



# Phytoplankton phenology indices in coral reef ecosystems: Application to ocean-color observations in the Red Sea

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## ABSTRACT

Phytoplankton, at the base of the marine food web, represent a fundamental food source in coral reef ecosystems. The timing (phenology) and magnitude of the phytoplankton biomass are major determinants of trophic interactions. The Red Sea is one of the warmest and most saline basins in the world, characterized by an arid tropical climate regulated by the monsoon. These extreme conditions are particularly challenging for marine life. Phytoplankton phenological indices provide objective and quantitative metrics to characterize phytoplankton seasonality. The indices i.e. timings of initiation, peak, termination and duration are estimated here using 15 years (1997–2012) of remote sensing ocean-color data from the European Space Agency (ESA) Climate Change Initiative project (OC-CCI) in the entire Red Sea basin. The OC-CCI product, comprising merged and bias-corrected observations from three independent ocean-color sensors (SeaWiFS, MODIS and MERIS), and processed using the POLYMER algorithm (MERIS period), shows a significant increase in chlorophyll data coverage, especially in the southern Red Sea during the months of summer NW monsoon. In open and reef-bound coastal waters, the performance of OC-CCI chlorophyll data is shown to be comparable with the performance of other standard chlorophyll products for the global oceans. These features have permitted us to investigate phytoplankton phenology in the entire Red Sea basin, and during both winter SE monsoon and summer NW monsoon periods. The phenological indices are estimated in the four open water provinces of the basin, and further examined at six coral reef complexes of particular socio-economic importance in the Red Sea, including Siyal Islands, Sharm El Sheikh, Al Wajh bank, Thuwal reefs, Al Lith reefs and Farasan Islands. Most of the open and deeper waters of the basin show an apparent higher chlorophyll concentration and longer duration of phytoplankton growth during the winter period (relative to the summer phytoplankton growth period). In contrast, most of the reef-bound coastal waters display equal or higher peak chlorophyll concentrations and equal or longer duration of phytoplankton growth during the summer period (relative to the winter phytoplankton growth period). The ecological and biological significance of the phytoplankton seasonal characteristics are discussed in context of ecosystem state assessment, and particularly to support further understanding of the structure and functioning of coral reef ecosystems in the Red Sea.

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## 1. Introduction

Coral reefs are among the most biologically diverse ecosystems on Earth. They occupy less than 0.1% of the world's oceans, yet they host 25% of all the marine species (Spalding, Ravilious, & Green, 2001). Coral reefs deliver valuable and vital ecosystem services. They offer coastal protection, employment (through fisheries, recreation, and tourism), and are a major source of food for millions of people around the world

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(Hoegh-Guldberg, 1999). However, coral reefs are fragile ecosystems, facing serious threats from global climate change, marine acidification, destructive and unsustainable fishing practices, and water-polluting land-use activities (Hoegh-Guldberg et al., 2007; Wilkinson, 1999).

The Red Sea hosts thriving coral reef communities that have adapted to one of the most saline and warm basins in the world (Belkin, 2009; Cantin, Cohen, Karnauskas, Tarrant, McCorkle, 2010). The Red Sea is also unique because of its partial isolation from the Indian Ocean, its arid tropical climate, and its prevailing wind system regulated by the monsoon (Halim, 1969). During summer, northwesterly winds predominate in the Southern part of the Red Sea from June to September

(i.e. summer NW monsoon period), while during winter, in the southern Red Sea, prevailing winds reverse to southeasterly from October to May (i.e. winter SE monsoon period). Over the northern part of the Red Sea (North of 20°N), wind blow persistently from north-northwest throughout the year (Patzert, 1972). The Red Sea is further characterized by warm water temperature, which has been shown to be rapidly increasing since the mid-90s (from 27.4 °C on average during 1985–2003 to 28.1 °C on average during 1994–2007; Raitso et al., 2011). These conditions make the Red Sea an excellent laboratory for studying the effects of the environment on marine organisms. In fact, the Red Sea has been shown not to be immune to the effects of global climate change and other disturbances, as evidenced by recent documentation of thermal coral bleaching events (e.g., Furby, Bouwmeester, & Berumen, 2013; Pineda et al., 2013) and historical crown-of-thorns starfish outbreaks (e.g., Riegl, Berumen, & Bruckner, 2013). Furthermore, in some regions, the majority of commercially targeted species have been overfished for decades (Jin, Kite-Powell, Hoagland, & Solow, 2012). High pressure is placed on apex predator populations, which are critical for ecological functioning in coral reefs, and are notably understudied in Red Sea ecosystems (Berumen et al., 2013; Spaet, Thorrold, & Berumen, 2012). For instance, the roving coral-grouper (*Plectropomus pessuliferus*) is highly targeted in the Red Sea, and a marked decline (or even disappearance in certain areas) has been shown, leading to the inclusion of this species into the International Union for Conservation of Nature (IUCN) Red List (Ferreira, Gaspar, Samoilys, Choat, & Myers, 2008).

Located at the base of the marine food web, phytoplankton support the functioning of coral reef ecosystems (Genin, Monismith, Reidenbach, Yahel, & Koseff, 2009; Wild, Jantzen, Struck, Hoegh-Guldberg, & Huettel, 2008; Wyatt, Lowe, Humphries, & Waite, 2010; Yahel, Post, Fabricius, & Genin, 1998), providing a source of food for many coral reef-associated organisms, including zooplankton, benthic grazers such as sponges (e.g., Richter, Wunsch, Rasheed, Kötter, & Badran, 2001; Yahel, Sharp, Marie, Häse, & Genin, 2003), bivalves (e.g., Yahel, Marie, Beninger, Eckstein, & Genin, 2009), and pelagic larvae (e.g., Erez, 1990; Johannes, 1978; Lo-Yat et al., 2011). In fact, the larvae of many marine species (including fish, crustaceans, mollusks and echinoderms) graze on phytoplankton during this vulnerable stage of their lives. Evidence that survival of gadoid fish larvae depends on the timing of the local spring bloom of phytoplankton has been demonstrated by Platt, Fuentes-Yaco, and Frank (2003) using a combination of satellite chlorophyll and in situ observations. Another characteristic example is the tight coupling reported between shrimp hatch time and the timing of remotely-sensed phytoplankton spring bloom peak in high-latitude ecosystems of the North Atlantic basin (Koeller et al., 2009). In a tropical ecosystem, Lo-Yat et al. (2011) have shown a significant positive relationship between remotely-sensed chlorophyll concentrations and the recruitment success of coral reef fish larvae. Assessing the phytoplankton phenology (timing of food availability) is important, as any changes may propagate up the marine food web, which may lead to trophic mismatch and alter the function of marine ecosystems (Edwards & Richardson, 2004).

To investigate trophic interactions in coral reef ecosystems, and to be able to detect anomalous trends or patterns, a comprehensive understanding of the seasonal variability (climatological cycle) of microscopic marine algae, phytoplankton, is required. In the Red Sea, general ecological research (Berumen et al., 2013) and long-term large-scale biological datasets are rare, with the latter mainly limited to satellite-based observations of ocean color (Acker, Leptoukh, Shen, Zhu, & Kempler, 2008; Brewin, Raitso, Pradhan, & Hoteit, 2013; Labiosa, Arrigo, Genin, Monismith, van Dijken, 2003; Raitso, Pradhan, Brewin, Stenchikov, & Hoteit, 2013). The color of the ocean is a good indicator of the primary photosynthetic pigment found in phytoplankton, chlorophyll (Sathyendranath & Platt, 1997). Over the past two decades, remote-sensing measurements of chlorophyll have provided unique information on surface marine phytoplankton, allowing us to monitor their distribution at high temporal and spatial resolution in coastal and open

oceans (Blondeau-Patissier, Gower, Dekker, Phinn, & Brandoc, 2014). Chlorophyll concentration varies seasonally following the growth and decline of phytoplankton populations, which define the phytoplankton growing period. A suite of indices has been proposed to quantify phytoplankton seasonality (Platt & Sathyendranath, 2008) and to provide support to investigations on the composition, structure and functioning of the marine ecosystem (Racault, Platt, et al., 2014). The study of timing of periodical growth of phytoplankton populations relates to phenology. Phenological indices include timings of initiation, peak, termination, and duration of phytoplankton growing period (e.g., Racault, Le Quééré, Buitenhuis, Sathyendranath, & Platt, 2012). Several methods have been proposed to estimate these indices (see Ji, Edwards, Mackas, Runge, & Thomas, 2010 for a review). Conventionally, the methods involve a threshold criterion, which provides a boundary value to delineate initiation and termination of a phytoplankton growing period. The threshold criterion can be estimated directly from the remotely-sensed chlorophyll time-series (Henson, Robinson, Allen, & Waniek, 2006; Racault et al., 2012; Siegel, Doney, & Yoder, 2002; Thomalla, Fauchereau, Swart, & Monteiro, 2011), or after fitting a density function to the chlorophyll time-series (Platt, White, Zhai, Sathyendranath, & Roy, 2009; Vargas, Brown, & Sapiano, 2009; Zhai, Platt, Tang, Sathyendranath, & Hernández Walls, 2011; Sapiano, Brown, Schollaert Uz, & Vargas, 2012; González Taboada & Anadón, 2014; Ardyna et al., 2014), or performing a cumulative summation of the chlorophyll concentration (Brody, Lozier, & Dunne, 2013). Although different methods and threshold criteria may yield similar results, the choice of method and threshold requires scrutiny of the shape of the phytoplankton seasonal cycle. A further caution is that phenological studies require data well distributed in time (i.e. with few missing data), to enable resolution of timings of seasonal events with sufficient precision (Cole, Henson, Martin, & Yool, 2012; Land, Shutler, Platt, & Racault, 2014; Racault, Sathyendranath, & Platt, 2014).

Using SeaWiFS (Sea-Viewing Wide Field-of-View Sensor) and MODIS (Moderate Resolution Imaging Spectroradiometer) observations of chlorophyll, Acker et al. (2008) described the general seasonal variability of phytoplankton in the northern Red Sea. Using MODIS data, Raitso et al. (2013) further examined phytoplankton seasonality in relation to environmental conditions in the Red Sea, and based on biological and physical characteristics, the latter authors proposed a partitioning into four provinces: the northern-Red Sea (NRS), the north-central Red Sea (NCRS), the south-central Red Sea (SCRS), and the southern Red Sea (SRS). However, in spite of the remarkable sampling coverage provided by remote-sensing technology, the presence of persistent clouds and atmospheric aerosol, sun-glint, and sensor saturation over sand have significantly limited data acquisition in the Arabian Sea and Red Sea regions. Specifically, during summer NW monsoon period, in addition to relatively frequent sand storms, the southern Red Sea is affected by hazy-cloudy conditions, which had prevented retrieval of useful remotely-sensed chlorophyll data until recently (Steinmetz, Deschamps, & Ramon, 2011). In 2014, the European Space Agency (ESA) Ocean-Colour Climate Change Initiative (OC-CCI) project produced and validated a consistent, stable, and error-characterized time-series of global ocean-color products based on merged SeaWiFS, MODIS and MERIS (MEdium Resolution Imaging Spectrometer) data (Hollmann et al., 2013; <http://www.esa-oceancolour-cci.org>). The progress made in the OC-CCI project has permitted improved coverage of remotely-sensed chlorophyll measurements in summer months in the Arabian Sea and southern Red Sea regions.

In the present study, we use ESA OC-CCI data to provide the first quantitative investigation of the phenology of phytoplankton in the Red Sea for both the winter and the summer growing periods. The research outcomes unfold as follows: 1) we assess and compare the spatial and temporal coverage of ocean-color observations from OC-CCI products with previously-available single-sensor SeaWiFS, MODIS and MERIS products; 2) we examine the performance of the ESA OC-CCI

chlorophyll retrieval algorithm in shallow reef-bound coastal waters of the Red Sea; 3) we develop an algorithm to compute a suite of phenology indices (i.e. timings of initiation, peak, termination and duration) in the entire Red Sea basin for the winter and summer phytoplankton growth periods; 4) we examine phytoplankton phenology in the vicinity of coral reefs, which we locate using sea-floor elevation data and the UNEP World Conservation Monitoring Centre (UNEP-WCMC) coral reefs dataset; 5) we analyze and compare phytoplankton phenological characteristics from open water areas located in the different ecological provinces and from coastal areas located in the vicinity of six large reef complexes; and 6) we review and discuss the significance of phytoplankton phenology for the functioning of coral reef ecosystems in the Red Sea.

## 2. Materials and methods

### 2.1. Datasets

#### 2.1.1. Sea-floor elevation data

ETOPO5 sea-floor elevation data on a 5-minute latitude/longitude grid were obtained from the National Oceanographic and Atmospheric Administration (NOAA) at: <http://www.ngdc.noaa.gov/mgg/global/etopo5.HTML>. To be coherent with the spatial resolution of remote-sensing data, the sea-floor elevation data were then re-gridded to 4 km resolution using linear interpolation.

#### 2.1.2. Coral reef distribution dataset

Global Distribution of Coral Reefs dataset (version 2010) compiled by the UNEP-WCMC was used to locate the position of coral reefs in the Red Sea. The dataset sources include the Millennium Coral Reef Mapping Project (IMaRS-USF (Institute for Marine Remote Sensing-University of South Florida) & IRD (Institut de Recherche pour le Développement), 2005) and the World Atlas of Coral Reefs (Spalding et al., 2001). The dataset was obtained in ArcGIS vector format at <http://data.unep-wcmc.org/datasets/13>. It was then converted to NetCDF (Network Common Data Form) and gridded at 4 km resolution using the GRASS GIS software (<http://grass.osgeo.org>).

#### 2.1.3. In situ chlorophyll data

Three independent datasets of in situ chlorophyll measurements were used to assess the performance of the ESA OC-CCI chlorophyll retrieval algorithm in reef-bound coastal and open waters of the Red Sea. The datasets, covering an extensive part of the study area, include: 1) In vivo fluorometric data on chlorophyll concentration collected from three research cruises during 2008, 2010 and 2011 (Brewin et al., 2013), as part of the Research Cruises expedition programme of the Red Sea Research Center (RSRC) of King Abdullah University of Science and Technology (KAUST); 2) In vivo Lidar fluorescence data on chlorophyll concentration in the Red Sea (Barbini et al., 2004) collected as part of the Mediterranean Sea, Indian and Pacific Oceans Transect (MIPOT) oceanographic campaign between Italy and New Zealand in November 2001; 3) In situ hyperspectral absorption and attenuation data collected on a flow-through system (Boss et al., 2013; Werdell, Proctor, Boss, Leeuw, & Ouhssain, 2013) during the Tara Oceans expedition in the Red Sea in January 2010. These in situ datasets were matched in time (day) and space (closest 4 km pixel) with the OC-CCI chlorophyll products and are presented in greater detail in Brewin et al., in review (Regional ocean-color chlorophyll algorithms for the Red Sea, paper in review in *Remote Sensing of Environment*, RSE-D-14-01018). Here, these datasets were used to evaluate the performance of OC-CCI chlorophyll products in shallow and deep-water regions, in the context of satellite estimates of phytoplankton phenology.

#### 2.1.4. Sea-surface temperature remote sensing data

Level-3 mapped data of night-time (11  $\mu\text{m}$ ) sea-surface temperature (SST) from Aqua MODIS were used in this study. The 8-day climatology (period 2003–2008) at 4 km resolution product was obtained from the NASA ocean color website (<http://oceancolor.gsfc.nasa.gov>). MODIS SST datasets have been already used to monitor seasonal changes in marine temperatures in the Red Sea (Raitsos et al., 2013).

#### 2.1.5. Ocean-color remote sensing data

Version 1 of ESA Ocean Colour CCI product comprising merged and bias-corrected MERIS, MODIS and SeaWiFS data was used in this study. OC-CCI reflectance data are processed using SeaDAS atmospheric-correction algorithm for SeaWiFS and MODIS (Fu, Baith, & McClain, 1998), and POLYMER algorithm for MERIS (Steinmetz et al., 2011). The level-3 mapped chlorophyll data were acquired at 4 km and 1-day resolution from <http://www.esa-oceancolour-cci.org> for the period Sep. 1997 to Dec. 2012. The data were then re-gridded to 8-day. This resolution provides a relatively high temporal resolution and limits missing data, which are important to improve accuracy and precision in the estimation of phenological indices (Racault et al., 2012; Racault et al., 2014).

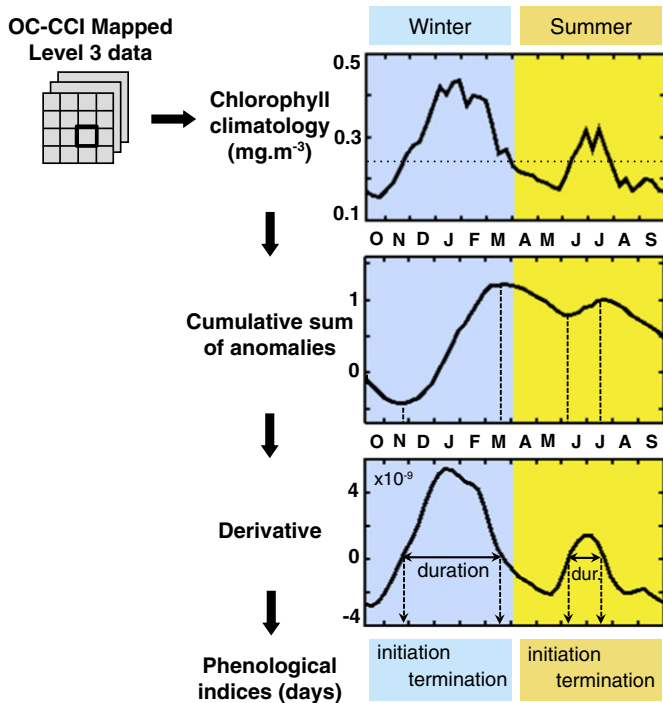
Level-3 mapped data of chlorophyll were downloaded at 8-day resolution for the period 2003–2010 for each individual sensor SeaWiFS, MODIS and MERIS from the NASA ocean color archive (<http://oceancolor.gsfc.nasa.gov/>). MODIS (SeaDAS processing version 2013.1) and MERIS (SeaDAS processing version 2012.1) chlorophyll data were downloaded at 4 km resolution, while SeaWiFS (SeaDAS processing version 2010.0) data were downloaded at 9 km resolution (i.e. the highest available resolution for the mapped product in the archive).

Chlorophyll climatologies were computed and used as follows: 1) a ~15-year climatology Sep 1997 to Dec 2012 of OC-CCI chlorophyll at 4 km and 8-day resolution was constructed and used to estimate phenological indices (this is the longest continuous ocean-color record available); 2) 8-year climatologies from Jan 2003 to Dec 2010 of OC-CCI, SeaWiFS, MODIS, and MERIS chlorophyll at 9 km and 8-day resolution were calculated and used in the comparison of data availability (this period corresponds to the time when all three satellite sensors SeaWiFS, MODIS and MERIS were coincidentally recording data; see Fig. 2 in Results and discussion section). Linear interpolation was applied to perform spatial re-gridding from 4 km to 9 km. The periods used to calculate the climatologies were chosen to maximize the data coverage (i.e. 1997–2012, see assessment in Section 2.2) and to ensure overlap between recordings of individual sensors (i.e. 2003–2010).

### 2.2. Estimation of phenological indices

The threshold criterion method proposed and implemented herein has been developed to encompass the large variability of shapes in the phytoplankton growth encountered in the Red Sea. Preliminary investigation of the characteristics of the phytoplankton seasonal cycle in the Red Sea was performed using published literature based on in situ (Genin, Lazar, & Brenner, 1995; Levanon-Spanier, Padan, & Reiss, 1979) and remote-sensing observations (Acker et al., 2008; Labiosa et al., 2003; Raitsos et al., 2013), and by direct examination of the OC-CCI chlorophyll dataset. A threshold criterion method, such as presented in Racault et al. (2012), was adopted to estimate phenological indices of timing of initiation, peak, termination and duration of phytoplankton growth during both winter and summer growing seasons (Fig. 1). First, the 8-day chlorophyll climatology (period 1997–2012) is calculated, providing complete coverage of the annual cycle at every pixel in the Red Sea (i.e. no missing data). Then, for the winter and summer seasons, the timings of initiation and termination are defined as the times when chlorophyll concentration rises above and falls below a threshold criterion of median plus 10%. The median is computed at every pixel from the annual cycle (based on the 8-day chlorophyll climatology). The same threshold criterion is used to estimate the





**Fig. 1.** Schematic representation of the method proposed to estimate phytoplankton phenological indices during winter (October to May) and during summer (June to September). The example time-series is from the pixel located at 41°E, 19°N. The chlorophyll climatology is computed based on the OC-CCI 8-day composites from Sep. 1997 to Dec. 2012. The median + 10% threshold is calculated from the climatology. The anomalies are estimated as chlorophyll minus the threshold criterion and the cumulative sum of the anomalies is calculated. The derivative of the cumulative sum of anomalies is then computed. The times when the derivative is equal to zero are used to identify the timings of initiation and termination of the phytoplankton growing periods. Note that the timing of peak is searched between the timings of initiation and termination using the chlorophyll climatology. The same threshold criterion (median + 10%, dashed line in the top panel) is applied to estimate both winter and summer growing periods. Further details on the method are provided in the main text (Section 2.2).

timings of initiation and termination during both the winter and summer seasons. The duration is calculated as the number of 8-day composites between the timings of initiation and termination. The timing of peak is detected as the time when maximum chlorophyll concentration is reached within the duration of a growing period.

Different threshold criteria were tested and limited sensitivity was observed in the estimations of timings of initiation and termination when thresholds ranged within median plus 5 to 15%. A lower threshold criterion may lead the algorithm to detect earlier (later) timings of initiation (termination) and in extreme cases, a nearly continuous growing period throughout the year may be estimated. On the other hand, a higher threshold criterion may exceed the maximum value in one of the growing periods relative to the other, with the result that the phenological algorithm may recover only one growing period.

The procedure to estimate the phenological indices is as follows (see Fig. 1 for a schematic representation). First, we subtracted the threshold criterion  $s$  from the 8-day climatology chlorophyll time-series  $Chl$ :

$$A = Chl - s. \quad (1)$$

The 8-day chlorophyll anomaly time-series is denoted  $A$ . When chlorophyll concentration is above the threshold,  $A > 0$ , and when chlorophyll concentration is below the threshold,  $A < 0$ . In the weeks preceding the commencement of the phytoplankton growing period, short pulses of chlorophyll associated with synoptic variability may be observed in the time-series. In these conditions, examination of

cumulative chlorophyll anomalies  $C$  helps to avoid spurious estimation of timing of initiation:

$$C = \sum_i A_i, \quad (2)$$

$A_i$  is the chlorophyll anomaly summed over 8-day periods  $i$ , starting at the beginning of the climatology. This method builds upon the benefits of using the cumulative sum approach (Brody et al., 2013; Greve, Prinage, Zidowitz, Nast, & Reiners, 2005), which has been shown to reduce potential “noise” or short pulses of chlorophyll data without filtering out any of the signal. Furthermore, the cumulative sum of anomalies method (Lozowski, Charlton, Nguyen, & Wilson, 1989) also allows us to identify persistent periods of chlorophyll increase above the threshold criterion  $s$ , which are used here to delineate a phytoplankton growing period. In the time-series of cumulative chlorophyll anomalies  $C$ , the growing periods appear as trends (i.e., an increasing trend when chlorophyll concentrations are above the threshold and a decreasing trend when chlorophyll concentrations are below the threshold; Fig. 1). Therefore, to estimate the timing when the trend changes direction, which corresponds to the time when chlorophyll rises above or falls below the threshold, we computed the derivative  $D$  of the cumulative anomalies of chlorophyll:

$$D = \frac{dC}{dt}. \quad (3)$$

The derivative time-series  $D$  was smoothed by applying a 3-week running average and the timings of initiation and termination were then readily estimated as the times when the derivative  $D$  was equal to zero. The phenology algorithm in the present study has been developed to estimate the main phytoplankton growing period in each season (i.e. winter and summer). If, during a season, a short pulse (<16 days) of chlorophyll reached concentrations above the threshold, then it would be detected by the algorithm but not output as a phenology metric.

The timings of initiation of phytoplankton growth were identified within two time intervals in the calendar year: 1) September to February for the winter growing period (phytoplankton growth had been demonstrated to peak between October and April, Raitso et al., 2013); and 2) March to August for the summer growing period. The time boundaries selected to identify the timings of initiation were chosen in relation to the seasonal SST regime (González Taboada & Anadón, 2014), as a mean to connect variations in chlorophyll concentration to changes in environmental conditions. Indeed, temperature has been demonstrated as an important factor involved in the regulation of phytoplankton community structure (e.g., Bouman et al., 2003) and physiological processes (e.g., Saux Picart, Sathyendranath, Dowell, Moore, & Platt, 2014). Using the cumulative sum of anomalies climatological (modulo) time-series, the timings of termination were identified in the time intervals as follows: a) for a time-series characterized by two growing periods, the first timing of termination was identified in the time interval between the timing of initiation of the first growing period and the timing of initiation of the second growing period. Then, the second timing of termination was identified between the timing of initiation of the second growing period and the timing of initiation of the first growing period; b) for a chlorophyll time-series characterized by only one growing period, the timing of termination was identified as the next date when the derivative is equal to zero (after the timing of initiation). Finally, the timing of peak was searched in the time interval between the timings of initiation and termination using the chlorophyll climatology.

An evaluation of the cumulative sum of anomalies threshold method proposed here to estimate phytoplankton phenological indices, shows that two growing periods (winter and summer) were identified in 75% of the chlorophyll time-series available, whereas a single growing

period (either in winter or in summer) was identified in the remaining 25% of the chlorophyll time-series. In the present study, the main limit of the method to detect of one to two growing periods is dictated by the shape of the annual chlorophyll time-series (i.e. whether chlorophyll concentrations rise and fall above the threshold criterion during both the winter and summer growing seasons). The advantage to use the same threshold for both the winter and summer growing periods is that the inter-seasonal variability in the estimated phenological metrics can then be compared.

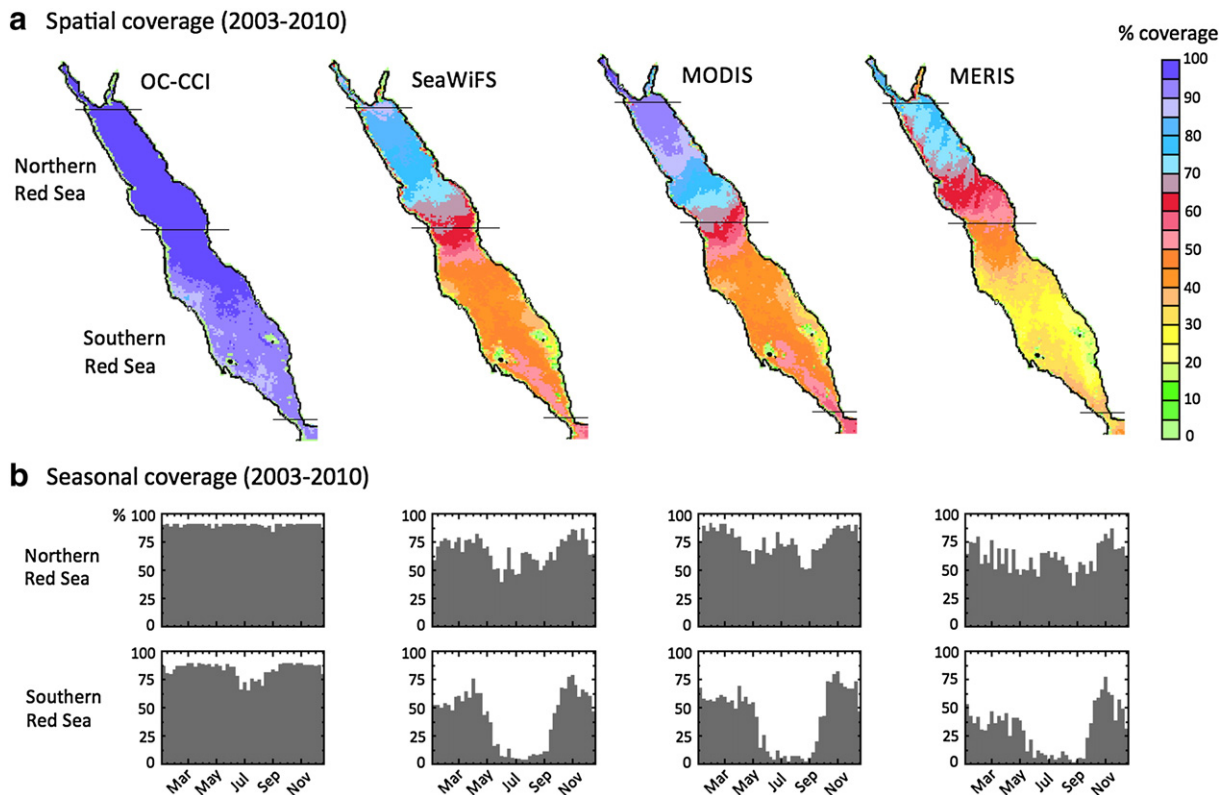
### 3. Results and discussion

#### 3.1. Availability and performance of ocean-color remote-sensing observations in the Red Sea

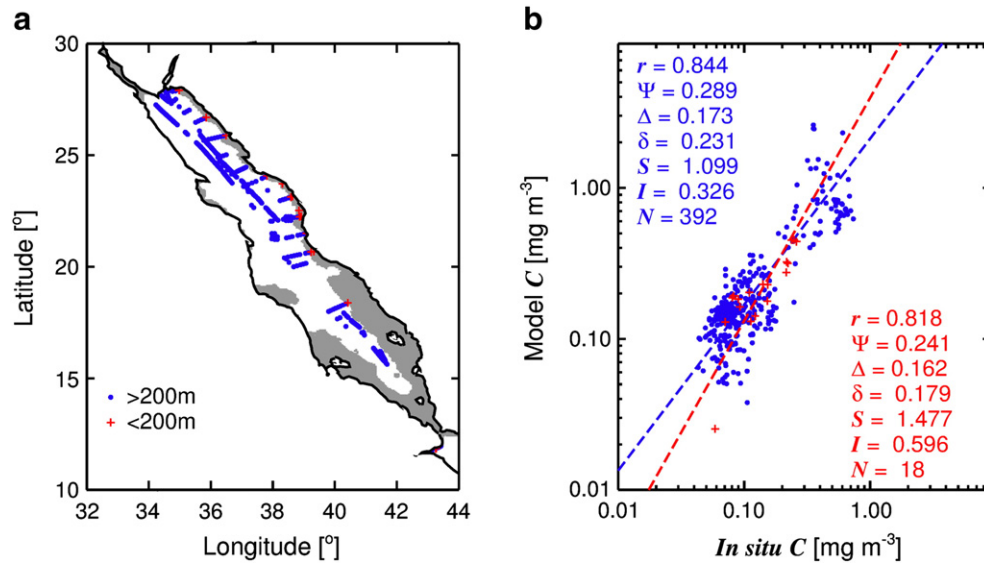
Ocean-color observations from OC-CCI show a significant increase in spatial and temporal coverage compared with the results from single sensors SeaWiFS, MODIS and MERIS over the period 2003–2010 (Fig. 2). The climatological coverage of OC-CCI chlorophyll data is  $\geq 75$ –80% throughout the year in the entire Red Sea basin, whereas from single sensors, in the Northern Red Sea, the coverage is around 65% (or less) throughout the year (with the exception of MODIS showing a coverage of nearly 75% during the winter months Nov–Jan). In the southern Red Sea, the climatological coverage from single sensors is around 50–55% (or less) in winter months, and this coverage drops to nearly 0% in the summer months (Jun–Sep). The extremely low data coverage in the southern Red Sea region reported from the single sensors is due to cloudy conditions and transport of dust particles in the atmosphere from the surrounding deserts during the northeast and southwest monsoon periods. In addition, the typical extreme heat during the summer months may induce hazy conditions. Hence, the region is particularly challenging for atmospheric-correction algorithms

when processing MERIS, MODIS and SeaWiFS images, and pixels are flagged as of low quality in the SeaDAS atmospheric correction (Acker et al., 2008; Raitos et al., 2013). The large increase in ocean-color data coverage especially in the southern regions and summer months observed in the OC-CCI dataset is due to the application of POLYMER algorithm to MERIS imagery (Steinmetz et al., 2011). The POLYMER algorithm is based on two models: 1) a polynomial atmospheric model representing the atmospheric scattering from large particles, aerosols, semi-transparent clouds and residual sun glint; and 2) a bio-optical ocean water reflectance model. These two models render the POLYMER algorithm robust to the effects of sun glint, semi-transparent clouds, and atmospheric dust aerosols, which occur during the monsoon, particularly in the Southern Red Sea region. This has led to a large increase in ocean-color data coverage in the OC-CCI chlorophyll data and permitted us for the first time to resolve the seasonal cycle of phytoplankton in summer months in the southern Red Sea. Finally, increased data coverage is highly beneficial, particularly to improve accuracy and precision in the estimation of phenological indices (Racault et al., 2014). The chlorophyll climatology estimated from the extended OC-CCI dataset (1997–2012) has permitted us to extract a complete annual cycle (i.e. no missing data in the climatology) at every pixel in the Red Sea.

Prior to analyzing phytoplankton seasonal cycle and estimating phenological indices, the performance of the OC-CCI chlorophyll retrieval algorithm was evaluated in shallow, reef-bound coastal waters of the Red Sea. The relationship between  $\log_{10}$ -transformed chlorophyll from in situ and satellite measurements was statistically significant ( $r = 0.82$  with  $p < 0.05$  and  $\Psi = 0.24$ , where  $r$  is the Pearson correlation coefficient,  $p$  relates to the significance of this correlation ( $p < 0.05$  mean statistically significant) and  $\Psi$  refers to the root-mean-square-error) in coastal and/or shallow reef waters (i.e. waters with a depth  $< 200$  m; Brewin et al., 2010; Uitz, Claustre, Morel, & Hooker, 2006), and comparable to the results in



**Fig. 2.** Spatial and seasonal coverage of chlorophyll data for the period 2003–2010 for the merged product ESA OC-CCI (data processed using POLYMER atmospheric correction algorithm for MERIS, and using SeaDAS for SeaWiFS and MODIS), and the sensors SeaWiFS, MODIS and MERIS (data processed using SeaDAS atmospheric correction algorithm for all three sensors). a) Spatial coverage is computed as the relative percent number of 8-day chlorophyll composites available during 2003–2010; b) Seasonal coverage corresponds to the relative percent number of chlorophyll data available for each 8-day period during 2003–2010 over the Northern and Southern Red Sea regions.



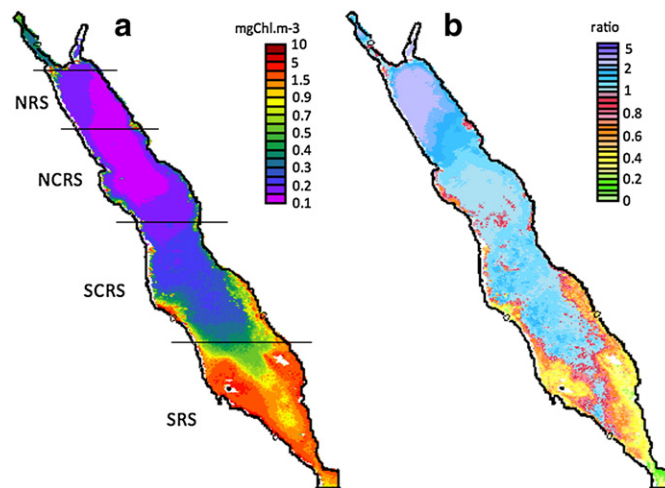
**Fig. 3.** a) Distribution of in situ and satellite match-up data used in the study. Red cross indicates match-up data located in water shallower than 200 m and blue dots indicate match-up data located in waters deeper than 200 m (the 200 m isobaths is used to delineate shallow reef-bound coastal waters, Brewin et al., 2010; Uitz et al., 2006); b) Scatter plots of the match-ups between in situ and satellite-derived chlorophyll (C) in the Red Sea using the OC-CCI algorithm for shallow waters (<200 m in red) and deeper waters (>200 m in blue). Univariate statistical tests were used and applied to  $\log_{10}$ -transformed chlorophyll data following Brewin et al. (2013). This included: the Pearson correlation coefficient ( $r$ ); the root-mean-square-error ( $\Psi$ ); the average bias between model and measurement ( $\delta$ ); the centre-pattern (or unbiased) root mean square error ( $\Delta$ ); the slope ( $S$ ) and intercept ( $I$ ) of a Type-2 regression; and the number of retrievals ( $N$ ).

deeper open-ocean waters ( $r = 0.84$  and  $\Psi = 0.29$ , Fig. 3). Previous comparison of match-ups of chlorophyll concentrations, measured in situ and from satellites, over extensive areas of the Red Sea indicated similar results to other standard chlorophyll products in the global ocean (Brewin et al., 2013). In optically complex waters, remotely-sensed chlorophyll data have known limitations, especially as yellow substances and non-algal particulate matter may not co-vary in a predictable manner with chlorophyll (Morel & Gentili, 2009). These factors may influence the retrieval of chlorophyll concentration, generally resulting in an overestimation (Blondeau-Patissier, 2009). However, not all high chlorophyll values in shallow waters are necessarily biased, as large coral reef complexes may be sources of either nutrients or chlorophyll-rich detritus that enhance phytoplankton production near the reefs (Genin et al., 2009;

Labiosa et al., 2003; Levanon-Spanier et al., 1979; Sakka, Legendre, Gosselin, Niquil, & Delesalle, 2002; Wild et al., 2008; Wyatt et al., 2010; Yahel et al., 1998). The performance of the OC-CCI chlorophyll data in the Red Sea (both coastal and deep-waters, Fig. 3) is comparable with the performance of standard chlorophyll algorithms using large globally-representative datasets (see Brewin et al., In press, their Fig. 4), lending support to the use of the OC-CCI chlorophyll data to estimate phytoplankton phenology in the Red Sea.

### 3.2. Phytoplankton seasonality in the Red Sea

Based on the remotely-sensed observations from ESA OC-CCI, the northern provinces of the Red Sea (NRS and NCRS) exhibit



**Fig. 4.** Phytoplankton biomass in the Red Sea. a) Annual mean (climatology 1997–2012) chlorophyll concentration (in  $\text{mg.m}^{-3}$ ) with the four provinces partitioning (Raitos et al., 2013) overlaid: northern-Red Sea (NRS), north-central Red Sea (NCRS), south-central Red Sea (SCRS), and southern Red Sea (SRS); b) ratio of the mean chlorophyll concentration during winter SE monsoon (October to May) to the mean chlorophyll concentration during summer NW monsoon (June to September). When the ratio is greater than 1, the chlorophyll during the winter period is higher than the mean chlorophyll during the summer period.

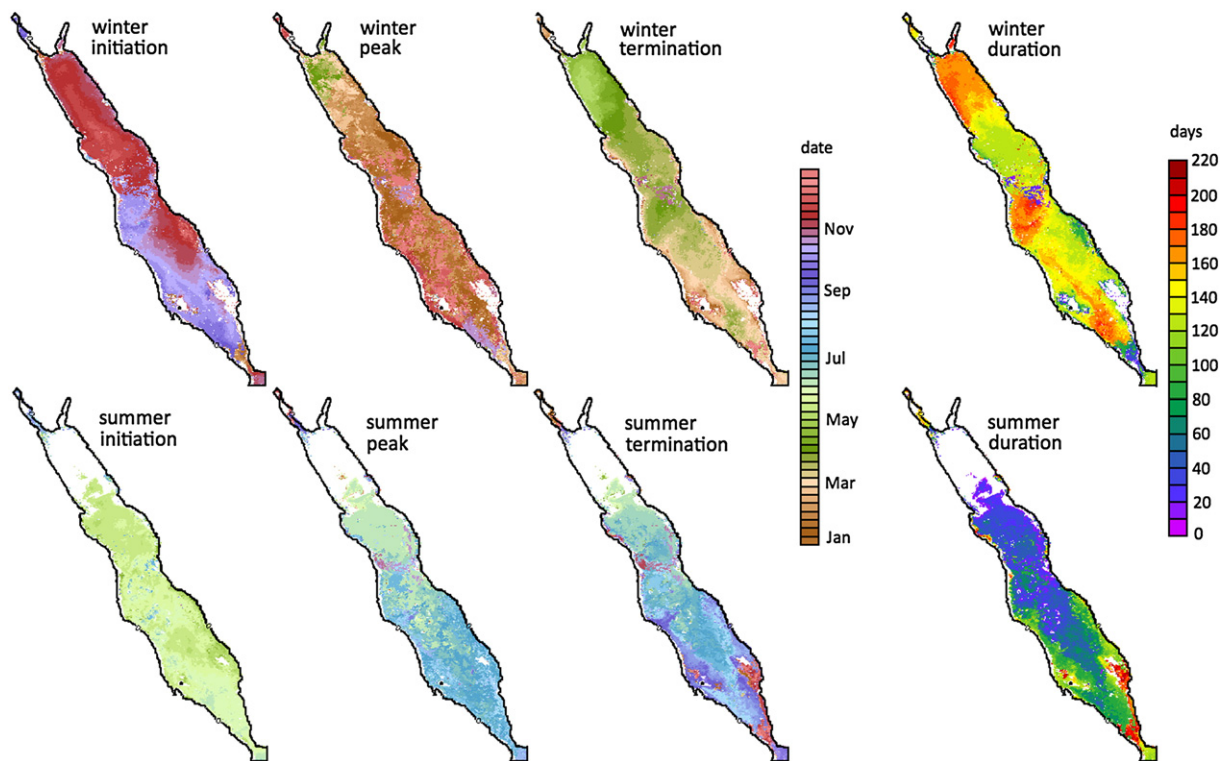


lowest chlorophyll concentrations of  $0.2 \text{ mg.m}^{-3}$  on average, which is characteristic of oligotrophic waters (Fig. 4a) and the southern provinces exhibit higher chlorophyll concentrations — an average of  $0.4 \text{ mg.m}^{-3}$  in SCRS, and  $1.7 \text{ mg.m}^{-3}$  in SRS. This increase in chlorophyll concentration from north to south is consistent with previous observations in the basin based on remotely-sensed chlorophyll concentration (Acker et al., 2008; Raitos et al., 2013) as well as in situ chlorophyll and total pigment concentrations (McGill & Lawson, 1966 in Halim, 1969; Weikert, 1981). In the latter studies, phytoplankton pigment concentrations have been further reported to display a marked contrast between the two monsoon periods. Given the unique coverage provided by the OC-CCI dataset in the Red Sea region (Fig. 2), we investigated the ratio between mean chlorophyll concentrations during the winter SE monsoon period (October to May) and the summer NW monsoon period (June to September). Overall, it appears that in most of the open waters of the basin, average chlorophyll concentrations are higher during the winter compared with those during the summer (i.e. ratio greater than one; Fig. 4b). The highest ratio is found in the northern Red Sea province with values of nearly 2.5. An interesting result is that the ratio tends to be lower than one (or close to) in the waters adjacent to large reef complexes along the coast of the Red Sea (see Fig. 6 for position of coral reefs). This phenomenon is particularly apparent in the southern provinces of the Red Sea, where ratio values as low as 0.3 are observed, indicating that chlorophyll concentrations in reef-bound coastal waters are higher during the summer period compared with the winter period. Higher (or nearly equal) chlorophyll concentrations in reef-bound coastal waters during the warm period (Jun–Sep) have also been reported previously from in situ measurements in the Gulf of Aqaba (Levanon-Spanier et al., 1979, their Fig. 9; Yehel et al., 1998, their Table 2).

### 3.2.1. Phenological indices in open waters of the Red Sea

During winter, phytoplankton seem to flourish initially during the month of October in the southern part of the Red Sea (Fig. 5). The timing

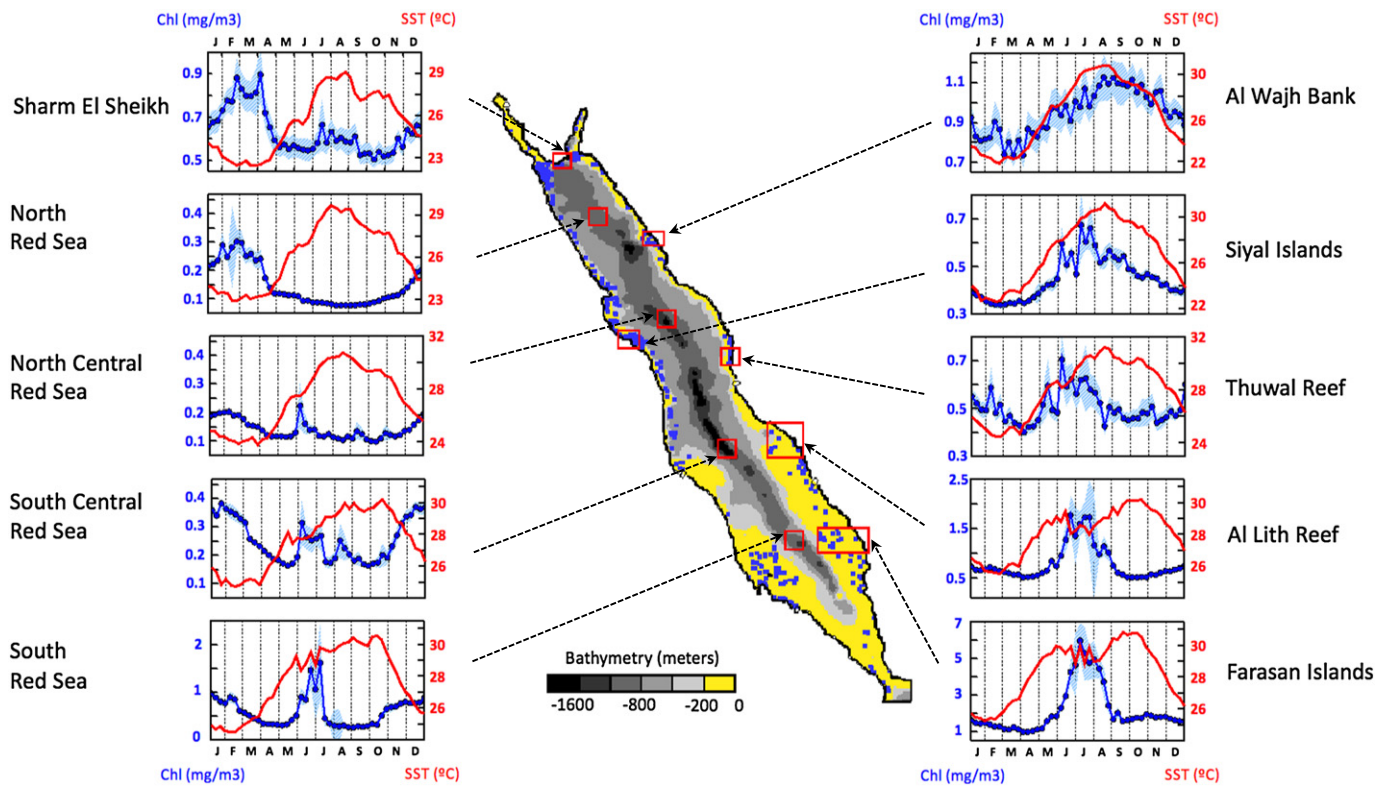
of initiation is slightly delayed in the northern part, with the winter phytoplankton growing period commencing in November. Chlorophyll concentrations peak between mid-December to mid-January throughout the entire Red Sea, except in the northwestern region of the NRS, which shows the latest peak timing at the end of March, and the south-eastern region of the SRS, where the peak timing appears to occur earlier, around the beginning of October; coinciding with the intrusion of colder nutrient-rich waters from the Gulf of Aden (Churchill, Bower, McCorkle, & Abualnaja, *In press*; Mill & Post, 1981; Sofianos & Johns, 2003). The termination of the phytoplankton winter growing period occurs generally earlier in the southern part of the Red Sea (termination in March) compared with that in the northern part of the Red Sea (termination in April). These patterns of chlorophyll concentrations can be attributed to distinct physical regimes occurring in the Red Sea during winter: 1) in the northern part, nutrients are mainly supplied through convective vertical mixing (Labiosa et al., 2003; Mill & Post, 1981; Sofianos & Johns, 2003; Triantafyllou et al., 2014), which can sustain relatively long phytoplankton growing periods between 140 to 180 days; and 2) in the southern part, nutrient-rich waters are transported from the Gulf of Aden into the Red Sea, through horizontal advection driven by south-easterly winds (Churchill et al., *In press*; Mill & Post, 1981; Sofianos & Johns, 2003; Yao, Hoteit, Pratt, Bower, Zhai, et al., 2014), sustaining phytoplankton growth for a duration of approximately 120 to 160 days (Fig. 5, Table 1). In the central region of the Red Sea (around  $20^\circ\text{N}$ ), an intermediate zone of wind convergence develops from October to April (Morcos, 1970), with the winds blowing from the north in the SSE direction, and from the south in the NNW direction (Patzert, 1972). These winds and the associated surface circulation trigger an initiation of the winter phytoplankton growing period (Raitos et al., 2013), which is apparent in October (Fig. 5). The central region is further characterized by a north-south SST gradient (Yao, Hoteit, Pratt, Bower, Zhai, et al., 2014) and displays phytoplankton growing periods longer than 140 days in the central-west domain (Fig. 5, Table 1).



**Fig. 5.** Phenology of phytoplankton in the Red Sea. Winter (upper panels) and summer (lower panels) phytoplankton growing periods. White color indicates that no phytoplankton growing period was detected based on the threshold criterion method (see Materials and methods section for further information). The short duration ( $<20$  days) reported in winter in the central part of the Red Sea are due to basin-wide anti-cyclonic eddies forming in the region (Johns, Jacobs, Kindle, Murray, & Mike, 1999), that generate short and particularly intense eddy-driven phytoplankton blooms, which may locally mislead the phenology detection algorithm implemented in this study (however, this feature is apparent for a small number of pixels and does not impede on the analysis of the phenology of phytoplankton in the entire basin).







**Fig. 6.** Phytoplankton and sea-surface temperature seasonality in open and reef-bound coastal waters in the Red Sea. Central map: sea-floor elevation in the Red Sea with the distribution of coral reefs shown as blue squares (source: Global Distribution of Coral Reefs dataset, 2010). Side plots: Climatologies of sea-surface temperature (SST, in red) and chlorophyll concentration  $\pm 1$  SE (in blue and shaded blue, respectively) at four open water sites and six reef complexes. The standard error (SE) represents the standard deviation of the error of the chlorophyll climatological mean calculated over the period 1997–2012.

suggests that the influence of coastal upwelling on phytoplankton dynamics may be limited to the Gulf of Aqaba and to the coastal waters adjacent of the entrance of the Gulf.

The large and prominent winter phytoplankton growing period compared with the limited or suppressed summer phytoplankton growth estimated in Sharm El Sheikh reef area is likely to have important implications for trophic interactions and pelagic-benthic coupling, which may turn to be quite different from what is occurring at other reef systems (see other case studies in the sections below, for which reef systems present equal or more prominent phytoplankton growth in summer compared with winter). Ecological studies have reported asynchronous spawning patterns for *Acropora* coral species in the Gulf of Aqaba (Shlesinger & Loya, 1985), while at other Red Sea reef complexes, these coral species appear to display synchronous spawning patterns (Hanafy, Aamer, Habib, & Baird, 2010; Bouwmeester, Khalil, De La Torre, & Berumen, 2011; Bouwmeester et al., 2014).

**3.2.2.2. Siyal Islands.** The Siyal islands reef area is located in the coastal waters of Sudan on the west side of the Red Sea around the latitudes 22.4–22.9°N in the north central Red Sea province. In the deeper central waters of this province, chlorophyll concentration appears higher in the winter period compared to the summer period (Fig. 4b). In contrast, the waters located in the vicinity of the Siyal Islands reef complex exhibit higher chlorophyll concentrations in summer (Fig. 4b; average ratio winter to summer chlorophyll for the Siyal Islands reef area is 0.77).

During the summer period, phytoplankton growth period appears to commence between 17th and 24th May on average in the reef area, and to reach peak chlorophyll concentration ( $0.8 \text{ mgChl.m}^{-3}$ ) between 12 and 19th July (Table 1). Timing of the chlorophyll peak precedes the SST maximum by approximately one and a half month (Fig. 6). This

summer phytoplankton growing period may last ~140 days (nearly four months and a half), which is the longest duration of the six reefs examined in the present study.

During the winter period, phytoplankton growth is estimated to commence during 1st to 8th November, which is approximately one month after the estimated termination of the summer phytoplankton growth. Winter maximum chlorophyll concentration of  $0.4 \text{ mg.m}^{-3}$  is reached between 11th and 18th December (Table 1). At the Siyal Islands reef complex, winter phytoplankton growth is characterized by lower maximum chlorophyll concentration (nearly 2 times lower) and shorter duration (2 months shorter) compared with the summer growing period (Fig. 6).

Sudanese offshore coral reefs appear to harbor a significantly greater biomass than other reefs in the Red Sea (Spaet, 2014). These reefs seem to hold the only significant and persistent populations of large apex predators such as sharks and coral-groupers, although the reef structure and benthic communities appear to be similar to those of offshore reefs directly across the Red Sea (e.g., near Thuwal, see below) (M. L. Berumen, unpublished data). It remains unclear whether the longer duration of the summer phytoplankton growing period (i.e. longer period of food availability) has an influence on the trophic interactions at Siyal Islands reefs, but this warrants further investigation into the structure of the food web in these reefs relative to other Red Sea reefs.

**3.2.2.3. Al Wajh Bank.** The Al Wajh Bank reef complex is located on the east coast between the latitudes 25.4 and 25.8°N, at the boundary between the north Red Sea and the north central Red Sea provinces. It is a significant area for ecological conservation. The Al Wajh Bank consists of a large, shallow lagoon area nearly completely enclosed by reefs with very few openings to deeper water. Although tidal range in the Red Sea is generally very low, the volume of water contained inside this lagoon,

combined with the few channels opening to the adjacent deeper water, generates strong tidal currents in and out of the lagoon. At this reef complex, high chlorophyll concentrations had been previously reported (Acker et al., 2008). We further observe that chlorophyll concentrations are nearly equal or higher in summer compared with winter (Fig. 4b; average ratio winter to summer chlorophyll for the Wajh bank reef area is 0.98). The summer chlorophyll concentration follows the SST regime closely (Fig. 6).

During the warm period in summer, the timing of initiation of the phytoplankton growing period occurs between 4 and 11th July, which is noticeably later compared with the other reefs examined in the present study (which all display an apparent common timing of initiation between 17 and 24th May; Table 1). Maximum peak of  $1.5 \text{ mgChl.m}^{-3}$  is reached later (compared with the other reefs), at the end of the summer months during the period 14th to 21st September, when SST is warmest (Fig. 6, Table 1). The summer phytoplankton growing period appears as one of the longest with a duration of ~130 days (approximately four months).

High chlorophyll concentration is also observed in the winter period, reaching maximum concentration of  $0.9 \text{ mgChl.m}^{-3}$  between 9 and 16th January. Establishing a clear delineation between summer and winter phytoplankton growing periods at this reef complex was not straightforward using a threshold criterion method. In fact, the winter growing period is estimated to commence between 17 and 24th November, which is just one week after the estimated termination of the summer growing period in 9–16th November (Table 1). Winter chlorophyll concentration remains above the threshold criterion for 110 days (approximately three and a half months). Consequently, phytoplankton tends to be characterized by continuously elevated chlorophyll concentration (> median plus 10%) from mid-July until mid-February (Table 1, Fig. 6).

Many anecdotal reports suggest that the Al Wajh Bank is an important site for several types of megafauna, such as manta rays. Mantas are directly dependent on planktivorous food sources, and in the Red Sea appear to be reasonably site-restricted (Braun, 2013; Braun, Skomal, Thorrold, & Berumen, 2014). Hence, the hydrodynamic features and extended phytoplankton growing season observed at the Al Wajh Bank could have follow-on effects to larger zooplankton communities and subsequently to planktivorous filter feeders such as mantas. Further investigation will be required to elucidate this trophic linkage.

**3.2.2.4. Thuwal Reefs.** The complex of Thuwal reefs is situated between  $21.9$  and  $22.4^\circ\text{N}$  within the north central Red Sea province. In the waters adjacent to the reefs, a main winter and a secondary summer phytoplankton growing period are observed (Table 1, Fig. 6d). The amplitude of the phytoplankton blooms shows limited difference between winter ( $0.5 \text{ mgChl.m}^{-3}$ ) and summer ( $0.9 \text{ mgChl.m}^{-3}$ ), with winter to summer chlorophyll ratio of 1.1 on average in the area (Fig. 4b). However, the duration of the growing period is nearly double in winter, estimated to last ~120 days (approximately four months) compared to summer, estimated to last ~65 days (just over two months; Table 1).

During the winter period, phytoplankton growth period appears to commence during 16–24th October and chlorophyll concentration tends to peak between 1st and 8th January, when SST is cooler and nutrients are brought up to the surface by vertical mixing from the deeper water in the central part of the basin. During the summer period, phytoplankton growth shows an apparent start between 17 and 24th May, similar to most of the reefs examined in the present study, but the peak timing appears to occur noticeably earlier between 18 and 25th June (Table 1). Hence, in the area, summer chlorophyll peak precedes summer SST maximum by approximately two months (Fig. 6).

The Thuwal reefs complex is the location of a large and increasing number of research projects on coral reef ecosystems (Mervis, 2009). Understanding the local phytoplankton phenological cycle will provide useful information for a wide range of in situ experiments related to fishing practices (e.g., Jessen, Roder, Villa Lizcano, Voolstra, & Wild,

2013), observational studies of reef grazers (e.g., Khalil, Cochran, & Berumen, 2013) and further investigations on biogeochemical responses following coral mass spawning (e.g., Bouwmeester et al., 2014).

**3.2.2.5. Al Lith Reefs.** The Al Lith reef complex is located in the south central Red Sea province within the latitude band  $19.2$ – $20.2^\circ\text{N}$ . It forms the northern end of the extensive Farasan Banks reef complex. Al Lith is emerging as a system of reefs with great importance on the eastern Red Sea coast, with value in both economic and research contexts. The site hosts manta rays (Braun et al., 2014) and the only known whale shark (*Rhincodon typus*) aggregation site in the Red Sea (Berumen, Braun, Cochran, Skomal, & Thorrold, 2014). Whale sharks, like manta rays, are large planktivorous fishes, and thus dependent on phytoplankton–zooplankton seasonal cycles.

During the summer period, the phytoplankton bloom begins around 17–24th May (as in most of the reefs examined in the present study; Table 1), and reaches maximum chlorophyll concentration of  $2.5 \text{ mg.m}^{-3}$  in the period 12–19th July, when the increase in SST shows an apparent pause (Table 1, Fig. 6). This slow down in SST warming is likely to be caused by anti-cyclonic eddies, which form in the region and generally show strongest activity during summer (Fanning, Carder, & Betzer, 1982; Johns, Jacobs, Kindle, et al., 1999; Mill & Post, 1981; Yao, Hoteit, Pratt, Bower, Köhl, et al., 2014; Zhan et al., 2014). The eddies may provide a source of nutrients through re-suspension and thus, sustain phytoplankton growth for a duration of ~100 days (just over three months; Table 1).

A winter phytoplankton growing period is also observed at Al Lith reefs, starting to develop during the period 9–16th November. Maximum winter chlorophyll concentration of  $0.8 \text{ mg.m}^{-3}$  is reached during the period 9–16th January when SST is lowest (Fig. 6). The winter phytoplankton growing period lasts for ~90 days (approximately three months); possibly sustained by an inflow of cold nutrient-rich waters from the Gulf of Aden (Churchill et al., In press; Yao, Hoteit, Pratt, Bower, Zhai, et al., 2014).

Al Lith reefs appear to support two phytoplankton growing periods of equivalent duration (Table 1) but with very different amplitudes (Table 1; and Fig. 4b, ratio of winter to summer chlorophyll concentrations is 0.65 on average in the area). These phytoplankton phenological characteristics may play an important role in the aggregation of large planktivorous fish such as manta rays and whale sharks. In another reef ecosystem of the Great Barrier in Australia, the aggregation and abundance of whale sharks (*Rhincodon typus*) have been reported to occur during increased phytoplankton productivity and mass coral spawning (Taylor, 1996; Wilson, Taylor, & Pearce, 2001). The phenological indices presented at Al Lith reefs may help to elucidate trophodynamic links between phytoplankton availability and megafauna abundance.

**3.2.2.6. Farasan Islands.** The Farasan Islands complex is one of the most extensive island groups in the Red Sea, and is located in the latitude bands  $16.4$ – $17.1^\circ\text{N}$  in the south Red Sea province. Many of the islands possess fringing coral reefs that show a notably different reef community composition compared with reefs from any of the other sites mentioned above. These reefs tend to have much more turbid water and a greater occurrence of macroalgae, both generally accepted as indications of greater local primary productivity. The summer peak chlorophyll concentration of  $7.1 \text{ mg.m}^{-3}$  observed in the Farasan Islands is among the highest of the Red Sea (Fig. 6).

The phytoplankton phenological characteristics in the coastal waters of the Farasan Islands are very similar to those observed in the Al Lith reefs area. During the summer NW monsoon, the phytoplankton growing period starts in 17–24th May (same as Al Lith reef) and peak chlorophyll concentration is reached during 20–27th July (which is one week later than in Al Lith reef; Table 1). The duration of the summer growing period is longer (compared with the Al Lith reef group), lasting ~125 days (just over three months and a half; Table 1).

During the winter SE monsoon, chlorophyll concentration also shows a seasonal increase, albeit of markedly lower magnitude compared with the summer peak (i.e. ratio of winter to summer chlorophyll concentrations is 0.45 on average in the area, indicating that the summer chlorophyll concentration are more than double that of the winter (Fig. 4b)). The timing of initiation of the winter phytoplankton growing period is around 30th Sep to 7th October and maximum chlorophyll concentration of  $2.1 \text{ mg} \cdot \text{m}^{-3}$  is generally reached during the period 25th November to 2nd December. The winter timings of initiation and peak at the Farasan Islands are nearly one month earlier than the timings of growing period observed in Al Lith reefs. Furthermore, the duration of the winter growing period is also longer (compared with Al Lith reefs), lasting ~90 days (approximately three months). Although this area is shallow (and satellite chlorophyll algorithms may tend to overestimate in such regions, Fig. 2), this province is also the most strongly influenced by the inflow of cold waters from Gulf of Aden (Sofianos & Johns, 2003; Yao, Hoteit, Pratt, Bower, Zhai, et al., 2014).

In terms of population genetics, the Farasan Islands reef habitats appear to be substantially different from the other reef areas considered in this study. Although very few studies have considered the patterns of gene flow between the Farasan Islands and numerous other sites in the broader Red Sea, the greater productivity observed in the Farasan Islands and more broadly over the southern Red Sea province is likely to have profound implications for larval connectivity. Indeed, using chlorophyll concentration as a proxy for an environmental gradient, recent work on patterns of gene flow in anemonefish and sponges species indicate that the Farasan Islands are somewhat distinct genetically from sites in the Red Sea further north (Giles, 2014; Nanninga, Saenz-Agudelo, & Berumen, 2014). Further investigations on these population gene flow patterns may be carried out in relation with their biophysical habitats, which could be characterized statistically using phenological indices.

#### 4. Conclusions and Perspectives

Phenological indices of timings of initiation, peak, termination and duration of the phytoplankton growing period estimated from ocean-color remote-sensing provide objective metrics to quantify phytoplankton seasonality in coral reef ecosystems. The phenological algorithm proposed in this study, based on a relative threshold method, is particularly relevant, as it allows us to encompass a broad diversity of shapes and concentration spectrum in chlorophyll seasonal cycles such as observed in open and reef-bound coastal waters in the Red Sea (Fig. 6). Specifically, the algorithm has been developed to resolve two phytoplankton growing periods per year. At each individual pixel, the winter and summer (if any) growing periods have been estimated relative to a unique threshold criterion, enabling inter-seasonal comparison of phenological indices. Furthermore, the index of duration of growing period (x-axis, with dimension [T]) carries information on phytoplankton availability that is conventionally contained in the measure of chlorophyll concentration (y-axis, with dimensions  $[\text{M L}^{-3}]$ ). Thus, phenological indices bear the significant advantage to be robust to errors in absolute values of chlorophyll, which may arise from standard (i.e. based on globally-representative relationships) and sensor-specific chlorophyll retrieval algorithms, particularly sensitive in the optically complex waters of the Red Sea. However, notwithstanding the importance of precision and accuracy of the remote-sensing chlorophyll retrieval algorithm, the OC-CCI chlorophyll product was tested here for open and reef-bound coastal waters of the Red Sea and showed comparable performance with standard global-ocean chlorophyll retrieval algorithms (Fig. 2). In addition, the large gain in the number of chlorophyll observations during summer months demonstrated here with the usage of the POLYMER atmospheric correction algorithm (Fig. 3), has permitted us to provide the first comprehensive investigation of the phenology of phytoplankton in the Red Sea during the winter and summer periods.

Open waters of the northern Red Sea province are generally characterized by one main growing period during the winter, between December and April, which is driven by convective mixing. In contrast, open and deeper waters located in the central and south Red Sea provinces display two distinct growing periods: a primary growing period in winter, lasting approximately 4–5 months, and a secondary growing period of shorter duration in summer, lasting approximately 1–2 months. In these provinces, the winter growing period commences in December and finishes in March, and the summer growing period is estimated to begin in June and end in August. The winter growing period generally reaches higher amplitude, while the summer growing period is characterized by relatively low chlorophyll concentration (except in the southern shallow waters of the Red Sea, see below). The reef-bound coastal waters in the central and southern provinces of the Red Sea display different chlorophyll seasonality (compared to the open waters): highest chlorophyll concentrations are reached in summer (up to five times higher than in the winter) and the timing of termination of the summer growing period appears to occur in September–October (which is one to two months later compared with the open waters of the central region, occurring in July, Fig. 5). The latter feature is also observed in the duration of the summer growing period, which is characterized by an apparent duration of 100 days or longer in the waters adjacent to the reefs compared with 50 days or shorter in the open and deeper water regions of the Red Sea. Thus, in reef-bound coastal waters, chlorophyll concentration appears to display larger peak in summer (relative to the winter peak), when SST is warmest and stratification is persistent (and nutrient supply from water circulation is likely to be minimum). This paradox suggests that nutrient sources from sediments (perhaps airborne deposition) and/or other organic compounds from coral reef complexes may be playing a considerable role in the enhancement of phytoplankton growth, and further investigations are required to shed light on this potential two-way mutual feedback mechanism between coral reefs and phytoplankton growth (Wild et al., 2008).

Monitoring and comprehensive analysis of phytoplankton seasonality, together with observations of reef fish spawning activity and larval survival, will allow us to improve our understanding of reef dwellers' recruitment strategies (Johannes, 1978) and to assess their sensitivities to environmental and climatic conditions (e.g., Wilson et al., 2001). Furthermore, the quantitative metrics provided by phytoplankton phenological indices should provide useful metrics to help us improve our understanding of coral reef biogeochemical pelagic-benthic coupling, aggregation of megafauna, and population gene flows (e.g., Nanninga et al., 2014). The phenological method proposed in this study may be extended to monitor inter-annual variability of phytoplankton seasonality in the Red Sea and in other reef ecosystems in the world oceans. It is anticipated that further studies - using advanced algorithms will enable serial assessment of the sensitivity of phytoplankton to environmental and climatic conditions (Raitos et al., in press) and the possible impact on marine trophic interactions. This essential information will support the development of conservation plans and responsible stewardship of coral reef ecosystems (McCook et al., 2009; McKinnon, Williams, & Young, 2014).

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through the ESA LearnEO! online educational resources at: <http://www.learn-eo.org/lessons/111/>.

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