

Answer to referee #2

We thank referee 2 for revising the manuscript and helping improving it. Please find below our answers to his comments:

Referee comment:

This is a review for the first revision of the manuscript, submitted by Zannoni et al. Overall, the manuscript has been improved in terms of scientific scopes and data presentation. However, I still believe that it needs to be significantly improved by addressing some inconsistencies to be considered for the final production.

The main weak point of the presented notion of data analysis is that the authors just focused on one period when the maximum OH reactivity was quantified (e.g. July 29th and the three-consecutive day after that). The justification was that this was the time period when local biogenic influences were dominant. Using a correlation analysis between temperature and missing OH reactivity, the authors argued the case (Figure 7). However, there are a couple of obvious observations that the authors need to address to substantiate this reasoning. First, it does not seem to anthropogenic influences did not change too much over the time by observing the temporal variations of AVOCs and others (probably mostly trace gases such as CO and NOX). The BVOC contribution to the total calculated OH reactivity changes significantly and the variance does not correlate with the notated wind direction shown in Figure 2. In this context, I would suggest that the authors need to discuss the obvious inconsistency in the level of assessed missing OH reactivity. For example, in Aug 8th, the BVOC contribution to the calculated OH reactivity is appeared to be the largest during the whole observational period. However, it does not seem to have any missing OH reactivity, which is inconsistent with the main conclusion of the current form of manuscript and requires further justification. Moreover, the duration between July 23rd and July 26th, the fraction or percentage of missing OH reactivity with respect to observed OH reactivity is appeared the highest although the BVOC contribution towards calculated OH reactivity was relatively small. This also requires for further reasoning. In this reason, I am strongly convinced that the data analysis should be redone for holistic discussion rather than just focused on one time period as the current manuscript is framed. One notable difference I can tell the episode (July 26th to July 28th) when the authors focused on from the other period is high temperature and low wind speed, which may indicate a strong stagnation episode. This may allow accumulating uncharacterized/unknown oxidation products, I suspect.

Response:

We thank the reviewer for his/her comments and would like to answer the different points. At first, the reviewer mentions “this is a review for the first revision of the manuscript...” but we would like to note that this was the second revision of the manuscript and therefore we consider important to take into consideration all the previous reviews.

The referee is right in saying that our discussion has focused mostly on a specific period of the campaign, 27-29 July. We did so because for this period there was a clear influence of the oxidation products of biogenic VOCs, compared to the other periods, where the observed missing reactivity (23-26 July) or the absence of missing reactivity (2-5 August) requires a more speculative explanation. Concerning this specific point, we would like to note that the initial version of the manuscript had a longer discussion about the missing reactivity. Following the suggestion of the reviewer of that version, we shortened the discussion to remove the too speculative hypothesis and focused on the period where we could bring a clear argumentation for the missing reactivity. Nevertheless, we do not agree with: “the authors just focused

on one period...”(29th July and the three consecutive days after). Indeed, we would like to argue that: (i) the calculations of the difference between PTR-MS and GC results concerns the whole time series and therefore the discussion on the missing reactivity from the unmeasured primary terpenes is done for the whole campaign period; (ii) the temperature dependency relationship was investigated on a longer period than just 3 days (from 23 to 30 July) with a discussion contrasting the 23-26 and 27-29 periods.

We agree with the reasoning of the referee concerning the influence of BVOCs during 27-29 July due to local drivers: higher temperature and low wind speed have led to a stagnant episode where oxidation products have accumulated. This is shown in Figure 6 where the concentrations of the characterized oxidation products increased over time during 27-29 July. A figure (figure 4) is also added to the supplementary material where ratios between measured oxidation products and their primary precursors are depicted. The accumulation of oxidation products over night and during the episode of missing reactivity of 27-29 July is evidenced by the large values observed for (MVK+MACR)/isoprene and nopinone/ β -pinene during nighttime periods. The measured first-generation oxidation products may have led to the formation of more oxidized species that were not characterized, therefore to an increase of the missing reactivity during this period.

The referee is also right in pointing out that the calculated OH reactivity profile due to BVOCs changes significantly with BVOCs emissions variability, which is due to local drivers rather than wind sectors or anthropogenic episodes. Therefore, the difference between 27-29 July and 2-5 August is due to the different atmospheric conditions: higher temperature and lower wind speed have led to the accumulation of unmeasured oxidation products during 27-29 July. Indeed, during 2-5 August, we measured lower concentrations of oxidation products of BVOCs (Figure 6) and we did not observe the same increase in the ratio of oxidation product towards primary species as during 27-29 July, despite the large impact of the primary BVOCs on the calculated OH reactivity. Therefore, we did not see any missing reactivity during 2-5 August. We think that the case of the period 23-26 July is different. During these days, BVOCs emissions were not dominant as during the last part of the campaign. The air masses measured at the site travelled from the western sector (North of Spain and South of France) for a longer transport time (up to 48 hours) over the sea (Michoud et al., 2017), which suggests that unmeasured oxidation products, likely different from those produced during the local oxidation of BVOCs, could have reached the measuring site, leading to the observed missing reactivity.

In order to take into account the different points mentioned above, we added these elements to the manuscript. Please find the changes below:

1.1 Missing reactivity and air masses fingerprint

Figure 2 reports the time series of the total measured OH reactivity and calculated OH reactivity with their associated errors (35% and 20%, respectively). The largest significant discrepancy between reactivities occurred during July 23-30 (56% of missing reactivity, calculated as the difference in percentage between measured and calculated reactivity). We combined air mass backtrajectories and atmospheric mixing ratios of some common atmospheric tracers to determine the chemical fingerprint of the sampled air and to investigate the origin of the missing reactivity. We chose isoprene and pinenes for air masses influenced by biogenic activity, while propane and CO were used for those enriched in anthropogenic pollutants (see supplement). Maximum concentrations of anthropogenic pollutants were measured when the air masses originated from the North East sector: between July 21-23 and between July 31 and August 3, indicating weak pollution events coming from the industrialized areas of the Po Valley and Milan (Italy). The OH

reactivity during these events was at most 9 s^{-1} , no significant missing reactivity occurred, mainly because of the weakness of the events, and the deployment of instruments that were able to characterize a large number of AVOCs. On the other hand, biogenic activity was independent of the wind sector and showed a larger variability linked to local drivers, such as air temperature, solar irradiance and wind speed (Figure 6). Biogenic VOCs and their oxidation products had a large influence on the measured OH reactivity and potentially on the missing reactivity (Figures 2-5); however, periods characterized by a large contribution of BVOCs to the measured OH reactivity (27-29 July and 2-4 August) present some differences in missing reactivity, which will be discussed in the next section. There is not a clear trend between the measured and missing OH reactivities with the origin of air masses (Figure 2). Nevertheless, when VOCs were imported from the western sector, the air masses travelled over the sea with an average transport time of up to 48 hours, which can have favored the enrichment in oxygenated compounds through the atmospheric oxidation of primary VOCs initially present in these air masses before they left continents (Michoud et al., 2017).

1.2 Insights into the missing OH reactivity

The missing OH reactivity, obtained as the absolute difference between measured and calculated reactivity values, is reported in Figure 6. Only values higher than the detection limit of 3 s^{-1} are displayed in this figure. Significant missing reactivity ($2.8 \pm 2.2\text{ s}^{-1}$ on average over the whole campaign) can be observed from 23 to 29 July and more sporadically before 23 July (3 data points over a total of 41) and after 29 July (10 data points). To determine the causes of the reported missing OH reactivity we first investigated whether the VOC instruments deployed on the measurement site did not selectively detect some of the emitted BVOCs. We compared the total monoterpene concentration observed by PTR-MS to the summed monoterpenes concentration from GCs and calculated a concentration difference ranging from 0.2 to 0.6 ppbv (see supplement), the PTR-MS measurements being higher than the sum of speciated monoterpenes. Assuming a reaction rate coefficient with OH of $1.56 \times 10^{-10}\text{ cm}^3\text{ molecule}^{-1}\text{ s}^{-1}$ (weighted value estimated from monoterpenes emitted by the plants present at the site, see Bracho-Nunez et al., 2011) we can roughly estimate a value of $0.8\text{--}2.3\text{ s}^{-1}$ of missing OH reactivity due to these unspciated monoterpenes, which represents 30% of the reported missing reactivity on average. Therefore, unmeasured monoterpenes account for a significant fraction of the missing OH reactivity but do not fully explain the total missing reactivity.

The missing OH reactivity temperature dependency was used in previous studies (Di Carlo et al., (2004), Mao et al., (2012) and Hansen et al., (2014)) to determine whether terpenes (Di Carlo et al., 2004) or their oxidation products (Mao et al., 2012) could explain the missing OH reactivity. Terpene emissions depend on temperature following the expression, $E(T) = E(T_s) \exp[\beta(T - T_s)]$, where $E(T_s)$ is the emission rate at temperature T_s , β a temperature sensitivity factor, and T the ambient temperature. Di Carlo et al., (2004) found that the missing OH reactivity reported from a temperate forest in northern Michigan followed the same temperature dependency as terpene emissions, with $\beta = 0.11\text{ K}^{-1}$. Similarly, Mao et al., (2012) reported a β factor of 0.168 K^{-1} from a temperate forest in California. Since this value is higher than the upper limit of 0.144 usually observed for terpene emissions, the authors attributed the missing OH reactivity to unmeasured oxidation products of BVOCs, which was supported by box model simulations.

In the present study, we investigated the temperature dependency of the missing OH reactivity to determine whether BVOC oxidation products caused the missing reactivity for 23-26 and 27-29 July periods. We found a clear dependence ($\beta = 0.158\text{ K}^{-1}$ and $R^2 = 0.643$) between the missing OH reactivity and

temperature for some days of the campaign (27th, 28th, 29th), while no correlation ($\beta = 0.007 \text{ K}^{-1}$ and $R^2 = 0.001$) was found during 23-26 July, see Figure 7. Indeed, the lack of a clear temperature dependency and the influence of long range transport on air masses imported from the western sector, as demonstrated by Michoud et al., (2017) for the same data set suggest that the local oxidation of BVOCs was not the main cause of the missing reactivity observed during 23-26 July. In contrast, the similar temperature dependencies observed in the study of Mao et al., (2012) and during 27-29 July suggests that oxidation products of locally emitted BVOCs was the main cause of missing reactivity for this period. This interpretation is further confirmed by a significant increase of the concentrations of some BVOC oxidation products during 27-29 July (Figure 6) and the accumulation of these secondary species versus the primary biogenic compounds during nighttime (Figure 4 supplement).

Figure 6 shows the missing OH reactivity and volume mixing ratios of BVOCs and their oxidation products together with meteorological parameters, such as temperature, wind speed and solar irradiance. Compounds are reported using their protonated mass measured by PTR-MS, such as: m/z 69 (isoprene) and m/z 137 (monoterpenes) for primary emissions, m/z 71 (isoprene first generation oxidation products: Methyl Vinyl Ketone (MVK) + methacrolein (MACR) + possibly fragments of isoprene hydroxyperoxides (ISOPOOH)), m/z 139 (nopinone, β -pinene first generation oxidation product) and m/z 151 (fragment of pinonaldehyde, α -pinene first generation oxidation product) for the BVOC oxidation products. It is worth noting that m/z 111 and m/z 113 are also reported since these masses have been attributed to oxidation products of several terpenes. Indeed m/z 111, m/z 113 and m/z 151 were observed in chamber and field studies (Lee et al., 2006, Holzinger et al., 2005) as they are formed from the photo-oxidation of different terpenes; highest yields were attributed to terpenes also common to the Mediterranean ecosystem, such as myrcene, terpinolene, linalool, methyl-chavicol and 3-carene (Lee et al., 2006, Bracho-Nunez et al., 2011). We note that all the above-mentioned masses (with the exception of m/z 111 and m/z 113, for which no rate coefficient was found for the reaction of the unprotonated molecule with OH) have been taken into account in the calculated OH reactivity. The reported time series show that both primary BVOCs and most of the OVOCs resulting from their oxidation had a clear diurnal profile with highest values during midday, similar to that observed for the missing reactivity.

Most of the nights were associated with low mixing ratios of BVOCs (29 pptv on average for isoprene) and their oxidation products (40 pptv on average for m/z 71) and corresponded to very low or no missing reactivity. Nevertheless, we note that during some nights (26 to 27, 27 to 28, 29 to 30 July), some compounds (m/z 71, m/z 113) were also present in low amounts and a missing reactivity of about 4 s^{-1} was noticed. These nights were associated with low wind speed and high temperatures, indicating a stagnant episode that could have favored the accumulation of unmeasured oxidation products during 26-29 July (ratios of m/z 71-to- m/z 69 and nopinone-to- β -pinene were maximum 13.9 and 7.7, respectively, see supplement) as was already observed by night in an oak-forest canopy (Zannoni et al., 2016). The mixing ratios of the first- oxidation products of isoprene and monoterpenes (m/z 71, m/z 139) strongly increased after July 26 when the wind speed was lower and increased again after July 27 when air temperature increased. In contrast, during another period of BVOCs influence as 2-5 August, slightly lower air temperature was experienced, less accumulation of oxidation products (Figure 4 supplement) and lower concentrations of secondary species (m/z 139, m/z 151) were reported (Figure 6). These pieces of evidence also suggest that unmeasured, locally generated BVOC oxidation products contribute significantly to the missing OH reactivity during 27-29 July.

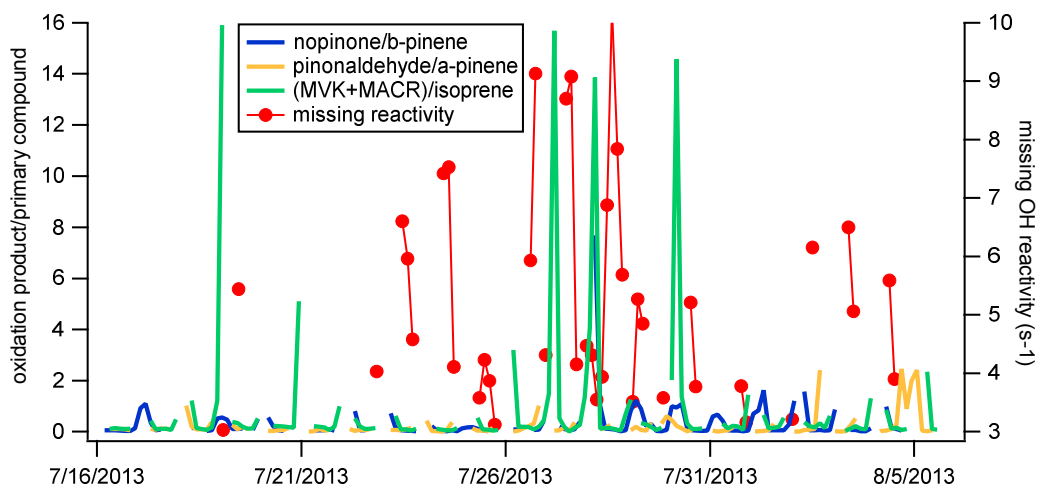


Figure 4. Missing OH reactivity and ratio between oxidation products and primary compounds for some biogenic molecules.

Minor comments:

Comment:

1) Recently in AMTD, the results of an OH reactivity intercomparison study has been published. It would better to describe the results in the perspectives of potential limitations of the analytical method.

Response:

A paragraph regarding the last intercomparison results was included in the methods section:

An intercomparison study carried out with other instruments based on the CRM technique and on laser induced fluorescence-based technique, showed that the measured OH reactivities agree among them (Fuchs et al., 2017). The same study showed the following limitations of CRM instruments compared to laser-based techniques: (i) higher limit of detection (2 s^{-1} vs. $< 1 \text{ s}^{-1}$), lower time resolution (10-15 minutes vs. 30 seconds to a few minutes), lower accuracy due to the required corrections to determine the final OH reactivity value (pseudo first order deviation and OH recycling for environments exposed to high NO_x concentrations). Additionally, CRM instruments underestimated the measured OH reactivity of known terpenes mixture, therefore if any missing reactivity is reported from terpenes-dominated environment this has to be seen as a lower limit of missing reactivity present.

Comment:

2) Page 10 Line 8: Please be consistent by either using hydroxyl radical reactivity or OH reactivity

Response:

Yes.

Comment:

3) Please define how you evaluate missing OH reactivity

Response:

We evaluated the missing OH reactivity as the difference between measured and calculated reactivity values, page 13 line 24:

The missing OH reactivity, obtained as the absolute difference between measured and calculated reactivity values, is reported in Figure 6. Only values higher than the detection limit of 3 s^{-1} are displayed in this figure.

Comment:

4) Page 14 Line 4 to Line 11: What about the other periods? The explanation for the dataset between 23rd to 26th is not clear enough. The authors need to discuss more specific how they have reach that conclusion.

Response:

Yes, we included the following paragraph: Indeed, the lack of a clear temperature dependency and the influence of long range transport on air masses imported from the western sector demonstrated by Michoud et al., (2017) for the same data set suggest that the local oxidation of BVOCs was not the cause of the missing reactivity during 23-26. In contrast, the similar temperature dependency with the study of Mao et al., (2012) during 27-29 July suggests oxidation products of BVOCs being the cause of the missing reactivity. This interpretation is further confirmed by the increase of concentrations of some BVOC oxidation products during 27-29 July (Figure 6) and the higher accumulation of the secondary species versus the primary biogenic compounds (Figure 4 supplement).

Comment:

5) Page 23 Tabel 1 it is “alpha” pinene and “beta” pinene not a and b-piene, repsectively.

Response:

Done.

Summertime OH reactivity from a receptor coastal site in the Mediterranean basin

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Abstract

Total OH reactivity, the total loss frequency of the hydroxyl radical in ambient air, provides the total loading of OH reactants in air. We measured the total OH reactivity for the first time during summertime at a coastal receptor site located in the western Mediterranean basin. Measurements were performed at a temporary field site located in the northern cape of Corsica (France), during summer 2013 for the project CARBOSOR (CARBOn within continental pollution plumes: SOurces and Reactivity) -ChArMEx (Chemistry-Aerosols Mediterranean Experiment). Here, we compare the measured total OH reactivity with the OH reactivity calculated from the measured reactive gases. The difference between these two parameters is termed missing OH reactivity, i.e., the fraction of OH reactivity not explained

by the measured compounds. The total OH reactivity at the site varied between the instrumental LoD (limit of detection= 3 s^{-1}) to a maximum of $17 \pm 6 \text{ s}^{-1}$ (35% uncertainty) and was $5 \pm 4 \text{ s}^{-1}$ (1σ standard deviation) on average. It varied with air temperature exhibiting a diurnal profile comparable to the reactivity calculated from the concentration of the biogenic volatile organic compounds measured at the site. For part of the campaign, 56% of OH reactivity was unexplained by the measured OH reactants (missing reactivity). We suggest that oxidation products of biogenic gas precursors were among the contributors to missing OH reactivity.

1 Introduction

Atmospheric photo-oxidation reactions are initiated by three main oxidants: the hydroxyl radical (OH), ozone (O_3) and the nitrate radical (NO_3). Among those, the OH radical is by far the most important, capable of reacting with the vast majority of chemical species in the troposphere (Levy, 1971). Photo-oxidation reactions are the most efficient cleansing processes occurring in the atmosphere, and constitute an important sink for reactive gases including volatile organic compounds (VOCs).

Total OH reactivity is the first-order total loss rate of the hydroxyl radical in the atmosphere due to reactive molecules. It is the total sink of OH, therefore representing a top-down measure of OH reactants present in ambient air.

Measurements of the total loss of OH and OH reactive gases are often coupled. The total reactivity of the latter is determined by summing each gas individual reactivity as the product of their atmospheric concentration and their reaction rate coefficient with OH. Here, this is referred to as calculated OH reactivity and comparisons between the calculated and the measured OH reactivity have showed that discrepancies in various environments exist (Di Carlo et al., 2004, Nölscher et al., 2016). The missing OH reactivity, namely the fraction of OH reactivity not explained by simultaneous measurements of reactive gases, has been associated with unmeasured compounds either primary emitted, secondary generated, or both (e.g. Sinha et al., 2010, Nölscher et al., 2012, Nölscher et al., 2013, Edwards et al., 2013, Hansen et al., 2014, Kaiser et al., 2016).

The Mediterranean basin comprises countries from three different continents and a population of 450 million inhabitants. Its climate is characterized by humid-cool winters to hot-dry summers, when the area is usually exposed to intense solar radiation and high temperatures. Forests, woodlands and shrubs occupy large areas of the region, which has rich biodiversity

and is the habitat to a high number of identified species (Cuttelod et al., 2008). The dominant airflow in summertime is driven from North to South; therefore, the basin is exposed to air masses coming from European cities and industrialized areas. Transported pollution and the intense local anthropogenic and biogenic activity result in high loadings of atmospheric gases, particles and complex chemistry (Lelieveld, 2002).

Climate model predictions indicate that the Mediterranean area will face unique impacts of climate change. Predictions show that this region will suffer higher temperatures and extended drought periods, thus affecting the strength and type of emissions which will further impact air quality and climate (Giorgi and Lionello, 2008). Finally, additional observations are useful for better predicting the future state of this region (Mellouki and Ravishankara, 2007).

In this study, we address the following scientific questions:

- 1) What proportion of the total reactive gases emitted and formed over the area do we know and can we detect?
- 2) Which species mostly influence the OH reactivity over this site within the basin?

To answer these questions, we measured the total OH reactivity at a receptor coastal site in the western Mediterranean basin during summer 2013. Measurements were part of an intensive fieldwork campaign aimed at investigating sources and sinks of gaseous constituents in the area (CARBOSOR, carbon within continental pollution plumes: Sources and Reactivity, within the ChArMEx project, Chemistry and Aerosols in a Mediterranean Experiment; charmex website: <http://charmex.lsce.ipsl.fr/>). We measured the total OH reactivity with the comparative reactivity method instrument (CRM) (Sinha et al., 2008) during 16/07/2013-05/08/2013 at the monitoring station of Erso, France. The field site was chosen for being: (i) far from anthropogenic sources; (ii) exposed to aged air masses of different origins, including air masses enriched in oxidation products transported from continental industrialized areas. Total OH reactivity here served to evaluate whether the ambient reactive gases were all identified or not. Specifically, it was useful to determine what kind of pollution event could be better captured through the instrumentation deployed at the site, assuming that a group of reactive gases traces a specific type of event (primary anthropogenic or biogenic emissions, secondary formation). Due to the high number of existing VOCs, OH reactivity also makes a powerful means for investigating VOC emissions and reactions. The following sections will describe the field site under study, the

methodologies used, our results of OH reactivity and insights into the unmeasured reactive gases.

2 Field site

The Ersa windfarm (42.97°N, 9.38°E, altitude 533 m) is located in the northern cape of Corsica (France), in the western Mediterranean basin (Figure 1). It is 2.5 km away from the nearest coast (West side) and 50 km away from the largest closest city and harbour Bastia (South side). It is located on a hill (533 m a.s.l.) and it is surrounded by the Mediterranean Sea on West, North and East sides. The site is exposed to air masses from continental areas especially France and northern Italy, with the harbours of Marseille and Genoa about 300 km away, and the industrialized areas of Milan and the Po valley 400 km away. Furthermore, the Mediterranean maquis, a shrubland biome typical of the Mediterranean region, densely surrounds the measurement station. The ground station consists of a long-term meteorology, trace gas concentrations, aerosol size and composition monitoring laboratory (measurements collected from 2012 to 2014), and temporary measurements of gases and aerosol properties over a total surface area of ~100 m² where instruments are distributed. Measurements of total OH reactivity and trace gases reported in this study were all performed within this area (see Figure 1 for details).

We measured the OH reactivity during two main periods: an intercomparison exercise for OH reactivity between two CRM instruments during 8/07/2013-13/07/2013 (see Zannoni et al., 2015), and the intensive ambient monitoring campaign, CARBOSOR during 16/07/2013-05/08/2013. Within the same project, instruments for measuring radicals, inorganic and organic compounds, aerosol chemical composition and physical properties, and meteorology were simultaneously deployed. The next section will provide an overview of the methods selected for this study.

3 Methods

3.1 Comparative Reactivity Method

We carried out measurements of total OH reactivity using a comparative reactivity method instrument assembled in our laboratory (CRM-LSCE from Laboratoire des Sciences du Climat et de l' Environnement, see Zannoni et al., 2015). In brief, the comparative reactivity method is based on the concept of producing a competition for in-situ generated OH radicals, between a reactive reference compound, in our case pyrrole (C₄H₅N), and ambient reactive

gases (Sinha et al., 2008). This is achieved by introducing a known amount of pyrrole diluted in zero air and N₂ in a flow reactor coupled to a Proton Transfer Reaction-Mass Spectrometer (PTR-MS, see Lindinger et al., 1998, De Gouw and Warneke, 2007). Pyrrole is chosen as a reference compound for its well characterized kinetics (Atkinson et al., 1984, Dillon et al., 2012), for not being present in the atmosphere at normal conditions, and for being easily detectable at the protonated m/z 68 (C₄H₅NH⁺) through PTR-MS without any interference. The Proton Transfer Reaction-Mass Spectrometer run at standard conditions (Pdrift = 2.2 mbar, E/N = 130 Td (1 Td=10⁻¹⁷ Vcm⁻¹), Tinlet = 60 °C) is the detector of choice for its real-time measurement capabilities and robustness over time (see also Nölscher et al., 2012b).

The CRM usual experimental procedure includes the following stages: monitoring of C0 wet/dry, followed by C1 dry or wet, C2 wet, and C3 ambient. With C0, C1, C2, C3 being the concentration of pyrrole detected with the PTR-MS, in order: after injection (C0), after photolysis of pyrrole (C1), after reaction with OH (C2), when ambient air is injected and the competition for OH radicals starts (C3). Switches between C2 (background pyrrole in zero air) and C3 (pyrrole in ambient air) result in modulations of the pyrrole signal, which are used to derive total OH reactivity values from the following equation:

$$R_{air} = \frac{(C3 - C2)}{(C1 - C3)} \cdot k_{pyrrole+OH} \cdot C1 \quad (1)$$

With $k_{pyrrole+OH}$ being the rate constant of reaction between pyrrole and OH= (1.20±0.16)×10⁻¹⁰ cm³ molecule⁻¹ s⁻¹ (Atkinson et al., 1984, Dillon et al., 2012).

During the whole campaign, we ran systematic quality check controls on the instrument (see supplementary material).

Sampling was performed through a 3 m long, 1/8'' OD PFA sampling line at a flow rate of 0.25 sL/min with a residence time of the sample of 3 s. The sampling line was covered and kept at ambient temperature and installed at about 1.5 m above the trailer where the CRM was installed. A PTFE filter was placed at the inlet of the sampling line to avoid sampling particles. Some highly-reactive chemical species (i.e. sesquiterpenes) may have been lost before reaching the reactor due to wall losses in the sampling line and/or filter surface.

We recorded PTR-MS data using a dwell time of 20 s for pyrrole, with a full cycle of measurements every 30 s. We switched between C2 and C3 every 5 minutes, resulting in a data point of reactivity every 10 minutes. Each data point of reactivity obtained from eq. (1)

was corrected for: (i) humidity changes between C2 and C3, (ii) deviation from the assumption of pseudo first order kinetics between pyrrole and OH, (iii) dilution of ambient air reactivity inside the reactor. A detailed description on how the correction factors were obtained and how the raw data were processed can be found in the publication of Zannoni et al., (2015). We did not account for OH recycling in our reactor due to nitrogen oxides ($\text{NO} + \text{NO}_2$) since ambient nitrogen monoxide (NO) was below 0.5 ppbv at the site (NO_2 below 2 ppbv), which is too low for interfering with the system. Tests performed in the laboratory after the campaign have demonstrated that the instrument is not subject to ozone interference. The impact on CRM measurements of OH recycling reactions observed during the oxidation of some ambient species (e.g. methylvinylketone and methacrolein (MVK+MACR), isoprene hydroxyhydroperoxides (ISOPOOH), aldehydes) was determined to be negligible due to the low concentrations of these species and the high HO_2 concentration in the CRM reactor, which disfavor unimolecular reactions.

The limit of detection (LoD) of CRM-LSCE was estimated to be $\sim 3 \text{ s}^{-1}$ (3σ) and the systematic uncertainty $\sim 35\%$ (1σ), including uncertainties on the rate coefficient between pyrrole and OH (8%), detector sensitivity changes and pyrrole standard concentration (22%), correction factor for kinetics regime (26%) and flows fluctuations (2%); see also Michoud et al., 2015. An intercomparison exercise with another CRM instrument carried out before the campaign demonstrated that the measured reactivities were in good agreement (linear least squares fit with a slope of one and R^2 value of 0.75). An intercomparison study carried out with other instruments based on the CRM technique and on laser induced fluorescence-based technique, showed that the measured OH reactivities agree among them (Fuchs et al., 2017). The same study showed the following limitations of CRM instruments compared to laser-based techniques: (i) higher limit of detection (2 s^{-1} vs. $< 1 \text{ s}^{-1}$), lower time resolution (10-15 minutes vs. 30 seconds to a few minutes), lower accuracy due to the required corrections to determine the final OH reactivity value (pseudo first order deviation and OH recycling for environments exposed to high NO_x concentrations). Additionally, CRM instruments underestimated the measured OH reactivity of known terpenes mixture, therefore if any missing reactivity is reported from terpenes-dominated environment this has to be seen as a lower limit of missing reactivity present.

3.2 Complementary measurements at the field site

Trace gases were measured using a broad set of techniques available at the site, including: Proton Transfer Reaction-Mass Spectrometry (PTR-Time of Flight MS, Kore Technology Ltd., UK), online and offline Gas Chromatography (GC-FID/FID and GC-FID/MS, Perkin Elmer), Liquid Chromatography (HPLC-UV, High Performance Liquid Chromatography-UV light detector), the Hantzsch reaction method (AERO-LASER GmbH, Germany) and wavelength-scanned cavity ring down spectrometer (WS-CRDS, G2401, Picarro, USA). The measured concentration and the reaction rate coefficients of each measured compound with OH were used to calculate the OH reactivity with eq. (2):

$$R = \sum_i k_{i+OH} \cdot X_i \quad (2)$$

With i being any measured compound listed in Table 1 and X its concentration.

Most of the chemical species whose reactivity was summed to determine the calculated reactivity were measured through PTR-MS and GC.

The sampling system for the PTR-MS consisted of a 5 m PFA sampling line, installed above the PTR-MS trailer (see Figure 1). The residence time in the sampling line was 4 s. We operated the PTR-MS at 1.33 mbar pressure and 40°C temperature in the drift tube for an E/N of 135 Td. The PTR-MS was calibrated every 3 days using certified mixtures of different VOCs: 15 VOCs (Restek, France), 9 VOCs (Praxair, USA), 9 OVOCs (Praxair, USA). More details on the calibration standards are available in Michoud et al. (2017). For this campaign, we only reported the masses that we clearly identified and for which we had a calibration standard. Some other masses, possibly corresponding to oxidation products of terpenes were also extracted from the mass spectra and tentatively quantified (e.g. m/z 99, 111, 113, 155). A visual inspection of the full mass spectra did not reveal any other abundant species that we did not consider. The GCs were calibrated twice at the beginning and at the end of the field campaign with certified gas mixtures: one including 29 VOCs (Praxair, USA), another including 29 NMHCs and three terpenes (NPL, UK). Total uncertainties from measurements (including precision and calibration procedure) were in the range 5-23% for compounds measured by PTR-MS and GC-FID, and in the range 5-14% for GC-MS.

Isoprene was measured by both PTR-MS and GC and the results correlated within the measurement uncertainty (slope and R^2 of the regression for 415 data points are 0.93 ± 0.03 and 0.77, respectively; see supplement). A small offset in the scatter plot (approximately 100

pptv) may indicate a small interference at m/z 69 for the PTR-MS measurements; therefore, we used data from GC to calculate the OH reactivity. We sampled individual monoterpenes with GC-FID, and on adsorbent tubes for the analysis with GC-MS, while we measured the total monoterpenes fraction by PTR-MS since the instrument cannot distinguish between structural isomers. Concentrations obtained from GC techniques were used to calculate the summed OH reactivity. As recently reported by Rivera-Rios et al., 2014, the m/z 71 measured by PTR-MS might also include the ISOPOOH which could have formed at the site and fragmented inside the PTR-MS. However, it is important for the reader to know that we did not separate the different components of the m/z 71, therefore the presence of ISOPOOH on m/z 71 is assumed based on the recent literature.

We refer to the manuscript of Michoud et al.(2017), for a detailed description of the PTR-MS, online GC and offline sampling on adsorbent cartridges on GC-FID/MS deployed at the site; while the formaldehyde, NO_x, O₃ analysers and WS-CRDS are briefly introduced in the following sections. Table 2 provides a summary of all techniques.

3.2.1 Hantzsch method for measuring formaldehyde

Formaldehyde (HCHO) was measured with a commercial instrument based on the Hantzsch reaction (Model 4001, AERO-LASER GmbH, Germany). Gaseous HCHO is stripped into a slightly acidic solution, followed by reaction with the Hantzsch reagent, i.e. a diluted mixture of acetyl acetone acetic acid and ammonium acetate. This reaction produces a fluorescent compound, which absorbs photons at 510 nm. More details are given in Dasgupta et al., (1988), Junkermann, (2009) and Preunkert et al., (2013).

Sampling was conducted through a 5 m long PTFE 1/4" OD line, with a 47 mm PFA in-line filter installed at the inlet and a flow rate of 1 L min⁻¹.

The liquid reagents (stripping solution and Hantzsch reagent) were prepared from analytical grade chemicals and ultrapure water according to the composition given by Nash, (1953) and stored at 4 °C on the field. The instrumental background was measured twice a day (using an external Hopcalite catalyst consisting of manganese and copper oxides) and calibrated three to four times a week using a liquid standard at 1.10⁻⁶ mol L⁻¹, i.e volume mixing ratio in the gaseous phase of about 16 ppbv. The calibration points were interpolated linearly in order to correct for sensitivity fluctuations of the instrument. The limit of detection was 130 pptv (2σ). The coefficient of variation, i.e the ratio of the standard deviation to the mean background

value, was estimated to be 0.4 %. Measurements of HCHO ran smoothly from the beginning of the campaign until 11 AM LT (local time) of 28/07/2013. At this time, an instrument failure occurred and measurements were stopped.

3.2.2 Chemiluminescence for measuring NO_x

A CRANOX instrument (Ecophysics, Switzerland) based on ozone chemiluminescence was used to measure nitrogen oxides (NO_x=NO+NO₂). Nitric oxide is quantified directly by the instrument while NO₂ is quantified indirectly after being photolytically converted to NO (conversion efficiency=86%). The instrument consists of a high performing two channel CLDs (Chemiluminescence Detectors) with pre-chambers background compensation, an integrated pump, a photolytic converter, an ozone generator and a calibrator. A control software handles and manages the different tasks. The detection limit is 50 pptv (3σ), and the time resolution is 5 minute.

3.2.3 Wavelength-scanned cavity ring down spectrometry (WS-CRDS) for measuring greenhouse gases

In-situ measurements of CO₂, CH₄, CO molar fractions at Ersa are part of the French monitoring network of greenhouse gases, integrated in the European Research Infrastructure ICOS (integrated carbon observation system). The air is sampled at the top of a 40 m high telecommunication tower (573 m), and is analyzed with a wavelength-scanned cavity ring down spectrometer (WS-CRDS, G2401, Picarro, USA). The analyzer is calibrated every 3 weeks with a suite of four reference standard gases, whose molar fractions are linked to the WMO (World Meteorological Organization) scales through the LSCE (Laboratoire des Sciences du Climat et de l'Environnement) reference scale. Measurements were corrected for an empirical correction which takes into account the dilution effect and pressure broadening effect. A humidifying bench was developed to humidify a certified concentration of a gas stream at different humidity levels (see Rella et al., 2013).

3.3 Air masses back-trajectories

The back-trajectories of the air masses were modelled with Hysplit (HYbrid Single-Particle Lagrangian Integrated Trajectory developed by the National Oceanic and Atmosphere Administration (NOAA) Air Resources Laboratory (ARL) (Draxler and Hess, 1998, Stein et al., 2015) for 48 h every 6 hours.

The back-trajectories were grouped according to their origin, altitude and wind speed, such as: 1.North-East, 2.West, 3.South, 4.North-West and 5. Calm-low wind speed/stagnant conditions. More details on the air masses origin and their photochemical age is available in Michoud et al., (2017).

4 Results

4.1 Total measured OH reactivity

The black line in Figure 2 represents the 3-h averaged measured OH reactivity. Here we report all data acquired during 16/07/2013- 05/08/2013, missing data points are due to minor instrumental issues and instrumental controls. Figure 2 also shows the temperature profile of ambient air (gray line, right axis). The OH reactivity varied between the instrumental LoD (3 s^{-1}) to $17 \pm 6 \text{ s}^{-1}$ (3-h averaged maximum value $\pm 35\%$ uncertainty). From the 10 minute time resolution data the highest value of OH reactivity was 22 s^{-1} , reached on 28/07/2013 during the afternoon, when the air temperature at the site was also exhibiting its maximum. During the whole field campaign, the average measured OH reactivity was $5 \pm 4 \text{ s}^{-1}$ (1σ). This value agrees with averaged values of OH reactivity collected during autumn 2011 in the South of Spain for southerly marine enriched air masses (Sinha et al, 2012). In contrast, higher OH reactivity was measured during spring 2014 in a Mediterranean forest of downy oaks, where the average campaign value was $26 \pm 19 \text{ s}^{-1}$ and the maximum value was 69 s^{-1} (Zannoni et al., 2016).

The OH reactivity and air temperature at the site in Corsica co-varied during the whole campaign, with highest values reached during daytime in the periods between July 26-28 and August 2-3. In Figure 2 the origin of the air masses reaching the field site is also reported. The dominant origin of the sampled air masses was west, indicating air masses that had travelled over the sea being possibly more aged. It is not evident that the variability of the OH reactivity is affected by the origin of the air masses; in contrast, air temperature seems to have played a major role. The diurnal pattern of OH reactivity for the whole campaign is reported in Figure 3. Here it is evident that the background value was about 4 s^{-1} during nighttime, it increased at 8:00 AM LT, peaked at 11:00 AM LT, reached a second maximum at 4:00 PM LT and finally decreased at 7:00 PM LT to reach its background value at 10:00 PM LT (local time GMT/UTC+2 hours). It is worth noting that the large amplitude of standard deviation bars (1σ) highlights the large diel variability.

4.2 Calculated OH reactivity and BVOCs influence

Table 1 provides the number and type of chemical species measured simultaneously with the OH reactivity. We determined the calculated OH reactivity from the concentrations and reaction rate coefficients with OH of the species reported in Table 1 using eq. (2). Please refer to Table 2 in the supplementary material for the reaction rate coefficients of the selected compounds with OH. A broad set of compounds were monitored at the site, herein classified as: anthropogenic volatile organic compounds (AVOCs, 41 compounds measured), biogenic volatile organic compounds (BVOCs, 7), oxygenated volatile organic compounds (OVOCs, 15) and others (3 species: CO, NO and NO₂). Please notice that we adopted this classification to simplify the discussion throughout the manuscript, for more details concerning the different sources of these compounds the reader can refer to the notes of Table 1. Figure 2 shows the time series of the summed calculated OH reactivity (thick blue line) and the contributions of each class of chemicals. The maximum of the summed calculated OH reactivity was 11 s⁻¹, and the 24-h averaged value was 3±2 s⁻¹ (1σ). As represented in Figure 3, the class of the biogenic compounds played an important role on the daytime OH reactivity. Here, the shape of the diurnal pattern of the measured reactivity is slightly shifted to the BVOCs OH reactivity, which suggests a possible influence from the oxidation products of biogenic molecules. The mean percentage contribution of each class of compounds to the summed calculated reactivity is determined for daytime (from 07:30 to 19:30, LT) and nighttime data (from 19.30 to 04.30 LT) in Figure 4. During daytime BVOCs contributed the largest fraction of OH reactivity (45%), followed by inorganic species (24%), OVOCs (19%) and AVOCs (12%). Interestingly, only 7 BVOCs had a higher impact than 41 AVOCs. This is explained by: i) the relatively high concentration of BVOCs (maximum values for isoprene and sum of monoterpenes is 1 and 1.5 ppbv, respectively), ii) the generally large BVOC reaction rate coefficients with OH (Atkinson and Arey, 2003) and iii) the relatively low concentration of AVOCs measured during the campaign. BVOCs accounted only for 5% of the total VOCs concentration, followed by AVOCs (15%) and OVOCs (79%) (mean campaign values, see also Michoud et al., 2017) which highlights the reactive nature of the measured BVOCs. During nighttime, BVOCs concentrations decreased (see Figures 2 and 3); CO and NO_x had the largest influence on OH reactivity (43%), followed by OVOCs (27%), AVOCs (23%) and BVOCs (7%). Particularly, CO and long-lived OVOCs and AVOCs constituted a background reactivity of ~ 2-3 s⁻¹, as also shown by the diurnal profiles reported in Figure 3.

1 Inside the BVOCs class, the total fraction of monoterpenes contributed more than isoprene to
2 the OH reactivity (Figure 5). During daytime, OH reactivity due to monoterpenes was
3 between 1.4 to 7.4 s⁻¹ and varied with air temperature, on the other hand, isoprene reactivity
4 with OH varied between 0.3-2.3 s⁻¹ (minimum and maximum values on 29/07/13 and
5 03/08/2013, respectively). In contrast with monoterpenes OH reactivity, the reactivity of
6 isoprene towards OH varied with both air temperature and solar irradiance. Overall, both
7 monoterpenes and isoprene OH reactivities had the characteristic diurnal profile observed for
8 their atmospheric concentrations. High concentrations depended on air temperature, solar
9 radiation as well as calm-low wind speed conditions. These results indicate a large impact of
10 BVOC oxidation on the local photochemistry.

11 The very reactive monoterpene α -terpinene had the largest contribution on OH reactivity
12 among the measured BVOCs (31%), followed by isoprene (30%), β -pinene (17%), limonene
13 (12%), α -pinene (8%), camphene (2%) and γ -terpinene (1%), over a total averaged daytime
14 reactivity due to BVOCs of 2 ± 2 s⁻¹ (1 σ), see Table 3. During the night, monoterpenes had a
15 larger impact than isoprene due to their known temperature dependency (Kesselmeier and
16 Staudt, 1999). α -terpinene was the most reactive-to-OH BVOC also during nighttime, see
17 Table 3. In terms of absolute values, α -terpinene had a maximum reactivity of 5.3 s⁻¹ on the
18 2nd of August at 2:00 PM LT, which is also when the maximum OH reactivity reported for the
19 whole class of BVOCs occurred. Remarkably, the mean concentration of this compound made
20 it the fourth most abundant BVOC measured, with isoprene being the first (35%), followed by
21 β -pinene (22%), α -pinene (15%), α -terpinene (13%), limonene (9%) and γ -terpinene (1%).
22 The α -terpinene volume mixing ratio was maximum 594 pptv, with an average value between
23 10:00 AM LT and 5:00 PM LT during the field campaign of 131 ± 110 pptv. Its short lifetime
24 is due to the high reaction rate coefficient towards OH, (as reported in literature, i.e. 3.6×10^{-10}
25 cm³ molecule⁻¹ s⁻¹, see Atkinson, (1986) and Lee et al., (2006)), which is more than three-fold
26 higher than the one of isoprene ($k_{\text{isoprene}+\text{OH}} = 1 \times 10^{-10}$ cm³ molecule⁻¹ s⁻¹, Atkinson, 1986). Very
27 little is reported in literature regarding its emission rates and ambient levels in the
28 Mediterranean region. Owen et al., (2001) measured α -terpinene from a few Mediterranean
29 tree species, including: *Juniperus phoenicea*, *Juniperus oxycedrus*, *Spartium junceum* L., and
30 *Quercus ilex*. Ormeno et al., (2007) published the α -terpinene content as 34.9 ± 2.3 $\mu\text{g/gDM}$ in
31 the leaves of *Rosmarinus officinalis*; shrubs of rosemary were present in large quantity around
32 our field site in Corsica.

4.3 Missing reactivity and air masses fingerprint

Figure 2 reports the time series of the total measured OH reactivity and calculated OH reactivity with their associated errors (35% and 20%, respectively). The largest significant discrepancy between reactivities occurred during July 23-30 (56% of missing reactivity, calculated as the difference in percentage between measured and calculated reactivity). We combined air mass backtrajectories and atmospheric mixing ratios of some common atmospheric tracers to determine the chemical fingerprint of the sampled air and to investigate the origin of the missing reactivity. We chose isoprene and pinenes for air masses influenced by biogenic activity, while propane and CO were used for those enriched in anthropogenic pollutants (see supplement). Maximum concentrations of anthropogenic pollutants were measured when the air masses originated from the North East sector: between July 21-23 and between July 31 and August 3, indicating weak pollution events coming from the industrialized areas of the Po Valley and Milan (Italy). The OH reactivity during these events was at most 9 s^{-1} , no significant missing reactivity occurred, mainly because of the weakness of the events, and the deployment of instruments that were able to characterize a large number of AVOCs. On the other hand, biogenic activity was independent of the wind sector and showed a larger variability linked to local drivers, such as air temperature, solar irradiance and wind speed (Figure 6). Biogenic VOCs and their oxidation products had a large influence on the measured OH reactivity and potentially on the missing reactivity (Figures 2-5); however, periods characterized by a large contribution of BVOCs to the measured OH reactivity (27-29 July and 2-4 August) present some differences in missing reactivity, which will be discussed in the next section. There is not a clear trend between the measured and missing OH reactivities with the origin of air masses (Figure 2). Nevertheless, when VOCs were imported from the western sector, the air masses travelled over the sea with an average transport time of up to 48 hours, which can have favored the enrichment in oxygenated compounds through the atmospheric oxidation of primary VOCs initially present in these air masses before they left continents (Michoud et al., 2017).

4.4 Insights into the missing OH reactivity

The missing OH reactivity, obtained as the absolute difference between measured and calculated reactivity values, is reported in Figure 6. Only values higher than the detection limit of 3 s^{-1} are displayed in this figure. Significant missing reactivity ($2.8 \pm 2.2 \text{ s}^{-1}$ on average over the whole campaign) can be observed from 23 to 29 July and more sporadically before

23 July (3 data points over a total of 41) and after 29 July (10 data points). To determine the causes of the reported missing OH reactivity we first investigated whether the VOC instruments deployed on the measurement site did not selectively detect some of the emitted BVOCs. We compared the total monoterpene concentration observed by PTR-MS to the summed monoterpenes concentration from GCs and calculated a concentration difference ranging from 0.2 to 0.6 ppbv (see supplement), the PTR-MS measurements being higher than the sum of speciated monoterpenes. Assuming a reaction rate coefficient with OH of $1.56 \times 10^{-10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ (weighted value estimated from monoterpenes emitted by the plants present at the site, see Bracho-Nunez et al., 2011) we can roughly estimate a value of 0.8-2.3 s^{-1} of missing OH reactivity due to these unspeciated monoterpenes, which represents 30% of the reported missing reactivity on average. Therefore, unmeasured monoterpenes account for a significant fraction of the missing OH reactivity but do not fully explain the total missing reactivity.

The missing OH reactivity temperature dependency was used in previous studies (Di Carlo et al., (2004), Mao et al., (2012) and Hansen et al., (2014)) to determine whether terpenes (Di Carlo et al., 2004) or their oxidation products (Mao et al., 2012) could explain the missing OH reactivity. Terpene emissions depend on temperature following the expression, $E(T) = E(T_s) \exp[\beta(T - T_s)]$, where $E(T_s)$ is the emission rate at temperature T_s , β a temperature sensitivity factor, and T the ambient temperature. Di Carlo et al., (2004) found that the missing OH reactivity reported from a temperate forest in northern Michigan followed the same temperature dependency as terpene emissions, with $\beta = 0.11 \text{ K}^{-1}$. Similarly, Mao et al., (2012) reported a β factor of 0.168 K^{-1} from a temperate forest in California. Since this value is higher than the upper limit of 0.144 usually observed for terpene emissions, the authors attributed the missing OH reactivity to unmeasured oxidation products of BVOCs, which was supported by box model simulations.

In the present study, we investigated the temperature dependency of the missing OH reactivity to determine whether BVOC oxidation products caused the missing reactivity for 23-26 and 27-29 July periods. We found a clear dependence ($\beta = 0.158 \text{ K}^{-1}$ and $R^2 = 0.643$) between the missing OH reactivity and temperature for some days of the campaign (27th, 28th, 29th), while no correlation ($\beta = 0.007 \text{ K}^{-1}$ and $R^2 = 0.001$) was found during 23-26 July, see Figure 7. Indeed, the lack of a clear temperature dependency and the influence of long range transport on air masses imported from the western sector, as demonstrated by Michoud et al., (2017) for

the same data set suggest that the local oxidation of BVOCs was not the main cause of the missing reactivity observed during 23-26 July. In contrast, the similar temperature dependencies observed in the study of Mao et al., (2012) and during 27-29 July suggests that oxidation products of locally emitted BVOCs was the main cause of missing reactivity for this period. This interpretation is further confirmed by a significant increase of the concentrations of some BVOC oxidation products during 27-29 July (Figure 6) and the accumulation of these secondary species versus the primary biogenic compounds during nighttime (Figure 4 supplement).

Figure 6 shows the missing OH reactivity and volume mixing ratios of BVOCs and their oxidation products together with meteorological parameters, such as temperature, wind speed and solar irradiance. Compounds are reported using their protonated mass measured by PTR-MS, such as: m/z 69 (isoprene) and m/z 137 (monoterpenes) for primary emissions, m/z 71 (isoprene first generation oxidation products: Methyl Vinyl Ketone (MVK) + methacrolein (MACR) + possibly fragments of isoprene hydroxyperoxides (ISOPOOH)), m/z 139 (nopinone, β -pinene first generation oxidation product) and m/z 151 (fragment of pinonaldehyde, α -pinene first generation oxidation product) for the BVOC oxidation products. It is worth noting that m/z 111 and m/z 113 are also reported since these masses have been attributed to oxidation products of several terpenes. Indeed m/z 111, m/z 113 and m/z 151 were observed in chamber and field studies (Lee et al., 2006, Holzinger et al., 2005) as they are formed from the photo-oxidation of different terpenes; highest yields were attributed to terpenes also common to the Mediterranean ecosystem, such as myrcene, terpinolene, linalool, methyl-chavicol and 3-carene (Lee et al., 2006, Bracho-Nunez et al., 2011). We note that all the above-mentioned masses (with the exception of m/z 111 and m/z 113, for which no rate coefficient was found for the reaction of the unprotonated molecule with OH) have been taken into account in the calculated OH reactivity. The reported time series show that both primary BVOCs and most of the OVOCs resulting from their oxidation had a clear diurnal profile with highest values during midday, similar to that observed for the missing reactivity.

Most of the nights were associated with low mixing ratios of BVOCs (29 pptv on average for isoprene) and their oxidation products (40 pptv on average for m/z 71) and corresponded to very low or no missing reactivity. Nevertheless, we note that during some nights (26 to 27, 27 to 28, 29 to 30 July), some compounds (m/z 71, m/z 113) were also present in low amounts

1 and a missing reactivity of about 4 s^{-1} was noticed. These nights were associated with low
2 wind speed and high temperatures, indicating a stagnant episode that could have favored the
3 accumulation of unmeasured oxidation products during 26-29 July (ratios of $m/z\ 71$ -to- $m/z\ 69$
4 and nopinone-to-bpinene were maximum 13.9 and 7.7, respectively, see supplement) as was
5 already observed by night in an oak-forest canopy (Zannoni et al., 2016). The mixing ratios of
6 the first- oxidation products of isoprene and monoterpenes ($m/z\ 71$, $m/z\ 139$) strongly
7 increased after July 26 when the wind speed was lower and increased again after July 27
8 when air temperature increased. In contrast, during another period of BVOCs influence as 2-5
9 August, slightly lower air temperature was experienced, less accumulation of oxidation
10 products (Figure 4 supplement) and lower concentrations of secondary species ($m/z\ 139$, m/z
11 151) were reported (Figure 6). These pieces of evidence also suggest that unmeasured, locally
12 generated BVOC oxidation products contribute significantly to the missing OH reactivity
13 during 27-29 July.

14 5 Conclusions

15 The total OH reactivity was used in this study to evaluate the completeness of the
16 measurements of reactive trace gases at a coastal receptor site in the western Mediterranean
17 basin during three weeks in summer 2013 (16 July 2013-05 August 2013).

18 The OH reactivity had a clear diurnal profile and varied with air temperature, suggesting that
19 biogenic compounds were significantly affecting the local atmospheric chemistry. Ancillary
20 trace gases measurements confirmed that most of the daytime reactivity was due to biogenic
21 VOCs, including relevant contributions from oxygenated VOCs, while during nighttime
22 inorganic species and oxygenated VOCs exhibited the largest contribution. The OH reactivity
23 was on average $5\pm4\text{ s}^{-1}$ (1σ) with a maximum value of $17\pm6\text{ s}^{-1}$ (35% uncertainty). The
24 observed maximum is comparable to values of OH reactivity measured at forested locations
25 in northern latitudes (temperate and boreal forests as reported by Di Carlo et al., 2004, Ren et
26 al., 2006, Sinha et al., 2010, Noelscher et al., 2013, Kumar and Sinha 2014, Nakashima et al.,
27 2014). This finding highlights the importance of primary-emitted biogenic VOCs on the OH
28 reactivity, especially where air temperature and solar radiation are high; even though our site
29 was specifically selected for a focused study on mixed and aged continental air masses
30 reaching the basin.

31 A comparison between the measured OH reactivity and the summed reactivity calculated
32 from the measured species showed that on average 56% of the measured OH reactivity was

not explained by the gas measurements during July 23-29. During this period, the air masses came from the West (July 23-27) and the South (July 27-29); lack of pollution events, calm wind conditions and peaks of air temperature were registered at the field site (28 July). Specifically, during July 27-29 we also detected an increase in oxygenated VOC concentrations originating from the photo-oxidation of primary-emitted BVOCs, **presumably due to an accumulation favored by stagnant atmospheric conditions**. Highest yields of these oxidation products (m/z 111, m/z 113, m/z 151) have been attributed to terpenes, which are strongly emitted by Mediterranean ecosystems (Lee et al., 2006, Bracho-Nunez et al., 2011). We found that the missing reactivity was temperature dependent for the same period and such dependency suggests that unmeasured oxidation products of BVOCs, as similarly reported by the study of Mao et al., (2012), caused the missing reactivity.

The impact of biogenic VOCs and their oxidation products is weaker during 23-26 July, since the site received air masses from West, therefore from the marine sector that is less exposed to biogenic emissions. However, large values of missing OH reactivity were also observed, similar to those observed on 27-29 July. The lack of a temperature dependence of the missing reactivity for 23-26 suggests that species other than BVOC oxidation products locally formed were also contributing to the missing OH reactivity.

Mediterranean plants are known to emit large quantities of reactive BVOCs, including sesquiterpenes and oxygenated terpenes (Owen et al., 2001), which were not investigated during our fieldwork. It could be possible that these molecules, as well as their oxidation products, have also played an important role in the missing OH reactivity detected during the campaign.

Further studies with chemical and transport models to identify the important chemical functions of these oxygenated molecules, as well as the effects of long-range transport would be beneficial to provide further insights.

Finally, as the Mediterranean basin differs from side to side, (air masses reception as well as type of ecosystems) more intensive studies at different key spots, e.g. western vs. eastern basin and remote vs. periurban ecosystems, would be helpful for a better understanding of the atmospheric processes linked to the reactive gases over the Mediterranean basin.

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Table 1. Measured compounds (whose concentration was above the instrumental detection limits) and their adopted classification in the manuscript for calculating the OH reactivity. Anthropogenic VOCs, BVOCs and OVOCs stand respectively for anthropogenic, biogenic and oxygenated volatile organic compounds.

Species group	Species name
AVOCs	methane, ethane, propane, n-butane, n-pentane, n-hexane, n-octane, n-nonane, n-undecane, n-dodecane, 2-methylpentane, 2-methylhexane, 2,2- dimethylbutane, 2,2-dimethylpropane, 2,3- dimethylpentane, 2,4-dimethylpentane, 2,2,3-trimethylbutane, 2,3,4- trimethylpentane, cyclohexane, ethylene, propylene, 1-butene, 2-methyl-2-butene, 3-methyl-1-butene, 1,3-butadiene, <i>trans</i> -2-butene, <i>cis</i> -2-butene, 1-pentene, <i>trans</i> -2-pentene, <i>cis</i> -2-pentene, hexene, benzene* ¹ , toluene* ² , ethylbenzene, styrene, m-xylene, o-xylene, p-xylene* ³ , acetylene, 1-butyne, acetonitrile.
BVOCs	isoprene, α -pinene, β -pinene, D-limonene, α -terpinene, γ -terpinene, camphene.
OVOCs	Acetaldehyde* ⁴ , formic acid* ⁵ , acetone* ⁶ , acetic acid* ⁷ , mglyox, methyl ethyl ketone* ⁸ , propionic acid, ethyl vinyl ketone, butiric acid, nopinone, pinonaldehyde, methacrolein, methyl vinyl ketone, formaldehyde* ⁹ , methanol* ¹⁰ .
Others	NO, NO ₂ , CO.

*Compounds with anthropogenic and biogenic origin. 1,2,3. Misztal et al., 2015. 4. Millet et al., 2010. 5, 7. Paulot et al., 2011. 6. Jacob et al., 2002. 8. Yanez-Serrano et al., 2016 and Cappelin et al, 2017. 9. Fortems-Cheiney et al., 2012. 10. Jacob et al, 2005.

Table 2. Summary of the experimental methods deployed during the field campaign and needed for calculating the OH reactivity. The number of measured compounds includes the compounds below the instrumental detection limit (LoD).

Technique	Compounds measured	LoD (pptv)
PTR-MS	16 VOCs	7-500
GC- FID/FID	43 NMHCs C2-C12	10-100
GC-FID/MS	16 NMHCs (OVOCs+ C3-C7)	5-100
off-line GC-FID/MS	35 NMHCs C5-C16 + 5 aldehydes C6-C12	5-40
Hantzsch reaction	HCHO	130
CLD	NO _x	50
WS-CRDS	CO ₂ , CH ₄ , CO	1000

Table 3. Relative contributions of individually detected biogenic volatile organic compounds (BVOCs) to the total calculated OH reactivity BVOCs fraction. Daytime BVOCs OH reactivity accounted for a maximum value of 9 s⁻¹, on average it was 2±2 s⁻¹. Nighttime BVOCs OH reactivity fraction accounted for a maximum value of 0.5 s⁻¹, on average it was 0.1 s⁻¹.

BVOCs	Day (%)	Night (%)
α-pinene	7.7	20.7
β-pinene	16.5	16.1
limonene	12	11.4
camphene	1.5	3.1
α-terpinene	31.1	31.3
γ-terpinene	1.3	5
isoprene	30	12.5



Figure 1. Field site top-view, Corsica, France (42.97°N, 9.38°E, altitude 533 m). Measures: 1. PTR-MS, online and offline chromatography for trace gases analysis; 2. OH reactivity; 3. NO_x, O₃, aerosols composition and black carbon; 4. Meteo, and particles microphysics; 5. HCHO, trace gases and radicals; 6. CO, CO₂, CH₄; 7. Trace gases and particle filters; 8. Particles physics. The photo was taken during the installation of the instruments.

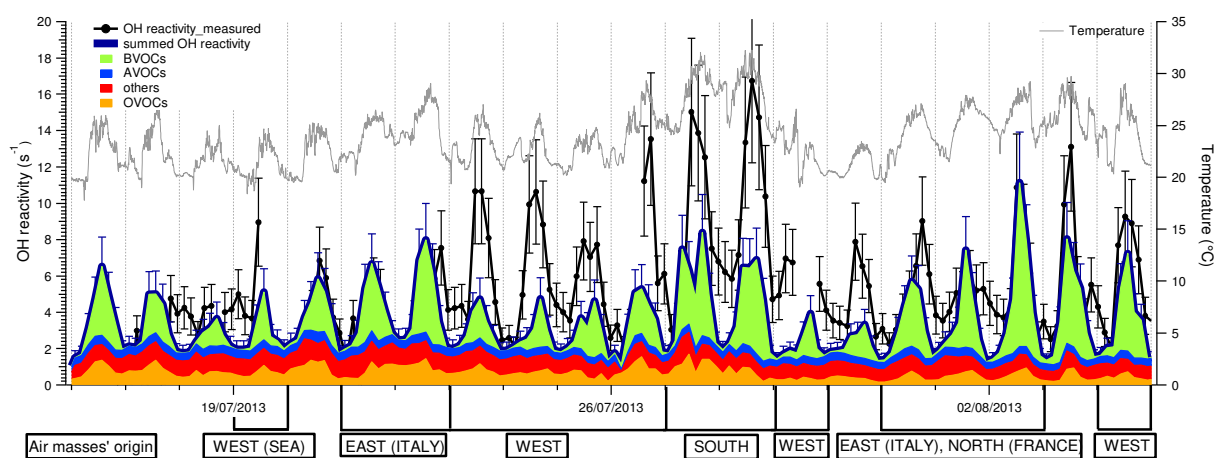


Figure 2. Three-hours averaged data of total OH reactivity measured and calculated from the measured gases. Summed OH reactivity is represented with the blue thick line and grouped as biogenic VOCs in green, anthropogenic VOCs in blue, oxygenated VOCs in orange and others in red. Others refer to carbon monoxide (CO) and nitrogen oxides (NO_x).

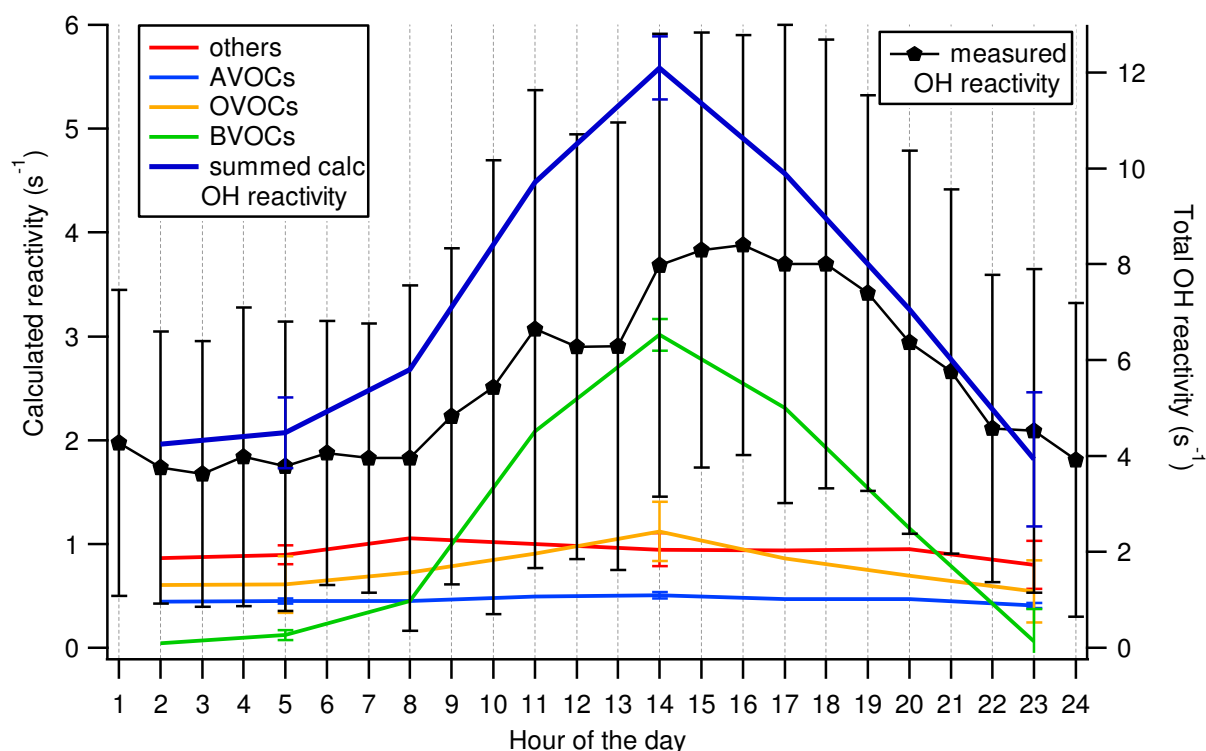


Figure 3. Diurnal patterns of measured (value with $\pm 1\sigma$, right axis) and calculated OH reactivity (left axis). Others, AVOCs, OVOCs, BVOCs are the contribution of CO and NO_x (others), anthropogenic volatiles, oxygenated volatiles and biogenic volatiles to the summed calculated OH reactivity.

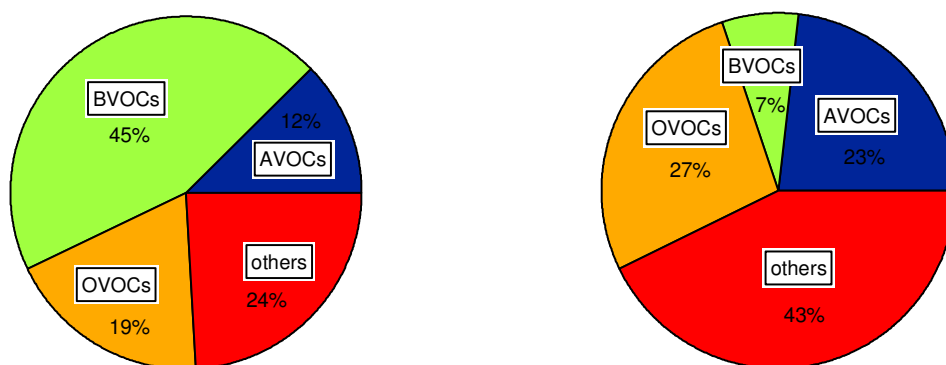


Figure 4. Daytime (left pie) and nighttime (right pie) contributions of the measured compounds to the calculated OH reactivity. Daytime data were collected between 07.30 and 19.30 while nighttime data were between 19.30 and 07.30. Summed OH reactivity during daytime was maximum 11 s^{-1} , on average $4 \pm 2 \text{ s}^{-1}$; while during nighttime it was maximum 3

s⁻¹, on average 2±0.4 s⁻¹. Biogenic VOCs (green), AVOCs (blue), OVOCs (orange) and others (red) stand for biogenic, anthropogenic, oxygenated volatile organic compounds and carbon monoxide and nitrogen oxides, respectively. During daytime, BVOCs, AVOCs, OVOCs and others contributions were 45%, 12%, 19%, 24%, respectively; while they were 7%, 23%, 27%, 43%, respectively during nighttime.

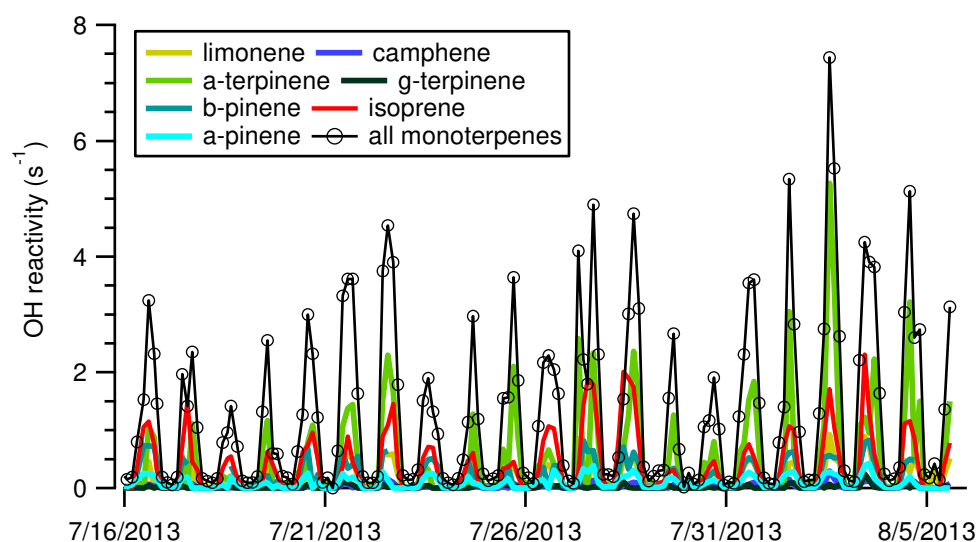


Figure 5. Absolute OH reactivity calculated for the measured biogenic compounds.

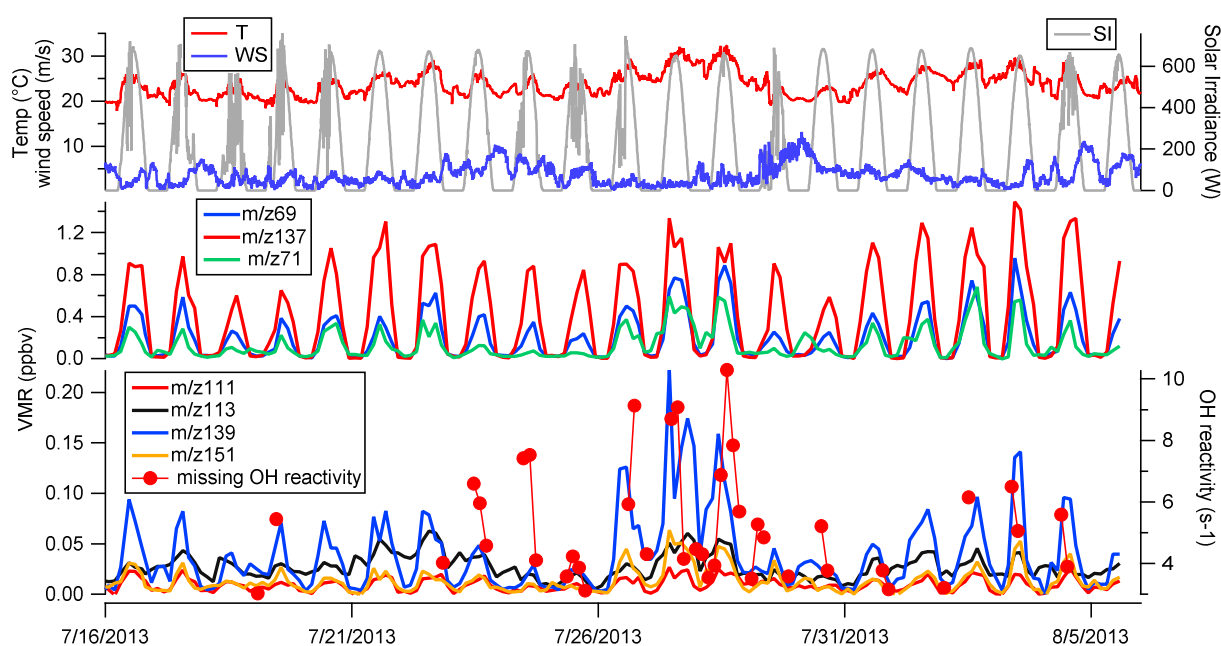


Figure 6. Volume mixing ratios (ppbv) of primary-emitted and secondary produced biogenic volatile organic compounds (BVOCs) measured by PTR-MS. Primary emitted BVOCs include: isoprene (m/z 69) and monoterpenes (m/z 137), oxidation products include: methyl vinyl ketone, methacrolein, possibly isoprene hydroperoxides MVK+MACR+ISOPOOH (m/z

71), nopinone (m/z 139), pinonaldehyde (m/z 151), m/z 111 and m/z 113. Top panel provides data of temperature, wind speed and solar irradiance. The lowest panel shows the variability of the missing OH reactivity, calculated as the difference between measured and calculated reactivity.

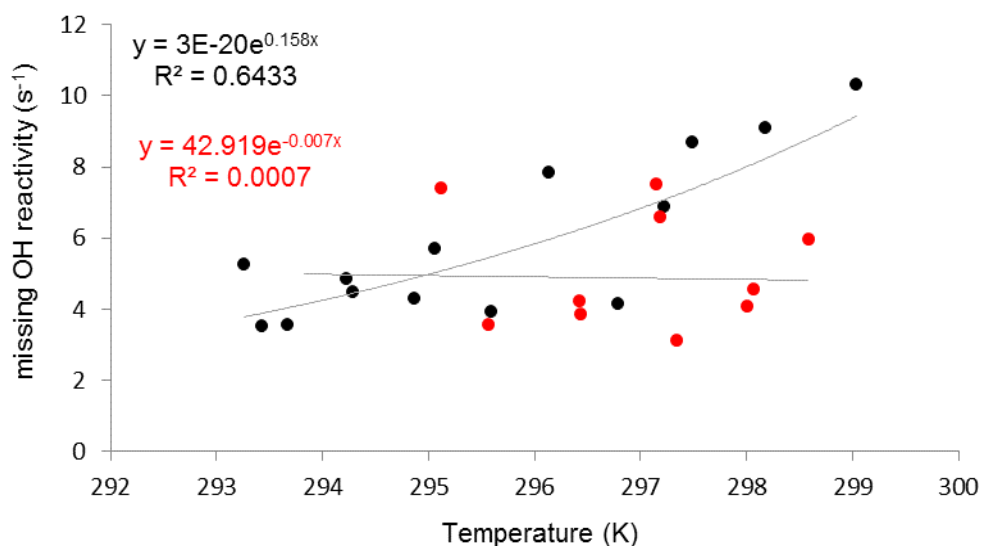


Figure 7. The difference between measured and calculated reactivity (missing OH reactivity) during July 23 -26 (red data points) and during July 27 -29 (black data points), dependence to temperature. The missing reactivity is fitted to $E(T)=E(293) \exp(\beta(T-293))$.