent UVR light from entering
133 from a lateral surface). The spectroradiometer (JAZ-EL 200, Ocean Optics, 830 Douglas Ave.,

134 Dunedin, FL, USA) was placed under the beaker and a full spectrum scan was recorded that
135 represented the UVR intensity on the bulk surface. The beaker was then filled step-by-step with the
136 appropriate volume of algal culture (of a certain cell density), such that with each step the culture
137 depth (z) increased by 1 cm. UVR intensity was recorded after a few seconds (3-5 s), as soon as the
138 UVR values had stabilised and prior to sedimentation of the cells. New aliquots were added using a 5
139 mL pipette, which provided sufficient flow to mix the entire liquid column. This assured a
140 homogeneous light pathway for UVR from the surface of the liquid to the sensor in the bottom of the
141 beaker for each sample. This method had been previously applied successfully to measure irradiance
142 extinction in a volume of water that was too shallow to use a radiometer [10] by placing a “vessel” on
143 top of a sensor and repeatedly refilling the vessel with water.

144
145 Each recorded spectrum was integrated at a radiant flux density (Watt m^{-2}) in the range of UV-B (280-
146 320 nm) and UV-A (320-400 nm) and plotted against z (Fig. 2) using Ocean Optics SpectrasuiteTM
147 software. The ranges for energy integration were chosen to fit the ranges of the integrative sensors of
148 Solar Light that were used in the second measuring system.

149
150 The second system used to quantify UVR intensity consisted of a PMA2100 Radiometer (Solar Light
151 Co., Oak Line, USA) equipped with two sensors (a UVA-2110 WP sensor integrating energy in the
152 320-400 nm wavelength range and a UVB-2106 WP sensor integrating energy in the 280-320 nm
153 range). The sensors were placed 55 cm away from a UV lamp equipped with either an Osram
154 HTC400-241 bulb (Fig. 1) or fluorescent tubes (Arcadia D3 Reptile Lamp T5, 39 W, 12% UVB, 30%
155 UVA). The algal suspensions were added to the upper part of a 50-mL Uthermöl chamber fixed over
156 the UVR sensors. The upper part of the chamber was a tube (95 mm in length, 25 mm in diameter)
157 made with plastic that was opaque to UVR wavelengths. A bit of silicon was used around the bottom
158 of the tube where the tube made contact with the glass surface of the sensors to avoid leaching of the
159 cell suspension. UVR intensity measurements were made before sequentially adding 5-mL suspension
160 aliquots to completely fill the column. Each aliquot increased the height of the suspension column by 1
161 cm, thus allowing UVA and UVB intensity data to be plotted as a function of depth.

162
163 **2.3. Experimental design.** To test the robustness, feasibility and usefulness of this new
164 methodological approach, various experimental setups that combined different algal strains or species

165 and measuring systems were tested (Tab. 1). To examine the role of cell density in UVR extinction,
166 three different (i.e., independently developed) cultures of *C. reinhardtii* CC125 at varying cell
167 densities were exposed to UVR using the fluorescent tubes previously described. UVR extinction was
168 measured in a borosilicate beaker using a spectroradiometer (Exp. 1, 2 and 8). See details of the whole
169 experimental setup in Figure 2. To assess the influence of different UV measuring devices on
170 extinction calculations, an integrative radiometer was used (Exp. 7). *C. reinhardtii* (137C+ 83.81) was
171 used in Experiments 3, 4, 5 and 6). UVR extinction was measured in a Uthermöl chamber using HTC
172 lamps and the integrative radiometer. The potential effects of using different UVR lamps, measuring
173 devices and vessels were tested in Experiment 9. Finally, to test the influence of smaller-sized algae on
174 UVR extinction, *Synechocystis* PCC 6803 wild type (Institute Pasteur, France) was used (Exp. 10). In
175 this case, UVR extinction was measured in borosilicate beakers using the PMA2100 Radiometer.

176
177 **2.4. Statistics and modelling.** The UVA and UVB extinction data from all experiments were
178 adjusted to the Beer-Lambert equation (Equation 1) using Sigma Plot 12.5 (Systat Software Inc., San
179 Jose, USA). Excel was used for fitting the linear models. To assess the range of cell density for which
180 our approach was most robust, r squared modelling terms were used (see Figure 3C).

181
182 **2.5. UVA and UVB extinction coefficients.** The coefficients k_{uva} and k_{uvb} were estimated by using
183 the UVA and UVB intensity values from all experiments in the Beer-Lambert equation. All k values
184 are shown in Table 1. Later, the resulting UVA and UVB extinction coefficients were represented as a
185 function of the OD_{750} of the cell suspension (used as a proxy of cell density) and adjusted to a
186 polynomial equation (see details in Fig. 3B). This was designed to eventually allow for the estimation
187 of extinction coefficients based on the OD_{750} of the cell suspensions (details in Section 3.2).

188
189 **2.6. Calculating average UVA and UVB intensity received by algal cultures.** The Beer-Lambert
190 equation, integrated from depth 0 to m (expressed in cm, with m being the total depth of the solution),
191 was used to calculate the average energy received by an algal cell in a cell suspension presenting a
192 certain k_{uva} or k_{uvb} and exposed to I_0 (i.e., UV intensity) at the surface of the liquid (Equation 2). The
193 resulting value (intensity integration) was divided by depth (expressed in cm to maintain homogeneity
194 in the units). This value (the average UVR intensity received by a cell in the suspension) was
195 multiplied by exposure time to obtain the UVR dose received by the algal cells.

196

197 3. Results and discussion

198

199 **3.1. UVR extinction coefficients.** UVB and UVA radiation intensity decreased exponentially with
200 increasing culture depth in all experiments (Fig. 3A), following the Beer-Lamberts equation (Eq. 1).
201 This behaviour was independent of the device used to measure UVR, cell density or the type of vessel
202 used. The extinction coefficients k_{uva} and k_{uvb} showed a linear relationship with cell density (measured
203 as OD₇₅₀), as shown in Fig. 4. Accordingly, the highest k_{uva} and k_{uvb} values were obtained for the
204 suspensions with the highest cell density (Tab. 1).

205

206 Values for k_{uva} ranged from 0.0076 to 1.86 cm⁻¹, whereas k_{uvb} ranged from 0.010 to 3.79 cm⁻¹. The
207 higher values of UVB extinction compared to UVA extinction were consistent with the weaker
208 absorption of UVA by biomolecules present in cells [28]. The k values determined in our study were in
209 the range of those determined for water with comparable concentrations of chlorophyll *a*. The k value
210 for wavelengths under 400 nm was between 1-16 m⁻¹ [29] for a freshwater lake, between 1-2 m⁻¹ for
211 water with a chl-*a* concentration of 0.5-0.6 µg l⁻¹ and approximately 1 m⁻¹ for water with a chlorophyll
212 *a* concentration of 0.3 µg l⁻¹ [30]. In our study, cell suspensions with chlorophyll concentrations
213 between 0.4-0.8 µg l⁻¹ had a k_{uva} between 2 and 4 m⁻¹.

214

215 The use of different UVR measuring devices affected the dose calculations, depending on the UV
216 range selected (see Fig. 4). As shown in Fig. 4, the k_{uva} values were quite similar, regardless of the
217 device used. The cultures that clearly differed from the others in terms of UVR dose (Cultures 5 and 6)
218 were those with the highest cell densities. These differences can be attributed to the fact that the
219 sensors on the integrative spectroradiometer only record part of the incident energy (approximately
220 64%). This effect was even more apparent for UVB radiation than for the UVA range. For these
221 cultures, k_{uvb} was between 30 and 80% higher than k_{uva} . This difference reflected the higher extinction
222 of UVB (i.e., lower penetration in the water column) in the cell suspensions (see k values in Fig. 4-A1
223 vs. A2).

224

225 Regarding the application of this approach to differently sized algae, Fig. 4 shows that extinction
226 coefficients from experiment 10 (using *Synechocystis sp.*) shown similar values and trend that other
227 experiments using *Chlamydomonas* and the same measuring setup (exp. 10). The content of pigment

228 per cell (Tab. 1) did not shown any kind of influence on the extinction coefficients; correlations
229 resulted in $r^2 = 0.03$ for k_{uva} and 0.01 for k_{uvb} .

230

231 For long-term UVR exposure, time-dependent variability in k is an important factor to consider. High
232 UVR intensities may cause bleaching of cell pigments or cell death, thereby modifying k values.

233 Therefore, when using the proposed method for long-term studies, time-dependent variation in k
234 should be assessed and integrated into the calculations. In our case, exposures up to two hours did not
235 modify the k values (results not shown).

236

237 **3.2. Assessing the cell range for which OD₇₅₀ is a reliable estimator of k_{uva} and k_{uvb} .**

238 Experiments focusing on the effects of UVR on algal cultures use homogeneous algal populations that
239 have similar physiological states, pigment contents and size. Based on this practice and the previously
240 described rationale for PAR studies (see Section 1.2.), UVR extinction under controlled conditions
241 would vary only with cell density or any of its proxies, such as OD₇₅₀ (see Fig. 4). Therefore, data for k
242 and OD₇₅₀ have been fitted to a polynomial equation (see details in Fig. 3B). Eventually, this model
243 could allow for the estimation of extinction coefficients based on the OD₇₅₀ of a cell suspension.

244

245 The r^2 values for all extinction curves were represented as a function of the OD₇₅₀ and fitted to a
246 polynomial inverse second order equation (see details in Fig. 3). This approach identified the OD
247 values for which the experimental data showed a good fit to the Beer-Lambert model. That values
248 ranged from OD₇₅₀ = 0.07 (defined by the intersection of the $r^2 = 0.95$ and the upper confidence
249 interval of the polynomial model) to OD₇₅₀ = 1. At these elevated OD, extinction curves start worsening
250 the fitting to Beer-Lambert model. That OD range corresponds to a cell density range of 2.5E5 to
251 3.2E6 cells mL⁻¹ (*C. reinhardtii*). Finally, we noted that the lowest cell densities showed a poor fit to
252 the Beer-Lambert equation (i.e., lower r^2 values). This point will be further discussed in the practical
253 considerations section.

254

255 **3.3. Comparison of UVR doses calculated using different approaches.** We calculated the UVR
256 dose received by a cell suspension as a function of cell density using three methods: a) measuring the
257 UVR intensity reaching the surface of the cell suspension, b) calculating k_{uva} and k_{uvb} using a
258 spectroradiometer (more accurate) and c) calculating k_{uva} and k_{uvb} using a radiometer (less accurate).

259 UVR intensity reaching the cell suspension was 9.54 and 1.56 mW cm⁻² for UVA and UVB,
260 respectively, for 1 hour in a beaker (10 cm depth). For these calculations, we used data from
261 Experiment 1 and Experiment 2 (see Tab. 1 for details), and the doses are shown in Figure 5. Without
262 considering UVR extinction, UVR dose was not dependent on cell density (black points on Fig. 5A
263 and 5B). In contrast, UVR dose based on k was dependent on cell density and had lower values. UVA
264 dose was overestimated by 1.4 to 20 times, depending on the cell density. The overestimation was
265 higher for the highest cell densities. UVB dose was overestimated by 1.8 to 39 times. Even if the
266 radiometer measurements underestimated UVR dose compared to the spectroradiometer measurements,
267 the values obtained with the radiometer method provided a more realistic estimate of UVR dose than
268 the estimates obtained by measuring UVR intensity at the surface of the liquid.

269
270 **3.4. Practical considerations.** At high cell densities, UVB radiation fell below the detection limit
271 at $z > 1$; and the extinction coefficients were calculated using only two data points ($z = 0$ cm and $z = 1$
272 cm). Accordingly, in these cases almost all UVA and UVB light (99%) was attenuated within the first
273 centimetres ($z < 10$ cm) of the suspensions (see details in Tab. 1, last two columns). It is thus important
274 to do not use cell densities higher of that leading to the complete UVR attenuation.

275
276 Another effect observed with the beakers was that at $z > 4$, UVA and UVB intensities slightly
277 increased. This was due to the “lense effect” of the beaker walls, but could only be observed under
278 very transparent media conditions, such as in the samples with very low cell densities. This may also
279 help explain the poor fit of the data to the Beer Lambert equation for these samples. When the beaker
280 was almost full, more light was diverted towards the sensor that, at a lower culture depth, would have
281 been absorbed by the walls of the beaker (or more likely, by the black tape with which the beaker was
282 wrapped). For that reason, some of the data points were excluded from the model fitting. This effect
283 would be most likely be specific to each particular experimental setup, depending on the distance of
284 the lamp from the vessel and the type of vessel used. For this reason, and the differences that may arise
285 from using the different UVR measuring devices in the market, it is always need to calibrate the
286 experimental setup used (see section 3.1).

287
288 The similarity between k values calculated in this study and those measured in real environments
289 showed that our approach provides much more realistic calculations of UVR dose than those based in
290 UVR intensity at the surface of the liquid. The accuracy of k calculations may be improved by

291 considering sensor shape corrections (e.g., the use of common plane sensors versus the spherical
292 sensors used in open waters), and applying calibrations for the different media encountered by the light
293 pathway from the lamp to the sensor, the differences with our method would be small. Nevertheless,
294 the simplicity and affordability of the our approach, and the robustness (shown using different
295 measuring setups, different algal species and different pigment concentrations) result in more precise
296 UVR dose response calculations, allowing for comparing results among various natural and in-vivo
297 studies. This experimental approach would be also useful for other than photosynthetic cell
298 suspensions.

299

300 **Acknowledgments.** This work was supported by the project “Linking Ecology and Ecotoxicology:
301 Response of River Biofilms to Simultaneous and Sequential Exposure to UV-Radiation and Cadmium”
302 of the Velux Foundation (Switzerland), by the Spanish Ministry of Economy and Competitiveness
303 (National Research Plan, Ref. BFU2010-22053), and by EUROPA Fellowship Program from Caja
304 Inmaculada (Aragón, Spain). We also thank Professor Ruben Sommaruga for his very useful
305 comments during the writing of this manuscript, and to Maria Pata for her advising on statistics.

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309 **References**

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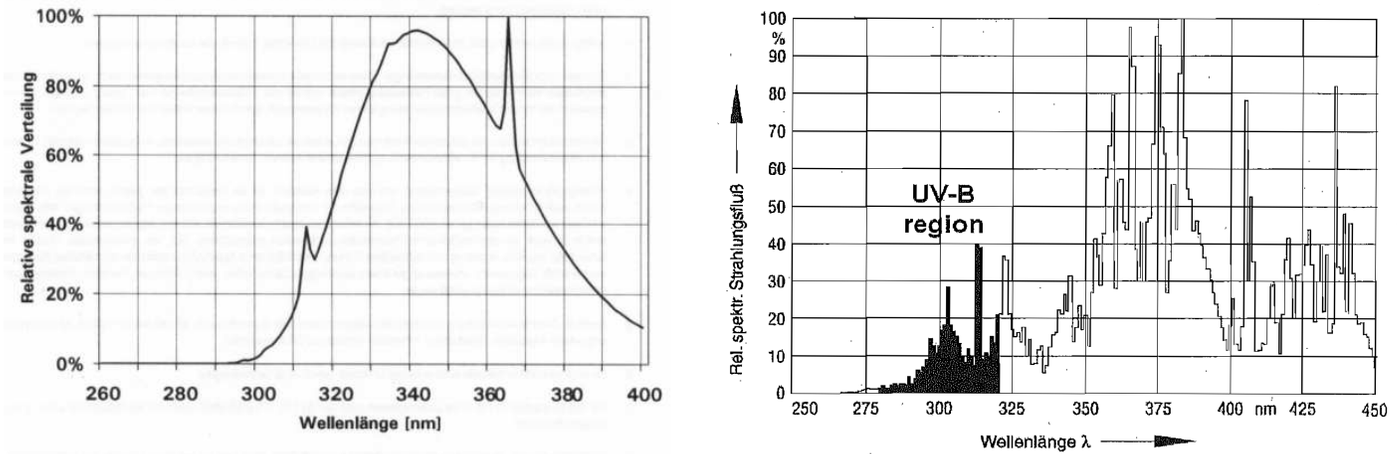
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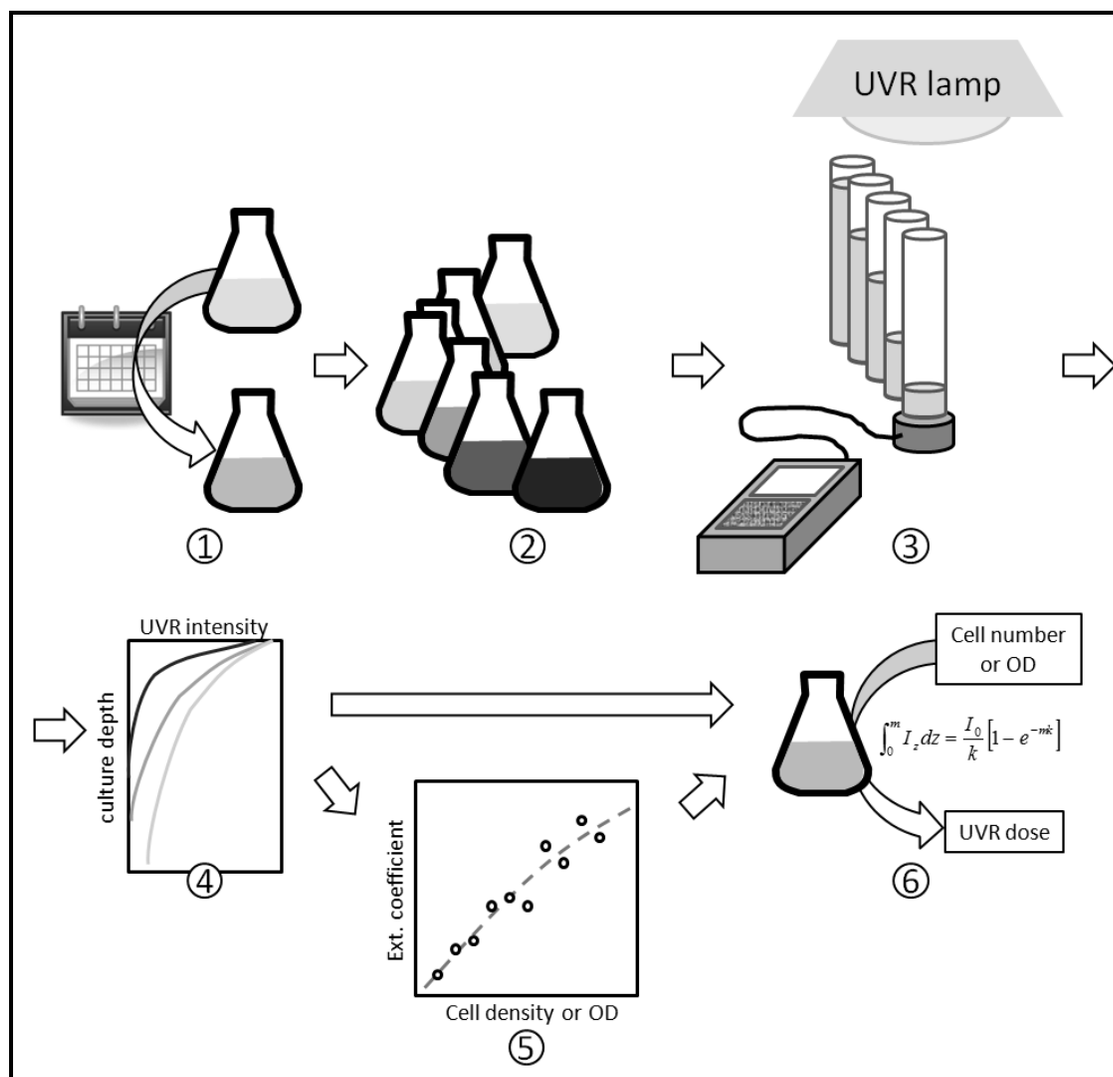
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403 **Figure 1.** Distribution of spectral power of the UV lamps used in this study. At left: ARIMED fluorescent lamps
404 (showing the 260-400 nm range). At right: OSRAM HTC bulbs (showing the 250-450 range and the UVB region,
405 in black).
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Figure 2. Diagram showing measurement setup and data processing. The method was applied to standardised

algal cultures (1). A dilution bank of the cultures is prepared (2). Extinction coefficients are measured by

sequentially adding fixed volumes of each algal dilution to a column exposed to UVR (3). Each time additional

volume is added to the column, the depth of the column increases, and UVA and UVB intensity is recorded.

The resulting UVR intensity data are plotted against depth and the Beer-Lambert equation is used to obtain

the extinction coefficient (k) for UVA and UVB (4). The k values were used to calculate the average intensity of

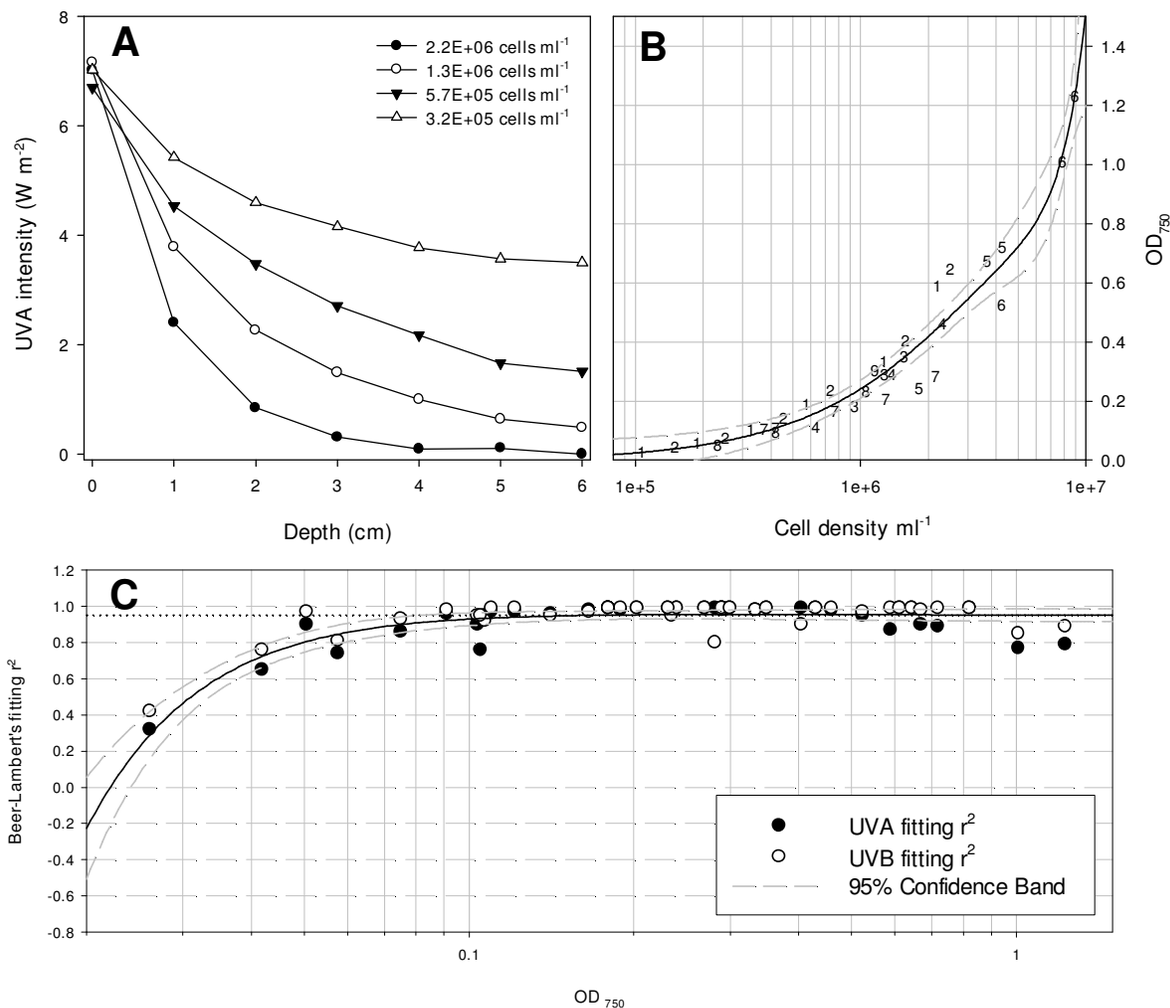
UVR received by a cell suspension (6). It is also possible to plot all k values calculated as a function of the

number of cells, or OD (5) and then fit these values to an appropriate model. The model could be used to

estimate the k of a given algal suspension.

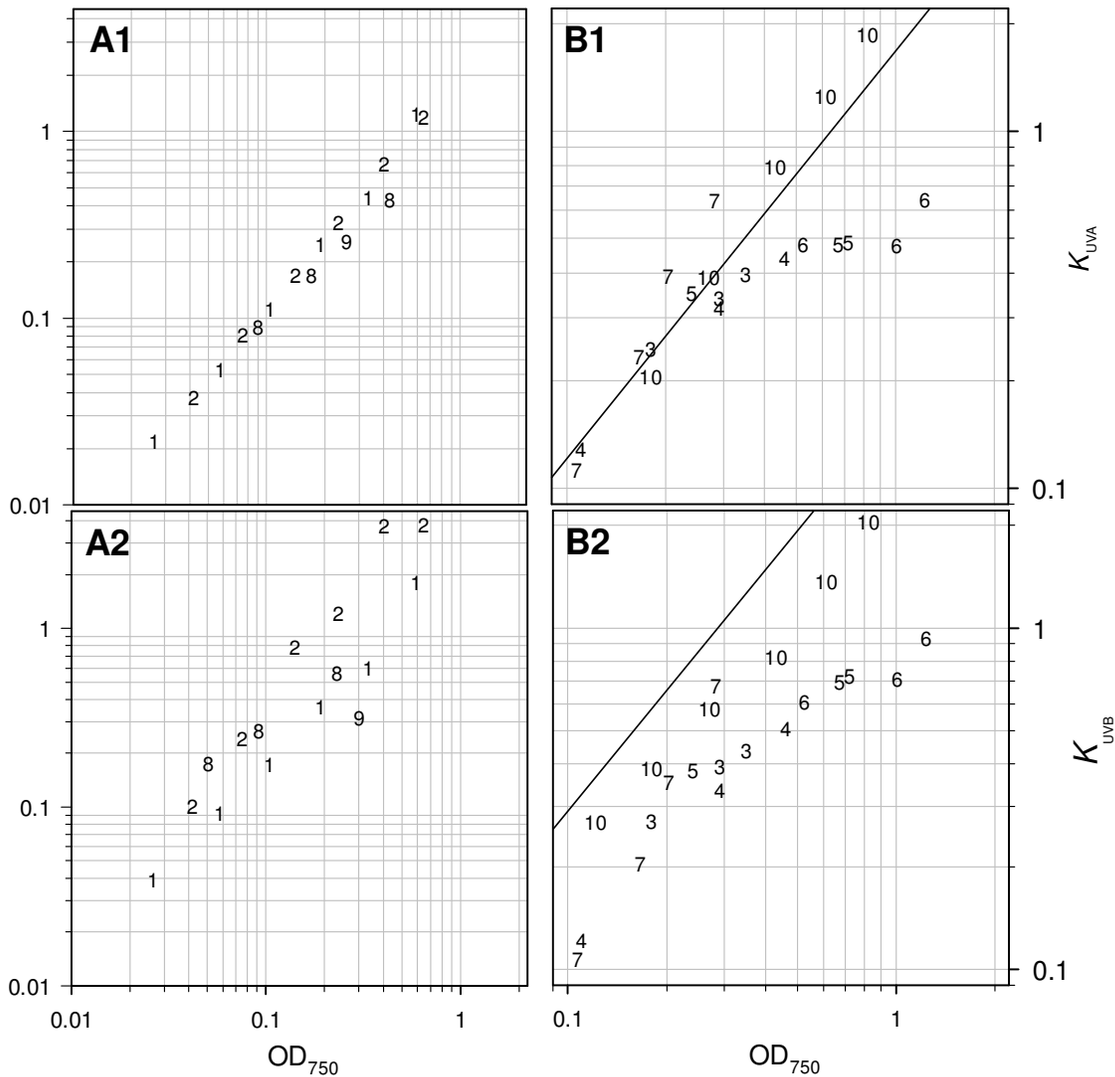
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 421 **Figure 3.** Graph A shows UVA extinction as a function of depth for four different cell densities.
 422 Values at 0 cm represent UVR intensity on the bulk surface. Graph B shows the correlation between
 423 OD_{750} and number of cells. The data showed a good fit ($r^2 = 0.94$) to a polynomial cubic equation (OD
 424 $= -0.0035 + 2.8\text{E}-7 \text{ cells} - 4.2\text{E}-14 \text{ cells}^2 + 2.8\text{E}-21 \text{ cells}^3$). Numbers (1 to 9) represent the different experiments
 425 (see Table 1). The black line represents the fitted model, and the long dashed grey lines are the 95%
 426 confidence intervals for that model. Graph C represents the r^2 fitted values of the Beer-Lambert
 427 equation for different values of OD (used as a proxy for cell density). The black dotted horizontal line
 428 represents $r^2 = 0.95$. White points represent k_{UVB} values and black points represent k_{UVA} values,
 429 regardless of the device used for measurement. The black line is the adjusted curve ($f(x) =$
 430 $y_0 + (a/x) + (b/x^2)$ with a r^2 of 0.75), with the 95% confidence intervals used as the criteria for selecting
 431 the applicability range of the method.

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Figure 4. Extinction coefficients for UVA (A1 and B1) and UVB (A2 and B2) calculated using a spectroradiometer (A1 and A2) or an integrative radiometer (B1 and B2) and represented as a function of the OD. Numbers represent the different experiments (see Tab. 1). For the purposes of comparison, the linear models of A1 and A2 are shown as lines in B1 and B2.

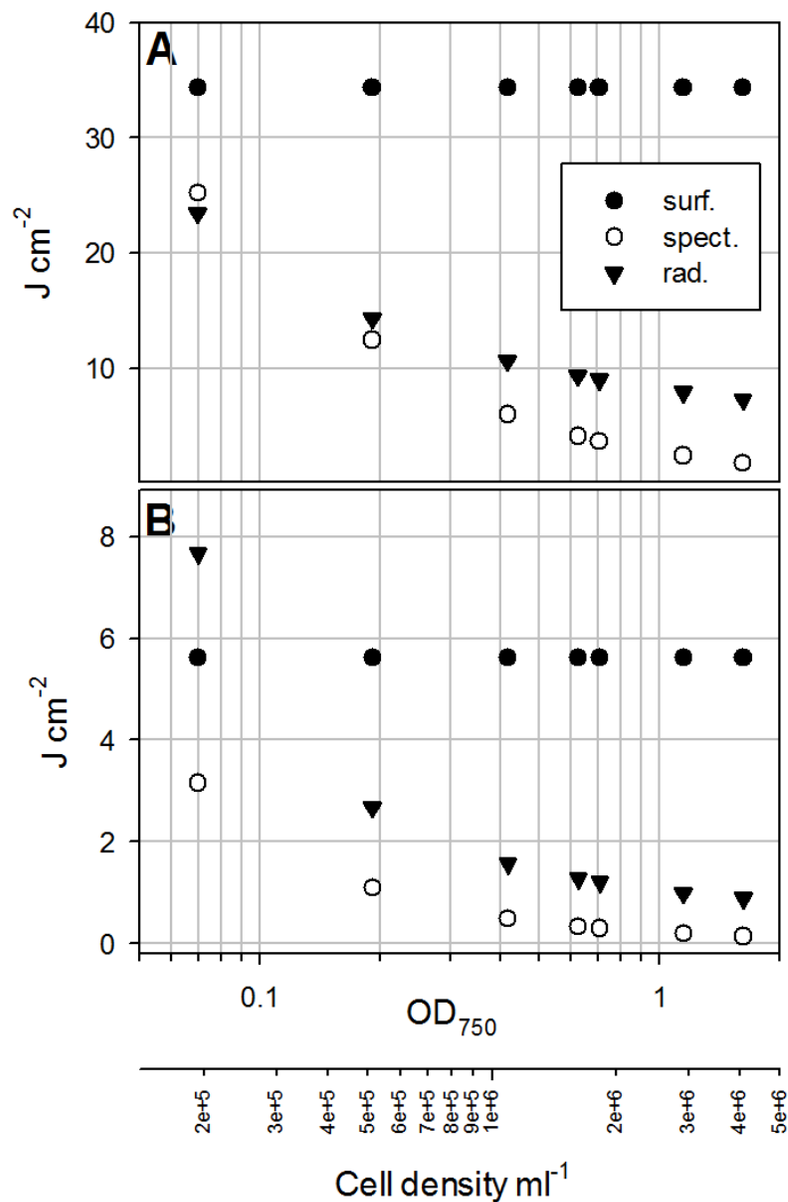
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Figure 5. Comparison of UVA (A) and UVB (B) calculated doses received by algal suspensions of varying cell densities based on different approaches. Surface measurements (surf.) are compared to methods based on k calculations using either a spectroradiometer (spect.) or radiometer (rad.). Results are represented as a function of the OD and cell density.

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$$452 \quad I_z = I_0 \cdot e^{-zk}$$

453 **Equation 1. Beer-Lambert's equation of light extinction.** I_z represents the light intensity at depth z (cm); I_0 is
454 the intensity at the surface of the bulk suspension; and k the extinction coefficient.

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$$I_z = I_0 \cdot e^{-zk}$$

$$\int_0^m I_z = I_0 \int_0^m e^{-zk}$$

$$= \frac{I_0}{-b} [e^{-zk}]_0^m$$

$$= \frac{I_0}{-b} [e^{-mk} - e^{-0k}]$$

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$$= \frac{I_0}{-b} [e^{-mk} - 1]$$

$$= \frac{I_0}{b} [1 - e^{-mk}]$$

458

$$\int_0^m I_z dz = \frac{I_0}{k} [1 - e^{-mk}]$$

459 **Equation 2.** The Beer-Lambert equation integrated for depths 0 to m (expressed in cm, with m being
460 the total depth of the solution) to allow for calculating the average energy received by each cell in a
461 given cell suspension with the coefficient k_{uva} or k_{uvb} and a UV intensity I_0 at the surface of the liquid.

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UVR	Measu.	Algae	Vessel	Exp.	OD ₇₅₀	K _{UVA}	r ²	K _{UVB}	r ²	pg chl cell ⁻¹	Z _{1%} UVA	Z _{1%} UVB
fluo	spectro	CC125	boro	1	0.5895	1.247	0.87	1.808	0.99	2.04	3.693	2.547
				1	0.3339	0.444	0.98	0.6	0.98	1.44	10.372	7.675
				1	0.1895	0.248	0.98	0.367	0.99	2.77	18.569	12.548
				1	0.1037	0.112	0.9	0.174	0.95	3.02	41.118	26.466
				1	0.0576	0.053	0.74	0.092	0.81	2.24	86.890	50.056
				1	0.0261	0.022	0.32	0.039	0.42	1.44	209.326	118.081
				2	0.6448	1.19	0.99	3.798	0.99	2.18	3.870	1.213
				2	0.4049	0.667	0.99	3.76	0.9	2.46	6.904	1.225
				2	0.2347	0.325	0.98	1.21	0.95	2.51	14.170	3.806
				2	0.1410	0.169	0.96	0.789	0.95	2.03	27.250	5.837
				2	0.0750	0.082	0.86	0.243	0.93	2.97	56.161	18.951
				2	0.0419	0.038	0.65	0.102	0.76	2.18	121.189	45.149
				8	0.2313	0.433	0.99	0.5608	0.99	-	10.635	8.212
				8	0.0911	0.1706	0.96	0.2683	0.98	-	26.994	17.164
8	0.0505	0.0897	0.9	0.1762	0.97	-	51.340	26.136				
8	0.0218	0.0779	0.83	0.2222	0.97	-	59.116	20.725				
fluo	radio	CC125	boro	7	0.2818	0.6393	0.99	0.6767	0.8	-	7.203	6.805
				7	0.2030	0.3923	0.99	0.3545	0.99	-	11.739	12.991
				7	0.1658	0.2332	0.98	0.2047	0.97	-	19.748	22.497
				7	0.1069	0.1124	0.95	0.1076	0.92	-	40.971	42.799
				7	0.1050	0.045	0.76	0.049	0.95	-	102.337	93.983
				7	0.1077	0.0076	0.59	0.0108	0.79	-	605.943	426.405
HTC	radio	137C	uterm	3	0.1800	0.2458	0.99	0.2724	0.99	1.32	18.735	16.906
				3	0.2900	0.3419	0.99	0.3934	0.98	1.53	13.469	11.706
				3	0.3500	0.3987	0.99	0.4393	0.99	1.10	11.550	10.483
				4	0.1100	0.1289	0.97	0.1224	0.99	1.03	35.727	37.624
				4	0.2900	0.3199	0.99	0.335	0.99	1.28	14.396	13.747
				4	0.4600	0.4404	0.98	0.5104	0.99	1.19	10.457	9.023
				5	0.2400	0.3531	0.99	0.3836	0.99	1.23	13.042	12.005
				5	0.6700	0.4817	0.9	0.6985	0.98	1.18	9.560	6.593
				5	0.7200	0.4887	0.89	0.7238	0.99	1.15	9.423	6.362
				6	0.5241	0.4809	0.95	0.6092	0.97	-	9.576	7.559
6	1.0095	0.4771	0.77	0.7099	0.85	-	9.652	6.487				
6	1.2299	0.645	0.79	0.9329	0.89	-	7.140	4.936				
fluo	spectro	137C	uterm	9	0.300	0.259	0.99	0.318	0.99	-	17.781	14.482
fluo	radio	Synech	boro	10	0.8219	1.8685	0.99	2.0554	0.99	-	2.465	2.241
				10	0.6123	1.2553	0.99	1.3719	0.99	-	3.669	3.357
				10	0.4302	0.7955	0.99	0.819	0.99	-	5.789	5.623
				10	0.2698	0.3889	0.99	0.5826	0.99	-	11.842	7.905
				10	0.1799	0.2051	0.99	0.3884	0.99	-	22.453	11.857
				10	0.1213	0.0898	0.97	0.2709	0.99	-	51.283	17.000
Models for k _{UVA}						Models for UVB						
k = 2.01 OD ₇₅₀ - 0.09 (r ² 0.97)						k = 5.36 OD ₇₅₀ - 0.19 (r ² 0.70)						
k = 0.18 ln(OD ₇₅₀) + 0.56 (r ² 0.91)						k = 0.31 ln(OD ₇₅₀) + 0.79 (r ² 0.96)						

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Table 1. Details of the six experimental setups (first column). UVR refers to the type of lamp used: *HTC* for halogen and *fluo* for fluorescence tubes; Measu. indicates the device used: *spectro* for the spectroradiometer and *radio* for the integrative radiometer; algae indicates the species; Vessel indicates the type of vessel used for measuring UVA/UVB extinction; Exp. indicates the use of different cultures (real replicates) and the sequence in which the experiments were performed; OD₇₅₀ indicates the optical density at 750 nm; and the k_{UVA}, k_{UVB} and their respective r² values are also shown (k units are cm⁻¹). It has been also shown the chlorophyll a concentration for selected experiments, expressed both as chlorophyll per volume (*chl a*) and as chlorophyll per cell (pg chl cell⁻¹). These last values have been calculated using the values obtained using the cell coulter (data not shown). The last two columns show the Z values at which 99% of the incident UVA and UVB have been attenuated. The last two rows in the table show the models used for calculating k as a function of OD by the method described in Section 3.2 of the manuscript.

476 **Highlights**

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478 • Extinction coefficients (k) for UVA and UVB were calculated using algal suspensions.479 • Under controlled growth conditions, k may be estimated using OD_{750} .

480 • Beer-Lambert's law definite integral was used to calculate average UVR intensity.

481 • UVR dose calculations were 2-40 times more accurate than values obtained with previous
482 approaches.

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