Editor Comments:

I agree with both reviewers that the manuscript has significantly improved in this revision. However, both reviewers are concerned that the authors have not addressed major comments from the first reviews adequately. The authors should carefully read the reviews and make sure that they discuss the effect of filling the gaps of data etc. Besides addressing the comments from both reviewers, the authors should also discuss how the different parameters of their approach were chosen. For example, for filling in the gaps they iteratively add 5% of vectors with missing data. Why 5%? Has this been chosen from parameter tuning? Also, how are the errors propagated in this approach? In later iterations, the results are based only little on observations and mainly on the model. This should be reflected in the error. I am looking forward to a revised revision of this manuscript, adequately dealing with all the reviewers' comments.

We have taken care to address the comments of the reviewers. The filling method has been more detailed and has also been submitted for a separate conference paper dealing with the completion of glider data profiles.


Parard et al. provide a revised version of their previously submitted manuscript. Having read and reviewed both versions, I am glad to say that the quality of the revised version immensely improved. The authors managed to sort out almost all language and editorial issues (besides a few typo's listed below) and now provide a very well structured introduction and method section. Furthermore, the authors added some additional results regarding the seasonal cycle of the sea surface pCO2 in the Baltic Sea – Overall, a great improvement.

There are however a few more issues, most of them raised in my first reviewers comment, which have not or only partly been addressed. Hence, I do still think that some major and minor comments need to be addressed before the manuscript can be accepted for final publication.

Major Comments:

While the authors have done some great effort to describe why time is used as an additional parameter, it is still unclear to the reader why. You provide a perfect example yourself: Even though in November and May the predictor data (SST, etc.) do have the same value, the pCO2 itself does not have the same value. Time itself does not change the pCO2. Imagine a steady Baltic Sea system where all potential and known pCO2 drivers are constant. In this system pCO2
would be constant in time as well. Its rather other processes that vary in time that are not considered as driver in your SOMLO model. However, this or these process(es) are not known (I assume) hence time replaces them. I am not suggesting here that you include more drivers, however, I do believe it needs to be clarified why time is important in your case.

We further explained and clarified the reason for the inclusion of the time of year as an input parameter in our method. (Lines 239 to 253 p 9)

.) In my last review I mentioned that figure captions are very short. More information in the figure caption is needed to understand the figures without constantly going back to the text to find the description. That interrupts the flow. Furthermore, please improve the resolution of figures 5, 13 and 15 as they are very difficult to read.

We did our best to improve the resolution of the figures.

.) I very much appreciate your effort in providing a PCA analysis, but I am not quite sure about the point you try to make. First of all, Figure 3 is very difficult to understand with the information you provide (although BG is a high quality journal, I am sure not every reader is familiar with a PCA analysis). The same counts for table 1. Besides that, table 1 illustrates that 76% of the variability can be explained by the first 3 loadings (or axes). Why not perform a PCA analysis a-priory and reduce the number of drivers to e.g. the first 3 loadings. This would strongly reduce your degrees of freedom while explaining the majority of all the variance in your system?

The problem we would face by keeping only 3 axes of the PCA is that the 24% of the total variation would remain unused. This much variation can be significant enough when classifying, and since the percentages after the first 2 axes are considered are of similar magnitude we did not wish to reduce it. The PCA section of the article has been further detailed to better reflect this (Line 254-267 p 9).

Minor comments:

.) in Section 2.2. you have 3 pCO2 sources, however, source 3 is not measured directly but calculated from pH and TALK. Due to the uncertainty in the calculation, have these data been considered more uncertain or have they been treated equally?

The third source of pCO2 is treated like the other source, we compared the data from the third source and the other when the position was less than 0.2 ° and the correlation is good with 0.98 and the standard deviation is 9 μatm due to upwelling event and 1.5 without upwelling event. So
we consider that the pCO2 from the tree sources can be treated equally. We add a comment in the paper (Lines 123 to 128 p 5)

.) Abstract line 6: Sasse et al 2013 (final version has been published in 2013)
We corrected this error. (line 6 p 1)

.) Abstract line 9: change “Sea” to “sea”
We corrected this error. (line 9 p1)

.) Abstract line 10: change “Depth” to “depth”
We corrected this error. (line 10 p 1)

.) Introduction lines 49-50: Some other, recent basin scale studies that could be mentioned are Schuster et al 2013 (BG), Nakaoka et al 2013 (BG) and Landschützer et al 2013 (BG)
We included these references. (lines 49 to 50 p 2)

.) section 2.3 line 147: I assume the +- sign at the MD estimate should be removed
We corrected this. (line 152 p 6)

.) section 2.3 line 156-157: “as it is taken into account when classifying and calculat in the MLR parameters” - How?
We rewrote this sentence. (lines 160 to 162 p6)

.) section 2.3 line 199: I would argue that “quite good” can be changed to “good” if you consider that previously you already calculated R values of 0.9 and more.
We corrected this error. (line 204 p 7)

.) section 2.3. line 203: This sentence is a repeat and can be deleted
We removed this sentence (line 207 p8)

.) section 2.4. line 240: change “PCO2” to “pCO2”
We corrected this error. (line 244 p 9)

.) section 3.2.1. line 368: You mention validation and testing data. Please provide a short description on their difference and how they are used later on
We added a sentence explaining this (Line 381 to 387 p15).

.) Figure 7 and Figure 8: I am a bit puzzled why there is such a big change in NPP, from a maximum of about 70 mgC/m2/yr in Figure 7 to 400 mgC/m2/yr in Figure 8. Please clarify
The 70 and 400 are not the value of NNP. It is a histogram so it represents the number of data on Y axis and the value on X axis. We changed the legend to clarify this.

.) Figure 12: there are a few “outliers” left with differences of more than 100 muatm. Have you considered a outlier removal criterion, e.g. Chavenets criterion?

Before to construct the map we already removed the outlier values in our data set. We tested the Chavenet’s criterion and we found that only 2 outliers remained with an α=0.1. We add a sentence about the outlier (Line 268 to 270 p 9)

.) section 3.2.3 line 458: you compare the model output to your estimates from 1998-2011 in Figure 14. In the text you say 1998-2009
The comparison is indeed made between 1998-2009. We changed the figure.

.) section 3.2.3 line 461 the word “strongly” is overconfident given an R value of 0.6
We corrected this error. (line 476 p 21)

.) section 4 line 507 and reference list: Landschützer et al 2013 has been published in BG (in the reference it still refers to the discussion paper – same as with Sasse 2013)
We corrected this error. (line 548 p25)

**Review of the revised version of Parard et al. [2014]**

Overall evaluation:

The manuscript has improved significantly and I am confident that it can be published after some revisions. However, some of my major points of criticism regarding the validation of the method were not addressed correctly in the revision. Please see below.

Accuracy of pCO2 map and errors in remotely sensed data:

In response to my initial review the authors state:

*The method, as presented compensates for these types of errors since, if the errors are recurrent and constant in nature, the method will “learn” them and produce results based on these recurring types of errors.*

This is exactly the problem: The errors are not recurrent and constant in nature!
The errors are random in nature. The validation provided for the individual parameters in section 2.3 clearly shows that we are not just dealing with a simple offset (otherwise the correlations
would all be 1.0). In the best case scenario, the random errors in the remote sensing data translate into random errors in the pCO2 that cancel each other out when averaged (e.g. as done for Figure 14). But not even this is guaranteed. Definitely do these random errors affect the accuracy of the pCO2(x,y,t) estimate and we need to know to what degree. A statement like “As for the chlorophyll, the bias is applied to all years so it does not affect the estimation of pCO2” (line 165) completely misses the point and does not provide the reader with confidence that the authors were seeking a thorough error estimate.

The validation presented in section 2.3 provides some estimate of the magnitude of the error for each parameter. A random noise with the standard deviation given by the validation can be added to each parameter and the SOMLO method can then be repeated with noisy parameters. This would provide at least a first order estimate of the additional error coming from the remote sensing of parameters.

In a similar manner my questions regarding the effect of filling in data gaps was not addressed by the authors. They state:

The completion of the missing data was a very important step in the reconstruction of the pCO2, since without it we would not have been able to apply an MLR and would have been forced to replace the missing value based on the average pCO2 value of the class to which the data would have been projected without being completed.

I understand that it was necessary. But my questions is how does it affect the accuracy of the method?

The validation presented in Figures 12+13 simply ignores errors implicitly introduced by the data and the data handling.

Repeating the statement from my first review:
Presenting a thorough error analysis for the pCO2 maps is as important as presenting the actual maps!

The random 90% / 5% / 5% separation of the data set for the purpose of validation needs to be repeated an appropriate number of times to make sure that the error estimate is robust and not just the result of a lucky or unlucky splitting. Following this review we have performed a number of tests and included many clarifications in the text in order to give better estimations and understanding of the propagation of errors in our method.
In regards to our comment on the method’s “learning” of errors, what we refer to is that by classifying our multi-dimensional data through SOMs, if there is a low level of noise which, when averaged is nullified, then each class is in fact taking into account this noise. Indeed each class’s referent vector can be considered to be generated, in the latter stages of the training where each class stops affecting each other class, by the average value of all the elements captured by that particular class. It is therefore able, when capturing a noisy data to correctly attribute it to the class it would have belonged or a neighboring class on the topological map. There will of course be some errors that are then induced by applying the MLR on the data, and we therefore have added some values to present how much the method is sensitive to noise and to the completion method.

As far as the separation of data, that was an error that had escaped us from previous versions and we have clarified that we do cross-validations by randomly selecting 90% data, training the method and then calculating the performances of SOMLO on the remaining 10%, iterating this process and presenting the average values of RMS we obtained. This has been clarified in the text (lines 381 to 387 p15)

The results of our analysis have been presented in section 3.2.4 of the article “Sensitivity Analysis”. (p22-23)

Figure quality:

The quality of the figures is still not acceptable! At the normal print-out size I am able to distinguish between individual pixels with my naked eye. Given the simplicity of the plots it should be no problem to view them in great resolution on 300% zoom if they were true vector graphics.

For example, I assume there is text in the upper left corner of the lower panel of Figure 5 but it is absolutely not recognizable.

Since the readability of figures was claimed by both reviewers I do not understand why the authors do such a poor job in improving them.

We did not notice any problems in our figures when inspecting them in our versions of the manuscript, however we increased the resolution on all the figures mentioned and they hopefully are of sufficient quality now.

Miscellaneous:
Please explain what you mean by “the non-linearity of the emission and absorption system”
We have clarified this in the document. (line 41 to 43 p2)

Please provide a reference.
We added a reference. (line 91 p3)

Do we need the black pixels over land?

The black pixel on the land are not necessary but we need a precise representation of coastline. When selecting the options in matlab to generate a really precise coastline, we also obtain a really good definition for the land topography (Figure 1). We opted to maintain the inland topography to preserve the high resolution of the coastlines.

Figure 1: Comparisons of the low resolution coast with data (left down) and without (left up) and the high resolution with data (right down) and without data (right up)
Maybe the location can be indicated in Figure 1.
We added the position with the black arrow (figure 1)

line 126: remove “from”
We removed this (line 131 p5)

line 209-210
“density measurements”? We made the statement clearer (line 213 p8)
line 248-257 and corresponding Figure 3:
The explanation provided is unclear. What is the distribution of the parameters in Figure 3 telling us?
We add information about this figure in the text (line 248 to 257 p 9)

line 291: The explanation is very good but what is “argmax(i)”?
It corresponds to the argument i that maximizes the following equation. We added a notation in the text. (line 303 p12)

line 451: remove one “)” We corrected this mistake (line 468 p 21).

Figure 15: aspect ratio is off (not to mention the resolution)
We improved this figure and hope that this has been resolved.

line 503: “This led to our having to complete ...”? We rewrote this sentence (line 540 to 543 p 25)
line 511-513:
I assume that spatial pCO2 variability might also be larger in the Baltic Sea compared to open-ocean. This would also lead to a larger error.
We agreed with the reviewer we added a sentence concerning this (line 549-552 p 25)

References: The correct spelling is “Lefèvre”
We corrected this (line 639 p28)

Please remove dispensable dots and letters in reference for Takahashi et al. [2009] and some other references.
We corrected all the references.
Remote sensing the sea surface CO$_2$ of the Baltic Sea using the SOMLO methodology

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Abstract. Studies of coastal seas in Europe have noted the high variability of the CO$_2$ system. This high variability, generated by the complex mechanisms driving the CO$_2$ fluxes, complicates the accurate estimation of these mechanisms. This is particularly pronounced in the Baltic Sea, where the mechanisms driving the fluxes have not been characterized in as much detail as in the open oceans. In addition, the joint availability of in situ measurements of CO$_2$ and of sea-surface satellite data is limited in the area. In this paper, we used the SOMLO (Sasse et al., 2013) methodology, which combines two existing methods (i.e., Self-Organizing-Maps and multiple linear regression) to estimate the ocean surface partial pressure of CO$_2$ (pCO$_2$) in the Baltic Sea from the remotely sensed sea surface temperature, chlorophyll, colored dissolved organic matter, net primary production, and mixed-layer depth. The outputs of this research have a horizontal resolution of 4 km and cover the 1998–2011 period. These outputs give a monthly map of the Baltic Sea on a very fine spatial resolution. The reconstructed pCO$_2$ values over the validation dataset have a correlation of 0.93 with the in situ measurements and a root mean square error of 38 µatm. Removing any of the satellite parameters degraded this reconstructed CO$_2$ flux, so we chose to supply any missing data using statistical imputation. The pCO$_2$ maps produced using this method also provide a confidence level of the reconstruction at each grid point. The results obtained are encouraging given the sparsity of available data, and we expect to be able to produce even more accurate reconstructions in coming years, given the predicted acquisition of new data.

1 Introduction

The ocean plays an important role in the global carbon budget. It acts as a major carbon sink for anthropogenic carbon dioxide (CO$_2$) emitted to the atmosphere from fossil fuel burning, cement
production, biomass burning, deforestation, and various land use changes. The ocean is currently slowing the rate of climate change, having absorbed approximately 30% of human emissions of CO₂ to the atmosphere since the industrial revolution (Stocker et al., 2013). The exchange of CO₂ between coastal environments and the atmosphere is a significant part of the global carbon budget (e.g. Borges et al., 2005; Chen and Borges, 2009; Laruelle et al., 2010). While continental shelves represent only 7% of the oceanic surface area and less than 0.5% of the ocean volume, the estimated overall sink of CO₂ in the continental shelf sea is ~0.22 Pg C yr⁻¹ (Laruelle et al., 2010), corresponding to 16% of the open oceanic sink (Takahashi et al., 2009). These estimates are subject to great uncertainty related to sparse data coverage in time and space. Monitoring the oceanic partial pressure of CO₂ (pCO₂) at monthly and seasonal time scales is essential for estimating the regional and global air–sea CO₂ fluxes and reducing this uncertainty. For technical and budgetary reasons, in situ measurements of marine pCO₂ are sparsely distributed in time and space. However, over the last decade, technical improvements and cooperation with the shipping industry have allowed for the installation of several autonomous monitoring systems aboard commercial vessels routinely crossing the ocean basins. These instruments make quasi-continuous measurements, allowing regional analysis of the highly variable spatial and temporal distributions of pCO₂ (e.g. Lefèvre et al., 2004; Lüger et al., 2004; Corbière et al., 2007; Schneider et al., 2003). The Baltic Sea, a semi-enclosed sea in Northern Europe, is relatively well monitored and has been studied for several decades (Meier et al., 2014). Despite the increased number of measurements made in the Baltic Sea, assessing the carbon fluxes in the Baltic Sea remains particularly challenging due to the nonlinearity of the emission and absorption system. This nonlinearity is complicated by a combination of varying salinity, varying river input of DOC, and large general variability due to the strong seasonal cycle in the region. Using new methodologies could generate additional information from the relatively limited number of existing measurement data. Neural network techniques are empirical statistical tools that somewhat resolve the nonlinear and often discontinuous relationships among proxy parameters without any a priori assumptions. In the past decade, several authors have reported the application of a neural network technique to basin-scale pCO₂ sea analysis (Lefèvre et al., 2005; Jamet et al., 2007; Friedrich and Oschlies, 2009; Telszewski et al., 2009; Landschützer et al., 2013; Nakaoka et al., 2013; Schuster et al., 2013), concentrating mainly on the North Atlantic Ocean. Most recently, Telszewski et al. (2009) successfully applied a neural network technique based on a Self-Organizing Map (SOM) to reconstruct the seawater pCO₂ distribution in the North Atlantic (10.5° to 75.5°N, 9.5°E to 75.5°W) for three years (i.e., 2004 to 2006) by examining the nonlinear/discontinuous relationship between pCO₂ sea and the ocean parameters of sea surface temperature (SST), mixed-layer depth (MLD), and chlorophyll a concentration (Chl). In this paper, we applied the SOMLO methodology (self-organizing multiple linear output), which creates an SOM classification of the available explicative oceanic parameters in the Baltic Sea and then calculates multiple linear regression (MLR) parameters for estimating the pCO₂ from the elements belonging to each class separately. The major benefit
of this methodology is that it allows the use of a linear model (i.e., the MLR) despite the nonlinear relationship between pCO$_2$ and its explicative parameters, by using the SOM classifier. The SOM classifier can determine the region of the multidimensional data space in which to perform the linear regression. If the classification is fine enough, each region will represent a single type of relationship between the pCO$_2$ and the explicative data, a type that can be reproduced by an MLR. Due to the temporal and spatial limitations of the in situ pCO$_2$ data, the satellite data can help estimate the air–sea CO$_2$ fluxes over the entire Baltic Sea. The satellite data have a higher spatial coverage of the Baltic Sea, allowing estimation of the pCO$_2$ from in situ data using the SOMLO method. Chierici et al. (2009) demonstrate that in the North Atlantic Ocean, SST, Chl, and MLD contributed significantly to the estimation of pCO$_2$ from a linear relationship. Based on this idea, we applied these parameters to the Baltic Sea, adding two other parameters, i.e., Net Primary Production (NPP) and Colored Dissolved Organic Matter (CDOM), that provide information about the biological activity occurring in summer. From this, we develop pCO$_2$ algorithms applicable to the Baltic Sea using in situ pCO$_2$ values; remotely sensed SST, Chl, and CDOM; modeled MLD and NPP; and time.

The manuscript is structured in four parts. First, we present a synopsis of the problem studied, including existing studies of pCO$_2$ reconstruction in other maritime regions. Next we present the available data and briefly describe the methodology used. In the third part of the article we present our results, namely, the topological maps obtained and the reconstructions performed with them. We conclude the article by discussing the results obtained and future possible improvements of the method used.

2 Materials and methods

2.1 Study area

The Baltic Sea is a semi-enclosed sea with limited exchange with the North Atlantic through the North Sea–Skagerrak system. Previous investigations of the Baltic Proper found large temporal and spatial variability of pCO$_2$. The amplitude of the annual pCO$_2$ cycle varies significantly depending on the region, ranging from 400 µatm in the northeastern Baltic Proper to 120 µatm in the transition areas to the North Sea (Schneider and Kaitala, 2006). The Baltic Sea receives significant river runoff from surrounding land (a total of approximately 15,000 m$^3$ s$^{-1}$ (Bergstrom, 1994) and net precipitation of approximately 1500 m$^3$ s$^{-1}$ (Omstedt et al., 2004). This large freshwater addition brings large amounts of nutrients and inorganic and organic carbon to the Baltic Sea basin (Omstedt et al., 2004; Hjalmarsson et al., 2008). The biogeochemical processes in the Baltic Sea marine environment are controlled mainly by the biological production and decomposition of organic matter occurring in the context of the region’s hydrography (Siegel and Gerth, 2012). Physical forcing controls the water transport, stratification, temperature, and salinity in the Baltic Sea; these factors then influence the nutrient and carbon distribution, thereby affecting biogeochemical processes. We divide the Baltic
Figure 1. The dashed red lines divide the Baltic Sea into three basins: CP: Central part, BB: Gulf of Bothnia, GF: Gulf of Finland. Monthly data are available from 1998 to 2011 in the Baltic Sea. The colorbar shows the pCO$_2$ values in $\mu$atm. The black arrow show the position of the SAMI sensor.

Sea into three basins, i.e., the central part (CP), the Gulf of Bothnia (GB), and Gulf of Finland (GF), as shown in Figure 1. The Baltic Sea has an average depth of 55 m and a maximum depth of 460 m at the Landsort Deep (Wesslander 2011).

2.2 pCO$_2$ observations

To compile the pCO$_2$ maps, we use measured data from three sources.

1. The Östergarnsholm site: This site is located next to the small island of Östergarnsholm in the central Baltic Sea and is further described by (Rutgersson et al. 2008; Norman et al. 2013a). The island is situated 4 km from the east coast of the larger island of Gotland. SST and pCO$_2$ are measured semi-continuously 4 m below the sea surface using a submersible autonomous moored instrument (SAMI) CO$_2$ sensor moored at a buoy 1 km southeast of the tower situate on the island. In addition, SST is also measured using a wave rider buoy (operated by the Finnish Meteorological Institute) at 0.5 m depth situated approximately 4 km southeast of the tower.
2. Cargo ship: This dataset derives from continuous measurements of the surface water $pCO_2$ made in the Baltic Sea using a fully automated measurement system deployed on a cargo ship. The Leibniz Institute for Baltic Sea Research, Warnemünde Germany (IOW—Institut für Ostseeforschung Warnemünde) has made continuous measurements of $pCO_2$ at 5 m depth aboard the cargo vessel Finnpartner. This ship crosses between Lübeck and Helsinki at a two-day interval, alternately crossing the eastern and western Gotland Sea (Schneider and Kaitala 2006 [Schneider et al.] 2009). Data from Finnpartner were acquired between July 2003 and December 2005.

3. Swedish Meteorological and Hydrological Institute (SMHI) database Svenskt Havsarkiv (SHARK): pH, (measured using the method of Grasshoff et al. (1999)) and Total Alkalinity (TA) (measured using potentiometric titration as described by Grasshoff et al. (1999)) are measured continuously at a monthly or semi-monthly resolution in the Baltic Sea at various stations. All measurements are made at a depth of 5 m depth. The uncertainty of the pH is ±0.03 pH units and of the TA is ±5% (Wesslander et al. 2009). $pCO_2$ is estimated from the pH, TA, salinity, and temperature measurements using the standard CO2SYS program (Lewis and Wallace 1998) with the equilibrium constant from (Weiss 1974) and (Merbach et al., 1973) as refitted by Dickson and Millero (1987) as in Wesslander et al. (2009). The $pCO_2$ estimation were compared with the other source of $pCO_2$ data, all the data were compared in time (less than 1 hour) and position (around less than 0.2°), the correlation coefficient (R) give 0.98 and the standard deviation (STD) is 9 µatm. The high standard deviation is explain by the presence of upwelling with high $pCO_2$ present near the coast on the SAMI sensor measurement, when the we remove the upwelling event the STD is 1.5 µatm.

2.3 Remote sensing data

The satellite data used in this study are from various sources. We use a monthly temporal resolution and a spatial resolution based on the lowest spatial resolution of our datasets, i.e., based on the lower spatial resolution CDOM dataset.

We obtain values for five parameters from various sources:

**SST** Sea Surface Temperature: Several datasets are used for SST, and we combine two types of datasets for 2007 and 2011. For 2005–2011, we use data from the Federal Maritime and Hydrographic Agency (BSH), which processed data from AVHRR-NOAA, and data from the Group for High-Resolution Sea Surface Temperature (GRHSST) dataset for the Baltic Sea, 2007–2011. The spatial resolution is 0.03 ° at a daily temporal resolution (http://podaac.jpl.nasa.gov/dataset/DMI-L4UHfnd-NSEABALTIC-DMI_OI). For 1998–2004, the data come from a reanalysis of the NOAA/NASA Advanced Very High Resolution Radiometer (AVHRR) data stream conducted by the University of Miami’s Rosenstiel School of Marine and Atmospheric Science (RSMAS) and the NOAA National Oceanographic Data Center (NODC). This
dataset consists of the monthly average SST (in °C) over the zone, with a spatial resolution of 4 km, extracted from version 5.2 of the AVHRR Pathfinder project (Casey et al., 2010) (http://www.nodc.noaa.gov/SatelliteData/pathfinder4km/). The various SST datasets were compared with measured SMHI temperature data at a monthly resolution from 1998 to 2011, giving a correlation coefficient (R) of 0.99 and a mean difference (MD) of 0.05°C between the two datasets. The difference observed between the measured and satellite data between 1998 and 2004 give a value (R = 0.99 and MD = 0.09°C) near the difference between 2005 and 2011 (R = 0.99 and MD = 0.14). The two SST datasets used between 2007 and 2011 were also compared with the SAMI sensor data at a daily resolution, giving a good correlation for the BSH data (R = 0.95 and MD = 0.06°C) and the GRHSST data (R = 0.95 and MD = 0.08°C).

Chl Chlorophyll a: This dataset consists of monthly averages from the following sensors: Sea-viewing Wide Field-of-view Sensor (SeaWiFS) (Sept. 1998–Dec. 2002) with 4-km spatial and monthly temporal resolutions and Moderate Resolution Imaging Spectroradiometer (MODIS-Aqua) (Jul. 2002–Jun. 2011) with 4-km spatial and monthly temporal resolutions (Casey et al., 2010). A lognormal distribution was assumed for the Chl data. Comparison with SMHI measurement data and in situ data (personal communication from Dr. Tiit Kutser) give R = 0.67 and MD = 7 mg m⁻³. The chlorophyll levels from the satellite data seem to be overestimated compared with the in situ data. Since we use the same dataset over the whole study period, this bias was learned during the classification process and and the MLR parameters are calculated considering that any further input will include that bias.

CDOM Colored Dissolved Organic Matter values come from MODIS-Aqua 4-km monthly average data. The CDOM index quantifies the deviation in the relationship between the CDOM and Chl concentrations, where 1.0 represents the mean relationship for Morel and Gentili (2009) case 1 waters, and values above or below 1.0 indicate an excess or deficit, respectively, in CDOM relative to the mean relationship. The algorithm and its application are fully described by Morel and Gentili (2009). In situ CDOM data (personal communication from Dr. Tiit Kutser) give a lower correlation coefficient and a low average difference (R = 0.48, MD = 2.3). As for the chlorophyll, the bias is applied to all years so it does not affect the estimation of pCO₂.

NPP: Net Primary Production values come from two data sources. The first dataset comes from the Environmental Marine Information System (EMIS): The EMIS model is depth integrated but allows for depth-dependent variability in the diffuse attenuation coefficient, which is calculated from a multiple-component semi-analytical inversion algorithm (Lee et al., 2005). The primary production calculation is based on the formulation obtained through dimensional analysis by Platt and Sathyendranath (1993). The photosynthetic parameters are assigned by the combined use of a temperature-dependent relationship for the maximum growth rate Ep-pley (1972) and a variable formulation to retrieve the C:Chl ratio following the empirical re-
relationship of Cloern et al. (1995). The EMIS dataset comprises monthly average values from October 1997 to September 2008. The second dataset, for 2009–2011, uses the Vertically Generalized Production Model (VGPM) of Behrenfeld and Boss (2006) as the standard algorithm. VGPM is a “chlorophyll-based” model that estimates net primary production from chlorophyll using a temperature-dependent description of chlorophyll-specific photosynthetic efficiency. For VGPM, net primary production is a function of chlorophyll, available light, and photosynthetic efficiency. VGPM uses MODIS-Aqua chlorophyll and temperature data, SeaWiFS photosynthetically active radiation (PAR) data, and estimates of the euphotic zone depth from a model developed by Morel and Berthon (1989) and based on chlorophyll concentrations. For the NPP for 2009–2011, the observed maximum value was limited to 10 to be comparable to the data for 1998–2008. Validation of NPP was difficult due to the number of data available in the area. Comparison of the two datasets gives similar values and seasonal cycles. We compared the seasonal cycles between 1998–2008 and 2009–2011, obtaining values on the same order of magnitude.

MLD Mixed-Layer Depth: There are also two sources for the MLD data. Monthly averages from 1998 to 2007 come from a 3D hydrodynamic model currently used at the Joint Research Centre/Institute for Environment and Sustainability (JRC/IES), i.e., the public-domain General Estuarine Transport Model (GETM; www.getm.eu), which has its roots in developments at the JRC/IES Burchard and Bolding (2002). GETM simulates the most important hydrodynamic and thermodynamic processes in coastal and marine waters and includes flexible vertical and horizontal coordinate systems. Different turbulence schemes are incorporated from the general ocean turbulence model (GOTM; www.gotm.net). For 2008–2011, we use data from the carbon-based production model at a monthly resolution (Behrenfeld et al., 2005). The MLD was estimated from SMHI temperature and salinity profile measurements using the density criterion of Boyer Montégut et al. (2004); the comparison between the model estimation and estimated SMHI MLD is good (R = 0.63 and MD = 17 m). Between 1998 and 2007, the correlation coefficient is higher (R = 0.8) than between 2008 and 2011 (R = 0.5). Nevertheless, the MD is lower for the second data source, i.e., 20 m versus 11 m. This can be explained by the available data coverage: 63% of the in situ data cover the 1998–2004 period. From 2008 to 2011, the maximal value is below 80 m between 1998 and 2007, so for the in situ data estimation, we replace every value above 80 m with “not a number.”

In the Baltic Sea, there are many gaps in the satellite data; this is due to the high proportion of coastal waters where satellite data are less reliable, and to the frequent large-scale cloud coverage. To increase the total number of our data, we used a monthly temporal resolution and a method to improve the spatial distribution of the data. For statistical analysis, the irregularly spaced density of the measurements were first uniformly resampled. To this end, Gaussian grinding was used, as described by Greengard and Lee (2004); Dutt and Rokhlin (1995). The data points of the original series are
convolved using a Gaussian kernel. As a result, the data points are smeared over their neighboring equi-spaced points, which are more densely distributed. This method produces more realistic values than does simple interpolation, particularly when there are many data gaps (Schomberg and Timmer 1995). There is no discontinuity between the different datasets, but NPP, CDOM, and Chl data are missing for January and December, so no reconstruction can be performed for these months.

2.4 Data available

The positions and values of all the in situ pCO$_2$ data are shown in Figure 1. We use the spatial resolution of the parameter with the lowest resolution for the final dataset chosen (i.e., CDOM). A monthly temporal resolution is used for this study. Rutgersson et al. (2008) demonstrate that the agreement between SAMI sensor data and the ship data from Finnpartner (near the mooring maximum of 23 km) is quite good. This good agreement is confirmed by the comparison between pCO$_2$ data from the SAMI sensor, the data surrounding the mooring (0.2°), and the other datasets. These analyses give a good correlation factor of 0.98. The in situ data are available mainly for the central basin, but the number of data for the Gulf of Bothnia is very low, coming from two SMHI stations. The in situ pCO$_2$ data are well distributed over the twelve months (Figure 2). January is the month for which the number of data is lower (i.e., below 80), but the other months have 110–155 data points each. In our case, each in situ data point is characterized by SST, Chl, CDOM, NPP, and MLD as well as information on the date the measurements were made. This temporal information was normalized by sine and cosine, as follows:

\begin{align}
\text{time}(\text{cosine}) &= \cos\left(\frac{\text{Days} \times 2\pi}{365}\right) \\
\text{time}(\text{sine}) &= \sin\left(\frac{\text{Days} \times 2\pi}{365}\right)
\end{align}

where Days represents the Julian day.

This definition of time is used to render the values continuous over the course of the year, sidestepping the artificial numerical transition from the last day of one year to the first day of the next, to be able to situate the process in relation to its seasonality.

Although time itself is not affecting the pCO$_2$, the inclusion of time (time(cosine) and time(sine)) as a parameter is important since, in the database, some situations have similar values for SST, Chl, CDOM, NPP, and MLD, but different pCO$_2$ values. For example, in May we had a situation (SST = 9.3 °C, Chl = 0.1 mg m$^{-3}$, CDOM = 4.3, NNP = 1.7 mg C m$^{-2}$ d$^{-1}$, and MLD = 10.8 m) whose parameters were very close to those of a situation in November (SST = 9.1 °C, Chl = 0.1 mg m$^{-3}$, CDOM = 4.3, NNP = 1.6 g C m$^{-2}$ d$^{-1}$, and MLD = 10.3 m); however, the pCO$_2$ values differed greatly, being pCO$_2$ = 214 and 444 μatm, respectively, during the two situations. This dissimilarity in the pCO$_2$ values informs us that these situations are generated by other drivers than SST, Chl, CDOM, NPP, and MLD. Since these drivers can be related to seasonal patterns, we included an information on the time of the year, as a proxy that allows us to fine tune our classification.
Figure 2. Histogram showing the number of observations in function of the month between 1998 to 2011.

Inclusion of two temporal values instead of only one, even though those are highly correlated, was unavoidable in order to preserve continuity in the values obtained when changing years.

A Principal Component Analysis (PCA) was conducted to highlight the importance of the parameters in the pCO\(_2\) variability (Figure 3). PCA is a method of analysis which involves finding the linear combination of a set of variables that has maximum variance and removing its effect, repeating this successively (Jolliffe, 2002). The percent of variance explained by each axis of the PCA is shown in Table 1. The results of this PCA indicate that the percentages of variance explained by all axes beyond the first do not differ greatly, with the notable exception of the last two axes, indicating that most parameters can be discriminant in the definition of the SOM states. The need to maintain the totality of these parameters is further demonstrated by the projection of the explicative parameters in the correlation circle, where all values are close to the boundaries of the correlation circle. This informs us that they are all tied to the phenomena explaining 62% of the total variation of our data. While we could reduce the dimension of the problem by using only the projection of these variables on the first few axes, we chose to maintain all the explicative parameters presented when applying SOMLO to estimate the pCO\(_2\) in order to be able to not lose any information when performing the SOM classifications.

In total, 1445 pCO\(_2\) data are used in this study, after having removed the outliers from our data-set. These 170 outliers were beyond three standard deviations away of at least one of the explicative parameters. All parameters (i.e., SST, Chl, CDOM, NPP, and MLD) are located around each pCO\(_2\) datum. In winter (i.e., October to March), more data are missing (Table 2 column 1), particularly for Chl, CDOM, and NPP, winter being the period when it is more difficult to measure or estimate these parameters. Between April and September, the number of missing SST, Chl, CDOM, and MLD data
Figure 3. Correlation scatterplot with the representation of parameters on axes 1 and 2. The first component plane contains 62% of the total variance of the system under study, and all parameters are close to the exterior of the circle, indicating that they are all important in order to invert the pCO$_2$.

Table 1. Percent of variance explained by each axis of the PCA.

<table>
<thead>
<tr>
<th>Axis number</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percent of variance</td>
<td>42</td>
<td>20</td>
<td>14</td>
<td>9</td>
<td>8</td>
<td>4</td>
<td>2.5</td>
<td>0.5</td>
</tr>
</tbody>
</table>

is relatively low compared with the total number of data (Table 2, column 2), i.e., fewer than 3% of the total. To increase the number of data available, we completed the data by training the topological map. Further details on this are presented in section 2.5.

2.5 Methodology

The relationship between pCO$_2$ and the environmental parameters is highly nonlinear: a slight variation in some of the environmental parameters could correspond to significant variations in pCO$_2$. We chose to use the SOMLO methodology, which combines two statistical approaches: self-organizing maps (SOMs) (Kohonen 1990) and linear regression. SOMs are a subfamily of neural network algorithms used to perform multidimensional classification. A defining characteristic of SOMs is that their classes can represent a Gaussian distribution centered around the typical profile of environmental parameters, if there is high discretization of the training dataset (Dreyfus 2005). We use
Table 2. Number of missing values for each parameter for the satellite data for the October–March and April–September periods. The numbers in parentheses indicate the total data points in each period.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>October–March (685)</th>
<th>April–September (814)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SST</td>
<td>28</td>
<td>0</td>
</tr>
<tr>
<td>Chl</td>
<td>202</td>
<td>24</td>
</tr>
<tr>
<td>CDOM</td>
<td>320</td>
<td>5</td>
</tr>
<tr>
<td>NPP</td>
<td>468</td>
<td>571</td>
</tr>
<tr>
<td>MLD</td>
<td>6</td>
<td>2</td>
</tr>
</tbody>
</table>

Figure 4. The various elements used in training a self-organizing map. On the left we have the dataset used to train the SOM, which discretizes it into classes. Each class contains a referent vector containing, for each parameter, the average value of the elements comprising it, and an index of the class that informs us of its location on the topological map.

This hypothesis to classify the environmental parameter dataset, and then estimate the parameters of a linear regression for each class. In the following section, we present a brief overview of the two statistical algorithms and their application to our datasets.

2.5.1 Self-organizing maps

Self-organizing topological maps (SOMs) are a clustering method based on neural networks. They cluster a learning dataset into a reduced number of subsets, called classes, with common statistical characteristics.

Generating a SOM requires the creation of a training database that contains homogenous vectors. After a training phase, we obtain a SOM. The term "map" corresponds to a 2D matrix that stores, for each class, its referent vector, $r_i$, which approximates the mean value of the elements belonging to
Figure 5. On top we have a representation of the topological map as a lattice, while on the bottom we have a projection of data in the data space (black circles), as well as the average values of each class (red circles). Adjacent classes on the SOM lattice correspond to adjacent areas in the multidimensional data space. The red lines indicate the connections between neighboring classes.

SOMs are also called self-organizing topological maps, “topological” indicating that the SOM training algorithm forces a topological ordering of the classes in the matrix, meaning that any two neighboring classes $C_i$ and $C_j$ on the map matrix have referent vectors $r_i$ and $r_j$ that are close in the Euclidean sense in the data space.

Let us consider a vector $x$ that is of the same dimensions and nature as the data used to generate the topological map; we can find the index of the class to which it is classified by choosing: index $= \text{argmax}_i (||x - r_i||)$, (where argmax is the argument maximal) therefore assigning it to the class whose referent is closest to it in the Euclidean sense (Figure 5). A classified vector $x$ will be represented by its class index, $C_{\text{index}}$. If we are trying to classify a vector that has some missing values, the comparison is performed between the existing values of $x$ and the corresponding values of each $r_i$.

As a version of the expectation–maximization algorithm, the SOM algorithm performs an iterative training. During the early phases of this training, the referent vectors of each class are strongly affected by the changes imparted on their neighbors’ referent vectors in order to capture the shape of the data cloud. Depending on the training parameters of the SOM, in the latter phases of the training, the effect of the neighboring vectors on the determination of the referent vector can be considered null. In these cases, each referent vector approximates, locally, the mean value of the...
multidimensional Gaussian random distribution that generated the training data assigned to that class (Dreyfus, 2005).

2.5.2 Multiple linear regression

Multiple linear regression (MLR) is a modeling method that expresses the value of one response variable, V (in our study pCO$_2$), as a linear function of other explicative variables, i.e., $X = X_1, X_2, \ldots, X_i$ (in our study SST, Chl, CDOM, NPP, MLD, time$_{sine}$, and time$_{cos}$). An MLR is generally performed either to interpret the relationship between the variable y and each of the other predictive variables $X_i$, or to predict, from a dataset of vectors containing the values of $X$, the corresponding value of y. In this paper, we used both aspects of MLR.

However, to perform MLRs in the present case, we had to take into account their limitations and the nature of the problem. Specifically, to perform an MLR we are obliged to assume that the relationship between the predictor variables and the response variable is linear. However, this is not the case in our datasets: pCO$_2$ is not linearly related to the variables presented when considering the entirety of the problem. However, as noted above in subsection 2.5.1 if we consider the classes created by the SOM, they are very localized regions of the combined explicative and response data space that can be considered to approximate, locally, the mean value of a multidimensional Gaussian random distribution. We therefore assume that, if performed in the reduced neighborhood of a SOM class, the relationships between pCO$_2$ and the explicative variables are linear.

3 Application and results

3.1 Statistical imputation

As described in section 2.4 both the satellite and measured data available for the application present missing values. To complete these datasets, we chose to use imputation methods similar to those described by Schafer and Graham (2002) and Malek et al. (2008). The main idea of these methods is to use the classifying abilities of the SOMs to regroup the data in typical situations and replace the missing explicative data values with the corresponding values of the referent vector of the class to which it belongs.

We first selected the database containing SST, Chl, CDOM, NPP, MLD, time$_{sine}$, and time$_{cosine}$. The vectors were sorted according to the number of values missing from each vector and noting the locations of these missing values. We chose all complete data vectors and the first 5% of the sorted vectors containing missing data and trained a SOM. We proceeded by replacing the missing values of these first 5% of vectors with the corresponding values of the referent vector of the class to which they each belong. We then included the next 5% of vectors with missing data in a new training dataset and created a new SOM. Based on this new SOM, we again filled the new missing values with the corresponding values of the referent vector of the class of the new SOM to which they each
Figure 6. A schematic of the imputation method used. We initially sort the data depending on the amount of missing values present, then progressively train SOMs on the dataset, steadily including more vectors for the training and completing and updating the missing data during each iteration. In addition, we deleted the values we added to the first 5% of vectors and replaced them with the values of the referent vector of the class of the new SOM to which they each belong. The training parameters of this method, such as the number of classes of the topological map and the number of iterations were selected by parameter tuning. We then continued iterating this process, updating the previously filled missing values with the values of their corresponding referents belonging to the most recently trained SOM, until all missing values were filled. The updating of the previously filled values allows the method to progressively incorporate information and to rectify A schematic of the imputation method used can be seen in Figure 6.

After this imputation of the missing data through iterative training, the reconstructed data represent the original data well. Figures 7 and 8 show data for six variables before and after the reconstruction, respectively. The main difference is observed for the values of Chl, where the peak of over 200 individuals occurs at 0 mg m\(^{-3}\) because, at the initialization of the imputation process, we decided to replace all null values. The repartition of pCO\(_2\) (Figure 8a) is very representative of the data variability with a large range of values. Some very high values occur during local events, such as coastal upwelling. Most of the data range between 180 \(\mu\text{atm}\) (value observed in summer) and 550 \(\mu\text{atm}\) (observed in winter). The SST (Figure 8b) is very representative of the variability in the Baltic Sea with a maximum occurring between July and September in all basins around 18°C (Siegel and Gerth [2012]). The NPP variability is fairly homogenous, except for the peak at 10 mg C m\(^{-2}\) d\(^{-1}\). This peak occurs because the first model providing us the NPP values has a set maximum of NPP
Figure 7. Histogram of a. pCO$_2$ and satellite data, b. SST, c. Chl, d. NPP, e. CDOM, and f. MLD available for the SOM before reconstruction. Y-axis represent the number of data and the X-axis the value of the parameters.

at 10 mg C m$^{-2}$ d$^{-1}$; therefore, the correction of the NPP satellite data takes this maximum into account.

The variability of chlorophyll results in one subset of the data with a low value and another subset with a value higher than 6 mg m$^{-3}$, which can be explained by the fact that the Baltic Sea is a narrow sea, with important coastal regions, and that two blooms take place, in spring and in summer. The chlorophyll value can be very high during these periods, and the reconstruction gives a mean value for this characteristic. A peak at 10 mg m$^{-3}$ is observed in the chlorophyll data, not due to the reconstruction but to the maximum value in the satellite data file. The low MLD occurs in summer, and in the model the minimum is 10 m deep, which appears to be around the minimum value observed in Figure 8f. Absorption by CDOM decreases with increased distance from the riverine sources, reaching a relatively stable absorption background in the open sea. Most of our CDOM data capture open sea conditions, so the values are quite low.

3.2 pCO$_2$ estimation

3.2.1 Topological map

We classified the explicative variables (i.e., SST, Chl, CDOM, NPP, MLD, time$_{SST}$, and time$_{CHL}$) into classes that share similar characteristics. In order to optimize the SOM map size for the method and to calculate the method’s performances we randomly sampled 90% of our completed dataset (1300
Figure 8. Histogram of a. pCO$_2$ and satellite data, b. SST, c. Chl, d. NPP, e. CDOM, and f. MLD available for the SOM after reconstruction. Y-axis represent the number of data and the X-axis the value of the parameters.

vectors) to be used for the training phase, keeping 10% (144 vectors) to be used for computing the performances of the method. We iterated this process, selecting many different random samplings for each map size, and selected the map size with the best average reconstruction when applying the SOMLO methodology.

At the end of our optimization, we selected a SOM consisting of 77 classes. The number of observations captured by each class ranges from 0 to 38 (Figure 9). The order of magnitude of the number of observations is constant throughout the SOM, and we can regard the classes as having spread in multidimensional space in order to accurately represent the data space of the explanatory parameters. The presence of classes that did not capture any elements can be justified as preventive: they preserve the topological aspect of the SOM by preventing classes that are not similar enough from becoming neighbors.

To estimate the average concentration of pCO$_2$ in each class, the measurements of pCO$_2$ associated with vectors consisting of SST, CDOM, NPP, MLD, and CHL components were presented to the already trained SOM as input data (Figure 10). The average value computed for the vectors belonging to each class corresponds to the average value of pCO$_2$ for that class.

In the final map, the distribution of pCO$_2$ is strongly dependent on the SST distribution, with low values of pCO$_2$ correlating with high values of SST (Figure 10). This is in agreement with the seasonal pCO$_2$ cycle, which is characterized by a large amplitude, ranging from a high value in
winter ($\approx 500 \mu\text{atm}$) and a low value in summer ($\approx 150 \mu\text{atm}$), described, for example, by Wesslander (2011). According to Schneider and Kaitala (2006), the high winter value of $\text{pCO}_2$ is a consequence of mixing with a deeper water layer enriched in $\text{CO}_2$, which is in agreement with the distribution of the MLD (Figure 10), with the higher value in winter and autumn correlating with the high value of $\text{pCO}_2$. High values can also be explained by the mineralization, which exceeds production in winter (Wesslander, 2011). Biological production starts in spring when sunlight and nutrients are sufficient. The chlorophyll begins to increase in March–April due to the spring phytoplankton bloom, which reduces the $\text{pCO}_2$ level during this period. The more intensive decrease occurs in April and May, which is consistent with the higher value of NPP (Figure 10). Studies in the central Baltic Sea identify two summer minima, the first in April/May and the second in July/August, resulting from a second production period. Higher variability is observed during this period, with a standard deviation between 39 $\mu$atm and 50 $\mu$atm for different regions (Wesslander, 2011; Schneider and Kaitala, 2006).

### 3.2.2 Linear regression in the neurons

To perform an MLR, we must assume that the relationship between the predictor variables and the response variable is linear. We could take this to be a valid hypothesis only when performing the MLR in the reduced neighborhood of a SOM class, where the relationship between $\text{pCO}_2$ and the explicative variables can be assumed to be linear.
For each class $j$ a separate training dataset was created containing all the vectors assigned to that class and to all its adjacent classes. Based on that dataset, we computed the linear regression coefficient parameters for every explicative parameter and for a constant value.

The calculated linear regression coefficient parameter values for each class are shown in Figure 11. Note that all parameters are important in specific regions of the SOM, having both positive and negative correlations in different classes.

More importantly, the fact that each parameter has a significantly varying range of values over the different classes demonstrates that each parameter is important in reconstructing the $pCO_2$ in the Baltic Sea, even though a parameter may be highly significant in some classes and relatively stable in other regions of the topological map.

The addition of vectors belonging to adjacent classes did not generally perturb the estimation of the coefficient parameters because, as seen in Figure 10, the values of all parameters are generally organized coherently on the map. The assumption that they are close in the data space is not as robust as it would have been had we solely considered the vectors belonging to each class but, given the limited number of data available for modeling this highly nonlinear and complex system, we would not have sufficient elements to correctly estimate the linear regression coefficients. Given the projected increase in available data in coming years, further applications of this approach will limit themselves to the elements belonging to each class.
3.2.3 Validation of reconstruction

To validate our results, we calculated the difference and standard deviation (std) between the value of pCO$_2$ reconstructed in each neuron and the observation defining that neuron (Figure 9). On average, the std is approximately 38 $\mu$atm and the difference observed is 25–30 $\mu$atm. By cross-validating a dataset (divided by 10 sequences), we obtained the following result with a mean std of 48 $\mu$atm with a variation of 32–57 $\mu$atm and R equal to 0.9 with a variation of 0.86–0.96. Nevertheless some points indicating higher values can be identified (shown in red in Figure 9). These values are explained by the positions of these points, which are at the edges of the cloud and therefore more likely to include outliers that disturb the estimation of the MLR coefficients. For the reconstruction of pCO$_2$, with this identifiable point, it is quite easy to organize a flag system. The flag can give information about the quality of the reconstructions based on the RMS errors of the neurons used for the reconstruction. The difference obtained for pCO$_2$ in each neuron ranges from 0 to 56 $\mu$atm (Table 3), but 58% of the values observed are under 30 $\mu$atm. The difference can be quite high for a parameter such as SST, with a maximum value of 1.9 °C, but most of the values are lower than 1°C and CDOM ranges from 0 to 5.15 (Table 3). The other parameters have quite low variability, such as MLD, which ranges from 0 to 9.7 m. The average is two to three times lower than the maximum value observed, which gives low value for all the satellite parameters.

The pCO$_2$ validation dataset gives a quite good correlation (R = 0.93) with the results of the reconstruction method (Figure 12), the root mean square (RMS) being 36.7 $\mu$atm: 12% of the validation data have a value higher than 20 $\mu$atm and 45% have a value between 20 $\mu$atm and 30 $\mu$atm (Figure

**Figure 11.** Distribution of the neural map of the coefficient from linear regression for each parameter: a. SST in °C, b. and d. time cosine and sine, respectively, c. Chl in mg m$^{-3}$, f. NPP in mg C m$^{-2}$, and g. MLD in m.
Table 3. Maximum and mean value observed in the difference between the data used for the trainee and the value in the neurons.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Maximum</th>
<th>Average</th>
</tr>
</thead>
<tbody>
<tr>
<td>pCO₂ (µatm)</td>
<td>56.4</td>
<td>29.15</td>
</tr>
<tr>
<td>SST (°C)</td>
<td>1.9</td>
<td>0.98</td>
</tr>
<tr>
<td>time(cos)</td>
<td>0.33</td>
<td>0.07</td>
</tr>
<tr>
<td>Chl (mg.m⁻³)</td>
<td>0.14</td>
<td>0.06</td>
</tr>
<tr>
<td>time(sin)</td>
<td>0.4</td>
<td>0.06</td>
</tr>
<tr>
<td>CDOM</td>
<td>5.15</td>
<td>0.06</td>
</tr>
<tr>
<td>NPP (mg.m⁻³)</td>
<td>1.19</td>
<td>0.25</td>
</tr>
<tr>
<td>MLD (m)</td>
<td>9.7</td>
<td>3.2</td>
</tr>
</tbody>
</table>

Figure 12. pCO₂ reconstructed in function of the measured. The red line represents the linear relationship. The data represents 10% of the total of data used for the validation dataset.

A reconstruction has been done using the satellite data from 1998 to 2011. The seasonal cycle of pCO₂ is well reproduced and in agreement with the results of other studies. The maximum is observed in winter with a pCO₂ of 437 µatm on average, while the level is 274 µatm in summer. These values are comparable to the averages estimated in the central Baltic Sea of 500 µatm in summer and 150 µatm in winter (Wesslander, 2011). The pCO₂ decreases in April due to the biological activity and increases slowly in September (Figure 14).

We also evaluate these results by comparing them with modeling results. The model output used in the present study is from a process-oriented biogeochemical ocean model in which the Baltic Sea
Figure 13. Difference between pCO$_2$ reconstructed and measured in function of time. The red crosses and black dotted lines represent differences greater than 20 µatm. The data represents 10% of the total of data used for the validation dataset is divided into 13 natural sub-basins (e.g. Omstedt et al., 2009; Norman et al., 2013b). The properties of each sub-basin are horizontally averaged and vertically resolved, and the various sub-basins are horizontally coupled to each other using strait flow models. The model is forced by meteorological gridded data with a 3-h temporal resolution and by river runoff and net precipitation data with a monthly resolution (Omstedt et al., 2005). To compare the output of the model with our results, we couple the 13 basins in optic to reduce the number to three basins, corresponding to the three basins defined in Figure 1. The modeled and estimated pCO$_2$ are compared for the entire Baltic Sea and for the three basins from 1998 to 2009 (Figure 14). The seasonal cycle for the entire Baltic Sea is well reproduced with a quite good correlation (R = 0.7) between the modeled and estimated pCO$_2$ values (Table 4), whose standard deviations differ by 74 µatm. The modeled and estimated pCO$_2$ values for the gulfs of Finland and Bothnia are not as correlated (R = 0.6), while the order of magnitude of the variability of pCO$_2$ in the Gulf of Finland as calculated with SOMLO is closer to the model estimate (122 µatm for the modeled and 142 µatm for the estimated pCO$_2$). This lower correlation could be due to the lower number of data in this region available for these basins. The central basin is well reproduced but the amplitude of the seasonal pCO$_2$ cycle is lower in the simulation. In the southwest part of the Baltic Sea, SOMLO underestimates the pCO$_2$ concentration by 60 µatm compared with the model. In the eastern and western parts of the basin, SOMLO produces good estimates of pCO$_2$ compared with the model with an average difference of 20 µatm. In Omstedt et al. (2009), the simulated pCO$_2$ agrees quite well with the calculated values based on observations in the Eastern Gotland Basin.

A simple flag was constructed to monitor the reconstruction quality and give an idea of the confidence in the estimated pCO$_2$. The difference between the estimated and neural values is computed. The flag equals 1 for classes in which the average difference is less than 20 µatm, equals 2 for
Table 4. Coefficient of correlation (R) between the modeled pCO$_2$ and the pCO$_2$ data estimated and the std for the model and the data.

<table>
<thead>
<tr>
<th>Basin</th>
<th>R</th>
<th>STD model (µatm)</th>
<th>STD data (µatm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baltic Sea</td>
<td>0.7</td>
<td>48</td>
<td>96</td>
</tr>
<tr>
<td>Central Basin</td>
<td>0.7</td>
<td>69</td>
<td>92</td>
</tr>
<tr>
<td>Bothnian Basin</td>
<td>0.6</td>
<td>44</td>
<td>101</td>
</tr>
<tr>
<td>Gulf of Finland</td>
<td>0.6</td>
<td>122</td>
<td>144</td>
</tr>
</tbody>
</table>

an average difference of 20–30 µatm, and equals 3 for higher average differences. In the example shown here, the flag values are high (i.e., 3), so the confidence in the reconstruction is low, but some points have flag values of 1 or 2 (Figure 15d, e, and f) so the reconstruction is more reliable. On the geographic map (Figure 15d, e, and f), the values of 4 correspond to the presence of ice, which is estimated using the satellite data of the National Snow and Ice Data Center based on NOAA level 3 data [Njoku (2007)]. The flag gives confidence in our reconstruction, for example, in March 2010 (Figure 15b), the southern portion of the map (i.e., the Bornholm and Arkona basins) shows lower pCO$_2$ values than does the northern portion and than in February (not show here). In March 2010, this region corresponds to a flag value of 2, which was attributed medium confidence. In July 2010, the flag value is quite good and the variability of pCO$_2$ seems to be in line with the monthly variability (Figure 15b and e). In September 2010, the value of pCO$_2$ has a good order of magnitude when the flag is 2 but seems slightly too high when there is a poor confidence (i.e., a flag value of 3) (Figure 15c and f).

In conclusion, the reconstruction of pCO$_2$ needs to be improved to increase the confidence in the reconstruction data, particularly in the gulfs.

3.2.4 Sensitivity analysis

In order to estimate the sensitivity of the reconstruction of the pCO$_2$ to noisy data, a white noise was added on all incomplete data before the reconstruction. We performed three tests by adding a white noise to each parameter which was related to the STD of that parameter. The tested configurations were 1*sigma, 2*sigma and 0.5*sigma.

The results as seen in 5 implicate that the method is sensitive to noisy data, since when increasing the noise we progressively get worse reconstructions. It is important to note that the values obtained when using 1*sigma give an average RMS of 77, and an average R coefficient of 0.80 which indicate that the method does not degrade too much.

Another area where errors might appear is the completion of missing data. In the dataset there were vectors with missing parameters. We kept only those missing up to 3 parameters. The difference observed between the pCO$_2$ measured and reconstructed in function of month and of number of missing parameters can be seen in Figure 16. We chose to present the data this way to also give
Figure 14. Comparison between modeled pCO\textsubscript{2} (dotted lines) and pCO\textsubscript{2} estimated using the SOM linear method (solid lines) for a. the Baltic Sea (BS, blue), b. the Gulf of Finland (GF, grey), c. the Gulf of Bothnia (BG, red), and d. the Central Basin (C, green).

Table 5. Coefficient of correlation (R) between the measured pCO\textsubscript{2} and the pCO\textsubscript{2} data estimated with SOMLO and the RMS in function of the sigma apply for the noise 0.5, 1 and 2 times the sigma.

<table>
<thead>
<tr>
<th>X SIGMA</th>
<th>R</th>
<th>RMS (µatm)</th>
<th>Numbers of tests</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5</td>
<td>0.85</td>
<td>61</td>
<td>701</td>
</tr>
<tr>
<td>1</td>
<td>0.80</td>
<td>77</td>
<td>733</td>
</tr>
<tr>
<td>2</td>
<td>0.59</td>
<td>134</td>
<td>709</td>
</tr>
</tbody>
</table>

a better understanding of the seasonal variation of the reconstruction. As seen in Table 6, the imputation method does introduce errors. When reconstructing the pCO\textsubscript{2} with data that have no missing data we obtain a correlation of 0.96 and an RMS of 25.7 µatm, while reconstructing the pCO\textsubscript{2} with data missing 3 parameters, the results are of less reliable with a correlation of 0.81 and an RMS of 51.4 µatm. The RMS we obtain in the case of the reconstruction using data that contain 1 missing value is higher than the one obtained when using data containing 2 missing values, while retaining a higher correlation. This could be due to the higher number of vectors containing only one missing value.

4 Discussion and conclusions

In this paper, we used the SOMLO methodology to reconstruct the pCO\textsubscript{2} from satellite data for the Baltic Sea. SOMLO was used to accommodate the nonlinearity of the mechanics driving the pCO\textsubscript{2}. 

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Figure 15. a, b, and c: reconstruction of the pCO$_2$ map and d, e, and f: the flag for each map. a,d. March 2010, b,e. July 2010 c,f. September 2010. The flag values correspond to: 1 = high confidence, 2 = medium confidence, and 3 = low confidence.

Figure 16. Absolute difference between pCO$_2$ reconstructed and pCO$_2$ measured in function of the month. The colorbar correspond to the number of empty parameter before the reconstruction (Blue : 0 missing data, light blue: 1 missing data, Yellow: 2 missing data, Red : 3 missing data)

It uses artificial neural networks to classify data into situations, and then performs a reconstruction by using an MLR in each class. The process involves classifying the explicative parameters (i.e., SST,
Table 6. Coefficient of correlation (R) between the measured pCO$_2$ and the pCO$_2$ data estimated and the std for the model and the data in function of the number of missing parameters.

<table>
<thead>
<tr>
<th>Number of parameter missing</th>
<th>R</th>
<th>RMS (µatm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.96</td>
<td>25.7</td>
</tr>
<tr>
<td>1</td>
<td>0.93</td>
<td>39.5</td>
</tr>
<tr>
<td>2</td>
<td>0.86</td>
<td>31.9</td>
</tr>
<tr>
<td>3</td>
<td>0.81</td>
<td>51.4</td>
</tr>
</tbody>
</table>

CDOM, Chl, time, NPP, and MLD) and then using the linear regression coefficients corresponding to that class in order to reconstruct the pCO$_2$. The satellite data used was also completed using an iterative process of SOM training.

We also performed a statistical analysis of the reconstructions obtained, which allowed us to add a flag to each class, informing us of the quality of the reconstruction obtained. This could both influence further numerical modeling of other phenomena depending on the pCO$_2$ and allow for an informed interpretation of the reconstructions obtained.

The current results obtained using this method, based on 1445 vectors, gave a high correlation coefficient of 0.93% and an RMS of 36 µatm. These results are promising given the conditions under which we obtained them since, in addition to having a limited number of in situ pCO$_2$ measurements, the co-localized satellite data were frequently incomplete.

In comparison, existing studies performed over the North Atlantic and North Pacific, based on a minimum of 10,000 data points (which take into account all the data from SOCAT) to a maximum of 800,000 data points (e.g., Friedrich and Oschlies [2009] Hales et al. [2012] Landschützer et al. [2013]. Friedrich and Oschlies [2009] obtained an RMS error (RMSE) of 19 µatm. A similar study over the totality of the Atlantic Ocean obtained an RMSE of 17µatm for independent time series [Landschützer et al. [2013], Hales et al. [2012] obtained an RMSE of 20 µatm with a correlation coefficient of 0.81. The RMSE we obtained here was higher than that obtained in a previous study of the Atlantic Ocean but, taking into account the much smaller number of data available, and a stronger spatial pCO$_2$ variability might in the Baltic Sea compared to open-ocean. the results presented are promising.

The organization of the values of various MLR coefficients over each class indicated that all the satellite data parameters are important to reconstructing the pCO$_2$ in the Baltic Sea, even if only in certain cases. Improved satellite data availability could therefore also improve the performance of our reconstruction.

This study could be further developed so as to reconstruct the spatial fields of pCO$_2$. Specifically, one could imagine a Bayesian approach that would select which class to use for the MLR by also taking into account the potential classes attributed to the neighboring grid points of a geographic
study area. This, however, remains dependent on the acquisition of additional in situ measurements to allow for the robust estimation of such Bayesian probabilities.

Many programs exist for the acquisition of new data. Data from the Östergarnsholm site are still being acquired; 2012 did not yield much data from this site, and the data from 2013 and 2014 still need to be validated. In time, the SMHI station could also supply additional data. The cargo ship transect data are not yet available for 2012–2014, but these measurements will continue, and some data will soon be available. Data are also being gathered from ferries sailing the Gothenburg–Kemi–Oulu–Lübeck–Gothenburg route. This Gothenburg transect is weekly (see [http://www.hzg.de/imperia/md/content/ferryboxusergroup/presentations/fb-ws2011_karlson.pdf](http://www.hzg.de/imperia/md/content/ferryboxusergroup/presentations/fb-ws2011_karlson.pdf)). The first tests of these data were conducted in 2010 and 2011, so some data should soon be available. In addition, new measurements of pCO\textsubscript{2} began in 2012 at the Utö Atmospheric and Marine Research Station ([see http://en.ilmatieteenlaitos.fi/GHG-measurement-sites#Uto](http://en.ilmatieteenlaitos.fi/GHG-measurement-sites#Uto)).

Given the amount of new data soon to be available, we remain optimistic that comprehension and statistical modeling of pCO\textsubscript{2} in the Baltic Sea will continue to improve in coming years.

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