Optical Classification of Lower Amazon Waters Based on In Situ Data and Sentinel-3 Ocean and Land Color Instrument Imagery

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1. Introduction

The flow of the water, starting with rainfall and headwater streams, carries particulate and dissolved materials from land to ocean, driving the transport and cycling of a vast suite of biogeochemical components. The physical-biogeochemical properties of the water carried by rivers are in constant change while traveling downstream to the coastal ocean due to interactions among the landscape, wetlands, weather and climate forcing, and in situ processes [1–3]. In situ sampling captures the properties at a specific point location and time. But it represents a considerable challenge for an effective and representative monitoring at regional scales, considering the large distance between sampling stations and the usually inconstant nature of field sampling, particularly in remote regions [4,5]. The intrinsic transience of river properties is more evident when using remotely sensed satellite observations, which have the advantage of capturing the dynamic and heterogeneous nature of surface waters [6]. In situ sampling obviously remains crucial since it is a required step for validation and robust analysis of remote sensing observations.

The measured upwelling radiance \( L_u \) from a water body is an apparent optical property and depends on both the physical properties of the water and its constituents.
and the light field conditions [7]. The $L_d$ reflects the inherent optical properties (IOPs), like absorption and scattering, of its optically significant constituents and are diagnostic of the dynamics of phytoplankton (phy), non-algal particles (NAP), and colored dissolved organic matter (CDOM) [8,9]. Over the years, empirical relationships between absorption coefficients ($a$) and biogeochemical parameters, such as $a_{\text{phy}}$ and chlorophyll-$a$ (chl $a$) [10]; and $a_{\text{nap}}$ and suspended particulate matter (SPM) [11], have been documented for ocean and coastal waters. Moving to inland waters, more studies are needed to confirm if these relationships are still reliable [12–14]. In the same way, subproducts of absorption coefficient measurements, such as the slope of $a_{\text{CDOM}}$ and $a_{\text{nap}}$ ($S_{\text{CDOM}}$ and $S_{\text{nap}}$, respectively), also require more studies on their applicability to retrieve qualitative information about the dissolved material and particles present in an optically complex water body [14,15]. Indeed, the slope $S_{\text{CDOM}}$ assessed over different wavelength intervals has been shown to be a relevant indicator of dissolved organic matter molecular weight [11,16,17], while $S_{\text{nap}}$ can provide information on the relative contribution of organic and mineral particles in the absorption at coastal waters [11].

Depending on the concentration of biogeochemical constituents, the color of Amazon River waters has been categorized as white, black, or clearwater [18]. White waters are classically associated with rivers characterized by high concentration of sediments, as the Amazon, Madeira, and Solimões rivers; black waters are related to rivers with high concentrations of CDOM, such as the Negro River; and clearwater, as observed in the Tapajós and Xingu rivers in the Lower Amazon region, are associated with high transparency and high rates of primary production. This type of characterization of freshwater optical water types (OWTs) presented by Sioli [18], however, is archetypal and does not consider the seasonality that changes river discharge patterns and, consequently, the water biogeochemical and bio-optical properties.

A synthetic representation of water optical characteristics and variability can be performed through the definition of OWTs which has been a recurrent research topic in water color remote sensing since the pioneering work by Jerlov in marine waters [19]. The increasingly large in situ radiometric, bio-optical, and biogeochemical database over the years, especially in oceanic and coastal regions, provides robust classifications [12,20–23]. In inland waters, the same data collection is more substantial in the northern hemisphere supporting more reliable classifications [20,21]. In tropical regions, especially in the southern hemisphere, efforts have been made in the Amazon region [13,20,24], but much remains to be done in terms of classifying tropical water types.

In a global analysis, Spyrakos et al. [20] have classified Amazon River waters within a single OWT, characterized by high reflectance across a wide interval of the spectrum, in association with a strong SPM dominance. However, the Amazon optical data utilized to develop the aforementioned classification scheme were limited to a collection of spectra and bio-optical constituents sampled at the Great Lake of Curuai floodplain, while no information about seasonal variability was presented for the region. Similarly, Silva et al. [24] performed a classification on overall Brazilian waters, including Amazonian lakes (e.g., Great Lake Curuai floodplain) and tributary rivers (e.g., Tapajós River), but still did not consider the seasonal variability. Seasonal variability of Amazon OWTs was considered in a study presented by Martinez et al. [13]. The authors proposed eight classes of different water types based on in situ samples acquired at Óbidos, in the state of Pará, which is roughly 900 km from the river mouth and marks the downstream boundary for tidal influence on the fluvial system [25].

The Ocean and Land Color Instrument imagery (OLCI) aboard the Sentinel 3 mission was launched in 2016, following the end of the Medium Resolution Imaging Spectrometer (MERIS) mission in 2012. The monitoring of inland waters, including rivers and lakes, are an important task of the OLCI mission and their spectral bands were developed with this concern [26]. Since its launch, OLCI has been used to monitor different aspects of inland waters studies, like river flow [27], inland water quality monitoring [28,29], and atmospheric correction over inland waters [30].
In this context, the objectives of this study for the lower Amazon River and its major tributaries are to: (i) identify optical water types; (ii) characterize each typology based on optically significant constituents and light absorption properties; and (iii) assess the OWT spatial and seasonal distributions. We hypothesized that mainstem and tributary waters would be bio-optically distinct and that seasonally variable hydrology would exert an important control on OWT classifications across both mainstem and tributary waters. The identification of OWTs is carried by a cluster analysis of in situ remote sensing reflectance ($R_{rs}$) spectra [20,22]. Then, the cluster specific spectra can be applied to remote sensing imagery, providing information on the spatial distribution of the identified OWTs [31]. Bio-optical properties can be used to characterize the reflectance spectra clustered in an OWT, providing qualitative information on the bio-optical characteristics associated with $R_{rs}$ based optical typology.

OWT classification can further provide a framework constraining the development of a suitable inversion algorithm for specific bio-optical environments, as already illustrated from several previous studies that have focused on the development of class specific inversion algorithms in coastal [21,22] and inland waters [21].

2. Materials and Methods

2.1. Study Area

The Lower Amazon study region comprises the area along a 900 km transect between the upstream boundary at Óbidos (01°55.141′S, 55°31.543′W) and the Amazon River mouth near the city of Macapã (Figure 1A). The wet season in the study area starts in November, with the peak in March (Figure 1B). The rising water discharge season starts at the end of December and reaches the high-water level peak in May. In June, the water level starts to fall, and the minimum is observed in November [32,33]. According to previous studies in this region [34,35], the seasonal discharge months are defined as (Figure 1C): rising water (January–March); high water (April–June); falling water (July–September); and low water (October–December).

Samples were collected at stations along the Amazon mainstem, in a series of expeditions by the research vessel M/V Mirage, at Óbidos, below Santarem, Almeirim, and two primary channels near the geographical river mouth by Macapã. At Óbidos, the width of the Amazon River is 2.3 km, and, at Macapã, each channel has a width of 11 km. The width of the mainstem at the Lower Amazon is stable and has little variation along the seasonal discharge [36].

The location of Óbidos is crucial for monitoring the dynamics of the Amazon basin. It is the furthest downstream monitoring gauge of the Brazilian National Water Agency (ANA) where the mainstem has already received the contribution from large rivers, such as the Solimões, Negro, and Madeira, and from other smaller tributaries. Downstream of Óbidos, the Amazon River receives the inputs from two large tributaries: the Tapajós and Xingu rivers. The downstream boundary of our study was the north and south channels near Macapã, which are the last two well-constrained channels near the Amazon River mouth (00°05.400′ S, 51°03.200′ W and 00°09.415′ S, 50°37.353′ W, respectively). In addition to the mainstem, samples were collected in Tapajós, Xingu, Paru, Jari, and Jaraçu rivers, as well as at the Great Lake of Curuai floodplain (Figure 1A). These rivers are classified as clearwater rivers (Figure 1E,F), according to Sioli [18], generally presenting high rates of primary production when compared with turbid Amazon River waters (Figure 1D) [35]. Sampling of Tapajós and Xingu rivers was performed upstream of their confluence with the Amazon River at a “lake-like” portion of these rivers, where the river becomes similar to a lentic system. The sampling at clearwater (CW) rivers was always performed near the outfall portion of the river and, therefore, under the Amazon River influence. The magnitude of the Amazon River influence over CW rivers varies with the river size and its proximity to the Amazon River mouth where semi-diurnal flow reversals are observed [37,38]. The seasonality of river discharge also moderates the influence of the mainstem upon the tributaries.
The dataset used in this study consists of in situ measurements of optically significant constituents and inherent optical properties acquired from 2014–2017 during different discharge seasons. The sampling period of 2014–2017 was marked by exceptional climate events. The year of 2014 was characterized by a record flood in the Amazon Basin that started with a La Niña event in 2012 increasing the rainfall regime and was followed by a positive sea surface temperature anomaly in the Tropical Atlantic south of the equator, and in the Western Tropical Pacific [39–42]. During the rising water season of 2014, the rainfall within the south-western Amazon basin was 80–100% above usual records [41], with a consequently higher river discharge (Figure 1C).

Conversely, the year of 2015–2016 was marked by a severe drought caused by an El Niño period (Figure 1C), especially in the eastern Amazon basin [43]. The anomalously low water levels may have caused a disconnection of the Amazon River from some streams and floodplains potentially resulting in lower inputs of organic matter to the river’s mainstem. The disconnection of floodplains and streams from the mainstem caused by the river’s receded water level typical of drought periods has also been noted by other authors [40].

2.2. In Situ Data
2.2.1. Above Water Radiometry

Above water radiometry was measured with a portable hyperspectral spectroradiometer FieldSpec® (ASD Inc., Boulder, CO, USA) which collects radiance \( L(\lambda, \mu W \text{ m}^{-2} \text{ sr}^{-1}) \) in the range of 350 to 1100 nm (bandwidth 1 nm) and a field-of-view of 25°. The acquisition geometry followed Mobley [44] recommendations to avoid shadows and sunglint contamination in the measurements. The radiance measurements were conducted between 09:00 a.m. to 04:00 p.m. local time. Total water leaving radiance \( L_w \), sky radiance \( L_{sky} \), and the radiance from a white panel Spectralon reference \( L_g \) were consecutively measured 10 times. The latter parameter was used to estimate the downwelling irradiance \( E_d \) (Equation (1)):

\[
E_d(\lambda) = L_g(\lambda) f_c \pi, \quad (1)
\]

where \( f_c \) is a correction factor estimated in laboratory by the ratio of a standard Spectralon reference that remains in the laboratory by the Spectralon panel used during fieldwork.

The remote sensing reflectance \( R_{rs} \) is computed according to Equation (2):

\[
R_{rs} = \frac{L_u}{E_d} = \frac{L_u - \rho_{air-river} L_{sky}}{E_d}, \quad (2)
\]

where \( L_u \) is the upwelling radiance that reaches the sensor, and \( \rho_{air-river} \) is a sky glint correction coefficient at the air-sea interface.

There are several methods in the literature to correct the optical signal from sunglint interference. In the present study, we used the approach of Ruddick et al. [45], which is indicated for turbid to highly turbid waters. The \( \rho_{air-river} \) is a function of wind and cloud cover (Equations (3) and (4)).

\[
\frac{L_{sky}(\lambda = 750)}{E_D(\lambda = 750)} \geq 0.05 \rightarrow \rho_{air-river} = 0.0256, \quad (3)
\]

or

\[
\frac{L_{sky}(\lambda = 750)}{E_D(\lambda = 750)} < 0.05 \rightarrow \rho_{air-river} = 0.0256 + 0.00039W + 0.000034W^2, \quad (4)
\]

where \( W \) is the wind measured concurrently with the radiometric measurements.

The residual glint or white offset correction was not performed for the spectra dataset. According to Ruddick et al. [46], this correction is usually based on NIR spectrum (e.g., \( R_{rs}(780) \) and \( R_{rs}(720) \)) assuming that its shape is largely determined by pure water absorption. In very turbid waters, like in the Amazon River, the NIR assumption is not verified, and the NIR spectrum still has a high water signal.
2.2.2. Bio-Optical Measurements

Surface water samples for bio-optical measurements were acquired at all stations and filtered aboard the ship up to one hour after collection under gentle vacuum (<5 mm Hg), according to Mitchell et al.’s [47] protocol. Sub-samples were later processed in the laboratory for the determination of light absorption coefficients by total particulate matter \(a_p(\lambda) \text{ m}^{-1}\) and CDOM \(a_{\text{CDOM}}(\lambda) \text{ m}^{-1}\) using a dual-beam Shimadzu UV-2450 spectrophotometer, equipped with an integration sphere, within the interval of 250–850 nm. Aliquots of surface water samples were also used for the determination of chl \(a\) and SPM concentration. The filters for \(a_p\) and chl \(a\) were stored in a portable ultrafreezer during the cruise and stored in liquid nitrogen until posterior analysis in the laboratory using a spectrophotometer and a fluorimeter, respectively.

Light Absorption by Particulate Matter

Duplicate surface water samples were filtered through 25 mm Whatman GF/F glass fiber filters (0.7 µm nominal pore size). Water filtration was performed in order to obtain enough coloration of the filter pad, varying the volume between 10–125 mL depending on the river turbidity. The \(a_p\) spectra were calculated using the transmittance-reflectance (T-R) method [48]. The T-R method is preferentially recommended for particle-rich waters with the correction of the light backscattering by the particle sample with an integrating sphere. The slope of the absorption due to non-algal particles \(a_{\text{nap}}(\lambda)\), \(S_{\text{nap}}\), computed over the 400–800 nm spectral domain \(S_{\text{nap}400–800}\) [49,50], potentially indicates the relative contribution of mineral and organic particles in the water [11]. In this study, we considered \(a_p\) as representative of NAP because \(a_{\text{phy}}\) contribution was very low for the Amazon and relatively low for CW rivers (<2% and <20% of \(a_p\) (443), respectively) that was under Amazon River influence.

Light Absorption by Colored Dissolved Organic Matter

After removing the larger suspended particles used to estimate \(a_p\), the remaining filtered surface water samples were filtered again using 0.2 µm polycarbonate membranes (Whatman Nuclepore, 25 mm). Samples were stored in pre-combusted glass bottles (450 °C, 6 h) wrapped with aluminum foil and kept under refrigeration (4 °C) until further laboratory analysis. CDOM samples were exposed to room temperature before spectrophotometric analysis in order to avoid any bias due to the thermal difference between the samples and the reference Milli-Q water. \(a_{\text{CDOM}}(\lambda)\) was calculated from absorbance measurements as follows (Equation (5)):

\[
a_{\text{CDOM}}(\lambda) = 2.303 \frac{A(\lambda)}{L},
\]

where \(A(\lambda)\) is the absorbance of the filtered water sample at the specific wavelength \(\lambda\), and \(L\) is the optical pathway of the quartz cell in meters (here, 0.1 m).

As recommended by Babin et al. [11], a baseline correction was applied to each spectrum by subtracting the mean absorbance in the range of 680–690 nm from the whole spectrum. The absorption spectral shape of CDOM is estimated using a linear least-squares regression of the logarithm of \(a_{\text{CDOM}}(\lambda)\) and reported in units of nm\(^{-1}\) [51] (Equation (6)):

\[
a_{\text{CDOM}}(\lambda) = a_{\text{CDOM}}(\lambda_0)e^{-S(\lambda-\lambda_0)},
\]

where \(a_{\text{CDOM}}(\lambda)\) is the absorption coefficient at wavelength \(\lambda\), \(a_{\text{CDOM}}(\lambda_0)\) is the absorption coefficient at a reference wavelength \(\lambda_0\), and \(S\) is the spectral slope in the spectral range from \(\lambda_0\) to \(\lambda\), with \(\lambda_0 < \lambda\).

Many studies have demonstrated the importance of the slope in the range of 275–295 nm \(S_{275–295}\) to assess the dissolved organic molecular weight [16,17,52,53]; therefore, this interval was adopted in this study.
The proportion of CDOM to the total absorption ($a_{CDOM}$: ($a_{CDOM} + a_p$)), and of particles to the total absorption ($a_p$: ($a_{CDOM} + a_p$)), except pure water, were also calculated at 443 nm.

### Chlorophyll-a Concentration

The concentration of chl $a$ ($\mu$g L$^{-1}$) was measured from 25 to 100 mL of water samples filtered using 25 mm Whatman GF/F glass fiber filters (0.7 $\mu$m nominal pore size). At the laboratory, triplicate samples of pigments were extracted from the filters after immersion in 10 mL of 90% acetone/Dimethyl sulfoxide (DMSO) solution (60/40 by volume, according to Shoaf and Lium [54]) for 24 h, in the dark, at $-10^\circ$C. The resulting extraction was analyzed using a calibrated bench Turner Designs 10-AU fluorimeter.

### Suspended Particulate Matter Concentration

Suspended particulate matter concentration (SPM, mg L$^{-1}$) was determined gravimetrically following the method of Van der Linde [55]. Before field work, 0.45 $\mu$m (nominal pore size) cellulose acetate filters (Millipore) were dried for 24 h, weighed ($W_1$), and placed in Petri plates for transportation. In the field, a volume ($V$) of 150 to 400 mL of surface water was filtered, and the filters for SPM were stored for further laboratory analysis. At the laboratory, SPM filters were dried for 72 h at 50 $^\circ$C and weighed ($W_2$) to determine SPM according to Equation (7):

$$\frac{W_1 - W_2}{V} \cdot \frac{W_1 - W_2}{V}.$$ (7)

The number of samples collected in the Amazon and CW rivers are shown in Table 1.

### Table 1. Total number of in situ data gathered during the years of 2014–2017 in the Lower Amazon region.

| Parameter          | Amazon River | CW Rivers | Total Samples |
|--------------------|--------------|-----------|---------------|
| $a_{CDOM}(\lambda)$ (m$^{-1}$) | 116          | 14        | 130           |
| $S_{CDOM}(\lambda)$ (nm$^{-1}$) | 116          | 14        | 130           |
| $a_p(\lambda)$ (m$^{-1}$) | 95           | 14        | 109           |
| $S_p(\lambda)$ (nm$^{-1}$) | 95           | 14        | 109           |
| SPM (mg L$^{-1}$) | 90           | 12        | 102           |
| chla ($\mu$g L$^{-1}$) | 85           | 12        | 97            |

#### 2.3. In Situ Rs Classification

#### 2.3.1. Spectra Normalization

Each $R_s$ spectrum was normalized by its integrated value [22, 23, 56] according to the Equation (8). The integration was computed using the trapezoidal method, over the 400–800 nm range.

$$r_n(\lambda) = \frac{R_{rs}(\lambda)}{\int_{\lambda_1}^{\lambda_2} R_{rs}(\lambda) d\lambda},$$ (8)

where $r_n(\lambda)$ (in units of nm$^{-1}$) is the normalized spectrum obtained by integration between $\lambda_1$ (400 nm) and $\lambda_2$ (800 nm).

The goal of the normalization is to reduce the first order variability of reflectance and focus on the reflectance spectral shape, hence removing amplitude differences caused by concentration variation. The normalization is a common step that precedes the optical classification procedure, as performed in previous studies [20, 22, 56, 57]. Vantrepotte et al. [22] highlighted the importance of using normalized reflectance spectra instead of raw $R_s$ to perform an optical water type classification. For example, if a water body has a high concentration of SPM (e.g., the Amazon River), the use of raw $R_s$ would lead to a
2.3.2. Optical Water Type Identification

Different supervised and unsupervised methods have been used for the classification of $R_{rs}$ with the aim of defining OWTs based upon differences in the magnitude and shape of the spectral curves [20–23,31,56–58]. Unsupervised classification can be used when a priori knowledge is not available, and $R_{rs}$ data is used to define the clusters, thereby being less subject to user errors [59]. The $k$-means and fuzzy c-means are known techniques among the unsupervised classification methods [12,13,20–22,24,31,57]. When comparing $k$-means versus fuzzy c-means and other unsupervised methods (e.g., Ward algorithm), the clustering results can be very similar [20,22,59]. Therefore, $k$-means was adopted in this study to cluster the data into homogeneous optical groups.

$k$-means technique (Equation (9)) requires a user-specified number of clusters ($k$), which are represented by their centroids [60]. The clusters are defined by measuring the square Euclidian distance of the centroids. For this, an initial mean spectra (“seed”) is arbitrarily specified for each of the $k$-clusters. Then, each spectra of the dataset are assigned to the class whose mean spectra value is closest to the seed, forming the first set of decision boundaries. A new set of cluster seed is then calculated from this classification, and the spectra is reassigned accordingly, until there is no significant change in the spectra assignments from one iteration ($i$) to the next [59]. The Silhouette width analysis [61] was used to determine the number of clusters, by measuring how similar an individual point is to its own cluster.

$$\Delta \mu(i) = \sum_{k=1}^{K} \left| \mu_i^k - \mu_i^{k-1} \right|.$$  

Another way to assess water color approached in this study was proposed by Vandermeulen et al. [62], by using the weighted harmonic mean of the $R_{rs}(\lambda)$ that create an index called Apparent Visible Wavelength ($AVW$), in units of nanometers (Equation (10)):

$$AVW = \frac{\sum_{i=\lambda_1}^{\lambda_n} R_{rs}(\lambda_i)}{\sum_{i=\lambda_1}^{\lambda_n} \frac{R_{rs}(\lambda_i)}{\lambda_i}} = \left( \frac{\sum_{i=\lambda_1}^{\lambda_n} \lambda_i^{-1} R_{rs}(\lambda_i)}{\sum_{i=\lambda_1}^{\lambda_n} R_{rs}(\lambda_i)} \right)^{-1}.$$  

The dynamical spectral index proposed by Vandermeulen et al. [62] does not require a training dataset and/or defining the number of classes a priori. Instead, the result is given as a gradient of wavelengths values that represent a quantitative descriptor of weighted mean color reflected from the water surface. Both classification $k$-means and the $AVW$ index were applied on the normalized $R_{rs}(\lambda)$ spectra and, therefore, focus on the spectral shape. The $AVW$ was used as a complement information, defining the thresholds of the determined classes.

2.4. Satellite Data

A set of 48 level 1 OLCI images from Sentinel-3 satellite (S3A) were downloaded from the Copernicus Online Data Access (https://coda.eumetsat.int/, accessed on 9 March 2021) and processed to level 2 and 3 using SeNtinel Applications Platform (SNAP) software, version 8.0.2. This imagery has a nominal spatial resolution of 300 m with a 14-bit radiometric resolution [26], and was processed to derive $R_{rs}$ data. Initially, four images with low cloud coverage (<10%) and representative of each discharge season (rising water—3 January 2018, high water—27 June 2020, falling water—25 July 2017, low water—7 December 2017) were selected. From those four starting images, other ±10 days images were also downloaded and processed, with a total of 48 images. The level 2 images were derived using the Case 2 Regional Coast Color processor (C2RCC), and the flagged bright areas were removed from the scenes. The C2RCC is an atmospheric correction method that has an overall good performance over inland water [63].

classification primarily based on the relative concentration of sediments, and, consequently, all other optically significant constituents contributions could be neglected.
The 490 nm, 620 nm, and 779 nm bands (B4, B7, and B16, respectively) were used for the classification approach. These bands were chosen by their potential for distinguishing classes, and, according to ESA [26], the band 620 nm is important to assess sediment loadings, while band 490 nm is relevant to assess chl [64]. The near infrared band at 779 nm is also relevant in waters with a high concentration of suspended sediments as in the study area [13].

2.4.1. Satellite $R_{rs}(\lambda)$ Classification

The Mahalanobis distance was used to determine the separability of the given classes upon S3A images. The utilization of the Mahalanobis distance to distinguish the spectra among the classes has been applied in previous ocean color studies [22,56,65]. For this, OLCI bands were simulated utilizing in situ $R_{rs}(\lambda)$ according to the spectral response function of the respective sensor bands. Later, the simulated B4, B7, and B16 bands were selected and normalized, as described in Equation 8. The normalized bands were then log-transformed. The same process was applied to OLCI B4, B7, and B16 bands, where the bands were normalized (Equation (8)) and log-transformed. The classification consisted of calculating statistically how much of a given unknown spectrum ($x$) belongs to a defined class according to the average spectrum, $\mu$, and the covariance matrix, $\Sigma$. The distance of the resulting input $x$ to a given class $ik$ was estimated by the squared Mahalanobis distance $\Delta^2_{ik}$ (Equation (11)):

$$\Delta^2_{ik}(x) = (x - \mu_{ik})^T \sum_{ik}^{-1} (x - \mu_{ik})$$  \hspace{1cm} (11)

where $T$ indicates the matrix transpose. The probability ($p$) that an unknown spectrum ($x$) belongs to a class $ik$ was estimated as (Equation (12)):

$$p(ik, x) = 1 - \frac{\gamma(n/2, \Delta^2_{ik}(x)/2)}{\Gamma(n/2)}$$  \hspace{1cm} (12)

where $\Gamma$ is the gamma function, $\gamma$ is the lower incomplete gamma function, and $n$ is the number of the bands (3) used in the classification.

The normalized belonging $p^*$ was calculated as (Equation (13)):

$$p^*(ik) = \frac{p(ik)}{\sum_{ic=1}^{NC} p(ik)}$$  \hspace{1cm} (13)

More details of how $p^*(ik)$ and $p(ik, x)$ were calculated can be found in Vantrepotte et al. [22]. The algorithm developed to calculate the Mahalanobis distance, as well as $p^*(ik)$ and $p(ik, x)$, was implemented in MATLAB programming language.

2.4.2. Satellite Pixels Labelling

The 48 classified OLCI images were temporally binned to represent the four discharge seasons. The $p^*(ik)$ of each image was averaged according to the respective season. The Mahalanobis classification was binned based on the mode, i.e., evaluating the maximum frequency of each class per pixel. The methodology used in this study is shown in Figure 2.
3. Results

3.1. General Bio-Optical Characterization of the Lower Amazon Region

Table 2 summarizes the basic statistics of all samples obtained across the Lower Amazon region, disregarding the seasonality. Values of $a_{\text{CDOM}}(443)$ of Amazon River and CW rivers are in a similar range, although the coefficient of variation of CW $a_{\text{CDOM}}(443)$ (44%) are almost two-fold higher than the coefficient of variation of Amazon River $a_{\text{CDOM}}(443)$ (25%). $S_{275-295}$ had a higher average value ($0.0147 \pm 0.0014$ nm$^{-1}$) and a higher variability (10%) for CW river samples compared to Amazon River samples (3%).

|                      | Clearwater Rivers                      | Amazon River                   |
|----------------------|----------------------------------------|--------------------------------|
| $a_{\text{CDOM}}(443)$ (m$^{-1}$) | 2.01 ± 0.88 | 2.12 ± 0.53 |
| $S_{275-295}$ (nm$^{-1}$) $\times 10^{-2}$ | 1.47 ± 0.14 | 1.45 ± 0.05 |
| $a_p(443)$ (m$^{-1}$) | 1.68 ± 2.44 | 3.16 ± 2.11 |
| $S_p$ (nm$^{-1}$) $\times 10^{-2}$ | 1.05 ± 0.13 | 1.05 ± 0.08 |
| SPM (mg L$^{-1}$) | 14.0 ± 18.1 | 69.7 ± 33.8 |
| chla (µg L$^{-1}$) | 6.1 ± 7.7 | 1.4 ± 0.8 |
| $a_{\text{CDOM}}(443)$: $a_p(443)$ | 0.64 ± 0.19 | 0.44 ± 0.12 |
| $a_p$: ($a_{\text{CDOM}} + a_p$) (443) | 0.36 ± 0.19 | 0.56 ± 0.12 |
| $a_p(443)$: SPM | 0.11 ± 0.05 | 0.05 ± 0.01 |

|                      | Average ± SD | CV % | Range | Average ± SD | CV % | Range |
|----------------------|--------------|------|-------|--------------|------|-------|
| $a_{\text{CDOM}}(443)$ (m$^{-1}$) | 2.01 ± 0.88 | 44   | 0.74–3.53 | 2.12 ± 0.53 | 25   | 0.89–4.40 |
| $S_{275-295}$ (nm$^{-1}$) $\times 10^{-2}$ | 1.47 ± 0.14 | 10   | 1.32–1.85 | 1.45 ± 0.05 | 3    | 1.30–1.66 |
| $a_p(443)$ (m$^{-1}$) | 1.68 ± 2.44 | 145  | 0.37–9.44 | 3.16 ± 2.11 | 67   | 1.18–8.93 |
| $S_p$ (nm$^{-1}$) $\times 10^{-2}$ | 1.05 ± 0.13 | 12   | 0.88–1.30 | 1.05 ± 0.08 | 8    | 0.91–1.33 |
| SPM (mg L$^{-1}$) | 14.0 ± 18.1 | 129  | 1.8–67.9 | 69.7 ± 33.8 | 48   | 18.3–160.8 |
| chla (µg L$^{-1}$) | 6.1 ± 7.7 | 126  | 1.5–31.3 | 1.4 ± 0.8 | 57   | 0.5–6.3 |
| $a_{\text{CDOM}}(443)$: $a_p(443)$ | 0.64 ± 0.19 | 30   | 0.17–0.80 | 0.44 ± 0.12 | 27   | 0.17–0.64 |
| $a_p$: ($a_{\text{CDOM}} + a_p$) (443) | 0.36 ± 0.19 | 53   | 0.20–0.83 | 0.56 ± 0.12 | 21   | 0.36–0.83 |
| $a_p(443)$: SPM | 0.11 ± 0.05 | 45   | 0.06–0.24 | 0.05 ± 0.01 | 20   | 0.03–0.11 |

$SD = \text{Standard Deviation}; CV = \text{Coefficient of Variation}.$

The absorption coefficient of particulate matter ($a_p(443)$) is strongly correlated with SPM ($R^2 = 0.91$, $p < 0.05$, $N = 77$) (Figure 3), as previously reported by other studies in the Amazon basin [66]. The average ratio of $a_p(443)$ : SPM is higher for CW rivers (CW: 0.11 ± 0.05; Amazon: 0.05 ± 0.01; see Table 2), with a 45% coefficient of variation (Amazon River CV: 20%). $S_p$ showed similar values for Amazon and CW rivers ($S_p$: 0.010 nm$^{-1}$), with low variability (Amazon 8% and CW rivers 12%) (see Table 2).
3.2. Seasonal Absorption Budget

Absorption by CDOM is dominant with a large contribution to the total absorption in CW rivers ($a_{\text{CDOM}}(443) : a_{\text{CDOM}}(443) + a_p(443) = 0.64 \pm 0.19 \text{ m}^{-1}$). However, its variability is similar ($\geq 30\%$) to what is observed for the Amazon River (Table 2). In the Amazon River, although the absorption by particulate matter has a meaningful dominance, it is almost even with absorption by CDOM ($a_{\text{CDOM}} 44\%$; $a_p 56\%$; see Table 2). The balanced proportion among $a_{\text{CDOM}}$ and $a_p$ in the Amazon River is clearer when the absorption budget is assessed by hydrological season. Except for rising water, when $a_p$ is almost three-fold higher than $a_{\text{CDOM}}$, all other seasons are nearly equivalent (Table 3). The variability is also greater during the rising water period for CW rivers, probably due to the influence of the overflow of the Amazon River that brings CDOM and SPM into the tributaries (Table 3).

Table 3. Absorption budget of the Amazon River (Am) and clearwater rivers (CW) for rising water (RW), high water (HW), falling water (FW), and low water (LW) seasons.

| Season | $a_{\text{CDOM}} : (a_{\text{CDOM}} + a_p) (443)$ | $a_p : (a_{\text{CDOM}} + a_p) (443)$ |
|--------|---------------------------------|---------------------------------|
|        | Average ± SD | CV % | Average ± SD | CV % |
| RW     | Am 0.26 ± 0.08 | 31   | 0.74 ± 0.08 | 11   |
|        | CW 0.55 ± 0.26 | 47   | 0.45 ± 0.26 | 58   |
| HW     | Am 0.48 ± 0.09 | 19   | 0.52 ± 0.09 | 17   |
|        | CW 0.78 ± 0.03 | 4    | 0.22 ± 0.03 | 14   |
| FW     | Am 0.50 ± 0.07 | 14   | 0.50 ± 0.07 | 14   |
|        | CW 0.72 ± 0.06 | 8    | 0.28 ± 0.06 | 21   |
| LW     | Am 0.50 ± 0.05 | 10   | 0.50 ± 0.05 | 10   |
|        | CW 0.62 ± 0.06 | 10   | 0.38 ± 0.06 | 16   |

SD = Standard Deviation; CV = Coefficient of Variation.

3.3. Optical Classification

The advantage of using normalized spectra for clearly highlighting variation in water optical quality focusing on the shape of the $R_{rs}$ spectra rather than on variation in $R_{rs}$ amplitude (and thus on the turbidity level) is illustrated in Figure 4.
The Silhouette method indicated four as the optimal number of OWT classes to be used with the k-means clustering analysis of the in situ normalized \( R_{rs} \) between 400–800 nm (Figure 5A). Likewise, the AVW method presented the normalized \( R_{rs} \) as a gradient of wavelength values from which four central AVW could be detected (Figure 5B), according to the frequency distribution (histogram, not shown). For the k-means perspective, OWT k1 and k2 are representative of the Amazon River mainstem, OWT k3 is representative of the large CW tributaries (Tapajós and Xingu rivers), and k4 is representative of a mixture of other Amazonian water types, respectively. Herein, the OWT classes are named: k1 as Amazon Optical Water Type 1 (AOWT1), k2 as Amazon Optical Water Type 2 (AOWT2), k3 as Clear Optical Water Type (COWT), and k4 as Mixed Amazonian Optical Water Type (MAOWT) (Figure 4A).

Figure 5. (A) Average normalized \( R_{rs} \) spectra resultant from the k-means clustering analysis of the Lower Amazon region; (B) AVW values derived from the normalized in situ \( R_{rs} \) spectra (400–800 nm), yielding a range of values from 543 nm to 612 nm. The thicker lines with circle markers are referred to the central AVW according to the k-means classes.

The AOWT1 consisted in clustered in situ \( R_{rs} \) that were measured during the high, falling, and low water in the Amazon River (T1, T2, T3, and T5; see Figure 1C). The AVW of AOWT1 comprises the interval 584 nm \( \leq \) AVW \( \leq \) 598 nm, with the central wavelength at 590 nm (Figure 5B). Nevertheless, AOWT2 has an increasing reflectance in the same interval (green to NIR; see Figure 5) as a result of the immense input of sediments during the rising water season [66], with the central wavelength shifting to 605 nm in the range of 593 nm \( \leq \) AVW \( \leq \) 612 nm (Figure 5B). The AOWT2 class is primarily composed of water present during the period of rising water and high water seasons, corresponding to T4 and T6 field campaigns, respectively (Figure 1C).

The COWT class consists of \( R_{rs} \) spectra measured during the falling and low seasons in the large CW tributaries (Tapajós and Xingu rivers), in which the AVW comprises the interval 543 nm \( \leq \) AVW \( \leq \) 563 nm, with the central wavelength at 560 nm (Figure 5B). The in situ \( R_{rs} \) spectra clustered in the MAOWT corresponds to rising water and high water periods in the CW rivers. MAOWT reflectance spectrum showed a less prominent chl peak in comparison to COWT, even though the average measured chl concentration was higher in comparison to other OWs (Table 4). Another characteristic of MAOWT is a higher signal at the red-NIR spectrum when compared to COWT (Figure 5). The overflow of the Amazon River into CW rivers during the rising and high water seasons carries a large load of CDOM and SPM (Table 4). Therefore, the AVW of MAOWT is shifted to higher wavelengths in comparison to COWT, comprising the interval 569 nm \( \leq \) AVW \( \leq \) 581 nm, with the central wavelength at 575 nm (Figure 5B).
Table 4. Bio-optical characteristics of the four optical water types (OWT) identified for the Lower Amazon region.

|                  | AOWT1 | AOWT2 | COWT  | MAOWT |
|------------------|-------|-------|-------|-------|
|                  | Average ± SD | Range | Average ± SD | Range | Average ± SD | Range | Average ± SD | Range |
| $a_{\text{CDOM}}$ (443) (m$^{-1}$) | 2.2 ± 0.6 | 1.3–4.2 | 2.0 ± 0.2 | 1.6–2.7 | 1.0 ± 0.3 | 0.7–1.3 | 2.4 ± 0.6 | 1.3–3.1 |
| $S_{275-295}$ (nm$^{-1}$) $\times 10^{-2}$ | 1.44 ± 0.05 | 1.32–1.54 | 1.45 ± 0.03 | 1.40–1.50 | 1.70 ± 0.17 | 1.51–1.85 | 1.41 ± 0.09 | 1.32–1.57 |
| $a_p$(443) (m$^{-1}$) | 2.03 ± 0.69 | 1.18–4.74 | 6.44 ± 1.61 | 2.72–9.44 | 0.48 ± 0.12 | 0.37–0.64 | 0.89 ± 0.25 | 0.50–1.11 |
| $S_{440-600}$ (nm$^{-1}$) $\times 10^{-2}$ | 1.04 ± 0.06 | 0.91–1.25 | 1.05 ± 0.07 | 0.95–1.26 | 0.93 ± 0.04 | 0.88–0.97 | 1.14 ± 0.14 | 0.95–1.30 |
| SPM (mg L$^{-1}$) | 43.6 ± 9.9 | 28.3–68.2 | 95.2 ± 29.4 | 49.8–157.2 | 4.4 ± 2.7 | 1.8–6.8 | 10.0 ± 5.8 | 3.7–17.8 |
| chl$a$ (µg L$^{-1}$) | 1.5 ± 0.8 | 0.5–4.4 | 1.4 ± 0.4 | 0.5–2.1 | 5.1 ± 2.9 | 2.3–8.5 | 9.1 ± 11.2 | 1.8–31.3 |
| $a_{\text{CDOM}} - a_p$ | 0.51 ± 0.06 | 0.28–0.64 | 0.25 ± 0.07 | 0.17–0.50 | 0.67 ± 0.10 | 0.55–0.78 | 0.73 ± 0.05 | 0.66–0.80 |
| $a_{\text{CDOM}} - a_p$ (443) | 0.49 ± 0.06 | 0.36–0.72 | 0.75 ± 0.07 | 0.50–0.83 | 0.33 ± 0.10 | 0.22–0.45 | 0.27 ± 0.05 | 0.20–0.34 |

4. Discussion

4.1. CDOM Absorption

The Amazon River’s $a_{\text{CDOM}}$(443) values (2.12 ± 0.53 m$^{-1}$; see Table 2) are in the same range of values presented by Martínez et al. [13] ($a_{\text{CDOM}}$(440) = 3.3 ± 1.0 m$^{-1}$), and relatively lower when compared to values presented by Jorge et al. [67] ($a_{\text{CDOM}}$(420) $\sim$ = 3.25 ± 0.46 m$^{-1}$) for Amazon white waters. CW rivers showed a similar average $a_{\text{CDOM}}$(443) value (2.01 ± 0.88 m$^{-1}$; see Table 2) in comparison to values reported by Martínez et al. [13] ($a_{\text{CDOM}}$(440) = 1.8 ± 0.8 m$^{-1}$). The discrepancies between $a_{\text{CDOM}}$ values from different studies can be attributed to differences in each dataset size and representativeness, alternative filtering methods, chosen wavelengths, and protocol measurements (no baseline correction in Martínez et al. [13], for instance), as well as to different sampling locations and seasonality (e.g., Jorge et al. [67] sampled only during the rising water season in the Amazonian lakes).

High $S_{275-295}$ values generally indicate lower molecular weight dissolved organic matter (DOM) [17]. This potentially suggests that photobleaching has a larger impact on DOM composition in the CW rivers, which also receive less organic matter input from the terrestrial biosphere. A higher contribution of autochthonous DOM from phytoplankton in the CW rivers might also explain the observed variation [17,68], although a recent study demonstrated that $S_{275-295}$ is only slightly sensitive to the impact of phytoplankton-derived DOM [69]. The greater presence of phytoplankton in the CW rivers can be evidenced by high values of chl$a$ in comparison to the Amazon River (average of 6.1 and 1.4 µg L$^{-1}$, respectively; see Table 2). The higher variability of $S_{275-295}$ (CV: 10% for CW and 3% for the Amazon River; see Table 2), and other parameters in the CW rivers might be related to the location where the samples were collected, very close to the Amazon River mainstem, where turbid Amazon River waters may inflow into the smaller tributaries depending on the discharge season.

4.2. Particulate Matter Absorption

The slope of the regression between $a_p$ and SPM (0.053; see Figure 3) is much higher than what is reported for coastal waters (~0.031) [11,70], and for combusted particles (0.024) [71], possibly due to an abundance of organic particles. CW river $a_p$(443) values are related to low concentrations of SPM (Table 2). Amazon River samples with SPM > 100 mg L$^{-1}$ were mainly obtained during the rising water season, indicating that the CW rivers may have more organic particles and may be more influenced by the Amazon River mainstem seasonality.
The Amazon and CW rivers $S_p$ presented relatively lower values than what is usually reported for coastal and oceanic waters ($>0.011 \text{ nm}^{-1}$) [11,50,72–74]. Previous studies showed that $S_{nap}$ values of about $0.011 \text{ nm}^{-1}$ are typical of mineral dominated waters [11,71,73,75], and higher $S_{nap}$ ($>0.011 \text{ nm}^{-1}$) are related to organic particles [11,75]. Nevertheless, recent studies found an opposite trend in CDOM-rich waters, where steeper $S_{nap}$ are in fact associated with more inorganic suspended matter by DOM adsorption in mineral surfaces [14,15,49,70]. Considering the high values of CDOM in the study area, this situation may also apply in the Lower Amazon region, where the relatively lower values of $S_p$, in comparison to the literature dedicated to coastal and oceanic waters, are in fact related to organic particles.

4.3. Differences among the Optical Water Types of the Lower Amazon Region

The average spectral shape of the four resultant OWTs from the Lower Amazon are in general agreement with other studies in the Amazon Basin, with measurements upstream of Óbidos and at lakes and floodplains [13,67,76–79]. Besides the evident distinction between the Amazon and CW rivers, another prevailing difference is related to the seasonality of the in situ measured $R_{rs}$. In the Amazon River (AOWT1 and AOWT2), there is no clear chl a signal, and, although the reflectance signal is dominated by SPM, it is possible to observe that AOWT1 has a uniform signal over a wide part of the spectrum from the green to near-infrared wavelengths (NIR).

As mentioned, the AOWT2 comprises in situ $R_{rs}$ from rising water and high water (T4 and T6, respectively; see Figure 1C). The high water period in 2017 (T6) contrasts to 2014 (T1) because it approximately resembles a rising water season, as a consequence of the El Niño impact on river discharge in the previous year [43]. Before 2014, the discharge peak did not fluctuate as much as that observed for 2016 (Figure 1C). The rainfall normalization in 2016 flushed sediments and organic matter to the Amazon River that were stored in the floodplains and tributaries from the previous year. The impact of the 2015–2016 El Niño on bio-optical properties of the Lower Amazon region was also discussed by Valerio et al. [1]. It is worth noting that AOWT2 is the spectrum resulting from the impact of rain and increased discharge after a long drought period. This situation could be related to seasonality, as well as to exceptional climate events, during the study period.

The well-defined 560 nm peak of the COWT (Figure 5A) is related to the backscattering of phytoplankton cells [80], indicating chl a contribution, as can be evidenced in Table 4. Although MAOWT has a higher chl a in comparison to COWT (Table 4), its 560 nm peak is less evident. As can be evidenced in Table 4, MAOWT also has higher values of CDOM. When in abundance, CDOM can have a role in the light absorption of the shorter wavelengths of the visible spectrum [81], reducing the reflectance attributed to chl a, while SPM (Table 4) also causes a higher signal in the red-NIR wavelengths.

The standard deviation of the MAOWT chl a ($9.1 \pm 11.2 \text{ mg L}^{-1}$; see Table 4) indicates a large variability. Previous studies also documented a large variability of chl a and phytoplankton blooms during the rising and high water season within CW rivers (e.g., Tapajós River) [82,83]. The optimal balance between the input of nutrients from the surrounding watershed and light availability is likely the cause of this variation. The water clarity of these rivers allows sunlight to penetrate deep into the water column, promoting high rates of primary production.

4.4. Seasonal Distribution of OWT at the Lower Amazon Region

OWT varied by season. During the rising water season (Figure 6A), the Amazon River mainstem was mostly classified as AOWT2, while the larger CW rivers (Tapajós and Xingu), other smaller rivers, lakes, and floodplains were mostly classified as MAOWT. The classes AOWT1 and COWT were almost inexistent, with some small patches in floodplains (AOWT1) and at the Tapajós and Xingu rivers (COWT).
Figure 6. Inset, mean seasonal discharge of the Lower Amazon (2005–2015), with the seasonal sampling shown in red; Mahalanobis classification with the Amazon Optical Water Type 1 (AOWT1) in yellow, Amazon Optical Water Type 2 (AOWT2) in red, Clear Optical Water Type (COWT) in green, and Mixed Amazonian Water Type (MAOWT) in blue. (A) rising water image composite; (B) high water composite; (C) falling water composite; and (D) low water composite.

Bio-optical properties of AOWT2 indicate a considerable increase in SPM (95.2 ± 29.4 mg L$^{-1}$), $a_{p}(443) (6.44 ± 1.61$ m$^{-1}$), and in the relative contribution of $a_{p}$ to total absorption (75 %) in comparison with AOWT1 (Table 4). The impact of seasonality on the variability of radiometric and bio-optical properties of the Amazon basin was previously discussed by other studies in the region, especially during the rising water season when the Amazon River reaches its highest SPM concentrations [66,84–86]. This variability can impact the definition of regional bio-optical inversion models. For example, Valerio et al. [1] showed that the dissolved organic carbon (DOC) could not be retrieved using a single relationship between $a_{CDOM}(412)$ and DOC. During the rising water season and other river discharge peak situation (i.e., caused by exceptional climate events), there is a decoupling of the relationship of CDOM-DOC. Hence, future efforts for retrieving DOC using an OWT classification approach at the Lower Amazon region require considering AOWT2 specifically.

During the high water season, the Amazon River mainstem is predominantly occupied by AOWT1 (Figure 6B). The relative contribution of $a_{p}$ to total absorption is now lower than presented by AOWT2 (AOWT1 49%; AOWT2 75%; see Table 4), due to the decrease in SPM input (43.6 ± 9.9; see Table 4). Conversely, the relative contribution of $a_{CDOM}$ to total absorption increased (AOWT1 51%; AOWT2 25%; see Table 4). The COWT is observed in the classified OLCI composite for the high water season as the dominant class in Tapajós and part of the Xingu River, with the lower SPM concentration in comparison to other classes (4.4 ± 2.7 mg L$^{-1}$; see Table 4). The probability of pixel belonging to a MAOWT in lakes, floodplains, smaller rivers, and CW river margins is high.

During the falling water season, the probability of a pixel being classified as COWT increased for CW rivers, especially the Xingu River (Figure 6C). The phytoplankton primary production rates could increase due to the receding of the Amazon River pulse, leading to an enrichment of dissolved nutrients and decrease in turbidity [87]. In the Amazon River mainstem, AOWT1 is still dominant, while MAOWT is dominant in lakes, floodplains, smaller rivers, and CW river margins.
During the low water season, the transient status of the Amazon River is evidenced (Figure 6D). For 07/12/2017 (±10 days), it is possible to observe a decrease in AOWT1 presence at the upstream region, while AOWT1 is still dominant downstream. By December, the discharge in the Lower Amazon region increases at Óbidos (Figure 6D), carrying SPM from the Andes [88]. This volume of water loaded with SPM pushes AOWT1 towards the Amazon River mouth. Lakes and floodplains of the Amazon River upstream are dominated by AOWT2, and COWT start to shrink as evidenced in the Xingu River. Conversely, MAOWT presented dominance in the Xingu, likely due to the overflow of the Amazon River and the input of SPM.

During all hydrologic seasons, AOWT2 was dominant at the Amazon River mouth (Figure 6), indicating high SPM concentrations, and high relative contribution of \(a_p\) to total absorption (Table 4), probably caused by the tidal influence on the lower reaches of the Amazon River. Near the Amazon River mouth, the tide can completely reverse the flow toward the river upstream, increasing the water residence time, turbulence [34], and consequently, sediment resuspension in the water column.

5. Conclusions

The present study describes the bio-optical and radiometric variability along the Lower Amazon region, considering different discharge seasons. The Amazon River and CW rivers \(a_{\text{CDOM}}\) showed similar values (in average), in agreement with previous studies in the region. CW rivers \(a_{\text{CDOM}}\) and \(S_{275-295}\) presented a relatively higher variability, indicating the influence of the Amazon River overflow into the tributaries. Dissolved organic matter in the CW rivers had lower molecular weight compared to the Amazon River, as indicated by higher \(S_{275-295}\) values. This indicates a higher impact of photobleaching rates in CW rivers and less organic matter input from the terrestrial biosphere.

The absorption coefficient of total particulate matter had a good relationship with SPM concentration, showing the presence of high concentration of organic particles, especially in CW rivers. As expected, CW rivers had the lowest values of SPM, while the Amazon River had the higher values, especially during rising water. The variability of \(a_p(443)\): SPM at CW rivers is also higher, indicating the influence of the Amazon River mainstem into the tributaries. Further studies are required to better understand the observed variability of \(S_p\) in the Lower Amazon region. At this point, it is not clear if the slope approach was able to trace the relative proportion of organic and inorganic particles, while this potential has already been underlined for marine coastal waters [11].

The \(k\)-means clustering analysis identified four distinct OWT, and the analysis was complemented using the optical index AVW that presented the threshold of the optical classes. Our results showed that the OWT in the Lower Amazon region are strongly seasonally dependent and impacted by exceptional climate events. Briefly, AOWT1 dominance occurs during the high water and falling water season, in the Amazon River mainstem. During the end of the low water season, river discharge starts to rise, surging water is carried with SPM from the Andes, and AOWT2 start to dominate the upstream region, pushing AOWT1 towards the Amazon River mouth. During the rising water season, AOWT2 dominates the Amazon River mainstem. In the largest CW rivers (Tapajós and Xingu), COWT dominance starts to increase during the high water season, with the peak in the falling water season, later decreasing during the low water season. MAOWT is dominant and/or codominant in almost all of the Lower Amazon region, with a higher contribution in smaller tributaries and lakes.

For the Amazon River, not only seasonality is important but also extreme climate events, such as the record flood of 2014 and the severe drought of 2015–2016, that occurred during the study period. AOWT1 class was defined with the clustering of in situ measured \(R_s\) from falling water, low water, and high water from 2014 (T1), being representative of the general condition of the Amazon River spectrum, except for the rising water season. AOWT2 class was defined with the \(R_s\) spectra from rising water and high water from 2017 (T6). The discrepancies between high water of 2014 and 2017 may be associated with the
impact of the El Niño 2015–2016 on the Amazon River discharge. Hence, AOWT2 can also be related to the direct impact of rising discharge due to rainfall on the river optical signature after a long drought. The spectra rise from green to NIR can be associated with the considerable increase in SPM input.

This study corroborates that rising water season presents a specific optical signature when compared to the other hydrological seasons, being a hallmark for the Lower Amazon region. Although rising waters occur during the first trimester of the year, river discharge is impacted by climate events and, as a result, can make it difficult to delineate the seasonality. Therefore, the k-means classification, as well the optical index AVW proposed here, can be used to monitor the nature of the water masses of the Lower Amazon region, considering the influence of unusual events. Among those, we can cite distinctive hydrological seasonal patterns associated with climate events, or other anomalous conditions potentially caused by anthropogenic impacts. Our study showed that water type classes derived from k-means clustering analysis can be used to assess bio-optical properties of the Lower Amazon region. Similarly, the AVW optical index can also be used for this purpose, especially if a proper central wavelength in the AVW range is considered in the spectra gradient. The classification presented here can provide a framework to the development of remote sensing inversion models adjusted to the complexity of the Lower Amazon River waters.

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