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Increase of dissolved inorganic carbon and decrease in pH in near-surface waters in the Mediterranean Sea during the past two decades

Liliane Merlivat 1 , Jacqueline Boutin 1 , David Antoine 2,3 , Laurence Beaumont 4 , Melek Golbol 3 , and Vincenzo Vellucci 3

¹Sorbonne Université-CNRS-IRD-MNHN, LOCEAN, 75005 Paris, France

²Remote Sensing and Satellite Research Group, School of Earth and Planetary Sciences, Curtin University, Perth, Australia ³Sorbonne Université-CNRS, Laboratoire d'Océanographie de Villefranche, LOV, 06230 Villefranche-sur-Mer, France ⁴Division Technique INSU-CNRS, 92195 Meudon CEDEX, France

Correspondence: Liliane Merlivat (merlivat@locean.upmc.fr)

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Abstract. Two 3-year time series of hourly measurements of the fugacity of CO_2 (fCO_2) in the upper 10 m of the surface layer of the northwestern Mediterranean Sea have been recorded by CARIOCA sensors almost two decades apart, in 1995–1997 and 2013–2015. By combining them with the alkalinity derived from measured temperature and salinity, we calculate changes in pH and dissolved inorganic carbon (DIC). DIC increased in surface seawater by \sim 25 µmol kg⁻¹ and $fCO₂$ by 40 µatm, whereas seawater pH decreased by \sim 0.04 (0.0022 yr⁻¹). The DIC increase is about 15 % larger than expected from the equilibrium with atmospheric $CO₂$. This could result from natural variability, e.g. the increase between the two periods in the frequency and intensity of winter convection events. Likewise, it could be the signature of the contribution of the Atlantic Ocean as a source of anthropogenic carbon to the Mediterranean Sea through the Strait of Gibraltar. We then estimate that the part of DIC accumulated over the last 18 years represents \sim 30 % of the total inventory of anthropogenic carbon in the Mediterranean Sea.

1 Introduction

The concentration of atmospheric carbon dioxide $(CO₂)$ has been increasing rapidly over the 20th century. As a result, the concentration of dissolved inorganic carbon (DIC) in the near surface ocean increases, which drives a decrease in pH in order to maintain a chemical equilibrium. These changes have complex direct and indirect impacts on marine organisms and ecosystems (Gattuso and Hansson, 2011). Empirical methods to estimate the anthropogenic $CO₂$ penetration in the ocean since the industrial revolution have improved over the past few decades (Chen and Millero, 1979; Gruber et al., 1996; Sabine et al., 2008; Touratier and Goyet, 2004, 2009; Woosley et al., 2016). As the concentration of anthropogenic carbon, Cant, cannot be distinguished from the natural background of DIC through total DIC measurements, these methods are based on the analysis of different chemical properties of the water column. Direct estimates of the anthropogenic $CO₂$ absorption in the sea surface layers are difficult due to the large natural variability driven by physical and biological phenomena. Bates et al. (2014) have extracted the trend from the large variability based on the analysis of a long time series (monthly or seasonal sampling). For the global surface ocean, Lauvset et al. (2015) have used the Surface Ocean $CO₂$ Atlas (SOCAT) database (Bakker et al., 2014) combined with an interpolation method. Estimates of anthropogenic storage in the Mediterranean Sea differ by about a factor of 2 (Huertas et al., 2009; Touratier and Goyet, 2009). In addition to the anthropogenic signal, oceanic DIC can also be the signature of a strong inter-annual variability. In the North Atlantic, for instance, McKinley et al. (2011) have shown that the long-term trend emerges only after more than 25 years because of natural variability.

A high-frequency sampling of the seawater carbon chemistry at the air-water interface over extended periods of time is useful in assessing the trends and variability of DIC. In

Figure 1. The area of the northwestern Mediterranean Sea showing the southern coast of France, the Island of Corsica, the main current branches (gray arrows), the location of the DYFAMED site (43[°]25′ N, 7[°]52′ E, red star) [\(http://doi.org/10.17882/43749,](http://doi.org/10.17882/43749) last access: 14 September 2018) and the BOUSSOLE buoy $(43°22'N,$ 7°54' E, black star) in the Ligurian Sea.

this paper we analyse two 3-year time series of hourly fugacity of $CO₂$, ($fCO₂$), measured with autonomous CARIOCA sensors (Copin-Montégut et al., 2004; Merlivat and Brault, 1995) in 1995–1997 and 2013–2015 at two nearby locations in the northwestern Mediterranean Sea (Fig. 1). Using measured fCO_2 , temperature (T) and salinity (S) , we derive the other variables of the carbonate system (pH and DIC). The experimental setting is first described, and the recent data obtained over the 2013–2015 period are presented. Combined with the 1995–1997 measurements previously published (Hood and Merlivat, 2001), we estimate the decrease in pH and the increase in DIC. The results are discussed with respect to the contributions of the exchange with atmospheric $CO₂$, the possible impact of vertical mixing and recent estimates of the transport of anthropogenic carbon from the Atlantic Ocean over a 18-year period.

2 Material and methods

2.1 The BOUSSOLE and DYFAMED sites

Data collection was carried out at the BOUSSOLE site $(43°22' N, 7°54' E)$ in 2013–2015 (Antoine et al., 2006, 2008) and at the DYFAMED site $(43°25' N, 7°52' E)$ in 1995–1997 (Marty et al., 2002). These sites are 3 nautical miles apart, both located in the Ligurian Sea, one of the basins of the northwestern Mediterranean Sea (Fig. 1). The water depth is \sim 2400 m. The prevailing ocean currents are usually weak $(< 20 \,\mathrm{cm\,s^{-1}})$, because these sites are in the central area of the cyclonic circulation that characterises the Ligurian Sea. The two sites surrounded by the permanent geostrophic Ligurian frontal jet flow are protected from coastal inputs (Antoine et al., 2008; Heimbürger et al., 2013; Millot, 1999). Monthly cruises are carried out at the same location.

2.2 Analytical methods

At DYFAMED, $fCO₂$ measurements at 2 m were provided by an anchored floating buoy fitted with a CARIOCA sensor. At BOUSSOLE, measurements were carried out from a mooring normally dedicated to radiometry and optical measurements where two CARIOCA sensors were attached. Both monitored $fCO₂$ hourly at the 3 and 10 m depths (although only one of the two depths was equipped with a functional sensor at some periods); S and T were monitored at the same two depths using a Seabird SBE 37-SM MicroCat instrument. The CARIOCA sensors were adapted to work under pressure in the water column. They were swapped about every 6 months, with serviced and calibrated instruments replacing those which were previously deployed. The accuracy of CARIOCA $fCO₂$ measurements by the spectrophotometric method based on the optical absorbance of a solution thymol blue diluted in seawater is estimated at 2 µatm during both periods. Hood and Merlivat (2001) have reported agreement between the $fCO₂$ measured by CARIOCA buoys, similar to the one deployed at DYFAMED, with ship-based measurements during a number of field programs and with an accuracy of 2 µatm and a precision of 5 µatm.

At Boussole, newly designed $fCO₂$ sensors have been calibrated using in situ seawater samples taken at 5 and 10 m depths during the monthly servicing cruises to the mooring. The total alkalinity (Alk) and DIC of the samples were determined by potentiometric titration using a closed cell according to the method developed by (Edmond, 1970). Certified reference materials (CRMs) supplied by A. G. Dickson (Scripps Institution of Oceanography, San Diego, USA) were used for calibration (Dickson et al., 2007). The accuracy is estimated at 3μ mol kg⁻¹ for both DIC and Alk. fCO_2 is calculated using the dissociation constants of Mehrbach refitted by Dickson and Millero (Dickson and Millero, 1987; Mehrbach et al., 1973) and as recommended by Álvarez et al. (2014) for the Mediterranean Sea. Uncertainty in $fCO₂$ derived from an individual sample is expected to be on the order of 5 µatm (Millero, 2007). About eight samples have been used to calibrate each CARIOCA sensor so that the uncertainty of the absolute calibration of each $fCO₂$ CARIOCA sensor is estimated at 1.8 µatm. In addition, we observe that the standard deviation of the difference between the CARI-OCA $fCO₂$ and $fCO₂$ computed with the monthly discrete samples (Fig. 2b) is equal to 4.4 µatm, consistent with the expected precision on CARIOCA $fCO₂$ of 5 µatm. Alk and S of the 56 samples taken at BOUSSOLE are linearly corre-

Figure 2. Inter-annual variability of CARIOCA data on the BOUSSOLE mooring; left column represents a function of time and right column a function of months for a given year (blue for 2013, green for 2014 and red for 2015). (a, d) T, (b, e) fCO_2 and (c, f) $fCO_2 @ 13 °C$. In (a–c), the dotted lines indicate the period affected by stratification and internal waves (26 July to 1 October 2014 and 8 July to 1 October 2015). In (b), the open circles correspond to $fCO₂$ data derived from DIC and alkalinity measurements of samples taken at 5 and 10 m. In (e), the thin lines indicate fCO_{2atm} . Note that the colour code on (d–f) is different from (a–c).

lated according the following relationship;

Alk (μ mol kg⁻¹) = 87.647*S* − 785.5. (1)

The standard deviation of the Alk data around the regression line is equal to $4.4 \,\text{\mu}$ mol kg⁻¹ ($r^2 = 0.89$).

3 Results

3.1 The BOUSSOLE mooring (2013–2015) time series

Temperature and $fCO₂$ were measured from February 2013 to February 2016. All seasons were well represented, with missing data only in May–July 2013. For some periods, simultaneous measurements were made at the 3 and 10 m depths (Fig. 2a–c).

The range of temperature (Fig. 2a) extends from 13° C in winter up to 27 °C in summer, followed by progressive cooling in fall. The coldest temperature, 13 ◦C, results from the

winter vertical mixing with the deeper Levantine Intermediate Water (LIW) marked by extremes in temperature and salinity (Copin-Montégut and Bégovic, 2002). Temperature provides the main control of the seasonality of $fCO₂$, from 350 µatm to more than 550 µatm in summer 2013 (Fig. 2b). The fugacity of $CO₂$ in seawater is a function of temperature, DIC, alkalinity, salinity and dissolved nutrients. In the oligotrophic surface waters of the Mediterranean Sea, the effect of nutrients may be neglected. Temperature and DIC have the strongest influences. By normalising $fCO₂$ to a constant temperature, the temperature effect can be removed and changes in $fCO₂$ resulting from changes in DIC can be more easily identified. Figure 2c shows the variability of $fCO₂$ normalised to the constant temperature of 13 °C, $(fCO₂@13)$ using the equation of Takahashi et al. (1993). The underlying processes that govern the seasonal variability of $fCO₂@13$ are successive winter mixing, biological activity (organic matter formation and remineralisation) and

the deepening of mixed layer in fall (Bégovic and Copin-Montegut, 2002; Hood and Merlivat, 2001). Biological processes account for the decline in $fCO₂@13$ observed from March–April to late summer; the ensuing increase in surface $fCO₂@13$ is associated with the deepening of the mixed layer in the fall or convection in winter, as the vertical distribution of $fCO_2@13$ at DYFAMED shows a maximum in the 50–150 m layer. Where a large remineralisation of organic matter occurs, the productive layer is mostly between 0 and 40 m (Copin-Montégut and Bégovic, 2002). The contribution of air–sea exchange is not significant (Bégovic and Copin-Montegut, 2002). Over the period 2013–2015, the air– sea $CO₂$ flux from the atmosphere to the ocean surface is equal to -0.45 mol m⁻² yr⁻¹.

During summer 2014, large differences between measurements at 3 and 10 m were observed (Fig. 2a–c between dashed lines). A detailed analysis of the temporal variability during that period underscores the role of inertial waves at the frequency of 17.4 h that create the observed differences between the 2 depths of observations, with the deeper waters being colder and enriched in $fCO₂@13$. Temperature and $fCO₂@13$ variability is dominated by inertial waves. In particular, from 15 to 26 August 2014, the difference in T between the two depths is as large as 7.6 (5.1 \degree C on average). $fCO₂$ decreases on average by 32.7 µatm, corresponding to an increase in $fCO₂@13$ equal to 42.8 µatm.

The 2013–2015 seasonal and inter-annual variability of T , $fCO₂$ and $fCO₂@13$ is illustrated in Fig. 2–f. The larger inter-annual changes in temperature (Fig. 2d) are observed during summer, both at 3 and 10 m depth, while over February and March, a constant value of $13\degree C$ is observed as the result of vertical mixing with the LIW. A very large interannual variability of $fCO_2@13$ is observed for $T < 14 °C$ (Fig. 2f). This is associated with the winter mixing at the mooring site, which is highly variable from year to year. Winter mixed-layer depth (MLD) varies between 50 and 160 m, at the top of the LIW over the 2013–2015 period (Coppola et al., 2016). The variable depth of the winter vertical mixing causes the difference in $fCO₂@13$, as $fCO₂$ increases with depth (Copin-Montégut and Bégovic, 2002). The deepening of the MLD is driven by episodic and intense mixing processes characterised by a succession of events lasting several days, which are related to atmospheric forcing (Antoine et al., 2008) and lead to increase in $fCO₂@13$. Figure 2e illustrates the solubility control of the variability of $fCO₂$, as $fCO₂$ increases when T increases. Another cause of the inter-annual variability of fCO_2 for $T \sim 14\degree C$ is the timing of the spring increase in biological activity, which differs by a month between years. For instance, the increase happened at the beginning of April in 2013, when $T \sim 15-16$ °C and by mid March in 2014, when $T \sim 14$ °C. Another cause is the deepening of the mixed layer due to the fall cooling, which varies by a month between years.

3.2 Decadal changes in hydrography

3.2.1 Sea surface temperature changes

Monthly mean values of temperature have been computed for the two 3-year periods, 1995–1997 and 2013–2015. In 1995– 1997, $fCO₂$ and T at 2 m were measured with CARIOCA sensors installed on a buoy at DYFAMED (Hood and Merlivat, 2001). The mean annual temperature of hourly CAR-IOCA data is equal to 18.21 ◦C. For 2013–2015, temperature measurements made on the BOUSSOLE mooring at 3 and 10 m have been used. For the April to September time interval, there are only data at the 3 m depth. In addition, temperature data measured half hourly at 0.7 m at a nearby meteorological buoy (43°23' N, 7°50' E) [\(https://doi.org/10.](https://doi.org/10.6096/hymex.azurbuoy.thermosalinograph.20100308) [6096/hymex.azurbuoy.thermosalinograph.20100308,](https://doi.org/10.6096/hymex.azurbuoy.thermosalinograph.20100308) last access: 19 September 2018; Rolland and Bouin, 2010) have been used (Fig. 3d). Mean annual temperatures are equal to 18.29 and 17.97 ◦C, respectively, based on the meteorological buoy and the BOUSSOLE mooring data. The two sets of data differ essentially during July and August, with the temperatures at 3 m colder than those at 0.7 m, which indicates a thermal gradient between the two depths during summer. Therefore, for 2013–2015, we select the mean annual value computed with the meteorological buoy, $18.29\degree C$, as a better representation of the sea surface. This value is close to 18.21 ◦C computed for 1995–1997. Thus, no significant change in SST is found between the 2 decades, with a mean value equal to $18.25 \degree C$.

3.2.2 Sea surface salinity changes

The mean value of salinity and the standard error of the mean computed from 56 samples taken at BOUSSOLE in 2013– 2015 are respectively 38.19 and 0.02. In 1998–1999, ship measurements of surface salinity were made during monthly cruises at the DYFAMED site (Copin-Montégut et al., 2004). The mean salinity and the standard error of the mean of this set of 19 data are respectively 38.21 and 0.03. Thus, there is no significant salinity change between the two decades.

3.3 Decadal changes in $fCO₂@13$

3.3.1 Time series of $fCO₂@13$ in 1995–1997 and 2013–2015

The two time series of high-frequency data were analysed in order to quantify the change in $fCO₂@13$ 2 decades apart at the sea surface. To account for the inter-annual seasonal variability as well as irregular sampling, we performed an analysis of the change in $fCO₂@13$ as a function of SST (Fig. 3a and b). For the 2013–2015 data set, we excluded summer data measured at 10 m depth as they were not representative of the surface mixed layer due to a strong stratification. Much larger $fCO_2@13$ values are observed at low temperatures than at high temperatures, the decrease being

Figure 3. (a) $fCO_2@13$ as a function of temperature for hourly data in 2013, 2014 and 2015. The yellow dots indicate mean $fCO_2@13$. (b) is similar to (a) but represents all hourly data in 1995–1997 (black) and in 2013–2015 (red). (c) is similar to (b) but represents average values per 1 °C interval (standard deviation as dotted line). The difference between the two periods is also displayed by the dashed blue curve (scale on the right axis; the mean difference over all SST is represented by the horizontal blue line). (d) Mean monthly sea surface temperature for 1993–1995 (black curve for CARIOCA sensors), 2013–2015 (green for CARIOCA sensors) and 2013–2015 (red for meteorological buoy). Corresponding mean annual values are indicated by dotted lines.

similar and strongly nonlinear for the two studied periods. As described in Sect. 3.1, large values at low temperatures result from mixing with enriched deep waters during winter, and low values for 26–28 ◦C temperatures occur at the end of summer after the biological drawdown of carbon. An increase in $f \text{CO}_2 \text{@} 13$ between the two periods is evident across the range of temperatures.

3.3.2 Trend analysis and statistics

To quantify the change in $fCO₂@13$ between the two data sets, we proceed as follows: data are binned by $1 \degree C$ temperature intervals, thereby removing any potential seasonal weighting, especially towards the 13–14 °C winter months temperature. The measurements made in this temperature interval represent about 25 % of the total number of data for both periods. For each of the fourteen $1 \degree C$ steps, the mean and standard deviation of hourly $fCO₂@13$ measurements are reported in Table 1 and in Fig. 3c. The mean temperature within each of the 1[°] steps differ for the two periods, as the distribution of individual measurements are not identical.

For both data sets, a monotonic relationship between $fCO₂@13$ and T is observed with correlation coefficients respectively equal to −0.861 and −0.857. The difference in $fCO_2@13$ between the two periods, $dfCO_2@13$, is derived in each temperature step, seen in the difference between column 2 and 6 of Table 1. The variability of this difference is estimated as the quadratic mean of the standard deviation in each time series. Both values are reported in Table 1, column 9 and 10 and in Fig. 3c. The distribution of each $dfCO_2@13$ values around the mean over all SST of $dfCO₂@13$ seems random and indicates no trend of dependency with SST (Fig. 3c). This suggests that the processes that control the seasonal variation of $fCO₂@13$ at the sea surface have not changed over the last two decades.

We have estimated the uncertainties in the estimates of the difference $dfCO_2@13$ with two methods. Firstly, the arithmetic mean of d $fCO₂@13$ is equal to 33.17 µatm, with a standard deviation (SD) and standard error (SE), respectively equal to 6.29 µatm and 1.68 µatm. A 95 % confidence interval is thereby achieved within 1.96 SE, i.e. 3.29 µatm. A second approach consists of computing a weighted average of the mean of $dfCO_2@13$. In this case, mean weighted value of $dfCO_2@13$ over the whole range of temperatures is estimated, the weights being equal to the variance of $dfCO_2@13$ in each temperature step. The value of $dfCO_2@13$ is equal to 32.70 µatm. The weighted SD, and the associated SE, of the 14 data points are respectively

	Time interval 1995–1997				Time interval 2013–2015				Temporal change	
T^1 \circ C	fCO ₂ $@13$ uatm	N	Standard deviation µatm	$T^{\, 1}$ $\rm ^{\circ}C$	fCO ₂ $@13$ µatm	N	Standard deviation µatm	dfCO ₂ @13 uatm	Standard deviation uatm	
13.45	331.58	1212	28.09	13.55	363.14	6869	18.07	31.56	33.40	
14.45	305.28	495	26.02	14.43	337.16	3270	16.65	31.87	30.89	
15.37	281.54	447	9.62	15.57	321.10	3112	11.09	39.56	14.68	
16.44	274.43	182	8.53	16.42	313.79	1818	11.09	39.36	13.99	
17.58	275.54	190	7.04	17.56	306.83	1528	14.65	31.29	16.25	
18.47	277.34	300	9.04	18.45	296.57	2621	10.95	19.23	14.20	
19.62	265.43	342	15.58	19.41	291.84	1406	13.45	26.40	20.59	
20.50	258.08	529	14.15	20.50	293.16	1135	18.21	35.08	23.06	
21.56	271.15	239	12.98	21.54	297.96	1200	20.41	26.82	24.19	
22.49	250.75	742	13.66	22.49	290.27	2385	18.57	39.52	23.05	
23.57	252.22	320	13.00	23.47	296.92	747	21.77	44.70	25.36	
24.41	245.85	506	7.08	24.40	280.44	959	14.82	34.59	16.43	
25.50	250.06	215	10.77	25.53	284.05	456	14.81	33.99	18.31	
26.42	256.29	279	6.24	26.29	286.71	249	11.23	30.42	12.85	

Table 1. Distribution of temperature, $f \text{CO}_2 \text{@13}$, and increase $df \text{CO}_2 \text{@13}$ data binned by 1 °C temperature interval for the two periods 1995–1997 and 2013–2015.

equal to 4.85 and 1.30 µatm. A 95 % confidence interval is achieved within 2.54 µatm. The difference between the two mean $dfCO_2@13$ estimates is 0.47 µatm, well below SE. In the following, we have chosen the former method.

3.4 Changes of seawater carbonate chemistry in surface waters

We estimated the DIC and pH changes related to the increase in $f \text{CO}_2 \text{@} 13$ measured at the sea surface 18 years apart, assuming a mean salinity equal to 38.2, a mean alkalinity equal to 2562.3 µmol kg^{-1} following Eq. (1) and a mean in situ temperature (T) equal to 18.25 °C. The dissociation constants of Mehrbach refitted by Dickson and Millero (Dickson and Millero, 1987; Mehrbach et al., 1973) were used. pH is calculated on the seawater scale. The uncertainty of $dfCO_2@13$, 3.3 µatm, has been propagated to compute the combined uncertainty in dDIC and dpH_{SWS}. The uncertainties in the equilibrium constants are neglected in this propagation of uncertainties. Likewise, an implicit assumption is that there is no systematic error on DIC and pH_{SWS} derived from $fCO₂@13$ between the two time periods; in particular, mean temperature and salinity remain the same (Sect. 3.2). This is further discussed in Sect. 4.1. We compute an increase in DIC and dDIC equal to $25.2 \pm$ 2.7 µmol kg⁻¹ (1.40 ± 0.15 µmol kg⁻¹ yr⁻¹) and a decrease in pH_{SWS} and dpH_{SWS} equal to -0.0397 ± 0.0042 pH_{SWS} $(-0.0022 \pm 0.0002 \,\text{pHsws} \,\text{yr}^{-1})$ (Table 2).

3.5 Changes in atmospheric and seawater $fCO₂$

The increase in atmospheric $fCO₂$ from 1995–1997 to 2013–2015 was computed from the monthly atmospheric $xCO₂$ concentrations measured at the Lampedusa Island station (Italy) $(35°31' N, 12°37' E)$ [\(http://ds.data.jma.go.jp/](http://ds.data.jma.go.jp/gmd/wdcgg/) [gmd/wdcgg/,](http://ds.data.jma.go.jp/gmd/wdcgg/) last access: 14 September 2018) (see Eq. 3 in Hood and Merlivat, 2001). Considering a mean annual in situ temperature equal to 18.25 °C and an atmospheric pressure of 1 atm, we derived a mean atmospheric $fCO₂$ equal to 355.3 ± 0.8 µatm for 1995–1997 and 389.6 ± 0.9 µatm for 2013–2015, which is an increase of 34.3 ± 2.3 µatm (95 % confidence interval) (Table 2). At this temperature, the change in $fCO₂$ at the sea surface is 41.4 ± 4.1 µatm. Thus the contribution of the increase in atmospheric $CO₂$ is responsible for $84\% \pm 5\%$ of the increase in $fCO₂$ measured in the surface waters. With the same salinity and alkalinity as previously, the measurement of the corresponding change in surface DIC, assuming air–sea equilibrium, would be 20.8 ± 1.3 μmol kg⁻¹ (Table 2).

4 Discussion

4.1 Time change in surface alkalinity

High-frequency measurements of $fCO₂$ and temperature over two periods of 3 years that are 2 decades apart have allowed the computation of an increase in DIC equal to 25.1 ± 2.3 µmol kg⁻¹ assuming no change in alkalinity. In the range of salinity of the BOUSSOLE samples, 37.9 to 38.5, the alkalinity values computed with Eq. (1) are larger than those predicted by the relationship established for the DYFAMED site, with a mean difference equal to 10 ± 2 µmol kg−¹ (Copin-Montégut and Bégovic, 2002). In both cases alkalinity measurements were made with a potentiometric method using the certified reference material

	dfCO ₂ ^a $@13$ µatm	dfCO ₂ ^a $@T$ uatm	dDIC ^a μ mol kg ⁻¹	$dpHsws$ ^c pH unit	dfCO ₂ $@T$ annual μ atm yr $^{-1}$	dDIC annual μ mol kg ¹ yr ¹	$dpHsws$ ^c annual pH unit yr ¹
Sea surface	33.2 ± 3.3	41.4 ± 4.1	25.2 ± 2.7	-0.0397 ± 0.0042	2.30 ± 0.23	1.40 ± 0.15	-0.0022 ± 0.0002
Atmosphere Lampedusa data		34.3 ± 2.3	20.8 ± 1.3^{b}		1.91 ± 0.13	1.15 ± 0.07	
dfCO ₂ @T _{air} / $dfCO$ ₂ QT_{sea}		0.83 ± 0.10	0.83 ± 0.09				

Table 2. Seasonally detrended long-term and annual trends of seawater carbonate chemistry and atmospheric composition.

T, mean annual temperature equal to 18.25 °C; ^a change from 1995–1997 to 2013–2015; ^b dDIC_{ant}; ^c dpH_{SWS} computed at T.

supplied by A. G. Dickson for calibration. It is difficult to identify the cause for a possible change in alkalinity between the two periods that are 18 years apart, while no salinity change has been observed. At a coastal site 50 km away from DYFAMED, Kapsenberg et al. (2017) have measured an increase in alkalinity unrelated to salinity over the period from 2007 to 2015. They attribute it to changes in freshwater inputs from land. However, based on data from Coppola et al. (2016), alkalinity in the upper 50 m at DY-FAMED did not change significantly from 2007 to 2014 $(3.204 \,\mathrm{\mu mol \,kg^{-1}}$, $P = 0.0794$, $r^2 = 0.08$). Thus, we cannot conclude whether the difference observed at DYFAMED and BOUSSOLE between the two periods is real or an artifact of measurement techniques. As a sensitivity test, we compute the expected changes in DIC and pH from 1995–1997 to 2013–2015 for a mean alkalinity increase in 10 μ mol kg⁻¹. We obtain annual changes of dDIC = $+0.46$ µmol kg⁻¹ yr⁻¹ and dpH = -0.0001 pH unit yr⁻¹, which are well below errors estimated in Sect. 3.4. Hence, such a change in alkalinity does not significantly affect the increase in DIC and the decrease in pH shown in Table 2.

4.2 Drivers of the temporal change in DIC in surface waters

The increase in sea surface DIC from 1995–1997 to 2013– 2015 is 25.2 ± 2.7 µmol kg⁻¹ (Table 2), whereas the expected contribution due to ocean uptake of anthropogenic $CO₂$ is 20.8 ± 1.3 µmol kg⁻¹. In order to interpret the difference between these two values, we examine potential changes that may result from the inter-annual variability in local physical and biological processes or anthropogenic carbon invasion from lateral advection of Atlantic waters.

4.2.1 Natural variability

Time series of the mixed layer depth (MLD) show a strong variability in winter at inter-annual scale. During the two periods, 1995–1997 and 2013–2015, the winter MLD never exceeded 220 m, whereas values over 300 m were observed in 1999 and especially in February and March 2006, with values close to 2000 m (Coppola et al., 2016; Pasqueron de Fommervault et al., 2015). These episodes of strong and deep vertical mixing must have entrained DIC-rich LIW in the surface waters. This entrainment could be causing an increase in DIC between the 1995–1997 and 2013–2015 periods. Monthly surface samples collected at the Dyfamed time series station between 1998 and 2013 indicate an increasing DIC trend of 1.35 µmol kg⁻¹ yr⁻¹. This value is uncertain $(r^2 = 0.05)$ because of the large seasonal variability displayed in the monthly samples (Gemayel et al., 2015). Nevertheless, this value is closer to the trend we calculated between the two periods, 1993–1995 and 2013–2015 $(1.40 \,\mu\text{mol kg}^{-1}\text{ yr}^{-1})$ than to the trend inferred from the atmospheric increase (1.15 µmol kg⁻¹ yr⁻¹). In the DYFAMED time series, we find no evidence that the strong increase in MLD observed during winters 1999 and especially 2006 resulted in a further increase in DIC.

The monthly cruises of the Dyfamed time-series study have also been analysed in order to investigate the hydrological changes and some biological consequences over the period 1995–2007 (Marty and Chiavérini, 2010). These authors show that extreme convective mixing events such as those recorded in 1999 and 2006 are responsible for large increases in nutrient content in surface layers and conclude that biological productivity is increasing, especially during the 2003–2006 period, which could lead to a larger consumption of carbon, i.e. a decrease in DIC.

4.2.2 Anthropogenic carbon exchange through the Strait of Gibraltar.

The concentration of oceanic anthropogenic carbon (C_{ant}) is not a directly measurable quantity. To estimate it, several empirical methods have been developed. Flecha et al. (2012) computed the anthropogenic carbon inventory in the Gulf of Cadiz. They used observations made during a cruise in October 2008 throughout the oceanic area covered by the Gulf of Cadiz and the Strait of Gibraltar to estimate C_{ant} with three methods, which are ΔC^* (Gruber et al., 1996), TrOCA (Touratier and Goyet, 2004; Touratier et al., 2007)

and φC_T^0 (Vázquez-Rodríguez et al., 2009). In the three cases, their results indicate a net import of Cant from the Atlantic towards the Mediterranean through Gibraltar.

Schneider et al. (2010), using the transit-time distribution method applied to a data set from a Mediterranean cruise in 2001, estimated a net anthropogenic carbon flux across the Strait of Gibraltar into the Mediterranean Sea of 3.5 Tg C yr^{-1} . Over the whole period from 1850 to 2001, this contribution of C_{ant} represents almost 10% of the total Cant inventory of the Mediterranean Sea. Accordingly, about 90 % must have been taken directly from an equilibrium with atmospheric $CO₂$. Based on a high-resolution regional model, Palmiéri et al. (2015) computed the anthropogenic carbon storage in the Mediterranean basin. They concluded that 75 % of the total storage of C_{ant} in the whole basin comes from the atmosphere and 25 % from net transport from the Atlantic through the Strait of Gibraltar. The findings of these two studies support our estimated change in DIC of (17 ± 10) % in addition to the direct contribution of air–sea exchange suggesting that it could result from the anthropogenic carbon input from the Atlantic Ocean towards the Mediterranean basin.

Huertas et al. (2009) and Schneider et al. (2010) report DICant surface concentrations respectively equal to 65– 70 µmol kg−¹ at the Strait of Gibraltar in the years 2005– 2007 and close to 65 µmol kg^{-1} in the western basin in 2001. We extrapolate these figures to the year 2014, assuming a mean increased rate of DIC equal to 1.4μ mol kg⁻¹ yr⁻¹ as previously computed (Table 2). Taking into account the increase in DIC_{ant} equal to 25.2 µmol kg⁻¹ between 1995– 1997 and 2013–2015, we estimate that the contribution of the change in DIC_{ant} over the last 18 years represents \sim 30 % of the total change since the beginning of the industrial period $(t\geq 1800)$.

4.3 Long-term trends in surface DIC and pH

The annual changes in DIC and pH_{SWS} calculated between 1995–1997 and 2013–2015 are respectively equal to $1.40 \pm$ 0.15 µmol kg⁻¹ and -0.0022 ± 0.0002 . At the DYFAMED site, at 10 m, Marcellin Yao et al. (2016) studied the time variability of pH over 1995–2011 based on measurements of T , S , Alk and DIC sampled approximately once a month. They computed a mean annual decrease in pH, $-0.003 \pm$ 0.001, which is not significantly different from our estimate. For the global surface ocean, Lauvset et al. (2015) have reported a mean rate of decrease in pH, -0.0018 ± 0.0004 for 1991–2011. This value is also within the limits of uncertainty of the pH change computed in our study.

Bates et al. (2014) examined changes in surface seawater CO₂-carbonate chemistry at the locations of seven ocean $CO₂$ time series that have been gathering sustained observations from 15 to 30 years with monthly or seasonal sampling. Six stations are located in the Atlantic and Pacific oceans in a latitudinal band between 10 and 68◦ N. The range of increasing and decreasing annual trends of DIC and pH extends from 0.93 ± 0.24 to 1.89 ± 0.45 µmol kg⁻¹ yr⁻¹ and -0.0014 ± 0.0005 to -0.0026 ± 0.0006 , respectively. The Revelle factor of surfaces waters vary from 9–10 in low latitudes to 12–15 in the subpolar time series sites, with higher Revelle factor values reflecting a reduced capacity to absorb atmospheric $CO₂$. The data show that the increase in DIC is not only controlled by the buffer capacity of the water but also by the compounding effects of changes in physical factors like the strengthening of winter mixing or larger air–sea uptake (Olafson et al., 2010).

The increase in DIC computed at DYFAMED is in the upper range of values reported in the other time series. A low Revelle factor of close to 10 characterises the Mediterranean Sea because of its warm and high-alkalinity waters. Moreover, as the result of a relatively short deep-water renewal time estimated to be 20–40 years in the western basin (Schneider et al., 2010), the waters of the Mediterranean Sea have a relatively high capacity to absorb anthropogenic $CO₂$ from the atmosphere and transport it to deep water.

The calculated decrease in pH in the surface water at DY-FAMED and in the global ocean are quite similar, despite the higher alkalinity of the Mediterranean Sea. Thermodynamic equilibrium calculations have highlighted the alkalinity's effect on the anthropogenic acidification of the Mediterranean Sea (Palmiéri et al., 2015). Their results show that, notwithstanding a higher total alkalinity, the average anthropogenic change in the surface pH of the Mediterranean Sea does not differ significantly from that of the average of global oceans.

5 Conclusion

High-frequency ocean $fCO₂$ measurements made by CAR-IOCA sensors have been used to calculate trends in $fCO₂$, DIC and pH over a period of two decades, notwithstanding the short-time and natural seasonal variability of these properties at the sea surface. We have estimated a large change in sea-surface carbonate chemistry, an increase in DIC and a decrease in pH. The computed increase in DIC is larger than the change expected from a chemical equilibrium with atmospheric $CO₂$. This could be the result of the strong interannual variability of the winter mixing observed between the two periods of 1993–1995 and 2013–2015. Likewise, our results support modelling work and analysis of vertical profiles measurements that suggest that the Atlantic Ocean contributes a substantial amount of anthropogenic carbon to the Mediterranean basin, (17 ± 10) %, which lies between the estimates of 10 % (Schneider et al., 2010) and 25 % (Palmiéri et al., 2015).

Data availability. The time series data from Dyfamed (1995–1997) are available in the SOCAT v3 database (Bakker et al., 2016). The Boussole data (2013–2015) are available in the SEANOE database (Merlivat and Boutin, 2018).

Competing interests. The authors declare that they have no conflict of interest.

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