Hydrogen peroxide in the marine boundary layer over the southern Atlantic during the OOMPH cruise in March 2007

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Abstract

In the OOMPH (Ocean Organics Modifying Particles in both Hemispheres) project a ship measurement cruise took place in the late austral summer from 1 to 23 March 2007. The French R/V Marion Dufresne sailed from Punta Arenas, Chile (70.85° W, 53.12° S) to La Reunion island (55.36° E, 21.06° S) across the southern Atlantic Ocean. In-situ measurements of hydrogen peroxide, methylhydroperoxide and ozone were performed and are compared to simulations with the atmospheric chemistry global circulation model EMAC. The model generally reproduces the measured trace gas levels, but underestimates hydrogen peroxide mixing ratios at high wind speeds, indicating too strong dry deposition to the ocean surface. An interesting feature during the cruise is a strong increase of hydrogen peroxide, methylhydroperoxide and ozone shortly after midnight off the west coast of Africa due to an increase in the boundary layer height, leading to downward transport from the free troposphere, which is realistically reproduced by the model.

1 Introduction

The oxidizing power of the lower atmosphere in the gas phase is defined by the concentrations of the hydroxyl radical (OH), ozone (O₃), the nitrate radical (NO₃), halogen radicals (e.g. ClO, BrO, IO) and hydrogen peroxide (H₂O₂), an important oxidizer in the liquid phase (Thompson, 1992). The dominant oxidizing agent is OH, whose primary source is the photolysis of O₃ and subsequent reaction of the formed O¹D-atom with water vapor (Levy, 1971). The major sinks of OH are reactions with carbon monoxide (CO), methane (CH₄), and volatile organic compounds (VOC) yielding peroxy radicals (HO₂ and RO₂). The fate of these peroxy radicals strongly depends on the concentrations of nitrogen oxides (NOₓ, NO plus NO₂). In semi-polluted and polluted regions with NOₓ levels in excess of several tens of pptv, the peroxy radicals predominantly react with NO, yielding NO₂ and recycling OH. In these environments the subsequent photol-
ysis of NO\textsubscript{2} yields ozone, with NO\textsubscript{x} acting as a catalyst. In low NO\textsubscript{x} environments, such as the marine boundary layer, the peroxy radicals undergo self-reactions (HO\textsubscript{2} + HO\textsubscript{2} and RO\textsubscript{2} + HO\textsubscript{2}) yielding H\textsubscript{2}O\textsubscript{2} and organic peroxides (e.g. CH\textsubscript{3}OOH from methane oxidation), and also destroying ozone (HO\textsubscript{2} + O\textsubscript{3} and OH + O\textsubscript{3}). The peroxides serve as reservoir species for the HO\textsubscript{x} (OH plus HO\textsubscript{2}) radicals, which can be recycled by photolysis or reaction with OH. Hydrogen peroxide is also an important oxidizing agent in the liquid phase, notably of sulfur dioxide. Since many peroxides are water soluble, physical removal processes (deposition to surfaces and washout in rain events) strongly influence the oxidizing power of the lower atmosphere.

In the marine boundary layer at low NO\textsubscript{x} concentrations the concentrations of H\textsubscript{2}O\textsubscript{2}, ROOH and O\textsubscript{3} are strongly coupled, since their formation and destruction compete for the HO\textsubscript{x} radicals. In order to model oxidation processes in this environment, formation and destruction of peroxides have to be accurately described, including the physical removal processes.

Previous measurements of peroxides (H\textsubscript{2}O\textsubscript{2} and ROOH) in the marine boundary layer in the 1980s and 1990s have been summarized in the review article by Lee et al. (2000). Since this review additional observations in the marine boundary layer have been reported in the literature (Junkermann and Stockwell, 1999; Weller et al., 2000; Kieber et al., 2001; O’Sullivan et al., 2004; Chang et al., 2004; Stickler et al., 2007). These observations indicate highest mixing ratios (> 500 pptv) of H\textsubscript{2}O\textsubscript{2} in the tropics (Slemr and Tremmel, 1994; Heikes et al., 1996; O’Sullivan et al., 1999, 2004; Junkermann and Stockwell, 1999; Weller et al., 2000) and decreasing concentrations toward higher latitudes in both hemispheres, reaching 200–300 pptv south of 40° in the Southern Hemisphere (Slemr and Tremmel, 1994; O’Sullivan et al., 1999, 2004; Junkermann and Stockwell, 1999; Weller et al., 2000). In general mixing ratios are about a factor of two higher in the Northern Hemisphere than at corresponding latitudes in the south (O’Sullivan et al., 1999). A significant dissimilarity between the different ocean basins has not been observed, while higher H\textsubscript{2}O\textsubscript{2} mixing ratios have been observed in continental outflow (e.g. Heikes et al., 1996).
The mixing ratios of the most abundant organic peroxide CH$_3$OOH show similar behavior as H$_2$O$_2$ in the marine boundary layer, with highest levels in the tropics and decreasing towards the poles. Also the absolute mixing ratios are comparable, yielding H$_2$O$_2$/CH$_3$OOH ratios close to 1 in air masses not affected by recent rainout (Lee et al., 2000).

Here we describe in-situ ship-based observations of O$_3$, H$_2$O$_2$ and a proxy for CH$_3$OOH in the marine boundary layer of the southern Atlantic Ocean in the austral late summer of 2007. These measurements are compared to the atmospheric chemistry global circulation model EMAC (Jöckel et al., 2006, 2010). Section 2 describes the methods (measurement principles and model) used, while the observations and model comparisons are described and discussed in Sect. 3. The final section summarizes the findings of this study.

2 Methods

2.1 OOMPH cruise MD160

As part of the OOMPH (Ocean Organics Modifying Particles in both Hemispheres) project a measurement cruise took place in the late austral summer from 1 to 23 March 2007. The French R/V Marion Dufresne sailed from Punta Arenas, Chile (70.85° W, 53.12° S) to La Reunion island (55.36° E, 21.06° S) crossing the southern Atlantic between the east coast of South America to the southern Indian Ocean east of South Africa between 20° W, 60° S and 35° E, 35° S (Fig. 1). During the first part of the cruise at high southern latitudes, cold air was encountered from the Antarctic continent. During this part of the cruise, cloud cover was extensive. Further north, temperatures increased together with solar radiation intensity and photolysis frequencies. The wind was generally from the west, with wind speeds varying between calm conditions and gale force winds up to 33 ms$^{-1}$. The average wind varied between more than 10 ms$^{-1}$.
during the first part and 8 ms$^{-1}$ during the second part of the campaign (Fig. 2). Details of the cruise can be found in Williams et al. (2010) and Hosaynali Beygi et al. (2011).

2.2 Trace gas measurements

Data used in this study were obtained by two in-situ instruments mounted in a temperature controlled container placed on the foredeck of the ship (see Fig. 3 in Hosaynali Beygi et al., 2011). Air was sampled from the top of an atmospheric mast (10 m above the deck, 20–25 m above the sea surface) through 17.1 m 1/2 Teflon tubes, shielded from sunlight by a black cover. The inlet was designed as a bypass with a total flow of 24 slm (retention time 3.4 s) sustained by a membrane pump. From the bypass inlet small flows were directed to the in-situ instruments inside the container via short 1/4 Teflon lines.

Hydrogen peroxide ($\text{H}_2\text{O}_2$) was measured with a commercial analyzer (AL2001 CA, Aero Laser, Garmisch Partenkirchen, Germany) based on wet chemical dual enzyme detection scheme described by Lazarus et al. (1985, 1986). Gaseous peroxides are sampled in a buffered (potassium hydrogen phthalate/NaOH) sampling solution (pH 5.8) in a glass stripping coil at a flow of 3 slm. The sampling efficiency for $\text{H}_2\text{O}_2$ was determined several times in the field and was always higher than 0.8. After passing the sampling coil the degassed liquid peroxide solution is divided into two channels and subsequently reacts with p-hydroxyphenyl acetic acid (POPHA) and horseradish peroxidase. The reaction with hydrogen peroxide, organic hydroperoxides and organic peroxides yields a fluorescent dye (6,6$'$-dihydroxy-3,3$'$-biphenyldiacetic acid) in stoichiometric quantities that is subsequently detected via fluorescence spectroscopy at 400–420 nm after excitation at 326 nm with a Cd ray lamp in a detection cell. Since the detection scheme is unspecific, the $\text{H}_2\text{O}_2$ concentration is determined from the difference of the two channels, with channel A measuring all peroxides (ROOH), while channel B measures ROOH – $\text{H}_2\text{O}_2$ after selective destruction of $\text{H}_2\text{O}_2$ via addition of catalase (efficiency > 95 % as determined in the field) prior to the reaction with POPHA.
Thus the difference between both channels provides the $\text{H}_2\text{O}_2$ concentration, while channel B provides an unspecific measurement of all organic hydroperoxides and organic peroxides. Nevertheless, assuming that methylhydroperoxide (MHP; $\text{CH}_3\text{OOH}$) is the most abundant organic peroxide in the remote marine boundary layer, as shown by previous measurements (e.g. Heikes et al., 1996), we obtain an upper limit assuming that ROOH consists of MHP only. The organic peroxide data is subsequently corrected for the lower sampling efficiency of $\text{CH}_3\text{OOH}$ compared to $\text{H}_2\text{O}_2$. The time resolution (10–90%) of the instrument is 30 s.

The in-field calibration of the instrument involves regular zero gas measurements (scrubbed ambient air after passage through cartridges filled with silica gel and hopcalite (Infiltec, Speyer, Germany)), liquid calibrations (liquid $\text{H}_2\text{O}_2$ standard of 35.5 µg L$^{-1}$) and gas phase calibrations with a $\text{H}_2\text{O}_2$ permeation tube (30% $\text{H}_2\text{O}_2$ in a glass flask temperature controlled to 40°C providing a calibration gas concentration of 6.38 ppbv). The detection limit of the instrument was determined from the 1σ variability of the in-field zero measurements performed every 2.5 h, estimated at 25 pptv. The total uncertainty determined from the precision (1σ variability of 9 in-field gas phase and liquid calibrations), the uncertainty of the standard, the inlet transmission and an ozone interference correction was about 12–13%. During the campaign the inlet transmission was determined twice by adding the gas phase standard at the top of the inlet line. Comparison of two calibrations directly in front of the analyzer yielded a transmission that decreased from 67% at the beginning of the campaign to 57% towards the end. The instrument has also been used (in combination with a constant pressure inlet) for airborne measurement of $\text{H}_2\text{O}_2$ in the free troposphere over the rainforest in South America (Stickler et al., 2007) and over Europe (Klippel et al., 2011).

The instrument used to measure ozone (together with NO and NO$_2$) is a high resolution (1 s) and highly sensitive 3-channel chemiluminescence detector (CLD, ECO-Physics CLD 790 SR, Duernten, Switzerland). The instrument and its performance characteristics during this campaign have been described in detail in a previous publication on the $\text{NO}_x$/$\text{O}_3$ photostationary state by Hosaynali Beygi et al. (2011). The total
uncertainty for the O$_3$ channel was determined from the 2σ deviation of the in-field calibrations (ozone calibrator model TE49C, Thermo Instruments, Germany) and the accuracy of the standard, estimated at 1% (Hosaynali Beygi et al., 2011).

2.3 Atmospheric chemistry model EMAC

Simulations of trace gas mixing ratios along the ship cruise were performed using the EMAC (ECHAM/MESSy Atmospheric Chemistry) global circulation model (Jöckel et al., 2010). EMAC uses the Modular Earth Submodel System (MESSy; Jöckel et al., 2005) to link multi-institutional sub models describing atmospheric processes interacting with oceans, land and human influences. For this study EMAC was applied in the T42L90MA-resolution (2.8° × 2.8° resolution in latitude and longitude, 90 vertical levels up to 0.01 hPa), using results from the lowest model level (~ 30 m) for comparison with measurements. The model was sampled (spatial bilinearly interpolated) along the ship track at every time step (i.e. 12 min) using the SD4 submodel (Jöckel et al., 2010), without any temporal interpolation. The meteorology was nudged to the operational ECMWF analysis. Tropospheric gas-phase and heterogeneous chemistry was calculated with the sub-model MECCA (Module Efficiently Calculating the Chemistry of the Atmosphere; Sander et al., 2005), aqueous-phase chemistry in cloud droplets and wet scavenging with the sub-model SCAV (Tost et al., 2006) and primary emissions and dry deposition of trace gases and aerosols with the sub-models ONLEM, OFFLEM, TNUDGE and DRYDEP (Kerkweg et al., 2006a, b). Previous results of a model comparison with airborne H$_2$O$_2$ measurements in the free troposphere have been discussed in Klippel et al. (2011).
3 Results and discussion

3.1 Data processing

For the present analysis the original observations were averaged over 12 min time intervals to be coherent with the model output time stepping. The time scale used is UTC, thus leading to deviations from solar noon of −80 min for the most westerly point (20° W) of the ship track to 140 min for the most easterly (35° E). Unfortunately due to the prevailing westerly winds the instruments often measured air polluted by the ship exhausts, notably when the wind was from the sector between 55 and 275°. This stack air contained very high concentrations of NO (up to 200 ppbv in individual plumes), leading to complete titration of O₃, while H₂O₂ was not affected at all on the short time scales involved, as has been observed previously (Weller et al., 2000). In total 53% of the observations were effected by stack emissions. Therefore we used Oₓ (O₃ + NO₂) for the experimental data, to deduce the original O₃ concentration that would have occurred without NO-titration to NO₂. In general, the difference between Oₓ and O₃ is marginal due to the very low NOₓ mixing ratios of less than 20 pptv in the remote marine boundary layer over the southern Atlantic and less than 200 pptv east of South Africa (Hosaynali Beygi et al., 2011). Thus the error in O₃ using Oₓ in all cases is smaller than 2%. From the model only the original O₃ data are used.

3.2 Distribution of measured and simulated trace gases

Figure 3 shows the time series for measured and simulated ozone mixing ratios. Observed O₃ varies between 17 and 25 ppbv, with lower values in the south-western Atlantic and higher values closer to Africa. The model qualitatively reproduces this gradient with a slight tendency to underestimate O₃ mixing ratios during the first part of the cruise (10–14 March) and a slight overestimation between 15 and 17 March. During the last part of the cruise after 17 March both model and observations agree quite well. Overall, the mean observed and simulated O₃ are 20.3 ± 1.8 ppbv and 19.7 ± 2 ppbv, re-
spectively. A least-square fit between simulated (y axis) and observed (x axis) ozone yields a slope of 0.98 ± 0.01 and an offset of −0.43 ± 0.21 ppbv. The regression coefficient $R^2$ is rather low (0.2). Taking into account that observed O₃ measurements are affected by background NO₂, the agreement between observations and model results is quite satisfactorily. Observations and model results are in good agreement with observations by Helmig et al. (2012) during the GasEx 2008 cruise in the southern Atlantic at 50° S between 65° W and 35° W in March 2008, who reported an average O₃ mixing ratio of 18.3 ppbv. Similar O₃ mixing ratios of the order of 20 ppbv were also observed in the southern Atlantic south of 40° S during two Polarstern cruises in November 1990 (Slemr and Tremmel, 1994) and October/November 1994 (Junkermann and Stockwell, 1999). As discussed in Lelieveld et al. (2004) seasonal variations of O₃ in the latitude band between 40 and 60° S are rather small, with slightly lower values in austral summer. For the period between 1977 and 2002 the calculated O₃ trend for this latitude band is 0.17 ± 0.08 ppbv year⁻¹ (Lelieveld et al., 2004), indicating a moderate increase of approximately 1 ppbv between 2002 and 2007. Based on the limited number of observations during OOMPH a confirmation of the continuation of this trend is not possible.

Time series of measured and observed H₂O₂ are shown in Fig. 4. During the first part of the cruise (10 to 14 March) observed H₂O₂ mixing ratios indicate small variability at levels between 200 and 300 pptv. During this period the model significantly underestimates observed H₂O₂, by more than a factor of two. After 14 March in the eastern part of the South Atlantic H₂O₂ mixing ratios tend to increase both in the observations and the model simulations, with the model overestimating observations during the first part, in particular on 15 March, while later on measurements and model results agree quite well. A period of relatively high H₂O₂ (in excess of 1 ppbv) was observed and simulated after midnight on 16 March, southwest of the African coast (see Fig. 1). On average observed and simulated H₂O₂ were 350 ± 220 pptv and 310 ± 240 pptv, respectively. Slope and intercept of a least-square regression analysis are 2.04 ± 0.04 and −0.27 ± 0.04 ppbv, respectively. The regression coefficient $R^2$ is 0.46. The imperfect
agreement here between model results and observations is due to the rather limited dynamical range of mixing ratios and the strong offset during the first part of the cruise. This is also the case for O₃. As can be deduced from the time series (Fig. 4) the model tends to reproduce trace gas levels over the Southern Atlantic. The measured H₂O₂ mixing ratios are comparable to previously reported observations south of 35° S in the Atlantic lower troposphere, being approximately 200–300 pptv (Slemr and Tremmel, 1994; Junkerman and Stockwell, 1999; Weller et al., 2000).

Observations of CH₃OOH mixing ratios are shown in the time series in Fig. 5 along with EMAC simulations. The temporal evolution is similar to H₂O₂, with low values (less than 200 pptv) during the first part of the cruise, a strong mixing ratio increase starting on midnight of 16 March and lower concentrations afterwards. The relative change is reproduced by the model, with a general tendency to overestimate CH₃OOH mixing ratios except during the period when the high peak was encountered on 16 March. Average measured CH₃OOH mixing ratios are 280 ± 250 pptv, while the model calculates a mean value of 450 ± 190 pptv. The offset between model and observations is clearly shown in the least-square regression analysis that yields an intercept at 0.18 ± 0.05 ppbv and a slope of 0.37 ± 0.12 at an R² of 0.6. In general the observed mixing ratios are in good agreement with reported levels in the literature (Slemr and Tremmel, 1994; Junkerman and Stockwell, 1999; Weller et al., 2000).

With the exception of the mixing ratio peaks on 16 March, the variability of O₃, H₂O₂ and CH₃OOH is small across the South Atlantic basin between South America and South Africa. The slightly northward orientation of the ship track results in a steady increase in temperature and photolysis rates (Hosaynali Beygi et al., 2011) leading to increasing photochemical activity that is likely responsible for the slight increase of the mixing ratios of the three species from the south-western to the north-eastern part of the South Atlantic. As shown in a backward trajectory analysis presented in Hosaynali Beygi et al. (2011) the air mass origin did not change throughout the cruise. All air masses encountered during the cruise originated in the western Antarctic Peninsula.
during the preceding week and had not been in contact with land surfaces for at least 5 days.

3.3 Discussion

Based on $\text{H}_2\text{O}_2$ mixing ratios and the comparison between observations and model results, one can differentiate three different periods in Fig. 4: from 10 to 14 March the model significantly underestimates the $\text{H}_2\text{O}_2$ mixing ratios, while the agreement is much better during the final phase of the campaign (after 17 March). In between these two periods $\text{H}_2\text{O}_2$ mixing ratios show a strong increase to well above 1 ppbv that is well reproduced by the model.

Similar behavior is also observed for ozone (Fig. 3). The model also tends to underestimate $\text{O}_3$ mixing ratios during the early phase of the campaign, while the agreement is much better in the second half of the campaign. On the other hand, the $\text{CH}_3\text{OOH}$ mixing ratios are almost always overestimated by the model (Fig. 5). Considering the ratio between $\text{CH}_3\text{OOH}$ and $\text{H}_2\text{O}_2$ there is an even stronger discrepancy (Fig. 6). Over the whole campaign, the observed ratio varies between 0.5 and 1 with a mean value and standard deviation of $0.8 \pm 1.1$, while the model predicts a time dependent ratio between 2 and 4 during the first half of the campaign and lower values, in good agreement with the observations, during the second half of the campaign. Hence the main problem seems to be that the model underestimates $\text{H}_2\text{O}_2$ during the early phase of the campaign, due to either an underestimation of the $\text{H}_2\text{O}_2$ production or an overestimation of the sinks. Gas phase $\text{H}_2\text{O}_2$ in the marine boundary layers stems from the recombination of two HO$_2$ radicals. As shown by Hosaynali Beygi et al. (2011), EMAC reproduces observed HO$_2$ levels during the whole campaign and indicates similar levels for HO$_2$ and CH$_3$O$_2$, the precursors of CH$_3$OOH. Given that the precursor levels are simulated realistically by EMAC, it is very unlikely that an underestimation of the peroxide production is responsible for the $\text{H}_2\text{O}_2$ underestimation during the first half of the campaign. This indicates that the discrepancies are due to an overestimation of the $\text{H}_2\text{O}_2$ sinks in the model during this period. Photochemical sinks (reaction with OH and $\text{H}_2\text{O}_2$ pho-
tolysis) are also unlikely causes, since the model also reproduces OH concentrations (Beygi et al., 2011) and radiation intensities (not shown). This leaves physical removal processes, such as rainout and dry deposition to the surface, as most likely causes. Although the model predicts some rain events, they are not particularly extensive during the first phase, when the problems occur. A striking observation is that the wind speeds are significantly higher during the first part of the campaign, being well above 10 m s\(^{-1}\) up to 15 March (Fig. 1), while they are generally lower in the second half of the campaign. The dry deposition in EMAC is simulated according to Kerkeweg et al. (2006) following Wesley (1989). It strongly depends on the wind speed, which determines the transfer velocity to the ocean surface. The loss itself is determined by the solubility of the species under investigation, defined by its Henry’s law coefficient. Based on this parameterization one expects increasing dry deposition losses going from CH\(_3\)OOH (only moderately soluble) to H\(_2\)O\(_2\) (highly soluble). Ozone has an intermediate role, since its solubility is in between these two extremes. The deposition velocity calculated by the model for O\(_3\) does not depend on the wind speed and is about 0.5 cm s\(^{-1}\), indicating that the deposition loss is limited by the solubility of O\(_3\). On the other hand, the H\(_2\)O\(_2\) deposition velocity is a strong function of wind speed, linearly increasing from \(\sim 5\) cm s\(^{-1}\) at a wind speed of 5 m s\(^{-1}\) to about 18 cm s\(^{-1}\) at 10 m s\(^{-1}\). This indicates that the deposition loss for this highly soluble species is limited by the transfer velocity to the ocean surface. Accordingly, due to the absence of low clouds and precipitation during the campaign (for the considered period) dry deposition is the dominant loss process for H\(_2\)O\(_2\) in the model, even during the day. During the night dry deposition is the only loss mechanism, as photochemical destruction ceases. The model indicates that during daytime the contribution of dry deposition to total H\(_2\)O\(_2\) loss varies between 98% (noon values) during the first part of the campaign (11 to 14 March) and about 90% during the second half (after 15 March), due to a combination of decreasing dry deposition loss due to decreasing wind speeds and a simultaneous increase by enhanced photochemical activity at lower latitudes. We performed a sensitivity study with EMAC, limiting the dry deposition velocity to 5 cm s\(^{-1}\) (SR1), the value calculated for
the period with lowest wind speed (5 m s\(^{-1}\)). The green line in Fig. 4 shows that this leads to an increase in simulated H\(_2\)O\(_2\) mixing ratios by approximately 50 % (mean calculated mixing ratio: 460 ± 350 pptv). It should be mentioned that globally this effect is strongest in the marine boundary layer in regions of high wind speed, notably in the latitude band between 40 and 60° of the storm tracks in both hemispheres. Outside of these regions the effect is much smaller and leads to increases in the H\(_2\)O\(_2\) mixing of less than 20 %. Thus it appears that the differences between model simulations and H\(_2\)O\(_2\) observations are due to a model overestimation of dry deposition to the ocean at high wind speed. This hardly affects CH\(_3\)OOH and O\(_3\) whose deposition loss is limited by their solubility and thus independent of wind speed, while H\(_2\)O\(_2\) is much more strongly affected. The importance of the deposition parameterization, being a critical process in the simulation of H\(_2\)O\(_2\) in the lower troposphere was also emphasized by Chang et al. (2004), who performed sensitivity studies with a single column model to simulate observations from PEM-Tropics B.

Another interesting feature of the H\(_2\)O\(_2\) time series in Fig. 4 is the strong increase of H\(_2\)O\(_2\) in the night from 15 to 16 March. Shortly before midnight the H\(_2\)O\(_2\) mixing ratio increases strongly from ~ 0.7 ppbv up to about 1.5 ppbv, and decreasing to the previous mixing ratios before noon on 16 March. The sudden increase occurs together with a similar increase in CH\(_3\)OOH (Fig. 5), which is well reproduced by EMAC. Since the sudden increase in H\(_2\)O\(_2\) and CH\(_3\)OOH occurs during nighttime a photochemical source can be excluded. During this phase of the campaign the ship was south of the southern tip of Africa. One explanation for the sudden increase could be a change in air mass origin. Heikes et al. (1996) observed a significant increase of marine boundary layer H\(_2\)O\(_2\) mixing ratios north of 20° N in continental outflow. Trajectory calculations for the OOMPH campaign were presented in the supplements to the paper by Hosaynali Beygi et al. (2011). They indicate, however, no change in air mass origin during the period between 15 and 17 March, so that this option can be dismissed.

Due to the dry deposition close to the ocean surface, both H\(_2\)O\(_2\) and CH\(_3\)OOH are expected to exhibit an increase in mixing ratios with height. Aircraft observations over
the ocean indeed show maxima of both species above the marine boundary layer (Heikes et al., 1996; O’Sullivan et al., 1999, 2004; Stickler et al., 2007). Thus transport from above the boundary layer can be a source of \( \text{H}_2\text{O}_2 \) and \( \text{CH}_3\text{OOH} \) for the marine boundary layer. Observations of the boundary layer height were not made during the OOMPH campaign, thus we have to rely on model results. The curtain plot in Fig. 7 shows a time series of the vertical profiles for \( \text{O}_3 \) (upper panel) and \( \text{H}_2\text{O}_2 \) (lower panel). Superimposed is the temporal evolution of the boundary layer height calculated by EMAC. Shortly before the event the simulation indicates a very shallow boundary layer (\( \sim 200 \text{ m} \)), that starts to increase around midnight on 16 March. Hence the increase in \( \text{H}_2\text{O}_2 \) is related to an increase in boundary layer height, and downward mixing of air masses that have not been affected by deposition and thus having higher mixing ratios of peroxides. Thus it seems that nighttime transport of free tropospheric air into the marine boundary layer is responsible for the concentration increase during the period from 15 to 16 March. This vertical redistribution process should also affect other species with a positive altitude gradient, in particular ozone. Model profiles indicate that the mixing ratio gradient for \( \text{O}_3 \) is much smaller than for \( \text{H}_2\text{O}_2 \) (upper panel in Fig. 7). This is consistent with the moderate change in \( \text{O}_3 \) mixing ratios during this event. Note that the diurnal variation of the boundary layer height during 17–19 March is related to the proximity to the African continent. Due to the limited resolution of the model the data points here are extrapolated between an oceanic and a continental cell, leading to a diurnal evolution of the boundary layer height that resembles that of a continental boundary layer instead of the marine boundary layer.

Finally, the difference in simulated and observed absolute mixing ratios of \( \text{CH}_3\text{OOH} \) needs to be addressed. Figure 5 indicates that although the model reproduces the relative changes quite well, the absolute values are off by a factor of about 2, with the model being consistently higher. The relative difference is higher in the beginning and lower towards the end of the campaign, but the absolute difference remains about 200 pptv. There are several potential explanations for this discrepancy. First, as mentioned in Sect. 2.2 \( \text{CH}_3\text{OOH} \) was not directly measured, but inferred from the ROOH...
signal, assuming that CH$_3$OOH is the only organic hydroperoxide and that the sampling efficiency can be calculated according to Lee et al. (2000). An experimental verification of the sampling efficiency was not performed (contrary to H$_2$O$_2$) since no CH$_3$OOH gas phase source was available. If the actual sampling efficiency was lower than the calculated 60%, this could close the gap between observations and model results. Additionally, the sampling efficiency would have to be lower at the beginning of the campaign and higher later on, since a simple multiplication by a factor cannot account for the rather constant absolute difference over the campaign. Therefore, although an error in the sampling efficiency cannot be excluded, it is unlikely the only source of the discrepancy between model and observations.

As discussed above, Hosaynali Beygi et al. (2011) demonstrated that EMAC very well reproduces the HO$_2$ measurements made during the OOMPH cruise. Since the precursor for H$_2$O$_2$ is simulated correctly by the model, one can assume that the source strength for H$_2$O$_2$ is correctly simulated. The model predicts CH$_3$O$_2$ concentrations that are similar to HO$_2$, but unfortunately measurements of CH$_3$O$_2$ were not made. Thus it is not possible to validate the model predicted precursor concentrations for CH$_3$OOH. But as discussed in Hosaynali Beygi et al. (2011) the simulated CH$_3$O$_2$ levels are quite realistic. Recently Fittschen et al. (2014) posed that the reaction of CH$_3$O$_2$ with OH radicals at low NO$_x$ concentrations can be a significant sink of methylperoxy radicals and thus could reduce CH$_3$OOH formation. In a sensitivity study (SR2) this reaction was included in the chemistry code of EMAC (Bossolasco et al., 2014). Additionally, we considered that in addition to CH$_3$OOH the reaction of CH$_3$O$_2$ with HO$_2$ also produces HCHO to some extend (Ayers et al., 1997). The green line in Fig. 6 indicates that including these additional reaction pathways reduces the mixing ratio of CH$_3$O$_2$ in the marine boundary layer of the South Atlantic by about 30%, yielding an average mixing ratio of 300 ± 110 pptv compared to the observed 180 ± 50 pptv. It seems that the missing reaction between CH$_3$O$_2$ and OH is responsible for a largest part of the CH$_3$OOH overestimation by the model, in particular during the second half of the cruise when OH concentrations are high. In general, the contribution of this reaction increases with
increasing OH concentration, which leads globally to a maximum impact in the tropical lower troposphere over the Pacific and Indian Oceans. The contribution of the HCHO channel in the reaction of CH$_3$O$_2$ with HO$_2$ is rather constant with a branching ratio of about 10%. Nevertheless, due to the uncertainties in both measurements and modeling the explanation for the difference in CH$_3$OOH simulation vs. observations remains ambiguous.

By combining both the reduced H$_2$O$_2$ deposition velocity and the reduced CH$_3$OOH source due to the competing reaction of CH$_3$O$_2$ with OH in an additional sensitivity simulation (SR3) the simulated CH$_3$OOH/H$_2$O$_2$ ratio is reduced to a mean value of 1.01 ± 1.06 (green line in Fig. 6), much closer to the observed value (0.8 ± 1.1).

4 Summary and conclusions

Hydrogen peroxide, MHP and ozone have been measured in the marine boundary layer over the South Atlantic Ocean during the austral summer in 2007. Observed mixing ratios are consistent with values reported in the literature. Simulations with the atmospheric chemistry model EMAC indicate that it qualitatively reproduces the observations very well. It also captures the downward mixing of high concentrations of H$_2$O$_2$ and CH$_3$OOH during a nighttime increase in boundary layer height south of the African continent. Quantitatively, the model tends to overestimate H$_2$O$_2$ mixing ratios during the first part of the cruise. During this part we experienced very high wind speeds, in excess of 15 m s$^{-1}$. Later on during the cruise much lower wind speeds were encountered for which the model accurately reproduces observed H$_2$O$_2$ mixing ratios. A similar tendency has been observed for ozone. The most likely reason for the discrepancy is that the model parameterization of trace gas deposition to the ocean surface tends to overestimate the dry deposition loss of the soluble and reactive H$_2$O$_2$ at high wind speeds. This was confirmed by a model sensitivity study with limited (at 5 cm s$^{-1}$) deposition velocity. MHP, which is less soluble and reactive, is not as much of affected by dry deposition, although EMAC significantly overestimates its mixing ra-
This overestimation is rather constant over the campaign and indicates an offset of approximately 200 pptv. The reasons for this discrepancy are not easy to identify, since the measurements provide less stringent constraints than for H$_2$O$_2$. MHP was estimated from a total ROOH signal, assuming MHP being the only ROOH component and a sampling efficiency of 60% compared to H$_2$O$_2$. Fittschen et al. (2014) recently suggested that a competing reaction of the CH$_3$O$_2$ radicals with OH can significantly diminish the source of MHP in the marine boundary layer, in particular at the low NO$_x$ concentrations observed during OOMPH. A sensitivity study with EMAC indicates that this reaction reduces the CH$_3$O$_2$ mixing ratio by about 30%, increasing with decreasing latitude due to the meridional gradient in OH radical concentrations. Although this is a significant change, bringing CH$_3$OOH closer to observations, it appears to be insufficient to fully remove the discrepancy between simulated and observed CH$_3$OOH mixing ratios during OOMPH.

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References


Chang, W., Lee, M., and Heikes, B. G.: One-dimensional photochemical study of H$_2$O$_2$, CH$_3$OOH, and HCHO in the marine boundary layer during Pacific Exploratory Mission in


Kerkweg, A., Buchholz, J., Ganzeveld, L., Pozzer, A., Tost, H., and Jöckel, P.: Technical Note: An implementation of the dry removal processes DRY DEPosition and SEDimenta-
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Figure 1. Ship track of the R/V *Marion Dufresne* over the southern Atlantic. The track is color coded with time.
Figure 2. Time series of observed and simulated temperature and wind speed.
Figure 3. Time series of observed (red) and simulated (blue) ozone mixing ratios.
Figure 4. Time series of observed (red) and simulated (blue) hydrogen peroxide mixing ratios. In green a sensitivity simulation (SR 1) of the model is shown with reduced dry deposition velocity (see text for details).
**Figure 5.** Time series of observed (red) and simulated (blue) methyl hydroperoxide mixing ratios. The green line shows a sensitivity simulation (SR2) including the reaction of CH$_3$O$_2$ with OH (see text for details).
**Figure 6.** Simulated (blue) and observed (red) CH$_3$OOH to H$_2$O$_2$ ratio. The green line shows a sensitivity simulation (SR3) including reduced H$_2$O$_2$ dry deposition velocity and the reaction of CH$_3$O$_2$ with OH (see text for details).
Figure 7. EMAC simulation of $O_3$ (top) and $H_2O_2$ (bottom) vertical profiles vs. time. Superimposed is the height of the planetary boundary layer. The increase in boundary layer height in the night from 15 to 16 March is associated with an increase in peroxide mixing ratios.