A neural network-based estimate of the seasonal to inter-annual variability of the Atlantic Ocean carbon sink

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Abstract. The Atlantic Ocean is one of the most important sinks for atmospheric carbon dioxide (CO2), but this sink has been shown to vary substantially in time. Here we use surface ocean CO2 observations to estimate this sink and the temporal variability from 1998 through 2007 in the Atlantic Ocean. We benefit from (i) a continuous improvement of the observations, i.e. the Surface Ocean CO2 Atlas (SOCAT) v1.5 database and (ii) a newly developed technique to interpolate the observations in space and time. In particular, we use a two-step neural network approach to reconstruct basin-wide monthly maps of the sea surface partial pressure of CO2 (pCO2) at a resolution of 1° × 1°. From those, we compute the air–sea CO2 flux maps using a standard gas exchange parameterization and high-resolution wind speeds. The neural networks fit the observed pCO2 data with a root mean square error (RMSE) of about 10 µatm and with almost no bias. A check against independent time-series data and new data from SOCAT v2 reveals a larger RMSE of 22.8 µatm for the entire Atlantic Ocean, which decreases to 16.3 µatm for data south of 40° N. We estimate a decadal mean uptake flux of −0.45 ± 0.15 Pg C yr−1 for the Atlantic between 44° S and 79° N, representing the sum of a strong uptake north of 18° N (−0.39 ± 0.10 Pg C yr−1), outgassing in the tropics (18° S–18° N, 0.11 ± 0.07 Pg C yr−1), and uptake in the subtropical/temperate South Atlantic south of 18° S (−0.16 ± 0.06 Pg C yr−1), consistent with recent studies. The strongest seasonal variability of the CO2 flux occurs in the temperature-driven subtropical North Atlantic, with uptake in winter and outgassing in summer. The seasonal cycle is antiphased in the subpolar latitudes relative to the subtrpocs largely as a result of the biologically driven winter-to-summer drawdown of CO2. Over the 10 yr analysis period (1998 through 2007), sea surface pCO2 increased faster than that of the atmosphere in large areas poleward of 40° N, while in other regions of the North Atlantic the sea surface pCO2 increased at a slower rate, resulting in a barely changing Atlantic carbon sink north of the Equator (−0.01 ± 0.02 Pg C yr−1 decade−1). Surface ocean pCO2 increased at a slower rate relative to atmospheric CO2 over most of the Atlantic south of the Equator, leading to a substantial trend toward a stronger CO2 sink for the entire South Atlantic (−0.14 ± 0.02 Pg C yr−1 decade−1). In contrast to the 10 yr trends, the Atlantic Ocean carbon sink varies relatively little on inter-annual timescales (±0.04 Pg C yr−1; 1σ).

1 Introduction

Over the last two decades, the Atlantic Ocean (44° S to 79° N and west of 19° E) has taken up about 0.49 ± 0.11 Pg C yr−1 on average (Schuster et al., 2013). This makes the Atlantic Ocean one of the most important sinks for atmospheric CO2, and especially for anthropogenic CO2 (Sabine et al., 2004; Mikaloff Fletcher et al., 2006, 2007; Gruber et al., 2009).
The Atlantic sink estimate was taken from a recent Regional Carbon Cycle Assessment and Processes (RECCAP) synthesis by Schuster et al. (2013), where the authors reviewed different methodologies to estimate the air–sea CO₂ fluxes and provided a “best” estimate. The methods included estimates derived (i) using ocean surface partial pressure of CO₂ (pCO₂) measurements (Takahashi et al., 2009), (ii) from ocean general circulation models that include a full representation of the oceanic carbon cycle (e.g. Le Quéré et al., 2007; Graven et al., 2012; Doney et al., 2009), (iii) inversions of ocean interior carbon measurements (e.g. Gruber et al., 2009) and (iv) from inversions of atmospheric CO₂ (e.g. Gurney et al., 2008). For the “best” estimate, only the observational-based estimates were used, however, i.e. those relying on pCO₂ observations and those derived from the ocean interior carbon observations through an inversion approach. These methods gave rather similar long-term mean values for the whole basin, although with substantial sub-basin differences.

The Atlantic Ocean sink varies substantially by season, which is in part driven by seasonal variations in surface ocean pCO₂ (Takahashi et al., 1993, 2002; Schuster et al., 2013), but is also affected by seasonal variations in surface ocean winds and atmospheric CO₂. Surface ocean pCO₂ varies over a wide range above and below the atmospheric pCO₂, with much of the seasonal amplitude dominated by temperature in the subtropical regions in both hemispheres (Gruber et al., 2002; Takahashi et al., 2002; Sarmiento and Gruber, 2006), explaining the summer maximum in surface ocean pCO₂. In contrast, biological processes acting in synergy with ocean mixing and circulation dominates the seasonal pCO₂ cycle in equatorial and high latitude regions (poleward of 40°) (Takahashi et al., 1993; Bennington et al., 2009), explaining the summer minimum in oceanic pCO₂. Due to their opposite phasing, these two drivers cancel each other out along the regime boundaries at around 40° (Takahashi et al., 2002), leading to a minimum in the seasonal amplitude there. Schuster et al. (2013) identified a broad agreement among independent seasonal flux estimates in the temperature driven sub-tropics, but not elsewhere.

Long-term trends and inter-annual variability of the Atlantic carbon sink represent a source of substantial disagreement between the different methodologies and studies. Using surface ocean pCO₂ observations, Schuster and Watson (2007) argued for a decrease in the North Atlantic carbon sink and a reduction in the season amplitude in both the subtropical and temperate North Atlantic from the mid-1990s to the mid-2000s. They linked this reduction to the large changes that occurred in the climate mode of the North Atlantic over this period, i.e. a shift of the North Atlantic Oscillation (NAO) from very positive phases in the early 1990s to negative and near-zero phases in the mid-2000s. Corbière et al. (2007) supported this conclusion on the basis of their observations from the subpolar gyre over the 1993 to 2003 period, pointing out that the larger than expected increase in the observed pCO₂ is mainly a result of rapid warming. Further support for a decreasing North Atlantic sink comes from Lefèvre et al. (2004), Lüger et al. (2006), Olsen et al. (2006), and Schuster et al. (2009), although each study analysed different regions and periods and also used different methods to determine trends. Schuster et al. (2009), for example, analysed data from 1990 to 2006 in the eastern subpolar gyre and throughout most of the central North Atlantic, while Olsen et al. (2006) only focused on the Nordic Seas, but looked at a more extended period, i.e. from 1981 to 2002/2003.

Based on the results of a global ocean biogeochemistry model, Thomas et al. (2008) argued that this trend toward a smaller North Atlantic sink is transitory and is expected to rebound in the near-term future, i.e. that the decreasing sink strength is the result of “natural” variability, and should not be interpreted as a signal of anthropogenic climate change. They interpreted the decline in sink strength to be the result of a NAO-driven reduction in the transport of water by the North Atlantic Current into the eastern subpolar gyre. In a contrasting modelling study, Ullman et al. (2009) argued that the North Atlantic carbon sink actually increased from the mid-1990s to the mid-2000s. They proposed that the declining trend in the NAO from the early- to mid-1990s until the mid-2000s led to reduced convective mixing in the subpolar gyre, counteracting the impact of warming.

The initial year and period of data analysed for trends are crucial aspects to consider when resolving different perspectives (Gruber, 2009). McKinley et al. (2011) pointed out that when the surface ocean trends in pCO₂ are analysed over more than 25 yr, all regions in the North Atlantic exhibit trends that are not statistically different from the trend in atmospheric CO₂, implying no change in the sink/source strength. However, when the periods of analyses were shortened to 10 yr and the beginning and ends shifted, substantial trends emerged, largely reflecting inter-annual to decadal timescale variability.

A major challenge in detecting trends in the Atlantic Ocean carbon sink is due to the highly heterogeneous distribution of the surface ocean pCO₂ observations in time and space. Different approaches have been employed to overcome this limitation and to create basin or sub-basin wide estimates. These include the binning of data to 4° × 5° bins in latitude and longitude followed by an advection-based interpolation method (Takahashi et al., 1999, 2003, 2009), binning of data to large-scale biogeochemical provinces (McKinley et al., 2011), multi-linear regression models (e.g. Chierici et al., 2009; Peng and Wanninkhof, 2010), and neural network-based methods (e.g. Lefèvre et al., 2005; Friedrich and Oschlies, 2009; Telszewski et al., 2009; Sasse et al., 2013). Each of these approaches has its strengths and weaknesses. For example, the binning and interpolation scheme employed by Takahashi et al. (1999) is well suited for constraining monthly climatologies, however, its coarse resolution tends to smooth out small-scale features. However, the method is very robust and it is not sensitive to outliers. The


binning to large-scale biogeochemical provinces works well to determine long-term trends (McKinley et al., 2011), but its resolution is even more coarse. The multi-linear regression models allow very finely resolved estimates, however, the explained variance in these statistical models is often relatively low, causing substantial uncertainties in the estimated fields.

In order to investigate the variability of the sea surface \( pCO_2 \) and the resulting carbon flux, we develop a novel neural network based approach that overcomes most of the aforementioned issues which have limited previous studies. Our method is capable of capturing a large amount of variability due to the non-linear predictor-observation relationship on a fine \( 1^\circ \times 1^\circ \) spatial grid. The method determines the non-linear relationships between the surface ocean \( pCO_2 \) observations and a set of independent observations to produce basin-wide sea surface maps of \( pCO_2 \) on a monthly basis. Our network gathers information from similar ocean biogeochemical provinces and provides us with regional \( pCO_2 \) estimates, which we then use to investigate the changing distribution of the sea surface \( pCO_2 \) in the Atlantic Ocean. Our method relies on the assumption that the Atlantic Ocean carbon sink and its variability can be estimated as a function of proxy variables, which are subjectively chosen. We further rely on ocean carbon measurements in order to establish a correct relationship. We therefore benefit from the recent publication of the Surface Ocean CO\(_2\) Atlas (SOCAT), which provides the to-date largest global data set of surface ocean fugacity of CO\(_2\) observations (Pfeil et al., 2013; Sabine et al., 2013), which we converted to \( pCO_2 \). To evaluate our estimates, we compare our results to independent data from time-series stations and additional independent \( pCO_2 \) measurements. Our basin-wide \( pCO_2 \) maps provide a basis to calculate the air–sea fluxes for the entire Atlantic and to quantify the seasonal to inter-annual variability in its sink strength.

2 Data and methods

We combine two neural network methods to reconstruct the sea surface partial pressure of CO\(_2\) (\( pCO_2 \)) for the period January 1998 to December 2007 on a monthly \( 1^\circ \times 1^\circ \) resolution. In particular, we use a two-stage approach to establish numerical relationships between \( pCO_2 \) and a suite of independent predictors that are known to drive its variability. In the first stage, we use a neural network clustering algorithm to define a discrete set of biogeochemical provinces that share a common relationship between the independent variables and climatological \( pCO_2 \). In the second stage, we derive for each biogeochemical province a non-linear and continuous relationship between \( pCO_2 \) and the input parameters on the basis of a feed-forward network (FFN) method. This input–output relationship is then used to estimate surface ocean \( pCO_2 \) for each month and each pixel. This two-stage approach has the advantage over previously employed one-stage approaches (e.g. Friedrich and Oschlies, 2009; Teltszewski et al., 2009) that the pre-clustering of the data substantially reduces the variance of the \( pCO_2 \) that needs to be explained by neural network methods, resulting in substantially better fits.

For both stages we use global data, as this permits us to take advantage of the fact that biogeochemical provinces with limited coverage in a particular ocean basin can learn from observations in the same biogeochemical province in another ocean basin. Here, we evaluate and discuss only the results for the Atlantic Ocean. The resulting surface ocean \( pCO_2 \) distribution is then combined with corresponding atmospheric \( pCO_2 \) data and wind-speed based estimates of the gas transfer velocity to construct the mean and variability of the Atlantic Ocean carbon sink from 1998 through 2007.

2.1 Data

The gridded observations of the surface ocean fugacity of CO\(_2\) \( (fCO_2) \) from SOCAT v1.5 (Sabine et al., 2013) form the basis for our computations. This data set includes global observations over the period 1970 to 2007 and was homogenized by an extensive series of automatic and manual secondary quality controls (Pfeil et al., 2013). The \( fCO_2 \) data distribution in the Atlantic Ocean is highly skewed in time and space. The number of \( 1^\circ \times 1^\circ \) pixels with \( fCO_2 \) measurements per year varies from as low as \( \sim 180 \) per year in 1999 and 2000, to over 4000 in 2006 and 2007, with the last two years accounting for 40 % of all measurements. The global data in the other ocean basins are more homogeneously distributed in time. In contrast, the Atlantic has a good spatial coverage, while this is not the case for many of the other ocean basins.

The reported \( fCO_2 \) estimates were converted to \( pCO_2 \) using the formulation (see e.g. Körtzinger, 1999)

\[
pCO_2 = fCO_2 \cdot \exp \left( \frac{P_{surf}}{R} \cdot \frac{B + 2 \cdot \delta}{T} \right)^{-1},
\]

where \( P_{surf} \) is the total atmospheric surface pressure, \( B \) and \( \delta \) are virial coefficients (Weiss, 1974), \( R \) is the gas constant and \( T \) is the absolute temperature. National Centers for Environmental Prediction (NCEP) monthly mean sea level pressure was used for \( P_{surf} \) (Kalnay et al., 1996).

A crucial choice concerns the selection of the independent input variables used for the training of the networks. We chose sea surface temperature (SST), chlorophyll \( a \) concentration (CHL), ocean mixed layer depth (MLD), sea surface salinity (SSS) and the atmospheric CO\(_2\) mole fraction \( x_{CO_2,atm} \). These parameters represent physical, chemical and biological proxies determining the distribution of \( pCO_2 \) in the sea surface layer. For SST, we use the National Oceanic and Atmospheric Administration (NOAA) Optimum Interpolation (OI) sea surface temperature v.2 (Reynolds et al., 2002), for CHL...
the SeaWiFS mapped chlorophyll (SeaWiFSProject, http://oceancolor.gsfc.nasa.gov/cgi/i3), for MLD the mixed layer depth data from the Estimating the Circulation and Climate of the Ocean, Phase II (ECCO2) project (Menemenlis et al., 2008), for SSS the Simple Ocean Data Assimilation (SODA) sea surface salinity data (Carton and Giese, 2008) and for \( \delta^{13} \text{CO}_2 \) the monthly atmospheric \( \text{CO}_2 \) from GLOBALVIEW-CO2 (2011). Furthermore, the monthly \( p\text{CO}_2 \) climatology of Takahashi et al. (2009) is used as an additional input parameter for defining the biogeochemical provinces. Due to their strongly skewed distribution, MLD and CHL were log-transformed before use as predictor values.

We restricted our analyses to the time period from January 1998 to December 2007 due to the temporal limitations of the data we chose for our study. No satellite chlorophyll data are available before the 1997 launch of the SeaWiFS mission, and the \( \text{CO}_2 \) observations in SOCAT v1.5 extend to the year 2007.

Data with an original resolution finer than the required \( 1^\circ \times 1^\circ \) were averaged onto the desired grid, whereas input data with a coarser resolution were interpolated using a bilinear interpolation. We further took monthly averages of all inputs with a finer temporal resolution.

To highlight anomalies and year-to-year trends within our data sets, we further produced deseasonalized sets of our input variables by removing their long-term mean seasonal cycle from 1990 to 2010 (1998 to 2010 in the case of chlorophyll \( a \) and 1992 to 2010 in case of MLD and SSS) from the original data set.

In the next step the monthly \( 1^\circ \times 1^\circ \) input data are rearranged into 3 major data sets. Each of these data sets consists of input vectors \( (p_n) \) where the input data are organized as row vector elements, for example SST, log(MLD), SSS, and \( p\text{CO}_2 \) Takahashi for the self-organizing map input (SINP) data set, sampled at the same space-time point (Table 1). Two of these sets, SINP and the feed-forward network input 2 set (FINP2) are global sets and do not have a corresponding target data set (Table 1). Input vectors with empty vector elements, e.g. where no salinity data are available, were removed from these data sets. The third major set, the feedforward network input set (FINP), consists only of input vectors where corresponding SOCAT v1.5 observations, or targets \( (t) \), are available, i.e. they are subsampled in time at the locations where observations are available. In order to train the feed-forward network, two sub sets of the FINP set are created, namely the actual training (FITR) set and a validation (FITVAL) set (Table 1, Appendix A2).

Where no chlorophyll \( a \) satellite data are available, due e.g. to cloud cover or lack of sufficient light at high latitudes in winter time, we estimate the sea surface \( p\text{CO}_2 \) with the remaining input parameters. This applies to about 22 \% of all pixels and mainly concerns the high latitude oceans in winter. Since chlorophyll concentrations tend to be low and unvarying during these months, we expect that this choice has a relatively small influence on our results.

We use data from two sources to evaluate and validate our results. First, we extracted time-series data from the combined record from BATS (Bermuda Atlantic Time Series Station) and Hydrostasion “S” (Bates, 2007; Gruber et al., 2002) and the European Station for Time Series in the Ocean (ES-TOC) (e.g. González-Dávila et al., 2007). Second, we use 3065 additional data points within our study region and period from the updated SOCAT v2 database (Bakker et al., 2013), which were not included in version v1.5 and therefore constitute independent data.

To evaluate the sensitivity of the results with regard to the chosen data product, we further performed 4 sensitivity runs, namely (i) SR1 (sensitivity run 1) where we replaced the SODA sea surface salinity with the World Ocean Atlas 2009 (Antonov et al., 2010) sea surface salinity climatology, (ii) SR2, where we replaced the ECCO2 MLD product with the de Boyer Montegut (de Boyer Montegut et al., 2004) MLD climatology, (iii) SR3, where we used the SODA (Carton and Giese, 2008) sea surface temperature and (iv) SR4, where we excluded chlorophyll \( a \) as an input parameter.

### 2.2 Methods

We use a self-organizing map (SOM) method (Kohonen, 1987, 2001) to partition the global ocean into 16 biogeochemical provinces. Such a biogeochemical province is characterized by each of its location having a similar relationship among all considered input variables. Based on trial and error, we finally chose the climatological \( p\text{CO}_2 \) from Takahashi et al. (2009), SST, log(MLD) and SSS as our input variables for the SOM (SINP data set). We did not include chlorophyll, i.e. log(CHL), due to its many missing values.

A SOM is a neural network based cluster algorithm that can detect regularities within the provided input data and then learns to group them together. Similar input data, arranged as input vectors, are identified via their Euclidean distance towards the nodes (or neurons) of the network. The choice of 16 provinces represents a subjectively determined optimum between too many provinces with too little data but a high degree of correlation between the provinces, and too few provinces with a lot of data, but too high variance in the data. Details on the SOM method can be found in Appendix A1.

The provinces change in shape from one month to the next and further change slightly between years, i.e. they are not static, unlike conventional provinces or biomes. As we did not provide any additional time or space information to the SOM, these temporal variations emerge solely from the temporal variability of the input data. Despite their strong seasonal dynamics in space (Fig. 1a) and time (Fig. 1b), the estimated biogeochemical provinces exhibit a coherent large-scale behaviour, reflecting the well known oceanic structures such as the gyres, the equatorial regions, and the high-latitude North Atlantic. Figure 1a shows the mode of the
Table 1. Input and target vector elements for each subset used within our method. The subscript ds describes deseasonalized data, which are computed by subtracting the long-term mean seasonal cycle from the original data set as explained in the text. Additionally, log(CHL) was excluded from sets FINP, FITR, FIVAL and FINP2 to estimate $pCO_2$ when no satellite chlorophyll $a$ was available due to cloud cover.

| Set name | Elements $n$ of the $j$th input vectors $p_j^n$ | Targets ($t_j$) |
|----------|---------------------------------|-----------------|
| SINP     | SST, log(MLD), SSS, $pCO_2$, Takahashi | –               |
| FINP, FITR, FIVAL | SST, log(CHL), log(MLD), SSS, $xCO_2$, atm, SST$_{ds}$, CHL$_{ds}$, MLD$_{ds}$, SSS$_{ds}$, $xCO_2$$_{atm,ds}$ | $pCO_2$, SOCAT  |
| FINP2    | SST, log(CHL), log(MLD), SSS, $xCO_2$, atm, SST$_{ds}$, CHL$_{ds}$, MLD$_{ds}$, SSS$_{ds}$, $xCO_2$$_{atm,ds}$ | –               |

Fig. 1. Map of the biogeochemical provinces in the Atlantic Ocean identified by the self-organizing map (SOM) method: (a) province number of the mode (i.e. most frequent occurrence), Provinces 1 and 2 do not occur in the Atlantic Ocean. (b) The number of different provinces every pixel belongs to for each month from January 1998 to December 2007.

provinces, i.e. the province each pixel mainly belongs to from 1998 to 2007 and Fig. 1b shows the number of shifting provinces per pixel. These provinces vary in time and space mainly in accordance with the variability of the climatological $pCO_2$. In the tropics, and the high latitude North Atlantic, the climatological $pCO_2$ vary little seasonally and therefore the provinces remain fairly steady, with only minimal province shifts. In contrast, the gyre regions of both hemispheres exhibit much larger seasonal variability, hence pixels there undergo many more province changes. We find the largest shifts along the Gulf Stream, where certain regions change their province association up to 10 times.

As a second step we use a feed-forward network (FFN) method to reconstruct the non-linear relationship between our input variables and the target, i.e. $pCO_2$, separately for each of the 16 biogeochemical provinces. The FFN method is a type of backpropagation network method that is capable of approximating any function with a finite number of discontinuities (Demuth et al., 2008). Similar to multi-linear regressions, a feed-forward network adjusts coefficients to establish a relationship between inputs and targets. The adjustment of the coefficients follows an iterative process. The first iteration includes an initial guess, where the coefficients are randomly initialized, the estimates are computed and compared to the target observations. From there on the network goes backwards (hence the name backpropagation) and automatically re-adjusts the coefficients with the aim to reduce the mean squared error between estimates and targets. For each iteration, only a random subset of the data is used to train the network, while the remaining data are used for validation. The updating process of the coefficients is repeated until the network estimates derived from the validation set no longer improve significantly relative to the targets. The established relationship is further used to predict the $pCO_2$ for each point in time and space where no observations are
available. This process is explained in more detail in Appendix A2.

The feed-forward network was trained with the FINP data set that included all input variables including their deseasonalized representation (see Table 1). To this end the data set was split into the 16 ocean provinces (FINP$k$, FINP$2k$, with $k = 1, \ldots, 16$) and each of them was processed separately. Due to the temporal and spatial variability of the provinces and the heterogeneous spatiotemporal distribution of the $p$CO$_2$ data, large differences exist in the number of observations within the different provinces. However, our neural network fit does not show degeneration as a function of the data density, as shown in Sect. 3.1 for the temporal distribution, and the spatial heterogeneity of the data does not lead to any major hidden bias. Details on the settings used for the FFN can be found in Appendix A2.

The air–sea flux was then calculated from the estimated $p$CO$_2$ for each grid cell and month, using the quadratic wind speed dependence of the gas transfer velocity of Wanninkhof (1992) with the gas transfer coefficient from Sweeney et al. (2007) and monthly mean wind speeds from the cross-calibrated multi-platform (CCMP) product by Atlas et al. (2011). More details are provided in Appendix A3.

### 3 Results

#### 3.1 Residuals and validation

The combined SOM-FFN method obtains very good fits with an overall mean $r^2$ between the fitted $p$CO$_2$ and the gridded Atlantic Ocean SOCAT v1.5 data of 0.87 and a RMSE of about 10 µatm (Table 2). The overall bias is small (−0.10 µatm). These results apply also to each year individually, indicating that the temporally inhomogeneous data distribution does not have a measurable effect on the estimates for each year.

The residuals are not entirely randomly distributed in space. As shown in Fig. 2a, the temporal mean residuals in each pixel show generally low values in the open ocean region, but tend to increase toward the fronts. The strongest model-observation discrepancies occur in the equatorial Atlantic, along the Gulf Stream and North Atlantic Current as well as in the Norwegian, Greenland and the North seas, i.e. mostly in regions with relatively strong horizontal gradients in surface ocean $p$CO$_2$.

The standard deviation of the residuals (Fig. 2b) shows that the strongest temporal errors occur again in the high latitudes of the North Atlantic, in particular the Norwegian and North seas, as well as along the North American coastline and in the eastern South Atlantic between 0 and 30° S. This indicates that the model input parameters are not able to predict all the temporal variability occurring in these regions with known biogeochemical complexity.

#### 3.2 Validation with independent observations

To test the impact of the inhomogeneous distribution of the neural network input data and $p$CO$_2$ observations, we show the residuals calculated as the difference between the neural network $p$CO$_2$ estimates and the gridded SOCAT v1.5 $p$CO$_2$ observations (Fig. 3). The residuals are independent of the magnitude of the estimated $p$CO$_2$, and also do not show any dependence on the magnitude of the independent variables (Fig. 3). Each bin median of the residuals is close to zero, with the strongest spread occurring in the low $p$CO$_2$ bins around 275 µatm which coincides with low SST at around 5 °C and high log(CHL) concentrations at around −0.25 to 1.25 mg m$^{-3}$. Figure 3 further shows that large residuals, most of which stem from regions characterized by strong horizontal $p$CO$_2$ gradients, are independent of the data density.

In conclusion, the residuals indicate that the combined SOM-FFN method fulfills most tests for a good fit and does not contain any major hidden biases. In particular, there is no indication of a substantial degeneration of the fits as a function of data density, neither in time nor in space. Regions with high spatial or temporal variability are the least well fitted, while the fits for most of the less variable open ocean are very good.

| Period     | $r^2$ | RMSE [µatm] | Bias [µatm] | # data  |
|------------|-------|-------------|-------------|---------|
| 1998–2007  | 0.87  | 8.99        | −0.10       | 20\,003 |
| 1998       | 0.93  | 7.15        | −0.18       | 583     |
| 1999       | 0.89  | 9.35        | 1.62        | 186     |
| 2000       | 0.78  | 11.14       | −0.19       | 178     |
| 2001       | 0.83  | 11.48       | −0.87       | 566     |
| 2002       | 0.87  | 8.98        | 0.22        | 1941    |
| 2003       | 0.87  | 7.47        | −0.11       | 1963    |
| 2004       | 0.87  | 8.13        | −0.14       | 2729    |
| 2005       | 0.88  | 9.52        | −0.43       | 3575    |
| 2006       | 0.85  | 11.51       | −0.10       | 4280    |
| 2007       | 0.87  | 10.89       | 0.12        | 4002    |

Table 2. Statistical measures of the comparison of the neural network-based estimates of $p$CO$_2$ with the SOCAT v1.5 gridded observations in the Atlantic Ocean from 44° S to 79° N and west of 30° E.
Fig. 2. (a) Temporal mean residuals and (b) standard deviation of the residuals in µatm between neural network estimates and SOCAT v1.5 gridded observations (Sabine et al., 2013) for the period from 1998 to 2007. Pixels that have a value in (a) but not in (b) indicate where only 1 observation in time is available.

near monthly coverage over the time period estimated by the model. As we do not have estimates centred at the exact geographical position of both time-series stations, we interpolate the 4 closest surrounding grid boxes, weighted by their distance, to compare our results.

Figure 4 shows the comparison between the neural network estimates with both time series for the period between 1998 through 2007 and additionally the mean seasonal cycle within this period. While the phase of the seasonal cycle is captured fairly well, Fig. 4 shows that the neural network estimates in general underestimate the winter minima at Bermuda from January to April and the autumn maxima at ESTOC from August to November. This underestimation of the seasonal amplitude is likely linked to the early stopping approach (see Appendix A2) which was implemented to prevent our neural network from overtraining. The neural network estimates further show a decrease in the summer sea surface $pCO_2$ from 2005 onwards which is not seen in the ESTOC data. The decadal mean difference between BATS data and neural network estimates is 7.56 µatm with a RMSE of 17.53 µatm. Similar to BATS, the decadal mean difference between the estimates in this study and the ESTOC data is $-8.06$ µatm with a RMSE of 16.85 µatm.

As a last test, we use data from the recently updated SOCAT v2 database (Bakker et al., 2013), which provides new independent data points within our study period to validate the results. A total of 3065 gridded observations, spread over the entire Atlantic Ocean have been added for our study region from 1998 to 2007 representing 15 % of the total amount of data used to train our network. Figure 5 shows the temporal mean and standard deviation of the residuals, similar to Fig. 2. The largest misfit between our estimates and the additional observations can again be identified along the Gulf Stream and North Atlantic Current, confirming that our method has difficulties to fully capture all variability within this region. Overall, the neural network estimates have a RMSE of 22.83 µatm and a bias of 4.85 µatm. When we exclude data north of 40° N, where we obtain the largest misfits, the results improve with a RMSE of 16.29 µatm and a mean difference of $-1.12$ µatm similar to the numbers obtained from the independent time-series stations. This suggests that over most of the ocean, our method succeeds in predicting the observed $pCO_2$ at any given time and place to within about 20 µatm, and a bias of a few µatm.

3.3 Uncertainty of the air–sea flux

The uncertainty of the flux product stems from the error in the estimated $\Delta pCO_2$ and the uncertainty of the gas transfer coefficient (Takahashi et al., 2009). We estimate this uncertainty for the integrated flux over the 4 considered RECAP/ocean inversion regions (see Table 3 for region borders) rather than for each $1^\circ \times 1^\circ$ grid cell.

The $\Delta pCO_2$ estimate is subject to two main sources of errors, i.e. the error derived from discretizing the original observations into $1^\circ \times 1^\circ$ bins and the error of the neural network method to interpolate the data in time and space. For the discretizing error we use a value of 5 µatm as reported
Fig. 3. Residuals as a function of (a) fitted $pCO_2$, (b) atmospheric $pCO_2$, (c) sea surface temperature (SST), (d) natural log of surface chlorophyll (log(CHL)), (e) natural log of mixed layer depth (log(MLD)), and (f) sea surface salinity (SSS). The upper plot in each panel depicts the residuals, shown as a box-and-whiskers plot. The red line in the box shows the median, the blue box indicates the 25 and 75 percentiles and red crosses mark residuals outside this interval. The lower plot shows the relative number of observations within each bin.

by Sabine et al. (2013), while we adopt our RMSE value of about 10 µatm (see Table 2) for the interpolation error. When computing next the error of the mean over a larger-scale region, it would be inappropriate to assume that each of the estimates is independent, as these errors are spatially correlated.
Fig. 4. Validation of neural network results with pCO$_2$ (a, b) from the BATS Hydrostation “S” (Bates, 2007; Gruber et al., 2002) and (c, d) from the ESTOC (González-Dávila et al., 2007) time-series stations. In (a) and (c) the actual time-series data are compared, while in (b) and (d), the long-term mean seasonal cycle is evaluated. Grey shading shows the uncertainty of the neural network estimate based on its RMSE. Red shading shows the standard deviation of the mean seasonal cycle for each time-series station.

To estimate the discretization error associated with gridding for each RECCAP/ocean inversion region, we use the spatial decorrelation length scale of 400 km estimated by Jones et al. (2012) to compute the effective number of degrees of freedom. The uncertainty of the mean is then estimated by dividing the standard deviation by the square root of the effective degrees of freedom. This results in an uncertainty of between 1 and 2 µatm for the individual regions.

To estimate the spatial mean of the neural network error for each RECCAP/ocean inversion region, we estimate the spatial correlation by analysing the semi-variogram of the residuals (see e.g. Kalkhan, 2011) (for details see Appendix A4). While in some regions the spatial correlation of the residuals fall very quickly, the correlation of the residuals within one bin is substantial, yielding a substantial reduction in the effective degrees of freedom. The uncertainty between different regions ranges from 1 to nearly 4 µatm.

Adding the error from the gridding and the neural network together, and assuming a mean error of 0.2 µatm for the atmospheric pCO$_2$ (Takahashi et al., 2009), yields a total ∆pCO$_2$ error for the 4 regions of between 2 and 6 µatm. With a mean gas transfer rate in the Atlantic Ocean of 0.05 mol C m$^{-2}$ yr$^{-1}$ µatm$^{-1}$ this results in a flux error between 0.03 and 0.06 Pg C yr$^{-1}$ and an overall basin error of 0.07 Pg C yr$^{-1}$ calculated by standard error propagation.

Furthermore, following Sweeney et al. (2007), we assume a random error of 30% in the gas-transfer velocity. For our long-term mean estimate of the Atlantic Ocean ($-0.45$ Pg C yr$^{-1}$) the error due to the piston velocity uncertainty is 0.13 Pg C yr$^{-1}$. This results in a total uncertainty estimate for the Atlantic Ocean of ±0.15 Pg C yr$^{-1}$, or roughly 33%, with the largest contribution stemming from the uncertain gas transfer velocity.

3.4 Decadal mean pCO$_2$ and air–sea CO$_2$ flux

The lowest decadal mean sea surface pCO$_2$ values are found in the northern North Atlantic, especially the Labrador Sea, the Greenland Sea and the Norwegian Sea with pCO$_2$ below 320 µatm and in the midlatitudes, along the Gulf stream and North Atlantic Current and the South Atlantic south of 30°S (Fig. 6a). The highest pCO$_2$ values can be identified in the equatorial Atlantic, in the North Atlantic along the Caribbean Current and the tropical and subtropical South Atlantic northwards of 30°S. Further high values are estimated at 60°N around the Irminger basin and at 30°N in the subtropical North Atlantic along the Portugal Current, and in the eastern North Atlantic along the Canaries Current.

The decadal mean pCO$_2$ distribution from the neural network method is generally very similar to that estimated by Takahashi et al. (2009), with some important exceptions
Fig. 5. (a) Temporal mean residuals and (b) standard deviation of the residuals in µatm between neural network estimates and independent data points obtained from the SOCAT v2 gridded observations (Bakker et al., 2013) for the period from 1998 to 2007. Pixels that have a value in (a) but not in (b) indicate where only 1 observation in time is available.

Fig. 6. (a) Decadal mean surface ocean $p$CO$_2$ and (b) CO$_2$ flux density in mol C m$^{-2}$ yr$^{-1}$ for the Atlantic Ocean. Negative flux densities indicate CO$_2$ uptake by the ocean.

(Fig. 7). To produce this comparison plot, we first binned our estimates to the same resolution ($4^\circ \times 5^\circ$) as the original climatology of Takahashi et al. (2009). We then corrected our estimates to the year 2000 by subtracting 4.5 µatm on the basis of the assumption that the surface ocean follows the atmospheric trend of 1.5 µatm per year and the fact that our
estimate is centred around 2003. The strongest differences can be identified in the high latitudes of the North Atlantic within the Labrador Sea, the Greenland Sea and the Norwegian Sea. For the entire Atlantic we derive a mean difference of 0.38 µatm and a RMSE of 6.45 µatm.

Given the overall small bias and the low RMSE between the two very different methods to interpolate the data, it appears that the long-term mean surface ocean pCO2 can be very robustly estimated from the available observations. The CO2 flux density (Fig. 6b) largely follows the pCO2 pattern, although with some notable differences. Overall, the North Atlantic is a strong sink for atmospheric CO2 in the mid and high latitudes, whereas the lower latitudes act as a source for atmospheric CO2. The strongest CO2 uptake in the North Atlantic occurs along the Gulf Stream and the North Atlantic Current, as well as in the Labrador, Norwegian, and Greenland seas, and in the South Atlantic south of 30° S.

We estimate a decadal mean flux of −0.44 ± 0.15 Pg C yr⁻¹ for the Atlantic Ocean from 44° S to 76° N and 100° W to 19° E (−0.45 ± 0.15 Pg C yr⁻¹ from 44° S to 79° N and west of 30° E). This is in good agreement with Schuster et al. (2013), who provided a “best” estimate of −0.49 ± 0.11 Pg C yr⁻¹ (derived from the mean fluxes of the pCO2 climatology and the ocean inversion fluxes within the RECCAP project).

Table 3 lists the long-term mean fluxes for the Atlantic Ocean as well as the four individual ocean RECCAP/ocean inversion regions considered by Schuster et al. (2013). While the basin average flux is well within the uncertainty range of the best estimate from Schuster et al. (2013), the subtropical North Atlantic (18°–49° N) mean flux is just outside the uncertainty range of the RECCAP best estimate. In general, the neural network fluxes are closest to those of the pCO2 climatology of Takahashi et al. (2009) with the exception of the subtropical South Atlantic (44°–18° S) where the long-term mean flux is closest to the results of the ocean inversion and the ocean biogeochemical models. We estimate the main carbon sink region to be the high latitude North Atlantic with strong uptake throughout the year and a decadal average uptake of −0.20 ± 0.07 Pg C yr⁻¹.

Comparing the decadal mean flux of −0.45 ± 0.15 Pg C yr⁻¹ to the results derived from the sensitivity runs SR1–4 reveals that the choice of products does not significantly influence the long-term mean result. The decadal mean fluxes from the sensitivity runs range from −0.41 ± 0.14 (SR2) up to −0.48 ± 0.16 Pg C yr⁻¹ (SR4) and are therefore well within the estimated uncertainty range.

### 3.5 Seasonality

The 10 yr mean sea surface pCO2 seasonal cycle exhibits strong latitudinal differences (Fig. 8). While we find the weakest seasonal signals north of 60° N, south of 40° S and near the Equator from 10° S to 10° N, the temperate North Atlantic (40°–60° N) has a distinct seasonal cycle with high

\[
pCO2 \text{ from October to April and low values from May to September. The opposite cycle occurs in the subtropical North Atlantic between 10°–40° N and 10–40° S with low partial pressures in winter and a seasonal maximum in the warmer summer months.}
\]

The mean seasonal cycle of our neural network-based estimates of pCO2 agrees relatively well with the seasonal cycle estimated by Takahashi et al. (2009), but substantial differences exist at the regional level (Fig. 9). As above, we corrected our estimate to the year 2000 in order to have better comparable estimates. The strongest difference can be identified in the high latitude North Atlantic, where we estimate a stronger seasonal cycle with higher values in the winter and lower values in the summer, with differences of more than 10 µatm. In comparison, the differences throughout the Atlantic Ocean are mostly within the calculated RMSE of our method.

To determine the drivers behind the seasonal cycles, we split the long-term mean seasonal cycle at each grid cell into a thermal and into a non-thermal component (Sarmiento and Gruber, 2006; Takahashi et al., 2002, Eqs. 1, 2). The latter is driven by the seasonal changes in temperature and is computed on the basis of the well known temperature sensitivity

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![Fig. 7. Difference in the surface ocean pCO2 in µatm between the decadal mean neural network estimates (this study), corrected to the year 2000, and the estimates from the climatology of Takahashi et al. (2009). Positive differences indicate higher pCO2 for the neural network estimates. The neural network estimates have been interpolated to the (4° × 5°) grid of the Takahashi climatology.](image-url)
Table 3. Comparison of regional and basin-wide decadal mean CO$_2$ fluxes in Pg C yr$^{-1}$ from the neural network-based method (column VIII) compared to a range of other methods. This includes (I) the pCO$_2$ climatology of Takahashi et al. (2009) and the tier 1 methodologies described in Schuster et al. (2013) which include (II) ocean inversion (Gruber et al., 2009), (III) atmospheric inversion (Peylin et al., 2013), (IV) ocean biogeochemical models as well as observation based results including (V) a SOCAT v1.5 based multi-parameter regression (Schuster et al., 2013) and (VI) an estimate based on the pCO$_2$ database of Takahashi et al. (2009) updated by McKinley et al. (2011). (VII) lists the best estimate from combining the different RECCAP methodologies as described in Schuster et al. (2013).

| Region | (I) pCO$_2$ climatology [Pg C yr$^{-1}$] | (II) Ocean inversion [Pg C yr$^{-1}$] | (III) Atmospheric inversion [Pg C yr$^{-1}$] | (IV) OBGC models [Pg C yr$^{-1}$] | (V) SOCAT MPR database [Pg C yr$^{-1}$] | (VI) pCO$_2$ database [Pg C yr$^{-1}$] | (VII) RECCAP best estimate [Pg C yr$^{-1}$] | (VIII) Neural network [Pg C yr$^{-1}$] |
|--------|-------------------------------------|-----------------------------------|-----------------------------------------------|--------------------------------|---------------------------------|---------------------------------|----------------------------------|---------------------------------|
| 49–76° N (West of 19° E) | −0.23 ± 0.12 | −0.19 ± 0.06 | −0.28 ± 0.03 | −0.17 ± 0.02 | −0.07 ± 0.04 | −0.13 ± 0.09 | −0.16 ± 0.09 | −0.21 ± 0.06 | −0.07 ± 0.07 |
| 18–49° N | −0.19 ± 0.09 | −0.34 ± 0.08 | −0.31 ± 0.03 | −0.13 ± 0.03 | −0.18 ± 0.09 | −0.18 ± 0.09 | −0.24 ± 0.09 | −0.26 ± 0.09 | −0.19 ± 0.07 |
| 18° S–18° N | 0.11 ± 0.05 | 0.13 ± 0.06 | 0.12 ± 0.05 | 0.15 ± 0.06 | 0.10 ± 0.05 | 0.10 ± 0.05 | 0.12 ± 0.05 | 0.12 ± 0.05 | 0.07 ± 0.07 |
| 44–18° S | −0.10 ± 0.05 | −0.17 ± 0.05 | −0.13 ± 0.02 | −0.17 ± 0.01 | −0.25 ± 0.01 | −0.25 ± 0.01 | −0.21 ± 0.01 | −0.14 ± 0.01 | −0.16 ± 0.06 |
| Atlantic Ocean | −0.42 ± 0.05 | −0.56 ± 0.06 | −0.60 ± 0.02 | −0.32 ± 0.01 | −0.40 ± 0.12 | −0.40 ± 0.12 | −0.63 ± 0.12 | −0.49 ± 0.12 | −0.44 ± 0.07 |
| | ±0.17 ± 0.13 | ±0.07 ± 0.07 | ±0.16 ± 0.07 | ±0.11 ± 0.07 | ±0.11 ± 0.07 | ±0.11 ± 0.07 | ±0.11 ± 0.07 | ±0.11 ± 0.07 | ±0.15 ± 0.07 |

Fig. 8. Hovmöller plot of the zonally averaged long-term mean seasonal cycle of (a) pCO$_2$ in µatm and (b) the CO$_2$ flux density in mol C m$^{-2}$ yr$^{-1}$.

of pCO$_2$ (Takahashi et al., 1993, Eq. 2) and the non-thermal component is computed by the difference.

The seasonal cycles of the thermally and non-thermally driven partial pressures tend to cancel each other (Fig. 10), consistent with previous analyses (Sarmiento and Gruber, 2006; Takahashi et al., 2002), i.e. assuming a 4 % change in pCO$_2$ per unit change in SST, and employing the same SST product used for the network training. In detail, we find in both hemispheres the non-thermally driven pCO$_2$ starts to decrease due to increasing biological production and reduced vertical mixing resulting in increased stratification in the warmer months. The thermally driven seasonal cycle, however, follows the increase in sea surface temperature and causes an increase in the sea surface pCO$_2$ due to a reduced solubility. Comparing Fig. 10 with Fig. 8 reveals that the non-thermal seasonal cycle of the sea surface pCO$_2$ dominates over the thermally driven seasonal cycle polewards of 40° N. In contrast, the seasonal cycle in the subtropical North and South Atlantic is driven by the thermal pCO$_2$ component. The thermal and non-thermal driven seasonal cycle of the equatorial band and in the South Atlantic south of 35° S compensate each other, resulting in little variability within each band.

The seasonal cycle of the CO$_2$ flux is largely driven by the seasonal cycle of the sea surface pCO$_2$ with only modest modifications by the seasonal cycles of wind and solubility.
shows that
mates.
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well where observations exist.
at each year individually, suggesting that trends are captured
itive values indicate a rate of increase faster than of the atmo-
itudes in winter, where the sea surface
to a larger outgassing of CO$_2$ (Fig.
2
The strongest increase in ocean surface pCO$_2$ relative to
that in the atmosphere is found in the North Atlantic pole-
ward of 40$^\circ$ N along the Gulf Stream and North Atlantic Current. Here, the neural network suggests an increase in sea
surface pCO$_2$ of more than twice the atmospheric increase.

Metzl et al. (2010) investigated the sink trend over a similar
time period (2001–2008) in the North Atlantic subpolar gyre
(53–57.5$^\circ$ N, 45–35$^\circ$ W and 57.5–62$^\circ$ N, 40–25$^\circ$ W). These
authors found a particularly strong increase in the winter sea
surface fCO$_2$ of 5.8 ± 1.1 and 7.2 ± 1.3 µatm yr$^{-1}$. While
this is much stronger than suggested here (see Fig. 11) both
studies agree on the North Atlantic subpolar gyre having a
trend towards a stronger increase of the sea surface pCO$_2$
over the ~2000 to ~2007 period. McKinley et al. (2011)
found a similar trend towards a weaker undersaturation in
their subtropical seasonally stratified region around 40$^\circ$ N for
the period 1993 until 2005, but did not identify a significant
trend in the subpolar gyre over the same period. This may
reflect differences in the time period, as their analyses with
an earlier start, i.e. pre-1990, suggest also a trend toward a
weaker undersaturation.

Decadal trends in surface ocean pCO$_2$ in the Labrador Sea
and some parts of the Icelandic Seas were much smaller than
that in the atmosphere, leading to an overall relatively small
trend for the entire region north of 40$^\circ$ N. As the low lati-
itudes of the North Atlantic (0–40$^\circ$ N) have close to zero trend
relative to that of the atmosphere, the entire North Atlantic
pCO$_2$ trend is also very close to that of the atmosphere, i.e.
1.80 ± 0.77 µatm yr$^{-1}$ versus 1.90 ± 0.34 µatm yr$^{-1}$ for the
atmosphere.

As expected from the uptake of anthropogenic CO$_2$ by the
surface ocean, the majority of the ocean pCO$_2$ trend stems
from the non-thermal part, i.e. the increase in surface ocean
DIC (dissolved inorganic carbon) (1.46 ± 1.75 µatm yr$^{-1}$).
However, the overall warming trend in the North Atlantic
over this decade further enhances the ocean pCO$_2$ trend quite
considerably (with on average stronger than atmospheric in-
creases of 2.25 ± 0.77 µatm yr$^{-1}$ from 40 to 60$^\circ$ N and east of
50$^\circ$ W and 4.03 ± 3.18 µatm yr$^{-1}$ at 60–70$^\circ$ N and 0–10$^\circ$ W).
Splitting the trend into thermal and non-thermal component
shows on average a linear trend of 1.46 ± 1.75 µatm yr$^{-1}$
for the non-thermal components, while the thermally driven
trend is on average 0.37 ± 1.47 µatm yr$^{-1}$. However, given
their uncertainty, we cannot statistically distinguish both
trends from zero. The small difference in the increase in sur-
face ocean pCO$_2$ relative to the atmosphere results in an al-
most steady strength of the Atlantic carbon sink in time north
of the Equator (−0.01 ± 0.02 Pg C yr$^{-1}$ decade$^{-1}$).

Trends for the South Atlantic show a weaker increase in
the sea surface pCO$_2$ relative to that in the atmosphere
with the exception of the eastern South Atlantic and parts
along the South American coast. On average, surface ocean
pCO$_2$ increased only by 0.98 ± 0.97 µatm yr$^{-1}$ over the 1998
through 2007 period, resulting in a carbon sink increase of
−0.14 ± 0.02 Pg C yr$^{-1}$ decade$^{-1}$. Similar to the North At-
lantic, the non-thermal component of the pCO$_2$ with an av-
ge trend of 0.76 ± 1.30 µatm yr$^{-1}$ appears to be stronger
compared to 0.19 ± 0.79 µatm yr$^{-1}$ of the thermal com-
ponent, but given their uncertainty, both trends are again in-
distinguishable from zero.

Fig. 9. Zonally averaged difference in the long-term mean seasonal cycle of the surface ocean pCO$_2$ between our neural network estimates (this study), corrected to the year 2000 and those from the climatology of Takahashi et al. (2009). Positive differences indicate higher partial pressures of CO$_2$ in the neural network-based estimates.
Fig. 10. Mean seasonal cycle of the zonally averaged sea surface $pCO_2$ driven by (a) the non-thermal component, such as changes in circulation, mixing, and biology, and (b) the thermal component, i.e. the changes in temperature. (c) and (d) show the non-thermal and thermal components respectively for the Takahashi et al. (2009) climatology. The decadal mean $pCO_2$ has been added to both components.

Taking the North and South Atlantic together, the trend over the entire study region is one toward a stronger sink over the 10 yr period with an overall mean trend of $1.46 \pm 0.76 \mu$atm yr$^{-1}$ and a flux trend of $-0.15 \pm 0.04$ Pg C yr$^{-1}$ decade$^{-1}$.

The sensitivity runs reveal that trend estimates are barely influenced by the choice of the input data product, with the exception of SR2, i.e. where we had replaced the ECCO2 MLD product with the de Boyer Montegut (de Boyer Montegut et al., 2004) MLD climatology. While $pCO_2$ trends are statistically indistinguishable between our neural network estimate ($1.46 \pm 0.76 \mu$atm yr$^{-1}$) and SR1–4 ($1.42 \pm 0.59$, $1.25 \pm 0.48$, $1.48 \pm 0.77$ and $1.37 \pm 0.73 \mu$atm yr$^{-1}$ respectively), this is not always true for the fluxes. Here, SR2 reveals a flux trend ($-0.26 \pm 0.03$ Pg C yr$^{-1}$ decade$^{-1}$) outside the uncertainty interval of our neural network estimate ($-0.15 \pm 0.04$ Pg C yr$^{-1}$ decade$^{-1}$).

It is not possible to conclude from our data whether the 10 yr trends identified here are part of truly long-term trends (30 yr or longer), or whether they are part of decadal timescale fluctuations (Thomas et al., 2008; Gruber, 2009; McKinley et al., 2011). The most recent studies by McKinley et al. (2011) and Fay and McKinley (2013) suggest the latter to be the case. The authors show that short-term trends on timescales similar to this study are strongly influenced by the chosen start and end year and strongly reflect climate mode signals such as ENSO and NAO. However, reported 50 yr trends in heat storage (Levitus et al., 2012) and interior ocean oxygen changes in the North Atlantic (Stendardo and Gruber, 2012) indicate that this region has been subject to multi-decadal changes, particularly in the subpolar gyre. It is also tempting to point out that the resulting pattern of a decreasing sink in large areas of the North Atlantic and increasing sink in the South Atlantic appears to be mirrored in the observation of a faster rate of accumulation in anthropogenic CO$_2$ in the South compared to the North Atlantic (Wanninkhof et al., 2010). One needs to be careful, though, as the surface ocean trends are for the sum of natural and anthropogenic CO$_2$, while the ocean interior trends are for anthropogenic CO$_2$ only.
Fig. 11. Linear trends (a) in sea surface $p$CO$_2$ relative to that in the atmosphere and (b) in the CO$_2$ flux over the period 1998–2007. The relative trend in sea surface $p$CO$_2$ was computed by subtracting the atmospheric mean trend. Areas with cross-hatch indicate where the trend is outside the 95% confidence level ($p \geq 0.05$). Trends were derived by first applying a 12 month running mean to each pixel to deseasonalize the data and then calculating the slope of a linear fit to these deseasonalized data.

The largest year-to-year variability in sea surface $p$CO$_2$ is found within the North Atlantic north of 40° N and in the eastern equatorial and South Atlantic. In contrast, the subtropics in both hemispheres show much less year-to-year variability.

Integrating our monthly air–sea CO$_2$ flux estimates for each year over the Atlantic Ocean reveals the largest annual mean flux differences during the second half of our study period (Fig. 12a), where annual mean fluxes range from $-0.39 \pm 0.13$ Pg C yr$^{-1}$ in 2001 up to $-0.56 \pm 0.18$ Pg C yr$^{-1}$ in 2006. Figure 12b illustrates the inter-annual variabilities (IAV) for both hemispheres and the entire Atlantic Ocean. The IAV, calculated as a 12 month running average, is fairly constant from 1998 to 2004 with a weak flux decrease in the Northern Hemisphere counterbalanced by a weak increase in the Southern Hemisphere. After 2004 the Atlantic Ocean sink increases mainly due to increases in the Southern Hemisphere. The standard deviations of the IAV (calculated as a 12 month running average and further detrended) for the Atlantic Ocean north of the Equator, south of the Equator and the entire basin are $\pm 0.02$ Pg C yr$^{-1}$, $\pm 0.02$ Pg C yr$^{-1}$ and $\pm 0.04$ Pg C yr$^{-1}$ respectively, which indicates only limited inter-annual variability in both hemispheres on a basin scale. This Atlantic Ocean low variability is further confirmed by the sensitivity runs, ranging from $\pm 0.03$ Pg C yr$^{-1}$ (SR1, SR2) to $\pm 0.03$ Pg C yr$^{-1}$ (SR3, SR4).
±0.04 Pg C yr⁻¹ (SR3, SR4), indicating that our result is not sensitive with regards to the data choice. This shows that the main findings are statistically indistinguishable from those derived without chlorophyll a (SR4), indicating the possibility to expand the analysis period back in time in future studies.

Inter-annual variability of the sea surface pCO₂ in the North Atlantic has previously been linked to variations in the NAO (e.g. Gruber et al., 2002; Schuster and Watson, 2007; Thomas et al., 2008). The NAO is the dominant large-scale climate mode in the Atlantic Ocean (e.g. Hurrell, 1995) and impacts sea surface pCO₂ via changes in the driving parameters. During positive NAO phases, sea surface temperature shows a tripole pattern with cold anomalies in the subpolar region and warm anomalies in the midlatitudes and corresponding changes in vertical mixing and nutrient supply (Marshall et al., 2001). Corbière et al. (2007) found an increase in sea surface pCO₂ in the subpolar gyre between the mid-1990s and mid-2000s linked to an increase in SST due to a shift from positive to negative NAO. We investigate the effect of the NAO by focusing on two 10° × 10° boxes, one located in the subtropical North Atlantic (20–30° N and 40–50° W) and the other in the subpolar North Atlantic (50–60° N and 30–40° W). Figure 13 illustrates the pCO₂ estimates and their anomalies for each box together with the NAO index. We find a weak but significant (p ≤ 0.05) positive correlation in the subpolar region (R = 0.32) and negative correlation in the subpolar box (R = −0.31). This pattern is consistent with that identified by Thomas et al. (2008) on the basis of a modelling study (see also summary by Gruber (2009) and recent multi-model analysis by Keller et al. (2012), and the available time-series analyses (e.g. Gruber et al., 2002; Bates, 2007) from the BATS site). The correlation patterns are derived from the neural network estimates, hence the NAO signal stems from the signal of the input data. Clearly, an important driver are the NAO-associated SST anomalies, but strongly modified by the various physical and biogeochemical changes that are driven by the NAO-induced changes in heat fluxes and wind stress (see e.g. Keller et al., 2012).

4 Summary and conclusions

We have developed a novel two-step neural network approach to diagnose monthly ocean surface pCO₂ fields from 1998 through 2007 in the Atlantic Ocean using the SOCAT v1.5 pCO₂ measurement network. Independent testing indicates our estimates are accurate to within 22.8 µatm for the entire Atlantic Ocean, which decreases to 16.2 µatm for data south of 40° N. Our study suggests a decadal mean CO₂ flux from 1998 through 2007 of −0.45 ± 0.15 Pg C yr⁻¹ for the Atlantic Ocean from 44° S to 79° N and west of 30° E. This result is in good accordance with the recent assessment from the RECCAP project (Schuster et al., 2013). We find the strongest seasonal variability in our predicted sea surface pCO₂ and air–sea fluxes within the subtropics in the Northern and Southern hemispheres, i.e. the zones where the temperature effect dominates the seasonal cycle of sea surface pCO₂. Trends in sea surface pCO₂ suggest that in large areas poleward of 40° N the rate of increase in oceanic pCO₂ was faster than in the atmosphere, leading to a regional weakening of the carbon sink strength. However, this is counter-balanced on the basin scale by weaker surface ocean pCO₂ trends elsewhere in the North Atlantic. The South Atlantic, in contrast, shows an increasing carbon sink strength throughout the study period. In total, the Atlantic Ocean carbon sink increased by −0.15 ± 0.04 Pg C yr⁻¹ decade⁻¹. The standard deviation of the inter-annual variability of the spatially integrated CO₂ flux within the study period in the Atlantic Ocean was ±0.04 Pg C yr⁻¹ with both low inter-annual variability in the Southern Hemisphere (±0.02 Pg C yr⁻¹) and in the Northern Hemisphere (±0.02 Pg C yr⁻¹).

It would be beneficial to extend the study period to further investigate responses to climate modes such as the NAO and to investigate multi-decadal variabilities. The absence of chlorophyll a permits us to prolong our analysis period and this option will be explored in subsequent work. However, chlorophyll a is a simple, but important proxy representing the relation between biology and pCO₂ and our results provide no evidence that chlorophyll can be neglected.
when considering longer timescales. Chlorophyll \( a \) is available from models before the launch of satellite observations, but the products to-date have not achieved sufficient reliability as yet.

Our product shows that the data collection and synthesis effort of the marine carbon community makes it possible to investigate the seasonal to inter-annual variability of the ocean carbon sink on a basin-scale based on observations. Future measurements are expected to increase the accuracy of such observation based estimates, and further improvements of the methods used to model the observations will result in providing better historical estimates and more accurate products for these important fluxes.

Appendix A

A1 Dividing the global ocean into biogeochemical provinces using a self-organizing map

A map with 16 neurons was chosen, organized on a 2-dimensional \( 4 \times 4 \) point hexagonal grid. This means that the input data are clustered into 16 neurons, which represent the 16 biogeochemical provinces. The term neuron refers to a processing unit, which consist of a weight vector, where each element of the weight vector corresponds to one input parameter.

In our case each weight vector consists of 4 elements (SST, log(MLD), SSS, \( pCO_2 \), Takahashi), representing its coordinates and the distance between 2 neurons is calculated via a distance function. These processing units are initially spread over a 2-dimensional field, in our case in a hexagonal formation, forming a single layer of neurons. Our experience has shown, however, that the choice of neuron topology does not have a significant effect on the final province distribution.

The use of neurons, their initialization and their distance relation describes the biggest difference towards other clustering algorithms, e.g. \( k \) means clustering. For our study, the Euclidean distance between a neurons weight vector and the input vectors of the SINP data set was used for the distance function. The weight matrix \( (W_{mn=16,n=4}) \), which is formed by the 16 neurons with their 4 vector elements, was randomly initialized.

After initialization, the training vectors are introduced to the SOM with each training parameter as 1 element of the vector. For the \( j \)th input vector \( p_{i}^{j} \) with the length \( n \) the Euclidean distances to each of the \( i = 1, \ldots, 16 \) neurons represented by each row \( i \) of the weight matrix are calculated:

\[
d_{i}^{j} = \sqrt{\sum_{l=1}^{n} (W_{l,i} - p_{l}^{j})^2},
\]

where \( d_{i}^{j} \) comprises a vector containing the Euclidean distances to each neuron \( i \) of the input vector \( p_{i}^{j} \). The smallest element of the distance vector, i.e. the shortest distance element, marks the distance towards the closest neuron, called the winning neuron. The neuron \( i \), gets updated by moving towards the average position of all the training vectors \( j \) it was identified as a winner, or a close neighbour of the winner. This sort of training is called “batch training”. This is done by adjusting the \( i \)th row of the initial weights matrix after the iterative step \( q \) following Vesanto et al. (2000):

\[
W_{i,n}(q + 1) = \frac{\sum_{j=1}^{q} S(d_{\text{neighbour}} - d_{i}^{j}) \cdot p_{n}^{j}}{\sum_{j=1}^{q} S(d_{\text{neighbour}} - d_{i}^{j})},
\]

where \( W_{(i,n)} \) is the \( i \)th row of the input weight matrix, \( d_{i}^{j} \) is the Euclidean distance between the neuron \( i \) and the presented \( j \)th input vector and \( d_{\text{neighbour}} \) is the neighbourhood radius. \( S \) describes the step function. Neighbouring neurons will only be updated if \( S(d_{\text{neighbour}} - d_{i}^{j}) > 0 \). During the training of our SOM we decrease \( d_{\text{neighbour}} \) in 2 steps from an initial coarse training phase where we update the surrounding nearest neighbours that lie within a neighbourhood radius of 3, which lasts for the first 100 iterations, to a final fine training phase where only the winning neuron is updated.

After \( q = 200 \) iterations (presenting all our input vectors to the SOM 200 times) we stop the training and the input vectors are re-introduced, without updating the SOM. In return every vector receives the neuron number \( i \) of the winning neuron, where \( d_{i}^{j} = \min(d_{m}^{j}) \), until every training vector is labelled with a number between 1 and 16, representing the province the vector belongs to. Since every training vector has a geographical location we can now divide the global ocean into these 16 provinces, i.e. the 16 biogeochemical provinces.

We forced the relative weights of the input data toward the climatological \( pCO_2 \) data, in order to minimize the variance of \( pCO_2 \) within each biogeochemical province. To do so, we did not normalize our input data, with the exception of MLD, which we log-transformed (Table 1). As a consequence, the range between the lowest and highest value of \( pCO_2 \) is one order of magnitude larger than that for SST, and about another order of magnitude larger than that for the remaining input parameters (log(MLD), SSS). This was done to reduce the biases in the second stage of the fitting, i.e. in the feed-forward network. As a consequence, the biogeochemical provinces follow the seasonal pattern of the \( pCO_2 \) climatology, meaning that the seasonality of \( pCO_2 \) at any given location will be mostly determined by the seasonal changes of the biogeochemical provinces and to a lesser degree by the seasonal cycle of the input data in the second state. In addition, owing to the climatological nature of the used \( pCO_2 \) data, there are little inter-annual shifts in the distribution of the biogeochemical provinces.
A2 Reconstructing the sea surface $pCO_2$ using a
feed-forward network

Our feed-forward network uses 2 layers of neurons, 1 hidden
layer of neurons using a sigmoid transfer function and 1 linear output layer. The hidden layer response to the input
vector $p^k_{n}$ can be written as follows:

$$a_i = \frac{2}{1 + \exp(-2 \cdot (W_{i,n} \cdot p^k_{n} + b_i))} - 1,$$  \hspace{1cm} (A3)

where $a_i$ is the hidden layer response of $i$th neuron. The response of all neurons $m$ then forms the output vector of the
hidden layer $a_m$. This vector serves as input for the second
layer of neurons, the linear output layer.

$$pCO^j_{2, est} = W_m \cdot a_m + b$$ \hspace{1cm} (A4)

Eqs. (A3) and (A4) show how the network calculates the
scalar output $pCO^j_{2, est}$ for the $j$th input vector $p^k_{n}$ in 2 steps,
each step referring to 1 layer of the network. In the hidden
layer the input vectors are multiplied with the weight matrix
of the hidden layer $W_{m,n}$ and added to the layer bias vector $b_m$. The output vector of the hidden layer $a_m$ is created using
a tangent sigmoid transfer function (Eq. (A3)), that computes
elements of $a_m$ in the range from $-1$ to $+1$. Similar to the
SOM, the size of $W_{m,n}$ is determined by the size $n$ of the
input vector and the number $m$ of neurons in the hidden layer.

The length of $b_m$ and $a_m$ is as well determined by the number
of neurons $m$. In the linear output layer $a_m$ is processed
the same way as $p^k_{n}$ in the hidden layer, with the exception
that the output layer only consists of 1 neuron to produce one
scalar $pCO_2$ estimate ($pCO^j_{2, est}$) for every input vector. Fur
thermore, the linear output layer allows $pCO^j_{2, est}$ to have any
value between $-infinity$ and $+infinity$. During the training the
weights and biases of each layer get iteratively adjusted to
minimize the error between the network output $pCO^j_{2, est}$ and
the scalar target element $t^j$ from the SOCAT v1.5 database
that corresponds to $p^k_{n}$. Therefore, the network can only be
trained by those input vectors which do have co-located ob
servations.

Before the training starts, the training vectors with corre
sponding observations from our FINP$^k$ data set are provided
to initialize the network layer size. The size of the 2 network
layers is determined by the number of neurons and the number
of input vector elements $n$. We randomly generate the
networks initial weights and biases. An important parameter
that has to be provided before training starts is the number
of neurons. Too few neurons are not able to reproduce
realistic results, whereas too many neurons decrease the
computational performance and cause over-fitting and therefore
the network is not able to generalize (Demuth et al., 2008).

Since the number of inputs and targets varies per province,
we cannot provide one best number of neurons to use for all
16 provinces. We therefore perform a pre-training, increasing
the number of neurons in the first or hidden layer parabolically
starting from 2 neurons up to a number where the ratio
between of the training sample size to the number of weights
does exceed 30. Amari et al. (1997) proposed this ratio to
prevent artificial neural networks from over-fitting.

During every pre-training process the FINP$^k$ set is divided
into 2 separate sub-sets. The first (FITR$^k$) is used to train
the network and the second (FIVAL$^k$) is used for validation
within the method. Amari et al. (1997) suggested an optimal
split ($r_{opt}$) between training and validation data as a function
of the modifiable parameters $m$:

$$r_{opt} = \frac{1}{\sqrt{2 \cdot m}}.$$ \hspace{1cm} (A5)

Modifiable parameter refers to the weights and biases of
the network. During every pre-training process the FITR$^k$
training vectors and the corresponding FITR$^k$ targets are in
trduced to the network and the weights and biases are iter
atively updated in the direction where the performance func
tion, which is the mean squared error between network out
puts $pCO^j_{2, est}$ and FITR$^k$ targets $t^j$, decreases most rapidly.
Our feed-forward network uses the Levenberg–Marquardt
(Marquardt, 1963) algorithm to update weights and biases
in every iteration step to reduce the mean squared error be
tween outputs and targets. The application of the algorithm
in neural networks is described in more detail in Hagan and
Menhaj (1994); Hagan et al. (1996).

After every iteration of each pre-training the network is
validated by using the FIVAL$^k$ sub-set. The updated weights
and biases are used to simulate outputs from the FIVAL$^k$
inputs and the mean squared error between these outputs and
the FIVAL$^k$ targets is calculated. Every pre-training of the
network stops automatically when 6 consecutive iterations
do not reduce the network’s error on the FIVAL$^k$ targets to
prevent the network from over-fitting. After the pre-trainings
with increasing number of neurons we select the one where
the mean squared error of the validation data set FIVAL is a
minimum and receive the optimal number of neurons for the
actual training process.

During the actual training process the number of neurons is
adjusted according to the best pre-training performance for
each of the 16 provinces separately. We perform 10 train
ings where we randomly pick validation data according to
Eq. (A5) out of the entire pool of observations to avoid over
fitting of the network output. After every training we use the
trained network to simulate $pCO^j_{2, est}$ from the FINP$^k$
data set and average the output of the 10 training cycles, to end up
with 1 estimate for our time period from 1998 through 2007
for each province. After 16 FFN runs we can now combine
our results of the 16 provinces to retrieve our $pCO_2$ estimates
from 1998 to 2007 on a global $1^\circ \times 1^\circ$ grid.
Fig. A1. Empirical semi-variograms of randomly chosen ensembles of the residuals as a function of distance in (a) the subpolar North Atlantic with a median decorrelation length of 430 km, (b) subtropical North Atlantic with a median decorrelation length of 166 km, (c) the equatorial Atlantic with a median decorrelation length of 530 km and (d) the South Atlantic with a median decorrelation length of 9 km. Region borders are listed in Table 3.

A3 Air–sea CO$_2$ flux calculation

We calculate the air–sea flux density in mol C m$^{-2}$ yr$^{-1}$ for each month and $1^\circ \times 1^\circ$ pixel from

$$ F_{CO_2} = -k_w \cdot S_{CO_2}(1 - f_{ice}) \cdot (p_{CO_2,\text{atm,wet}} - p_{CO_2}), \quad (A6) $$

where $S_{CO_2}$ is the mainly temperature driven solubility of CO$_2$ (calculated in mol C m$^{-3}$ µatm$^{-1}$) and $k_w$ is the gas transfer velocity (calculated in m yr$^{-1}$) and where the flux is defined positive upward, i.e. outgassing is positive, and uptake is negative. $f_{ice}$ refers to the percent of ice cover within a region derived from Rayner et al. (2003). For the gas transfer velocity (here calculated in cm h$^{-1}$) we decided to use the formulation of Wanninkhof (1992) with the scaling factor of Sweeney et al. (2007), i.e.

$$ k_w = 0.27 \cdot (Sc/660)^{-\frac{1}{2}} \cdot u^2, \quad (A7) $$

where $Sc$ the dimensionless Schmidt number and $u$ the monthly mean CCMP wind speed (Atlas et al., 2011) at a height of 10 m above the sea surface.

The solubility of CO$_2$ is calculated according to Weiss (1974) and the Schmidt number according to Wanninkhof (1992) using the same SST and SSS data we used for the training of our network.

The partial pressure of atmospheric CO$_2$, i.e. $p_{CO_2,\text{atm,wet}}$, was computed from the dry air mixing ratio $x_{CO_2}$ of GLOBALVIEW-CO2 (2011), taking into account the water vapour correction according to Dickson et al. (2007):

$$ p_{CO_2,\text{atm,wet}} = x_{CO_2,\text{atm}} \cdot [P_{\text{atm,surf}} - P_{H_2O}], \quad (A8) $$

where $P_{\text{atm, surf}}$ is the sea-level pressure from NCEP (Kalnay et al., 1996), and $P_{H_2O}$ describes the water vapour pressure.
A4  Accounting for spatial autocorrelation in flux uncertainty analysis

For each RECCAP/ocean inversion region (see Table 3 for region borders), we first divide the residuals into 5 randomly chosen mutually exclusive ensembles, with the exception of the subtropical North Atlantic, where we use 10 ensembles, due to the larger amount of data. For each ensemble, we compute the semi-variance of the residuals and their point-to-point Haversine distance matrix, and then fit an exponential function of the form

\[ a + b \cdot \exp \left( -\frac{x}{c} \right) \]  

(A9)

to the semi-varioigram in order to estimate the correlation length (parameter \( c \)) between the residuals. We find that the semi-varioigrams are very sensitive to extreme values of the residuals, forcing us to use Chauvenet’s criterion to reject them prior to the computation and the fit. By using several different ensembles per region, we will account for the potential biasing effect of their removal.

Figure A1 shows the semi-variograms of all ensembles in the Atlantic Ocean. Correlation lengths of the residuals vary between 9 km, where the ensembles are well below the distance between 2 neighbouring grid boxes, and 532 km. However, in all cases the semi-variogram shows a large lag 0 correlation, (semi-variance at 0 distance varies between 20 and 60 \( \mu \text{atm}^2 \) within the different ensembles in the different regions) indicating the residuals within one grid cell are correlated with each other, leading to a substantial reduction of the degrees of freedom.

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