Atmospheric occurrence, transport and deposition of polychlorinated biphenyls and hexachlorobenzene in the Mediterranean and Black Seas

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Received: 27 February 2014 – Accepted: 24 March 2014 – Published: 14 April 2014

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Published by Copernicus Publications on behalf of the European Geosciences Union.
Abstract

The Mediterranean and Black Seas are unique marine environments subject to important anthropogenic pressures due to direct and indirect loads of atmospheric inputs of organochlorine compounds (OCl) from primary and secondary sources. Here we report the results obtained during two east-west sampling cruises in June 2006 and May 2007 from Barcelona to Istanbul and Alexandria, respectively, where gas phase and aerosol samples were collected. Both matrices were analyzed for 41 polychlorinated biphenyls (PCBs), including dioxin-like congeners, and hexachlorobenzene (HCB). The values reported in this study for gas phase HCB and $\sum_{41}^{41}$PCB (LOD to 418.3 pg m$^{-3}$ and from 81.99 to 931.6 pg m$^{-3}$ respectively) are in the same range of those reported in former studies, possibly suggesting a limited decline in their atmospheric concentrations during the last decade for the Mediterranean region due to land base OCl sources. There is a clear influence of the direction of the air-mass on the atmospheric concentrations of PCBs, with higher concentrations when the air mass was from southern Europe, and the lowest concentrations for air masses coming from the SW Mediterranean and Atlantic Ocean. PCBs and HCB are close to air–water equilibrium for most sampling periods, thus resulting in low atmospheric deposition fluxes at open sea. This is consistent with the oligotrophic character of the Mediterranean Sea with a small influence of the biological pump capturing atmospheric PCBs. Therefore, degradation of gas-phase PCBs by OH radicals is estimated to be the main loss process of atmospheric PCBs during their transport over the Mediterranean Sea. Conversely, atmospheric residence times of HCB are predicted to be very long due to a lack of atmospheric degradation and low depositional fluxes due to concentrations at air–water equilibrium.

1 Introduction

Among the thousands of anthropogenic compounds emitted to the atmosphere, chemicals embraced within the group of the Persistent Organic Pollutants (POPs) have...
obtained a relevant attention due to their persistence, bioaccumulation potential and proved toxic effects (United Nations, 2014). Organochlorine compounds (OCl), such as polychlorinated biphenyls (PCBs) and hexachlorobenzene (HCB), are among the POPs that have been studied extensively due to their ubiquitous occurrence, and potential for long range atmospheric transport, being a paradigm of globally distributed pollutants. In addition, the wide range of physical-chemical properties of PCBs makes them useful as surrogate of the behavior of other semi-volatile and hydrophobic POPs, including those which presence in the environment is relatively recent and thus not yet properly quantified at regional and global scales. Although international regulations such as the Stockholm Convention have banned or restricted the global emissions of many OCls, the consequent expected decline trend in their atmospheric concentrations has not been as fast as predicted, and has varied upon location/region, time period and specific compounds (Hung et al., 2005). In fact, current atmospheric levels of these chemicals in the atmosphere may be explained as a result of the interplay of different factors (Nizzetto et al., 2010a, b): (i) primary sources as a direct result of the intentional production and usage or accidental releases of materials containing OCls; (ii) secondary emission sources as the consequence of re-volatilization of previously released OCls from environmental reservoirs, such as sea–air or soil–air diffusive exchange (Brevik et al., 2002) and (iii) the removal processes which are also particular for each compound (Wania and Daly, 2002; Dachs et al., 2002). Nevertheless, there are still large uncertainties regarding the different factors controlling the atmospheric occurrence and deposition of OCls, especially when referring to the marine atmospheric environment due to the scarcity of the available data and the difficulties associated with the generation of reliable new measurements.

The persistence and subsequent long half-lives in air make OCls prone for long range atmospheric transport before their removal by deposition or reaction with OH radicals (Bidleman, 1999; Anderson and Hites, 1996; Mandalakis et al., 2003; Jurado and Dachs, 2008). The distribution of the compounds in the atmosphere between the aerosol and gas phases will determine to great extent the relative importance of these
loss processes. The inputs of OCI to the open sea come from the combination of three depositional processes: dry deposition driven by particle settling (Jurado et al., 2004), wet deposition through scavenging of OCIs by rain or snow (Jurado et al., 2005), and especially diffusive air–water exchange (Jaward et al., 2004; Jurado et al., 2004). Atmospheric deposition supports OCIs accumulation in biota at open sea, where their harmful effects could threaten the marine ecosystem, and humans through different exposure routes. In consequence, it is important to gain knowledge of the occurrence and deposition of POPs in the marine background atmosphere in order to quantify their inputs, dynamics, and impacts in the oceanic environment.

The Mediterranean region is surrounded by Europe and Africa where highly developed industrial countries in the North stand in contrast to the developing countries in the Southern Basin. Historically, the Mediterranean and Black Seas have been recipients of significant anthropogenic pressures of direct and indirect loads of OCIs from intensive industrial and agricultural activities (Tolosa et al., 1997; Turley et al., 1999). However, the atmospheric measures of OCI concentrations in the Mediterranean Sea region are scarce. For the open sea these have been focused on polychlorinated dibenzo dioxins and furans (Castro-Jiménez et al., 2010), while for other OCI the previous studies have only covered local coastal sites (Mandalakis et al., 2002; Garcia-Flor et al., 2009; Castro-Jiménez et al., 2010). Therefore, the main objectives of this work are: (i) to provide a large dataset of PCB and HCB concentrations in the open Mediterranean and SW Black Sea atmosphere; (ii) to quantify the atmospheric deposition of PCBs and HCB to the Mediterranean Sea, and (iii) evaluate the factors and processes affecting the occurrence and deposition of PCBs and HCB in the Mediterranean marine atmosphere.
Materials and methods

2.1 Sample collection

Air samples were collected on board the R/V Garcia del Cid during the two THRESH-OLDS sampling cruises carried out in June 2006 and May 2007. In both cruises, Barcelona was the initial and final port, with Istanbul and Alexandria being the intermediate stops, respectively. Transects covered an extensive area within a year of difference allowing for a good spatial coverage of the different Mediterranean Basins (Fig. 1).

The air samples for OCl analysis were taken using a high-volume air sampler (MCV, Barcelona, Spain) operating at a flow rate of 40 m$^3$ h$^{-1}$ located above the bridge (at 6–7 m.a.s.l). On average, samples were collected within twelve hours, and the mean sampled air volume was of 500 m$^3$, ranging from 230 m$^3$ to 950 m$^3$. In addition, a second high-volume sampler was operated simultaneously to collect samples with larger sampled volumes allowing for the determination of mono-ortho dioxin-like PCB (300–1600 m$^3$) (see Supplement A). Both high-volume air samplers were operated connected to a weather station to avoid smokestack and minimize interference from any potential ship contamination. The air was drawn through a precombusted Quartz Filter (QM-A; Whatman, 8 × 10 inches) to collect aerosol bound compounds and then circulated through a polyurethane foam (PUF) to collect chemicals present in the gas phase. All samples were stored in freezers at −20°C until analysis. Preceding sampling, PUF were rinsed with acetone : hexane (1 : 1, v/v), then Soxhlet extracted for 24 h and dried into a vacuum desiccator.

Throughout the campaign, meteorological parameters, including temperature, wind speed and wind direction, were measured using the meteorological station located on the deck of the ship. In order to examine the origin of the air masses for each sampling period and region, air mass back-trajectories were calculated using the HYSPLIT model of NOAA. 48 h back-trajectories were determined every 6 h for 15 and 100 m of altitude at the start and end locations of each sampling transect (see Supplement B).
2.2 Chemical analysis

Prior extraction, all samples were spiked with PCB 65 and PCB 200 which were used as surrogate standards. PUFs were Soxhlet extracted with acetone : hexane (1 : 1, v/v) for 24 h. The extracts were rotary evaporated to 1 mL and purified on a 3% Milli-Q grade water deactivated alumina column (3 g) with a top layer of anhydrous sodium sulfate. Each column was eluted with 5 mL of hexane. The extract was concentrated to 0.5 mL by vacuum rotary evaporation, transferred to a 1.7 mL amber vial with isooctane and evaporated to 150 µL under a gentle nitrogen stream. At this step, 5 ng of PCB 30 and PCB 142, used as internal standards, were added to the extract. QM-A filters were freeze-dried, weighed and Soxhlet extracted with dichloromethane : methanol (2 : 1, v/v) for 24 h. The extracts were rotary evaporated to 1 mL, solvent-exchanged to hexane and purified by column chromatography as indicated for gas phase samples. The resultant extract was treated analogously as for gas-phase samples.

All samples were analyzed for 41 PCB congeners and HCB by gas chromatography coupled to an electron capture detector operating with splitless injection mode. The separation was achieved with a 60 m × 0.25 mm i.d. × 0.25 µm capillary column. The oven temperature was programmed from 90 °C (holding time 1 min) to 190 °C at 2 °C min⁻¹, and finally 310 °C at 3 °C min⁻¹ keeping the final temperature for 18 min. Compounds were quantified by the internal standard procedure (see Supplement C for instrumental detection limits).

In addition, a selected group of samples with higher sampled volumes were also used to quantify 8 mono-ortho PCB congeners (PCB 105, PCB 114, PCB 118, PCB 123, PCB 156, PCB 157, PCB 167 and PCB 189). The analysis of this group of samples containing 18 gas and aerosol phase samples (see Supplement A) was performed by isotope dilution high-resolution gas chromatography coupled to a high resolution mass spectrometer (HRGC-HRMS) on the basis of EPA 1668 method (Castro-Jiménez et al., 2010).
2.3 Quality assurance and quality control

The analytical procedure was validated by determining the recovery rates of the surrogates for each sample (see Supplement C). Moreover, a total of ten blank samples (five for PUF, five for QM-A filters) were collected to test the background contamination during the sample preparation, transportation, treatment and analysis. These blanks were processed in parallel to the samples (see Supplement C).

2.4 Estimation of the atmospheric deposition of PCBs and HCBs

The dry deposition fluxes \( F_{\text{DD}} \), ng m\(^{-2}\) d\(^{-1}\) were calculated as:

\[
F_{\text{DD}} = v_{\text{D}} \cdot C_{\text{AP}}
\]  

where \( C_{\text{AP}} \) is the volumetric concentration of PCBs or HCB in the aerosol phase (ng m\(^{-3}\)) and \( v_{\text{D}} \) is the deposition velocity of particles (m d\(^{-1}\)). Due to the lack of field measurements of \( v_{\text{D}} \) over the Mediterranean Sea and during the THRESHOLDS cruises reported here, we assume a value of 0.2 cm s\(^{-1}\) as representative of marine aerosol deposition velocity. This value is slightly above the average value measured in the Atlantic Ocean (Del Vento and Dachs, 2007a), but it is slightly lower than those measured at a coastal Western Mediterranean station (Del Vento and Dachs, 2007b) and within the range of what was calculated for the coastal Eastern Mediterranean (Mandalakis et al., 2005).

The diffusive air–water exchange fluxes \( F_{\text{AW}} \), ng m\(^{-2}\) d\(^{-1}\) were estimated by:

\[
F_{\text{AW}} = k_{\text{AW}} \left[ \frac{C_{\text{G}}}{H'} - C_{\text{W}} \right]
\]  

where \( k_{\text{AW}} \) is the air–water mass transfer coefficient (m d\(^{-1}\)), \( C_{\text{G}} \) (pg m\(^{-3}\)) and \( C_{\text{W}} \) (pg m\(^{-3}\)) are the OCl concentration in the gas and dissolved phases, respectively, and \( H' \) is the temperature and salinity corrected dimensionless Henry’s law constant. \( C_{\text{W}} \)
were measured simultaneously to the atmospheric concentrations and are discussed elsewhere (Berrojalbiz et al., 2011). For determining both diffusive air–water exchange fluxes and air–water fugacity-ratios (see below), $C_W$ was corrected by the dissolved organic carbon as reported elsewhere (Garcia-Flor et al., 2005). Details on estimation methods for $k_{AW}$ are explained elsewhere (Jurado et al., 2004). The nonlinear influence of wind speed has been taken into account by correcting $k_{WA}$ considering a Weibull distribution of wind speeds (Simó and Dachs, 2002).

The fugacity approach has been used to further evaluate the significance of the direction of the diffusive air–water exchange ($f_G/f_W$) and estimated by,

$$\frac{f_G}{f_W} = \frac{C_G}{C_W H'}$$

Due to the uncertainty in $H'$ values, when $f_G/f_W$ ranges from 0.3 to 3, gas and dissolved phases can be assumed to be close to equilibrium, when $f_G/f_W > 3$ there is a net absorption, and when $f_G/f_W < 0.3$ there is a net volatilization.

### 3 Results and discussion

#### 3.1 Atmospheric occurrence of PCBs and HCB

A summary of atmospheric concentrations in the gas ($C_G, N = 34$) and aerosol phases ($C_{AP}, N = 30$) for HCB and individual PCBs is given in Table 1 (complete data set in Supplement D). The atmospheric concentrations of the 8 dioxin-like PCB congeners are summarized in Table 3 for the gas ($C_G^*, N = 18$) and aerosol phase ($C_{AP}^* = 18$) (see Supplement D for complete data set). The sum of the measured 41 PCB congeners ($\sum_{41}\text{PCB}$) ranged from 82.0 pg m$^{-3}$ to 931.6 pg m$^{-3}$, with a mean value of 297.3 pg m$^{-3}$ for the gas phase in the Mediterranean Sea atmosphere. Likewise, for the samples collected in the SW Black Sea, the $\sum_{41}\text{PCB}$ gas phase concentration ranged from 206.2 pg m$^{-3}$ to 644.2 pg m$^{-3}$, with a mean concentration of 421.6 pg m$^{-3}$. The highest
$C_G$ for the $\sum_{41}$ PCB corresponded to the sample taken between Barcelona and Mallorca in 2007 (T2) followed by the sample taken in the Black Sea in 2006 (STIVc). The lowest $\sum_{41}$ PCB concentrations were measured in transects from diverse locations in the open Mediterranean Sea (T4, T9 and T17). There was not a significant ($p < 0.05$) east-west trend in $\sum_{41}$ PCB gas phase concentrations for the Mediterranean Sea, and there were no statistically significant differences between those samples taken in 2006 and 2007, nor between the Mediterranean and Black Seas. The lack of significant differences between sub-basins and years, points out to other factors driving the observed variability in concentrations (see Sect. 3.3).

In order to compare PCB levels measured in this study with those reported in former works, the seven ICES PCB congeners (28, 52, 118, 101, 153, 138, and 180) were selected (Table 2). $\sum_{7 \text{ICES}}$ PCBs ranged from 27 to 259.4 pg m$^{-3}$ in the Mediterranean Sea’s gas phase, and from 98.70 to 166.15 pg m$^{-3}$ in the Black Sea, with mean values of 88.1 and 136.9 pg m$^{-3}$, respectively. On average, $\sum_{7 \text{ICES}}$ PCBs account for 34% of $\sum_{41}$ PCB in both regions. The $\sum_{7 \text{ICES}}$ PCBs gas phase concentrations are in agreement with previous measurements made in North-Western Mediterranean Sea in 2002–2003 (Garcia-Flor et al., 2009) and in the Eastern Mediterranean Sea in 1999–2001 (Mandalakis et al., 2002). The $\sum_{41}$ PCB concentrations reported in this study are also in the same range of what was found by Iwata and coworkers (sum of 40 congeners) (Iwata et al., 1993). This comparison suggests a limited decline in PCB atmospheric concentrations for the last 15 years in the Mediterranean Sea region. There are no previous reports of the atmospheric occurrence of PCBs for the Black Sea region. In addition, the concentrations reported here for the Mediterranean atmosphere are not significantly different from those in the subtropical Atlantic Ocean (Jaward et al., 2004; Gioia et al., 2008). Nevertheless, in this study the variability of concentrations of individual congeners is higher, probably caused by the influence of the surrounding continents and the more variable atmospheric circulation patterns over the Mediterranean.

Aerosol phase $\sum_{41}$ PCB concentrations varied from 0.8 pg m$^{-3}$ to 35.7 pg m$^{-3}$ in the Mediterranean Sea and from 8.3 pg m$^{-3}$ to 19.4 pg m$^{-3}$ in the Black Sea, with average...
values of 14.3 pg m\(^{-3}\) and 19.8 pg m\(^{-3}\), respectively. The maximum volumetric concentrations of aerosol phase \(\sum_{41}\) PCB were found in the western Mediterranean Sea, whereas the lowest levels were obtained in one of the samples collected in W. Mediterranean (T3), and three samples from the Levantine Basin (T11, T12 and T17). When the aerosol phase \(\sum_{41}\) PCB concentrations were normalized by the concentration of aerosols (\(C_{\text{AP}}\) given as ng g\(^{-1}\)), these ranged from 21.5 ng g\(^{-1}\) to 653.8 ng g\(^{-1}\) with an average value of 35.5 ng g\(^{-1}\) for the Mediterranean Sea, while Black Sea aerosol phase concentrations varied from 70.5 ng g\(^{-1}\) to 35.6 ng g\(^{-1}\), exhibiting a mean concentration of 91.7 ng g\(^{-1}\). The highest aerosol phase \(\sum_{41}\) PCB concentrations, as normalized by the aerosol mass, were found in samples taken between Barcelona and Mallorca (T1 and T2), while lowest values were also detected in sample T3 and in samples taken in the Levantine Basin. Therefore, the western Mediterranean Sea presented higher concentrations for particle bound \(\sum_{41}\) PCB than the eastern basin, either normalized by filtered volume or amount of particles, being Barcelona and Mallorca the regions showing higher \(C_{\text{AP}}\) values. Particle bound \(\sum_{\text{ICES}}\) PCBs varied from 0.4 pg m\(^{-3}\) to 17.9 pg m\(^{-3}\) in the Mediterranean Sea and from 4.5 pg m\(^{-3}\) to 11.55 pg m\(^{-3}\) in the Black Sea, accounting for 47 % and 57 % of the \(\sum_{41}\) PCB respectively. These concentrations are in the lower range than those concentrations reported for low populated coastal regions of the North western Mediterranean Sea (Garcia-Flor et al., 2009) and in the same range of the levels measured in an Eastern Mediterranean Sea coastal station (Mandalakis et al., 2005).

The average levels of mono-ortho substituted PCBs (dioxin-like congeners) in both gas and aerosol phases were generally lower than for the rest of PCB congeners, with average concentrations ranging from 0.03 pg m\(^{-3}\) to 1.53 pg m\(^{-3}\), and from 0.001 pg m\(^{-3}\) to 0.2 pg m\(^{-3}\), respectively (Table 3). The concentrations in all the cases were fairly constant, showing non-significant spatial variations neither in the gas nor in the aerosol phases. The long sampling time periods for this subset of samples (4 days) may have produced an averaging of the air masses from different origins leading to the observed lack of variability. There is no data available for mono-ortho PCBs congeners
in the literature for the Mediterranean Sea. However, the values reported here were generally lower than the atmospheric concentrations reported for the same congeners in a coastal Mediterranean lagoon (1–7 and 0.1–0.6 pg m\(^{-3}\) gas and aerosol phase levels, respectively) (Castro-Jiménez et al., 2011).

The gas phase concentrations of HCB varied from LOD to 418.3 pg m\(^{-3}\) (mean of 54.4 pg m\(^{-3}\)), with higher concentrations in the Western than in the Eastern Mediterranean. The highest \(C_G\) for HCB was detected in sample STIIa collected between Mallorca and Sardinia. The average HCB concentrations for the Mediterranean and Black Seas (59.2 pg m\(^{-3}\) and 42.0 pg m\(^{-3}\) respectively) are in agreement with measures performed by other studies in the Mediterranean region (Albaiges et al., 2005) and the Atlantic Ocean (Jaward et al., 2004; Lohmann et al., 2009), even though we observed a larger variability in concentrations.

The aerosol phase HCB varied from non-detected to 0.7 pg m\(^{-3}\), with an average value of 0.1 pg m\(^{-3}\). Samples with no HCB were located in the Levantine Basin and Black Seas whereas the maximum value was measured in the Marmara Sea, near the city of Istanbul. Particle normalized aerosol phase HCB concentrations did not show any significant spatial variations with the exception of two peak values in samples collected between Barcelona and Mallorca, and near Istanbul. There are no previous studies of aerosol phase HCB for the Mediterranean atmosphere.

### 3.2 Aerosol bulk composition and aerosol–gas partitioning of PCBs and HCB

A Principal Component Analysis (PCA) was performed for congener groups to assess the variability of PCBs in the aerosol phase (see Supplement E). When using volumetric concentrations (pg m\(^{-3}\)), the first two principal components (PC1 and PC2) explaining 41% and 30% of the variability, respectively, separated samples in a gradient of concentrations (see Supplement E). In addition, two groups of samples were differentiated from the rest of samples, both of them characterized by the highest \(C_{AP}\) levels. This discrimination between sample groups was also well defined when the PCA was generated with particle-normalized concentrations (in ng g\(^{-1}\)). The first group was
comprised by samples located between Barcelona and Mallorca (T1 and T2), consistent with the influence of the air masses coming from the industrialized and populated coast of Catalonia (NW Mediterranean). The content of the organic and black carbon in atmospheric particles, which favors elevated concentrations of POPs in the aerosol phase, was relatively higher for these samples ($f_{OC}$ of 0.11 and $f_{BC}$ of 0.017), and within the range of previously described coastal/urban aerosols (Dachs and Eisenreich, 2000; Querol et al., 2009). The second set of samples differentiated by the PCA analysis included samples collected between Mallorca and Sardinia, with air masses associated to back trajectories indicating an Atlantic/North-African origin (ST1b, STIIb, T4 and T5). Saharan dust events occurred when these samples were collected (see Supplement E). Other studies have described high concentrations of POPs associated to dust events (Garrison et al., 2013).

Gas-particle partitioning is an important mechanism affecting POPs’ fate and transport within the atmosphere, since dominant atmospheric depositional processes and the relative contribution of different degradation mechanisms differ between the gas and aerosol phase compounds (Anderson and Hites, 1996; Mandalakis et al., 2003). The aerosol–gas partition coefficient ($K_P$) is given by:

$$K_P = \frac{C_{AP}}{C_G \cdot TSP} = \frac{f_{OC}}{\rho_{OCT}} \left( \frac{\gamma_{OCT} \cdot MW_{OCT}}{\gamma_{OC} \cdot MW_{OC}} \right) K_{OA}$$  \hspace{1cm} (4)$$

where TSP is the concentration of total suspended particles (µg m$^{-3}$), $f_{OC}$ is the fraction of organic carbon in the aerosol, $\gamma_{OCT}$ and $\gamma_{OC}$ are the activity coefficients of PCBs in octanol and the aerosol organic carbon, respectively, $MW_{OCT}$ (gmol$^{-1}$) and $MW_{OC}$ (gmol$^{-1}$) are the molecular weight of octanol and aerosol organic carbon, and $K_{OA}$ is the temperature corrected octanol-air partition coefficient. Figure 3 (and Supplement E) shows the correlation between $K_P$ and $K_{OA}$ assuming that the value of $\gamma_{OCT}MW_{OCT}/\gamma_{OC}MW_{OC}$ equals unity, an assumption discussed below.

There was a significant correlation ($p < 0.05$) between $K_P$ and $K_{OA}$ when considering HCB and all the measured PCBs, but it only explained 26% of the variability and the
slope was significantly lower than one (Supplement E). Nevertheless, if only HCB and tri- to penta-chlorinated PCB congeners were considered, the fitted least square regression line explained 75% of the variability and the slope was close to unity (Fig. 3). Conversely, congeners with a higher chlorinated degree did not show the same trend, since the $K_P$ values for these compounds did not further increase as the corresponding $K_{OA}$ increased. PCB congeners with six or more Cl fitted to a regression line with a slope significantly less than unity (Fig. 3).

For aerosol–gas partitioning close to equilibrium, the slope of the correlation log $K_P - \log K_{OA}$ is expected to have a value near unity. Therefore, a slope lower than unity in the relationship between $K_P$ and $K_{OA}$ could be explained by differential values in $\gamma_{OC} M_{OC} / \gamma_{MW} M_{MW}$ for the different congeners, which in turn, assuming the same value of $M_{OC}$, could be driven by the differential values of $\gamma_{OC}$ depending of the compound as suggested elsewhere (Goss et al., 1998). In addition to varying values of $\gamma_{OM}$ however, it is likely that there is a lack of aerosol–gas equilibrium for the higher chlorinated congeners, with aerosol phase concentrations lower than those in equilibrium. The kinetics of sorption is faster for the less chlorinated PCB congeners and HCB than for the more chlorinated congeners. Moreover, air–water exchange diffusion for the highly chlorinated congeners is also slower resulting in a potential disequilibrium driven by short-term increases in gas phase concentrations and/or decreases in aerosol concentrations. In consequence, for the sampling periods when there is a net volatilization of PCBs from underlying seawater, gas phase concentrations could be increased due to recent emissions of PCBs from the sea, which may not be yet equilibrated with the aerosol phase. Likewise, for the sampling periods where there is a net deposition of PCBs (higher fugacity in air than in water), emissions of sea spray aerosol that has not yet been equilibrated with the gas phase could also lead to shallow slopes.

### 3.3 Factors affecting the atmospheric occurrence of PCBs and HCB

A Principal Component Analysis (PCA) was carried out for congener groups (sum of the PCB congeners with the same degree of chlorination) in the gas phase (see
Supplement F). The two first principal components (PC1 and PC2), explaining 35% and 23% of the variability respectively, separate samples in a gradient of concentrations. Samples T4, T9, T17 and STI show the lowest $C_G$ for $\sum_{41}^\text{PCB}$, while T2, T22, T25, STV and T12 where those samples with the highest $C_G$.

For the sampling period of this study, the temperature influence on PCBs and HCB gas phase concentrations was evaluated using the Clausius–Clapeyron equation (see Supplement F). Nevertheless, unlike land-based studies, the results did not show any significant correlation between gas phase concentrations and temperature for any of the compounds, which is in agreement with other measurements in oceanic regions (Mandalakis et al., 2002; Lohmann et al., 2009). On one hand, the temperature range during the cruises was possibly too tight to highlight the temperature dependence on the partial pressure of the compounds. The seawater temperatures showed a narrow range of variability (20°C ±3). On the other hand, this temperature dependence is more marked in land where temperature dependent reemissions from other compartments such as soils are much stronger than in the atmosphere of the open Mediterranean, where air and water are from close to equilibrium to a net deposition. Indeed, after long range atmospheric transport, the temperature dependence of gas phase concentrations observed in source regions is lost due to dilution and depositional processes (Wania et al., 1998; Galbán-Malagón et al., 2013). Even though temperature alone does not influence significantly the gas phase concentrations, it contributes to the third principal component (Table S11 of Supplement F) of the statistical analysis explaining 15% of the variability.

In order to examine the influence of the origin of air masses on $C_G$, back-trajectories were evaluated for these two sub-sets of samples with extreme values of $C_G$. The Mediterranean region has a complicated atmospheric circulation pattern characterized by a strong northern component, which is particularly pronounced during the warm periods of the year. This is due to differential heating between the land of North-Africa (almost bare soil), the Mediterranean waters and the land of Southern Europe.
with vegetation cover (UNEP, 2002). This circulation pattern is reflected by the back-trajectories for the sample transects in this study (see Supplement B).

Figure 4 shows the air-mass back-trajectories corresponding to the samples with the lowest $C_G$ (lower panels in Fig. 4) and higher $C_G$ (upper panels of Fig. 4) for the $\Sigma_{41}$ PCB. The lowest $C_G$ values corresponded to sampling periods with predominant marine influenced air masses. Conversely, the highest concentrations were associated to land based air masses from the north (Europe). These trends can be explained as the result of the contribution of POPs’ primary and secondary sources, in addition to the depositional and degradative processes. On one hand, when air masses encounter land surfaces they potentially could receive OCls from the terrestrial reservoirs. Populated coastal regions and large industrial areas are possible sources of PCBs to the atmosphere (Garcia-Flor et al., 2009) where large reservoirs of these chemicals still remain. For example, soils are the major environmental reservoir of previously emitted and re-deposited POPs (Dalla Valle et al., 2005), and could be acting as a secondary source of OCI to the atmosphere (Jones et al., 1999). Volatilization of PCBs and HCB from soils is enhanced during warm periods and they could be an important source of OCls to the regional atmosphere (Cabreroiz et al., 2011a, b). Therefore, the higher PCB concentrations for the sampling periods dominated with air masses from Europe could be driven by these Southern European OCI sources. On the other hand, the lowest concentrations were associated with air masses of marine origin (Fig. 4). During atmospheric transport, a dilution process of OCI levels occurs as distance from the source increases. The strength of this dilution mechanism is modulated by depositional and degradative fluxes influencing air masses traveling over the sea and can be assessed calculating the atmospheric residence times for the different compounds.

### 3.3.1 Atmospheric residence times over Mediterranean Sea

The burden and fate of PCBs and HCB in the marine atmospheric environment can be affected by a number of physical and biogeochemical variables that in turn will be determining the atmospheric residence times of the compounds. The atmospheric resident

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times \( (R, d) \) were calculated following the approach proposed elsewhere (Jurado and Dachs, 2008):

\[
R = \frac{C_{\text{TOTAL}}}{F_{\text{OH}} + F_{\text{DD}} + F_{\text{WD}} + F_{\text{AWdep}} - F_{\text{AWvol}}} \cdot \text{AML} \tag{5}
\]

where \( C_{\text{TOTAL}} \) is the total concentration of POPs in the atmosphere (gas and aerosol phase), AML is the considered atmospheric mixed layer height, and \( F_{\text{OH}}, F_{\text{DD}}, F_{\text{WD}}, F_{\text{AWdep}} \) and \( F_{\text{AWvol}} \) are the atmospheric OH degradation flux, atmospheric dry deposition flux, atmospheric wet deposition flux, diffusive deposition flux and diffusive volatilization flux, respectively. The volatilization from the under-laying sea surface is the only input of OCls to the atmosphere of the sea regions away from the shore. Wet deposition process was in this case not considered due to lack of rainfall events during the sampling cruise. On average, \( R \) ranged from 1.2 days for less chlorinated compounds (PCB 18), to 3.9 days for more chlorinated congeners (PCB 187), whereas HCB atmospheric resident time was much higher, up to 53 days due to the extreme persistence of HCB in the atmosphere and close to equilibrium air–water conditions.

To evaluate the critical process affecting the magnitude of the residence times for the different compounds, Eq. (5) was parameterized as (see Supplement F for details):

\[
R = \left[ \frac{1}{r_{\text{OH}} \cdot \text{AML}(1 - \varnothing)} + \frac{1}{v_{\text{DD}} \cdot \varnothing} + \frac{1}{k_{\text{AW}} \cdot \frac{(1 - \varnothing)}{H'}} - \frac{1}{(k_{\text{AW}} - k_{\text{ADW}}) \cdot \frac{(1 - \varnothing)}{H'}} \right] \cdot \text{AML} \tag{6}
\]

where \( \varnothing \) is the fraction of OCI bound to the aerosols in the atmosphere, \( r_{\text{OH}} \) is the compound specific OH radical degradation decay rate \( (\text{d}^{-1}) \), and \( k_{\text{ADW}} \) is the air-deep water mass transfer coefficient estimated as reported elsewhere (Dachs et al., 2002).

The latter is strongly dependent on the magnitude of the biological pump removing PCBs from surface waters and the lower atmosphere. Below we discuss each of these removing processes of PCBs and HCB from the lower atmosphere.
3.3.2 Atmospheric degradation due to OH radicals

OCl degradation in the atmosphere is dominated by the reaction with hydroxyl radicals and occurs mainly in the gas-phase (Anderson and Hites, 1996; Jurado and Dachs, 2008). The degradation decay rate can be calculated as follows:

\[ r_{OH} = k_{OH} \cdot C_{OH} \]  \hspace{1cm} (7)

where \( C_{OH} \) (molecules cm\(^{-3}\)) is the atmospheric concentration of OH radicals and \( k_{OH} \) (cm\(^3\) molecules\(^{-1}\) s\(^{-1}\)) is the degradation rate constant. Gas phase PCB congeners containing five and six chlorine atoms showed the highest relative abundance in the gas phase, accounting for 50% and 60% of \( \sum_{41} \) PCB in the Mediterranean and Black Seas atmosphere, respectively. This contrasts with other studies that have reported higher abundance of dichlorinated and trichlorinated congeners in oceanic atmospheres (Jaward et al., 2004; Gioia et al., 2008). The described decrease in the relative abundance of the less chlorinated compounds could be explained by PCB depletion due to OH radicals present in the atmosphere, a process that is relatively faster for the less chlorinated congeners (Anderson and Hites, 1996) as indicated by the corresponding degradation rate constant. Mandalakis and coworkers (Mandalakis et al., 2002) have already suggested this process to explain the compound profile found in a Eastern Mediterranean atmosphere, where a relative decrease of dichlorinated and trichlorinated congeners was observed in comparison to tetrachlorinated and pentachlorinated congeners during the summer period and periods of high OH radical concentrations (Mandalakis et al., 2002, 2003). HCB degradation rate constant on the contrary are 1 order of magnitude lower than for PCB, thus resulting in longer \( R \) values.

3.3.3 Atmospheric deposition of PCBs and HCB

Figure 5a shows the dry deposition fluxes for the different Mediterranean sub-basins. Due to the assumption of a unique value of \( v_D \) for all sampling events, the spatial pattern
of $F_{DD}$ equals that of the aerosol phase concentrations (Fig. 2), with higher $F_{DD}$ values for the Western Mediterranean Basin. As commented above, higher aerosol phase concentrations of PCBs were also found associated to dust events. The association of PCBs and other POPs to dust aerosols has been poorly characterized in term of their deposition fluxes, but large dust particles may deposit with $v_D$ values much higher than those assumed here. Therefore, it is possible that we are underestimating the dry deposition fluxes associated to dust driven deposition by one order of magnitude, and thus overestimating the atmospheric resident times for periods with Saharan dust events.

Figure S15 shows the box-plot of $f_G/f_W$ for each PCB congener. Most PCB congeners are found very close to dynamic air–water equilibrium, with a predominance of net diffusive deposition over volatilization episodes. Figure 5 shows the air–water diffusive fluxes (Fig. 5b) and their comparison with the dry deposition fluxes, evidencing the greater importance of diffusive exchanges due to the fact that PCBs and HCB are mostly found in the gas phase. In fact, the amount of PCBs and HCB associated with particles in our study was very low and on average accounted for the 6% of the total atmospheric concentration. This is in agreement with previous studies performed in the same region (Mandalakis et al., 2002; Garcia Flor et al., 2009) and in other oceanic regions (Galbán-Malagón et al., 2012). Nevertheless, while the eastern basin presents a clear predominance of gas phase PCBs (91–99.6%), the variability in the western basin is higher due to higher aerosol loads for some sampling periods, with a maximum of particle bound PCBs of 24% of the total atmospheric PCBs.

### 3.3.4 Biogeochemical characteristic of the underlying water column

Water column biogeochemistry plays a critical role in controlling the dissolved phase concentration, and thus the air–water gradient of POP concentrations (Dachs et al., 2002). Previous studies have already assessed the role of the biological pump as one of the key mechanisms governing the sink of hydrophobic and persistent pollutants, such as PCBs or polybrominated diphenyl ethers (PBDEs) in the oceanic (Dachs et al., 2002).
2002; Jaward et al., 2004; Galbán-Malagón et al., 2012) and lacustrine environments (Dachs et al., 1999; Nizzetto et al., 2012). In fact, air–water exchange and partitioning to/uptake by planktonic organisms (particularly phytoplankton) and other pools of particulate organic matter could act as coupled processes in aquatic environments (Dachs et al., 2002). Once atmospheric POPs enter the dissolved phase through air–water diffusion exchange, they partition to particles and/or planktonic organisms and they can be removed from the surface waters and delivered to the deep ocean by sinking particles and/or zooplanktonic vertical migrations. Driven by these losses, air–water dis-equilibria can be enhanced, which is particularly important for hydrophobic chemicals and in situations with higher amounts of biomass. A companion study (Berrojalbiz et al., 2011) showed field evidence of the relevance of these processes as a driver of PCB water column concentrations in the Mediterranean Sea.

The described scenario is consistent with the results found here, where for the time periods with the higher plankton biomass, the gas phase concentrations were at the lower end of those observed (Fig. S16). However, most of the sampling periods correspond to areas with biomass within a very narrow range and low biomass values that does not allow for further assessment of this interaction between water column biogeochemistry and the atmospheric occurrence of POPs in the Mediterranean. Unlike other oceanic regions, the Mediterranean Sea is oligotrophic (Regaudie-de-Gioux et al., 2009), and thus the biological pump has a much lower potential to deplete the dissolved concentrations in surface waters and drive air and water out of equilibrium as has been observed for the North Atlantic and Southern Ocean (Galbán-Malagón et al., 2012, 2013). Conversely, close to air–water equilibrium conditions for PCBs have also been described for the oligotrophic Atlantic and Pacific oceans (Gioia et al., 2008; Zhang et al., 2010), consistent with the results reported here.
3.3.5 Seasonal changes on the relative contribution of processes controlling OCI atmospheric burden

Among the different loss processes affecting the atmospheric residence times of PCBs, in this study performed during summer months there is a clear dominance of the influence of OH radical degradation in the atmosphere over the rest of the processes. Figure 6 shows the integrated atmospheric loss fluxes (kg month$^{-1}$) of PCBs and HCB due to reaction with OH radicals and atmospheric deposition. The oligotrophic characteristics of the Mediterranean surface waters do not pose importance to the contribution of the biological pump that could potentially alter air–water diffusive exchange toward a net deposition flux, thus atmospheric deposition during the summer is less important as a depletion mechanism for atmospheric concentrations of PCBs than degradation due to reaction with OH radicals. Nevertheless, during the spring season, the Mediterranean Sea is subjected to important planktonic blooms (Turley et al., 2000) with significantly higher planktonic biomass where it is likely to produce an increase in the relative importance of the biological pump on the decrease of the atmospheric resident times of POPs. Likewise, wet deposition processes not measured in this work should also be considered to be an important variable in more rainy seasons of the year and for some Mediterranean sub-basins not sampled during the thresholds cruises.

The thermic variations along the year may also affect the contributions of different processes to atmospheric residence times. Atmospheric concentrations of semi-volatile organic compounds have been shown to be strongly dependent on ambient temperature due to its influence on environmental partitioning (Wania et al., 1998; Lee et al., 1998; Gouin et al., 2002). Therefore, it is likely that changes in the fractionation of OCIs between gas and aerosol phase could lead to an increase in the dry deposition, while changes in $H'$ values may favor POP air–water diffusion towards a net deposition flux. Moreover, OH radical degradation is strongly dependent on the temperature and radiation (Anderson and Hites, 1996; Mandalakis et al., 2002) and thus, it is expected
to decrease in great extent during winter season (OH radical concentration is 5 to 10 times lower than in summer periods) (Spivakovsky et al., 2000).

4 Conclusions

The present study has generated an extensive and unique data set of gas and aerosol phase concentrations of OCls in the Mediterranean atmosphere. Concentrations are lower than those reported over the adjacent continents, but there is a large variability in concentrations. Higher concentrations were observed for periods with air masses from the northern basin, while the minimum concentrations were observed when the air masses had a marine origin. Dry deposition, air–water diffusive fluxes and degradation with OH radical fluxes have been estimated. There is a close to air–water equilibrium for most periods and PCB congeners, even though there is a net deposition in some periods. Degradation with OH radicals is the major loss process of PCBs during the summer months. Conversely, the atmospheric residence times of HCB are long due to a lack of effective atmospheric degradation and deposition.

Supplementary material related to this article is available online at http://www.atmos-chem-phys-discuss.net/14/9747/2014/acpd-14-9747-2014-supplement.pdf.

Acknowledgements. This work was funded by the European Union and Spanish Ministry of Science and Innovation through the THRESHOLDS projects. N. Berrojalbiz acknowledges a pre-doctoral fellowship from the Basque Government.
References


Wania, F. and Daly, G. L.: Estimating the contribution of degradation in air and deposition to the deep sea to the global loss of PCBs, Atmos. Environ., 36, 5581–5593, 2002.


Table 1. Summary of the PCB and HCB concentrations in the Mediterranean atmosphere.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Gas phase (pgL⁻¹)</th>
<th>Aerosol samples (ngg⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mediterranean Sea (N = 30)</td>
<td>Black Sea and Marmara Sea (N = 4)</td>
</tr>
<tr>
<td></td>
<td>Median (Mean)</td>
<td>Range</td>
</tr>
<tr>
<td>HCB</td>
<td>28.1 (59.2) (nd–418.3)</td>
<td>42.0 (49.5) (13.7–100.2)</td>
</tr>
<tr>
<td>PCB 17</td>
<td>4.0 (7.2) (nd–29.4)</td>
<td>10.3 (22.7) (0.1–70.1)</td>
</tr>
<tr>
<td>PCB 18</td>
<td>9.6 (21.6) (nd–170.3)</td>
<td>11.9 (16.9) (nd–43.9)</td>
</tr>
<tr>
<td>PCB 28</td>
<td>3.1 (9.9) (nd–72.8)</td>
<td>18.4 (24.2) (6.0–54.0)</td>
</tr>
<tr>
<td>PCB 31</td>
<td>nd</td>
<td>nd</td>
</tr>
<tr>
<td>PCB 33</td>
<td>10.7 (16.8) (nd–52.1)</td>
<td>5.6 (22.9) (nd–80.4)</td>
</tr>
<tr>
<td>PCB 44</td>
<td>nd</td>
<td>nd</td>
</tr>
<tr>
<td>PCB 49</td>
<td>6.7 (11.3) (nd–47.7)</td>
<td>7.1 (12.6) (nd–36.2)</td>
</tr>
<tr>
<td>PCB 52</td>
<td>12.4 (19.2) (nd–101.0)</td>
<td>nd (4.8)</td>
</tr>
<tr>
<td>PCB 70</td>
<td>0.7 (4.0) (nd–16.4)</td>
<td>2.2 (6.6) (nd–22.2)</td>
</tr>
<tr>
<td>PCB 74</td>
<td>6.0 (8.4) (nd–62.1)</td>
<td>5.1 (7.9) (nd–27.1)</td>
</tr>
<tr>
<td>PCB 82</td>
<td>5.8 (11.8) (nd–74.3)</td>
<td>15.6 (37.1) (8.1–109.3)</td>
</tr>
<tr>
<td>PCB 87</td>
<td>8.6 (14.4) (nd–74.3)</td>
<td>15.6 (37.1) (8.1–109.3)</td>
</tr>
<tr>
<td>PCB 95</td>
<td>nd (6.9) (nd–85.3)</td>
<td>nd (1.8) (nd–72)</td>
</tr>
<tr>
<td>PCB 99/101</td>
<td>8.0 (18.4) (nd–123.1)</td>
<td>13.3 (19.4) (12.6–32.2)</td>
</tr>
<tr>
<td>PCB 105/132</td>
<td>4.3 (12.5) (nd–105.5)</td>
<td>9.5 (8.3) (nd–14.4)</td>
</tr>
<tr>
<td>PCB 110</td>
<td>3.5 (5.6) (nd–23.4)</td>
<td>10.5 (10.0) (5.8–16.7)</td>
</tr>
<tr>
<td>PCB 118</td>
<td>4.3 (5.8) (nd–17.8)</td>
<td>4.7 (4.2) (nd–7.3)</td>
</tr>
<tr>
<td>PCB 128</td>
<td>nd (1.01) (nd–6.1)</td>
<td>4.5 (4.0) (nd–6.9)</td>
</tr>
<tr>
<td>PCB 138</td>
<td>14.9 (17.3) (1.2–56.0)</td>
<td>22.6 (26.9) (20.0–42.2)</td>
</tr>
<tr>
<td>PCB 149</td>
<td>15.3 (20.4) (nd–98.9)</td>
<td>23.5 (30.5) (17.5–56.8)</td>
</tr>
<tr>
<td>PCB 151</td>
<td>0.3 (4.2) (nd–44.6)</td>
<td>2.3 (2.8) (nd–6.6)</td>
</tr>
<tr>
<td>PCB 153</td>
<td>19.3 (20.1) (nd–68.4)</td>
<td>30.9 (32.8) (24.0–45.4)</td>
</tr>
<tr>
<td>PCB 156/171</td>
<td>nd (1.0)</td>
<td>nd</td>
</tr>
<tr>
<td>PCB 158</td>
<td>nd (0.5) (nd–4.5)</td>
<td>2.0 (1.8) (nd–3.3)</td>
</tr>
<tr>
<td>PCB 169</td>
<td>nd (0.7)</td>
<td>nd</td>
</tr>
<tr>
<td>PCB 170</td>
<td>6.1 (6.8) (nd–17.8)</td>
<td>13.0 (12.1) (7.9–14.4)</td>
</tr>
<tr>
<td>PCB177</td>
<td>4.7 (5.9) (nd–15.0)</td>
<td>11.0 (10.2) (6.2–12.3)</td>
</tr>
<tr>
<td>PCB 180</td>
<td>16.9 (18.6) (nd–51.3)</td>
<td>32.8 (32.2) (21.0–42.1)</td>
</tr>
<tr>
<td>PCB 183</td>
<td>5.2 (5.8) (nd–23.7)</td>
<td>7.1 (7.1) (4.9–9.5)</td>
</tr>
<tr>
<td>PCB 187</td>
<td>10.6 (11.7) (3.4–50.3)</td>
<td>15.6 (13.2) (nd–21.54)</td>
</tr>
<tr>
<td>PCB 191</td>
<td>1.6 (2.3) (nd–12.6)</td>
<td>4.4 (4.4) (2.8–5.8)</td>
</tr>
<tr>
<td>PCB 194</td>
<td>2.3 (2.4) (nd–7.6)</td>
<td>4.0 (4.2) (3.7–4.9)</td>
</tr>
<tr>
<td>PCB 195</td>
<td>nd</td>
<td>nd</td>
</tr>
<tr>
<td>PCB 199(199)</td>
<td>4.9 (4.3) (nd–10.35)</td>
<td>8.2 (8.2) (6.2–10.4)</td>
</tr>
<tr>
<td>PCB 205</td>
<td>nd (0.1) (nd–1.7)</td>
<td>nd</td>
</tr>
<tr>
<td>PCB 206</td>
<td>nd</td>
<td>nd</td>
</tr>
<tr>
<td>PCB 208</td>
<td>nd (0.1) (nd–2.9)</td>
<td>nd</td>
</tr>
<tr>
<td>PCB 209</td>
<td>nd</td>
<td>nd</td>
</tr>
</tbody>
</table>

∑7PCBs (pgL⁻¹) = 88.5 (96.4) (27.7–259.4) 136.9 (134.7) (98.7–166.1) 46.2 (66.1) (107.7–324.8) 56.4 (58.7) (41.3–80.9)

∑41PCBs (pgL⁻¹) = 237.6 (297.3) (82.0–931.6) 417.9 (421.5) (206.2–644.1) 92.9 (140.7) (21.4–653.8) 80.4 (91.7) (70.5–135.6)

nd (not detected).
Table 2. Comparison of the gas phase concentrations of the 7 ICES PCB congeners in different locations of Mediterranean and Black Sea (mean ± SD) obtained in the present study with those reported previously.

<table>
<thead>
<tr>
<th>PCB</th>
<th>Atlantic Ocean</th>
<th>Western Mediterranean Sea</th>
<th>Eastern Mediterranean Sea</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Jaward et al. (2004)</td>
<td>Gioia et al. (2008)</td>
<td>This study</td>
</tr>
<tr>
<td></td>
<td>Sample size: (N) = 49</td>
<td>(N) = 2</td>
<td>(N) = 3</td>
</tr>
<tr>
<td>PCB 28</td>
<td>12 ± 10</td>
<td>8.45 ± 7.</td>
<td>4.3 ± 5.4</td>
</tr>
<tr>
<td>PCB 52</td>
<td>6 ± 5</td>
<td>3.05 ± 1.8</td>
<td>5.2 ± 4.5</td>
</tr>
<tr>
<td>PCB 101</td>
<td>1.4 ± 4</td>
<td>1 ± 0.3</td>
<td>9.0 ± 12.1</td>
</tr>
<tr>
<td>PCB 118</td>
<td>1.5 ± 1.5</td>
<td>2.1 ± 0.4</td>
<td>4.5 ± 0.4</td>
</tr>
<tr>
<td>PCB 138</td>
<td>2.9 ± 2.3</td>
<td>1.95 ± 0.6</td>
<td>5.4 ± 3.6</td>
</tr>
<tr>
<td>PCB 153</td>
<td>6.1 ± 5</td>
<td>0.6 ± 0.1</td>
<td>5.6 ± 5.</td>
</tr>
<tr>
<td>PCB 180</td>
<td>1.2 ± 0.87</td>
<td>3.15 ± 1.2</td>
<td>2.3 ± 4.1</td>
</tr>
</tbody>
</table>
Table 3. Mono-ortho substituted PCB congeners in gas and aerosol phase in pg m\(^{-3}\).

<table>
<thead>
<tr>
<th>PCB</th>
<th>Gas phase ((N = 18))</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th>Aerosol phase ((N = 18))</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Average</td>
<td>Median</td>
<td>Min</td>
<td>Max</td>
<td>Average</td>
<td>Median</td>
<td>Min</td>
<td>Max</td>
<td>Average</td>
<td>Median</td>
</tr>
<tr>
<td>105</td>
<td>0.590</td>
<td>0.576</td>
<td>0.348</td>
<td>1.183</td>
<td>0.080</td>
<td>0.056</td>
<td>0.011</td>
<td>0.303</td>
<td>0.075</td>
<td>0.067</td>
</tr>
<tr>
<td>114</td>
<td>0.075</td>
<td>0.067</td>
<td>0.041</td>
<td>0.142</td>
<td>0.009</td>
<td>0.009</td>
<td>0.001</td>
<td>0.021</td>
<td>0.004</td>
<td>0.002</td>
</tr>
<tr>
<td>118</td>
<td>1.530</td>
<td>1.468</td>
<td>0.938</td>
<td>3.030</td>
<td>0.204</td>
<td>0.133</td>
<td>0.029</td>
<td>0.825</td>
<td>0.004</td>
<td>0.002</td>
</tr>
<tr>
<td>123</td>
<td>0.030</td>
<td>0.029</td>
<td>0.015</td>
<td>0.056</td>
<td>0.004</td>
<td>0.002</td>
<td>0.001</td>
<td>0.017</td>
<td>0.004</td>
<td>0.002</td>
</tr>
<tr>
<td>156</td>
<td>0.291</td>
<td>0.300</td>
<td>0.153</td>
<td>0.452</td>
<td>0.075</td>
<td>0.049</td>
<td>0.011</td>
<td>0.184</td>
<td>0.009</td>
<td>0.007</td>
</tr>
<tr>
<td>157</td>
<td>0.033</td>
<td>0.030</td>
<td>0.020</td>
<td>0.064</td>
<td>0.009</td>
<td>0.007</td>
<td>0.002</td>
<td>0.025</td>
<td>0.003</td>
<td>0.022</td>
</tr>
<tr>
<td>167</td>
<td>0.144</td>
<td>0.144</td>
<td>0.075</td>
<td>0.217</td>
<td>0.033</td>
<td>0.022</td>
<td>0.005</td>
<td>0.093</td>
<td>0.010</td>
<td>0.008</td>
</tr>
<tr>
<td>189</td>
<td>0.031</td>
<td>0.032</td>
<td>0.016</td>
<td>0.046</td>
<td>0.010</td>
<td>0.008</td>
<td>0.002</td>
<td>0.029</td>
<td>0.010</td>
<td>0.008</td>
</tr>
</tbody>
</table>
Fig. 1. Location of the atmospheric samples and spatial distribution of gas phase concentrations of PCBs (pg m\(^{-3}\)).
Fig. 2. Spatial distribution of PCBs concentrations in the aerosol phase (pg m\(^{-3}\) and pg m\(^{-3}\)).
Fig. 3. Aerosol-particle partitioning of PCBs and HCB. Log–log relationship between aerosol–gas partition coefficient ($K_p$) and octanol-air partitioning coefficient ($K_{OA}$) for PCBs and HCB. $K_{OA}$ values were corrected for temperature.
Fig. 4. Air mass back-trajectories (48 h) of selected samples at two different heights (15 m in red and 100 m in blue) obtained with the HYSPLIT model. Upper panels' back-trajectories correspond to samples with the higher PCB concentrations (T2, T19, T22, T25, STIVa, STIVb and STIVc) whereas the lower panels correspond to air masses with lower \( C_G \) (T4, T9 and T17).
Fig. 5. Atmospheric deposition of PAHs: (a) dry deposition fluxes for the different Mediterranean Basins; (b) diffusive air–water exchange for the different Mediterranean Basin.
Fig. 6. Comparison of losses of atmospheric PCBs and HCB due to atmospheric degradation and reaction with OH radicals during the summer. Fluxes are given as integrated for the whole Mediterranean Basin.