Variations of light absorption by suspended particles with chlorophyll a concentration in oceanic (case 1) waters: Analysis and implications for bio-optical models

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Abstract. Spectral absorption coefficients of total particulate matter $a_p(\lambda)$ were determined using the in vitro filter technique. The present analysis deals with a set of 1166 spectra, determined in various oceanic (case 1) waters, with field chl a concentrations $(\langle\text{chl}\rangle)$ spanning 3 orders of magnitude (0.02–25 mg m$^{-3}$). As previously shown [Bricaud et al., 1995] for the absorption coefficients of living phytoplankton $a_{ph}(\lambda)$, the $a_p(\lambda)$ coefficients also increase nonlinearly with (chl). The relationships (power laws) that link $a_p(\lambda)$ and $a_{ph}(\lambda)$ to (chl) show striking similarities. Despite large fluctuations, the relative contribution of non-algal particles to total absorption oscillates around an average value of 25–30% throughout the (chl) range. The spectral dependence of absorption by these non-algal particles follows an exponential increase toward short wavelengths, with a weakly variable slope (0.011 ± 0.0025 nm$^{-1}$). The empirical relationships linking $a_p(\lambda)$ to (chl) can be used in bio-optical models. This parameterization based on in vivo measurements leads to a good agreement with a former modeling of the diffuse attenuation coefficient based on in situ measurements. This agreement is worth noting as independent methods and data sets are compared. It is stressed that for a given (chl), the $a_p(\lambda)$ coefficients show large residual variability around the regression lines (for instance, by a factor of 3 at 440 nm). The consequences of such a variability, when predicting or interpreting the diffuse reflectance of the ocean, are examined, according to whether or not these variations in $a_p$ are associated with concomitant variations in particle scattering. In most situations the deviations in $a_p$ actually are not compensated by those in particle scattering, so that the amplitude of reflectance is affected by these variations.

1. Introduction

Radiative transfer studies have established analytical links between the apparent optical properties (AOP) of the ocean [Preisendorfer, 1961] and its inherent optical properties (IOP), namely, the absorption and scattering properties of the water column [e.g., Gordon et al., 1975; Kirk, 1981; Mobley, 1994]. Therefore, when developing bio-optical models, such as for predicting the propagation of the radiant energy within the ocean or interpreting ocean color observations, there is a need to express the bulk IOPs or the contributions of the various substances to each of these coefficients as functions of their content in seawater. In oceanic “case 1” waters, the substances determining the IOPs, besides water itself, are (1) living phytoplankton; (2) associated “non-algal” particles, such as biog enous detritus and heterotrophic organisms; and (3) colored dissolved organic matter (CDOM) resulting from biological activity. Therefore analytical bio-optical models should ideally include parameterizations of IOPs for each of these components.

This has generally not been the case because of the variability in the individual spectral signatures of these various components as well as in the natural relationships existing between their concentrations. Prieur and Sathyendranath [1981] first proposed a parameterization of the total (particulate + dissolved) absorption coefficient at 440 nm, $a(440)$, that was derived from measurements of the diffuse attenuation coefficient $K_d$ (see notation section). Direct parameterizations of $K_d(\lambda)$, which is related to $a(\lambda)$, have also been proposed for the visible spectrum [Smith and Baker, 1978, Morel, 1988; Gordon et al., 1988]. These various studies have shown that either $a$ or $K_d$ vary nonlinearly with the chlorophyll a concentration in the water body (hereinafter denoted as (chl)). This nonlinearity was generally ascribed to the increasing proportion of non-algal particles with decreasing chl a concentration. Besides non-algal particles, $a$ and $K_d$ also include the contribution of CDOM, which has to be taken into account [Baker and Smith, 1982]. While this contribution is generally thought to be weak in the open ocean, its importance relative to that of other compartments is poorly known. Finally, the $K_d$ coefficient also depends on scattering, which may contribute to the nonlinear character of the relationship between $K_d$ and (chl) [Kirk, 1981; Gordon et al., 1988].

A further step in bio-optical models is to consider separately the particulate and dissolved compartments, where the particulate compartment includes both living phytoplankton and non-algal matter. Direct spectral measurements of particulate absorption $a_{ph}(\lambda)$ using the filter technique [Trüper and Yentsch, 1967] have been performed more and more frequently, allowing variations in these coefficients to be studied [Mitchell and Kieber, 1988; Garver et al., 1994; Cleveland, 1995]. Such measurements also allow the respective contributions of the living phytoplankton $a_{ph}$ and non-algal particles $a_{nonp}$ (with $a_p = a_{ph} + a_{nonp}$) to be estimated, using either experimental
Table 1. Cruises in Which Absorption Data Were Collected

<table>
<thead>
<tr>
<th>Cruise</th>
<th>Location</th>
<th>Period</th>
<th>Water Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>PACIPROD</td>
<td>Peru upwelling</td>
<td>Aug.–Sept. 1986</td>
<td>M</td>
</tr>
<tr>
<td>CHLOMAX</td>
<td>Sargasso Sea</td>
<td>Sept.–Oct. 1987</td>
<td>O</td>
</tr>
<tr>
<td>LIDAR 89</td>
<td>St. Lawrence estuary and gulf</td>
<td>Sept. 1989</td>
<td>E, M</td>
</tr>
<tr>
<td>LIDAR 90</td>
<td>St. Lawrence estuary and gulf</td>
<td>July 1990</td>
<td>E, M</td>
</tr>
<tr>
<td>TOMOFRONT</td>
<td>northwestern Mediterranean</td>
<td>April 1990</td>
<td>M</td>
</tr>
<tr>
<td>MEDIPROD 6 B</td>
<td>southwestern Mediterranean</td>
<td>July 1990</td>
<td>E, M</td>
</tr>
<tr>
<td>EUMELI 4</td>
<td>tropical North Atlantic</td>
<td>June 1992</td>
<td>E, M, O</td>
</tr>
<tr>
<td>FLUPAC</td>
<td>equatorial and subequatorial Pacific</td>
<td>Sept.–Oct. 1994</td>
<td>O, M</td>
</tr>
<tr>
<td>OLIPAC</td>
<td>equatorial and subequatorial Pacific</td>
<td>Nov. 1994</td>
<td>O, M</td>
</tr>
<tr>
<td>MINOS</td>
<td>eastern and western Mediterranean</td>
<td>May 1996</td>
<td>O, M</td>
</tr>
</tbody>
</table>

M, O, and E denote mesotrophic, oligotrophic, and eutrophic, respectively.

[e.g., Kishino et al., 1985] or numerical [e.g., Morrow et al., 1989; Bricaud and Stramski, 1990] techniques. Systematic studies [Bricaud et al., 1995; Cleveland, 1995] have shown that \( a_{\phi}(\lambda) \) varies in a nonlinear manner with \( \langle \text{chl} \rangle \); as a result of changes in pigment composition and in the package effect with the trophic state of waters. Furthermore, the exponent in the power law that links \( a_{\phi} \) to \( \langle \text{chl} \rangle \) (e.g., 0.67 at the blue maximum [Bricaud et al., 1995]) is very close to that previously found for \( K_d \) (0.71 [Morel, 1988]); note that \( K_d \) depends on \( a_{\phi} \) rather than solely on \( a_{\phi} \). The similarity of these exponents suggests that the relative contribution of nonalgal particles to absorption does not vary markedly with \( \langle \text{chl} \rangle \) and is not the source of the nonlinear variations in total absorption, as previously assumed [e.g., Gordon et al., 1988].

It is therefore timely to examine how the particulate absorption coefficients vary with \( \langle \text{chl} \rangle \) and to what extent the nonalgal particulate matter contributes to total particulate absorption. The consistency of the variations in \( a_{\phi}(\lambda) \) and \( K_d(\lambda) \) with \( \langle \text{chl} \rangle \) will also be examined. Finally, particular attention will be paid to the natural variability of particulate absorption, with its implications for bio-optical models, particularly those aimed at predicting the diffuse reflectance of the ocean.

2. Data and Methods

The present data set includes 783 samples previously used for studying the variations in phytoplanktonic absorption [Bricaud et al., 1995] and collected during eight cruises in various areas of the world ocean. From the original data set, 32 samples have been removed as containing nonbiological detrital matter, so that only case 1 waters are included in this data set. In addition, the present data set also includes 383 new samples, collected during the following three recent cruises: (1) the FLUPAC cruise in September–October 1994 (Joint Global Ocean Flux Study (JGOFS) France program) [Le Borgne and Gesbert, 1995], where measurements were performed in the oligotrophic and mesotrophic waters of the equatorial Pacific, along a meridian transect at 165°E (from 15°S to 6°N) and a transect along the equator between 167°E and 150°W (84 spectra); (2) the OLIPAC cruise in November 1994 (JGOFS France program [Mounin and Coste, 1996]), also in the equatorial Pacific, along a transect at 150°W between 16°S and 1°N (184 spectra); and (3) the MINOS cruise in May 1996 in waters ranging from ultraoligotrophic waters (eastern Mediterranean Sea) to mesotrophic waters in the Western Basin (115 spectra). Information concerning the various cruises is summarized in Table 1. Only samples collected in the 0 to 200-m layer have been considered in the present data set.

Methods are described in detail by Bricaud et al. [1995]. Briefly, particulate absorption coefficients \( a_{\phi}(\lambda) \) were determined using the glass fiber filter (GF/F) technique, except for the FLUPAC cruise, where the “glass-slide technique” [Allali et al., 1995] was used. All \( a_{\phi} \) spectra were set to 0 at 750 nm. Although this may be not strictly true for detrital matter, this assumption is believed to minimize possible differences between sample and reference filters. Then the spectra measured using the GF/F technique were corrected for the path length amplification effect, using the algorithm given by Bricaud and Stramski [1990] for the previous data set and that given by Allali et al. [1997] for the three additional cruises. This latter algorithm, derived from measurements in the equatorial Pacific and providing slightly lower values than the Bricaud and Stramski algorithm (by 15% for an optical density of 0.3), is thought to be more appropriate for oligotrophic waters (see discussion by Allali et al. [1997]). Finally, the contributions of phytoplankton \( a_{\phi}(\lambda) \) and nonalgal matter \( a_{\text{non}}(\lambda) \) to total particulate absorption were determined either experimentally [Kishino et al., 1985] or by numerical decomposition [Bricaud and Stramski, 1990; see Bricaud et al., 1995, Table 1]. For the three recent cruises, numerical decomposition was always used (for a few samples the method of Kishino et al. was also applied for verification). It is recalled here that Kishino et al.’s method does not provide \( a_{\phi}(\lambda) \) and \( a_{\text{non}}(\lambda) \) sensu stricto but, rather, the absorption coefficients of pigmented matter (at least that extractable in methanol) and nonpigmented matter.

Figurant concentrations were measured using various techniques (high pressure liquid chromatography (HPLC), spectrophotometry, fluorometry, spectrophotometry). The HPLC method was considered as the reference, and for the previous data set some corrections were made to account for systematic divergences between HPLC and other methods [see Bricaud et al., 1995]. For the three recent cruises, measurements were always made using the HPLC technique, according to the procedure described by Vidussi et al. [1996]. Absorption coefficients \( a_{\phi}, a_{\text{non}}, a_{\text{non}} \) were converted into chi-specific coefficients \( a_{\phi}, a_{\phi}, a_{\text{non}} \) by normalizing to the chi concentration (including divinyl-chl a and pheophytin a, if any).

3. Results and Discussion

3.1. Relationship Between Particulate Absorption (at 440 nm) and Chlorophyll a Concentration

The particulate absorption coefficient \( a_{\phi}(440) \) exhibits a clear trend of increasing nonlinearly with \( \langle \text{chl} \rangle \) (see log-log plot in Figure 1a), as observed previously for the phytoplankton.
absorption coefficient [Bricaud et al., 1995]. As a first approximation, power functions are appropriate to represent these variations. For the wavelength 440 nm, where \(a_p(\lambda)\) and \(a_d(\lambda)\) are near their maximum (the spectral values will be considered later), least squares fits provide the following relationships:

\[
a_p(440) = 0.0520 \langle \text{chl} \rangle^{0.615}, \quad r^2 = 0.91, \quad N = 1166 \quad (1)
\]

\[
a_d(440) = 0.0378 \langle \text{chl} \rangle^{0.627}, \quad r^2 = 0.90, \quad N = 1166 \quad (2)
\]

where \(r^2\) is the determination coefficient and \(N\) is the number of samples. Note that (2) slightly differs from the expression previously given by Bricaud et al. [1995] as it is derived from a larger data set (1166 versus 815 samples).

The nonlinear character of (1) appears very similar to that previously found by Morel [1988] for the non-water diffuse attenuation coefficient \((K_d - K_w)\), hereinafter referred to as \(K_{\text{bio}}\):

\[
K_{\text{bio}}(440) = 0.1041 \langle \text{chl} \rangle^{0.707}, \quad r^2 = 0.94
\]

although the \(K_{\text{bio}}\) values are, on average, twice as high as the \(a_p\) values, as a consequence of the contribution of absorption by CDOM and of the effect of scattering.

Equations (1) and (2) can be straightforwardly transformed into

\[
a_p^*(440) = 0.0520 \langle \text{chl} \rangle^{-0.365} \quad (1')
\]

\[
a_d^*(440) = 0.0378 \langle \text{chl} \rangle^{-0.373} \quad (2')
\]

where the starred quantities represent the chl-specific coefficients \((\text{m}^2\text{mg}^{-1})\). Nonlinear relationships similar to (1') were previously found by Tentsch and Phinney [1989] at 440 nm and by Garver et al. [1994] for particulate absorption averaged over the visible spectrum (but only for \(\langle \text{chl} \rangle\) values lower than 1 mg m\(^{-3}\)).

The exponents in (1) and (2) are remarkably close, and the variations of \(a_p(440)\) and \(a_d(440)\) versus \(\text{chl}\) are nearly parallel (Figure 1), which suggests that at this wavelength, absorption by nonalgal particles is roughly proportional to total absorption. This is also suggested by the relationship directly established between \(a_{\text{nap}}(440)\) and \(\text{chl}\):

\[
a_{\text{nap}}(440) = 0.0124 \langle \text{chl} \rangle^{0.724}, \quad r^2 = 0.73 \quad (3)
\]

in which the exponent is not appreciably different from those in (1) and (2) (note, however, the much lower determination coefficient for \(a_{\text{nap}}\)). According to (1) and (2), \(a_p(440)\) is higher than \(a_d(440)\) by, on average, a factor 1.37 over the entire \(\langle \text{chl} \rangle\) range (from 1.33 when \(\langle \text{chl} \rangle = 0.02\) mg m\(^{-3}\) to 1.41 when \(\langle \text{chl} \rangle = 25\) mg m\(^{-3}\)). In other words, living phytoplankton and nonalgal particles would contribute, on average, 73% and 27%, respectively, to total particulate absorption at 440 nm.

It is worth noting that the scatter for \(a_p\) (actually high throughout the whole \(\langle \text{chl} \rangle\) range) is not larger than that for \(a_d\), as the determination coefficients on the log-transformed data are almost identical (0.90). Residuals are nearly normally distributed around the regression lines, which allows confidence intervals to be computed for delimiting the natural variability in these coefficients. The dashed lines in Figure 1a delimit the prediction intervals at a 90% confidence level and can be considered as realistic upper and lower limits. According to these prediction intervals, the \(a_p(440)\) values may vary by a factor of 2.9 for a given \(\langle \text{chl} \rangle\) value. Oligotrophic waters (with \(\langle \text{chl} \rangle < 0.1\) mg m\(^{-3}\)) actually encompass this range, while eutrophic waters span a more restricted range. This might be the result of geographical diversity, which was larger for oligotrophic waters than for eutrophic waters in our data set. For mesotrophic waters, some groups of points stand apart, particularly from the EUMELLI 4 cruise (above the upper limit).

Figure 1. (a) Variations of the absorption coefficients of particles at 440 nm, \(a_p(440)\), as a function of \(\langle \text{chl} \rangle\), for various regions of the world ocean (see Table 1). All depths have been pooled together. The regression line (equation (1)) is shown as a solid line; the dashed lines delimit the prediction interval at a 90% confidence level. Here \(\langle \text{chl} \rangle\) represents the \((\text{chl} a + \text{DIF-chl} a + \text{pheo} a)\) concentration. (b) Same as Figure 1a, but for the absorption coefficient of living phytoplankton \(a_p(440)\). The equation of the regression line is (2). (c) Same as Figure 1a, but for the samples located within the first optical depth \(z = z_0/4.6\), where \(z_0\) is euphotic depth). The equation of the regression line is (4) (solid line). The regression line and prediction interval corresponding to the whole data set (as in Figure 1a) are also shown (dotted and dashed lines) for comparison.
and data from the PACIPROD and CHILOMAX cruises (below the lower limit).

A similar range of variation (a factor of 2.95) is observed for \( a_p(440) \) (Figure 1b). Groups of points outside these limits are also observed for the EUMELI 4 and CHLOMAX cruises, which means that for these particular samples, anomalously high or low \( a_p \) values can be attributed to algae (by virtue, for instance, of a particular pigment composition) and not to non-algal particles. Samples from the mesotrophic site during the EUMELI 4 cruise were characterized by an exceptionally high concentration of cyanobacteria [Morel, 1997] and therefore by a high zeaxanthin content relative to (chl). In contrast, most \( a_p(410) \) data from the PACIPROD cruise are low but not “anomalous,” so that the very low \( a_p \) values appear to result from a weak contribution of nonalgal particles to total absorption.

When computing the prediction intervals throughout the visible spectrum (not shown), it is observed that for a given (chl) value, \( a_p(\lambda) \) varies by a factor of 2.8 to 4 according to wavelength, except in the green part of the spectrum where it becomes even higher (likely because of the increasing experimental uncertainty affecting \( a_p \) in this spectral domain). Such relatively high scatter around the mean \( a_p(\lambda) \) versus (chl) relationships may originate from geographical or seasonal differences. From a restricted data set, Carder et al. [1991] discerned different tendencies between temperate and subtropical waters. In contrast, Cleveland [1995] did not detect distinct patterns for these regions but only for subpolar (Arctic) waters, in agreement with previous findings of Mitchell and Holm-Hansen [1991] for Antarctic waters. For the South Atlantic Bight shelf waters, Nelson and Guarda [1995] observed season-dependent relationships between \( a_p(443) \) and (chl).

As the present data set is built up from different cruises in various locations and periods, discriminating regional and seasonal trends, if any, is hazardous. In addition, the various cruises cover very distinct ranges of (chl), some of which are rather restricted, which hampers robust cruise-by-cruise relationships to be established. Therefore our purpose here is not to examine the existence of such regional/seasonal relationships. Rather, the aim is to extract general trends for a variety of case 1 waters, to assess the natural variability around these trends, and to evaluate the impact of this variability upon bio-optical modeling.

### 3.2. Variations of Particulate Absorption in the Near-Surface Layer

With reference to the particular problem of modeling the diffuse reflectance \( R(\lambda) \), the analysis of the \( a_p \) versus (chl) variations has to be restricted to the layer accessible to remote sensing observations, i.e., the layer corresponding to the first optical depth [Gordon and McCluney, 1975]. This depth can be simply computed as \( z_c/4.6 \), where \( z_c \), the euphotic depth, is either determined from the measured downwelling photosynthetically available radiation (PAR) profile when available or estimated from the vertical profile of (chl) [Morel, 1988]. When the data set is restricted to this near-surface layer (260 samples out of 1166), the variations of \( a_p(440) \) versus (chl) can be described by the relationship

\[
a_p(440) = 0.0519 \times \text{chl}^{0.590}, \quad r^2 = 0.92
\]

which does not significantly differ from (1) (Figure 1c). It can also be observed that with the same exceptions as in Figure 1a (samples from EUMELI 4 and some PACIPROD and CHLOMAX samples), the lines corresponding to the prediction interval at the 90% confidence level for the whole data set still represent realistic upper and lower limits for the near-surface layer.

Larger differences between the \( a_p \) values within and below the near-surface layer might be expected within the spectral bands of some pigments such as the nonphotosynthetic carotenoids (e.g., zeaxanthin, diatoxanthin, diadinoxanthin), as it is well known that these pigments are more abundant (relative to (chl)) within the surface well-lit layer than in deeper waters. As these pigments have a broad absorption band centered on the domain 430–500 nm [e.g., Bidigare et al., 1990], this hypothesis was tested by comparing the \( a_p(490) \) versus (chl) relationships for the near-surface layer and for the deeper layer. Somewhat unexpectedly, they were found to be rather close (with \( a_p(490) = 0.0330 \times \text{chl}^{0.2594} \) for the near-surface layer and \( a_p(490) = 0.0345 \times \text{chl}^{0.651} \) for the deeper layer). Similar results are obtained when considering \( a_p(490) \) instead of \( a_p(440) \). This suggests that the nonphotosynthetic pigments influence only weakly the \( a_p \) (and \( a_p \)) versus (chl) relationships near the surface, probably because the ratio of total (photosynthetic + nonphotosynthetic) carotenoids to (chl) does not vary greatly throughout the euphotic zone (see, e.g., Allali et al. [1997] for the OIIPAC cruise).

### 3.3. Contribution of Nonalgal Particles to Total Particulate Absorption

As suggested by the similarity of the exponents in (1) and (2), the ratio of nonalgal absorption to total particulate absorption at 440 nm, \( a_{n\text{ap}}(440)/a_p(440) \), does not reveal a regular trend as a function of (chl) (Figure 2a). Waters from the equatorial and subequatorial Pacific (FLUPAC and OLIPAC cruises) exhibit particularly low \( a_{n\text{ap}}(440)/a_p(440) \) values, mostly between 0.01 and 0.15. In contrast, oligotrophic Mediterranean waters (MINOS cruise) generally show high values for this ratio (up to 0.8). Except for these cases, most of the points lie between 0.15 and 0.60, with a cluster around 0.25–0.30, whatever the (chl) range, as previously inferred from the comparison between (1) and (2). It could be argued that this rather large variability originates from a varying contribution of nonalgal particles throughout the water column. When restricting the analysis to the first optical depth, however, the range observed above still remains unchanged (Figure 2b).

This range in \( a_{n\text{ap}}(440)/a_p(440) \) is consistent with that observed for temperate, subtropical, and subpolar regions by Cleveland [1995] (approximately 0.2–0.8). This study already emphasized that the proportion of absorption by nonalgal particles exhibited no simple trend as a function of (chl). Note that this author did not observe a particularly low content in nonalgal particles for waters from the equatorial Pacific. In contrast, the \( a_p \) spectra that we measured in these waters were close in shape to those of “pure cultures,” which indicates that waters were dominated by living phytoplankton [Allali et al., 1997]. Since for these cruises, \( a_{n\text{ap}}(\lambda) \) was obtained from \( a_p(\lambda) \) by numerical decomposition, an artifact could be suspected. During the OLIPAC cruise, however, \( a_{n\text{ap}}(\lambda) \) was also determined experimentally for several samples, so that it could be confirmed that the low \( a_{n\text{ap}} \) values were not an artifact of the decomposition. It appears therefore that no simple relationship between the contributions of nonalgal particles and phytoplankton to absorption can presently be proposed, which is not surprising, considering the variety and complexity of the
3.4. Spectral Dependence of Absorption by Nonalgal Particles

Absorption by nonalgal particles has long been recognized to vary approximately according to an exponential function of the form [e.g., Yentsch, 1962; Kirk, 1980; Roessler et al., 1989]:

\[ a_{\text{nap}}(\lambda) - a_{\text{nap}}(\lambda_0) \exp \left(-S(\lambda - \lambda_0)\right) \]  \hspace{1cm} (5)

This expression is found adequate for the present measurements (Figure 3). For the cruises during which the method of Kishino et al. [1985] was applied, the values of the slope \( S \) can be determined by fitting \( a_{\text{nap}}(\lambda) \) to (5). When applying this procedure to the experimental \( a_{\text{nap}}(\lambda) \) values in the spectral domain 380–700 nm, the coefficient of determination \( r^2 \) is higher than 0.90 in most cases (262 out of 267 measured \( a_{\text{nap}}(\lambda) \) spectra). The \( S \) values obtained by numerical decomposition are not considered here, because it was shown previously that, in contrast to the retrieved \( a_\phi(\lambda) \) and \( a_{\text{nap}}(\lambda) \) values, the retrieved \( S \) values are sensitive to the assumptions used for the decomposition (see discussion and Figure 9 of Bricaud and Stramski [1990]).

The \( S \) values estimated from measured \( a_{\text{nap}} \) spectra range between 0.008 and 0.016 nm\(^{-1}\), with an average value of 0.011 nm\(^{-1}\) (standard deviation 0.0025, \( N = 267 \)). Average values of \( S \) are 0.012 (±0.002), 0.011 (±0.002), and 0.010 (±0.001) nm\(^{-1}\) for each of the three (chl) decades (0.01–0.1, 0.1–1, 1–10 mg m\(^{-3}\)), respectively. These values are in good agreement with those reported by Roessler et al. [1989] from their measurements off the San Juan Islands (40 spectra), as well as with those they compiled from previously published works (\( S = 0.011 + 0.002 \) nm\(^{-1}\) in both cases). The variations in this spectral dependence also show some similarities with CDOM, although the slope values appear to be, on average, lower for nonalgal particles. Carder et al. [1989] observed \( S \) values for CDOM varying from 0.011 to 0.017 nm\(^{-1}\), with the high values in oligotrophic surface waters, related to a larger fraction of fulvic acids, and the low values in areas of senescent blooms [see also Green and Blough, 1994].

3.5. Parameterization of the Chl-specific Particulate Absorption Spectra

This parameterization consists of fitting the relationship between \( a_\phi(\lambda) \) and \( \langle \text{chl} \rangle \) at each wavelength to a power function

\[ a_\phi(\lambda) = A_\phi(\lambda)^{\langle \text{chl} \rangle E_\phi(\lambda)} \]  \hspace{1cm} (6)

which can be transformed into

\[ A_\phi(\lambda)^{\langle \text{chl} \rangle E_\phi(\lambda)} = A_\phi(\lambda)^{\langle \text{chl} \rangle E_\phi(\lambda)}^{-1} \]  \hspace{1cm} (6')

The values of \( A_\phi(\lambda), E_\phi(\lambda) \), and the coefficients of determination on the log-transformed data \( r^2 \) are shown in Figure 4, as compared to those \( (A_\phi, E_\phi) \) obtained for \( a_\phi \). The numerical values of these various coefficients (at 2-nm intervals) can be provided on request. Note that \( A_\phi \) and \( E_\phi \) correspond to \( A \) and \( (1 - B) \) of Bricaud et al. [1995, equation (1)]; the numerical values are actually slightly different, as they have been derived from an enlarged data set.

As noted for the 440-nm wavelength, the values of the exponents \( E_\phi(\lambda) \) and \( E_\phi(\lambda) \) are very close throughout the 400- to 520-nm domain, indicating again that the contribution of nonalgal particles to total particulate absorption shows no particular trend with \( \langle \text{chl} \rangle \). As shown by the coefficients of determination (Figure 4, bottom), the scatter of the \( a_\phi \) versus \( \langle \text{chl} \rangle \) relationship is very similar to that of the \( a_\phi \) versus \( \langle \text{chl} \rangle \) relationship throughout this wavelength domain. The \( E_\phi \) and \( E_\phi \) values are also in good agreement from 640 to 700 nm. In the green domain, mainly between 520 and 600 nm, they diverge, and the \( E_\phi \) values become lower, indicating a stronger nonlinearity.

The difference between the \( A_\phi \) and \( A_\phi \) spectra, very slight

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{figure2.png}
\caption{(a) Variations of the ratio of nonalgal absorption to total particle absorption at 440 nm \( (a_{\text{nap}}/a_\phi)(440) \) as a function of \( \langle \text{chl} \rangle \). Symbols are the same as in Figure 1. (b) Same as Figure 2a, but for the samples located within the first optical depth.}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{figure3.png}
\caption{Spectra of the chl-specific absorption coefficient of nonalgal particles, \( a_{\text{nap}}(\lambda) \), obtained using the method of Kishino et al. [1985]. All spectra are shown to illustrate the range of variation of \( a_{\text{nap}} \).}
\end{figure}
Figure 4. Spectral values of the numerical coefficients (top) $A_p$ and (middle) $E_p$ in the power function representing the variations of $a_p(\lambda)$ as a function of $\langle \text{chl} \rangle$ (equation (6)) and (bottom) coefficients of determination on the log-transformed data, $r^2$. The corresponding values for the absorption coefficients of living phytoplankton $a_d(\lambda)$ are shown as dashed lines for comparison.

within the red absorption band, increases toward the blue domain and expresses, as expected, the increasing contribution of nonalgal particles to total absorption with decreasing wavelength. As the exponents $E_p(\lambda)$ and $E_d(\lambda)$ are very close in the blue and red domains, the ratio $A_d(\lambda)/A_p(\lambda)$ can be directly used to evaluate the contribution of phytoplankton to particulate absorption at various wavelengths. For the present data set the contribution of algal (found earlier to be 73%, on average, at 440 nm) turns out to be 88% at the red absorption maximum (675 nm) and drops to 55% at 400 nm, where the contribution of nonalgal particles is the highest (Figure 4, top).

Figure 5 shows the mean $a_p^*(\lambda)$ spectra computed for various $\langle \text{chl} \rangle$ values (solid lines) and compared to $a_d^*(\lambda)$ spectra (dashed lines), obtained by using the above parameterizations. The higher relative contribution of nonalgal particles at shorter wavelengths is revealed by the slight deformation of the spectral shape of $a_p^*$ compared to $a_d^*$. It is remarkable, however, that even for very low $\langle \text{chl} \rangle$ values, the shape of the $a_p^*$ spectra remains strongly influenced by phytoplanktonic pigments. This feature may be overemphasized by our data set because of the numerical dominance of Pacific oligotrophic waters, where the contribution of nonalgal particles seems to be exceptionally weak. However, it is believed to be real and consistent with the fact that in most other areas (with the exception of the MINOS cruise), the relative contribution of nonalgal particles to absorption is not higher in oligotrophic waters than in eutrophic waters (Figure 2). Note also the progressive appearance of an absorption shoulder around 390–600 nm when waters become more oligotrophic (decreasing $\langle \text{chl} \rangle$); this feature originates from the increasing occurrence of divinyl-chl $b$ present in prochlorophytes.

3.6. Comparison With the Parameterization of the Diffuse Attenuation Coefficient

It is worthwhile examining whether the present parameterization of particulate absorption as a function of $\langle \text{chl} \rangle$ is consistent with that established for the diffuse attenuation coeffi-
cient for downwelling irradiance $K_d(\lambda)$ by Morel [1988]. This former parameterization,

$$K_d(\lambda) = K_w(\lambda) + \chi(\lambda) (chl)^{a_d(\lambda)}$$ (7)

where $K_w$ represents the diffuse attenuation coefficient of pure seawater, was based on a statistical analysis of $K_d(\lambda)$ values derived from downwelling irradiance spectra. Therefore the experimental methods used for providing $K_d(\lambda)$ and $a_d(\lambda)$ are completely different. The data sets are also almost independent, from a geographical viewpoint, as only the PACIFRIP cruise is included in both data sets (21 out of the 178 $K_d$ spectra, 30 out of the 1166 $a_d$ spectra). On the basis of the same approach, a new statistical analysis that includes also the $K_d$ measurements acquired during the cruises EUMELI 3, EUMELI 4, OLIPAC and MINOS (207 additional spectra) was recently performed (A. Morel et al., manuscript in preparation, 1998), and led to a new set of coefficients for (7). These two sets (actually rather close) are used simultaneously below.

To examine whether the parameterizations of $a_d$ and $K_d$ are consistent, we have computed $K_d(\lambda)$, using an empirical formula derived by Kirk [1981]. This computation is described in the appendix. The $K_d$ values computed in this way can then be compared with those parameterized by Morel [1988] and A. Morel et al. (manuscript in preparation, 1998). Such a comparison was already attempted by Morel [1988]; however, for lack of available data, the chl-specific absorption coefficients of phytoplankton were assumed to be independent of (chl), and the contributions of nonnalgal particles and CDOM were ignored. Here the same approach can be used again in a more realistic way.

Since for very low (chl), the $K_d$ values are governed almost exclusively by pure seawater, a more meaningful comparison is performed by examining the differences $(K_d - K_w)(\lambda) = K_{bio}(\lambda)$. The $K_{bio}$ spectra provided by both approaches have been compared for three (chl) values (Figure 6). For low and moderate (chl) values (0.1 and 1 mg m$^{-3}$) the spectral shape is reasonably well reproduced throughout the spectrum, except below 440 nm, and in the 560- to 620-nm domain. It is recalled that the spectral values of $a_d$ have to be considered with caution in the green part of the spectrum because of the low coefficients of determination (see Figure 4). For high (chl) values (10 mg m$^{-3}$) the reconstructed values below 500 nm become significantly lower than those provided by (7); also, the blue maximum tends to disappear (especially in the presence of CDOM), while it remains visible on the spectrum provided by (7). The values in the red part of the spectrum (around 670 nm) are in good agreement, not only for low (chl) values, where water absorption predominates, but also for (chl) values as high as 10 mg m$^{-3}$.

In spite of the spectral divergences noted above, it is important to note that, over the whole (chl) range, the $K_{bio}$ values experience a variation by a factor of about 20 and that these variations in amplitude are correctly reproduced by (10–13). This agreement between the absolute values is comforting; the present uncertainty concerning the actual contribution of CDOM to total absorption prevents further progress in the comparison. It can be concluded, however, that the parameterizations of $K_d(\lambda)$ and $a_d(\lambda)$ are basically consistent for a (chl) range covering oligotrophic to eutrophic waters.

3.7. Implications for Diffuse Reflectance Modeling

3.7.1. Impact of the variations in absorption. Models aimed at predicting ocean color from water content generally rely upon unique relationships between absorption (or diffuse attenuation), scattering, and (chl). It is therefore necessary to examine how the natural variability in $a_d$ may affect the diffuse reflectance of the ocean. With this purpose a simple reflectance model was operated, using the relationship

$$R(\lambda) = f(0.5b_w(\lambda) + [b_{bp}(\lambda) b_p(\lambda)]a(\lambda))$$ (8)

where $(b_{bp}/b_p)$ represents the backscattering efficiency of particles; $f$ is the proportionality factor between $R$ and the backscattering-to-absorption ratio and varies with the (chl) content, the Sun elevation, and wavelength [Morel and Gentili, 1996]. The model was operated here with the Sun at zenith. The $a(\lambda)$ coefficients were computed using (11) and (12) (see appendix); $y$ in (12) was given the value 0.2, and the $a_d(\lambda)$ coefficients

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**Figure 6.** Comparison of the nonwater diffuse attenuation spectra ($K_{bio}$ in m$^{-1}$) obtained by using (7) with two sets of numerical coefficients (solid lines) and (10–13) (dashed and dotted lines), for three values of the chlorophyll concentration. The $y$ values indicated are those used in (12). The $a_d$ spectra provided by (b) for the same chlorophyll concentrations are also shown for comparison.
were parameterized as in (6). The numerical coefficients $A_p(\lambda)$ and $E_p(\lambda)$ computed for the whole euphotic zone were preferred to those obtained for only the near-surface layer (they are not statistically different and are thought to be more robust because they are derived from a fourfold larger data set). The particle scattering coefficient $b_p(\lambda)$ was computed by applying (13) and (13') and subtracting the molecular scattering coefficients $b_m(\lambda)$. Finally, the particle backscattering efficiency, $(b_p/b_m)$, was made varying with (chl) as in the paper by Morel [1988].

In order to assess the impact of the residual variability in $a_p(\lambda)$ (with respect to the mean $a_p(\lambda)$ versus (chl) relationships), the reflectance spectra were also computed using the upper and lower limits of these coefficients (defined as previously from the prediction intervals at a 90% confidence level). This computation was performed for two values of (chl), 0.1 and 10 mg m$^{-3}$ (Figure 7a). As expected, the impact of this variability is rather limited at low (chl), where molecular absorption predominates. It becomes much higher at high (chl), about a factor of 3 over the blue-green domain, for 10 mg m$^{-3}$. In the red part of the spectrum this effect is again partially masked by the increasing contribution of water absorption.

For the sake of comparison, the effect of the possible changes in the CDOM concentration has been examined (Figure 7b). The absorption coefficient at 440 nm, $a_{ps}(440)$, was varied as previously, from 0 to 20% of the $(a_p + a_{ps}(440))$ coefficient (equation (12)). At low (chl) (0.1 mg m$^{-3}$), the changes in CDOM absorption and variability in particulate absorption appear to have a similar impact upon reflectance (limited essentially to the blue wavelengths). At high (chl) (10 mg m$^{-3}$) the effect of changing CDOM absorption becomes negligible compared to the effect of variability in particulate absorption.

**3.7.2. Impact of the combined variations in absorption and scattering.** The above computations assume that particle scattering at 660 nm follows perfectly (13'). In reality, the $b_p(660)$ values also show natural variability around this mean relationship [see Loiselle and Morel, 1998], comparable to that observed for particulate absorption. Therefore it is of interest to know whether deviations of particulate absorption (with respect to the $a_p$ value provided by the parameterization) are associated or not to deviations in particle scattering or, in other words, whether variations in $a_p$ and $b_p$ might counterbalance or conversely cumulate their effects upon reflectance.

This was examined for the cruises where simultaneous measurements of $a_p(\lambda)$ and the attenuation coefficient at 660 nm, $c(660)$ (measured in situ with a Sea Tech transmissimeter [Bartz et al., 1978]) were available. The $c(660)$ values were corrected for pure seawater attenuation by subtracting the deep local value [see Loiselle and Morel, 1998] and then converted into $b_p(660)$ by subtracting the measured $a_p(660)$ value (in general, a few percent of the $c(660)$ value). For each station and depth the relative deviations (in percent) of $b_p(660)$ and $a_p(440)$ from their average value were computed as:

$$\delta b_p(660) = 100\left[\frac{b_p(660) - b_{pm}(660)}{b_{pm}(660)}\right]$$

$$\delta a_p(440) = 100\left[\frac{a_p(440) - a_{pm}(440)}{a_{pm}(440)}\right]$$

where $b_{pm}(660)$ and $a_{pm}(440)$ were obtained using (13') and (1), respectively (note that $\delta b_p(660)$ values close to -100% correspond to samples at the bottom of the euphotic zone, where $b_p(660)$ approaches 0).

Figure 8, divided in four panels, demonstrates that several situations occur. (1) In Panels I and II, where $\delta b_p(660)$ and $\delta a_p(440)$ are of identical sign, low and high $a_p$ values are associated with low and high $b_p$ values, respectively. Therefore their variations should partially compensate for each other (totally for those samples lying on the 1:1 line in Figure 8) and should have a limited impact upon diffuse reflectance (provided that there is no change in the backscattering efficiency for these samples). This is the case for most samples of the OLIPAC cruise, as well as some from EUMELI 3, EUMELI-4, and MINOS. For most samples from the OLIPAC cruise the "low $b_p$—low $a_p$" values (with respect to the mean values) are easily explained by their exceptionally low content in nonalgal particles. On the opposite side, the "high $b_p$—high $a_p$" values observed during the MINOS cruise (corresponding mainly to surface samples) indicate a high detrital content comparatively to (chl). (2) In Panels III and IV, in contrast, the deviations in $a_p$ and $b_p$ are opposite (low and high $a_p$ values are associated to high and low $b_p$ values, respectively). In such situations the effects of variations in $a_p$ and $b_p$ upon diffuse reflectance are cumulative and should induce reflectance variations still larger than those predicted in Figure 7a. This is the case for some EUMELI 3 and OLIPAC samples and for most MINOS and FIMFI 1 samples. For those samples the high $a_p(440)$ values mainly result from the pigment composition, as also demonstrated by the high $a_{ps}(440)$ values (not shown; see
Rabin et al. [1996] for FLMFI I 4. This particular pigment composition (abundance of zeaxanthin) has no direct effect on $b_p(660)$, but it may indicate the presence of phytoplanktonic populations of small size, such as prochlorophytes, which are characterized by very low chl-specific scattering coefficients [Morel et al., 1993].

With regard to the diversity of these situations, the variability shown in Figure 7a may probably be considered as realistic, to the extent that variations in $a_p$ are very seldom compensated by those in $b_p$. These results emphasize the difficulty of relying upon the absolute values of reflectance, when, for instance, attempting to retrieve the chlorophyll content from satellite data via a semianalytical reflectance model. Even when ignoring other sources of uncertainty in reflectance models, such as the spectral dependence of particle scattering, the variations of the backscattering efficiency with the (chl) content, and the contribution of CDOM to total absorption, the unexplained variability in $a_p(\lambda)$ and $b_p(\lambda)$ may be responsible for large variations in the amplitude of reflectance, particularly for moderate or high chlorophyll contents.

Reflectance values, however, do not vary in an erratic way from one wavelength to the other, as demonstrated, for instance, by the positive correlation between $\delta a_p(440)$ and $\delta a_p(550)$ (Figure 9, panels I and II). This emphasizes the interest of using reflectance ratios instead of absolute values of reflectance when retrieving the chlorophyll concentration from satellite data. In such algorithms the impact of the variability in $a_p$, upon the chlorophyll retrieval is significantly reduced (actually cancelled for the samples on the 1:1 line in Figure 9). It is observed in Figure 9, however, that for a nonnegligible number of samples (panels III and IV), some low/high values of $a_p(440)$ coexist with high/low values of $a_p(550)$. In such cases the impact of the variations in particle absorption is obviously not reduced when using reflectance ratios.

4. Conclusions

The former analysis of the natural variations in absorption by phytoplankton [Briceaud et al., 1995] was performed mainly in view of modeling primary production, and its results were accounted for in a revised version of the photophysiological parameters involved in such a modeling [Morel et al., 1996]. The present analysis dealing with total particulate absorption is mainly aimed at providing an input parameter for modeling apparent optical properties, such as diffuse attenuation or reflectance. It also assesses the respective contributions of algal and nonalgal particles to particulate absorption in waters with various trophic regimes.

Both parameterizations (for $a_p$ and $a_{ph}$) should be used only when bearing in mind their limitations: (1) they are applicable to case 1 waters only and not valid outside the (chl) range corresponding to the present data set; (2) because of the large possible individual deviations from the average relationships, they can in no way be operated on a sample-by-sample basis; and (3) in spite of the rather large data set, these parameterizations are not intended to be universal, in particular, because some peculiar situations (e.g., polar waters) have not been explored, and the data set was not appropriate to detect regional or seasonal trends, if any.

The present analysis demonstrates that the $a_p(\lambda)$ coefficients can vary by more than 1 order of magnitude when the chl a concentration varies by 3 orders of magnitude (0.02 to 25 mg m$^{-3}$). Such variations definitely have to be taken into account when developing analytical (or semianalytical) bio-optical models. The actual impact of these variations upon diffuse reflectance values, for instance, is predictable; however, two other input parameters in reflectance models, absorption by CDOM and particle backscattering efficiency, still remain very poorly documented. While the G/F technique for particulate absorption measurements has been widely used during the last decade, so that this parameter is now rather well known, absorption by CDOM and particle backscattering was very seldom determined at sea because of instrumental limitations. The development and increasing use of new in situ optical devices such as spectral absorption meters [Bruce et al., 1996] equipped with filters and backscattering sensors [Maffione and Dona, 1997] should improve the situation.

As a first approximation, the nonlinear $a_p$ versus (chl) and $a_{ph}$ versus (chl) relationships parallel each other, indicating that the contribution of nonalgal particles to total absorption oscil-
lates around an average value (~25-30%), whatever the chlorophyll content in case 1 waters. This does not contradict the findings of previous studies showing that the carbon-to-chl ratio increases with decreasing chl [Hobson et al., 1973; Losel and Morel, 1998], but this indicates that the increase in this ratio is not necessarily reflected by a relative increase in non-algal absorption.

The contribution of non-algal absorption to total absorption actually exhibits large variations around its average value and also shows exceptionally low values in some areas (e.g., in the equatorial Pacific); therefore modeling optical properties for the non-algal compartment remains uncertain. As the scatter observed on the $a_p(\lambda)$ versus (chl) relationships is not higher than that for $a_o$, it appears that, at least as a first step, absorption by particles can be represented globally in bio-optical models (i.e., without increasing the level of complexity by modeling separately the absorption coefficients of the algal and non-algal compartments).

The residual variability in particulate absorption (as well as in phytoplanktonic absorption), with respect to the average relationships, is rather large and probably conceals specific and valuable information. Further work is needed to analyze this information. In our present state of knowledge this variability represents a limitation for reflectance prediction, especially because deviations in $a_p$ are generally not compensated by concomitant deviations in particle scattering, so that the amplitude of reflectance is affected. Using spectral reflectance ratios, instead of absolute values of reflectance, greatly cancels out this limitation.

Appendix

Examining whether the parameterizations of $a_p(\lambda)$ and $K_d(\lambda)$ are consistent implies that the contributions to $K_d$ of absorption by CDOM and particle scattering are accounted for. For this we use the approximative formula proposed by Kirk [1981]:

$$K_d = (a^2 + 0.256 a b) \exp(0.14 \lambda)$$  \hspace{1cm} (10)

where $a$ and $b$, the total absorption and scattering coefficients, respectively, and $K_d$ are represented at the midpoint of the euphotic zone. Note that this simplified formula does not account for the influence of Sun elevation. The total absorption coefficient is expressed as

$$a(\lambda) = a_o(\lambda) + a_p(\lambda) + a_{d}(440) \exp \left[ -0.014(\lambda - 440) \right]$$  \hspace{1cm} (11)

in which $a_o(\lambda)$ is the absorption coefficient of pure seawater [Pope and Fry, 1997]. The term $a_{d}(440) \exp [-0.014 (\lambda - 440)]$ represents absorption by locally produced CDOM, which decreases exponentially with wavelength, with an average slope of 0.014 [Bricaud et al., 1981]. The contribution of CDOM to total absorption is not well documented. Prieur and Sathyendranath [1981] suggested that the absorption coefficient of non-algal matter at 440 nm contributes 20% of the total absorption coefficient at this wavelength. However, this amount included the contribution of non-algal particles, so that the actual contribution of dissolved yellow substance is probably lower. A review of existing measurements for oceanic waters [Kirk, 1994] shows that the absorption coefficient of yellow substance at 440 nm varies roughly from -0 to 0.1 m$^{-1}$ from clear to eutrophic waters and is likely to vary for a given particulate content. Here the absorption coefficient of CDOM at 440 nm was tentatively represented as

$$a_{o}(440) = y (a_o + a_{d}(440))$$  \hspace{1cm} (12)

with $y = 0$, 0.1 or 0.2, so that it is allowed to increase with the total absorption coefficient at various rates; the inclusion of $a_o$ implies that a CDOM “background” remains, even in the clearest waters. The total scattering coefficient was represented as

$$b(\lambda) = b_o(\lambda) + [b_p(660)(660/\lambda)]$$  \hspace{1cm} (13)

with

$$b_p(660) = 0.252 (\mathrm{chl})^{0.635}$$  \hspace{1cm} (13')

in which $b_o(\lambda)$ is the scattering coefficient of pure seawater. Equation (13') was derived by Losel and Morel [1998] from a statistical analysis of 610 recent measurements of in situ attenuation within the upper homogeneous layer (their set 2 and 3); the second part of the bracketed term in (13) makes particle scattering spectrally varying as $\lambda^{-1}$ [e.g., Gordon and Morel, 1983]. The $b_o$ coefficients are those provided by Morel [1974]. Note that for the range of (chl) values examined in the comparison (0.1-10 mg m$^{-3}$; see Figure 3), the ratio $b_o/a$, as provided by (11) (13), varies from 3 to 15, which is in the domain of the validity of (10) ($b/a < 30$).

Notation

Some of these notations differ from those used by Bricaud et al. [1995] in order to conform with those recommended by the Joint Global Ocean Flux Study Task Team for Photosynthetic Measurements [Sakshaug et al., 1997].

- $\lambda$: wavelength, nm.
- $\langle \mathrm{chl} \rangle$: chlorophyll $a +$ divinyl chl $a$ + pheophtin $a$ concentration, mg m$^{-3}$.
- $a$: absorption coefficient of the water body, m$^{-1}$.
- $a_o$: absorption coefficient of pure seawater, m$^{-1}$.
- $a_p$: absorption coefficient of particles, m$^{-1}$.
- $a_{\phi}$: absorption coefficient of living phytoplankton, m$^{-1}$.
- $a_{\text{nap}}$: absorption coefficient of non-algal particles, m$^{-1}$.
- $a_{\text{ds}}$: absorption coefficient of colored dissolved organic matter, m$^{-1}$.

- $a^*_p$, $a^*_o$, $a^*_\text{nap}$: chlor-specific absorption coefficients, m$^2$ mg$^{-1}$.
- $S$: slope of the exponential function representing the spectral dependence of $a_{\text{nap}}$ (equation (5)), nm$^{-1}$.
- $A_{\rho}$, $F_{\rho}$: numerical constants in the $a_\rho$ versus $\langle \mathrm{chl} \rangle$ relationship (equation (6)).
- $A_\phi$, $E_\phi$: numerical constants in the $a_\phi$ versus $\langle \mathrm{chl} \rangle$ relationship.
- $K_d$: diffuse attenuation coefficient (for downwelling irradiance), m$^{-1}$.
- $K_w$: diffuse attenuation coefficient of pure seawater, m$^{-1}$.
- $K_{\text{bio}}$: Nonwater diffuse attenuation coefficient $(K_d - K_o)$, m$^{-1}$.
- $b$: scattering coefficient of the water body, m$^{-1}$.
- $b_o$: scattering coefficient of pure seawater, m$^{-1}$.
- $b_p$: scattering coefficient of particles, m$^{-1}$.
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