

1 Coastal Atmosphere & Sea Time Series (CoASTS) and Bio-Optical mapping of
2 Marine optical Properties (BiOMaP): the CoASTS-BiOMaP dataset.

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ABSTRACT

The *Coastal Atmosphere & Sea Time Series* (CoASTS) and the *Bio-Optical mapping of Marine optical Properties* (BiOMaP) programs produced bio-optical data supporting satellite ocean color applications for more than two decades. Specifically, relying on the Acqua Alta Oceanographic Tower (AAOT) in the northern Adriatic Sea, from 1995 till 2016 CoASTS delivered time series of marine water apparent and inherent optical properties, in addition to the concentration of major optically significant water constituents. Almost concurrently, from 2000 till 2022 BiOMaP produced equivalent spatially distributed measurements across major European Seas. Both, CoASTS and BiOMaP applied standardized instruments, measurement methods, quality control schemes and processing codes to ensure temporal and spatial consistency to data products. This work presents the CoASTS and BiOMaP near surface data products, named CoASTS-BiOMaP, of relevance for ocean color bio-optical modelling and validation activities.

26 1. Introduction

27 The validation of primary (*i.e.*, radiometric) and derived (*e.g.*, phytoplankton pigments
28 concentration) satellite data products, as well as the development of bio-optical algorithms
29 linking radiometric data to the inherent optical properties or to the concentration of natural water
30 optically significant constituents, require accurate and comprehensive *in situ* bio-optical
31 measurements (*e.g.*, see Werdell and Bailey 2007). Anticipating this need for the Sea-Wide
32 Field-of-View (SeaWiFS) ocean color mission, during the 90s several measurement programs
33 were established to gather bio-optical data representative of the world marine waters. Among
34 these, the *Coastal Atmosphere & Sea Time Series* (CoASTS) and the *Bio-Optical mapping of*
35 *Marine optical Properties* (BiOMaP) measurement programs implemented by the Marine
36 Optical Laboratory (Belward et al. 2022) of the Joint Research Center (JRC) in collaboration
37 with a number of European institutions, produced comprehensive *in situ* bio-optical
38 measurements of relevance for satellite ocean color applications. While CoASTS benefited of the
39 Acqua Alta Oceanographic Tower (AAOT) in the northern Adriatic Sea to generate time-series
40 data at a fixed coastal site (Berthon et al. 2002; Zibordi et al. 2002), BiOMaP relied on
41 oceanographic ships to collect spatially distributed measurements across various European Seas
42 (Berthon et al. 2008, Zibordi et al. 2011). Both CoASTS and BiOMaP endorsed standardization
43 of instruments, measurement methods, quality control schemes and processing codes to enforce
44 consistency to temporally and spatially distributed data products. It is still recognized that some
45 of the measurement methods primarily implemented for optically complex coastal waters, may
46 not warrant the desirable high accuracy in oligotrophic clear waters.

47 Overall, CoASTS and BiOMaP data extend over a period exceeding two decades and
48 constitute a unique dataset for bio-optical investigations across a variety of water types with
49 potential application to climate change studies. Objective of this work is to introduce the
50 CoASTS and BiOMaP derived data products relevant for satellite ocean color applications.
51 Specifically, the near-surface data products with spectral values restricted to key ocean colour
52 center-wavelengths are presented together with a description of the measurement and data
53 reduction methods.

54 55 2. The CoASTS and BiOMaP programs

56 CoASTS and BiOMaP were conceived as complementary programs: CoASTS focused on
57 the generation of time-series of reference data from a fixed coastal site showing significant
58 seasonal cycles and moderate bio-optical complexity (Berthon et al. 2002); conversely BiOMaP
59 covered a variety of marine regions exhibiting very diverse bio-optical regimes, but with limited
60 temporal representativity (Berthon et al. 2008).

61 The use of an oceanographic tower as logistic platform for comprehensive optical and
62 bio-geochemical measurements, when compared to oceanographic ships, does not allow for
63 spatially extended observations. However, it offers the unique opportunity of a very stable
64 measurement platform enabling easy control of the deployment geometry of optical instruments
65 with respect to the structure. Specifically, regardless of sea state, the use of the AAOT as
66 measurement platform made possible deploying optical sensors relying on tower–sensor–Sun
67 geometry favouring the application of corrections for the minimization of potential
68 superstructure perturbations in radiometric data (Zibordi et al. 1999, Doyle and Zibordi 2002).

69 CoASTS measurements are representative of marine frontal regions exhibiting
70 occurrence of waters with optical properties largely determined by phytoplankton and its
71 degradation components (*i.e.*, Case 1 waters), as wells as optically complex waters characterized

72 by moderate concentrations of sediments and coloured organic matter (CDOM), with bio-optical
73 variability determined by the impact of local currents, seasonal changes in biological regimes
74 and rivers discharge (Berthon et al. 2002).

75 CoASTS measurements took place with monthly occurrence since 1995. However, from
76 2001 and up to the end of the measurement program in 2016, the frequency of field
77 measurements was reduced to one every 2-3 months. Each comprehensive data collection —
78 called a *station* — included in-water optical and hydrographic profiles, seawater samples at
79 different depths (*i.e.*, near surface, 8 m and 14 m), meteorological data, and visual observations
80 of cloud cover and sea state. CoASTS comprises 176 field campaigns leading to 883
81 measurement stations. Still, only CoASTS campaigns and stations performed from December
82 1998 onward (*i.e.*, 125 and 617, respectively) fulfil the requirement of measurements
83 standardization.

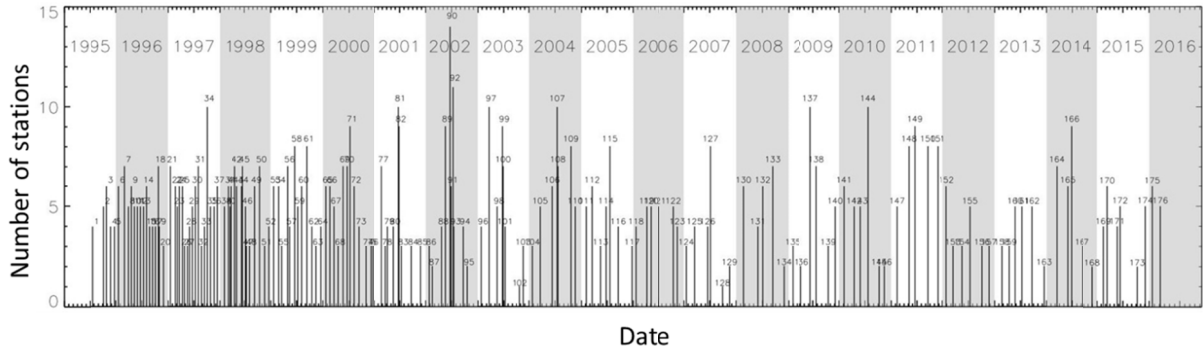
84 Spatially distributed measurements are best possible using oceanographic ships. Because
85 of this, BiOMaP measurements were performed relying on research vessels across a variety of
86 bio-optical regions (see Berthon et al. 2008): the Baltic Sea exhibiting waters dominated by a
87 high concentration of CDOM; the Adriatic Sea, Black Sea, North Sea (including the English
88 Channel), Ligurian Sea, Iberian Shelf and the Greenland Sea, characterized by a variety of
89 optically complex waters determined by diverse concentrations of CDOM and suspended
90 particulate matter (*SPM*); the Eastern and Western Mediterranean oligotrophic and mesotrophic
91 Seas with optical properties largely determined by phytoplankton and its degradation
92 components.

93 BiOMaP, encompassing 36 bio-optical oceanographic campaigns and 1915 measurement
94 stations, started in 2000 and ended in 2022. It is mentioned that some measurements from 33
95 BiOMaP stations performed in the Black Sea during 2011 were included in an independent
96 dataset constructed to support the validation of satellite data products (Valente et al. 2016).

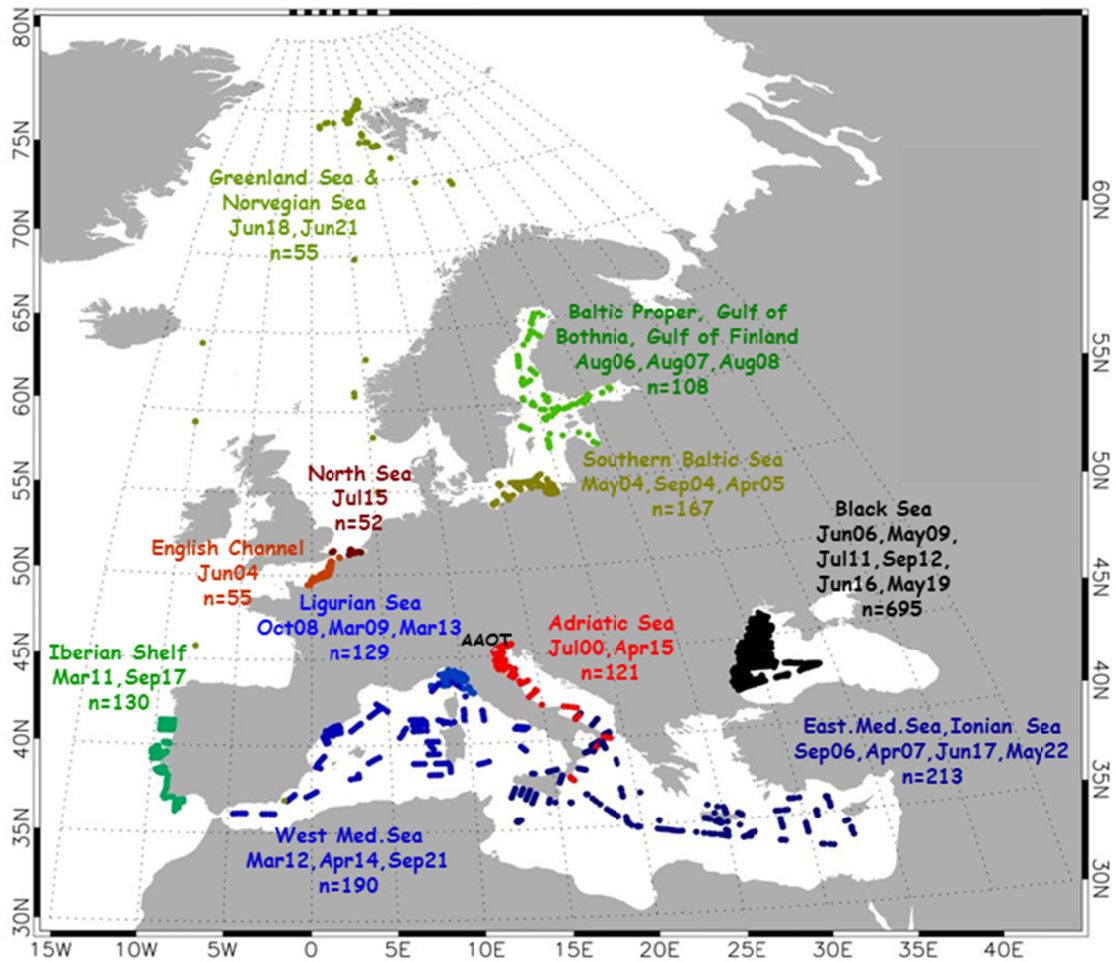
97 As already anticipated, measurement consistency between the CoASTS and BiOMaP
98 programs was achieved using identical field and laboratory instrumentation, and applying the
99 same consolidated methods, quality control schemes and processing codes. Consequently,
100 BiOMaP measurements performed during each station exhibit equivalence with those of
101 CoASTS, except for restricting the collection of water samples to the near surface. Finally,
102 superstructure perturbations in BiOMaP radiometric data were avoided by operating optical
103 radiometers on free-fall profilers deployed at some distance from ships (IOCCG 2019).

104 Figure 1 shows the temporal evolution of CoASTS measurement campaigns and the number
105 of stations per campaign. These latter were largely benefitting of sea state conditions allowing
106 access to the tower. Figure 2 shows the overall distribution of BiOMaP stations across the
107 various European Seas.

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 111 Figure 1. CoASTS measurement campaigns (176 total, 125 since December 1998) and stations
 112 (883 total, 637 since December 1998) completed between 1995 and 2016.
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 115 Figure 2. BiOMaP oceanographic campaigns (36) and measurement stations (1915) performed
 116 between 2000 and 2022.

117 Table 1. The CoASTS measurement program: campaign identifiers, marine region, years,
 118 number of stations, research platform, collaborating institution.
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Campaign ID	Location	Year	Stations #	Research platform	Collaborating Institution
V03-V99	Northern Adriatic Sea (AAOT)	1998-2011	481	<i>Acqua Alta Oceanog. Tower (AAOT)</i>	Italian National Research Council (IT)
W01-W28	Northern Adriatic Sea (AAOT)	2011-2016	136	<i>Acqua Alta Oceanog. Tower (AAOT)</i>	Italian National Research Council (IT)

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Table 2. The BiOMaP measurement program: campaign identifiers, marine regions, year, number of stations, research vessels, collaborating institutions.

Campaign ID	Region	Year	Stations #	Research vessel	Collaborating Institution
A01	Adriatic Sea (ADRS)	2000	55	<i>R/V Friuli-Venezia Giulia (FVG)</i>	University of Trieste (IT)
A02	Adriatic Sea (ADRS)	2014	66	<i>R/V Minerva-1</i>	Italian National Research Council (IT)
B01	Baltic Sea (BLTS)	2004	52	<i>R/V Oceania</i>	Institute of Oceanology (PL)
B02	Baltic Sea (BLTS)	2004	52	<i>R/V Oceania</i>	Institute of Oceanology (PL)
B03	Baltic Sea (BLTS)	2005	63	<i>R/V Oceania</i>	Institute of Oceanology (PL)
B04	Baltic Sea (BLTS)	2006	23	<i>R/V Aranda</i>	Institute of Marine Research (FI)
B05	Baltic Sea (BLTS)	2007	38	<i>R/V Aranda</i>	Institute of Marine Research (FI)
B06	Baltic Sea (BLTS)	2008	47	<i>R/V Aranda</i>	Institute of Marine Research (FI)
E01	Eastern Med. Sea (EMED)	2006	62	<i>R/V Urania</i>	Italian National Research Council (IT)
E02	Eastern Med. Sea (EMED)	2007	69	<i>R/V Urania</i>	Italian National Research Council (IT)
E03	Eastern Med. Sea (EMED)	2017	51	<i>R/V Minerva-1</i>	Italian National Research Council (IT)
E04	Eastern Med. Sea (EMED)	2022	31	<i>R/V Philia</i>	Hellenic Centre for Marine Research (GR)
I01	Iberian Shelf (IBSH)	2011	68	<i>NRP Almirante Gago Coutinho</i>	Portuguese Hydrographic Institute (PT)
I02	Iberian Shelf (IBSH)	2017	62	<i>NRP Almirante Gago Coutinho</i>	Portuguese Hydrographic Institute (PT)
K01	Black Sea (BLKS)	2006	93	<i>R/V Akademik</i>	Institute of Oceanology (BG)
K02	Black Sea (BLKS)	2009	73	<i>R/V Akademik</i>	Institute of Oceanology (BG)
K03	Black Sea (BLKS)	2009	40	<i>R/V Akademik</i>	Institute of Oceanology (BG)
K04	Black Sea (BLKS)	2011	38	<i>R/V Mare Nigrum</i>	National Institute of Marine Geology and Geocology (RO)
K05	Black Sea (BLKS)	2011	24	<i>R/V Akademik</i>	Institute of Oceanology (BG)

K06	Black Sea (BLKS)	2011	59	<i>R/V Akademik</i>	Institute of Oceanology (BG)
K07	Black Sea (BLKS)	2012	93	<i>R/V Akademik</i>	Institute of Oceanology (BG)
K08	Black Sea (BLKS)	2012	14	<i>R/V Akademik</i>	Institute of Oceanology (BG)
K09	Black Sea (BLKS)	2016	54	<i>R/V Akademik</i>	Institute of Oceanology (BG)
K10	Black Sea (BLKS)	2016	83	<i>R/V Akademik</i>	Institute of Oceanology (BG)
K11	Black Sea (BLKS)	2019	80	<i>R/V Akademik</i>	Institute of Oceanology (BG)
K12	Black Sea (BLKS)	2019	44	<i>R/V Akademik</i>	Institute of Oceanology (BG)
L01	Ligurian Sea (LIGS)	2008	41	<i>R/V Alliance</i>	Undersea Research Center (NATO)
L02	Ligurian Sea (LIGS)	2009	63	<i>R/V Alliance</i>	Undersea Research Center (NATO)
L04	Ligurian Sea (LIGS)	2013	25	<i>R/V Alliance</i>	Undersea Research Center (NATO)
N01	English Channel & North Sea (NORS)	2004	55	<i>R/V Côtes de la Manche</i>	Université du Littoral Côte d'Opale (FR)
N02	North Sea (NORS)	2015	52	<i>R/V Belgica</i>	Royal Belgian Institute of Natural Sciences (BE)
O01	Western Med. Sea (WMED)	2012	73	<i>R/V Urania</i>	Italian National Research Council (IT)
O02	Western Med. Sea (WMED)	2014	64	<i>R/V Urania</i>	Italian National Research Council (IT)
O03	Western Med. Sea (WMED)	2021	53	<i>R/V Garcia del Cid</i>	Institute of Marine Science (ES)
P01	Greenland Sea (GRLS)	2018	15	<i>R/V Alliance</i>	Undersea Research Center (NATO)
P03	Greenland Sea ¹ (GRLS)	2021	40	<i>R/V Alliance</i>	Italian Hydrographic Institute (IT)

¹ It includes stations from the Norwegian Sea.

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3. Measurements overview

CoASTS and BiOMaP core data comprise *in situ* and laboratory measurements performed on samples prepared in the field. The firsts include:

- a. Multispectral profiles of upwelling nadir radiance $L_u(z,\lambda)$, downward irradiance $E_d(z,\lambda)$, and upward irradiance $E_u(z,\lambda)$, where z indicates the depth and λ the center-wavelength of each spectral band;
- b. Multispectral above-water downward irradiance $E_s(t,\lambda)$ acquired during in-water profiling (where t is the time corresponding to the depth z) and diffuse sky irradiance $E_i(t,\lambda)$ acquired at the end of each station with an irradiance sensor operated in conjunction with a rotating shadow band;
- c. Multispectral profiles of beam attenuation $c(z,\lambda)$, absorption $a(z,\lambda)$ and backscattering $b_b(z,\lambda)$ coefficients, commonly restricted to the first 25 m depth for BiOMaP and 15 m for CoASTS;
- d. Profiles of water temperature $T_w(z)$ and salinity $S_w(z)$, also restricted to the first 25 m depth for BiOMaP and 15 m for CoASTS;

- 143 e. Meteorological data including wind speed W_s in addition to cloud cover C_c and sea state
144 S_s observations.
145 The laboratory measurements performed on field samples, complementary to the *in situ* ones,
146 are:
147 f. Spectral *in vivo* particulate absorption coefficients $a_{ph}(\lambda)$ for the pigmented and $a_{dt}(\lambda)$ for
148 the non-pigmented particles;
149 g. Spectral CDOM absorption coefficient $a_{ys}(\lambda)$;
150 h. Phytoplankton pigments concentration;
151 i. Suspended particulate matter concentration *SPM*.

152 153 **4. Measurements and data reduction methods**

154 Information on measurement methods and data reduction are summarized in the following
155 subsections.

156 157 **4.1 Radiometric products**

158 CoASTS in-water radiometric measurements of $L_u(z,\lambda)$, $E_d(z,\lambda)$, $E_u(z,\lambda)$ were performed
159 with the Wire-Stabilized Profiling Environmental Radiometer (WiSPER) using Satlantic
160 (Halifax, Canada) OCR/OCI-200 multispectral radiometer series. Concurrently, above-water
161 $E_s(t,\lambda)$ and $E_i(t,\lambda)$ measurements were also collected with OCI-200 radiometers. In the case of
162 BiOMaP, the equivalent measurements were performed using miniPRO and microPRO Satlantic
163 custom designed free-fall profilers equipped with OCR/OCI-200 or alternatively OCR-507
164 multispectral radiometers. All radiometric quantities were measured with 6 Hz acquisition rate at
165 spectral bands relevant for ocean color applications with 10 nm bandwidth and nominal center-
166 wavelengths at 412, 443, 490, 510, 555, 665 and 683 nm. WiSPER data were gathered with a
167 deployment speed of 0.1 m s^{-1} . Conversely, the deployment speed of the free-fall systems
168 generally varied in the range of approximately $0.3\text{-}0.4 \text{ m s}^{-1}$. The collection of in-water
169 radiometric measurements with low tilt and as close as possible to the surface, was always
170 attempted to ensure best retrieval of subsurface radiometric values through the extrapolation of
171 profile data.

172 The regular absolute radiometric calibration of field optical radiometers was performed at
173 the JRC Marine Optical Laboratory using 1000W FEL lamps traceable to the National Institute
174 of Standards and Technology (NIST) or alternatively the National Physical Laboratory (NPL).
175 While CoASTS radiometers were re-calibrated on a six-monthly basis, BiOMaP radiometers
176 were calibrated before and after each oceanographic campaign. Regular inter-calibrations
177 between the JRC Marine Optical Laboratory and the National Aeronautics and Space
178 Administration (NASA) performed within the framework of the Ocean Color component of the
179 Aerosol Robotic Network (AERONET-OC), ensured continuous verification of the accuracy of
180 the calibration process (Zibordi et al. 2021).

181 Data pre-processing included: *i.* the application of absolute calibration coefficients and
182 immersion factors for in-water radiometers (Zibordi et al. 2004; Zibordi 2006); *ii.* the removal of
183 in-water and in-air data exhibiting tilt higher than 5° (this was confidently established from 2009
184 for BiOMaP $E_s(t,\lambda)$ and $E_i(t,\lambda)$); *iii.* limited to BiOMaP, the composition of successive profile
185 data typically collected within a 5 min interval to create multi-cast combined profiles to increase
186 the number of measurements per unit depth and consequently improve the precision of
187 extrapolated values; and *iv.* the correction of in-air irradiance data for the non-cosine response of
188 collectors (see Zibordi and Bulgarelli 2007). Additional corrections for sensors non-ideal

189 performance, such as out-of-band response or temperature dependence, were not implemented
 190 being considered minor for the multispectral instruments applied.

191 In agreement with consolidated protocols (*e.g.*, see IOCCG 2019), the impact of illumination
 192 changes in profile data were minimized through normalization of each radiometric quantity with
 193 respect to above–water downward irradiance $E_s(t, \lambda)$ simultaneous to the in-water measurements.
 194 Specifically, the normalization aimed at producing radiometric quantities as if they were taken at
 195 the same time t_0 at each depth z , where t_0 was chosen to coincide with the beginning of the
 196 acquisition sequence during each cast or multi-cast.

197 The sub-surface quantities $L_u(0^-, \lambda)$, $E_u(0^-, \lambda)$ and $E_d(0^-, \lambda)$ were then determined at the
 198 depth $z_0 = 0$ (identified by 0^-) as the exponentials of the intercepts resulting from the least-
 199 squares linear regressions of $\ln \mathfrak{S}(z, \lambda)$ versus z within the extrapolation interval $z_0 - z_l$, where
 200 $\mathfrak{S}(z, \lambda)$ indicates either $L_u(z, \lambda)$, $E_d(z, \lambda)$ or $E_u(z, \lambda)$ normalized with respect to $E_s(t, \lambda)$ at matching
 201 times. The extrapolation interval was chosen on a profile-by-profile basis with the aid of
 202 absorption and scattering profile data to identify the depths z_0 and z_l , generally comprised within
 203 0.3 and 5 m and best satisfying the requirement of linear decay with depth of the log-transformed
 204 radiometric values. It is pointed out that the application of linear extrapolations to log-
 205 transformed data to determine sub-surface radiometric values, alternative to use of non-linear
 206 exponential extrapolations (see D’Alimonte et al 2012), was suggested by the objective to ensure
 207 consistency with existing radiometric data datasets.

208 Extreme outliers in the $z_0 - z_l$ depth interval generally due to major wave focusing and
 209 shadowing effects, were excluded from the extrapolation process by removing points exhibiting
 210 distance higher than $3 \cdot \sigma$ from the linear regression line, where σ is the standard deviation of the
 211 differences between data points and regression line. This filtering process is mostly effective in
 212 the presence of a relatively small number of points in the extrapolation layer. The application of
 213 a very slow deployment speed in the case of CoASTS radiometric data and the application of the
 214 multi-cast method for BiOMaP data, ensured the availability of hundreds of measurements in
 215 each selected extrapolation interval. This restricts the application of the $3 \cdot \sigma$ filter to a few
 216 extreme values without significantly impacting the precision of the extrapolated data.

217 The $L_u(0^-, \lambda)$ and $E_u(0^-, \lambda)$ data products were corrected for self-shading and potential
 218 bottom perturbations (Zibordi et al. 2002). Additionally, limited to CoASTS data collected in
 219 the vicinity of the AAOT, corrections were also applied for perturbations due to the deployment
 220 structure (Doyle and Zibordi 2002, Doyle et al. 2003). BiOMaP data, generally collected at
 221 distances from the ship of approximately 15–30 m, did not require corrections for the
 222 perturbations by the deployment structure.

223 In addition to $L_u(0^-, \lambda)$, $E_u(0^-, \lambda)$ and $E_d(0^-, \lambda)$, further retrieved data products are the
 224 slopes of the regression fits $K_{\mathfrak{S}}(\lambda)$ (*i.e.*, $K_L(\lambda)$, $K_u(\lambda)$ and $K_d(\lambda)$) in the extrapolation interval,
 225 *i.e.*, the diffuse attenuation coefficients. These $K_{\mathfrak{S}}(\lambda)$ values and particularly $K_d(\lambda)$, may exhibit
 226 underestimated values due to the impact of wave focussing in the near surface water layer. This
 227 effect is expected to be more pronounced for radiometric profiles collected in clear waters during
 228 clear sky conditions.

229 Derived radiometric data products are then the remote sensing reflectance $R_{rs}(\lambda)$

$$230 \quad 231 \quad 232 \quad R_{rs}(\lambda) = L_w(\lambda) / E_s(\lambda) \quad (1)$$

233 and the normalized water–leaving radiance $L_{wn}(\lambda)$

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$$L_{wn}(\lambda) = R_{rs}(\lambda)E_0(\lambda), \quad (2)$$

where $E_s(\lambda)$ refers to the value measured at time t_0 , $E_0(\lambda)$ is the extra-atmospheric solar irradiance (Thuillier et al. 2003) at the mean sun-earth distance, and $L_w(\lambda)$ is the water-leaving radiance, *i.e.*, the radiance leaving the sea and quantified just above the surface through the extrapolation process, given by

$$L_w(\lambda) = 0.544 L_u(0^-, \lambda). \quad (3)$$

where the factor 0.544 accounts for the radiance reduction across the water surface due to the change in the refractive index at the air-water interface, as determined assuming that the refractive index of seawater is independent of wavelength (Austin 1974). It is acknowledged that the values of $L_w(\lambda)$ determined with Eq. 3 [assuming a flat sea surface](#) exhibit differences well within $\pm 1\%$ with respect to the values computed accounting for the spectral dependence of the water refractive index in the spectral range of interest (Voss and Flora 2017).

Finally, supplementary derived quantity is the Q -factor at nadir $Q_n(0^-, \lambda)$ determined by the ratio of $E_u(0^-, \lambda)$ to $L_u(0^-, \lambda)$ spectrally fitted to a quadratic function in the 412-555 nm spectral interval to minimize the impact of calibration and extrapolation uncertainties. The ratio of fitted $Q_n(0^-, \lambda)$ to $E_u(0^-, \lambda) / L_u(0^-, \lambda)$ provides a basic approach to evaluate the relative consistency of the $E_u(0^-, \lambda)$ and $L_u(0^-, \lambda)$ multispectral values (*e.g.*, any appreciable bias affecting a single spectral value leads to a spectral inconsistency in $E_u(0^-, \lambda) / L_u(0^-, \lambda)$).

The quantities $R_{rs}(\lambda)$ and $L_{wn}(\lambda)$, due to the normalization with respect to $E_s(\lambda)$, benefit of a first correction for changes in illumination conditions with sun zenith, sun-earth distance and atmospheric transmittance (Mueller and Austin 1995). The additional correction performed through the application of the $C_{f/Q}(\theta_0, \lambda, \tau_a, IOP)$ factors to $L_{wn}(\lambda)$ and analogously to $R_{rs}(\lambda)$, accounts for in-water bi-directional effects, and leads to the determination of the final $L_{WN}(\lambda)$ and $R_{RS}(\lambda)$ data products. The $C_{f/Q}$ factors are a function of the water inherent optical properties IOP (absorption and back-scattering coefficients), the atmospheric optical properties conveniently expressed through the aerosol optical depth τ_a and the sun zenith angle θ_0 . These correction factors were determined applying the tabulated values proposed by Morel et al. (2002) for Case 1 waters with $IOPs$ solely expressed as a function of total chlorophyll-*a* concentration ($Chla$) as determined from water samples for each measurement station. It is acknowledged that this correction may be affected by large uncertainties when applied to optically complex waters. Still, the inclusion of both $L_w(\lambda)$ and $E_s(\lambda)$, as well as spectral values of the water inherent optical properties, would allow any potential user of the CoASTS-BiOMaP data set to implement alternative solutions for the determination of $L_{WN}(\lambda)$ and $R_{RS}(\lambda)$.

An estimate of the uncertainties for CoASTS and BiOMaP L_{WN} and similarly R_{RS} data, was attempted and discussed in various publications (Zibordi and Voss 2010, Zibordi et al. 2011) accounting for the major uncertainties characterizing: *i.* absolute calibration coefficients and immersion factors; *ii.* correction factors for shading perturbations; *iii.* correction factors for in-water bidirectional effects; *iv.* the determination of $E_s(\lambda)$; *v.* the quantification of $E_0(\lambda)$ when ignoring actual bandwidths; *vi.* the extrapolation process for the computation of sub-surface data; and *vii.* finally, environmental stability as a result of wave perturbations and changes in illumination conditions and seawater optical properties during profiling. In the specific case of moderate optically complex waters such as those characterizing CoASTS measurements, the uncertainties affecting L_{WN} and R_{RS} are expected to approach 5% in the blue green spectral

281 region and 7% in the red. In agreement with analyses performed for alternative *in situ*
282 radiometric methods (Gergely and Zibordi 2014), the above relative uncertainties may become
283 significantly larger in the blue spectral region for data products from marine regions
284 characterized by high water absorption such as the Baltic Sea.

285 Quality indices for radiometric products were determined during data processing in view
286 of supporting an evaluation of their accuracy. These include: *i.* the ratio $Q_R(412) = Q_n(0^-,412) /$
287 $Q_n(1,412)$ of Q_n values determined at 0^- and 1 m depths at the 412 nm center-wavelength, whose
288 significant deviation from 1 suggests issues in the extrapolation of sub-surface values; *ii.* the
289 coefficient of variation $CV_{E_s}(412)$ of in-air downward irradiance for the extrapolation interval,
290 whose high value indicates significant perturbations by ship movement or changes in
291 illumination conditions during profiling; *iii.* the diffuse to direct ratio of above-water downward
292 irradiance $R_d(412)$, whose high values indicates poor illumination conditions likely due to high
293 sun zeniths or cloudiness; *iv.* the index $R_i(412) = E_s(412) / [1.04 E_d(412)]$, whose significant
294 deviation from 1 indicates inconsistency between in-air and in-water measurements of the
295 downward irradiance; and finally *v.* the index $K_i(490)$ determined by the difference between
296 $K_d(490)$ and the corresponding value for pure water $K_w(490)$ set to 0.0212 m^{-1} (Smith and
297 Baker 1981), whose negative value identifies radiometric data products (mostly related to clear
298 waters and clear sky conditions) significantly challenged by wave perturbations.

299

300 **4.2 Absorption and attenuation from profile data**

301 Beam attenuation $c_{t-w}(z, \lambda)$ and absorption $a_{t-w}(z, \lambda)$ coefficients, excluding the contribution of
302 pure seawater, were determined from measurements performed using AC9s instruments from
303 WET Labs Inc. (Philomath, Oregon) with 25 cm path-length and nine spectral bands 10 nm wide
304 at the 412, 440, 488, 510, 555, 630, 650, 676 and 715 nm center-wavelengths. The values of
305 $c_{t-w}(z, \lambda)$ and $a_{t-w}(z, \lambda)$, in agreement with the scheme proposed by the instrument manufacturer
306 (WET Labs 2006), were corrected for the effects of differences in temperature T_w and salinity S_w
307 between field measurements and laboratory calibrations. These corrections were performed using
308 $T_w(z)$ and $S_w(z)$ profile data simultaneous to the AC9 ones.

309 AC9 absorption coefficients need correction for the non-completely reflective surfaces of
310 the absorption measurement tube, which prevents the collection of the whole scattered light and
311 naturally leads to an overestimate of $a_{t-w}(z, \lambda)$. This correction was performed by removing a
312 variable percentage of the scattering coefficient $b_{t-w}(z, \lambda)$ estimated from the difference between
313 $c_{t-w}(z, \lambda)$ and $a_{t-w}(z, \lambda)$ at each λ , assuming the absorption coefficient of particulate and dissolved
314 material is zero at the reference wavelength $\lambda_0 = 715$ nm and the shape of the volume scattering
315 function is spectrally independent (Zaneveld et al. 1992). Recent investigations showed this
316 correction method may lead to significant underestimates of $a_{t-w}(z, \lambda)$. Still, it was used in the
317 CoASTS-BiOMaP data processing because alternative promising correction methods such as that
318 proposed by Roettgers et al. (2013) may not be universally applicable (Stockley et al. 2017).
319 Nevertheless, the potential for applying alternative scattering corrections is allowed by including
320 in the dataset the absorption values at 715 nm, $a_{t-w}(z, 715)$, not corrected for the scattering offset
321 ($a_{t-w}(z, 715)$ would be zero when corrected).

322 The additional correction for the finite acceptance angle of the detector, which would need
323 additional field measurements of the volume scattering phase function (Boss et al. 2009) not
324 included among the CoASTS and BiOMaP core data, could not be implemented.

325 In addition to regular instrument calibration and maintenance by the manufacturer,
326 systematic AC9s pure water offsets were determined during each CoASTS campaign and at the

327 beginning and completion of each BiOMaP campaign in agreement with best practices for field
 328 operation. This offset accounts for any instrument response change while the AC9s are operated
 329 in their actual deployment configuration. The absorption and scattering offsets between the
 330 reference manufacturer calibrations and those performed in the field were applied as correction
 331 values. In the presence of appreciable offsets between successive field calibrations performed
 332 during the same campaign, differences were linearly interpolated over time.

333 Automated quality control was applied to each data record to verify the spectral and spatial
 334 (*i.e.*, vertical) consistency aiming at identifying those measurements affected by perturbations
 335 caused by bubbles or large particles flowing into the AC9 measurement chambers (*i.e.*, mostly
 336 individual spikes independently affecting $c_{t-w}(z,\lambda)$ or $a_{t-w}(z,\lambda)$ measurements especially in the
 337 surface layer). Specifically, $c_{t-w}(z,\lambda)$ and $a_{t-w}(z,\lambda)$ spectra exhibiting pronounced differences with
 338 respect to those characterizing the mean of profile spectra determined through a spectral
 339 consistency test, or pronounced changes with respect to depth at any λ identified through a
 340 spatial consistency test, were removed. The statistical parameters characterizing such a filtering
 341 process were tuned for profile data typical of individual campaigns in view of minimizing the
 342 potential for removing valid measurements.

343 The quality controlled $c_{t-w}(z,\lambda)$ and $a_{t-w}(z,\lambda)$ data were successively binned at 1 m
 344 resolution and retained when the depth d_b assigned to the center of the bin determined from the
 345 mean of the actual depths of individual measurements satisfies the condition $d_b = d_n \pm 0.25 \cdot d_i$,
 346 where d_n is the nominal depth of the center of the bin and d_i the bin width. The $c_{t-w}(\lambda)$ and
 347 $a_{t-w}(\lambda)$ values included in the CoASTS-BiOMaP dataset are the binned values tentatively
 348 corresponding to an average depth of 1 m.

349 A minimum uncertainty of 0.005 m^{-1} is assumed to affect AC9 measurements
 350 (Twardowski *et al.* 2001). Still, Stockley *et al.* (2017) showed that these values are largely
 351 underestimated especially in highly scattering waters in the blue-green spectral bands.

352

353 **4.3 Backscattering from profile data**

354 *In situ* vertical profiles of backscattering coefficients $b_b(z,\lambda)$ were determined using
 355 measurements performed with HydroScat-6 instruments from HOBI Labs Inc. (Tanque Verde,
 356 Arizona) in six bands 10 nm wide at the 442, 488, 510, 555, 620 and 676 (or 671) nm center-
 357 | wavelenghts. The values of $b_b(z,\lambda)$ were derived applying the conversion factor $\chi=1.08$ to
 358 measurements of the volume scattering function $\beta(z,\psi,\lambda)$ performed at the sole scattering angle
 359 $\psi=140^\circ$ (Maffione and Dana 1997). The derived backscattering values were successively
 360 corrected for the water scattering and absorption applying the factor

361

$$362 \quad \sigma_c(z, \lambda) = \exp[k_e(\lambda)(a(z, \lambda) + 0.4b(z, \lambda))] \quad (4)$$

363

364 where $a(z,\lambda)$ and $b(z,\lambda)$ (with $b(z,\lambda)=c(z,\lambda)-a(z,\lambda)$) were obtained from AC9 measurements
 365 adding the pure water absorption and scattering coefficients, respectively, while the instrument
 366 specific spectral factors $k_e(\lambda)$ were those determined by the manufacturer during the initial
 367 calibration. Salinity corrections were applied considering *Fresh water* b_{bw} from Morel (1974) for
 368 the Black Sea and Baltic Sea measurements, and the *Salt water* b_{bw} from Morel (1974) for the
 369 other measurements. This solution, with respect to the use of actual salinity values may lead to
 370 misestimates of b_{bb} generally not exceeding 2% at 443 nm for the Baltic Sea and Black Sea.

371 The accuracy of the applied equation 4 was questioned by Doxaran *et al.* (2016). However, their
 372 newly derived relationship was determined from $b_b(550)$ values comprised in the 0-2.5 m^{-1} range while

373 the CoASTS and BiOMaP $b_b(555)$ values are lower than 0.1 m^{-1} with $a_{t-w}(555)$ not exceeding 1.0 m^{-1} . In
374 this interval the equation proposed by Doxaran et al. (2016) does not appear to closely fit the plotted data
375 (see their Fig. 5b). Because of this, still acknowledging their work, the processing equations originally
376 proposed by Maffione and Dana (1997) for HydroScat-6 were applied

377 Equivalent to AC9 measurements, automated quality control was also applied to $b_b(z,\lambda)$
378 data to remove measurements exhibiting poor spectral and spatial (*i.e.*, vertical) consistency (*i.e.*,
379 mostly individual spikes affecting $b_b(z,\lambda)$ at a single λ). By tuning the parameters defining the
380 filtering process, spectra of $b_b(z,\lambda)$ exhibiting extreme differences with respect to the mean of
381 profile spectra, or very high changes with depth at any λ , were removed. Quality controlled
382 $b_b(z,\lambda)$ data were also binned at 1 m resolution adopting the same criteria applied for $a(z,\lambda)$ and
383 $c(z,\lambda)$. The $b_b(\lambda)$ values included in the CoASTS-BiOMaP dataset are the binned values
384 tentatively corresponding to an average depth of 1 m.

385 The quality index defined by the difference between $b_b(488)$ and the corresponding value
386 $b_{bw}(488)$ is included in the dataset to identify those measurements mostly collected in very
387 clear waters challenged by measurement uncertainties. The values of $b_{bw}(488)$, set equal to
388 0.001603 m^{-1} or alternatively equal to 0.001233 m^{-1} for the sole Black Sea and Baltic Sea data,
389 were determined from those provided in Morel (1974) fitted according to Twardowski et al.
390 (2007).

391 Annual factory calibrations performed at HOBILabs were complemented by pre-field
392 laboratory verifications performed at the JRC Marine Optical Laboratory. These laboratory
393 verifications aimed at correcting for HysdroScat-6 response changes between factory
394 calibrations.

395 Whitmire et al. (2007) estimated minimum uncertainties of 0.0007 m^{-1} for measurements of
396 $b_{bp}(z,\lambda)$ (*i.e.*, $b_b(z,\lambda)$ minus the backscattering of pure water) performed with HydroScat-6
397 instruments. Still, also in this case, actual uncertainties are expected to be much larger.

399 **4.4 Absorption of particulate matter determined from discrete water samples**

400 In vivo absorption coefficients $a_p(z,\lambda)$ of aquatic particles from water samples at discrete
401 depths z were determined using the Transmission and Reflection (*T-R*) method proposed by
402 Tassan and Ferrari (1995). This method was shown appropriate for any particle type, including
403 highly back-scattering mineral particles or highly absorbing sediments. The method was
404 implemented on a Perkin Elmer Lambda-19 and from 2004 on a Lambda-950, dual beam
405 spectrometers equipped with integrating spheres.

406 Samples of particles were collected filtering water volumes on Whatman GF/F glass fibre
407 filters with nominal pore size of $0.7 \text{ }\mu\text{m}$. Samples from the field were preserved in liquid
408 nitrogen until laboratory analysis. The absorption coefficient $a_p(z,\lambda)$ of the equivalent particle
409 suspension in the 400-750 nm spectral range with 1 nm resolution was determined from

$$410 \quad a_p(z,\lambda) = 2.3 A_s(z,\lambda) (F_a / V_w(z))^{-1} \quad (5)$$

413 where $V_w(z)$ is the volume of filtered water, F_a the filter clearance area and $A_s(z,\lambda)$ the equivalent
414 particle suspension absorbance obtained with the *T-R* method.

415 The pigmented $a_{ph}(z,\lambda)$ and non-pigmented $a_{dp}(z,\lambda)$ fractions of the particulate absorption
416 coefficient $a_p(z,\lambda)$ were obtained bleaching the sample using a solution of sodium hypochlorite
417 (NaClO). The solution rapidly acts on pigment molecules and slowly on detritus making possible

418 a selective analysis of the two absorption components. A description of the bleaching technique
419 is presented in Tassan and Ferrari (1995) and in Ferrari and Tassan (1999).

420 Focused studies on the accuracy of the *T-R* method are given in Tassan and Ferrari
421 (1995) and in Tassan et al. (2000). Still, comprehensive uncertainty estimates for $a_{ph}(z,\lambda)$ and
422 $a_{dp}(z,\lambda)$ are not available. Nevertheless, dedicated analysis addressed the repeatability of in vivo
423 particulate absorption measurements performed with the *T-R* method (see Zibordi et al. 2002).
424 These investigated: *i.* repeated analysis of the same sample (*i.e.*, each sample was analysed
425 twice) and *ii.* the analysis of duplicate samples (*i.e.*, duplicates obtained from the same water
426 volume). Results for repeated analysis of the same samples showed mean absolute percent
427 differences of $2.9\pm 2.3\%$ at 443 nm with mean $a_p(443) = 0.082\pm 0.042 \text{ m}^{-1}$, increasing up to
428 $7.4\pm 6.0\%$ at 555 nm with mean $a_p(555) = 0.023\pm 0.011 \text{ m}^{-1}$. These differences are attributed to: *i.*
429 method sensitivity, and *ii.* variations in the mechanical re-positioning of the sample in front of
430 the aperture of the integrating sphere combined with spatial non-homogeneities of the particles
431 distribution on the filter.

432 The analysis of duplicate samples showed mean absolute percentage differences of
433 $8.9\pm 5.9\%$ at 443 nm with mean $a_p(443) = 0.090\pm 0.049 \text{ m}^{-1}$ and of $9.8\pm 7.0\%$ at 555 nm with
434 mean $a_p(555) = 0.024\pm 0.012 \text{ m}^{-1}$. The former differences, increased by a few percent with
435 respect to those given for the repeated analysis of samples, are justified by: *i.* unavoidable
436 differences in replicates due to inhomogeneity affecting the particles distributions on filters; and
437 also *ii.* inhomogeneity in the distribution of particles in the water volumes used to produce the
438 samples. It is mentioned that an intrinsic error in the estimate of the actual particle absorption
439 coefficients results from the application of GF/F filters with nominal pore size of $0.7 \mu\text{m}$. In fact
440 these filters do not allow bacteria and the fraction of mineral particles with diameter lower than
441 $0.7 \mu\text{m}$ to be accounted for. However, the absorption of these small mineral particles is generally
442 negligible compared to the total absorption, while the absorption of bacteria is almost 10 times
443 lower than that of algal cells and 5–10 times lower than that of cyanobacteria (Morel and Ahn
444 1990). The $a_{ph}(z,\lambda)$ and $a_{dp}(z,\lambda)$ measurements included in the CoASTS-BiOMaP dataset refer to
445 water samples collected at approximately 1 m depth.

446

447 **4.5 Absorption of CDOM determined from discrete water samples**

448 The absorption coefficient $a_{ys}(z,\lambda)$ of CDOM at depth z was determined applying the
449 method detailed in Ferrari et al. (1996) using a Perkin Elmer Lambda-12 and from 2010 a
450 Lambda-35 dual-beam spectrometers. Samples were prepared by filtering water volumes on
451 Millipore $0.22 \mu\text{m}$ pore size cellulose filters and adding a solution of 10 g l^{-1} of NaN_3 to the
452 filtered water to prevent bacteria growth (typically 1 ml of the solution was added to 100 ml of
453 filtered water).

454 CDOM samples were preserved at approximately 4°C in an amber glass bottle until
455 laboratory analysis. The spectrometric measurements, generally carried out within a few days
456 from the completion of the measurement campaign, were performed with 1 nm resolution in the
457 350-750 nm spectral region. Measurements were performed placing a 10 cm quartz cuvette
458 containing pure milli-Q water in the optical path of the reference beam, and a 10 cm quartz
459 cuvette containing the CDOM sample in the optical path of the sample beam. It is acknowledged
460 that the 10 cm path-length systematically applied for the analysis of CoASTS and BiOMaP field
461 samples, naturally challenges the accuracy of measurements characterized by low CDOM
462 absorption such as those from the Eastern Mediterranean Sea.

463 The spectral absorption coefficient $a_{ys}(z,\lambda)$ was computed from the measured absorbance
464 $A_{ys}(z,\lambda)$ resulting from the difference between the sample absorbance and the reference
465 absorbance (Ferrari et al., 1996), as

$$466 \quad a_{ys}(z,\lambda) = 2.3 A_{ys}(z,\lambda) L_c^{-1} \quad (6)$$

467 where L_c is the pathlength of the cuvette.

468 Assuming CDOM does not absorb in the red [and following community recommendations](#)
469 [\(see Section III in IOOCG 2019\)](#), the absorption coefficients are corrected for the background
470 offset by subtracting to $a_{ys}(z,\lambda)$ the mean of $a_{ys}(z,\lambda_i)$ spectral values for λ_i in the 670-680 nm
471 interval.

472 Comprehensive uncertainty values are also not available for $a_{ys}(z,\lambda)$. Still, the
473 repeatability of $a_{ys}(z,\lambda)$ measurements (see Zibordi et al. 2002) was also investigated through: *i.*
474 repeated analysis of the same samples; and *ii.* the analysis of duplicate samples. The repeated
475 analysis of the same samples showed average absolute percent differences varying as a function
476 of the absorption value from $10.1\pm 7.3\%$ at 412 nm with mean $a_{ys}(z,412) = 0.168\pm 0.037 \text{ m}^{-1}$ up to
477 $24.2\pm 19.8\%$ at 555 nm with mean $a_{ys}(z,555) = 0.015\pm 0.005 \text{ m}^{-1}$. These differences are mostly
478 ascribed to the precision of the method. The analysis of the duplicate samples showed expected
479 augmented average absolute percent differences when compared to repeated analysis of samples
480 varying from $12.1\pm 6.3\%$ at 412 nm with mean $a_{ys}(z,412) = 0.175\pm 0.038 \text{ m}^{-1}$ and up to
481 $30.3\pm 23.8\%$ at 555 nm with mean $a_{ys}(z,555) = 0.018\pm 0.005 \text{ m}^{-1}$. The latter increased values are
482 largely justified by differences between samples.

483 It is finally mentioned that the use of $0.22 \mu\text{m}$ pore size filters to produce CDOM
484 samples, when the $0.7 \mu\text{m}$ pore size filters are applied for the quantification of particle
485 absorption coefficients, suggests that the overall absorption budget cannot be fully resolved. In
486 fact, as already anticipated, bacteria and very small mineral particles having size between 0.2 and
487 $0.7 \mu\text{m}$, are not included in the absorption analysis. Still, this missing contribution to the overall
488 absorption budget is expected to be minor.

489 As per $a_{ph}(\lambda)$ and $a_{dp}(\lambda)$, also the $a_{ys}(\lambda)$ measurements included in the CoASTS-BiOMaP
490 dataset refer to water samples collected at approximately 1 m depth.

491 **4.6 Pigments concentration**

492 Phytoplankton pigment concentrations were determined by using High Performance Liquid
493 Chromatography (HPLC) with the method proposed by Van Heukelem and Thomas (2005).
494 Exceptions are the samples collected before 2000 for which the method proposed by Jeffrey et al.
495 (1997) was applied.

496 The analysis were performed on samples of particulate matter retained on GF/F filters with a
497 nominal pore size of $0.7\mu\text{m}$: this choice is justified by the diameter of living phytoplankton cells
498 generally higher than $1 \mu\text{m}$ (Stramsky and Kiefer, 1991). After filtration, samples were preserved
499 in liquid nitrogen until laboratory analysis.

500 Following Van Heukelem and Thomas (2005), the samples were transferred to vials with 3
501 ml 95% acetone and vitamin E as internal standard. Samples were then disrupted using a vortex
502 mixer, sonicated on ice, extracted at 4°C for 20 h, and mixed again. The samples were
503 successively filtered through $0.2 \mu\text{m}$ Teflon syringe filter into HPLC vials and placed in the
504 cooling rack of the HPLC system. Buffer and sample were injected in the HPLC (Shimadzu LC-

508 10A or alternatively an HP-1100, systems) in the 5/2 ratio using a pre-treatment program and
509 mixing in the loop before injection.

510 The list of pigments systematically analysed at the JRC Marine Optical Laboratory or
511 alternatively at DHI A/S (Hørsholm, Denmark) includes: chlorophyll a (resulting from the sum
512 of divinyl- and monovinyl-chlorophyll a), chlorophyll b, chlorophyll c₁+c₂, chlorophyllide a,
513 fucoxanthin, diadinoxanthin, β-carotene, zeaxanthin, alloxanthin, 19'-butanoyloxyfucoxanthin,
514 19'-hexanoyloxyfucoxanthin and diatoxanthin.

515 Various inter-comparisons of HPLC methods performed within the framework SeaWiFS
516 HPLC Analysis Round-Robin Experiments (SeaHARRE) organized by NASA with the JRC
517 participation, demonstrated the capability of various laboratories to achieve differences lower
518 than 6% in the determination of total chlorophyll *a* concentration *Chla* (*i.e.*, the sum of
519 chlorophyll *a* and chlorophyllide *a*) and lower than 25% for the other ancillary pigments
520 characterizing marine waters (Hooker et al. 2010). The analysis of CoASTS and BiOMaP shows
521 *Chla* values always higher than 0.03 - 0.04 μg l⁻¹. This may suggest some quantification limit for
522 the methodology applied to determine pigments concentration.

523 Consistent with $a_{ph}(z,\lambda)$, $a_{dp}(z,\lambda)$ and $a_{ys}(z,\lambda)$, measurements of *Chla* were performed on
524 water samples collected at approximately 1 m depth.

525

526 4.7 Suspended particulate concentration

527 The concentration of suspended particulate matter, *SPM*, was obtained from the net weight
528 of the particulate material collected on filters following the method detailed in Van der Linde
529 (1998) as an evolution of that proposed by Strickland and Parsons (1972). Samples were
530 produced by filtering volumes of water on GF/F filters with 0.7 μm nominal pore size previously
531 baked at 450 °C for 1 hour, pre-washed, dried for 1 hour at 75 °C and finally pre-weighed on a
532 electrobalance. After water filtration, the filters (*i.e.*, filtration area and border) were washed
533 with distilled water and stored at -18 °C for successive laboratory analysis. Before final
534 weighting, the filters were dried at 75 °C for 1 hour, and then temporarily stored in a desiccator.

535 The concentration of *SPM* was calculated from

536

$$537 \quad SPM(z) = [(W_f(z) - W_s(z)) - w_b]V(z)^{-1} \quad (7)$$

538

539 where $W_f(z)$ is the weight of the filter before filtration, $W_s(z)$ is the weight of the filter after
540 filtration, $V(z)$ is the volume of the filtered water and w_b is a correction term introduced to
541 account for variations in the weight of the filter sample due to changes in environmental
542 conditions between the two weightings steps. The values of w_b were determined from *blank*
543 filters (*i.e.*, GF/F filters completely conditioned, not used for water filtration, but exposed to the
544 same processes of the sample filters: transportation to the measurement site and back, storage in
545 the freezer, drying). The w_b values applied in Eq. 7, are the differences between the average final
546 weight of *blank* filters and their original average weight.

547 *SPM* values included in the CoASTS-BiOMaP data set are generally obtained from the
548 average of duplicate samples. In the case of large differences between duplicates (*i.e.*, tentatively
549 exceeding 20%) the *SPM* value from one of the two samples is used prior investigating the
550 surface and integrity of the samples, and also verifying the consistency of their values with AC9
551 measurements from close stations.

552 The use of GF/F filters with 0.7 μm nominal pore size for *SPM* analysis leads to an
553 underestimate of total suspended matter due to the loss of particles with diameter lower than 0.7

554 μm . However, it is recognized that the filter rinsing for salt removal and the filter conditioning
555 after filtration before final weighting, can induce errors certainly much larger than the mass of
556 particles with diameter lower than $0.7 \mu\text{m}$.

557 An analysis of measurement repeatability performed with duplicate samples showed mean
558 percent difference equal to $13.9 \pm 13.4\%$ with mean $SPM(z) = 0.86 \pm 0.40 \text{ mg}\cdot\text{l}^{-1}$. The largest
559 differences between duplicate samples (*i.e.*, larger than 30%) were observed with values of
560 $SPM(z)$ lower than approximately 0.5 mg l^{-1} . This is explained by the intrinsic uncertainty
561 affecting sample preparation (*i.e.*, water sample non-homogeneity and filter rinsing).

562 As for other quantities determined from the analysis of water samples, also the SPM values
563 included in the CoASTS-BiOMaP dataset refer to samples collected at approximately 1 m depth.
564

565 **4.8 Salinity and temperature**

566 Profiles of salinity $S_w(z)$ and temperature $T_w(z)$ measurement were performed with SBE
567 19-plus Conductivity-Temperature-Depth (CTD) sensors from Sea-Bird Scientific (Bellevue,
568 Washington). These devices were calibrated by the manufacturer approximately on a two-year
569 basis. Uncertainties are tentatively expected to be within 0.01% for salinity and 0.01°C for
570 temperature.

571 Equivalent to $a(z, \lambda)$, $c(z, \lambda)$ and $b_b(z, \lambda)$ profiles, automated quality control was also
572 applied to $S_w(z)$ and $T_w(z)$ data to remove measurement artefacts. By trimming filtering parameters
573 to individual campaigns, values of $S_w(z)$ and $T_w(z)$ exhibiting extreme changes with respect to
574 depth, were removed. Quality checked $S_w(z)$ and $T_w(z)$ data were binned at 1 m resolution
575 adopting the same criteria already applied for $a(z, \lambda)$, $c(z, \lambda)$ and $b_b(z, \lambda)$. The values associated
576 with the first bin, tentatively representing the 1 m depth, are included in the CoASTS-BiOMaP
577 dataset.

578

579 **4.9 Meteorological and environmental observations**

580 Among the meteorological quantities and observations recorded during each measurement
581 station, the wind speed W_s , sea state S_s and cloud cover C_c are included in the data set.
582

583 **5. The near-surface CoASTS and BiOMaP dataset (CoASTS-BiOMaP)**

584 CoASTS-BiOMaP data are accessible at <https://doi.org/10.1594/PANGAEA.971945> in
585 tabular form and include the near-surface data products from CoASTS and BiOMaP
586 measurements of relevance for the validation of satellite ocean color data and the development of
587 bio-optical algorithms. All spectral data products are restricted to the nominal center-
588 wavelengths 412, 443, 490, 510, 555, 665 nm, unless diversely specified. CoASTS data products
589 are only provided from December 1998 when full standardization of measurements and
590 processing was put in place. In addition, station data were excluded from CoASTS-BiOMaP
591 when the $L_{WN}(\lambda)$ or $K_d(\lambda)$ radiometric products did not satisfy basic quality control criteria by
592 exhibiting spectra with unexplained shape or amplitude. Furthermore, poor quality of data
593 products other than radiometric, implied the exclusion of their individual values from the dataset.

594 Table 3 provides a comprehensive list of the quantities included in the CoASTS-BiOMaP
595 dataset: each one is identified by a convenient symbol, a brief description and its physical units.
596 A summary of the average values of the major bio-optical and hydrographic quantities
597 determined for the various marine regions is provided in Table 4. These are: the diffuse
598 attenuation coefficient K_d at 490 nm, the water absorption coefficient (from discrete sample

599 analysis, pure water contribution excluded) a at 490 nm, the backscattering coefficient (water
600 contribution included) b_b at 488 nm; the concentrations of total chlorophyll-a $Chla$ and
601 suspended particulate matter SPM ; and the salinity S_w . All quantities exhibit ample differences
602 across the various marine regions. Notable, variations in $K_d(490)$ exceed one order of magnitude
603 between the Eastern Mediterranean (EMED) and the Baltic Sea (BLTS) waters (*i.e.*, $K_d(490)$
604 varies from 0.037 to 0.494 m^{-1}).

605 Figure 3 displays BioMaP and CoASTS $L_{WN}(\lambda)$ spectra for the different marine regions.
606 These spectra clearly indicate diverse bio-optical features for the different regions. They span
607 from the highly oligotrophic Eastern Mediterranean Sea (EMED) showing maximum values in
608 the blue region, to the optically complex Baltic Sea (BLTS) dominated by the presence of high
609 concentrations of CDOM as expressed by low values of L_{WN} in the blue spectral region. Between
610 these, there are marine regions exhibiting diverse bio-optical complexity due to different
611 concentration of optically significant constituents. Notably, some spectra from the North Sea
612 (NORS) indicate the presence of relatively high concentration of sediments, while spectra from
613 the Black Sea (BLK) and the northern Adriatic Sea (AAOT) suggest bio-optical conditions
614 determined by the presence of various concentrations of SPM and CDOM determining L_{WN}
615 maxima at the 510 or 555 nm center-wavelengths.

616 Table 5 provides the mean spectral values and related standard deviations of $Q_n(\lambda)$ for the
617 various marine regions as determined from radiometric profiles performed during near clear sky
618 conditions determined by $Cc \leq 1/4$. These naturally exhibit some spectral dependence varying
619 with the water type. For instance, $Q_n(\lambda)$ from the Eastern Mediterranean Sea (EMED) exhibit
620 almost spectrally constant mean values approaching 4 sr in the 412-555 nm spectral interval and
621 of approximately 5 sr at 665 nm. Conversely, regions such as the northern Adriatic Sea (AAOT)
622 exhibit mean values approaching 4.5 sr with some spectral dependence in the 412-555 nm
623 spectral region, and also a mean value of 5 sr at 665 nm.

624 Figure 4 displays the $a_{ph}(\lambda)$ spectra for the CoASTS and BioMaP regions. Notable is the
625 increase in the values of mean $a_{ph}(443)$ from 0.007 m^{-1} for the Eastern Mediterranean Sea
626 (EMED) to 0.191 m^{-1} and 0.220 m^{-1} for the Baltic Sea (BLTS) and North Sea (NORS),
627 respectively. The peculiar spectra shown by North Sea stations off the Belgian coast exhibiting
628 a_{ph} values higher at 412 nm than at 443 nm (see panel for NORS data in Fig. 4), are explained by
629 high concentrations of pheophytin leading to an increase of the absorption coefficient toward 412
630 nm.

631 Figure 5 displays the comparison of the near surface absorption coefficients (pure water
632 excluded) determined from AC9 measurements at the center-wavelength of 443 nm, $a_{t-w}(AC9)$,
633 versus the equivalent absorption coefficients determined from water samples, $a_{t-w}(\text{sample}) =$
634 $a_{ph}(443) + a_{di}(443) + a_{ys}(443)$. Results suggest an increasing underestimate of $a_{t-w}(AC9)$ and larger
635 relative scattering with a decrease in absorption. This is highlighted by the scatter plots of data
636 from the Eastern Mediterranean Sea (EMED) exhibiting an underestimate exceeding 70% with
637 values of $a_{t-w}(\text{samples})$ generally lower than 0.1 m^{-1} . Conversely, the Baltic Sea (BLTS) shows
638 outstanding agreement between the compared quantities with absorption values comprised in the
639 range of 0.2-1.2 m^{-1} . These mean differences between $a_{t-w}(AC9)$ and $a_{t-w}(\text{samples})$ absorption
640 values could be explained by an incomplete correction of the perturbing effects due to finite
641 acceptance angle of the detector, the non-fully reflective surface of the AC9 absorption chamber
642 (*i.e.*, the two short 25 cm path-length tubes) and also by the non-negligible absorption of
643 particles at the reference wavelength $\lambda_0 = 715$ nm applied for scattering corrections.

644

645 Table 3. The CoASTS-BiOMaP data set: quantities identified by symbols, description of
646 quantities and related units.

Symbol	Description	Units	Details
<i>Station ID</i>	Station identifier	Code	<i>Gcessit</i> ⁽¹⁾
<i>Date&Time</i>	Date and time	GMT	<i>yyyy-mm-ddT hh:mm:ss</i> ⁽²⁾
<i>Lon</i>	Longitude	Degrees	
<i>Lat</i>	Latitude	Degrees	
<i>Sz</i>	Sun zenith	Degrees	
<i>Sa</i>	Sun azimuth	Degrees	
$L_u(\lambda)$	Upwelling radiance at depth 0 ⁻	$W m^{-2} nm^{-1} sr^{-1}$	At nominal λ_s ⁽³⁾
$E_d(\lambda)$	Downward irradiance at depth 0 ⁻	$W m^{-2} nm^{-1}$	At nominal λ_s ⁽³⁾
$E_u(\lambda)$	Upward irradiance at depth 0 ⁻	$W m^{-2} nm^{-1}$	At nominal λ_s ⁽³⁾
$K_L(\lambda)$	Diffuse att. coeff. from $L_u(z,\lambda)$	m^{-1}	At nominal λ_s ⁽³⁾
$K_d(\lambda)$	Diffuse att. coeff. from $E_d(z,\lambda)$	m^{-1}	At nominal λ_s ⁽³⁾
$K_u(\lambda)$	Diffuse att. coeff. from $E_u(z,\lambda)$	m^{-1}	At nominal λ_s ⁽³⁾
$E_s(\lambda)$	Downward irradiance at depth 0 ⁺	$W m^{-2} nm^{-1}$	At nominal λ_s ⁽³⁾
$Q_n(\lambda)$	Q -factor an nadir at depth 0 ⁻	sr	At nominal λ_s ⁽³⁾
$R_{RS}(\lambda)$	Remote sensing reflectance at depth 0 ⁺	sr^{-1}	At nominal λ_s ⁽³⁾
$L_{WN}(\lambda)$	Normalized water-leaving rad. at depth 0 ⁺	$W m^{-2} nm^{-1} sr^{-1}$	At nominal λ_s ⁽³⁾
$Q_R(412)$	Ratio of $Q_n(412)$ at depth 0 ⁻ to $Q_n(1,412)$ at 1 m depth	–	Introduced to best support the use of $Q_n(\lambda)$ (large deviations from 1 may indicate extrapolation issues)
$R_d(412)$	Ratio of the diffuse $E_i(412)$ to direct [$E_s(412) - E_i(412)$] above-water downward irradiance at 412 nm	–	
$CV E_s(412)$	Coefficient of variation $E_s(412)$	%	
$R_i(412)$	Ratio of the above-water downward irradiance $E_s(412)$ to the in-water downward irradiance $E_d(412)$ multiplied by 1.04	–	
$K_i(490)$	Diffuse attenuation coefficient $K_d(490)$ minus the diffuse attenuation coefficient of pure sea water $K_w(490)$ assumed constant and equal to 0.0212	m^{-1}	Introduced to best support the exploitation of data (a negative value may suggest extrapolation challenged by wave perturbations)
$a_{ph}(\lambda)$	Absorption coefficient by pigmented particles at 1 m depth	m^{-1}	At nominal λ_s ⁽³⁾
$a_{dt}(\lambda)$	Absorption coefficient by non-pigmented part. at 1 m depth	m^{-1}	At nominal λ_s ⁽³⁾
$a_{ys}(\lambda)$	Absorption coefficient by CDOM at 1 m depth	m^{-1}	At nominal λ_s ⁽³⁾
$a_{t-w}(\lambda)$	Absorption coefficient from AC9 at 1 m depth	m^{-1}	At AC9 λ_s ⁽⁴⁾ . The values $a_{t-w}(715)$ are not corrected for the scattering offset. If corrected, their values would be zero.
$c_{t-w}(\lambda)$	Beam attenuation coefficient from AC9 at 1 m depth	m^{-1}	At AC9 λ_s ⁽⁴⁾
$b_b(\lambda)$	Backscattering coefficient from HydroScat-6 at 1 m depth	m^{-1}	At HydroScat-6 λ_s ⁽⁵⁾
$b_b(488) - b_{bw}(488)$	Backscattering coefficient $b_b(488)$ minus	m^{-1}	Introduced to best support the

	the backscattering coefficient of pure sea water $b_{bw}(488)$ assumed constant and equal to 0.001603 m^{-1} or alternatively 0.001233 m^{-1} for the sole Black Sea and Baltic Sea measurements		exploitation of data (a negative value may indicate measurements challenged by significant uncertainties)
<i>Chla</i>	Total chlorophyll- <i>a</i> concentration at 1 m depth ⁽⁶⁾	$\mu\text{g l}^{-1}$	
<i>SPM</i>	Suspended particulate matter concentration at 1 m depth	mg l^{-1}	
T_w	Temperature of seawater at 1 m depth	$^{\circ}\text{C}$	
S_w	Salinity of seawater at 1 m depth	‰	
W_s	Wind speed	m s^{-1}	
S_s	Sea state	0-9	WMO scale
C_c	Cloud cover	0-4	Octa/2

647 ¹ *G* indicates the site or geographic region (*V* and *W* for AAOT, *A* for Adriatic Sea, *B* for Baltic Sea, *E* for Eastern
648 Mediterranean Sea, *K* for Black Sea, *L* for Ligurian Sea, *N* for North Sea, *O* for Western Mediterranean Sea, *I* for
649 Iberian Shelf, *P* for Greenland Sea), while *cc* indicates the campaign number for the specific region, *ss* the station
650 number and *ii* the cast number.

651 ² The letters *yyyy* indicate the year, *mm* the month, *dd* the day *hh*, the hours and *mm* the minutes.

652 ³ Nominal center-wavelengths for radiometric data products are 412, 443, 490, 510, 555 are 665 nm.

653 ⁴ Center-wavelengths for AC9 data products are 412, 440, 488, 510, 555, 630, 650, 676, and 715 nm.

654 ⁵ Center-wavelengths for HydroScat-6 data products are 442, 488, 510, 555, 620, and 676 (or 671) nm.

655 ⁶ Total chlorophyll-*a* concentration indicates the sum of chlorophyllide-*a*, monovinyl- and divinyl-chlorophyll-*a*.

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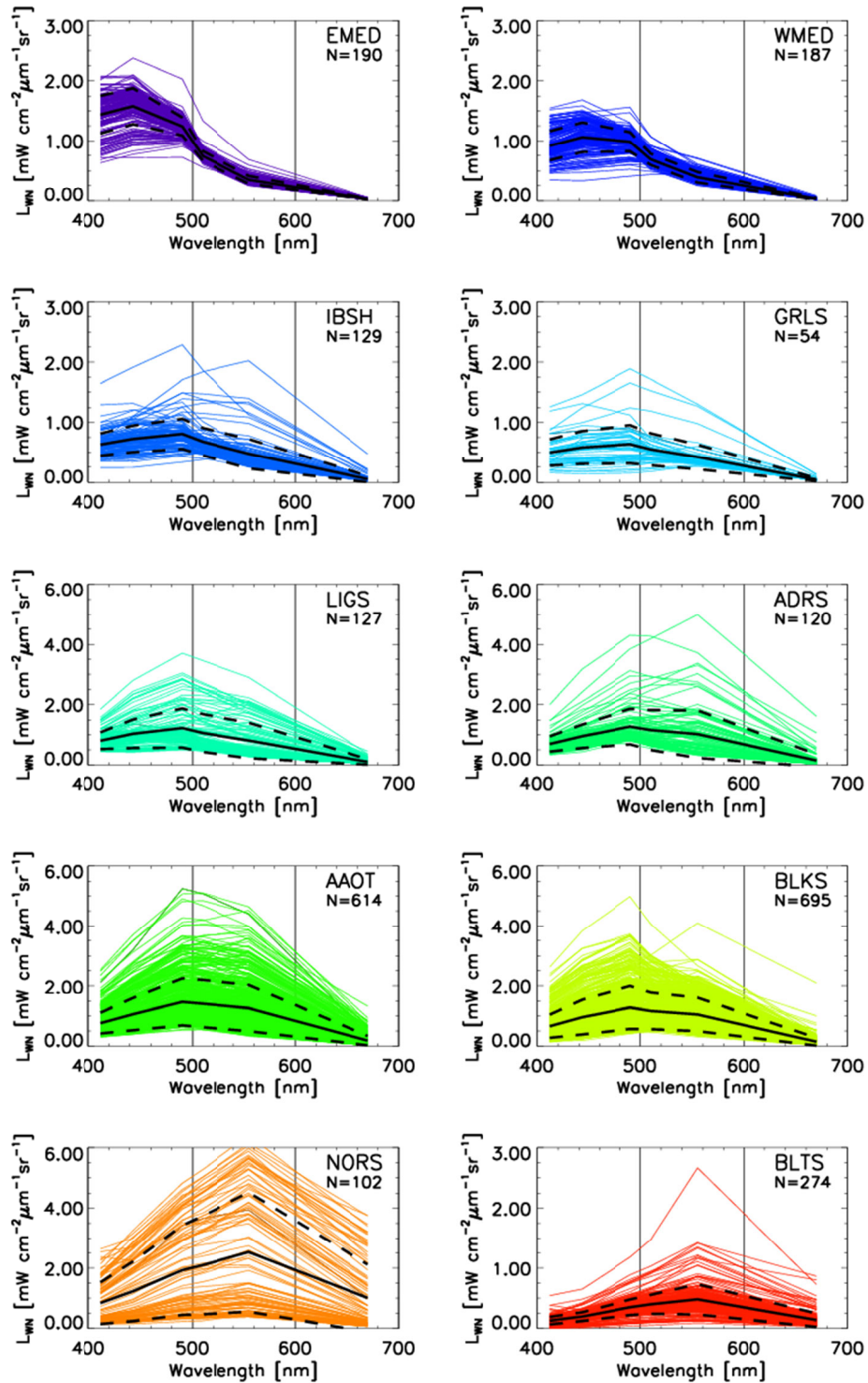
658 Table 4. Mean \pm standard deviations of quantities describing the bio-optical and hydrographic
659 characteristics of the CoASTS and BioMaP marine regions: the diffuse attenuation coefficient K_d
660 at 490 nm; the seawater absorption coefficient (excluding pure water contribution) a determined
661 from discrete sample analysis at 490 nm; the backscattering coefficient (including pure water
662 contribution) b_b at 488 nm; the concentrations of total chlorophyll-*a* *Chla* and suspended
663 particulate matter *SPM*; and finally the salinity S_w .

664

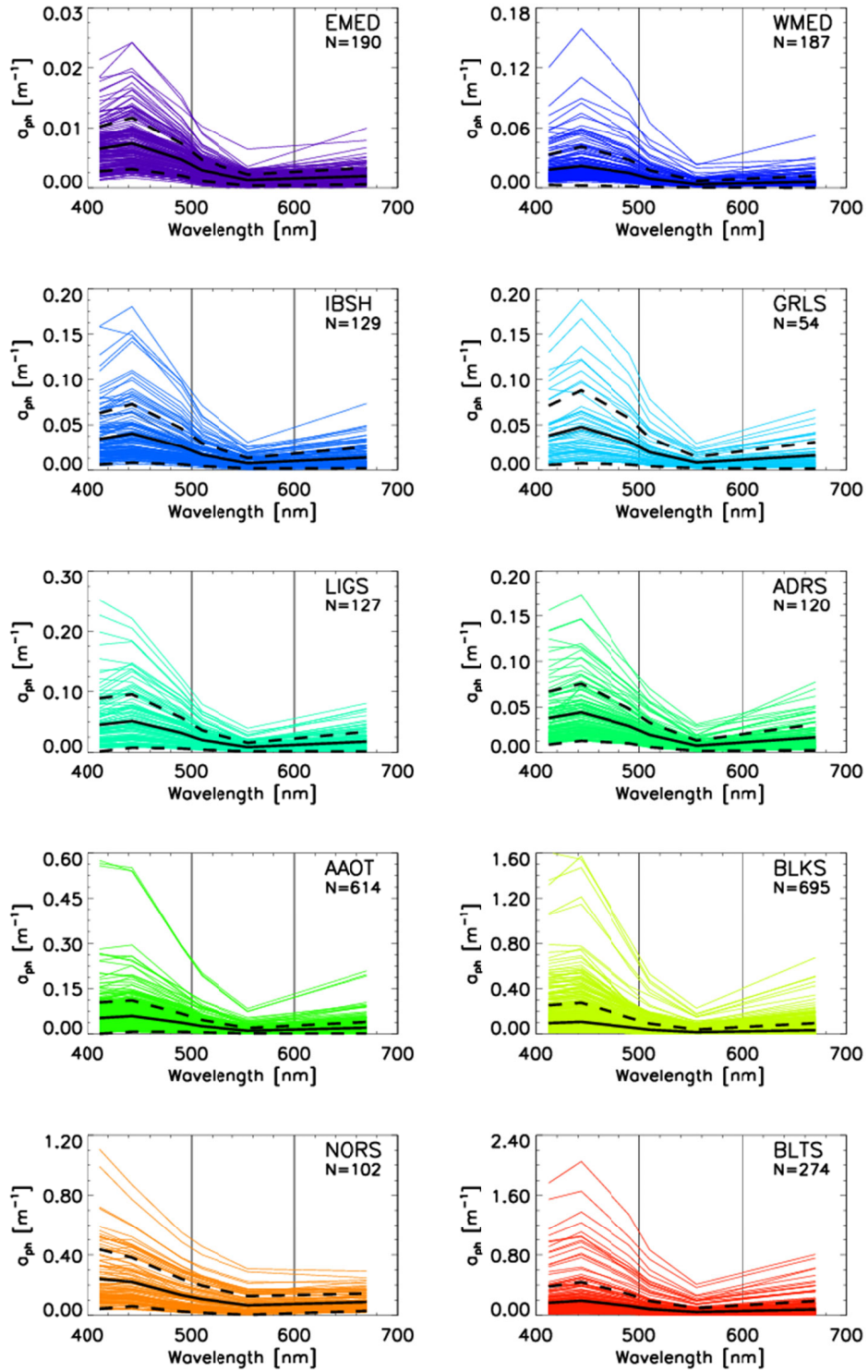
Region	$K_d(490)[\text{m}^{-1}]$	$a(490) [\text{m}^{-1}]$	$b_b(488)[\text{m}^{-1}]$	<i>Chla</i> $[\mu\text{g l}^{-1}]$	<i>SPM</i> $[\text{mg l}^{-1}]$	$S_w[\text{‰}]$
EMED	0.037 \pm 0.022	0.031 \pm 0.012	0.0026 \pm 0.0007	0.09 \pm 0.08	0.27 \pm 0.45	38.6 \pm 0.7
WMED	0.046 \pm 0.025	0.040 \pm 0.019	0.0032 \pm 0.0009	0.30 \pm 0.37	0.30 \pm 0.22	37.8 \pm 0.4
IBSH	0.084 \pm 0.049	0.073 \pm 0.033	0.0040 \pm 0.0023	0.81 \pm 0.83	0.53 \pm 0.39	36.0 \pm 0.2
GRLS	0.097 \pm 0.062	0.082 \pm 0.032	0.0039 \pm 0.0021	0.94 \pm 1.04	0.64 \pm 0.28	34.0 \pm 1.6
LIGS	0.110 \pm 0.079	0.079 \pm 0.045	0.0078 \pm 0.0067	0.93 \pm 0.85	0.71 \pm 0.57	37.7 \pm 1.0
ADRS	0.141 \pm 0.125	0.085 \pm 0.059	0.0090 \pm 0.0067	1.25 \pm 1.32	1.14 \pm 1.45	35.6 \pm 2.3
AAOT	0.176 \pm 0.102	0.099 \pm 0.053	0.0121 \pm 0.0073	1.28 \pm 1.13	1.25 \pm 0.76	34.9 \pm 2.3
BLKS	0.219 \pm 0.254	0.131 \pm 0.130	0.0093 \pm 0.0066	1.62 \pm 3.13	1.17 \pm 1.24	16.6 \pm 1.8
NORS	0.876 \pm 0.864	0.377 \pm 0.346	0.0197 \pm 0.0160	4.23 \pm 2.27	9.96 \pm 12.52	33.7 \pm 1.4
BLTS	0.494 \pm 0.409	0.308 \pm 0.269	0.0107 \pm 0.0084	4.99 \pm 8.04	1.53 \pm 1.71	6.2 \pm 1.4

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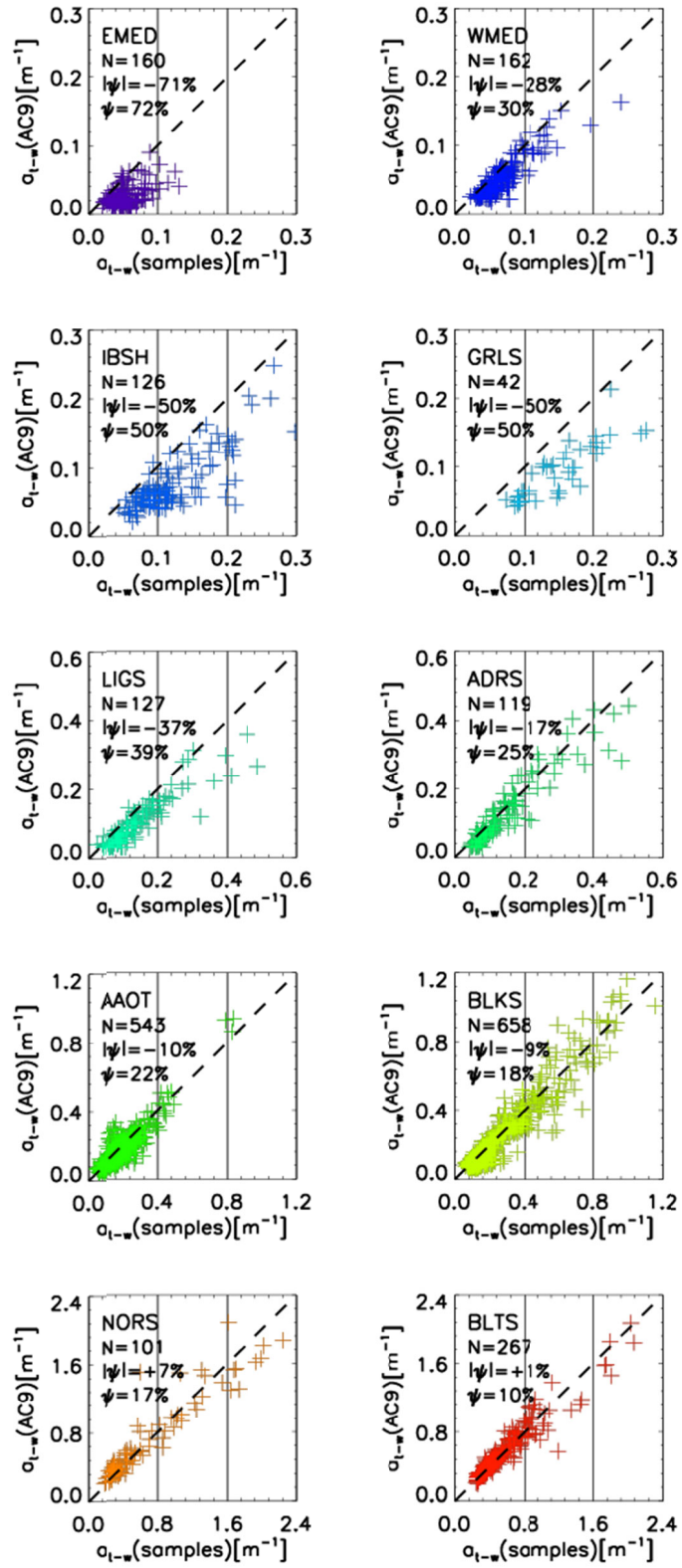


667
 668 Figure 3. Spectra of $L_{WN}(\lambda)$ for the CoASTS and BioMaP geographic regions (see Table 1 for
 669 acronyms). N indicates the number of spectra. The continuous black lines indicate mean values
 670 while the dashed lines indicate ± 1 standard deviation. For convenience, the spectra are plotted in
 671 units of $\text{mW cm}^{-2} \mu\text{m}^{-1} \text{sr}^{-1}$.



672

673 Figure 4. Spectra of $a_{ph}(\lambda)$ for the CoASTS and BioMaP marine regions. N indicates the number
 674 of spectra. The continuous black lines indicate mean values while the dashed lines indicate ± 1
 675 standard deviation.



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677

678 Figure 5. Scatter plot of AC9 derived $a_{t-w}(AC9)$ and laboratory measurements performed on
 679 water samples $a_{t-w}(\text{samples})$, of the water absorption coefficient (water excluded) determined at
 680 the 443 nm center-wavelength for the diverse CoASTS and BioMaP marine regions. N indicates
 681 the number of samples while $|\psi|$ and ψ indicate the mean of absolute (unsigned) percent
 682 differences and the mean of (signed) percent differences, respectively.

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 684

685 Table 5. Spectral values of $Q_n(\lambda)$ in units of sr at the 412, 443, 490, 510, 555 and 670 nm center-
 686 wavelengths for the CoASTS and BioMaP marine regions, determined from in-water radiometric
 687 profiles performed with cloud cover $C_C \leq 1/4$.

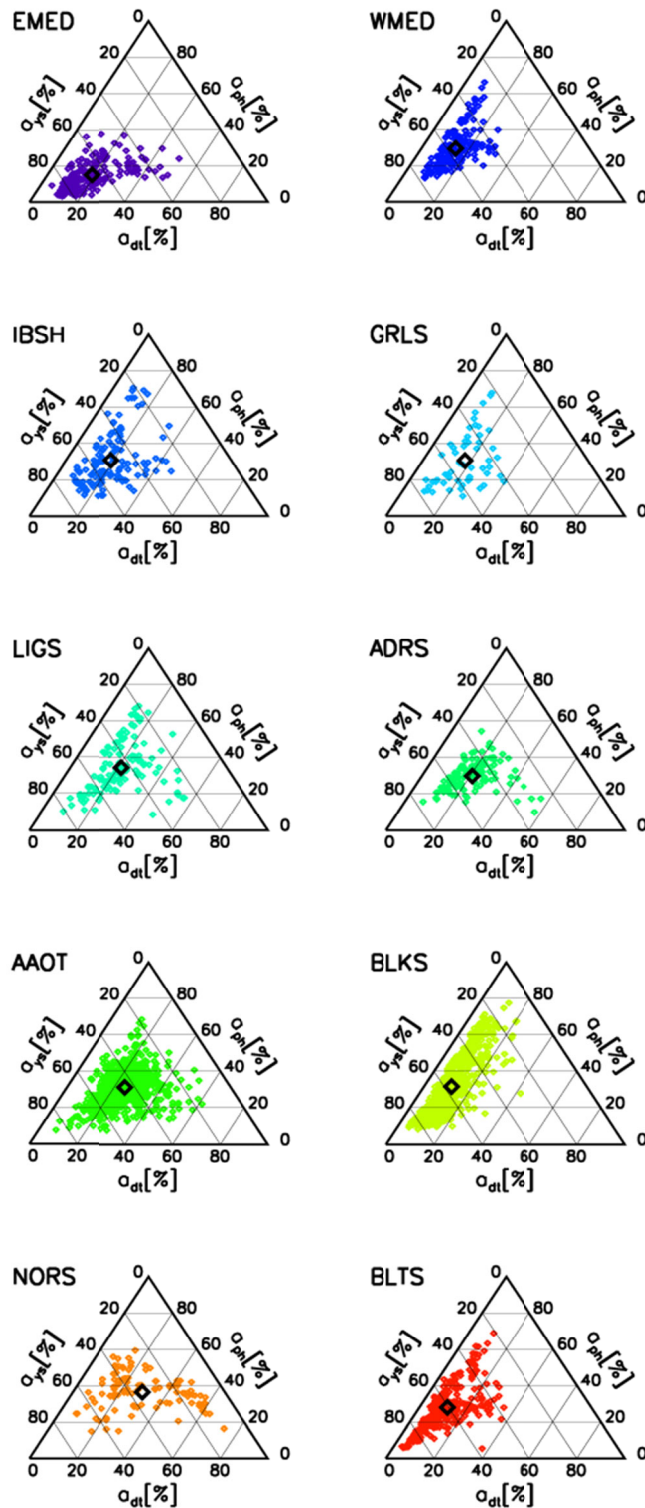
Region	412	443	490	510	555	670
EMED (N=127)	3.89±0.33	3.90±0.36	3.88±0.42	3.87±0.45	3.84±0.54	4.90±1.12
WMED (N=100)	4.08±0.36	4.14±0.41	4.20±0.46	4.21±0.48	4.19±0.52	4.96±0.76
IBSH (N=87)	4.18±0.37	4.22±0.38	4.26±0.43	4.26±0.45	4.24±0.51	4.58±0.59
GRLS (N=11)	3.97±0.33	4.08±0.37	4.14±0.38	4.12±0.37	4.00±0.34	4.18±0.38
LIGS (N=53)	4.52±0.40	4.54±0.36	4.57±0.36	4.59±0.38	4.66±0.44	5.14±0.58
ADRS (N=71)	4.47±0.65	4.39±0.57	4.33±0.54	4.34±0.55	4.40±0.62	4.98±0.95
AAOT (N=372)	4.56±0.56	4.43±0.51	4.33±0.49	4.33±0.50	4.41±0.58	5.02±0.84
BLKS (N=401)	4.51±0.54	4.49±0.57	4.47±0.59	4.47±0.59	4.47±0.59	5.06±0.80
NORS (N=27)	4.70±0.60	4.71±0.57	4.69±0.54	4.67±0.53	4.60±0.50	4.90±0.48
BLTS (N=87)	4.93±0.69	5.09±0.74	5.18±0.78	5.16±0.76	4.99±0.66	5.20±0.86

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690 Figure 6 shows trilinear (ternary) plots of the absorption coefficients $a_{ys}(443)$, $a_{dt}(443)$ and
 691 $a_{ph}(443)$, expressed in percent of the total absorption (*i.e.*, with respect to $a_{ys}(443)+a_{dt}(443)+$
 692 $a_{ph}(443)$), displayed with values increasing in the counter-clockwise direction (Harris 1999).
 693 These results exhibit very few cases characterized by dominance of absorption by particles with
 694 a_{ph} and a_{dt} values close to the upper and lower right apexes, respectively. Conversely, most of the
 695 cases indicate dominance of absorption by coloured dissolved organic matter: see the a_{ys} values
 696 near the lower left apex). This is particularly evident for the oligotrophic waters of the Eastern
 697 Mediterranean Sea (EMED), and by the patterns characterizing the oligotrophic-mesotrophic
 698 waters of the Western Mediterranean Sea (WMED), the optically complex water of the Black
 699 Sea (BLKS) and the highly absorbing waters of the Baltic Sea (BLTS).

700 The specific results shown for the Mediterranean Sea (*i.e.*, EMED and WMED), which
 701 may suggest inconsistency with the definition of Case-1 waters (IOCCG 2000), are supported by
 702 an independent study from Pérez et al. (2016).

703



704
 705 Figure 6. Trilinear (ternary) plots of the absorption coefficients a_{ys} , a_{dt} and a_{ph} expressed in
 706 percent of the total absorption (*i.e.*, with respect to $a_{ys}+a_{dt}+a_{ph}$) at the 443 nm center-wavelength.
 707 The empty black square indicates the mean of the plotted values.

708 Parameters determined from the exponential fit versus wavelength of $a_{dt}(\lambda)$ and $a_{ys}(\lambda)$, and
 709 the power law fit of $b_b(\lambda)$ versus wavelength, are provided in Tables 6-8. Specifically, the
 710 spectral values of $a_{dt}(\lambda)$ and $a_{ys}(\lambda)$ were fitted within the 412-665 nm spectral interval using

$$711 \quad a_{dt}(\lambda) = A_{dt} \exp(-S_{dt}(\lambda - 412)) + B_{dt} \quad (8)$$

713 and

$$714 \quad a_{ys}(\lambda) = A_{ys} \exp(-S_{ys}(\lambda - 412)) + B_{ys}, \quad (9)$$

715 where A_{dt} and A_{ys} indicate the absorption coefficients fitted at 412 nm, S_{dt} and S_{ys} the slope of
 716 the exponential function, and, B_{dt} and B_{ys} account for the background.

717 Conversely, the spectral values of $b_b(\lambda)$ at the center-wavelengths $\lambda=442, 488, 510, 550$ and
 718 620 nm (excluding 676 or [671](#) nm due to potential perturbations by fluorescence), were fitted
 719 using

$$720 \quad b_b(\lambda) = A_b (\lambda/442)^{-S_b}, \quad (10)$$

721 where A_b indicates the backscattering coefficient at 442 nm and S_b the slope of the power law
 722 function.

723 Table 6 shows mean values of the slope S_{dt} varying from 0.009 nm⁻¹ for the Eastern
 724 Mediterranean Sea (EMED) up to 0.013 for the North Sea (NORS). Values of the bias B_{dt}
 725 naturally increase with A_{dt} : the largest value of $B_{dt}=0.067$ m⁻¹ is observed for the North Sea
 726 (NORS) that also exhibits the highest value of $A_{dt}=0.288$ m⁻¹. Residuals R_{dt} , which also
 727 increase with A_{dt} , are quite minor suggesting a general good performance of the exponential
 728 fitting function.

729 Table 7 shows mean values S_{ys} varying from 0.012 nm⁻¹ for the Eastern Mediterranean Sea
 730 (EMED) up to 0.019 nm⁻¹ for the Baltic Sea (BLTS). The systematic negative biases B_{ys} across
 731 all marine regions are likely explained by the choice of zeroing the original spectra of absorption
 732 coefficients using values averaged in the 670-680 nm spectral interval. High residuals B_{ys} of
 733 0.029 m⁻¹ are observed for the Baltic Sea (BLTS). This is explained by a decreased performance
 734 of Eq. 9 when fitting spectra of absorption coefficients exhibiting values approaching or
 735 exceeding 1 m⁻¹ at 412 nm. Still, all residuals B_{ys} expressed in percent of A_{ys} vary between 0.3
 736 and 0.5%, except for the East Mediterranean Sea (EMED) showing a value of 0.9%.

737 As expected, also the values of S_b largely vary across the CoASTS and BiOMaP marine
 738 regions: in particular they exhibit values of 2.97 μm^{-1} for the East Mediterranean Sea (EMED),
 739 2.06 μm^{-1} for the Iberian Shelf (IBSH) and 0.74 μm^{-1} for the North Sea (NORS). This is likely
 740 explained by an increase of the average particles size when going from the oligotrophic East
 741 Mediterranean Sea to the eutrophic and more sediment loaded North Sea.

742 Figure 7 and 8 shows the distribution of *Chla* and *SPM* across the CoASTS and BiOMaP
 743 marine regions. The very low concentrations characterizing the oligotrophic waters of the
 744 Eastern Mediterranean Sea (EMED) exhibiting mean *Chla* values of 0.09 $\mu\text{g l}^{-1}$ and mean *SPM*
 745 values of 0.27 mg l⁻¹, are remarkable. Conversely, *Chla* exhibits mean values in the range of 4-5
 746 $\mu\text{g l}^{-1}$ for both the North Sea (NORS) and Baltic Sea (BLKS), while for the same marine regions
 747 *SPM* shows mean values of 9.96 and 1.53 mg l⁻¹, respectively. A log-normal distribution of both
 748 *Chla* and *SPM* is generally confirmed for the CoASTS and BiOMaP data.

753 Figure 9 displays the scatter plots of $b_{bp}(488)/b_p(488)$ versus $Chla$, where $b_p(488)$ is
754 determined by the difference between $c_{t-w}(488)$ and $a_{t-w}(488)$, while $b_{bp}(488)$ is determined from
755 $b_b(488)$ by subtracting the scattering coefficient of water $b_w(488)$ from Morel (1974). Results are
756 consistent with those shown by Twardowski et al. (2001) for a variety of experimental data, with
757 $b_{bp}(488)/b_p(488)$ typically varying between 0.003 and 0.025. Exception are some very low
758 values of $b_{bp}(488)/b_p(488)$ for EMED data likely explained by large measurement uncertainties.
759 Coherent with published results is also the generally higher value and higher scatter of
760 $b_{bp}(488)/b_p(488)$ in correspondence of low $Chla$ concentrations.

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762

763 Table 6. Parameters A_{dt} , S_{dt} and B_{dt} of the exponential fitting function (see Eq. 8) applied to the
764 values of $a_{dt}(\lambda)$. The quantity R_{dt} indicates the spectral average of absolute differences (*i.e.*,
765 residuals) between actual and fitted values.

766

Region	$A_{dt} [m^{-1}]$	$S_{dt} [nm^{-1}]$	$B_{dt} [m^{-1}]$	$R_{dt} [m^{-1}]$
EMED (N=190)	0.010±0.007	0.009±0.002	0.002±0.001	0.0000
WMED (N=186)	0.009±0.004	0.012±0.001	0.003±0.001	0.0000
IBSH (N=129)	0.024±0.022	0.011±0.001	0.006±0.005	0.0001
GRLS (N=54)	0.024±0.014	0.012±0.002	0.007±0.004	0.0000
LIGS (N=126)	0.032±0.026	0.011±0.002	0.007±0.004	0.0001
ADRS (N=120)	0.042±0.057	0.012±0.001	0.009±0.011	0.0000
AAOT (N=614)	0.048±0.031	0.012±0.001	0.009±0.005	0.0000
BLKS (N=692)	0.034±0.057	0.011±0.002	0.005±0.008	0.0001
NORS (N=102)	0.288±0.377	0.013±0.001	0.067±0.094	0.0005
BLTS (N=274)	0.095±0.125	0.011±0.002	0.011±0.017	0.0003

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769 Table 7. Parameters A_{ys} , S_{ys} and B_{ys} of the exponential fitting function (see Eq. 9) applied to the
770 values of $a_{ys}(\lambda)$. The quantity R_{ys} indicates the spectral average of absolute differences (*i.e.*,
771 residuals) between actual and fitted values.

772

Region	$A_{ys} [m^{-1}]$	$S_{ys} [nm^{-1}]$	$B_{ys} [m^{-1}]$	$R_{ys} [m^{-1}]$
EMED (N=182)	0.056±0.025	0.012±0.004	-0.005±0.007	0.0004
WMED (N=183)	0.059±0.019	0.013±0.003	-0.002±0.002	0.0002
IBSH (N=129)	0.093±0.036	0.014±0.003	-0.004±0.005	0.0004
GRLS (N=54)	0.107±0.027	0.014±0.003	-0.004±0.003	0.0003
LIGS (N=126)	0.091±0.052	0.014±0.004	-0.004±0.004	0.0004
ADRS (N=120)	0.114±0.058	0.016±0.002	-0.002±0.002	0.0003
AAOT (N=592)	0.132±0.059	0.017±0.004	-0.003±0.005	0.0003
BLKS (N=693)	0.205±0.122	0.017±0.002	-0.004±0.003	0.0005
NORS (N=102)	0.280±0.094	0.017±0.002	-0.004±0.002	0.0007
BLTS (N=274)	0.606±0.330	0.019±0.001	-0.004±0.003	0.0029

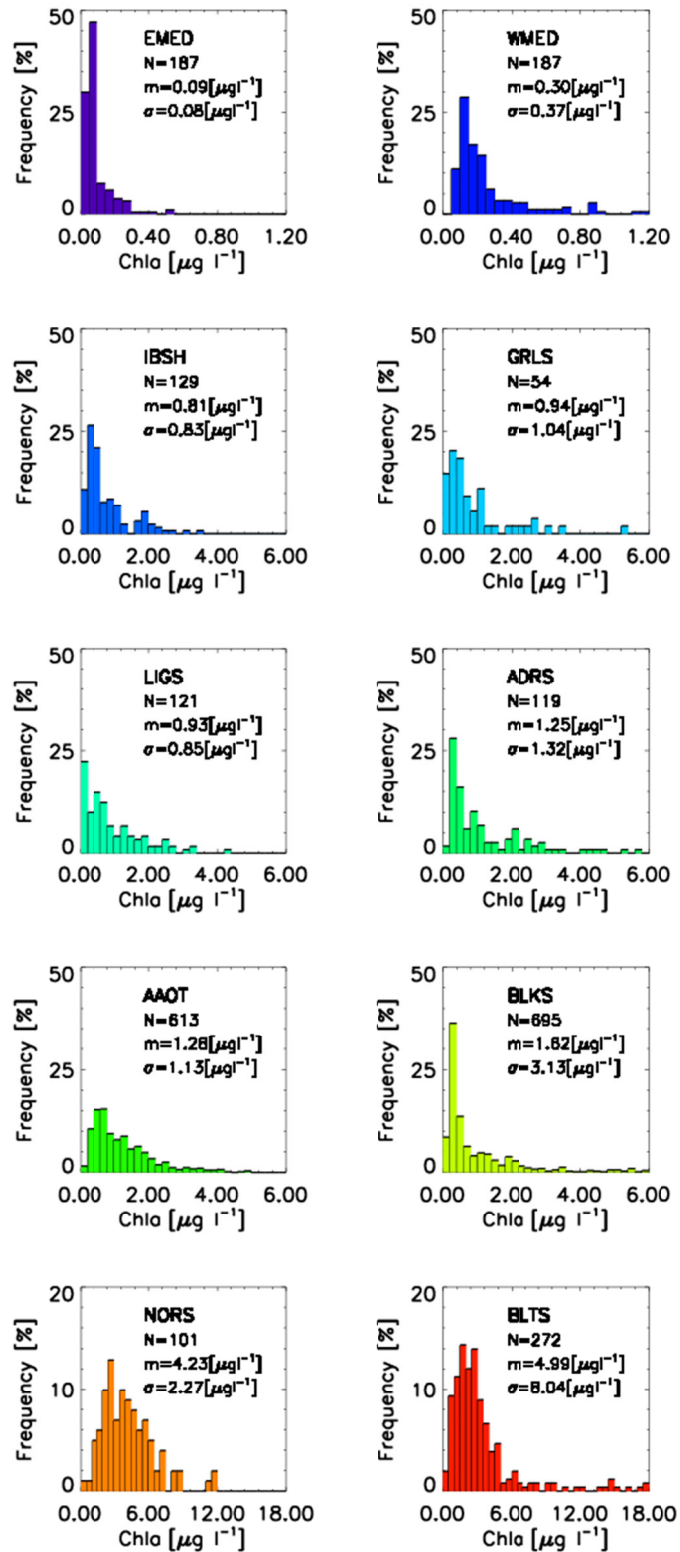
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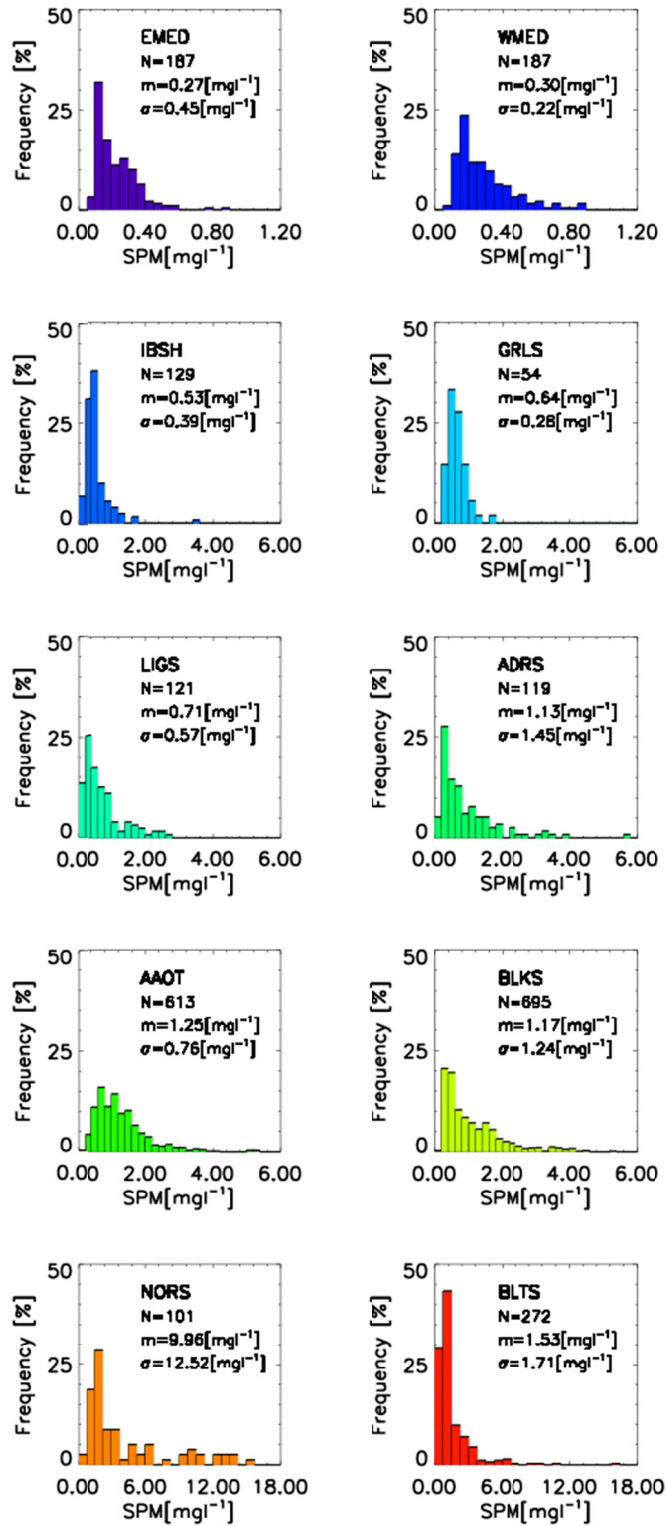
775 Table 8. Parameters A_b and S_b of the power law fitting function (see Eq. 10) applied to the values
 776 of $b_b(\lambda)$ at $\lambda= 443, 488, 510, 555$ and 620 nm for the CoASTS and BioMaP marine regions. The
 777 quantity R_b indicates the spectral average of absolute differences (*i.e.*, residuals) between actual
 778 and fitted data.

Region	A_b [m^{-1}]	S_b [μm^{-1}]	R_b [m^{-1}]
EMED (N=184)	0.0034±0.0008	2.97±0.56	0.0001
WMED (N=186)	0.0041±0.0009	2.54±0.42	0.0001
IBSH (N=127)	0.0051±0.0025	2.06±0.55	0.0002
GRLS (N=52)	0.0048±0.0024	2.25±0.33	0.0001
LIGS (N=126)	0.0091±0.0072	1.83±0.64	0.0002
ADRS (N=111)	0.0103±0.0071	1.74±0.57	0.0002
AAOT (N=479)	0.0136±0.0078	1.35±0.42	0.0004
BLKS (N=534)	0.0126±0.0077	1.99±0.53	0.0006
NORS (N=57)	0.0207±0.0157	0.74±0.38	0.0005
BLTS (N=256)	0.0118±0.0082	1.15±0.49	0.0003

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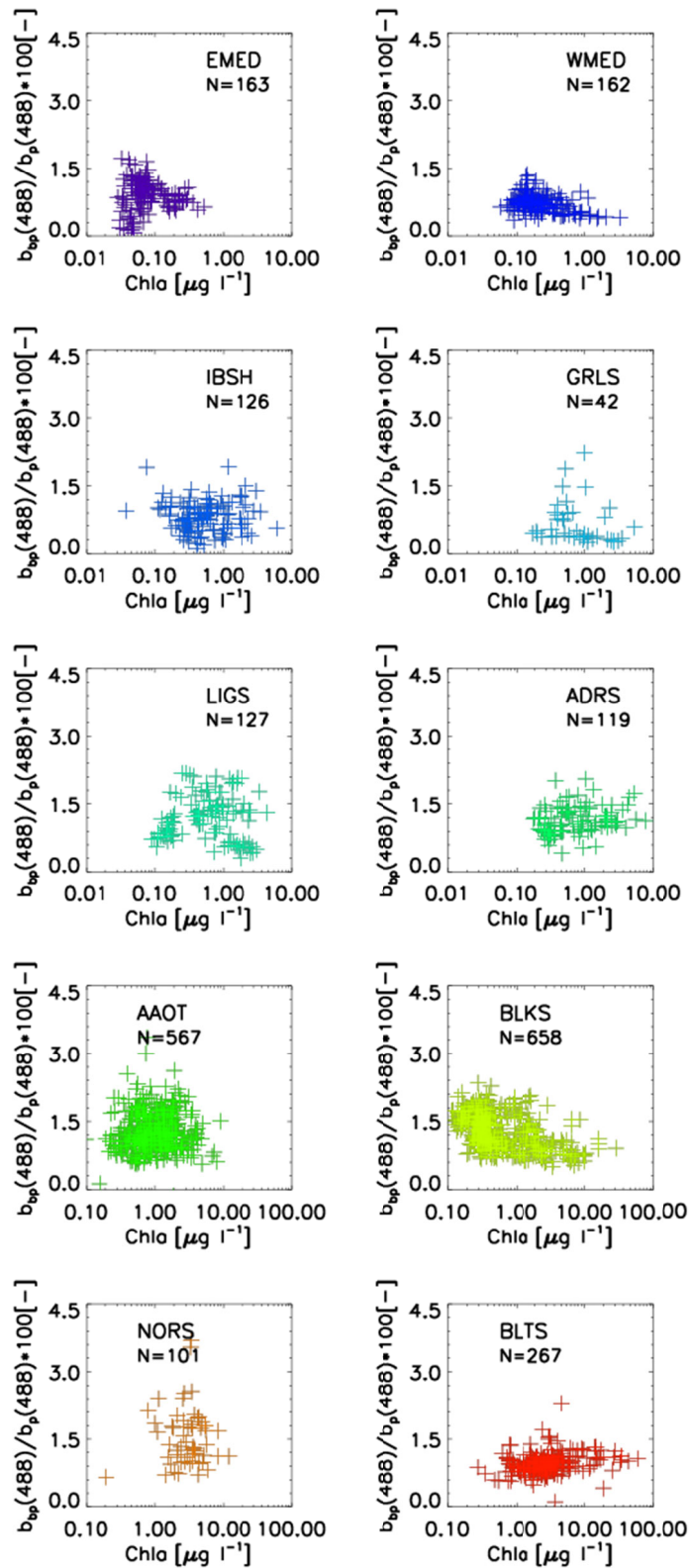


781
 782 Figure 7. Frequency distribution of *Chla* across the CoASTS and BioMaP marine regions. N
 783 indicates the number of stations, *m* the mean values and σ the standard deviation.



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Figure 8. Frequency distribution of *SPM* across the CoASTS and BioMaP marine regions. N indicates the number of stations, *m* the mean values and σ the standard deviation.



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Figure 9. Scatter plot $b_{bp}(488)/b_p(488)$ and $Chla$ for the diverse CoASTS and BioMaP marine regions. N indicates the number of samples.

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Table 9. Mean values of the *Chla* specific absorption coefficient a_{ph}^* at 443 nm.

Region	$a_{ph}^*(443)$ [m^2mg^{-1}]
EMED (N=210)	0.090±0.029
WMED (N=190)	0.083±0.018
IBSH (N=129)	0.062±0.050
GRLS (N=54)	0.063±0.018
LIGS (N=121)	0.065±0.021
ADRS (N=119)	0.053±0.034
AAOT (N=613)	0.052±0.022
BLKS (N=695)	0.084±0.046
NORS (N=101)	0.061±0.108
BLTS (N=272)	0.047±0.014

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Table 9 provides the mean specific absorption coefficients $a_{ph}^*(443)$ determined by the ratio of $a_{ph}(443)/Chla$ across the various CoASTS and BioMaP marine regions. These mean values of $a_{ph}^*(443)$ vary from 0.047 m^2mg^{-1} in the Baltic Sea (BLT) to 0.090 m^2mg^{-1} in the Eastern Mediterranean Sea. It is warned that these latter values could be challenged by increased relative uncertainties in the determination of both $a_{ph}(443)$ and *Chla*.

6. Summary and conclusions

The CoASTS and BiOMaP measurement programs led by the JRC Marine Optical Laboratory benefitting of the collaboration of a number of European institutions and various funding programs, were conceived to support satellite ocean color applications. Between 1995 and 2022 the two programs produced time-series at the AAOT site in the northern Adriatic Sea and geographically distributed bio-optical measurements across the major European Seas. The measurements delivered by the two programs beyond December 1998 include identical quantities and are characterized by standardization of measurement methods, instruments, data processing and quality assurance/control schemes.

This work introduced the CoASTS-BiOMaP data set comprising the near surface data products from the CoASTS and BiOMaP measurement programs of major relevance for satellite ocean color validation activities and bio-optical modelling.

7. Author contributions

Both authors, Giuseppe Zibordi and Jean-François Berthon, who implemented and co-led the CoASTS and BiOMaP programs, contributed to the generation of the data set and to the writing of the manuscript. Giuseppe Zibordi was a JRC Scientific Officer since the conception and up to the end of the CoASTS and BiOMaP programs.

8. Competing interests

Both authors declare no competing interest.

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825 **9. Data availability**

826 Interested researchers can download the CoASTS-BiOMaP data set at <https://doi.pangaea.de/10.1594/PANGAEA.971945> (Zibordi and Berthon, 2024). The original field
827 measurements leading to the creation of this data set are currently not publicly available.
828 However, they can be obtained from the authors upon a reasonable request.
829

830

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834 Linde, Lukasz Jankowski, Lyudmila Kamburska, Davide D'Alimonte, Marco Talone, Pietro
835 Sciuto, Iliaria Cazzaniga, Jean Verdebout, Elisabetta Canuti, Alessandro Marchetti, Violeta
836 Slabakova, Natalia Slabakova, Carolina Sa', Simone Colella, Gianluca Volpe, Seppo Kaitala,
837 Jukka Seppala, Aleksandra Mazur.

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839 [submitted manuscript](#).

840

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846 Administration (NASA), the European Space Agency (ESA), the Romanian Space Agency
847 (ROSA), the Institute of Oceanology of the Bulgarian Academy of Sciences, the Institute of
848 Oceanology of the Polish Academy of Sciences, the Finnish Environment Institute, the Italian
849 National Research Council, the Portuguese Hydrographic Institute, the Italian Hydrographic
850 Institute, the Royal Belgian Institute of Natural Sciences, the Hellenic Centre for Marine
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858 **12. References**

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1002 **Appendix A: Acronyms**

1003	AAOT	Acqua Alta Oceanographic Tower
1004	ADRS	Adriatic Sea
1005	AERONET-OC	Ocean Color component of the Aerosol Robotic Network
1006	BiOMaP	Bio-Optical mapping of Marine Properties
1007	BLKS	Black Sea
1008	BLTS	Baltic Sea
1009	CDOM	Colored Dissolved Organic Matter
1010	CoASTS	Coastal-Atmosphere and Sea Time-Series
1011	CTD	Conductivity, temperature and depth
1012	EMED	Eastern Mediterranean Sea
1013	GRLS	Greenland Sea
1014	HPLC	High-Pressure Liquid Chromatography
1015	IBS	Iberian Shelf
1016	LIGS	Ligurian Sea
1017	JRC	Joint Research Center
1018	NASA	National Aeronautics and Space Administration
1019	NIST	National Institute of Standards and Technology
1020	NORS	North Sea
1021	NPL	National Physical Laboratory
1022	SeaWiFS	Sea-viewing Wide Field-of-view Sensor
1023	WMED	Western Mediterranean Sea
1024	WiSPER	Wire-Stabilized Profiling Environmental Radiometer
1025		