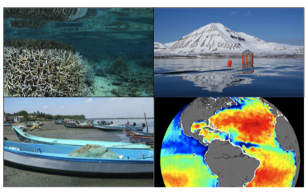


Salinity from Space Unlocks Satellite-Based Assessment of Ocean **Acidification**

Peter E. Land,**,† Jamie D. Shutler,‡ Helen S. Findlay,† Fanny Girard-Ardhuin,§ Roberto Sabia,^{||} Nicolas Reul,§ Jean-Francois Piolle,§ Bertrand Chapron,§ Yves Quilfen,§ Joseph Salisbury,^{||} Douglas Vandemark,^{||} Richard Bellerby,^{||} and Punyasloke Bhadury

 $^{^{}abla}$ Department of Biological Sciences, Indian Institute of Science Education and Research-Kolkata, Mohanpur 741 246, West Bengal India



Approximately a quarter of the carbon dioxide (CO₂) that we emit into the atmosphere is absorbed by the ocean. This oceanic uptake of CO₂ leads to a change in marine carbonate chemistry resulting in a decrease of seawater pH and carbonate ion concentration, a process commonly called "Ocean Acidification". Salinity data are key for assessing the marine carbonate system, and new space-based salinity measurements will enable the development of novel space-based ocean acidification assessment. Recent studies have highlighted the need to develop new in situ technology for monitoring ocean acidification, but the potential capabilities of space-based measurements remain largely untapped. Routine measurements from space can provide quasi-synoptic, reproducible data for investigating processes on global scales; they may also be the most efficient way to monitor the ocean surface. As the carbon cycle is dominantly controlled by the balance between the biological and solubility carbon pumps, innovative methods to exploit existing satellite sea surface temperature and ocean color, and new satellite sea surface salinity measurements, are needed and will enable frequent assessment of ocean acidification parameters over large spatial

1. INTRODUCTION

Each year global emissions of carbon dioxide (CO₂) into our atmosphere continue to rise. These increasing atmospheric concentrations cause a net influx of CO2 into the oceans. Of the roughly 36 billion metric tons of CO₂ that is emitted into our

atmosphere each year, approximately a quarter transfers into the oceans. This CO₂ addition has caused a shift in the seawater carbonate system, termed ocean acidification (OA), resulting in a 26% increase in acidity and a 16% decrease in carbonate ion concentration since the industrial revolution.² Recently there has been recognition that this acidification is not occurring uniformly across the global oceans, with some regions acidifying faster than others.^{3,4} However, the overall cause of OA remains consistent: the addition of CO₂ into the oceans, and as such, it remains a global issue. Continual emissions of CO2 into the atmosphere over the next century will decrease average surface ocean pH to levels which will be deleterious to many marine ecosystems and the services they provide.⁵

While the seawater-carbonate system is relatively complex, two parameters have been suggested as pertinent to the monitoring and assessment of OA through time and space. These are pH (the measure of acidity) and calcium carbonate (CaCO₃) mineral saturation state, with aragonite generally considered to be an important CaCO₃ mineral to be monitored because of its relevance to marine organisms (e.g., corals) and its relative solubility. Thermodynamically, CaCO₃ is stable when the saturation state (an index of the concentrations of calcium and carbonate ions) is greater than one and becomes unstable when seawater becomes undersaturated with these ions (saturation <1). While there is significant variability between types of organism, there is ample experimental evidence that many calcifying organisms are sensitive to OA,6 and that thresholds exist below which some organisms become stressed and their well-being and existence becomes threatened. Increasingly evidence suggests that the physiology and behavior of calcifying and noncalcifying organisms can be impacted by increasing OA,8 with cascading effects on the food chain and protein supply for humans,³ and alterations to the functioning of ecosystems and feedbacks to our climate.9

In 2012 the Global Ocean Acidification Observing Network (GOA-ON, www.goa-on.org) was formed in an attempt to bring together expertise, data sets and resources to improve OA

Published: January 8, 2015



1987

[†]Plymouth Marine Laboratory, Prospect Place, The Hoe, Plymouth PL1 3DH, U.K.

[‡]University of Exeter, Penryn Campus, Cornwall TR10 9FE, U.K.

[§]Institut Francais Recherche Pour ĹExploitation de la Mer, Pointe du Diable, 29280 Plouzané France

Telespazio-Vega U.K. for European Space Agency (ESA), ESTEC, Noordwijk, The Netherlands

¹Ocean Processes Analysis Laboratory, University of New Hampshire, Durham, New Hampshire 3824, United States

^{*}Norwegian Institute for Water Research, Thormøhlensgate 53 D, N-5006 Bergen, Norway

monitoring. At present, OA monitoring efforts are dominated by in situ observations from moorings, ships and associated platforms. While key to any monitoring campaign, in situ data tend to be spatially sparse, especially in inhospitable regions, and so on their own are unlikely to provide a comprehensive, robust and cost-effective solution to global OA monitoring. The need to monitor and study large areas of the Earth has driven the development of satellite-based sensors.

Increasingly, as in situ data accumulate, attempts are being made to use in situ hydrographic data 10-13 and/or remotely sensed data 14,15 to provide proxies and indicators for the condition of the carbonate system, enabling data gaps to be filled in both space and time. The increased availability of in situ data creates a substantial data set to develop and test the capabilities of satellite-derived products, and we suggest that the recent availability of satellite-based salinity measurements provides new key insights for studying and assessing OA from space.

2. THE COMPLEXITIES OF THE CARBONATE SYSTEM

The oceanic carbonate system can be understood and probed through four key parameters: total alkalinity (TA), dissolved inorganic carbon (DIC), pH, and fugacity of CO_2 (f_{CO2}). The latter may be replaced with the related partial pressure of CO₂, p_{CO2} , from which f_{CO2} can be calculated, and the two are often used interchangeably. In principle, knowledge of any two of these four is sufficient to solve the carbonate system equations. However, overdetermination, the process of measuring at least three parameters, is advantageous.

The relationships between the different carbonate system parameters are fundamentally driven by thermodynamics, hence influenced by temperature and pressure, and knowing these is fundamental for calculating the carbonate system as a whole.¹⁶ Water temperature is the major controller of the solubility of CO₂, ¹⁷ so seasonal changes in sea temperature can, depending on the region, be significant for driving changes in f_{CO2} (and consequently DIC and pH). Salinity affects the coefficients of the carbonate system equations. Hence to solve the equations, it is necessary to estimate temperature, salinity and pressure along with carbonate parameters.

The ratio between ions (the constituents of salinity) will tend to remain constant anywhere in the global oceans, resulting in a strong relationship between TA and salinity. 18 Unfortunately, a universal relationship between TA and salinity does not apply in certain regions, for instance in areas influenced by freshwater outflows from rivers, or areas where calcification and/or CaCO₃ dissolution occurs, such as where calcifying plankton are prevalent.¹⁹ In these regions, it is therefore critical to gain additional local knowledge. For example, different rivers will have different ionic concentrations (and therefore different TA concentrations) depending on the surrounding geology and hydrology

For DIC, f_{CO2} (or p_{CO2}), and pH, the other important process is biological activity. ¹⁹ Removal or addition of CO_2 by plankton photosynthesis or respiration can be a significant component of the seasonal signal.²⁰ Biological activity, in turn, is driven by factors such as nutrient dynamics and light conditions, which again are regionally specific. Measurements of chlorophyll (a proxy for biomass) and/or oxygen concentration can be useful for interpreting the biological component of the carbon signal.

The combination of these processes means that it is extremely challenging to produce a global relationship between any component of the carbonate system and its drivers. To enable us to understand these dynamics, extrapolation from collected data points to the global ocean is needed, and along with model predictions, empirical relationships and data sets are important and need to be studied and developed. OA needs to be assessed using these relationships on a global scale, but regional complexities, particularly where riverine and coastal processes dominate, ^{21,22} cause significant challenges for global empirical relationships.

3. CURRENT IN SITU APPROACHES AND CHALLENGES

Laboratory measurements are the gold standard for assessing the carbonate system in seawater, with accuracy far in excess of that achievable from satellites. ^{23–25} However, research vessel time is expensive and limited in coverage, so autonomous in situ instruments are also deployed, for example, on buoys, with less accuracy. 26 A notable example is the Argo network of over 3000 drifters, which measure temperature and salinity throughout the deep global ocean. Interpolation of Argo data is much less challenging than for most in situ measurements. Argo is the closest in situ data have come to the global, synoptic measurements possible with satellites, but shallow or enclosed seas are not represented (there are as yet no Argo instruments in the open Arctic Ocean). Table 1 lists more examples. Of the four

Table 1. In Situ Datasets and Programs than Can Be Used for the Development and Validation of OA Remote Sensing Algorithms

data set name and reference	temporal period	geographic location	variables	no. of data points
SOCAT v2.0 ²⁷	1968-2011	global*	$f_{\rm CO2}$, SSS, SST	6 000 000+
LDEO v2012 ²⁸	1980-present	global*	$p_{\rm CO2}$, SSS, SST	6 000 000+
GLODAP ²⁹	1970-2000	global	TA, DIC, SSS, SST, Nitrate	10 000+
CARINA AMS v1.2 ³⁰	1980-2006	Arctic	TA, DIC, SSS, SST	1500+
CARINA ATL v1.0 ³¹		Atlantic		
CARINA SO v1.1 ³²		Southern Ocean		
AMT ³³	1995-present	Atlantic	$p_{\rm CO2W}$, SSS, SST, Chl, pH	1000+
NIVA Ferrybox ³⁴	2008-present	Arctic	p _{CO2W} , TA, DIC, SSS, SST	1000+
OWS Mike ³⁵	1948-2009	Arctic	TA, DIC, SSS, SST, Chl	1000+
RAMA Moored buoy array ³⁶	2007-present	Bay of Bengal	SSS, SST	1000+
ARGO buoys ³⁷	2003-present	global	SSS, SST	1 000 000+
OOI ³⁸	2014 onward	global (six sites)	p_{CO2} , SSS, SST, nitrate	new program
SOCCOM ³⁹	2014 onward	Southern Ocean	SSS, SST, pH, nitrate	new program

key parameters, only f_{CO2} (or p_{CO2}) and pH are routinely monitored in situ. As yet there are limited capabilities to measure DIC and TA autonomously, hence these parameters must be measured either in a ship-based laboratory or on land.

4. POTENTIAL OF SPACE BASED OBSERVATIONS

4.1. Advantages and Disadvantages. While it has proven difficult to use remote sensing to directly monitor and detect changes in seawater pH and their impact on marine organisms, satellites can measure sea surface temperature and salinity (SST and SSS) and surface chlorophyll-a, from which carbonate system parameters can be estimated using empirical relationships derived from in situ data. Although surface measurements may not be representative of important biological processes, for example, fish or shellfish, observations at the surface are particularly important for OA because the change in carbonate chemistry due to atmospheric CO₂ occurs in the surface first. Thus, satellites have great potential as a tool for assessing changes in carbonate chemistry.

SST has been measured from space with infrared radiometry since the 1960s, but the data are only globally of sufficient quality for climate studies since 1991. Satellite measurements of chlorophyll-*a* in the visible are more recent, starting in 1986 and delivering high quality global data since 1997. Both measurements are made globally at high spatial and temporal resolution, but with data gaps due to effects such as cloud, which can greatly affect data availability in cloudy regions. SST is measured in the top few microns, and chlorophyll-a is generally measured to depths around 1–100 m, depending on water clarity. Data quality can be affected by many issues, for example, adjacent land or ice may affect both SST and chlorophyll-a retrievals, and suspended sediment may affect chlorophyll-a retrievals.

Only since 2009 has a satellite-based capability for measuring SSS existed. Increasing salinity decreases the emissivity of seawater and so changes the microwave radiation emitted at the water surface. ESA Soil Moisture and Ocean Salinity (SMOS) and NASA-CONAE Aquarius (launched in 2009 and 2011 respectively, both currently in operation), are L-band microwave sensors designed to detect variations in microwave radiation and thus estimate ocean salinity in the top centimeter. The instruments are novel and the measurement is very challenging, and research is ongoing to improve data quality. 42 The instruments can measure every few days at a spatial resolution of 35-100 km, but single measurements are very noisy, so the instantaneous swath data are generally spatially and temporally averaged over 10 days or a month, with an intended accuracy around 0.1-0.2 g/kg for monthly 200 km data. A particular issue close to urban areas is radio frequency interference from illegal broadcasts, which are gradually being eliminated but still result in large data gaps, particularly for SMOS. The signal can be affected by nearby land or sea ice, and the sensitivity to SSS decreases for cold water, by about 50% from 20 to 0 °C.⁴³

With these challenges, a central question is whether satellite SSS can bring new complementary information to in situ SSS measurements such as Argo for assessing OA. Direct comparisons 44,45 indicate differences of 0.15–0.5 g/kg in a 1° region over 10–30 days. The two are difficult to compare directly, however, as Argo measures 5 m or more from the surface, so some differences are expected even in the absence of errors, especially where the water column is stratified. A better strategy might be to compare their effectiveness in estimating OA. How the uncertainties propagate through the carbonate system calculations is the subject of ongoing research.

Despite biases and uncertainties, satellite measurements of SSS in the top centimeter contain geophysical information not detected by Argo. 46,47 In addition, Argo coverage can be much poorer than satellite SSS in several regions such as the major western boundary or equatorial currents and across strong oceanic fronts. The use of interpolated Argo products presents an additional source of uncertainty due to the interpolation scheme. Satellite SSS can also resolve mesoscale spatial structures not resolved by Argo measurements, and unlike Argo, satellites provide a synoptic "snapshot" of a region at a given time.

Regular mapping of the SSS field with unprecedented temporal and spatial resolution at global scale is now possible from satellites. The impact of using satellite SSS for carbonate system algorithms can now be tested, where previously there was a reliance on climatology, in situ or model data. For example, this provides the means to study the impact that freshwater influences (sea ice melt, riverine inputs and rain) can have on the marine carbonate system. The use of satellite SSS data will also allow evaluation of the impact on the carbonate system of the inter- and intra-annual variations in SSS.

Recent advances in radar altimetry (e.g., Cryosat-2 and Sentinel 1 satellites and sensors) are already enabling significant improvements in satellite sea-ice thickness measurements.⁵⁰ Thin sea ice thickness can now also be determined from SMOS, complementing altimeter estimates mostly valid for thick sea ice. ⁵¹ Sea ice thickness is important for OA research as it indicates whether ice is seasonal or multiyear, supporting the interpretation of carbonate parameters. Altimetry is also used to measure wind speeds and increases the coverage of scatterometer estimates in polar regions. It provides higher-resolution (along track) estimates of surface wind stress, which can potentially be used to indicate regions of upwelling. Wind-driven upwelling causes dense cooler water (with higher concentrations of CO₂ and thus more acidic) to be drawn up from depth to the ocean surface. This upwelling can have significant impacts on local OA and ecosystems, 4,52 especially at eastern oceanic boundaries. 53,54

It is important to emphasize that the use of Earth observation data to derive carbonate parameters should not be seen as a replacement for in situ measurement campaigns, especially due to the current reliance on empirical and regional algorithms. Earth observation algorithms need calibration and validation with in situ data such as those taken by GOA-ON, and if the carbonate system response changes over time, empirical and regional algorithms tuned to previous conditions may become less reliable.

4.2. Algorithms for Estimating Carbonate Parameters.

The four key OA parameters (pCO₂, DIC, TA, pH) are largely driven by temperature, salinity and biological activity, allowing empirical relationships to be developed using in situ measurements of OA parameters. Table 2 shows a range of published algorithms based on such relationships, while Figure 1 shows

Table 2. Example Regional Algorithms for Each Carbonate Parameter Illustrating the Variable Dependencies. Chl is Chlorophyll-*a* and MLD is Mixed Layer Depth

parameter	dependencies	region and references
pCO_2	SST	global, ⁵⁶ Barents Sea ⁵⁷
	SST, SSS	Barents Sea, ⁵⁸ Caribbean ¹⁴
	SST, Chl	North Pacific ⁵⁹
	SSS, Chl	North Sea ⁶⁰
	SST, SSS, Chl	North Pacific ⁶¹
	SST, Chl, MLD	Barents Sea ⁶²
TA	SSS	Barents Sea ⁵⁷
	SST, SSS	global, 18,63 Arctic 15
	SSS, nitrate	Global ⁵⁵
DIC	SST, SSS	Equatorial pacific ⁶⁴
	SST, SSS, Chl	Arctic ¹⁵
pН	SST, Chl	North Pacific 10

Environmental Science & Technology

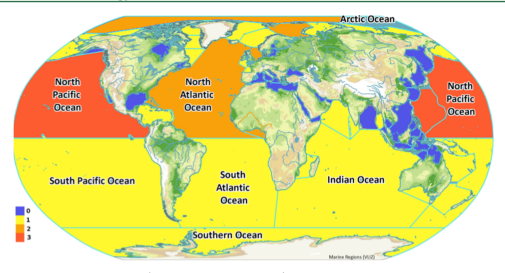


Figure 1. Number of key carbonate parameters (f_{CO2} or p_{CO2} , TA, DIC, pH) for which regional algorithms exist in the literature that can be implemented using just satellite Earth observation data. Regions are indicative of open ocean areas, as implementation of algorithms in coastal areas may be problematic.

their geographical coverage. Both illustrate that most of the literature has focused on the northern basins of the Pacific and Atlantic and the Arctic, especially the Barents Sea, with all other regions only attracting algorithms for a single parameter or none at all. 55

NOAA's experimental Ocean Acidification Product Suite (OAPS) is a regional example of using empirical algorithms with a combination of climatological SSS and satellite SST to provide synoptic estimates of sea surface carbonate chemistry in the Greater Caribbean Region. 14 p_{CO2} and TA were derived from climatological SSS and satellite SST, then used to calculate monthly estimates of the remaining carbonate parameters, including aragonite saturation state and carbonate ion concentration. In general the derived data were in good agreement with in situ measured data (e.g., mean derived TA = $2375 \pm 36 \mu mol$ kg⁻¹ compared to a mean ship-measured TA = 2366 \pm 77 μ mol kg⁻¹). OAPS works well in areas where chlorophyll-a is low, however in regions of high chlorophyll-a, where net productivity is likely to perturb the carbonate system, and in areas where there are river inputs, the approach tends to underestimate aragonite saturation state, for example.²¹

A quite different approach is the assimilation of satellite data into ocean circulation models.⁶⁵ The model output carbonate parameters can then be used directly. This allows satellite-observed effects to be extended below the water surface, albeit with the uncertainties inherent in model data. Here we seek to assess the direct use of satellite data through empirical algorithms to improve OA estimates.

4.3. Regions of Interest for Earth Observation. *Arctic Seas.* It is increasingly recognized that the Polar Oceans (Arctic and Antarctic) are particularly sensitive to OA. Lower alkalinity (and thus buffer capacity), enhanced warming, reduced sea-ice cover resulting in changes in the freshwater budget, ⁶⁷ and nutrient limitation make it more vulnerable to future OA. Retreating ice also provides increased open water for air-sea gas exchange and primary production. ⁶⁹

The remote nature of the Arctic Ocean provides difficulties for collecting in situ data sets, with limited ship, autonomous vehicle and buoy access, and in situ data collection during winter months is often impossible. Therefore, the use of remote sensing techniques is very attractive, if sufficient in situ data can be found

to calibrate satellite algorithms, and if the challenges of Arctic remote sensing can be overcome. These waters are very challenging regions for satellite remote sensing. For instance, low water temperatures reduce the sensitivity range of SSS sensors, ⁴³ and sea ice can complicate retrievals of SSS and chlorophyll-a. ^{70,71} Improvement in the accuracy of high latitude satellite SSS is expected soon by combining observations from SMOS, Aquarius and the upcoming SMAP sensor, all polar-orbiting L-band radiometers.

The Bay of Bengal. This region is clearly a focus of current OA research with unique characteristics due to the large freshwater influence. The flow of fresh water from the Ganges Delta into Bay of Bengal (42 000 m³/sec) represents the second greatest discharge source in the world. Additionally, rainfall along with freshwater inputs exceeds evaporation, resulting in net water gain annually in the Bay of Bengal. Collectively these provide an annual positive water balance that reduces surface salinity by 3-7 g/kg compared to the adjacent Arabian Sea, 72,73 resulting in distinctly different biogeochemical regimes.⁷⁴ Biogeochemically, the Indian Ocean is one of the least studied and most poorly understood ocean basins in the world.⁷⁴ This is particularly true for the Bay of Bengal where a relatively small number of hydrographic sections and underway surface observations have been undertaken, despite the notable influence of freshwater on particle dynamics, air-sea carbon flux and surface carbonate chemistry. 75-79 North of 15° S, TA increases relative to salinity, 80 indicating the presence of an important land source that can broadly affect acidification dynamics.

To date there is little work on acidification dynamics and air sea exchange of CO_2 in the Bay of Bengal. $^{81-83}$ In 2013, the Bay of Bengal Ocean Acidification (BOBOA) Mooring was deployed for the first time in Bay of Bengal (15°N, 90°E) by PMEL (NOAA) and the Bay of Bengal Large Marine Ecosystem Program (BOBLME). Data from the buoy will improve our understanding of biogeochemical variations in the open ocean environment of the Bay of Bengal.

It is an open question whether SSS can be used to estimate TA in the Bay of Bengal. An important step toward answering this question would be to investigate the spatial variability of the TA to salinity relationship in the region. Use of satellite SSS in the region is also challenged by heavy radio frequency interference.

The Greater Caribbean and the Amazon plume. The reefs in the Greater Caribbean Region are economically important to the US and Caribbean nations with an estimated annual net value of US\$3.1-4.6 billion in 2000.84 At least two-thirds of these reefs are threatened from human impacts including OA. The skeleton of a coral is made of aragonite and the growth of their skeletons is reduced by OA,⁶ and numerous studies have shown a net decline in coral calcification (growth) rates in accordance with declining CaCO₃ saturation state. 85 The waters of the Greater Caribbean region are predominantly oligotrophic and similar to the subtropical gyre from which it receives most of its water.¹⁴ While the often shallow water environments of coral reefs and the plethora of small islands can make it challenging for Earth observation instruments to collect reliable data, the oligotrophic nature and the similarities in water type across the whole region make it ideal for the development of novel products. This region therefore provides an ideal case study to develop and evaluate algorithms representative of a shallow, oligotrophic environment.

The Amazon plume, south of the Greater Caribbean, is the largest freshwater discharge source in the world (209 000 m 3 /sec). It can cause SSS decreases of several units many hundreds of kilometers from land, and has an area that seasonally can reach $10^6~{\rm km}^2$. These characteristics make it an ideal case study for testing and evaluating remote sensing algorithms, particularly to study the space-time resolution trade-offs using SSS sensors.

5. FUTURE OPPORTUNITIES AND FOCUS

The Copernicus program is a European flagship initiative, worth more than €7 billion, which aims to provide an operational satellite monitoring capability and related services for the environment and security.86 The launch of the Sentinel-1A satellite in 2014 signaled its start. Of the five Sentinel satellite types, Sentinels 2 and 3 are most appropriate for assessment of the marine carbonate system. 87–89 These satellites will provide chlorophyll-a and SST with unprecedented spatial and temporal coverage. The development of higher spatial resolution geostationary sensors that continually monitor chlorophyll-a and SST over the same area of the Earth also holds much potential for the future of OA assessment and research. 90 These satellites and sensors are able to provide 10 or more observations per day, allowing the study of the effect of tidal and diurnal cycles on OA. The societal importance of measuring and observing the global carbon cycle was further highlighted with the launch of the NASA Orbiting Carbon Observatory (OCO-2) in 2014. This satellite and its sensors are designed to observe atmospheric CO2 concentrations, but its potential for marine carbon cycle and OA is likely to be a focus of future research.

SMOS and Aquarius have recently passed their nominal lifetimes, with SMOS now extended until 2017. Based on the lifetimes of previous satellite Earth observation sensors, they may well operate until the early 2020s. NASA's SMAP satellite, to be launched in January 2015, should provide short-term continuity. The development of the technology and the clear importance of monitoring ocean salinity are likely to support the development of future satellite sensors. Also, historical time series data from alternative microwave sensors hold the potential for a 10+ year time series of satellite based SSS observations, ⁹¹ and this sort of measurement record is likely to extend into the future as it forms the basis of a global SSS monitoring effort.

In summary, satellite products developed up to now in the OA context have been regional, empirical or derived with a limited variety of satellite data sets, rendering an effort to systematically

exploit remote sensing assets (capitalizing on the recent advent of satellite salinity measurements) absolutely timely. To-date there is only regional application of satellite SST to address the issue of assessing OA, ⁶² along with two nonpeer-reviewed attempts to calculate carbonate system products using satellite SSS data. ^{92,93} Supported by good in situ measurement campaigns, especially in places with currently poor in situ coverage such as the Arctic, satellite measurements are likely to become a key element in understanding and assessing OA.

AUTHOR INFORMATION

Corresponding Author

*E-mail: peland@pml.ac.uk.

Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Funding

This work was funded by the European Space Agency Support to Science Element Pathfinders Ocean Acidification project (contract No. 4000110778/14/I-BG).

Notes

The authors declare no competing financial interest.

Biography

Peter Land is a remote sensing scientist at Plymouth Marine Laboratory (PML), specializing in atmosphere-ocean gas exchange and carbonate chemistry. Jamie Shutler is an oceanographer and former European Space Agency (ESA) fellow specializing in atmosphere-ocean gas exchange at the University of Exeter. Helen Findlay is an oceanographer at PML specializing in ocean acidification and carbonate chemistry. Fanny Girard-Ardhuin is a remote sensing scientist specializing in sea ice at l'Institut Français de Recherche pour l'Exploitation de la Mer (Ifremer). Nicolas Reul is a remote sensing scientist at Ifremer and member of the SMOS scientific team. Jean-Francois Piolle is a computer scientist at Ifremer. Bertrand Chapron leads remote sensing research at Ifremer. Yves Quilfen is an altimetry remote sensing scientist at Ifremer. Joseph Salisbury and Douglas Vandemark are oceanographers at the University of New Hampshire focusing on biogeochemistry and ecology in coastal areas. Richard Bellerby is a chemical oceanographer at the Norwegian Institute for Water Research, a member of the GOA-ON executive committee, and leader of the AMAP and SCAR ocean acidification working groups. Punyasloke Bhadury is a coastal ecologist at the Indian Institute of Science Education and Research-Kolkata. Roberto Sabia is a specialist in remote sensing of salinity working for

ACKNOWLEDGMENTS

This work was enabled by European Space Agency (ESA) Support to Science Element (STSE) Pathfinders Ocean Acidification project (contract No. 4000110778/14/I-BG). The authors gratefully acknowledge the assistance of Diego Fernandez (STSE programme manager).

■ REFERENCES

(1) Stocker, T. F.; Qin, D.; Plattner, G. K.; Tignor, M.; Allen, S. K.; Boschung, J.; Nauels, A.; Xia, Y.; Bex, V.; Midgley, P. M. Climate Change 2013: The Physical Science Basis. Intergovernmental Panel on Climate Change, Working Group I Contribution to the IPCC Fifth Assessment Report (ARS); Cambridge Univ. Press: New York, 2013.

(2) Fabry, V. J.; Seibel, B. A.; Feely, R. A.; Orr, J. C. Impacts of ocean acidification on marine fauna and ecosystem processes. *ICES J. Mar. Sci.* **2008**, *65* (3), 414–432.

- (3) Turley, C.; Eby, M.; Ridgwell, A. J.; Schmidt, D. N.; Findlay, H. S.; Brownlee, C.; Riebesell, U.; Fabry, V. J.; Feely, R. A.; Gattuso, J. P. The societal challenge of ocean acidification. *Mar. Pollut. Bull.* **2010**, *60* (6), 787–792.
- (4) Feely, R. A.; Sabine, C. L.; Hernandez-Ayon, J. M.; Ianson, D.; Hales, B. Evidence for upwelling of corrosive "acidified" water onto the continental shelf. *Science* **2008**, 320 (5882), 1490–1492.
- (5) Bellerby, R. G. J. UN biodiversity and OA report. http://www.cbd.int/ts.
- (6) Kroeker, K. J.; Kordas, R. L.; Crim, R.; Hendriks, I. E.; Ramajo, L.; Singh, G. S.; Duarte, C. M.; Gattuso, J. P. Impacts of ocean acidification on marine organisms: Quantifying sensitivities and interaction with warming. *Global Change Biol.* **2013**, DOI: 10.1111/gcb.12179.
- (7) Salisbury, J.; Green, M.; Hunt, C.; Campbell, J. Coastal acidification by rivers: A threat to shellfish? *Eos, Trans. Am. Geophys. Union* **2008**, 89 (50), 513.
- (8) Widdicombe, S.; Spicer, J. I. Predicting the impact of ocean acidification on benthic biodiversity: What can animal physiology tell us? *J. Exper. Mar. Biol. Ecol.* **2008**, *366* (1), 187–197.
- (9) Ridgwell, A.; Schmidt, D. N.; Turley, C.; Brownlee, C.; Maldonado, M. T.; Tortell, P.; Young, J. R. From laboratory manipulations to Earth system models: Scaling calcification impacts of ocean acidification. *Biogeosciences* **2009**, *6* (11), 2611–2623.
- (10) Nakano, Y.; Watanabe, Y. W. Reconstruction of pH in the surface seawater over the north Pacific basin for all seasons using temperature and chlorophyll-a. *J. Oceanogr.* **2005**, *61* (4), 673–680.
- (11) Juranek, L. W.; Feely, R. A.; Peterson, W. T.; Alin, S. R.; Hales, B.; Lee, K.; Sabine, C. L.; Peterson, J. A novel method for determination of aragonite saturation state on the continental shelf of central Oregon using multi-parameter relationships with hydrographic data. *Geophys. Res. Lett.* 2009, 36 (24), L24601.
- (12) Midorikawa, T.; Inoue, H. Y.; Ishii, M.; Sasano, D.; Kosugi, N.; Hashida, G.; Nakaoka, S.-i.; Suzuki, T. Decreasing pH trend estimated from 35-year time series of carbonate parameters in the Pacific sector of the Southern Ocean in summer. *Deep Sea Res.*, Part I 2012, 61, 131–139.
- (13) Bostock, H. C.; Mikaloff Fletcher, S. E.; Williams, M. J. M. Estimating carbonate parameters from hydrographic data for the intermediate and deep waters of the Southern Hemisphere Oceans. *Biogeosci. Discuss.* **2013**, *10* (4), 6225–6257.
- (14) Gledhill, D. K.; Wanninkhof, R.; Millero, F. J.; Eakin, M. Ocean acidification of the greater Caribbean region 1996–2006. *J. Geophys. Res.* **2008**, *113* (C10), C10031.
- (15) Arrigo, K. R.; Pabi, S.; van Dijken, G. L.; Maslowski, W. Air-sea flux of CO_2 in the Arctic Ocean, 1998–2003. *J. Geophys. Res.* **2010**, 115 (G4), G04024.
- (16) Dickson, A. G.; Goyet, C. Handbook of Methods for the Analysis of the Various Parameters of the Carbon Dioxide System in Sea Water, 1992; Vol. 2.
- (17) Weiss, R. F. Carbon dioxide in water and seawater: The solubility of a non-ideal gas. *Mar. Chem.* **1974**, 2 (3), 203–215.
- (18) Lee, K.; Tong, L. T.; Millero, F. J.; Sabine, C. L.; Dickson, A. G.; Goyet, C.; Park, G. H.; Wanninkhof, R.; Feely, R. A.; Key, R. M., Global relationships of total alkalinity with salinity and temperature in surface waters of the world's oceans. *Geophys. Res. Lett.* **2006**, *33*, (19).
- (19) Smith, S. V.; Key, G. S. Carbon dioxide and metabolism in marine environments. *Limnol. Oceanogr* **1975**, 20 (3), 493–495.
- (20) Sarmiento, J. L.; Gruber, N. Ocean Biogeochemical Dynamics; Cambridge University Press, 2006; Vol. 503.
- (21) Gledhill, D. K.; Wanninkhof, R.; Eakin, C. M., Observing ocean acidification from space. *Oceanography* **2009**, 22.
- (22) Sun, Q.; Tang, D.; Wang, S. Remote-sensing observations relevant to ocean acidification. *Int. J. Rem. Sensing* **2012**, 33 (23), 7542–7558.
- (23) Dickson, A. G., The carbon dioxide system in seawater: Equilibrium chemistry and measurements. In *Guide to Best Practices for Ocean Acidification Research and Data Reporting*, Riebesell, U.; Fabry, C. J.; Hansson, L.; Gattuso, J.-P., Eds.; European Commission: Brussels, 2011; pp 17–40.

- (24) DicksonA. G.SabineC. L.ChristianJ. R.Guide to Best Practices for Ocean CO₂ Measurements, PICES Special Publication 3, 2007
- (25) Byrne, R. H. Measuring Ocean Acidification: New Technology for a New Era of Ocean Chemistry. *Environ. Sci. Technol.* **2014**, 48 (10), 5352–5360.
- (26) Martz, T. R.; Connery, J. G.; Johnson, K. S. Testing the Honeywell Durafet® for seawater pH applications. *Limnol. Oceanogr. Methods* **2010**, *8*, 172–184.
- (27) Bakker, D. C. E.; Hankin, S.; Olsen, A.; Pfeil, B.; Smith, K.; Alin, S. R.; Cosca, C.; Hales, B.; Harasawa, S.; Kozyr, A. An update to the surface ocean CO₂ Atlas (SOCAT version 2). *Earth Syst. Sci. Data* **2014**, DOI: 10.5194/essd-6-69-2014.
- (28) Takahashi, T.; Sutherland, S. C.; Kozyr, A. Global Ocean Surface Water Partial Pressure of CO₂ Database: Measurements Performed During 1957–2012 (Version 2012), ORNL/CDIAC-160, NDP-088(V2012); Carbon Dioxide Information Analysis Center, Oak Ridge National Laboratory, U.S. Department of Energy: Oak Ridge, TN, 2013.
- (29) Key, R. M.; Kozyr, A.; Sabine, C. L.; Lee, K.; Wanninkhof, R.; Bullister, J. L.; Feely, R. A.; Millero, F. J.; Mordy, C.; Peng, T. H., A global ocean carbon climatology: Results from Global Data Analysis Project (GLODAP). *Global Biogeochem. Cycles* **2004**, *18*, (4).
- (30) CARINA group. Carbon in the Arctic Mediterranean Seas Region— The CARINA Project: Results and Data, Version 1.2; Carbon Dioxide Information Analysis Center, Oak Ridge National Laboratory, U.S. Department of Energy: Oak Ridge, TN, 2009.
- (31) CARINA group. Carbon in the Atlantic Ocean Region—The CARINA Project: Results and Data, Version 1.0.; Carbon Dioxide Information Analysis Center, Oak Ridge National Laboratory, U.S. Department of Energy: Oak Ridge, TN, 2009.
- (32) CARINA group. Carbon in the Southern Ocean Region—The CARINA Project: Results and Data, Version 1.1.; Carbon Dioxide Information Analysis Center, Oak Ridge National Laboratory, U.S. Department of Energy: Oak Ridge, TN, 2010.
- (33) Robinson, C.; Holligan, P.; Jickells, T.; Lavender, S. The Atlantic Meridional Transect Programme (1995–2012). *Deep Sea Res., Part II* **2009**, *56* (15), 895–898.
- (34) Yakushev, E. V.; Sørensen, K. On seasonal changes of the carbonate system in the Barents Sea: Observations and modeling. *Mar. Biol. Res.* **2013**, *9* (9), 822–830.
- (35) Skjelvan, I.; Falck, E.; Rey, F.; Kringstad, S. B. Inorganic carbon time series at Ocean Weather Station M in the Norwegian Sea. *Biogeosciences* **2008**, *5*, 549–560.
- (36) McPhadenM. J.MeyersG.AndoK.MasumotoY.MurtyV. S. N.RavichandranM.SyamsudinF.VialardJ.YuL.YuW.RAMA: The Research Moored Array for African—asian—australian Monsoon Analysis and Prediction, 2009
- (37) ARGO Argo part of the integrated global observation strategy. http://www.argo.ucsd.edu (14/12/2014),.
- (38) OOI Ocean Observatories Initiative. http://oceanobservatories.org (14/12/2014),.
- (39) SOCCOM, Southern Ocean Carbon and Climate Observations and Modeling. http://soccom.princeton.edu (accessed December 14, 2014).
- (40) Merchant, C. J.; Embury, O.; Rayner, N. A.; Berry, D. I.; Corlett, G. K.; Lean, K.; Veal, K. L.; Kent, E. C.; Llewellyn-Jones, D. T.; Remedios, J. J., A 20 year independent record of sea surface temperature for climate from Along—Track Scanning Radiometers. *J. Geophys. Res.: Oceans* **2012**, *117*, (C12).
- (41) McClain, C. R.; Feldman, G. C.; Hooker, S. B. An overview of the SeaWiFS project and strategies for producing a climate research quality global ocean bio-optical time series. *Deep Sea Res., Part II* **2004**, *S1* (1), 5–42.
- (42) Font, J.; Boutin, J.; Reul, N.; Spurgeon, P.; Ballabrera-Poy, J.; Chuprin, A.; Gabarró, C.; Gourrion, J.; Guimbard, S.; Hénocq, C. SMOS first data analysis for sea surface salinity determination. *Int. J. Rem. Sens.* **2013**, *34* (9–10), 3654–3670.
- (43) Font, J.; Camps, A.; Borges, A.; Martín-Neira, M.; Boutin, J.; Reul, N.; Kerr, Y. H.; Hahne, A.; Mecklenburg, S. SMOS: The challenging sea

- surface salinity measurement from space. *Proc. IEEE* **2010**, 98 (5), 649–665.
- (44) Boutin, J.; Martin, N.; Reverdin, G.; Morisset, S.; Yin, X.; Centurioni, L.; Reul, N. Sea surface salinity under rain cells: SMOS satellite and in situ drifters observations. *J. Geophys. Res.: Oceans* **2014**, 119 (8), 5533–5545.
- (45) Reul, N.; Chapron, B.; Lee, T.; Donlon, C.; Boutin, J.; Alory, G. Sea surface salinity structure of the meandering Gulf Stream revealed by SMOS sensor. *Geophys. Res. Lett.* **2014**, *41* (9), 3141–3148.
- (46) Boutin, J.; Martin, N.; Reverdin, G.; Yin, X.; Gaillard, F. Sea surface freshening inferred from SMOS and ARGO salinity: Impact of rain. *Ocean Sci.* **2013**, *9*, 183–192.
- (47) Sabia, R.; Klockmann, M. Fernández-Prieto, D.; Donlon, C., A first estimation of SMOS-based ocean surface T-S diagrams. *J. Geophys. Res.: Oceans* **2014**, *119* (10), 7357–7371.
- (48) Hosoda, S.; Ohira, T.; Nakamura, T. A monthly mean dataset of global oceanic temperature and salinity derived from Argo float observations. *JAMSTEC Rep. Res. Dev* **2008**, *8*, 47–59.
- (49) Reul, N.; Fournier, S.; Boutin, J.; Hernandez, O.; Maes, C.; Chapron, B.; Alory, G.; Quilfen, Y.; Tenerelli, J.; Morisset, S. Sea surface salinity observations from space with the SMOS satellite: A new means to monitor the marine branch of the water cycle. *Surv. Geophysics* **2014**, 35 (3), 681–722.
- (50) Laxon, S. W.; Giles, K. A.; Ridout, A. L.; Wingham, D. J.; Willatt, R.; Cullen, R.; Kwok, R.; Schweiger, A.; Zhang, J.; Haas, C. CryoSat-2 estimates of Arctic sea ice thickness and volume. *Geophys. Res. Lett.* **2013**, *40* (4), 732–737.
- (51) Kaleschke, L.; Tian-Kunze, X.; Maaß, N.; Mäkynen, M.; Drusch, M., Sea ice thickness retrieval from SMOS brightness temperatures during the Arctic freeze-up period. *Geophys. Res. Lett.* **2012**, *39*, (5).
- (52) Mathis, J. T.; Pickart, R. S.; Byrne, R. H.; McNeil, C. L.; Moore, G. W. K.; Juranek, L. W.; Liu, X.; Ma, J.; Easley, R. A.; Elliot, M. M., Storminduced upwelling of high pCO₂ waters onto the continental shelf of the western Arctic Ocean and implications for carbonate mineral saturation states. *Geophys. Res. Lett.* **2012**, *39*, (7).
- (53) Mahadevan, A.; Tagliabue, A.; Bopp, L.; Lenton, A.; Memery, L.; Lévy, M. Impact of episodic vertical fluxes on sea surface pCO₂. *Philos. Trans. R. Soc., A* **2011**, 369 (1943), 2009–2025.
- (54) Mahadevan, A. Ocean science: Eddy effects on biogeochemistry. *Nature* **2014**, *506*, 168–169.
- (55) Takahashi, T.; Sutherland, S. Climatological Mean Distribution of pH and Carbonate Ion Concentration in Global Ocean Surface Waters in the Unified pH Scale and Mean Rate of Their Changes in Selected Areas, OCE 10-38891; National Science Foundation: Washington, D. C., USA, 2013.
- (56) Goddijn-Murphy, L. M.; Woolf, D. K.; Land, P. E.; Shutler, J. D.; Donlon, C. Deriving a sea surface climatology of CO₂ fugacity in support of air-sea gas flux studies. *Ocean Sci. Discuss.* **2014**, *11*, 1895–1948.
- (57) Årthun, M.; Bellerby, R. G. J.; Omar, A. M.; Schrum, C. Spatiotemporal variability of air—sea CO < sub> 2</sub> fluxes in the Barents Sea, as determined from empirical relationships and modeled hydrography. *J. Mar. Syst.* **2012**, *98*, 40–50.
- (58) Friedrich, T.; Oschlies, A., Basin-scale pCO₂ maps estimated from ARGO float data: A model study. *J. Geophys. Res.: Oceans* **2009**, *114*, (C10).
- (59) Ono, T.; Saino, T.; Kurita, N.; Sasaki, K. Basin-scale extrapolation of shipboard pCO₂ data by using satellite SST and Chla. *Int. J. Rem. Sens.* **2004**, 25 (19), 3803–3815.
- (60) Borges, A. V.; Ruddick, K.; Lacroix, G.; Nechad, B.; Asteroca, R.; Rousseau, V.; Harlay, J., Estimating pCO_2 from remote sensing in the Belgian coastal zone. *ESA Spec. Publ.* **2010**, *686*.
- (61) Sarma, V. V. S. S.; Saino, T.; Sasaoka, K.; Nojiri, Y.; Ono, T.; Ishii, M.; Inoue, H. Y.; Matsumoto, K., Basin-scale pCO₂ distribution using satellite sea surface temperature, Chl a, and climatological salinity in the North Pacific in spring and summer. *Global Biogeochem. Cycles* **2006**, 20, (3).
- (62) Lauvset, S. K.; Chierici, M.; Counillon, F.; Omar, A.; Nondal, G.; Johannessen, T.; Olsen, A. Annual and seasonal fCO₂ and air—sea CO₂

- fluxes in the Barents Sea. J. Mar. Syst. 2013, DOI: 10.1016/j.jmarsys.2012.12.011.
- (63) Millero, F. J.; Lee, K.; Roche, M. Distribution of alkalinity in the surface waters of the major oceans. *Mar. Chem.* **1998**, *60* (1), 111–130.
- (64) Loukos, H.; Vivier, F.; Murphy, P. P.; Harrison, D. E.; Le Quéré, C. Interannual variability of equatorial Pacific CO_2 fluxes estimated from temperature and salinity data. *Geophys. Res. Lett.* **2000**, 27 (12), 1735–1738.
- (65) Anderson, D.; Sheinbaum, J.; Haines, K. Data assimilation in ocean models. *Rep. Prog. Phys.* **1996**, *59* (10), 1209.
- (66) Steinacher, M.; Joos, F.; Frölicher, T. L.; Plattner, G. K.; Doney, S. C. Imminent ocean acidification in the Arctic projected with the NCAR global coupled carbon cycle-climate model. *Biogeosciences* **2009**, *6* (4), 515–533.
- (67) Peterson, B. J.; Holmes, R. M.; McClelland, J. W.; Vörösmarty, C. J.; Lammers, R. B.; Shiklomanov, A. I.; Shiklomanov, I. A.; Rahmstorf, S. Increasing river discharge to the Arctic Ocean. *Science* **2002**, 298 (5601), 2171–2173.
- (68) Shadwick, E. H.; Trull, T. W.; Thomas, H.; Gibson, J. A. E., Vulnerability of polar oceans to anthropogenic acidification: Comparison of arctic and antarctic seasonal cycles. *Sci. Rep.* **2013**, 3.
- (69) McGuire, A. D.; Anderson, L. G.; Christensen, T. R.; Dallimore, S.; Guo, L.; Hayes, D. J.; Heimann, M.; Lorenson, T. D.; Macdonald, R. W.; Roulet, N. Sensitivity of the carbon cycle in the Arctic to climate change. *Ecol. Monogr.* **2009**, *79* (4), 523–555.
- (70) Zine, S.; Boutin, J.; Font, J.; Reul, N.; Waldteufel, P.; Gabarró, C.; Tenerelli, J.; Petitcolin, F.; Vergely, J. L.; Talone, M. Overview of the SMOS sea surface salinity prototype processor. *IEEE Trans. Geosci. Rem. Sens.* **2008**, 46 (3), 621–645.
- (71) Bélanger, S.; Ehn, J. K.; Babin, M. Impact of sea ice on the retrieval of water-leaving reflectance, chlorophyll *a* concentration and inherent optical properties from satellite ocean color data. *Rem. Sens. Environ.* **2007**, *111* (1), 51–68.
- (72) Varkey, M. J.; Murty, V. S. N.; Suryanarayana, A. Physical oceanography of the Bay of Bengal and Andaman Sea. *Oceanogr. Mar. Biol.: Annu. Rev.* **1996**, *34*, 1–70p.
- (73) Vinayachandran, P. N.; Murty, V. S. N.; Ramesh Babu, V. Observations of barrier layer formation in the Bay of Bengal during summer monsoon. *J. Geophys. Res.: Oceans* **2002**, *107* (C12), SRF-19.
- (74) International CLIVAR Project Office Understanding The Role Of The Indian Ocean In The Climate System—Implementation Plan For Sustained Observations; International CLIVAR Project Office: 2006.
- (75) Sarma, V. V. S. S.; Krishna, M. S.; Rao, V. D.; Viswanadham, R.; Kumar, N. A.; Kumari, T. R.; Gawade, L.; Ghatkar, S.; Tari, A. Sources and sinks of CO_2 in the west coast of Bay of Bengal. *Tellus B* **2012**, *64*, 10961.
- (76) Madhupratap, M.; Gauns, M.; Ramaiah, N.; Prasanna Kumar, S.; Muraleedharan, P. M.; De Sousa, S. N.; Sardessai, S.; Muraleedharan, U. Biogeochemistry of the Bay of Bengal: Physical, chemical and primary productivity characteristics of the central and western Bay of Bengal during summer monsoon 2001. *Deep Sea Res., Part II* 2003, 50 (5), 881–
- (77) Ittekkot, V.; Nair, R. R.; Honjo, S.; Ramaswamy, V.; Bartsch, M.; Manganini, S.; Desai, B. N. Enhanced particle fluxes in Bay of Bengal induced by injection of fresh water. *Nature* **1991**, *351* (6325), 385–387.
- (78) Ramaswamy, V.; Nair, R. R. Fluxes of material in the Arabian Sea and Bay of Bengal—Sediment trap studies. *Proc. Indian Acad. Sci., Earth Planet. Sci.* **1994**, *103* (2), 189–210.
- (79) Gomes, H. R.; Goes, J. I.; Saino, T. Influence of physical processes and freshwater discharge on the seasonality of phytoplankton regime in the Bay of Bengal. *Continental Shelf Research* **2000**, *20* (3), 313–330.
- (80) Sabine, C. L.; Key, R. M.; Feely, R. A.; Greeley, D. Inorganic carbon in the Indian Ocean: Distribution and dissolution processes. *Global Biogeochem. Cycles* **2002**, *16* (4), 1067.
- (81) Biswas, H.; Mukhopadhyay, S. K.; De, T. K.; Sen, S.; Jana, T. K. Biogenic controls on the air-water carbon dioxide exchange in the Sundarban mangrove environment, northeast coast of Bay of Bengal, India. *Limnolo. Oceanogr.* **2004**, *49* (1), 95–101.

- (82) PrasannaKumar, S.; Sardessai, S.; Ramaiah, N.; Bhosle, N. B.; Ramaswamy, V.; Ramesh, R.; Sharada, M. K.; Sarin, M. M.; Sarupria, J. S.; Muraleedharan, U. Bay of Bengal Process Studies Final Report; NIO: Goa, India, 2006; p 141.
- (83) Akhand, A.; Chanda, A.; Dutta, S.; Manna, S.; Hazra, S.; Mitra, D.; Rao, K. H.; Dadhwal, V. K. Characterizing air—sea CO₂ exchange dynamics during winter in the coastal water off the Hugli-Matla estuarine system in the northern Bay of Bengal, India. *J. Oceanogr.* **2013**, 69 (6), 687–697.
- (84) Burke, L. M.; Maidens, J. Reefs at Risk in the Caribbean; World Resources Institute: Washington, DC, 2004.
- (85) Langdon, C.; Atkinson, M. J., Effect of elevated pCO₂ on photosynthesis and calcification of corals and interactions with seasonal change in temperature/irradiance and nutrient enrichment. *J. Geophys. Res.: Oceans* **2005**, *110*, (C9).
- (86) Aschbacher, J.; Milagro-Pérez, M. P. The European Earth monitoring (GMES) programme: Status and perspectives. *Rem. Sens. Environ.* **2012**, *120*, 3–8.
- (87) Berger, M.; Moreno, J.; Johannessen, J. A.; Levelt, P. F.; Hanssen, R. F. ESA's sentinel missions in support of Earth system science. *Rem. Sens. Environ.* **2012**, *120*, 84–90.
- (88) Drusch, M.; Del Bello, U.; Carlier, S.; Colin, O.; Fernandez, V.; Gascon, F.; Hoersch, B.; Isola, C.; Laberinti, P.; Martimort, P. Sentinel-2: ESA's optical high-resolution mission for GMES operational services. *Rem. Sens. Environ.* **2012**, *120*, 25–36.
- (89) Donlon, C.; Berruti, B.; Buongiorno, A.; Ferreira, M. H.; Féménias, P.; Frerick, J.; Goryl, P.; Klein, U.; Laur, H.; Mavrocordatos, C. The global monitoring for environment and security (GMES) sentinel-3 mission. *Rem. Sens. Environ.* **2012**, *120*, *37*–57.
- (90) IOCCG. http://www.ioccg.org/sensors/GOCI.html (accessed August 27, 2014).
- (91) Reul, N.; Saux-Picart, S.; Chapron, B.; Vandemark, D.; Tournadre, J.; Salisbury, J., Demonstration of ocean surface salinity microwave measurements from space using AMSR-E data over the Amazon plume. *Geophys. Res. Lett.* **2009**, *36*, (13).
- (92) Sabia, R.; Fernández-Prieto, D.; Donlon, C.; Shutler, J.; Reul, N. In A Preliminary Attempt to Estimate Surface Ocean pH from Satellite Observations; IMBER Open Science Conference: Bergen, Norway, 2014.
- (93) Willey, D. A.; Fine, R. A.; Millero, F. J. Global surface alkalinity from Aquarius satellite. In *Ocean Sciences Meeting*, Honolulu, HI, 2014.