Oxidation processes in the eastern Mediterranean atmosphere: evidence from the modelling of HOx measurements over Cyprus

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Abstract. The Mediterranean is a climatically sensitive region located at the crossroads of air masses from three continents: Europe, Africa, and Asia. The chemical processing of air masses over this region has implications not only for the air quality but also for the long-range transport of air pollution. To obtain a comprehensive understanding of oxidation processes over the Mediterranean, atmospheric concentrations of the hydroxyl radical (OH) and the hydroperoxyl radical (HO2) were measured during an intensive field campaign (CYprus PHotochemistry EXperiment, CYPHEX-2014) in the northwest of Cyprus in the summer of 2014. Very low local anthropogenic and biogenic emissions around the measurement location provided a vantage point to study the contrasts in atmospheric oxidation pathways under highly processed marine air masses and those influenced by relatively fresh emissions from mainland Europe.

The CYPHEX measurements were used to evaluate OH and HO2 simulations using a photochemical box model (CAABA/MECCA) constrained with CYPHEX observations of O3, CO, NOx, hydrocarbons, peroxides, and other major HOx (OH + HO2) sources and sinks in a low-NOx environment (< 100 pptv of NO). The model simulations for OH agreed to within 10 % with in situ OH observations. Model simulations for HO2 agreed to within 17 % of the in situ observations. However, the model strongly under-predicted HO2 at high terpene concentrations, this under-prediction reaching up to 38 % at the highest terpene levels. Different schemes to improve the agreement between observed and modelled HO2, including changing the rate coefficients for the reactions of terpene-generated peroxy radicals (RO2) with NO and HO2 as well as the autoxidation of terpene-generated RO2 species, are explored in this work. The main source of OH in Cyprus was its primary production from O3 photolysis during the day and HONO photolysis during early morning. Recycling contributed about one-third of the total OH production, and the maximum recycling efficiency was about 0.7. CO, which was the largest OH sink, was also the largest HOx source. The lowest HOx production and losses occurred when the air masses had higher residence time over the oceans.

1 Introduction

1.1 Air pollution and HOx chemistry

The chemical and photochemical processing of air pollutants, in conjunction with local emissions, meteorology, and atmospheric transport, strongly influences the air quality over a region. The regional air quality impacts human health, agriculture, the overall condition of the biosphere, and subsequently the climate. Studies attribute 2–4 million premature deaths globally to outdoor air pollution (Silva et al., 2013;
Lelieveld et al., 2015). Oxidants in the Earth’s atmosphere prevent the pollutants released into it from building up to toxic levels. These oxidants not only convert many toxic pollutants into less toxic forms, e.g. carbon monoxide (CO) to carbon dioxide (CO₂), but also help in their removal, e.g. nitrogen oxides (NOₓ) and sulfur dioxide (SO₂) are converted into soluble nitric acid (HNO₃) and sulfuric acid (H₂SO₄) respectively, although some toxic chemicals may still be formed during degradation of non-toxic ones. Most of the oxidation processes in the atmosphere proceed along reaction pathways initiated by the hydroxyl radicals (OH) during the day (Levy, 1971), making it the dominant chemical cleaning agent of the daytime atmosphere (Lelieveld et al., 2004). The dominant primary production of OH is via the photolysis of ozone (O₃) at ultraviolet wavelengths (Crosley, 1995), producing electronically excited O(¹D) atoms (Reaction R1). O(¹D) that escapes quenching (Reaction R2) reacts with water vapour in the atmosphere to produce OH radicals (Reaction R3).

\[
\begin{align*}
O_3 + h\nu & \rightarrow O(¹D) + O_2 \quad (R1, \lambda < 330\text{nm}) \\
O(¹D) + M & \rightarrow O(³P) + M(M = N_2, O_2) \quad (R2) \\
O(¹D) + H_2O & \rightarrow 2OH \quad (R3)
\end{align*}
\]

Because OH reacts rapidly with several biogenically and anthropogenically emitted gases (e.g. Reaction R4), its lifetime is typically a few hundred milliseconds in the troposphere. Oxidation limits these gases (methane (CH₄), CO, volatile organic compounds (VOCs)) from accumulating above trace levels in the troposphere. Reactions of OH with CO and hydrocarbons (HCs) (Reaction R4) also trigger quasi-instant formation of the hydroperoxyl radical (HO₂, via Reaction R5) in our O₂-rich atmosphere. In the presence of nitric oxide (NO) and O₃ (Reactions R6, R7), a chemical equilibrium is set between OH and HO₂ on a timescale of seconds. The sum of OH and HO₂ is known as HOₓ. The Reactions (R6) and (R7) maintain the oxidizing efficiency of the atmosphere by recycling OH. Simultaneously, in the presence of NOₓ (= NO + nitrogen dioxide (NO₂)), they also propel a reaction chain leading to O₃ formation via Reactions (R8) and (R9). Reactions (R8)–(R9) are the main chemical source of tropospheric O₃, which is an important oxidant, a greenhouse gas, and a major secondary pollutant.

\[
\begin{align*}
\text{CO} + \text{OH} & \rightarrow \text{CO}_2 + \text{H} \quad (R4) \\
\text{H} + \text{O}_2(=\text{M}) & \rightarrow \text{HO}_2(=\text{M}) \quad (R5) \\
\text{NO} + \text{HO}_2 & \rightarrow \text{NO}_2 + \text{OH} \quad (R6) \\
\text{HO}_2 + \text{O}_3 & \rightarrow \text{OH} + 2\text{O}_2 \quad (R7) \\
\text{NO}_2 + h\nu & \rightarrow O(³P) + \text{NO}(\lambda < 430\text{nm}) \quad (R8, \lambda < 430\text{nm}) \\
O(³P) + \text{O}_2(=\text{M}) & \rightarrow \text{O}_3(=\text{M}) \quad (R9)
\end{align*}
\]

OH also oxidizes anthropogenic and biogenic HCs emitted into the atmosphere, leading to peroxy radicals (RO₂). NO converts the RO₂ into HO₂, which again leads to OH recycling (Reaction R6). The HOₓ cycle can be terminated by the radical+radical interactions of RO₂ and HO₂ with and among themselves to form peroxydis (Reactions R10 and R11). Depending on NO levels, the RO₂ species can react through different channels. Under low-NOₓ conditions, e.g. in the clean marine boundary layer, when the reaction of RO₂ with NO is less significant than its reaction with HO₂, peroxy formation becomes a major sink for HO₂, further leading to OH and O₃ removal. Under higher NO levels, RO₂ would react with NO to form NO₂ (O₃ source), HO₂ (OH source), alkyl nitrates, and aldehydes (Reactions R12, R13, and R14). Further, there have been suggestions of auto-oxidation of RO₂ to produce peroxydis, yielding OH under atmospheric conditions (Crounse et al., 2013). These auto-oxidation mechanisms proceed through catalytic H shifts at a very fast rate (up to ~0.1 s⁻¹), e.g. conversion of RO₂ generated from a carbonyl compound into a highly functionalized dicarbonyl hydroperoxide compound (Reaction R15).

\[
\begin{align*}
\text{HO}_2 + \text{HO}_2 & \rightarrow \text{H}_2\text{O}_2 + \text{O}_2 \quad (R10) \\
\text{RO}_2 + \text{HO}_2 & \rightarrow \text{ROOH} + \text{O}_2 \quad (R11) \\
\text{RO}_2 + \text{NO} + \text{O}_2 & \rightarrow \text{R}’\text{CHO} + \text{HO}_2 + \text{NO}_2 \quad (R12) \\
\text{RO}_2 + \text{NO} & \rightarrow \text{RNO}_3 \quad (R13) \\
\text{NO}_2 + \text{OH} + \text{M} & \rightarrow \text{HNO}_3 + \text{M}(M = \text{N}_2, \text{O}_2) \quad (R14)
\end{align*}
\]

In polluted air when NOₓ concentrations become large enough that the reaction of NO₂ with OH dominates over other HOₓ sinks, the major loss of HO₂ is via the formation of nitric acid (HNO₃, Reaction R14), which is then removed from the atmosphere by wet and dry deposition. Some acid-forming reactions are also the starting point of particle formation in the atmosphere, e.g. via oxidation of SO₂. In sum, oxidation is a means by which major pollutants are processed and removed from the atmosphere, but with concomitant implications for air quality, agriculture, climate, and health. However, large uncertainties in atmospheric reaction pathways, rate coefficients, and measurements of atmospheric gases and radicals limit our ability to understand and predict the quality of ambient air.

1.2 Motivation for the campaign

The Mediterranean is a climatically sensitive region which is rapidly getting warmer and dryer over time (Lelieveld et al., 2012). The atmospheric chemistry over the region is complex, with regular exceedances of the European Union (EU) ozone air quality standard (Doche et al., 2014; Kalabokas et al., 2013) and a projected increase in summertime O₃ over the eastern Mediterranean (Lelieveld et al.,...
Cyprus is a small, remote island (area: 9251 km$^2$; population: $1.15 \times 10^6$; density: 119.2 people km$^{-2}$) in the eastern Mediterranean (south of mainland Europe) with very low local emissions. The non-methane volatile organic compounds (NMVOCs), NO$_x$, and SO$_x$ emitted by Cyprus and the European Union (EU28) were 6.8, 17.2, and 16.8 Gg and 6722, 7819, and 3083 Gg respectively for 2014 (European Union emission inventory report, 2017). Being located downwind of the mainland European emissions, Cyprus provides an ideal vantage point to study the impact of European emissions on the Mediterranean climate (Lelieveld et al., 2002). To study the atmospheric chemistry over the Mediterranean and understand the impact of emissions from different regions of Europe on air quality in Cyprus, an intensive field measurement campaign (the Cyprus Photochemistry EXperiment, hereafter CYPHEX) was conducted in Cyprus during the summer of 2014, wherein a comprehensive suite of trace gases was measured.

Cyprus is exposed to air masses influenced by emissions from southwestern European countries that are photochemically processed over the Mediterranean Sea. The Etesian winds during summer carry air masses loaded with emissions from industry and biomass burning over central and eastern European countries. Although emissions from Europe have been steadily decreasing over the past few decades (e.g. NO$_x$, SO$_x$, and NMVOCs decreased by nearly 50%, 80%, and 57% respectively during 1990–2013; European Union emission inventory report, 2017), these emissions are still substantial with respect to the global total. Apart from the influence of European emissions, Cyprus is strategically located at a crossroads of air masses from Asia, Africa, and the Atlantic (Kleanthous et al., 2014). Influence from all these different source regions, incorporating the impact of a plethora of HCs, nitrogen and sulfur species, and mineral dust on the atmospheric processing in this part of the world, has attracted several field campaigns (Carslaw et al., 2001; Berresheim et al., 2003). For example, the Mediterranean Intensive Oxidant Study (MINOS) was conducted to study the budget of atmospheric oxidants influenced by long-range transport of pollution in the summer of 2001 (Lelieveld et al., 2002; Gros et al., 2003; Salisbury et al., 2003). OH radicals were measured in the coastal boundary layer of Crete as part of the MINOS campaign and the major sources of OH were found to be photolysis of O$_3$ and recycling from HO$_2$ + NO reaction (Berresheim et al., 2003). Although HO$_2$ was not measured during MINOS, high levels of formaldehyde (HCHO), a major primary HO$_2$ source, were observed (Kormann et al., 2003). In the present study, we report simultaneous measurements of OH and HO$_2$ during the CYPHEX campaign and compare these observations with simulations from a photochemical box model constrained with O$_3$; CO; NO; NO$_2$; peroxides; several anthropogenic and biogenic HCs including important alkanes, alkenes, alkynes, aldehydes, and acids; and photolysis frequencies of important gases relevant to HO$_x$ chemistry. The main objective of this study is to identify the major chemical species and reaction pathways controlling the HO$_x$ chemistry over this climatically sensitive region.

## 2 Methods

### 2.1 Measurement site

The CYPHEX campaign was conducted at the northwest coast of Cyprus, on a hilltop about 650 m a.m.s.l., in Paphos District of the island and facing the sea about 5–8 km away on the west. The Akamas Peninsula National Park is about 25 km to the northwest of the measurement site (34°57′ N, 32°23′ E), with terrain descending rapidly to the northeast towards the town of Polis, with a population of about 2000. The area comprises sparse shrub-type vegetation with small Mediterranean trees comprising pines, junipers, olives, carob, pomegranates, and almonds. The area is weakly populated in a radius of about 20 km. The site is away from the major cities: Paphos with 90 000 people is 20 km south, Limassol with 235 000 people is 70 km southeast, Larnaca with 145 000 people is 110 km east, and Nicosia with 325 000 people is 90 km northeast (census of population October 2011; www.cystat.gov.cy, last access: 28 December 2017). The local wind direction is predominantly southwest (70% between 202.5° and 247.5°) (Meusel et al., 2016; Derstroff et al., 2017), bringing in humid air from the sea, with no immediate anthropogenic influence. Analysis of 5-day back trajectories show mainly two major air mass regimes (Hüser et al., 2017). Etesian winds influenced by fresh emissions from eastern–central Europe crossing Turkey and Greece, and mistral winds influenced by emissions from southwestern Europe but processed for a longer period over the Mediterranean (Fig. 1). During the summer of 2014, a weakened east–west pressure gradient led to weak and delayed Etesian winds (Tyrlis et al., 2015). The site is also influenced by local land–sea breezes.

### 2.2 HO$_x$ measurements during CYPHEX

Atmospheric OH and HO$_2$ were measured during CYPHEX using the HydrOxyl Radical measurement Unit based on fluorescence Spectroscopy (HORUS) instrument (Martinez et al., 2010). The set-up was based on the well-established laser-induced fluorescence – fluorescence assay by gas expansion (LIF-FAGE) technique for atmospheric OH measurements (Brune et al., 1995; Crosley, 1995; Hard et al., 1984). The laser system consisted of a tunable dye laser, which was pumped by a diode-pumped Nd:YAG laser (Navigator I J40-X30SC-532Q, Spectra Physics) pulsing at 3 kHz. The two laser assemblies were mounted on either side of a vertical plate, which was mounted on a rack inside a measurement container with dimensions of 6 m $\times$ 2.5 m $\times$ 2.5 m. The output laser radiation was split into a 9 : 1 ratio using
beam splitters, and channelled through 8 m optical fibers into the detection module for measurements of OH and H\textsubscript{2}O.

The detection system was mounted about 6 m above ground level (a.g.l.) on the top of a scaffolding tower, constructed close to the wall of an air-conditioned container adjacent to the laser unit. The container with the HORUS instrument was mounted on top of another similar measurement container with O\textsubscript{3}, CO, NO\textsubscript{x}, and peroxide instruments inside. The inlets for all the instruments measuring during CYPHEX were placed within 5 m of the HORUS detection unit on the tower. Ambient air was drawn in HORUS through a critical orifice with a pinhole size of 0.9 mm into a detection cell with about 4.3 mbar of pressure. This internal pressure was achieved by using a combination of a roots blower (M90 compressor from Eaton, USA) followed by a scroll pump (XDS35i, Edwards, USA). The resulting high volume flow ensured that the exchange of air illuminated by the laser is fast enough between two consecutive pulses, thus avoiding possible laser-induced interferences.

The interaction of the ambient air with the laser beam in the low-pressure detection cell was maximized using a White cell set-up, in which the laser light was reflected 32 times across the detection volume. The OH molecules were selectively excited by a 308 nm laser light pulsing at 3 kHz on resonance with the \( Q_1 (2) \) transition line (\( \Delta^2 \Sigma - \chi \Pi \), \( \nu' = 0 \leftarrow \nu'' = 0 \)). An etalon controlled by a stepper motor, in the dye laser set-up, caused the laser radiation to be tuned on and off resonance with the OH transition every 7 s to account for the OH fluorescence plus background signals and the background signals respectively, resulting in a time resolution of 14 s. The spectra from a reference cell, in which OH was produced by H\textsubscript{2}O thermolysis, was used to detect the \( Q_1 (2) \) line. The counts from the reference cell were used to sustain the on-resonance position while the etalon tethered around it. Although the pressure reduction in the detection cell results in smaller numbers of OH molecules, it improves the OH fluorescence quantum yield due to reduced collisional quenching of excited OH radicals. The lifetime of the excited OH extends beyond the Mie scattering by aerosols and the Raleigh scattering by smaller molecules/particles. These interfering background signals were discriminated by using time-gated microchannel plate detectors (Hamamatsu), enabling excitation of OH radicals and detection of fluorescence at the same wavelength. The excitation at 308 nm is advantageous to the excitation at 282 nm as interference from laser-generated OH due to O\textsubscript{3} photolysis is \( \sim 25 \) times smaller at 308 nm. A band-pass filter with a central wavelength of 308 \( \pm 0.75 \) nm and bandwidth of 5 nm (full width at half maximum, FWHM) in front of the detector prevented signals due to any interfering wavelengths from registering on the detector. The fluorescence decay was recorded with a time resolution of 4 ns and integrated for on-resonance and off-resonance periods, thus delineating the signal from the spectroscopic background. Further, the spectra of the measured atmospheric OH were compared with the one obtained from a reference cell ensuring no interferences from fluorescence of species like SO\textsubscript{2} and naphthalene.

The chemical background, originating from OH being internally produced during the transit time of about 3 ms from the orifice to the centre of the detection cell, was accounted for by removing ambient OH with an OH scavenger. This scavenger, 3–5 sccm of pure propane in a carrier flow of 8000 sccm synthetic air for CYPHEX, was introduced through an inlet pre-injector (IPI) mounted on top of the inlet nozzle (Novelli et al., 2014a; Hens et al., 2014). The scavenger amount was just sufficient to scavenge off \( \sim 95 \% \) of atmospheric OH as determined from propane titration experiments on site. IPI was connected to a blower that draws about 180–200 L min\(^{-1}\) of ambient air from the top of the IPI, the flow being monitored by a differential pressure sensor. The
OH scavenger was injected via eight 0.5 mm diameter holes 5 cm above the pinhole of the OH inlet into the centre of the air flow sampled by the IPI close to its hyperbolic minima (Novelli et al., 2014b). The hyperbolic internal shape of the IPI (maximum cross section of 35 mm and minimum cross section of 6 mm) helps to sample air that has minimum wall contact and flows at a high velocity through a small diameter, ensuring that the scavenger is well mixed with atmospheric air. The high flow also ensures minimum impact of the horizontal wind speed on the mixing efficiency. The IPI cycling was automated by a script resulting in repeated cycles of scavenger injection, flushing, and no injection. The scavenger was periodically injected every 2 min, resulting in alternating measurements of background OH and total OH, i.e. with and without scavenger injection. The difference between these two signals gives a measure of the atmospheric OH with a time resolution of 4 min. The wall losses in the IPI were periodically determined by physically dismounting the IPI for 5 min during measurements during different times of the day. On average, the background signal was 45 % of the total signal during the daytime (Fig. S1 in the Supplement). While the constituents of background OH can have a pronounced impact on atmospheric oxidation processes, especially over regions with high biogenic emissions (Mauldin et al., 2012), the study of the influence of background OH on oxidation pathways during CYPHEX is beyond the scope of the present study.

Atmospheric HO₂ was measured in a detection cell located 16 cm below the OH detection block. HO₂ measurements were achieved by injecting NO purified through a sodium hydroxide-coated silica (Sigma-Aldrich Ascarite) using a 1/8 in. OD, 0.035 in. ID coiled stainless steel loop with a diameter of about 3 cm, placed just above the HO₂ detection block. The NO was injected through 0.1 mm holes, leading to conversion of HO₂ into OH and detected by the same principle as in the first block. To minimize impact of RO₂ on OH formation (Fuchs et al., 2011; Hens et al., 2014), only about 30 % of atmospheric HO₂ was converted to OH using NO. The required NO flow, about 0.5 sccm of pure NO in 10 sccm of carrier flow during CYPHEX, was determined from titrations in ambient and calibration air. Further, tests were carried out with and without NO to ensure that there was no influence of HO₂ conversion on OH in the first axis. The measurement in the HO₂ detection cell yields the total HO₂ (OH + HO₂). HO₂ is calculated as the difference between HO₂ and the OH measured in the first axis, after accounting for the relative OH sensitivities of the two detection axes.

Due to low OH reactivity during CYPHEX, we expect the RO₂ production from the oxidation of HCs to be low. Hence the interference due to potential conversion of atmospheric RO₂ radicals into HO₂ radicals due to the NO injected to convert HO₂ into OH in the low-pressure detection volume is expected to be low compared to regions with high OH reactivities like the boreal forest in Finland. In order to reduce the conversion of RO₂ to HO₂, we used a reduced NO flow of 0.5 sccm resulting in ∼7 × 10^12 cm⁻³ of NO in the detection cell, thus converting only about 30 % of HO₂ radicals to OH radicals while simultaneously reducing the RO₂–HO₂ conversion efficiency. We did not conduct experiments to measure the conversion efficiencies of all possible RO₂ radicals in the atmosphere. To estimate the potential interference due to RO₂ radicals on our measured signal, we made model calculations using CAABA/MECCA in which most of the RO₂ radicals from higher HCs directly form HO₂ radicals after reaction with NO, skipping the reaction step of alkoxy radicals with O₂, which is slower at reduced pressure inside the instrument compared to ambient atmosphere.

The model is run at ∼4 hPa to see how OH and HO₂ radicals evolve with time in the low-pressure detection volume at different NO concentrations and is validated for calibration conditions (manuscript under preparation). The RO₂ radicals in the model are initialized with the concentrations generated from the model run for the base case (case III in Fig. 3) for our study while OH and HO₂ radicals are initialized with measured concentrations. In Fig. S2a, we show the evolution of OH and HO₂ radicals inside the detection cell after injection of NO (t = 0) for NO concentrations of 7.1 × 10¹⁴ and 1.71 × 10¹⁴ cm⁻³ respectively. The converted OH signal is detected after 6.6 ms (time of detection). For the high conversion efficiency case, the contribution of RO₂ to the OH signal at 6.6 ms is about 35 % or 31 % of the initial RO₂ mixing ratio. This value matches with estimates from a previous study by Hens et al. (2014). For the low conversion efficiency case that represents the CYPHEX measurement mode, the estimated contribution of RO₂ to the measured signal is about 12 % or 2.5 % of the initial RO₂ mixing ratio.

Further, we estimate that more than 50 % of the RO₂ interference is due to isoprene oxidation products. This is due to the fast conversion of isoprene-based hydroxy peroxy radicals towards HO₂ and OH. These isoprene-based hydroxy peroxy radicals have one of the largest conversion efficiencies of up to 90 % (Fuchs et al., 2011; Lew et al., 2018). Moreover, we see that the RO₂ interference does not increase with increasing terpene concentrations and is nearly constant at terpene levels greater than 80 pptv (Fig. S2b), mostly because during the course of the day HO₂ concentration increases during this time faster than terpene-based RO₂ concentration. This indicates that the RO₂ interference effects cannot explain the deviation of modelled HO₂ with respect to measurements at high terpene mixing ratios (discussed later in Sect. 3.3 and 3.4).

Calibration of the instrument for OH and HO₂ measurements is achieved by measuring the signals generated by known amounts of OH and HO₂ (Martinez et al., 2010) in a calibrator set-up. The calibrator was mounted on top of the OH inlet without the IPI. Known amounts of OH and HO₂ were produced by irradiating different concentrations of humidified air with 185 nm radiation produced by a Pen-Ray Hg lamp. The actinic flux density of the Hg lamp (Pen-
Ray line source, LOT-Oriel, Germany) used for the photolytic radical production was determined before and after the campaign using the actinometry method by N$_2$O photolysis (Martinez et al., 2010). Different H$_2$O mixing ratios were produced by mixing different combinations of humidified and dry air flows using mass flow controllers. The humidified air was generated by bubbling dry air into a container half filled with water maintained at 25–30 °C. The water mixing ratio in the humid air stream was measured with a LI-COR CO$_2$/H$_2$O-analyzer (Li-7000) based on the detection of differential absorption with an infrared spectrometer. The stability of the Li-7000 was ensured by calibrations with a dew point generator (Li-610 from LI-COR).

The precision and minimum detection limit for OH measurements during CYPHEX were determined from the variability in the background signals (Table 1). The chemical background, during propane injection, being larger than the spectroscopic background (off resonance), has a dominating impact on the precision. The precision for OH and HO$_2$ measurements was calculated to be 4.8 × 10$^{5}$ molecules cm$^{-3}$ and 0.39 pptv respectively for 4 min and 14 s time resolutions respectively. The accuracy for OH measurements was derived from the uncertainties in OH calibrations, which involve uncertainties in determination of lamp flux (traceable to a NIST spectroradiometer (Metcon GmbH) operating at 275–640 nm with a 2-π integrating hemispheric quartz dome. The spectroradiometer was calibrated prior to the campaign using a 1000 W NIST traceable irradiance standard. Photolysis frequencies were obtained with a charge-coupled device spectroradiometer (Metcon GmbH) operating at 275–640 nm with a 2-π integrating hemispheric quartz dome. The spectroradiometer was calibrated prior to the campaign using a 1000 W NIST traceable irradiance standard. Photolysis frequencies were calculated using molecular parameters recommended by the International Union of Pure and Applied Chemistry (IUPAC) and NASA evaluation panels (S. P. Sander et al., 2011; IUPAC, 2015). An automatic weather station (Vantage Pro2; Davis Instruments Corp., Hayward, CA) was used to measure temperature, pressure, wind direction and speed, solar radiation, and humidity with a time resolution of 1 min.

### 2.4 The CAABA/MECCA model

Due to the high reactivity of OH with a plethora of anthropogenic and biogenic HCs, the HO$_2$ chemistry of the atmosphere involves numerous reactions. To efficiently evaluate these reactions, several mechanisms of various complexity have been developed, e.g. the Regional Atmospheric Chemical Mechanism (RACM; Stockwell et al., 1997), the Master Chemical Mechanism (MCM; http://mcm.leeds.ac.uk, last access: 28 December 2017), the Mainz Isoprene Mechanism (MIM; Taraborrelli et al., 2009), the Carbon Bond Mechanism (Yarwood et al., 2005). For this study, we use the Module Efficiently Calculating the Chemistry of the Atmosphere (MECCA; S. P. Sander et al., 2011; R. Sander et al., 2005), which is an atmospheric chemistry module that contains a comprehensive set of gas- and aqueous-phase chemical reaction mechanisms covering tropospheric and stratospheric chemistry. The gas-phase chemistry in the present version contains 2664 species (including 40 dummy species to account for deposition) and 1670 reactions, including basic oxygen, hydroxyl, peroxy, alkyl and organic nitrogen-containing species due to the various mechanisms are indicated in Table 1. C$_2$–C$_4$ alkanes and alkenes (ethane: C$_2$H$_6$; ethene: C$_2$H$_4$; propane: C$_3$H$_8$; propene: C$_3$H$_6$; butane: C$_4$H$_10$ (i and n); butene: C$_4$H$_8$ (c and t)) were measured with a time resolution of 60 min and the mixing ratio represents an average over a sampling period of 20 min (Sobanski et al., 2016). Photolysis frequencies were obtained with a charge-coupled device spectroradiometer (Metcon GmbH) operating at 275–640 nm with a 2-π integrating hemispheric quart dome. The spectroradiometer was calibrated prior to the campaign using a 1000 W NIST traceable irradiance standard. Photolysis frequencies were calculated using molecular parameters recommended by the International Union of Pure and Applied Chemistry (IUPAC) and NASA evaluation panels (S. P. Sander et al., 2011; IUPAC, 2015). An automatic weather station (Vantage Pro2; Davis Instruments Corp., Hayward, CA) was used to measure temperature, pressure, wind direction and speed, solar radiation, and humidity with a time resolution of 1 min.
**Table 1.** Details of experiments conducted during CYPHEX (relevant to this study). PTR-TOF-MS: proton-transfer-reaction time-of-flight mass spectrometry. GC-FID: gas chromatography–flame ionization detector.

<table>
<thead>
<tr>
<th>Species</th>
<th>Measurement technique</th>
<th>Time resolution</th>
<th>Precision</th>
<th>Limit of detection</th>
<th>Total uncertainty</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydroxyl radical</td>
<td>LIF-FAGE-IPI</td>
<td>4 min</td>
<td>4.8 \times 10^5 molec cm^{-3}</td>
<td>1 \times 10^6 molec cm^{-3}</td>
<td>28.5 % (2σ)</td>
<td>Hens et al. (2014)</td>
</tr>
<tr>
<td>Hydroperoxyl radical</td>
<td>LIF-FAGE</td>
<td>14 s</td>
<td>0.4 pptv</td>
<td>0.8 pptv</td>
<td>36 % (2σ)</td>
<td>Martinez et al. (2010)</td>
</tr>
<tr>
<td>Ozone</td>
<td>UV photometry</td>
<td>1 min</td>
<td>2 ppbv</td>
<td>5 %</td>
<td></td>
<td>Li et al. (2015); Hosaynali Beygi et al. (2011)</td>
</tr>
<tr>
<td>Carbon monoxide</td>
<td>Room temperature Quantum cascade Laser spectrometer</td>
<td>1 s</td>
<td>0.4 ppbv</td>
<td>14.4 %</td>
<td></td>
<td>Li et al. (2015)</td>
</tr>
<tr>
<td>Nitric oxide</td>
<td>Chemiluminescence</td>
<td>5 s</td>
<td>5 pptv</td>
<td>20 pptv</td>
<td>20 %</td>
<td>Li et al. (2015); Hosaynali Beygi et al. (2011)</td>
</tr>
<tr>
<td>Nitrous acid</td>
<td>Long-path absorption photometry</td>
<td>30 s</td>
<td>4 pptv</td>
<td>10 %</td>
<td></td>
<td>Muesel et al. (2016)</td>
</tr>
<tr>
<td>Formaldehyde</td>
<td>Hantzsch reaction fluorescence</td>
<td>10 min</td>
<td>38 pptv</td>
<td>16 %</td>
<td></td>
<td>Kormann et al. (2016)</td>
</tr>
<tr>
<td>Hydrogen peroxide</td>
<td>Enzymatic reaction</td>
<td>10 min</td>
<td>14.9 pptv</td>
<td>16 %</td>
<td></td>
<td>Fischer et al. (2015)</td>
</tr>
<tr>
<td>Organic peroxides</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methyl hydroperoxide</td>
<td>High-performance liquid chromatography</td>
<td>12 min</td>
<td>25 pptv</td>
<td>9 %</td>
<td></td>
<td>This study</td>
</tr>
<tr>
<td>SO2</td>
<td>Chemical ionization mass spectrometer</td>
<td>10 mins</td>
<td>50 pptv</td>
<td>30 %</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C2−C4 (alkanes, alkenes)</td>
<td>GC-FID</td>
<td>60 min</td>
<td>&lt; 5 %</td>
<td>1–8 pptv</td>
<td>10 %</td>
<td>Sobanski et al. (2016)</td>
</tr>
<tr>
<td>Methane</td>
<td>GC-FID</td>
<td>1 min</td>
<td>2 %</td>
<td>20 ppbv</td>
<td>2 %</td>
<td>This Study</td>
</tr>
<tr>
<td>Isoprene</td>
<td>GC-MS</td>
<td>45 min</td>
<td>3.3 %</td>
<td>1 ppt</td>
<td>14.5 %</td>
<td>Destroff et al. (2017)</td>
</tr>
<tr>
<td>α-pinene</td>
<td></td>
<td></td>
<td>4.9 %</td>
<td>1 ppb</td>
<td>15 %</td>
<td></td>
</tr>
<tr>
<td>β-pinene</td>
<td></td>
<td></td>
<td>8.8 %</td>
<td>2 ppb</td>
<td>16.7 %</td>
<td></td>
</tr>
<tr>
<td>Limonene</td>
<td></td>
<td></td>
<td>4.2 %</td>
<td>1 ppb</td>
<td>14.7 %</td>
<td></td>
</tr>
<tr>
<td>Benzene</td>
<td>PTR-TOF-MS</td>
<td>1 min</td>
<td></td>
<td></td>
<td></td>
<td>Destroff et al. (2017)</td>
</tr>
<tr>
<td>Toluene</td>
<td></td>
<td></td>
<td>5.4 % (at ~ 280 pptv)</td>
<td>14 pptv</td>
<td>14 % (21 %)*</td>
<td></td>
</tr>
<tr>
<td>Isoprene oxid. pds.</td>
<td></td>
<td></td>
<td>4.9 % (at ~ 280 pptv)</td>
<td>12 pptv</td>
<td>14 % (20 %)*</td>
<td></td>
</tr>
<tr>
<td>Methyl ethyl ketone (MEK)</td>
<td></td>
<td></td>
<td>5.0 % (at ~ 260 pptv)</td>
<td>14 ppb</td>
<td>11 % (14 %)*</td>
<td></td>
</tr>
<tr>
<td>Methanol</td>
<td></td>
<td></td>
<td>3.8 % (at ~ 280 pptv)</td>
<td>16 ppb</td>
<td>11 % (16 %)*</td>
<td></td>
</tr>
<tr>
<td>Acetaldehyde</td>
<td></td>
<td></td>
<td>2.5 % (at ~ 2800 pptv)</td>
<td>242 ppb</td>
<td>37 % (41 %)*</td>
<td></td>
</tr>
<tr>
<td>Acetone</td>
<td></td>
<td></td>
<td>2.2 % (at ~ 1300 pptv)</td>
<td>85 ppb</td>
<td>22 % (27 %)*</td>
<td></td>
</tr>
<tr>
<td>Acetic acid</td>
<td></td>
<td></td>
<td>1.4 % (at ~ 2500 pptv)</td>
<td>97 ppb</td>
<td>10 % (17 %)*</td>
<td></td>
</tr>
</tbody>
</table>
| with high isoprene mixing ratios (< 1 nmol mol^{-1}) and low NO (< 40 pmol mol^{-1}) (Taraborrelli et al., 2012). MOM was first presented and used in the study by Lelieveld et al. (2016). It represents the gas-phase oxidation of more than 40 primarily emitted VOCs. The structure and the construction methodology mirrors that of the MCM. The oxidation mechanism for aromatics has been presented in Cabrera-Perez et al. (2016). The terpene oxidation scheme includes previous developments for modelling HOx field measurements (Taraborrelli et al., 2012; Hens et al., 2014; Nolscher et al., 2014). Most of the known and/or proposed HOx-recycling mechanisms under low-NO conditions are taken into account. Finally, isoprene chemistry follows, to a large extent, Peeters et al. (2014) and Jenkin et al. (2015), with modifications by Nolscher et al. (2014). Chemistry of the pinenes (monoterpenes) is a reduction of the MCM with modifications proposed in the past by Vereecken et al. (2007), Nguyen et al. (2009), Vereecken and Peeters (2012), and Capouet et al. (2008). For the numerical integration, MECCA uses the KPP software (Sandu and Sander, 2006). All gas-phase reactions are contained in a single chemical file (gas.eqn).

The most common method to efficiently evaluate a complex chemical environment is to use a zero-dimensional, photochemical box model. Here, we use version 3.8 of the box model CAABA (Chemistry As A Boxmodel Application), www.atmos-chem-phys.net/18/10825/2018/ Atmos. Chem. Phys., 18, 10825–10847, 2018
which is based on version 3.0 described by R. Sander et al. (2011). To apply the MECCA chemistry to atmospheric conditions, it is connected to the CAABA base model via the MESSy interface. The model is constrained by observed concentrations of the main reactive trace gases, O$_3$, CO, NO, NO$_2$, nitrous acid (HONO), HCHO, hydrogen peroxide (H$_2$O$_2$), methyl hydroperoxide (MHP, CH$_3$OOH), SO$_2$, C$_2$–C$_4$ alkanes and alkenes, isoprene, pinene (α and β), limonene, benzene, toluene, methane, methanol, acetonitrile, acetaldehyde, acetone, acetic acid, and photolysis rate constants. Model results were obtained by letting the model run into steady state for OH and HO$_2$ for each set of data points. The steady state was set to be achieved for OH when the relative changes in OH were less than 5 x 10$^{-7}$. Deposition velocities used in our scheme for some important species are provided in Table 2. The deposition is incorporated into the model scheme by converting a species into a dummy species according to its deposition velocity. As these dummy species will not participate in any further chemical reaction, their precursors are effectively removed from the model chemical scheme. The deposition velocities of peroxides and formic and nitric acid are based on Nguyen et al. (2015) while those for PINAL and PAN are based on Evans et al. (2000), using the average of the deposition rates for water, forest, and grass. The model scheme is included in the Supplement.

### 2.5 The FLEXPART model

In order to understand the impact of different emission sources on the atmospheric processing of air masses, the dynamical transport history of air parcels reaching Cyprus during the CYPHEX campaign was traced using FLEXPART 9.2 (Stohl et al., 2005). FLEXPART is a Lagrangian particle dispersion model that describes the transport and diffusion of tracers by computing the trajectories of a large number (ca. 10,000) of infinitesimally small tracer particles. For CYPHEX, trajectory simulations were performed at 3 h time intervals during 21–31 July 2014. For this, FLEXPART was run 120 h backward in time from the measurement site driven with analyses from ECMWF with 0.2° × 0.2° horizontal resolution and a temporal resolution of 1 h (Hüser et al., 2017). The particle density distribution of the tracer particles during this 12 h backward simulation provided information on their residence time in each grid cell of the defined geographical area (Fig. S3). Major transport routes of air reaching the site during CYPHEX were identified from the grid cells with higher residence times.

### 3 Results and discussion

#### 3.1 HO$_2$ measurements during CYPHEX and associated meteorological and chemical parameters

The variation in OH and HO$_2$ along with a few important chemical and radiation parameters relevant to HO$_2$ chemistry during 21–31 July (day of the year: 202–212) are shown in Fig. 2. The mean values of OH and HO$_2$ during 21–31 July were 2.2 × 10$^6$ and 2.87 × 10$^8$ molecules per cm$^3$ respectively. The mean OH concentration during peak noon hours ($J(O^1D) > 2 \times 10^{-5}$ s$^{-1}$) was 5.75 × 10$^6$ molecules per cm$^3$. While both OH and HO$_2$ showed a clear diurnal variation, the HO$_2$ / OH ratio decreased with enhancement in $J(O^1D)$, and was close to 100 at $J(O^1D) = 3 \times 10^{-5}$ s$^{-1}$. During the MINOS campaign in the summer of 2001 on Crete, an island in the central Mediterranean at similar latitude about 700 km west of the CYPHEX site, average OH levels were measured to be 3.6 – 6.7 × 10$^6$ molecules per cm$^3$ (Berresheim et al., 2003). The peak OH levels during MINOS reached twice the peak values observed during our study on Cyprus. During MINOS, OH peaked at about 2.1 × 10$^7$ molecules per cm$^3$, with O$_3$ and $J(O^1D)$ of about 60 ppbv and 2.8 × 10$^{-5}$ s$^{-1}$. These O$_3$ and $J(O^1D)$ values are not very different from CYPHEX values, indicating that the peak primary OH production would not be very different for MINOS and Cyprus. Although HO$_2$ was not measured during MINOS, the CO levels (mean values reaching close to 160 ppbv) were much higher (Heland et al., 2003) compared to CYPHEX, indicating the possibility of higher HO$_2$. Nevertheless, the absence of HO$_2$ measurements during MINOS impairs our ability to directly compare the OH chemistry from the two datasets for a regional perspective. However, the peak OH levels observed during CYPHEX are comparable to the midday OH concentrations predicted for the Finokalia Aerosol Measurement Experiments (FAME-08) at a remote coastal site on the island of Crete, Greece (Hildebrandt et al., 2010). The FAME measurements showed the presence of aged, highly oxygenated organic aerosols (OAs) during summer compared to winter despite being heavily influenced by continental air masses (53%) during summer and aged marine air masses (61%) during winter. The presence of highly oxygenated OA during summer was attributed to strong photochemical impacts due to higher O$_3$ and UV levels during summer, leading to double the values of midday OH concentrations during summer compared to winter (Hildebrandt et al., 2010).

While OH measurements during CYPHEX started on 12 July 2014 (day of year: 193), systematic HO$_2$ measure-
Figure 2. Measurements of OH, HO₂, and selected chemical and radiation parameters relevant to HOₓ chemistry during the CYPHEX campaign. Time is in UTC. Local time in Cyprus during summer is UTC+3.

3.2 Modelling simulations of CYPHEX HOₓ

Figure 3 shows simulations of OH and HO₂ using CAABA/MECCA in comparison with the measurements in ambient air. Due to the remote location of the measurement site, it is expected that the HOₓ chemistry would be representative for the background Mediterranean atmosphere. In general, NO levels were low during CYPHEX, with only ~2% of data above 1 ppbv NO and ~5% above 100 pptv of NO. The study is limited to NO below 100 pptv to exclude measurements affected by emissions from the local diesel generator and local automobile traffic. Under very remote conditions, the steady state of HOₓ would be achieved from a balance between HOₓ production (Reactions R1–R3) and HOₓ loss (Reaction R10). However, such a simplified condition is rarely achieved in the boundary layer as the presence of HCs leads to additional production and loss channels. A first model run was carried out constraining only O₃, CO, CH₄, NO, NO₂, HONO, HCHO and H₂O₂, H₂O, and photolysis frequencies to measured values (Fig. 3, case I). Thus, the major primary production channels for OH and HO₂ via photolysis of O₃, HONO, and HCHO are constrained by measured quantities, while the recycling of OH and HO₂ occurs via reactions with O₃, CO, CH₄, and NO only. However, in the absence of the contribution from important HCs, OH loss is underestimated, leading to an over-prediction of OH on the order of 88%. Further, HO₂ is also overestimated, on aver-
Figure 3. LIF-FAGE measurements of OH (a–d) and HO$_2$ (e–h) vs. model simulations with CAABA/MECCA. From left to right: model simulations using simple chemistry without NMHCs (case I), simulations including anthropogenic and biogenic hydrocarbons except terpenes (case II), after adding terpenes (case III), and after initializing with zero terpenes but with CH$_4$ increased to account for the additional measured terpene reactivity (case IV).

age, by more than 30% in this case (Fig. 3, case I) as loss via formation of hydroperoxides is under-represented due to the incomplete VOC representation, leading to too low RO$_2$ concentrations.

When we constrain the model with all measured species, the OH simulations are much improved with a model-to-measured ratio of 0.90 (Fig. 3 case III). However, HO$_2$ is now under-predicted by about 17%. The under-prediction of HO$_2$ is enhanced with increasing terpene ($\alpha$-pinene, $\beta$-pinene, and limonene) mixing ratios, with an average HO$_2$ model/measurement ratio of 0.72 for $>80$ pptv of measured terpenes. When terpenes are not included in the model, the HO$_2$ simulations are much improved (model/measured = 1.02). The mean daytime OH reactivity calculated for the measured major terpenes was about 0.06 s$^{-1}$. This value is only 3.7% of the calculated reactivity from the total of all measured species and about 10.7% of the OH reactivity ($k_{\text{OH}}$: Eq. 1) from CO. The OH reactivity calculated from the trace gases measured during CYPHEX varied between 1 and 2 s$^{-1}$ during the study period (Fig. 4), which is comparable to reactivity measurements made in the free troposphere (Mao et al., 2009). During the TORCH-2 campaign in Weybourne, England, and the DOMINO campaign in El Arenosillo, Spain, the total reactivity was generally < 5 s$^{-1}$ when air masses originated from the sea (Lee et al., 2009; Sinha et al., 2012). The OH reactivity measured in suburban regions is generally higher than 5 s$^{-1}$ (Yang et al., 2016). During CYPHEX, CO and CH$_4$ constituted more than half of the total calculated reactivity (Fig. 4). CO, CH$_4$, HCHO, C$_5$H$_8$, and CH$_3$CHO accounted for 35%, 17%, 13%, 9%, and 8% respectively of the calculated OH reactivity for daytime values during CYPHEX.

\[
k_{\text{OH}} = \sum (k_{\text{species}} \times C_{\text{species}}), \tag{1}
\]

where $k_{\text{species}}$ is the rate coefficient for reaction of a given species with OH and $C_{\text{species}}$ is the concentration of the same species.

As limonene chemistry is not included in the current model scheme, it is accounted for in the model in the form of additional $\alpha$-pinene after normalizing for their OH reactivities. When limonene is included as CH$_4$ instead of $\alpha$-pinene, the OH loss increases by about 1% compared to case II in which there were no terpenes in the model, which confirms the average OH reactivity of limonene at 0.02 s$^{-1}$, i.e. an additional 1% of the calculated reactivity from the measured species. Although limonene chemistry is different from $\alpha$-pinene in that it caters more towards secondary organic aerosol (SOA) formation, addition of limonene as $\alpha$-pinene increases HO$_2$ and OH losses by 3%–6%, which is well within the uncertainty of the measurements. Nevertheless, the degradation products from terpene oxidation (shown in Fig. 5) further act as sources and sinks for OH and HO$_2$, hence playing an important role in the overall HO$_x$ chemistry (Calogirou et al., 1999; Librando and Tringali, 2005;
Figure 4. Calculated OH reactivity of various chemical species measured during CYPHEX.

Figure 5. Contribution of different terpene-related reactions to the total HOX loss. Additional peroxy radicals are produced in the model by terpenes rather than organics, e.g., isoprene degradation. Pinene includes α-pinene and β-pinene. TerpRO2 includes only C8H13O3 (C85O2), C9H15O3 (C96O2), C9H15O4 (C97O4), C10H15O5 (C98O2), C10H15O5 (C106O2), C10H15O4 (C107O2), C6H9O5 (C104O2), C6H9O4 (PinOOH), and C5H7O4 (BPinOOH). TerpROOH includes only C10H16O4 (PinalOOH), C9H14O3 (C85OOH), C9H14O4 (C86OOH), C9H16O4 (C97OOH), C9H16O5 (C98OOH), C10H16O5 (C106OOH), and C9H5O4 (C511OOH). The names in brackets are MCM nomenclature. Only those species are selected whose contribution is at least 0.25% to the HOX sink.

Zhang et al., 2015; Wisthaler et al., 2001). Further, increasing or decreasing the terpene concentrations by their measured uncertainties (15%; Table 1) decreases or increases the OH and HO2 model/measured slope by 2%–3%, which is close to the errors on the regression slopes. Also, changing other gases like CO and C3H8 by their uncertainties (Table 1) has less than 3% impact on the model vs. measured slopes for OH and HO2. Nevertheless, when terpenes are included as CH4, both OH and HO2 simulations are much improved (Fig. 3, case IV). In this case, the terpene chemistry does not play any role in OH–HO2 simulations but the primary OH reactivity of terpenes is taken into account.
To access the model uncertainty, Monte Carlo simulations are carried out by selecting 228 data points from the input dataset, representing at least two occurrences of concentrations, i.e. 1–15, 25–75, and 85–99 percentiles of O₃, CO, NO, HCHO, HONO, C₅H₅, C₁₀H₁₆, and J(O(1D)). For each of these points, we performed 9999 Monte Carlo simulation runs. The Monte Carlo simulations are based on the method described in R. Sander et al. (2011), but additionally we varied the boundary conditions of the measured species by their ±1σ uncertainty (Table 1). The derived overall model uncertainty at 1σ is 17.4% and 10.5% for OH and HO₂ respectively.

Review studies show agreements between in situ observations of OH and HO₂ and box model calculations using different chemical schemes within a factor of 2 for both urban and marine (boundary layer) environments (Stone et al., 2012). During the North Atlantic Marine Boundary Layer Experiment (NAMBLEX) field campaign in the summer of 2002 at the Mace Head Atmospheric Research Station, OH was under-predicted early in the morning and in the late afternoon–early evening and over-predicted in the middle of the day based on simple steady-state calculations (Smith et al., 2006). However, photolysis of HONO was not considered due to the absence of HONO measurements during NAMBLEX. The authors contemplate that HONO photolysis and OH recycling could play crucial roles in explaining the discrepancy between measured and calculated OH during periods with a high solar zenith angle. Despite low-isoprene regimes (Lewis et al., 2005), inclusion of dimethyl sulfide (DMS) and C₅H₅ improved the calculated / measured ratio for OH from 1.13 ± 0.36 to 0.94 ± 0.39 during NAMBLEX. HO₂ was over-predicted but model performance improved by incorporating reactions with BrO, IO, and aerosol uptake; the best-case scenario for HO₂ modelled / measured was 1.26±0.36. During the Intercontinental Chemical Transport Experiment-A (INTEX-A), OH was predicted fairly well for the lowest 2 km over North America and the western Atlantic Ocean using a zero-dimensional, time-dependent photochemical box model developed at NASA Langley Research Center (Ren et al., 2008). In the study, HO₂ was over-predicted (model / measurement = 1.37). It is challenging to put the model–measurement agreement during CYPHEX in perspective with other studies because they vary with respect to chemical regimes, i.e. different amounts of NOₓ and HCs as well as the chemical mechanisms employed to simulate the HO₂ chemistry. Rohrer et al. (2014) compared the HO₂ chemistry from different field campaigns using a common modelling framework and point to buffering mechanisms that maintain OH in various NOₓ and VOC regimes. Ren et al. (2008) have noted that despite highly constrained measurement suites during various field campaigns, there are significant discrepancies between model predictions and actual measurements of OH and HO₂ in different environments. One possible reason for the discrepancy could be due to unmeasured atmospheric constituents. However, during CYPHEX we have been able to measure the major primary sources and sinks related to HOₓ chemistry, and simulations agree within uncertainty of OH and HO₂ measurements. Another plausible reason for discrepancy is the uncertainty in mechanistic pathways, branching ratios, and rate constants, etc., which are not known with sufficient accuracy. A recent study comparing seven different chemical mechanisms constrained with the same input data and boundary conditions revealed 25%–40% ambiguity in predictions of HO₂ and OH over the United States (Knote et al., 2015). As we highlight in the next section, the HO₂ chemistry related to terpenes is also associated with substantial uncertainty.

3.3 Reasons for decreased model accuracy when including the terpene chemistry

As discussed in the last section, we find that the under-prediction in HO₂ gradually increases with increasing terpene levels, being about 38% for the highest terpene levels observed (about 120 pptv). To ascertain if this under-prediction is related to terpene chemistry or some other chemical pathway, we compare two model runs, namely case III vs. case IV in Fig. 3. Case IV is achieved by initializing the model with zero terpenes, however keeping the terpene reactivity towards OH intact. The terpene reactivity, i.e. the product of individual terpene concentrations with their rate coefficient towards OH, is approximated in the model scheme in the form of additional CH₃OH (Fig. 3, case IV). Under this scenario, we find that the agreement between modelled and observed HO₂ again improves, to levels similar to case II without terpenes in the model (Fig. 3). The small difference of 6% with respect to OH prediction between cases II and IV (Fig. 3) also corroborates the minor primary reactivity of < 0.06 s⁻¹ for the sum of α-pinene, β-pinene, and limonene towards OH. However, up to 38% under-prediction in HO₂ for terpene concentrations over 80 pptv indicates the important role of secondary products formed during terpene oxidation in the HO₂ budget.

To quantify the magnitude of the impact of terpene degradation products on the HO₂ budget, we compared the HO₂ sinks from the two model cases: case III (Fig. 3, henceforth “terpene case”) and case II (Fig. 3, henceforth “isoprene case”), the difference being the presence and absence of α-pinene, β-pinene, and limonene. The estimation of HO₂ sinks from the two model cases is conspicuously marked by significant contribution of terpene degradation and oxidation products in the terpene case which are absent in the isoprene case. The additional reactions in the terpene case make up 6.3%–27.8% of the total HO₂ sinks, the contributions increasing with the terpene concentrations and peaking between 13:00 and 14:00 LT on clear days (Fig. 5). While the terpene degradation products contribute, on average, 15% to the total HO₂ loss, this value is 20.5% for terpenes > 80 pptv. The corresponding contribution of terpene-related peroxy radicals to the total HO₂ loss is 7.8% and
10.3% respectively, accounting for about 52% of the impact of the terpene degradation products, including peroxy radicals, pinonal, norpinonal, and peroxides, towards HO₃ loss. 

Since only about 3.7% of primary terpene reactivity with OH results in up to 28% of HO₂ loss, with terpene-generated peroxy radicals having more than 4 times the impact of the primary terpenes on the HO₂ loss, we examine a sequence of reactions starting with the oxidation of α-pinene to understand the genesis and propagation of these terpene-generated peroxy radicals (Reactions R15–R23). The reactions show that oxidation of α-pinene via OH and O₃ directly generates several peroxy radicals (Reactions R15–R16), which are sinks for HO₂ via their reactions with HO₂-forming hydroperoxides (e.g. Reactions R21–R22). These first-generation peroxy radicals also form other peroxy radicals (Reactions R17–R20), and the chain propagates to produce additional peroxy radicals contributing via formation of the respective hydroperoxide in reaction with HO₂, leading to overall HO₃ loss (e.g. Reaction R23).

Thus, a cascade of HO₃ scavengers mostly in the form of RO₂ species, at low NOₓ levels, starting with oxidation of primary terpenes, leads to nearly 28% HO₂ loss, resulting in up to 38% under-prediction of HO₂. The losses increase with increases in terpene levels. The major contribution towards the HO₃ loss comes from terpene-generated organic peroxy radicals (Fig. 5). However, there are major uncertainties regarding the chemical pathways and rate coefficients governing the chemistry of these peroxy radicals in the actual atmosphere and in the next section we examine several postulates based on the literature.

### 3.4 Terpene-generated organic peroxy radicals

A limitation in our understanding of peroxy radical oxidation pathways is the rate constant of peroxy radicals with NO. For most peroxy radicals, lumped rate coefficients are defined by the expression Eq. (2) (Rickard and Pascoe, 2009) as kinetic data are unavailable for most of the RO₂–NO reactions.

\[
K_{RO}\text{NO} = 2.54 \times 10^{-12} \exp(360/T) \cdot f
\]

Thus, \(K_{RO}\text{NO}\) is the product of the rate coefficient and an efficiency factor, \(f\). The values of \(f\) for different classes of RO₂ are given in Jenkin et al. (1997). The reaction of RO₂ with NO can proceed via two channels, one forming alkyl nitrate (RONO₂) and the other forming RO and NO₂ (Reactions R12–R13). Enhancing the RO₂–NO rate coefficient suppresses the peroxide-forming channel via HO₂. For our study period, increasing the \(K_{RO}\text{NO}\) even by a factor of 4 led to only a 6.8% increase in modelled HO₂ levels for terpenes > 80 pptv (Fig. 6).

During the summertime 1999 Southern Oxidants Study (SOS) at Nashville, Tennessee, agreement could be found between photochemical models and in situ measurements when the product of the branching ratio and rate constant for organic peroxide formation, via Reaction (R11), was reduced by a factor of 3–12 (Thornton et al., 2002). To test this hypothesis, we decreased the \(K_{RO}\text{HO₂}\) for terpenes by half and by one-fifth of its value considered in the model. The rate coefficient of RO₂ with HO₂ leading to ROOH is given by Eq. (3), where \(K_{RO}\text{HO₂}\) increases with the size of RO₂ radicals (Atkinson et al., 1999; Boyd et al., 1996) (http://mcm.leeds.ac.uk/MCMv3.3.1/categories/saunders-2003-4.6.4-gen-master.html?rnId=6942, last access: 28 December 2017).

\[
K_{RO}\text{HO₂} = 2.91 \times 10^{-13} \exp(1300/T) \\
\left[1 - \exp(-0.245n)\right]
\]

where \(n\) is the carbon number and \(T\) is the temperature.
Figure 6. HO$_2$ predictions using different chemical schemes which were employed to investigate the impact of peroxy radicals as a HO$_2$ sink. The x axis shows the sum of measured α-pinene, β-pinene, and limonene. The base case is the model simulation using measured species; $4 \times K_{RO2:NO}$ is a simulation in which the rates of selected terpene-related RO$_2$ species with NO are increased by a factor of 4; $0.2 \times K_{RO2:HO2}$ is a simulation in which the rates of these RO$_2$ species with HO$_2$ are reduced by a factor of 0.2; the autoxidation simulation includes an autoxidation scheme rapidly converting selected RO$_2$ species into ROOH. The last scheme refers to a simulation in which $K_{RO2:R’O2}$ is set to $4 \times 10^{-10}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ (close to the collision limit). The mean and sigma are calculated for 100 pptv bins along the x axis.

We find that decreasing the rate coefficient by a factor of 5 leads to better reconciliation with our observations than when decreasing it by a factor of 2. When $K_{RO2:HO2}$ was reduced by a factor of 5, the modelled HO$_2$ increased by only 9.9 % for higher terpene levels, i.e. α-pinene + β-pinene + limonene > 80 pptv (Fig. 6).

Recent studies have proposed inter- and intra-molecular hydrogen abstraction by peroxy radicals (autoxidation) to be effective reaction pathways leading to OH formation (Crounse et al., 2013). This abstraction is largely determined by the thermochemistry of the nascent alkyl radicals and thus is strongly influenced by neighbouring substituents; hence the rate increases rapidly when more oxygen-containing functional groups are involved. The H-shift rates reported in literature can be as fast as 0.1 s$^{-1}$ for atmospheric conditions. We incorporated a simple autoxidation scheme for some of the most important terpene-related RO$_2$ species as specified in Fig. 5 in our model. In this scheme, the specific terpene-generated RO$_2$ species whose contribution was at least 0.25 % to the HO$_x$ sink are converted into a corresponding ROOH at a very fast rate ($1 \times 10^4 \times K_{RO2:HO2}$), reaching up to 0.2 s$^{-1}$; e.g. C$_{10}$H$_{15}$O$_4$ (PINALO2) is converted to C$_{10}$H$_{16}$O$_4$ (PINALOOH). We find that this scheme is effective in increasing the model HO$_2$ levels by 24.2 % for terpene levels > 80 pptv (Fig. 6). Since we have used a very simple approach to test the impact of autoxidation in the model atmosphere, more studies are required for a mechanistic development for incorporating these schemes, namely which way the H shift will occur at which rate, which products will be formed, etc. The present exercise in implementing the autoxidation scheme is only meant to show its importance, effectiveness, and potential in atmospheric chemistry models.

Further, when we greatly increased the RO$_2$–R’O$_2$ reaction rates to $4 \times 10^{-10}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ (close to the collision limit), we found that the discrepancy of the modelled HO$_2$ loss with respect to the measurements with increasing terpenes decreases (Fig. 6) and the overall agreement between modelled and measured HO$_2$ is still good (Figs. 6, S4, and S5). The large deviation between observations and modelled HO$_2$ around midday on days 205, 208, and 210 occurs during periods with high terpene concentrations. On days 205 and 208, the simulation with the autoxidation scheme shows much better agreement with observations compared to the base case (Case III in Fig. 3) while on day 210, when terpene concentrations are much higher, even the autoxidation scheme fails to reproduce the HO$_2$ observations. In this case the simulation in which the rate coefficient of RO$_2$–R’O$_2$ reactions was increased close to the collision limit shows a much better agreement compared to both the base case and the autoxidation case.

3.5 HO$_x$ chemistry during CYPHEX: production and loss estimates

Using the optimal scenario from our model results, i.e. the one in which all measured species are included in the model (case III, Fig. 3), we have studied different processes impact-
ing HO$_x$ budgets. Since impact of terpene-generated RO$_2$ species was maximum on day 210 (29 July), we will ignore any special features on this day for our subsequent discussions using model case III, the terpene case. For CYPHEX, the major OH- as well as HO$_x$-producing channel is the reaction of atmospheric water vapour with O$_1^D$ generated from the photolysis of O$_3$ (Fig. 7). Peak daytime contributions of this channel towards OH production exceeded 45% for most of the days and about 60% on days 205, 208, and 209. The midday values coinciding with peak OH production on day 205 were marked by a conspicuous influence of aged air masses originating over southwestern Europe and considerably processed over the Mediterranean before reaching the site. The peak HO$_2$ values on this day were about 11% lower than the average peak HO$_2$ values during the study period. The peak HO$_2$ value for $J$(O$_1^D$) > 2.5 × 10$^{-5}$ s$^{-1}$ was 6.4 × 10$^8$ molec cm$^{-3}$ for the study period while this value was only 5.7 × 10$^8$ molec cm$^{-3}$ on day 205. While, on average, the recycled OH via reactions of HO$_2$ with NO and O$_3$ (Reactions R6–R7) contributed 33.6% to the total OH production, this value was about 6% lower for days 205–206. It may also be noted that although O$_3$ was very low on both days 205 and 206, with predominant influence of aged marine air, the contribution from O$_1^D$ + H$_2$O to the total OH production still exceeded 50%. Due to lower HONO mixing ratios, the fractional contributions of HONO photolysis towards peak OH production during midday on day 208 were significantly low at about 2.5%–3.5%. On all other
days, for which values are available, this channel contributed more than 6 % to peak OH production during noontime. The photolysis of HONO has the largest fractional contribution to the early morning OH production on day 211, reaching above 30 %. Overall during CYPHEX, the average daytime ($JO^1D > 0$) contribution to OH production from O$_3$ photolysis and subsequent reaction of O$_3^1D$ with water vapour was about 39.1 %, the average daytime contribution from HONO photolysis was 12.3 %, and recycled OH from reaction of HO$_2$ with O$_3$ and NO accounts for 15.2 % and 18.4 % of the total OH production respectively. The four major OH-producing channels (Fig. 7) contribute up to 95 % of daytime OH production on most occasions, with 85.3 % on average. In addition, the reactions of acyl peroxy radicals (RCO$_3$) with HO$_2$ contribute about 3.1 % to the OH production, while the photolysis of H$_2$O$_2$ and the ozonolysis of pinene contribute 1.95 % and 2 % respectively.

The single major sink of OH during CYPHEX was CO, followed by CH$_4$, HCHO, C$_5$H$_8$, CH$_3$CHO, and O$_3$, on average accounting 20.9 %, 10.0 %, 7.8 %, 5.1 %, 4.9 %, and
4.1 % of OH losses respectively. The reactions of various peroxides with OH to form peroxy radicals, e.g. Reaction (R24), contribute 8.4 % to the OH loss. Further, oxidation of CH$_3$OH, pinal, C2–C4 alkenes, $\alpha$-pinene, HO$_2$, NO$_2$, and CH$_3$O$_2$ by OH contributed 2.7 %, 2.4 %, 2.4 %, 2.2 %, 1.6 %, 1.4 %, and 1.4 % respectively. During days 205–206, the modelled OH loss (Fig. 7) as well as the calculated OH reactivity (Fig. 4) were the lowest of the study period. It is likely that the air masses arriving at the site during these days were already much processed, spending considerable time over the Atlantic and Mediterranean, leading to depleted OH reactivity. During this period of marine influence, the contribution of long-lived gases to the daytime OH loss increased by about 15 % while the contribution of shorter-lived gases like HCHO and CH$_3$CHO decreased by 34 % and 39 % respectively. HCHO concentrations are observed to be lowest between days 205 and 206, when the marine influence was the highest. The average lifetime of HCHO for the study period is calculated to be 0.86 days, and its concentrations are higher whenever air masses are influenced by southeastern Europe/Black Sea region.

The major source of HO$_2$ during CYPHEX was the oxidation of CO by OH, contributing on average 36.2 % of daytime HO$_2$ production values. On days 205 and 206, when the site experienced aged oceanic air masses and CO concentrations were the lowest of the study period, the peak contribution of CO to daytime HO$_2$ production was still about 40.4 %. This shows that the contribution from other HO$_2$ precursors like HCHO decreases much faster than CO because of their shorter lifetime as the air mass passes over the ocean, and because they are isolated from their emission sources. The contributions of HCHO towards HO$_2$ production varied between 8 % and 23 % with an average daytime contribution of 15.6 % and included oxidation via both OH and photolysis. The methoxy radical (CH$_3$O), formed during oxidation of CH$_4$ and CH$_3$O$_2$, was also a significant source of HO$_2$, with peak contributions reaching 23 % on some occasions and an average daytime contribution of 17.8 %. On average, the reactions of HO$_2$ with NO and O$_3$, recycling OH, contributed 23.3 % and 18.2 % respectively towards daytime loss for HO$_2$. HO$_2$ losses by its self-reactions and reactions with CH$_3$O$_2$ reached peak daytime values of over 40 % and on average accounted for 30.3 % of daytime HO$_2$ losses. Among these, the major contribution was from the HO$_2$ self-reactions, with peak values of over 30 % during most days and average values of 22.4 %. The reactions of peroxy radicals (RO$_2$ and RCO$_3$) with HO$_2$ contribute 24.6 % to the HO$_2$ radical loss, resulting in increased under-prediction of HO$_2$ by the model with increasing terpene concentrations (Fig. S6). HO$_2$ losses due to recycling via NO and O$_3$ and the HO$_2$ losses via reactions with itself and CH$_3$O$_2$ are complementary during a diurnal cycle. The self-reactions gain in importance and peak around midday as the peroxy radi-
cal concentrations increase with increasing photochemistry, while recycling reactions dominate during periods with a larger solar zenith angle. During CYPHEX, for small solar zenith angles with $J(\Omega_{1D}) > 2 \times 10^{-3}$ s$^{-1}$, the contribution of recycling and self-reactions to HO$_2$ loss was 36.2% and 32.7% respectively. Overall, it is found that the lowest HO$_2$ production and losses occurred in the highly processed air masses influenced by the marine boundary layer, mostly over the Mediterranean Sea, but sometimes as far as the Atlantic Ocean (days 205–206), with some influence from southwestern Europe/northern Africa (Fig. 7b and d).

### 3.6 Impact of NO$_x$ levels on HO$_x$ cycling

The close proximity of Cyprus to several countries with different socio-economic conditions makes its air quality vulnerable to the increases or decreases in primary and secondary air pollutants in one of these countries in Europe, Asia, and Africa. For example, increased NO$_x$ emissions in one of the southern European countries or high CO from forest fires in Ukraine can be quickly transported to Cyprus in a matter of hours to days and modulate the atmospheric processing over the region. Increased NO$_x$ emissions can impact the O$_3$–NO$_x$–VOC chemistry by changing the HO$_x$ cycling. Even though the study period was characterized by NO of mostly less than 100 pptv, decreasing NO$_x$ by 30% still decreased OH levels by 14%, these impacts being higher in the early morning and late afternoon. Conversely, increasing NO by 5 and 10 times caused 44% and 70% average enhancement in OH levels.

Because of its important role in O$_3$ production, NO$_x$ emissions have been the focus of air quality investigations during preceding decades. Apart from their crucial role in O$_3$ production, NO$_x$ levels also play a significant role in the self-cleaning capacity of the atmosphere due to their reactions with atmospheric HO$_2$ and RO$_2$. We test the impact of mainland Europe NO$_x$ emissions on the self-cleaning capability of the climatically vulnerable Mediterranean atmosphere using the CYPHEX dataset. The recycling efficiency for OH (REF-OH) is defined as the ratio of OH produced from secondary sources via reactions of HO$_2$ with NO and O$_3$ (Reactions R6–R7) to the OH produced from primary and secondary sources. The primary sources of OH considered in our calculation include reaction of H$_2$O with O$_{1D}$ produced from O$_3$ photolysis and the photolysis of HONO, H$_2$O$_2$, and organic peroxides. The contribution of the primary OH production from HONO photolysis is corrected by subtracting off the contribution from the recombination reaction of OH with NO.

While the present study is limited to NO < 100 pptv, modelling results show that REF-OH increases with an increase in NO levels (Fig. 8). The REF-OH increases from 0.28 at 10 pptv of NO to 0.42 at around 30 pptv of NO to 0.7 at 100 pptv of NO (Fig. 8). The REF-OH of 0.7 corresponds to a chain length of 2.8. The relationship of REF-OH with NO indicates that at higher NO (not observed for the CYPHEX data), the REF-OH would increase further. To understand if REF-OH would reach runaway conditions or saturate at a particular value, we chose several points along the REf–OH–NO curve and for each of these points, and we generated a hypothetical input dataset keeping all parameters constant but changing the NO$_x$ levels from 0.2 to 3 ppbv, while maintaining the original NO/NO$_2$ ratio (Fig. 8).

Figure 8 shows that in general the REf-OH increases to about 0.85 at 500 ppbv of NO. After this, it decreases gradually, related to the reaction of NO$_2$ with OH. Concurrently, the loss of HO$_2$ due to self-reactions and reaction of HO$_2$ with CH$_3$O$_2$ becomes much weaker compared to the loss of HO$_2$ to NO with increasing NO levels. The ratio of HO$_2$ losses via the radical–radical interactions (i.e. the reactions of HO$_2$ with HO$_2$ and CH$_3$O$_2$) to the HO$_2$ losses via the recycling channel (i.e. reactions of HO$_2$ with NO and O$_3$) decreases exponentially, becoming less than 1 at 35 pptv of NO and less than 0.5 at 70 pptv of NO and 0.01 at 1 ppbv NO. The secondary OH production also shows a similar pattern with respect to REF-OH, peaking around 500 ppbv for the different input datasets and decaying thereafter. The model generated OH peaks in this range but the HO$_2$ concentrations start to drop much earlier, from around 70 pptv of NO. At these low NO levels, HO$_2$ loss by peroxide–peroxide reactions is comparable to the channels recycling HO$_2$ into OH, which can then be removed through reaction, e.g. with NO$_2$. However, the recycling efficiency is only about 0.6 at 70 ppbv of NO. Budget analysis indicates that HO$_2$ production drops with increasing NO levels for this test dataset constrained to specific NO$_x$ levels. Overall, for the HC levels observed during CYPHEX, the self-cleaning capacity of the atmosphere peaks at around 500 ppbv of NO, which at a NO / NO$_2$ ratio of 0.2 is equivalent to about 3.0 ppbv NO$_x$. Increasing NO$_x$ emissions further would lead to decreased HO$_2$ levels and the recycling efficiency is unlikely to increase further. Conversely, decreasing NO levels below a few hundred parts per trillion by volume of NO would also decrease the recycling efficiency for the same HC levels, indicating that a small amount of NO$_x$ helps sustain the self-cleaning capacity of the atmosphere (Fig. 8).

### 4 Conclusions

Atmospheric OH and HO$_2$ were measured as part of a comprehensive atmospheric chemistry field measurement campaign conducted in Cyprus in the summer of 2014 to study the major processes impacting atmospheric oxidation and air chemistry in a relatively unpolluted coastal region, periodically influenced by long-range transport of European emissions. A comprehensive suite of atmospheric chemistry measurements obtained during CYPHEX enabled a detailed investigation of atmospheric oxidation processes, using a photochemical box model, under low-NO$_x$ conditions. The
box model (CAABA/MECCA with MOM chemistry) simulations for OH agreed to within 10 % with in situ OH observations. Model simulations for HO2 agreed to within 17 % of the in situ observations. However, the model strongly underpredicted HO2 at high terpene (α- and β-pinene, limonene) concentrations (> 80 pptv), this under-prediction reaching up to 38 % at the highest terpene levels due to loss of HO2 by reactions with terpene-generated peroxy radicals. Comparison of alternative reaction pathways to reduce this unrealistic HO2 loss showed that autoxidation can be an effective sink for the RO2 species generated from terpene oxidation. However, low-terpene and low-NO regimes, prevailing during CYPHEX, and the absence of limonene chemistry in the current model scheme, precluded a more rigorous analysis of the probable chemical pathways for RO2 degradation in the atmosphere. Further, there is evidence that the rate constant of CH3O2 with OH could be 2 times faster than used in the current models (Bossolasco et al., 2014). Applying this in general to other peroxides, the rate constants need to be revisited as they will have a non-negligible impact on the chemical composition of the atmosphere, especially in remote low-NOx environments where the peroxy lifetimes are relatively long. As already indicated in literature, there is large uncertainty in the rate coefficients of different RO2 species with NO and HO2 (King et al., 2001), which are scopes for future studies.

The radical budget analysis for CYPHEX showed that primary production of OH via photolysis of O3 constituted the main OH source, and peak daytime contributions from this channel exceeded 45 % for most of the days. The average daytime contribution from HONO photolysis was 12.3 % but it exceeded 30 % in the early morning on a few occasions. The recycled OH from reaction of HO2 with O3 and NO accounted for 14.5 % and 17.7 % of the total OH production respectively. The maximum observed recycling efficiency was about 0.7 for about 100 pptv of NO. CO was not only the single major daytime sink of OH, accounting for nearly one-fifth of OH loss during the study period, but also the single major HO2 production source. Despite low-NOx regimes during CYPHEX, NO contributed more than 23 % of peak daytime loss for HO2, reaching over 50 % on a few occasions. The lowest HOx production and losses occurred in the highly processed air masses with low OH reactivity, i.e. low precursor levels. These air masses were heavily influenced by the marine boundary layer, mostly by the Mediterranean Sea, but sometimes as far as the Atlantic Ocean (24–25 July, days 205–206 of the year) with some influence from southwestern Europe/northern Africa. Additionally, our results connote the need for deeper understanding of the reaction channels for organic peroxides in the atmosphere.

Data availability. Data are available at the EDMOND repository of the Max Planck Society (Mallik et al., 2018). If further information is needed, please send an email to hartwig.harder@mpic.de.

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