

Reactivity study of 3,3-dimethylbutanal and 3,3-dimethylbutanone: Kinetic, reaction products, mechanisms and atmospheric implications

Inmaculada Aranda¹, Sagrario Salgado^{1,2}, Beatriz Cabañas^{1,2}, Florentina Villanueva² and Pilar Martín^{1,2*}

¹Universidad de Castilla-La Mancha, Departamento de Química Física, Facultad de Ciencias y Tecnologías Químicas, Avda. Camilo José Cela s/n, 13071, Ciudad Real, Spain.

²Universidad de Castilla-La Mancha, Instituto de Investigación en Combustión y Contaminación Atmosférica (ICCA), Camino de los Moledores s/n, 13071, Ciudad Real, Spain.

*Corresponding author. Email address: mariapilar.martin@uclm.es

Abstract: 3,3-dimethylbutanal (33DMbutanal, $(\text{CH}_3)_3\text{CCH}_2\text{C}(\text{O})\text{H}$) and 3,3-dimethylbutanone (33DMbutanone, $(\text{CH}_3)_3\text{CC}(\text{O})\text{CH}_3$) are carbonyl compounds that could play a key role in tropospheric chemistry. To better understand the effects of carbonyl compounds in the atmosphere, a kinetic and mechanistic study was conducted on the degradation of 33DMbutanal and 33DMbutanone with atmospheric oxidants (Cl atoms, OH and NO_3 radicals). The kinetic experiments were performed at 710 ± 30 Torr and at room temperature (298 ± 5 K) using a relative method and FTIR (Fourier Transform Infrared Spectroscopy) to monitor the reactions. The rate coefficients (k in units of $\text{cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$) obtained were: $k_{\text{Cl}+33\text{DMbutanal}} = (1.27 \pm 0.08) \times 10^{-10}$, $k_{\text{Cl}+33\text{DMbutanone}} = (4.22 \pm 0.27) \times 10^{-11}$, and $k_{\text{OH}+33\text{DMbutanone}} = (1.26 \pm 0.05) \times 10^{-12}$. The reaction products were also determined using FTIR and GC-MS (Gas Chromatography/Mass Spectrometry). The main products observed were carbonyl compounds, including acetone, formaldehyde and 2,2-dimethylpropanal. In the presence of NO, nitrated compounds were also formed, and at high NO_2 concentrations peroxyacetyl nitrate (PAN) and peroxy-3,3-dimethylbutyryl nitrate were identified. Other unquantified compounds were multifunctional organic compounds and organic acid of low volatility. Both 33DMbutanal and 33DMbutanone degrade rapidly near emission sources with minimal impact on radiative forcing. However, they may contribute to tropospheric ozone, with a Photochemical Ozone Creation Potential (POCP) range of 15-69, and secondary organic aerosol formation, potentially worsening air quality and contributing to photochemical smog.

28 1 Introduction

29 Carbonyl compounds are a group of oxygenated volatile organic compounds (OVOCs) that are emitted into the
30 atmosphere from natural and anthropogenic sources (Bao et al., 2022), but they are also formed in the atmosphere
31 as oxidation products of other volatile organic compounds (VOCs) (Mellouki et al., 2015). It is well established that
32 OVOCs play an important role in the sequence of chemical reactions that leads to their further oxidation and
33 contributes to the tropospheric ozone formation in both polluted and remote environments with important effect on
34 health as is the case of formaldehyde and acetaldehyde (Zhou et al., 2023; Liu et al., 2022; Mellouki et al., 2015;
35 Calvert et al., 2011). In addition, large carbonyl compounds could influence on climate change altering the Earth's
36 radiation balance if they are strong infrared light absorbers and their atmospheric concentrations are sufficiently
37 high. Additionally, carbonyls could be an important source of aerosol which could further affect radiation balance
38 and be hazardous for health (Liu et al., 2022; Heald and Kroll, 2020).

39 The rising O₃ levels in mega-city clusters like Chinese cities underscore the critical need for effective control of
40 ambient carbonyls, significant precursors of O₃. Moreover, as intermediate products of hydrocarbon oxidation,
41 carbonyls likely play a pivotal role in minimizing the disparity between atmospheric reactivity in measurements and
42 simulations. Previous studies (Zhou et al., 2023; Liu et al., 2022; Mellouki et al., 2015; Calvert et al., 2011) have
43 provided valuable insights into carbonyls' presence, composition, origins, and impact on O₃ and SOA formation,
44 using a combination of field measurements, numerical simulations, and laboratory experiments. Nevertheless,
45 further research is still warranted to achieve a more comprehensive understanding of carbonyls' sources and sinks,
46 given the complexity of their emission and degradation processes (Liu et al., 2022).

47 In this work, the tropospheric reactivity of two carbonyls compounds whose reactivity is not yet completely
48 established, has been studied: 3,3-dimethylbutanal (33DMbutanal, (CH₃)₃CCH₂C(O)H) and 3,3-dimethylbutanone
49 (33DMbutanone, (CH₃)₃CC(O)CH₃). These two carbonyls are among the reaction products identified in the
50 atmospheric degradation of two alcohols (3,3-dimethyl-1-butanol and 3,3-dimethyl 2-butanol), whose reactivity has
51 been previously studied (by our research group) (Colmenar et al., 2020). On the other hand, 33DMbutanal has also
52 been detected as reaction product in the reaction of 2,4,4-trimethyl-1-pentanol with Cl atoms (Vila et al., 2020) and
53 it could be an intermediate in the synthesis of neotame, a sweetener (Tanielyan and Augustine, 2012). Industrially,
54 33DMbutanone, known as methyl tert-butyl ketone, is produced for use in fungicides, herbicides and pesticides (Liu
55 et al., 2022) and it might also be used as a solvent for the extraction of methylphenols from wastewater (Xiong et
56 al., 2018). Specifically, in the study of Byrne et al. (2018) 33DMbutanone has been identified as potential
57 replacements for hazardous volatile non-polar solvents such as toluene, due to the low toxicity and good solvation
58 characteristics. In addition to direct emissions, 33DMbutanone could be present in the atmosphere as a reaction
59 product of the gas-phase oxidation of 2,2-dimethylbutane and 3,3-dimethyl-2-butanol (Saunders et al., 2003; Jenkin
60 et al., 1997). Although there are no atmospheric concentration measurements for these compounds, their current and
61 potential uses justify the need to understand their atmospheric reactivity and degradation processes.

62 Specifically, for 33DMbutanal and 33DMbutanone few studies about their atmospheric reactivity have been reported
63 in the literature. In the case of the reaction of 33DMbutanal with OH radicals, the experimental rate coefficient has
64 been measured by Aschmann et al. (2010) and D'Anna et al. (2001). Only one study on the reaction products with
65 OH radicals has been reported by Aschmann et al. (2010). For the reaction of 33DMbutanal with NO₃ radicals two
66 kinetic studies are available in the literature (D'Anna et al., 2001; D'Anna and Nielsen 1997). Tadic et al., 2012 has

Con formato: Inglés (Estados Unidos)

67 reported the photochemical parameters of 33DMbutanal due to the importance of the photodissociation of aldehydes
 68 in the atmosphere since it could represent an important source of free radicals. To our knowledge, there are no data
 69 about the reaction of 33DMbutanal with Cl atoms.

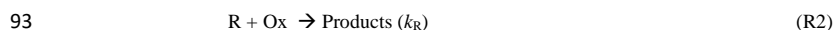
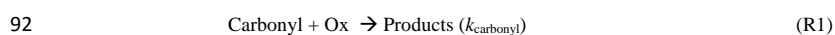
70 In the case of 33DMbutanone only a kinetic study with Cl atoms has been carried out (Farrugia et al., 2015) and two
 71 studies with OH radicals (Mapelli et al., 2023; Wallington and Kurylo 1987). In the studies of OH radicals, the rate
 72 coefficient has been obtained at different temperatures and at low pressure, using absolute methods. No studies have
 73 been carried out on the reaction products of Cl atoms and OH radicals with 33DMbutanone that could help to
 74 establish the reaction mechanisms.

75 Taking the above into consideration, the aim of this work is to complete the studies about the reactivity of
 76 33DMbutanal and 33DMbutanone to further understand their atmospheric chemistry in particular and the carbonyls
 77 reactivity in general. For this purpose, the kinetic study has been conducted for the reactions of 33DMbutanal and
 78 33DMbutanone with Cl atoms and 33DMbutanone with OH radicals using a relative method and FTIR (Fourier
 79 Transform Infrared Spectroscopy) technique as detection system. Additionally, for the reactions of 33DMbutanal
 80 with Cl atoms, OH and NO₃ radicals and for the reactions of 33DMbutanone with Cl atoms and OH radicals a
 81 complete reaction product study has been performed using FTIR and GC-MS (Gas Chromatography/Mass
 82 Spectrometry) techniques. This work is to date the first kinetic study reported in bibliography for the reaction of
 83 33DMbutanal with Cl atoms and the first study on reaction products and mechanisms for the reactions of
 84 33DMbutanone with Cl atoms and OH radicals, and 33DMbutanal with Cl atoms and NO₃ radicals. Additionally,
 85 this work includes a study on the reaction products for the reaction of 33DMbutanal with OH radicals in order to
 86 confirm the mechanism proposed by Aschmann et al. (2010).

87 2 Experimental Section

88 2.1 Rate coefficients determination: relative method

89 Rate coefficients have been determined using a relative rate method on the assumption that the organic compound
 90 (carbonyl: 33DMbutanal or 33DMbutanone), and the reference compound (R) are removed solely by their reactions
 91 with the oxidants (Ox: Cl or ·OH):



94 where k_{carbonyl} and k_{R} are the rate coefficients of the carbonyl and the reference compound, respectively. The kinetic
 95 treatment for the reactions (R1) and (R2) gives the following relationship:

$$96 \quad \ln \left(\frac{[\text{carbonyl}]_0}{[\text{carbonyl}]_t} \right) = \frac{k_{\text{carbonyl}}}{k_{\text{R}}} \times \ln \left(\frac{[\text{R}]_0}{[\text{R}]_t} \right) \quad (\text{I})$$

97 where $[\text{carbonyl}]_0$, $[\text{R}]_0$, $[\text{carbonyl}]_t$, $[\text{R}]_t$, are the initial concentrations and those at time t for the carbonyl and the
 98 reference compound, respectively. At least three reference compounds were employed for each studied reaction, and
 99 the experiments were performed in triplicate for each one. According to Eq. (I), a plot of $\ln([\text{carbonyl}]_0/[\text{carbonyl}]_t)$
 100 versus $\ln([\text{R}]_0/[\text{R}]_t)$ should give a linear fit with an intercept equal to zero. The slope of the plot corresponds to the

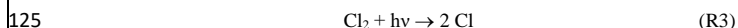
ratio of the rate coefficients ($k_{\text{carbonyl}}/k_{\text{R}}$). Therefore, the value of k_{carbonyl} can be determined if the rate coefficient of the reference compound (k_{R}) is known.

2.2 Experimental systems and procedure

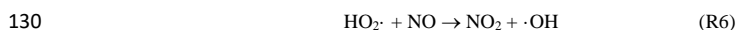
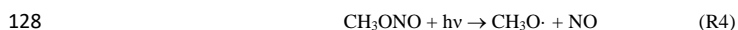
Experimental details can be found in previous publications (Aranda et al., 2021; Colmenar et al., 2020, 2018). Therefore, only a brief description is provided here.

Kinetic and product studies were performed at room temperature (298 ± 5 K) and atmospheric pressure (710 ± 18 Torr) employing a 50 L Pyrex® gas cell as reaction chamber, coupled to a FTIR spectrometer (Thermo, Nicolet 6700). The gas cell contains a multireflection system that allows a maximum optical path of 200 m (Saturn Series Multi-Pass cell). For the FTIR spectra collection, a total of 60 interferograms were co-added over 98 s, usually taken in the range of 650–4000 cm^{-1} with a resolution of 1 cm^{-1} . Additionally, for product identification, samples of the reaction mixture were collected via a port on the gas cell using a Solid Phase MicroExtraction (SPME) fiber (DVB/CAR/PDMS) as the pre-concentration passive sampling method. After the adsorption process (5–8 minutes), the fiber was transferred to the injection port of the gas chromatograph (GC) coupled to a Time-of-Flight Mass Spectrometer analyzer (MSTOF, AccuTOF GCv/Jeol). In the injection port of the gas chromatograph, the compounds were desorbed at 250 °C, separated and detected by GC-MSTOF. Two different capillary columns with the same characteristics were used: a TRB-1701 (Teknokroma, 30 m \times 0.32 mm \times 1 μm) and a Equity™ – 1701 (Supelco, 30 m \times 0.32 mm \times 1 μm). Once in the mass spectrometer, the compounds were ionized (Electron Ionization (EI) and/or Field Ionization (FI)) and fragmented to obtain their mass spectrum (MS). The chromatographic conditions used for the analysis were as follows: injection port, 250 °C; interface, 250 °C; oven initial temperature of 40 °C for 4 min; ramp, 25 °C min^{-1} to 120 °C, held for 10 min; second ramp, 20 °C min^{-1} to 200 °C, held for 2 min.

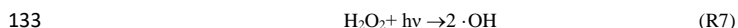
The oxidants were generated by photolysis or decomposition of a precursor (Finlayson-Pitts and Pitts 2000). The generation of Cl atoms was achieved through the photolysis of Cl_2 using radiation emitted by actinic lamps ($\lambda_{\text{max}} = 365$ nm).



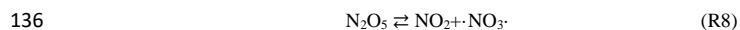
Hydroxyl radicals ($\cdot\text{OH}$) were generated by the photolysis of methyl nitrite (CH_3ONO) in air in the presence of NO , following the reaction sequence below:



Some experiments have been carried out using H_2O_2 as precursor of $\cdot\text{OH}$ and UV radiation ($\lambda=254$ nm) in a Quartz gas cell reactor.



Nitrate radicals (NO_3^\cdot) were generated through the thermal decomposition of dinitrogen pentoxide (N_2O_5) at room temperature, according to the following equilibrium



The kinetic experiments were conducted in a nitrogen atmosphere for reactions with Cl atoms, while synthetic air was used for reactions with OH radicals. All the experiments conducted for the study of products were carried out in synthetic air and in the absence of the reference compound.

The reactions were followed by measuring the absorbance of the characteristic IR bands of each organic compound (33DMbutanal/33DMbutanone and the reference compounds in the case of kinetic analysis) at different reaction times. The IR spectra were processed using OMNIC software, through a subtraction procedure (elimination) of the IR bands.

The concentration ranges (in ppm) used in the kinetic experiments were: 10-12 for 33DMbutanal and 33DMbutanone, 9-10 for 1-butene, 10-14 for propene, 35-40 for 2-methylpropene, 13-14 for isopropanol, 10-17 for cyclohexane, 5-14 for propanal, 9-10 for 2-methyl-2-butanol, 5-6 for ethyl formate, 11-15 for 1-butanol, 17-22 for Cl_2 , 15-20 for NO and 16-20 for methyl nitrite. In the case of the reaction product experiments, the typical concentration (in ppm) was: 10-14 for 33DMbutanal and 33DMbutanone; 22 for Cl_2 , 15-20 for NO, 16-20 for methyl nitrite, 30 for H_2O_2 and for 14-25 for N_2O_5 .

2.3 Materials

Information on the purity and supplier company of the reagents used to carry out the experiments is specified below: 33DMbutanal (95%), 33DMbutanone (97%), and the reference compounds: 1-butene and propene ($\geq 99\%$), 2-methylpropene ($\geq 99.5\%$), isopropanol (70% in H_2O), propanal (97%), cyclohexane (99.5%), 2-methyl-2-butanol ($\geq 99\%$), ethyl formate (97%), 1-butanol ($\geq 99\%$), 2,2-dimethylpropanoic acid (99%), 3,3-dimethylbutanoic acid (98%) from Sigma Aldrich. Acetone ($\geq 99.5\%$) from Supelco. 2,2-dimethylpropanal ($> 95\%$) from TCI. The precursors of the radicals were: methyl nitrite, synthesized in the laboratory according to the method of Taylor et al., (1980) and Cl_2 (99%) from Praxair. NO (98.5%) from Air Liquide, H_2O_2 (50 wt. % in H_2O , stabilized) from Sigma Aldrich. N_2 (99.999%) and synthetic air (99.999%) from Praxair. Dinitrogen pentoxide (N_2O_5) synthesized in the laboratory according to the procedure described by Schott and Davids (1958).

3 Results and discussion

3.1 Kinetic study

The reference compounds have been selected according to the following conditions: first that at least one active IR band does not overlap with those of the compound under study (33DMbutanal or 33DMbutanone) and second, that $0.1 \leq k_{\text{carbonyl}}/k_{\text{R}} \leq 10$. In addition, a series of experiments was carried out in order to evaluate possible heterogeneous reactions with the walls, reactions between the compound under study and the reference compound, photolysis of any of them and/or reactions with the oxidant precursor. To determine these loss processes, the initial spectrum of the each reactants was compared with the spectrum recorded after a long period (45 minutes, which is the typical time range for the kinetic study). The results of these experiments showed that under the experimental conditions

used in this work, the losses of the reactants due to these processes were negligible (< 3% photolysis and/or dark loss in the case of 33DMbutanal and 0% for 33DMbutanone).

The IR absorption bands used to follow the evolution of the different compounds were: 33DMbutanal, 2700 cm^{-1} ; 1-butene, 911 cm^{-1} ; propene, 878-942 cm^{-1} ; 2-methylpropene, 912 cm^{-1} ; isopropanol, 1070 and 1251 cm^{-1} ; cyclohexane, 2862 and 2933 cm^{-1} ; 33DMbutanone, 1137 cm^{-1} ; propanal, 2710 cm^{-1} ; 2-methyl-2-butanol, 883 cm^{-1} ; ethyl formate, 1192 and 1194 cm^{-1} ; 1-butanol, 1060 cm^{-1} .

The plots of $\ln([\text{carbonyl}]_0/[\text{carbonyl}]_t)$ versus $\ln([R]_0/[R]_t)$ for each reaction were generated according to Eq. (I). As an example, Fig. 1 shows the plot of this equation for the reaction of 33DMbutanal with Cl atoms, along with two of the reference compounds used. At least three reference compounds were used for each reaction, to ensure the accuracy of the determined value. The reference compounds used and the values of their rate coefficients are included in Table 1.

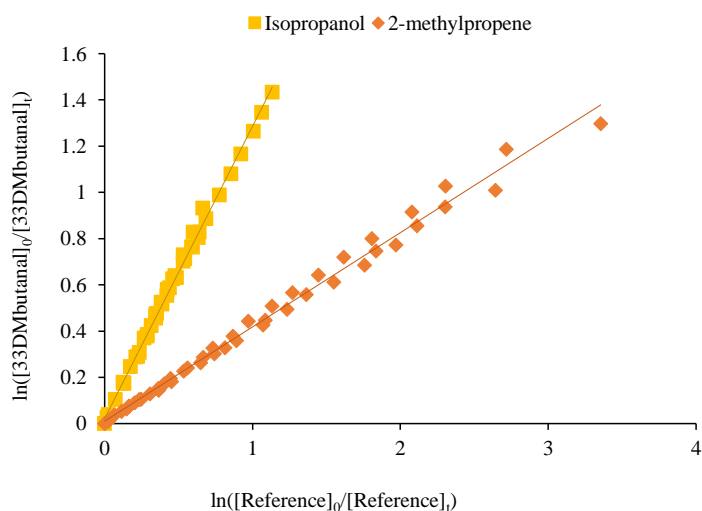


Figure 1. Plots of Eq. (I) for the reaction of 33DMbutanal with Cl atoms using two reference compounds.

The slopes of the plots correspond to the relationship k_{carbonyl}/k_R , knowing the value of k_R , the rate coefficients of carbonyls can be determined. It can be seen the good linear fit with an intercept close to zero, indicating the absence of secondary reactions. Plot of Eq. (I) depicting the reaction of 33DMbutanal with Cl atoms, using cyclohexane as a reference, is presented in Fig. 1Sa. Additionally, the Fig. 1S includes the plots for the reaction of 33DMbutanone with Cl atoms and OH radicals, along with all the reference compounds used. The plots show a linear fit with $r^2 \sim 0.99$ and low origin intercept values, indicating the absence of secondary reactions. The dispersion of the data observed in the -OH reaction with 33DMbutanone (Fig. 1Sc) is due to the complexity of the analysis, especially for slow reactions, and the overlapping of the bands of the precursor (methyl nitrite). The k_{carbonyl}/k_R and k_{carbonyl} obtained are shown in Table 1.

Table 1. Summary of relative and absolute rate coefficients for the reaction of 33DMbutanal with Cl atoms and 33DMbutanone with Cl atoms and OH radicals.

Reaction	Reference compound	$k_{\text{carbonyl}}/k_{\text{R}} \pm 2\sigma$	$^b k_{\text{carbonyl}} \pm 2\sigma$
33DMbutanal+Cl	2-methylpropene	0.38±0.01	1.26 ± 0.27
	$^a k_{\text{Cl}} = (3.3 \pm 0.7) \times 10^{-10}$	0.40±0.01	1.32 ± 0.28
		0.44±0.01	1.46 ± 0.31
	Isopropanol	1.38±0.02	1.20 ± 0.31
	$^a k_{\text{Cl}} = (0.87 \pm 0.23) \times 10^{-10}$	1.25±0.02	1.09 ± 0.28
		1.27±0.02	1.11 ± 0.29
	Cyclohexane	0.43±0.01	1.42 ± 0.22
	$^a k_{\text{Cl}} = (3.3 \pm 0.5) \times 10^{-10}$	0.36±0.01	1.19 ± 0.18
		0.42±0.02	1.40 ± 0.22
Weighted Average			1.27 ± 0.08
	Reference compound	$k_{\text{carbonyl}}/k_{\text{R}} \pm 2\sigma$	$^c k_{\text{carbonyl}} \pm 2\sigma$
33DMbutanone+Cl	Propanal	0.41 ± 0.02	5.10 ± 1.06
	$^a k_{\text{Cl}} = (12.5 \pm 2.5) \times 10^{-11}$	0.37 ± 0.01	4.62 ± 0.93
		0.42 ± 0.01	5.22 ± 1.06
		0.42 ± 0.01	5.31 ± 1.07
	2-methyl-2-butanol	0.42 ± 0.01	3.03 ± 0.76
	$^a k_{\text{Cl}} = (7.30 \pm 1.80) \times 10^{-11}$	0.43 ± 0.01	3.10 ± 0.78
		0.47 ± 0.01	3.40 ± 0.85
	Isopropanol	0.51 ± 0.02	4.41 ± 1.16
	$^a k_{\text{Cl}} = (8.7 \pm 2.3) \times 10^{-11}$	0.48 ± 0.02	4.21 ± 1.10
		0.49 ± 0.01	4.22 ± 1.10
	Ethyl formate	4.87 ± 0.36	4.97 ± 1.05
	$^a k_{\text{Cl}} = (1.0 \pm 0.2) \times 10^{-11}$	4.93 ± 0.18	5.03 ± 1.02
		4.51 ± 0.32	4.60 ± 0.98
Weighted Average			4.22 ± 0.27
	Reference compound	$k_{\text{carbonyl}}/k_{\text{R}} \pm 2\sigma$	$^d k_{\text{carbonyl}} \pm 2\sigma$
33DMbutanone+OH	Isopropanol	0.27 ± 0.01	1.39 ± 0.15
	$^a k_{\text{OH}} = (5.2 \pm 0.5) \times 10^{-12}$	0.22 ± 0.01	1.17 ± 0.12
		0.23 ± 0.02	1.18 ± 0.12
	2-methyl-2-butanol	0.49 ± 0.03	1.67 ± 0.51
	$^a k_{\text{OH}} = (3.4 \pm 1.0) \times 10^{-12}$	0.40 ± 0.02	1.38 ± 0.42
		0.56 ± 0.03	1.92 ± 0.59
	1-butanol	0.14 ± 0.01	1.27 ± 0.20
	$^a k_{\text{OH}} = (9.1 \pm 2.0) \times 10^{-12}$	0.16 ± 0.01	1.50 ± 0.24
		0.18 ± 0.01	1.68 ± 0.26
	Cyclohexane	0.18 ± 0.01	1.23 ± 0.13
	$^a k_{\text{OH}} = (6.7 \pm 0.7) \times 10^{-12}$	0.15 ± 0.01	1.00 ± 0.11
		0.25 ± 0.03	1.67 ± 0.26
		0.23 ± 0.02	1.52 ± 0.20
Weighted Average			1.26 ± 0.05

^aThe data of k_{R} (in units of $\text{cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$) are values recommended by McGillen et al., (2020). k_{carbonyl} (in units of $\text{cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$) is given in ^b 10^{-10} , ^c 10^{-11} , ^d 10^{-12} . The quoted errors in the $k_{\text{carbonyl}}/k_{\text{R}}$, (2σ), are twice the statistical errors from the regression analysis ($2 \times \sigma_{\text{slope}}$). The total absolute error $\sigma(k_{\text{carbonyl}})$ is a combination of the statistical errors from the regression analysis (σ_{slope}) and the quoted error in the value of the rate coefficient of the reference compound (σ_{R}). The final values of the rate coefficients and the associated error were calculated as weighted average (Colmenar et al., 2020).

Upon examining the rate coefficients and their associated uncertainties, it is evident that the majority of most of the experimental values fall within the expected error margins. Furthermore, the 50% difference observed between the rate coefficient values obtained for the reaction of 33DMbutanone with the OH radical is considered usual^{normal}, given the level of analytical complexity, particularly in the case of slow reactions. To the best of our knowledge, this is the first work where the rate coefficient for the reaction of 33DMbutanal with Cl atoms is determined. On the other hand, the values of the rate coefficients of the reaction of 33DMbutanone with Cl atoms and OH radicals have previously been determined with values of $(0.48 \pm 0.05) \times 10^{-10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ (Farrugia et al., 2015) for Cl reactions and $(1.21 \pm 0.05) \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ (Wallington and Kurylo, 1987) and $(1.2 \pm 0.2) \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ (Mapelli et al., 2023) for OH reaction. These data are in good agreement with the values obtained in this study, thereby contributing to the accurate determination of the rate coefficients.

In the case of 33DMbutanone reactions, the rate coefficient for Cl reactions ($4.22 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$) is one order of magnitude higher than the corresponding to OH reactions ($1.265 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$). This is the general trend observed in the atmospheric chemistry for the oxidation reactions of organic compounds; $k_{\text{Cl}} > k_{\text{OH}} > k_{\text{NO}_3}$ (Mellouki et al., 2015; Calvert et al., 2011; Atkinson, 2007). This behaviour can be explained by the higher reactivity and lower selectivity of Cl atoms compared to the OH radicals, where the site of attack determines its reactivity (Colmenar et al., 2020).

In Table 2, the rate coefficients for the reaction of different aldehydes and ketones in the butyl series with the main atmospheric oxidants have been tabulated to analyse the influence of ramifications on reactivity.

Table 2. Rate coefficients for aldehydes and ketones in the butyl series with key atmospheric oxidants.

Aldehydes			
Compounds	^a k_{Cl}	^b k_{OH}	^c k_{NO_3}
Butanal	$1.66 \pm 0.4^{\text{d}}$	$23.7 \pm 5^{\text{d}}$	$1.10 \pm 0.4^{\text{d}}$
2-methylbutanal	$2.16 \pm 0.32^{\text{e}}$	$33.3 \pm 13^{\text{d}}$	$2.67 \pm 0.8^{\text{d}}$
3-methylbutanal	$2.07 \pm 0.14^{\text{f}}$	$25.9 \pm 5^{\text{d}}$	$2.19 \pm 0.7^{\text{d}}$
3,3-dimethylbutanal	$1.27 \pm 0.08^{\text{g}}$	$21.4 \pm 9^{\text{d}}$	$1.77 \pm 0.4^{\text{d}}$
Ketones			
Compounds	^a k_{Cl}	^b k_{OH}	^c k_{NO_3}
2-butanone	$0.40 \pm 0.16^{\text{d}}$	$1.05 \pm 0.2^{\text{d}}$	
3-methylbutanone	$0.68 \pm 0.07^{\text{d}}$	$3.00 \pm 1.2^{\text{d}}$	$< 0.05^{\text{h}}$
3,3-dimethylbutanone	$0.48 \pm 0.05^{\text{d}}$	$1.21 \pm 0.5^{\text{d}}$	
	$0.42 \pm 0.03^{\text{g}}$	$1.26 \pm 0.1^{\text{g}}$	
		$1.2 \pm 0.2^{\text{i}}$	

k (in units of $\text{cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$), ^a 10^{-10} , ^b 10^{-12} , ^c 10^{-14} . ^dValues recommended in McGillen 2020. ^eAsensio et al., 2022.

^fBo et al., 2022. ^gThis work. ^hGlasius et al., 1997. ⁱMapelli et al., 2023

For butanals, the trend in rate coefficient values indicates that the presence of a methyl group influences to the reactivity, resulting in an increase in the rate coefficient compared to the compound without a methyl group. This could be attributed to the activation of the hydrogen atom at the α - position by a methyl group, as noted in the literature (Mellouki et al., 2015). Furthermore, the reactivity is influenced by both the position and quantity of methyl groups. Consequently, the activating influence exerted by the methyl group on the hydrogen bonded to the carbon adjacent to the aldehyde (α - position) is less pronounced when the methyl group occupies position 3 as opposed to position 2. Basically, the impact of the methyl group manifests as a short-range activating effect. Regarding the

227 impact of the number of methyl groups on reactivity, in the case of 33DMbutanal, the significant decrease in the
 228 value of rate coefficient with respect to 3-methylbutanal could be explained by an increase in steric hindrance,
 229 making the hydrogen abstraction process at the α - position less probable, thereby resulting in lower reactivity
 230 compared to the compound with one methyl group (3-methylbutanal). The decrease in the rate coefficient may also
 231 be related to the fact that many of the abstractable hydrogens are now primary.

232 Concerning butanones, no data are available for reactions with NO_3 radicals, with only one value for the upper limit
 233 of 3-methylbutan-2-one (Glasius et al., 1997). The fact that the reactions of ketones with NO_3 radicals are too slow
 234 complicates their experimental study and therefore, the determination of their rate coefficients. The available data
 235 of rate coefficients for reactions of butanones with Cl atoms and OH radicals, show again that the presence of a
 236 methyl group attached to a carbon in the α - position with respect to the carbonyl group activates the abstraction of
 237 one hydrogen atom from this carbon, resulting in an increase of the rate coefficient. A comparison of rate coefficients
 238 for 2-butanone and 3-methylbutan-2-one reveals this effect, particularly pronounced in $\cdot\text{OH}$ reactions. The presence
 239 of two methyl groups (33DMbutanone) produces a significant decrease in the value of the rate coefficient, which
 240 can be attributed to steric hindrance and a reduction in the number of secondary/tertiary abstractable hydrogens.

241 As can be observed in Table 2, the type of carbonyl group (aldehyde or ketone) also exerts a significant influence
 242 on the reactivity. The rate coefficients are generally one or two orders of magnitude higher for the aldehyde reactions
 243 compared to the reaction of ketones. The different reactivity of aldehydes and ketones with the main atmospheric
 244 oxidants has extensively been studied and documented in the literature (McGillen et al., 2020; Mellouki et al., 2015).
 245 The different reactivity observed in the reactions of atmospheric oxidant with saturated carbonyl compounds that
 246 are initiated by hydrogen abstraction, are due to the presence in the carbonyl compound of different types of
 247 hydrogens. In the aldehydic compounds there are two types of hydrogens that can be abstracted, the hydrogen
 248 directly attached to the carbonyl group (aldehydic hydrogen) and the hydrogen attached to the alkyl group (alkyl
 249 hydrogen), while in a ketone only alkyl hydrogens are present. The available kinetic and mechanistic data on the
 250 atmospheric degradation indicate that the H atom abstraction from the aldehydic group ($-\text{CHO}$) is more favoured
 251 than H atom abstraction from the C-H bonds of the alkyl chain. The rate coefficients obtained in this study for
 252 33DMbutanal and 33DMbutanone confirm this argument.

253 Additionally, it is well known that functional groups exert an activating or deactivating effect on reactivity,
 254 depending on the type of groups. The reactivity factors ($F(R)$; R =functional group) associated with the functional
 255 group to which a type of carbon is attached (primary (k_{prim}), secondary (k_{sec}) or tertiary (k_{tert})) can be quantified using
 256 the experimental kinetic database available in the literature. Consequently, rate coefficients for the reactions of
 257 33DMbutanal and 33DMbutanone with Cl atoms and OH and NO_3 radicals have been estimated with the SAR
 258 (Structure-Activity Relationship) predictive methods (Carter et al., 2021; Calvert et al., 2011; Jenkin et al., 2018;
 259 Kerdouci et al., 2014; Kwok and Atkinson, 1995). In the case of the two carbonyls compounds of this work, the only
 260 possibility of reacting is the abstraction of one hydrogen atom due to the absence of double bonds. The global
 261 abstraction rate coefficients can be calculated as $k_{\text{abs}} = 3(k_{\text{prim}}F(\text{C})) + k_{\text{sec}}F(\text{C})F(-\text{CHO}) + k_{-\text{COH}}F(\text{CH}_2)$ for
 262 33DMbutanal and $k_{\text{abs}} = 3(k_{\text{prim}}F(-\text{CR}_2\text{CO}-)) + (k_{\text{prim}}F(-\text{CO}-))$ for 33DMbutanone.

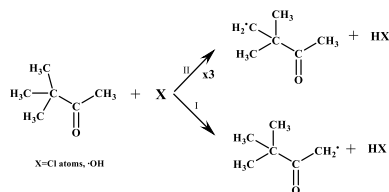
263 The rate coefficients (in $\text{cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$) and factors used to obtain the estimated rate coefficients for
 264 33DMbutanal and 33DMbutanone with Cl atoms are: $k_{\text{prim}}=2.84 \times 10^{-11}$, $k_{\text{sec}}=8.95 \times 10^{-11}$, $k_{-\text{COH}}=5.13 \times 10^{-11}$,

Con formato: Sin Superíndice / Subíndice

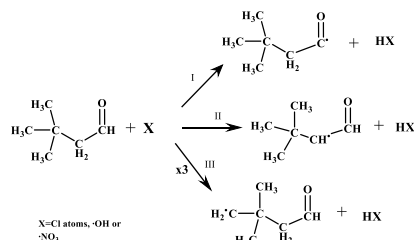
265 $F(C)=0.79$ proposed by Calvert et al., 2011, $F(-CHO)=0.4$ proposed by Carter et al., 2021, $F(-CR_2CO-)=0.563$ and
 266 $F(-CO)=0.037$ proposed by Farrugia et al., 2015. Therefore, the estimated rate coefficients for Cl reaction have been
 267 $k_{\text{estimated}} = 1.36 \times 10^{-10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ and $k_{\text{estimated}} = 0.49 \times 10^{-10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ for 33DMbutanal and
 268 33DMbutanone respectively. In the case of $\cdot\text{OH}$ reactions the rate coefficients have been estimated using the EPI
 269 (Estimation Programs Interface) Suite™, (US EPA), specifically the AOPWIN™. The rate coefficients estimated
 270 have been $k_{\text{estimated}}=2.21 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ for 33DMbutanal and $k_{\text{estimated}}=1.69 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ for
 271 33DMbutanone. For the reaction of NO_3 radicals only the estimated rate coefficient for 33DMbutanal have been
 272 obtained, $k_{\text{estimated}}=2.02 \times 10^{-14} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, using the data of Kerdouci et al., (2014). In all cases the estimated
 273 rate coefficients are very similar to the experimental values (see Table 1), indicating that the reactivity factors used
 274 for the estimations are well established. Reaction product studies and theoretical calculations of these reactions will
 275 help confirm the arguments presented above.

276 3.2 Products study and Mechanisms of reaction

277 The SAR method, explained above and used to estimate the rate coefficient, can also be used to define the branching
 278 ratios for hydrogen atom abstraction in the reaction of oxidant (Cl atoms, OH and NO_3 radicals) with a given
 279 saturated organic compound (Jenkin et al., 2018). In the Table 1S the percentages of hydrogen abstraction calculated
 280 for 33DMbutanone and 33DMbutanal are shown. For 33DMbutanone, the percentage of hydrogen abstraction from
 281 the $-\text{CH}_3$ group of the tert-butyl (channel II) is approximately 98% for Cl atoms and approximately 94% for OH
 282 radicals. These percentages are much higher than the percentages of hydrogen abstraction from the $-\text{CH}_3$ group in
 283 the α -position with respect to the carbonyl group (channel I) (~2% for Cl atoms and ~6 % for OH radicals). "x3"
 284 indicates three equivalent attack positions.



285
 286 For 33DMbutanal reactions, Aschmann et al. (2010) suggested that the primary pathway with OH radical involves
 287 hydrogen abstraction from the $-\text{CHO}$ group (channel I). However, the SAR predictions indicate that the main
 288 reaction channel varies with the oxidant type (see Table 1S). With Cl atoms, the major pathway is hydrogen
 289 abstraction from the $-\text{CH}_3$ group (~49%, channel III), followed by the $-\text{CHO}$ group (~30%, channel I), and the $-\text{CH}_2$ -
 290 group (~21%, channel II). For NO_3 radicals, the dominant pathway is hydrogen abstraction from the $-\text{CHO}$ group
 291 (~63%, channel I), with the $-\text{CH}_2$ - group (~37%, channel II) being the secondary channel. For OH radicals, the
 292 principal channel is hydrogen abstraction from the $-\text{CHO}$ group (~94%, channel I), followed by the $-\text{CH}_2$ - group
 293 (~4%, channel II), and $-\text{CH}_3$ group (~2%, channel III).



294

295 It is well established (Atkinson, 2007) that alkyl radicals, formed in the initial step of these reactions, rapidly react
 296 with O₂ to generate the corresponding peroxyradical (RO₂·). These RO₂ radicals can then follow various pathways.
 297 In the absence of NO, peroxyradicals primarily undergo two self-reaction processes: one leading to the formation of
 298 alkoxyradicals (RO₂· + RO₂· → 2RO· + O₂), and the other producing two molecules, such as hydroxy compound
 299 and carbonyl compound (RO₂· + RO₂· → hydroxy compound + carbonyl compound + O₂). Another significant
 300 process is the reaction of RO₂· with OH radicals and HO₂ radicals that undergoes different pathways (Bottorff et al.,
 301 2023; Berndt et al., 2022; Fittschen, 2019; Jenkin et al., 2019; Berndt et al., 2018; Winiberg et al., 2016).

302 In the presence of NO, the RO₂ radical may react to form alkoxyradicals and NO₂ (RO₂· + NO· → RO· + NO₂) or
 303 nitrated compounds (RONO₂), and in presence of large concentration of NO₂, RO₂· generates peroxy-nitrated
 304 compounds (ROONO₂) (pathway less favoured). Under typical tropospheric conditions, alkoxyradicals can react
 305 with molecular oxygen (O₂), undergo unimolecular decomposition or isomerize. The reaction of RO· radicals with
 306 O₂ is only possible if the carbon atom bearing the radical contains at least one hydrogen atom. Additionally, in the
 307 presence of NO and NO₂, alkoxyradicals can also form nitrated compounds (Atkinson, 2007).

308 The rate coefficients for unimolecular decomposition and isomerization processes have been estimated in this work,
 309 using the method by Vereecken and Peeters (2009, 2010). In the case of 33DMbutanone, the rate coefficients for
 310 isomerization were much lower than those for decomposition making isomerization products insignificant. The
 311 estimated decomposition rate coefficient to form acetone (1.7 × 10¹² s⁻¹) was much higher than to obtain butane-2,3-
 312 dione (6 × 10³ s⁻¹). For 33DMbutanal the decomposition rate is four times higher than that for the isomerization
 313 process.

314 Taking all the above into account, and to facilitate the qualitative analysis of reaction products from the reaction of
 315 33DMbutanone and 33DMbutanal with atmospheric oxidants, proposed reaction mechanisms are depicted in
 316 Schemes 1S and 2S in the supplementary material. The RO₂· + HO₂· reactions are excluded to avoid further
 317 complicating the mechanism for 33DMbutanal, as they are significant only in the absence of NO_x, that is, in remote
 318 unpolluted atmospheres.

319 In the FTIR experiments, the residual IR spectra of the reaction products (obtained after subtracting the spectra of
 320 33DMbutanone, 33DMbutanal, HCl, NO, NO₂, CH₃NO₂, N₂O₅, HNO₃, HNO₂, etc.) were compared with IR spectra
 321 of commercial samples or database spectra (Eurochamp 2020 database <https://data.eurochamp.org/data-access/spectra/> last access: 9 July 2024). In those cases where quantification was possible, concentration-time plots
 322 were made to obtain information on whether the products formed are primary or secondary. Additionally, plots of
 323 product concentrations versus carbonyl consumption were created to determine the yields of the products formed
 324 from the slopes of these plots.

325

SPME/GC-TOFMS with EI and/or FI ionization was used as a qualitative technique to complement FTIR in identifying reaction products. The chromatograms collected at different reaction times, show peaks at different retention times whose areas increase with the reaction time, indicating that they correspond to reaction products. A specific software tool of the mass spectrometer was used to generate chromatograms (from the original chromatograms) with a lower signal to noise ratio to improve peaks visualization and thereby facilitating the analysis of the experiments performed using this technique. For that, in the software it is specified the desired m/z, or m/z range, and then the chromatogram is created (displaying the ion intensity versus time) with peaks representing the compounds that correspond to the specified m/z (or m/z range). In Fig. 2S a comparison of an original chromatogram and a chromatogram generated with this tool are shown. This tool has been used for all SPME/GC-TOFMS chromatograms obtained in EI mode. Due to the characteristic of SPME sampling method, only a qualitative analysis was possible. Mass spectra were analysed using the NIST database or compared with commercial samples. In some cases, a high similarity index allowed positive identification, but most assignments were tentative based on fragmentation patterns and proposed reaction products in schemes 1S and 2S. When FI spectra were available, the assignment was based on the m/z of the molecular ion fragment.

Next a discussion of the results on reaction products with both analytical techniques for each of the studied compound is presented.

3.2.1 33DMbutanone

FTIR experiments

The identified and quantified reaction products were acetone ($\text{CH}_3\text{C}(\text{O})\text{CH}_3$) and formaldehyde (HCHO) for all reactions and nitrated compounds in those reactions carried out in the presence of NO and/or NO_2 . The nitrated compounds were attributed to alkoxy nitrates (RONO_2 , IR bands ~ 1663 , 1284 , 853 cm^{-1}) and peroxy nitrates (ROONO_2 , IR bands ~ 1720 , 1300 and 793 cm^{-1}) (Finlayson-Pitts and Pitts, 2000). A peroxy carbonyl nitrate such as PeroxyAcetyl Nitrate (PAN, $\text{CH}_3\text{C}(\text{O})\text{OONO}_2$, IR bands ~ 1830 , 1740 , 1300 and 793 cm^{-1}) was identified and quantified in the reactions of 33DMbutanone + Cl in the presence of NO after 3-5 minutes of reaction. Fig. 2 shows an example of residual spectra for the reactions of 33DMbutanone with Cl atoms in the absence and presence of NO. The figure includes reference spectra to confirm the formation of these compounds. Residual FTIR spectra for the reaction of 33DMbutanone with $\cdot\text{OH}$ in the absence and presence of NO, are presented in Fig. 3S.

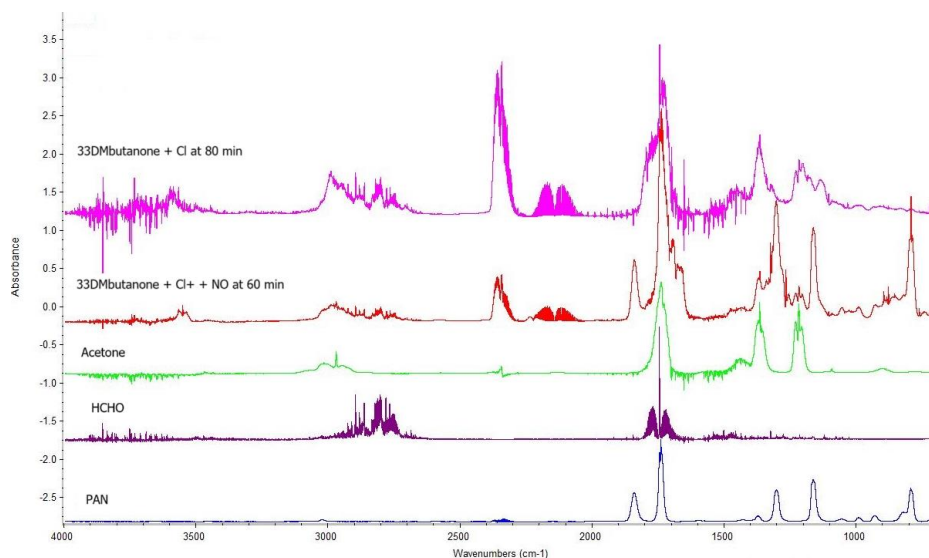
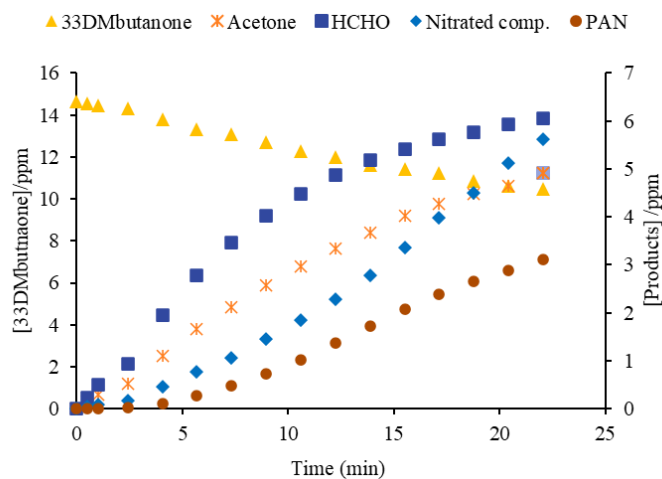


Figure 2. Residual FTIR spectra for the reactions of 33DMbutanone with Cl in the absence of NO (t=80 min) and in the presence of NO (t=60 min). Reference IR spectrum of HCHO, PAN (Eurochamp 2020 database <https://data.eurochamp.org/data-access/spectra/>) last access: 9 July 2024). Reference IR spectrum of acetone from a commercial sample. The spectra have been displaced for clarity.

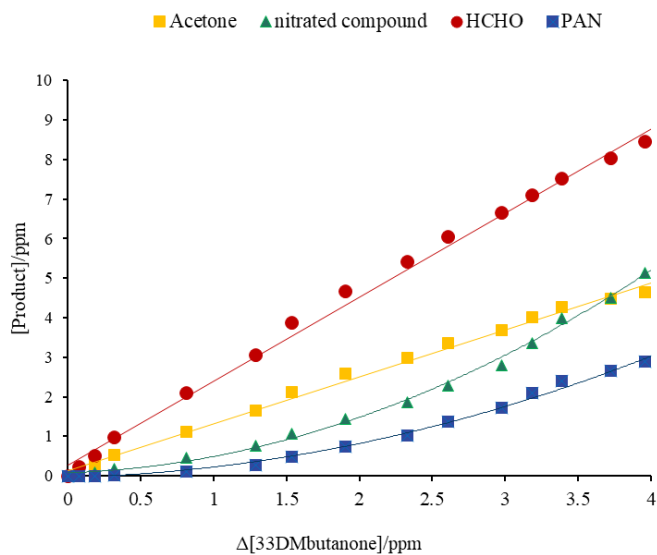
After removing the IR absorption bands of formaldehyde and acetone from the spectra in Fig. 2, the newly obtained residual spectra reveal the presence of IR absorption bands characteristic of various organic functional groups, such as carbonyl (-C(O)- $\sim 1745\text{-}1795\text{ cm}^{-1}$) and hydroxyl (-OH , $\sim 3600\text{ cm}^{-1}$) (see Fig. 4S). As it can be seen, all the spectra show IR bands around $3725\text{-}3500\text{ cm}^{-1}$, $3000\text{-}2750\text{ cm}^{-1}$ and 1780 cm^{-1} , indicating the formation of common reaction products in the reactions of 33DMbutanone with Cl and -OH . Other different IR bands (1136 , 1180 , 1364 cm^{-1}) are also identified, which would suggest different reaction products. Additionally, the IR bands shown in Fig. 4S are consistent with the multifunctional products proposed in Scheme 1S. Confirmation of these compounds is not possible due to their unavailability as commercial standards and the absence of reference spectra in existing infrared databases. Furthermore, the low intensity of their IR bands, likely resulting from low concentrations in the medium, combined with band overlap, hinders their identification.

To estimate the amount of nitrated compounds formed in the reactions with Cl atoms in the presence of NO, an average integrated absorption coefficient of $1.2 \times 10^{-17}\text{ cm}^2\text{ molecule}^{-1}$ was used for the IR range $1250\text{-}1330\text{ cm}^{-1}$ based on similar compounds (Tuazon and Atkinson, 1990). For PAN quantification, the reference spectrum (Eurochamp 2020 database, <https://data.eurochamp.org/data-access/spectra/>) last access: September 2024) has been used. Fig. 3 shows the concentration-time profiles of the products and 33DMbutanone for the reactions with Cl in the presence of NO.

Con formato: Superíndice



374
 375 Figure 3. Concentration-time profiles of the products and 33DMbutanone for the reaction with Cl atoms in the
 376 presence of NO.
 377 The trend of the acetone and HCHO profiles indicate that they are primary products, although the concentration of
 378 HCHO starts to decrease after 20 min of reaction, possibly due to secondary chemical reactions such as photolysis
 379 or by the reaction with the main oxidant. The profile of the nitrated compounds, especially PAN, shows a significant
 380 increase after 5 minutes of reaction, related to the rise in NO₂ concentration in the medium after that time.
 381 Fig. 4 shows an example of yield plots for the reactions of 33DMbutanone with Cl atoms in the presence of NO.



383 Figure 4. Plots of the reaction products formed versus the consumption 33DMbutanone in the reaction of Cl atoms
384 in the presence of NO.

385 The yields of nitrated compounds were estimated from the slopes of the plots showing linear behavior (using the
386 initial data) to avoid contributions from secondary chemistry. For PAN, the data used were from a $\Delta[33\text{DMbutanone}]$
387 of approximately 2 ppm (see Fig. 5S). In the case of HCHO, the yield has been recalculated using the formalism
388 published by Tuazon et al., 1986. The Fig.6S and Fig.7S of supplementary material show the concentration-time
389 profiles and yield plots for the reactions of 33DMbutanone with Cl and $\cdot\text{OH}$ in the absence of NO. The magnified
390 residual spectra (see Fig. 8S) show IR absorption bands that can be assigned to formic acid. Formic acid is mainly
391 produced through secondary reactions when formaldehyde is present, as evidenced by typical secondary
392 concentration time profile (see Fig. 9S).

393 In the reaction of 33DMbutanone with Cl atoms and OH radicals in the presence of NO, the acetone yield is uncertain
394 due to interference from the IR bands of the nitrated compounds. Additionally, the yields of HCHO and nitrated
395 compounds in the reaction with OH radical were not determined due to contributions from other sources such as the
396 precursor, methyl nitrite, and its degradation products (methyl nitrate and formaldehyde).

397 A summary of the estimated yields of reaction products, identified and quantified through FTIR analysis, is presented
398 in Table 3. The yield of total carbon in % is calculated with Eq. (II)

399
$$\text{Total carbon}(\%) = \sum_i \frac{n^{\circ} \text{ of carbon of product}_i}{n^{\circ} \text{ of carbon of carbonyl compound}} \times \text{molar yield}_i \quad (\text{II})$$

400

Table 3. Estimated yields (%) and total carbon of reaction products identified with FTIR analysis, and the reaction products tentatively assigned from SPME/GC-TOFMS analysis in the reactions of 33DMbutanone with Cl atoms and OH radicals in the absence and presence of NO.

FTIR				
Reaction	Yield acetone (%) ± 2σ	Yield HCHO ^{a,b} (%) ± 2σ	Yield nitrated comp (%) ± 2σ	Yield PAN (%) ± 2σ
33DMbutanone + Cl	65.0 ± 1.1	153.3 ± 7.0	-	-
	65.9 ± 1.8	178.1 ± 4.1	-	-
	69.0 ± 1.3	174.7 ± 3.6	-	-
	Average 66.6 ± 4.2	168.7 ± 27.0	-	-
Total carbon (%)			61	
33DMbutanone + Cl + NO	123.2 ± 3.2	212.4 ± 7.4	59.2 ± 1.9	101.0 ± 7.0
	111.2 ± 6.3	195.4 ± 5.8	56.6 ± 6.6	112.9 ± 1 8.4
	137.4 ± 11.0	199.9 ± 11.5	66.9 ± 6.1	95.5 ± 11.0
	Average 124.0 ± 26.2	202.6 ± 17.6	202.6 ± 17.6	103.1 ± 17.7
Total carbon ^c (%)			95.6	
33DMbutanone + ·OH ^d	31.4±2.0	59.5±2.4	-	-
	31.5±1.0	68.2±3.9	-	-
	34.1±1.6	73.5±4.2	-	-
	Average 32.3±3.0	67.0±14.0	-	-
Total carbon (%)			28	
33DMbutanone + ·OH + NO ^e	121.1 ± 6.7	-	-	-
	161.9 ± 19.7	-	-	-
	89.2 ± 6.4	-	-	-
	Average 93.0 ± 72.8	-	-	-
Total carbon (%)			47	
GC-TOFMS ^f				
Product (Retention time/min)	Cl	Cl + NO	OH + NO	
Acetone (2.2)	X	X	X	
Nitrated compound (2.6)			X	
Hydroxyacetone (2.9)	X			
Peroxyacetylnitrate, PAN (5.6)		X		
4-Hydroxybutan-2,3-dione (7)	X	X		
22DM3oxobutanal (8.3)	X	X		
Nitrated compound (9.1)		X	X	
Nitrated compound (9.7)		X		
4H33DMbutanone (10.3)	X			
Nitrated compound (14.6)		X	X	

^aYields have been estimated using the reference IR spectra from the Eurochamp database (Rodenas et al., 2017). ^bThe rate coefficient used to correct the concentration of formaldehyde has been $k_{Cl}=7.2 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ from IUPAC (2017).

^cNitrated compounds have not been accounted for total carbons. ^dOnly FTIR experiments using H₂O₂ as OH radical precursor.

^eExperiments using methyl nitrite as OH radical precursor. ^fThe positive identification and quantification were not possible due to the scarce of commercial samples. The quoted error in the individual yield (2σ) is two times the statistical errors from the regression analysis (2-×-σ_{slope}). The quoted error in the average yield (2σ) is two times the Standard deviation (2-×-σ).

22DM3oxobutanal (2,2-dimethylhydroxy-3-oxobutanal), 4H33DMbutanone (4-hydroxy-3,3-dimethyl-butan-2-one).

^aYields have been estimated using the reference IR spectra from the Eurochamp database (Rodenias et al., 2017). ^bThe rate coefficient used to correct the concentration of formaldehyde has been $k_{CI}=7.2 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ from IUPAC (2017). ^cNitrated compounds have not been accounted for total carbons. ^dOnly FTIR experiments using H_2O_2 as OH radical precursor. ^eExperiments using methyl nitrite as OH radical precursor. ^fThe positive identification and quantification were not possible due to the scarce of commercial samples. The quoted error in the individual yield (2σ) is two times the statistical errors from the regression analysis ($2-\times-\sigma_{\text{slope}}$). The quoted error in the average yield (2σ) is two times the Standard deviation ($2-\times-\sigma$). 22DM3oxobutanal (2,2-dimethylhydroxy-3-oxobutanal), 4H33DMbutanone (4-hydroxy-3,3-dimethylbutan-2-one).

Con formato: Fuente: Negrita

Con formato: Fuente: Negrita

Tabla con formato

Con formato: Fuente: Negrita

Con formato: Izquierda

Con formato: Izquierda

Con formato: Izquierda

Con formato: Izquierda

Con formato: Izquierda

Con formato: Izquierda

Con formato: Izquierda

Con formato: Izquierda

Con formato: Izquierda

Con formato: Izquierda

Con formato: Fuente: 9 pto

Con formato: Fuente: 9 pto

Table 3 shows that the total carbon recovery is less than 100%, however the residual FTIR spectra (Fig. 4S) indicate the formation of other reaction products not accounted for in the total carbon balance. In the reaction of 33DMbutanone with Cl atoms in the presence of NO the total carbon is 95% (without nitrated compounds yields). This high yield suggests possible overestimation of acetone and/or HCHO. In this regard, it should be noted that the calculated yields for acetone in reactions where nitrated compounds are generated have significant errors due to overlapping bands. On the other hand, as previously mentioned, the yields of the nitrated compound have been estimated using the average integrated absorption of similar compounds. Therefore, the yields of these nitrated compounds should also be interpreted with caution.

SPME/GC-TOFMS experiments

Fig. 5 shows an example of the SPME/GC-TOFMS chromatograms for the reactions of 33DMbutanone with Cl atoms and OH radicals. For the reaction with Cl atoms both EI and FI ionization modes were used. For the reaction with \cdot OH in the presence of NO, only the EI mode was used.

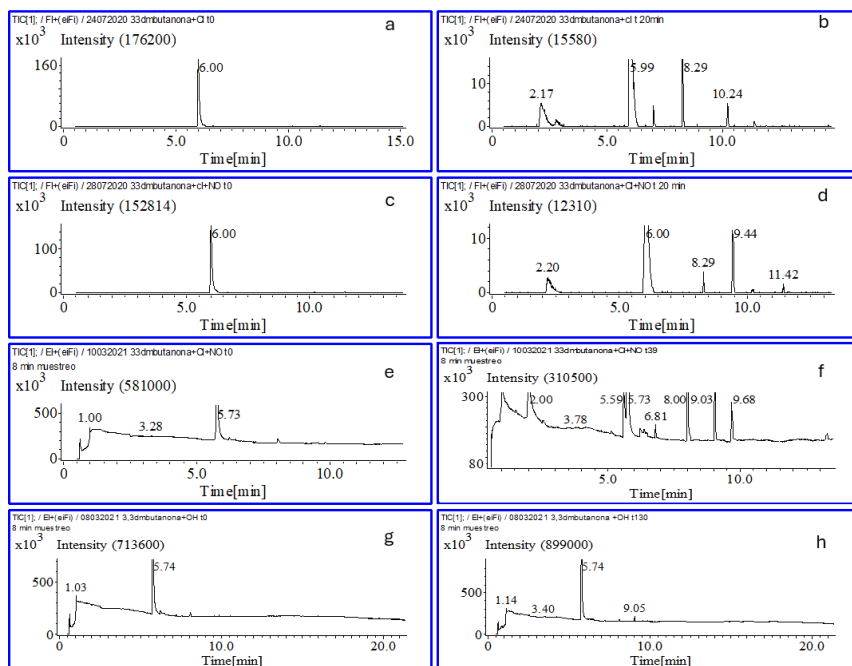


Figure 5. Example of the SPME/GC-TOFMS chromatograms for the reaction of 33DMbutanone + Cl atoms in the absence of NO and FI mode (reaction time t=0 min (a), t=20 min (b)) in the presence of NO and FI mode (reaction time t=0 min (c), t=20 min (d)) and in the presence of NO and EI mode (reaction time t=0 min (e), t=19 min (f)). Chromatograms (g) and (h) correspond to the reaction of 33DMbutanone + \cdot OH + NO at t=0 and t= 130 min of reaction respectively. **EI mode.**

The FI chromatogram in the absence of NO (b) shows different peaks compared to that in the presence of NO (d), with two common peaks (t_r ~2.20 and ~8.3 min). This indicates that the presence of NO influences the reaction mechanism. In different ionization modes ((d) and (f) chromatograms), also result in a different number of peaks

433 suggesting some products were not ionized with FI (for example, peak at t_r = 9.68 min). Additionally, the peak of
434 33DMbutanone in chromatograms (e-h), appears at shorter retention times than in chromatograms (a-d), due to the
435 use of a different chromatographic column. Taking into account this, the peaks at t_r =2.20, 6, 8.29 and 9.44 min,
436 showed in chromatograms (d) correspond to peaks at t_r =2.01, 5.73, 8.0 and 9.03 min in chromatograms (f).

437 The SPME/GC-TOFMS chromatograms for the reaction of 33DMbutanone with Cl and with \cdot OH in the presence
438 of NO (Fig. 10S) reveal both common peaks (t_r ~2 min, t_r ~ 8 min, t_r ~ 9.0 min and t_r ~14.6 min) and unique peaks
439 for each reaction (t_r ~2.5 min for \cdot OH, and t_r ~ 5.6 min, t_r ~ 6.8 min and t_r ~ 9.6 min for Cl) which is consistent with
440 the FTIR analyses. The mass spectra of all chromatographic peaks for the reaction products formed in the reactions
441 with Cl atoms and with OH radicals in the presence of NO are shown in Table 2S. Due to the low intensity of
442 chromatographic peak at t_r =8 min for the \cdot OH reaction, it was not possible to obtain its mass spectrum with enough
443 clarity.

444 The EI mass spectrum of each peak was analyzed using the NIST database of GC-MS. Except for acetone (98% of
445 similarity index), the similarity index for the assignment of the reaction products was below 15%. This low
446 similitude index may be due to low peak intensity (and thus the amount of product generated) or the absence of the
447 mass spectra of the reaction products in the NIST database, either because the compounds are non-commercial or
448 because they have not been previously reported in any bibliographic study. Without commercial samples, the
449 formation of proposed products could not be confirmed, so only tentative assignments were made. These
450 assignments were based on the m/z fragments of the mass spectra of these peaks, the expected m/z fragmentation
451 pattern for the reaction products from Scheme 1S and the IR absorption bands from the residual spectra from FTIR
452 experiments (Fig. 4S and Fig. 8S). In cases where the FI spectrum has been obtained, the assignment was also made
453 based on the m/z fragment of the molecular ion.

454 In the Cl reaction without NO, the identification of a product with a molecular ion of m/z =116 (at t_r =10.26 min, see
455 Table 2S) assigned to 4-hydroxy-3,3-dimethyl-2-butanone (4H33DM2butanone) is only explained by the self-
456 reaction of two RO_2 radicals, leading to the formation of two molecules. In this case, the co-product molecule to
457 4H33DM2butanone would correspond to a compound with a molecular ion of m/z =114 (at t_r =8.28 min), assigned
458 to 2,2-dimethyl-3-oxo-butanal (22DM3oxobutanal). This compound is also observed in the Cl reaction in the
459 presence of NO, formed by the reaction of the alkoxy radical with O_2 . The peaks at t_r =2.55 min with a m/z =72.08
460 and the peak at t_r =7.03 min with a molecular ion of m/z =102 (see Table 2S) were assigned to hydroxyacetone and
461 to hydroxybutan-2,3-dione respectively. The identification of these compounds indicates that the alkoxy radical also
462 undergoes isomerization processes (see Scheme 3S).

463 The peaks only observed in the reactions with NO (t_r =5.59, 9.04, 9.68, and 14.62 min for the Cl reaction, and t_r =2.55,
464 9.04, and 14.62 min for the OH reaction) must correspond to the nitrated compounds (peroxy and/or alkoxy nitrates)
465 proposed in scheme S1 and also observed in the residual spectra of the FTIR experiments (Fig. 2 and Fig. 3S). The
466 analysis of the mass spectra for the peaks at 2.55, 9.04, 9.68, and 14.62 min do not allow to identify clearly which
467 nitrated compound they correspond to Scheme 1S. The peak at 5.59 minutes is identified as peroxyacetyl nitrate
468 (PAN) which is unstable and decomposes in the injection port of the gas chromatograph. Upon decomposition, the
469 $CH_3-C(O)OO$ radical could fragment generating the ions $CH_3-C(O)^+$ (m/z = 43) and $CH_3-C(O)O^+$ (m/z = 59). These
470 fragments are like those observed in the EI mass spectrum (see Table 2S). There is not any reference mass spectrum

471 to compare, only two studies determined the chemical ionization mass spectrum of PAN (Phillips et al., 2013; Pate
472 et. al., 1976). Moreover, considering that PAN is clearly detected in the experiments conducted with the FTIR system
473 and that the mass spectrum corresponding to the chromatographic peak with a retention time of $t_r = 5.59$ min does
474 not account for the formation of any other compound proposed in Scheme 1S, it was concluded that this peak likely
475 corresponds to PAN.

476 The main reaction products tentatively assigned from SPME/GC-TOFMS analysis are shown in Table 3. The
477 SPME/GC-TOFMS experiments confirm the formation of acetone as identified and quantified in the FTIR
478 experiments. However, formaldehyde was undetectable due to the SPME sampling method. Other proposed products
479 include various multifunctional organic compounds (Hydroxybutan-2,3-dione, 22DM3oxobutanal,
480 4H33DM2butanone) and nitrated compounds with different carbon chain lengths.

481 Table 3 shows that the percentage of acetone and formaldehyde is higher in reactions with Cl and $\cdot\text{OH}$ in the presence
482 of NO compared to those without NO. This indicates that NO favors the formation of the alkoxy radical by reaction
483 of the peroxy radical that is chemically activated and undergo prompt decomposition to form acetone (Atkinson,
484 2007). In the absence of NO, the percentage of acetone is lower in the $\cdot\text{OH}$ reaction than in the Cl reaction, possibly
485 due to the formation of other products such as RO_2OH by the reaction of RO_2 with OH radical (Berndt et al., 2022;
486 Fittschen, 2019; Jenkin et al., 2019). This compound could not be identified due to the lack of a reference IR/MS
487 spectrum. The IR bands at $3700\text{--}3500\text{ cm}^{-1}$ in the residual spectrum of 33DMbutanone with $\cdot\text{OH}$ in the absence of
488 NO may be due to the OH stretching vibration in the ROO-OH molecule. The IR band at $\sim 1800\text{ cm}^{-1}$ for
489 33DMbutanone + Cl reaction (Fig. 2 and Fig. 4S) could correspond to the stretching vibration of the C=O (carbonyl
490 bond) in the acyl chloride. This compound can be formed by the reaction of RO_2 with Cl_2 or Cl atoms (Ren et al.
491 2018).

492 Considering that the formation of PAN can only be explained via channel II (hydrogen abstraction from the $-\text{CH}_3$
493 group of the tert-butyl group), which involves the decomposition of the initially formed alkoxy radical (2,2-
494 dimethyl-3-oxobutan-1-yloxy radical), and that the plot of PAN concentration versus the variation in 33DMbutanone
495 concentration shows a linear relationship with a slope close to one (indicating an estimated PAN yield of 100%, see
496 Fig. 5S), it could be concluded that the percentage of Cl attack on the $-\text{CH}_3$ group of the tert-butyl group is nearly
497 100%. This value closely with the 98% estimated by the SAR method for reactions with Cl (see Table 1S). For
498 reactions with $\cdot\text{OH}$, only acetone and HCHO were quantified but with the obtained yields do not confirm that 94%
499 of the reaction proceeds through channel II, as suggested by the SAR. However, considering that the rate coefficient
500 estimated by the SAR method is similar to the experimental one, channel II can be considered the main process.

501 Finally based on the kinetic results and the products observed in this study, the reaction mechanism for the
502 degradation of 33DMbutanone is shown in Fig. 6.

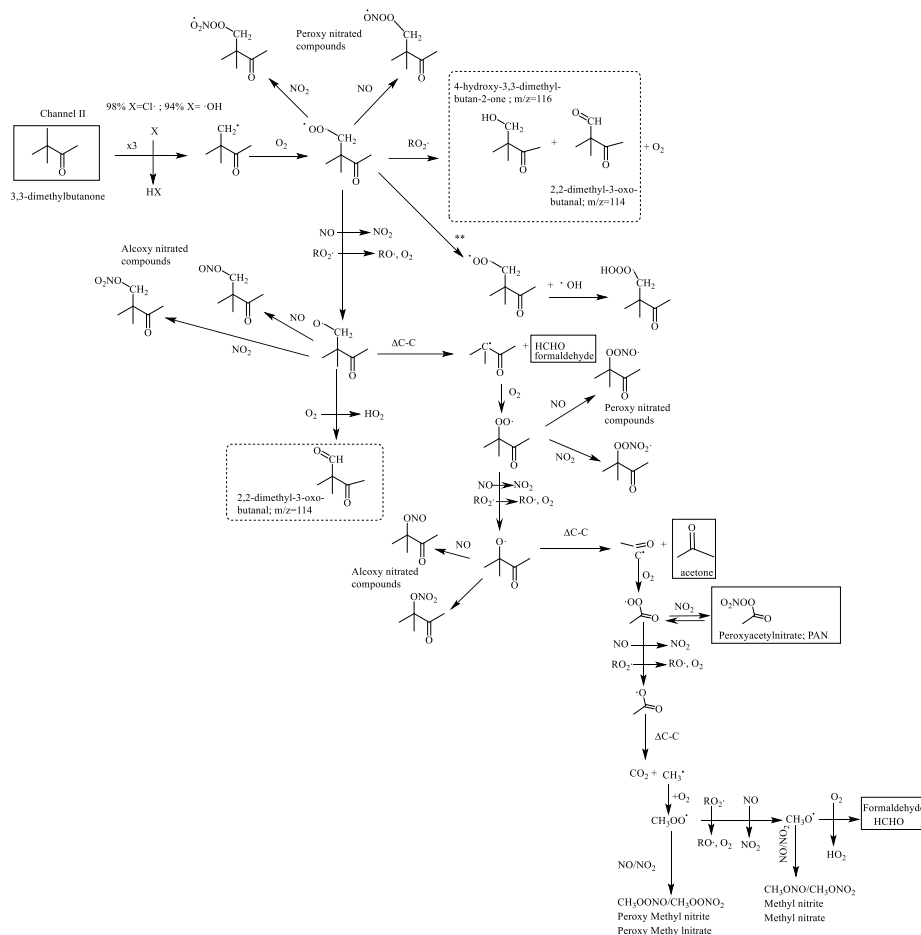


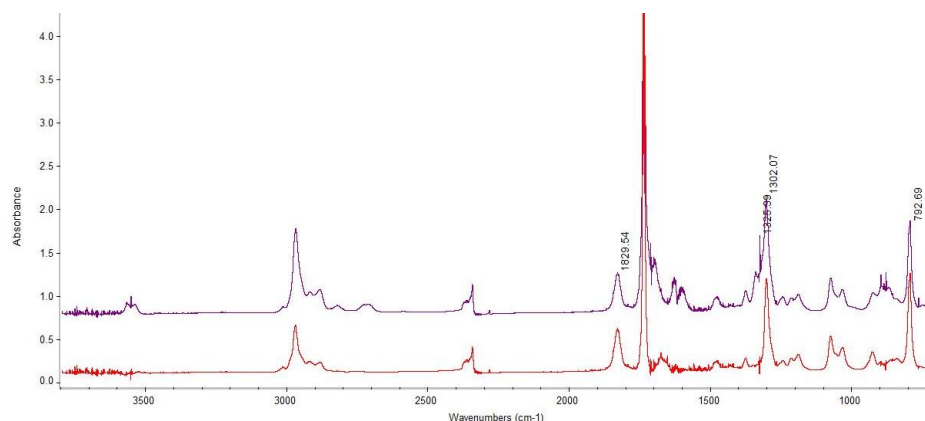
Figure 6. Mechanism proposed for the formation of the reaction products observed for 3,3-dimethylbutanone reactions with Cl atoms and OH radicals. The solid framed products correspond to products quantified by FTIR and dotted framed products correspond only to those products identified by SPME/GC-TOFMS. x3 indicates that there are 3 equivalent attack positions. ** Pathway proposed to reaction of 3,3-dimethylbutanone with OH in the absence of NO.

3.2.2 3,3-dimethylbutanal reaction

FTIR experiments

Fig. 11S shows an example of the residual IR spectra of the reaction of 3,3-dimethylbutanal with Cl atoms (in the absence and presence of NO), OH radicals and NO₃ radicals (obtained after subtracting the spectra of all known compounds). The reaction products identified and quantified from these residual spectra were: acetone, HCHO and 2,2-dimethylpropanal ((2,2-dimethylpropanal, (CH₃)₃CCHO)) for the reactions with Cl atoms, acetone for the reactions with OH radicals and nitrated compounds for the reaction with NO₃ radicals.

515 The IR bands of peroxy carbonyl nitrates observed in the residual spectra of NO₃ reactions (~1830, 1710, 1300 and
 516 790 cm⁻¹) could correspond to peroxy-3,3-dimethylbutyryl nitrate ((CH₃)₃CCH₂C(O)OONO₂) that is formed due to
 517 the large amount of NO₂ presents in the reaction mixture from the initial time. Fig. 7 shows the characteristic IR
 518 absorption bands of nitrated compounds formed in the reaction of 33DMbutanal with NO₃.



519
 520 Figure 7. FTIR spectrum of the reaction of 33DMbutanal (at 21 min) with NO₃ radical (upper). Residual FTIR
 521 spectrum (assigned to peroxy-3,3-dimethylbutyryl nitrate) after elimination of N₂O₅, HNO₃, 33DMbutanal and NO₂
 522 (lower).

523 In the reactions of 33DMbutanal with Cl atoms in the presence of NO and with the OH radicals at long reaction
 524 times, IR bands corresponding to a nitrated compound (similar to the one in the reaction with NO₃) have been
 525 detected. IR bands of N₂O₅ (a precursor of NO₃) have also been observed in the reaction with Cl atoms in the
 526 presence of NO as a consequence of the reaction between O₃ and NO₂ (see Fig. 12S). This indicates the formation
 527 of ozone in the degradation process of 33DMbutanal with Cl atoms in the presence of NO at long reaction times.
 528 The IR bands of the nitrated compounds show greater absorbance in the case of Cl reactions compared to the OH
 529 reactions, possibly due to an additional contribution from the 33DMbutanal reactions with NO₃. On the other hand,
 530 22DMpropanal has not been observed in the reactions of 33DMbutanal with OH and NO₃ radicals, likely due to the
 531 low concentration of this compound and the overlap of its characteristic IR bands with those of nitrated compounds.

532 After removing the IR absorption bands of the identified products in Fig. 11S, the residual spectra reveal the presence
 533 of IR absorption bands characteristic of carbonyl (~1700-1790 cm⁻¹), hydroxyl (~3600 and 1033 cm⁻¹) and organic
 534 acid/acyl chloride compound (~1800 cm⁻¹) (Fig. 12S). The amplified residual spectra of Fig. 12S (Fig.13S) shows
 535 clearly some IR absorption bands that appear at the same wavenumber indicating common reaction products. The
 536 IR band at 1105 cm⁻¹ shown in Fig. 13S again indicates the formation of formic acid.

537 The trends of the concentration-time profiles of the quantified products for the reaction of 33DMbutanal with Cl
 538 atoms (Fig. 14S), suggest that they are primary products in the early stages (linear trend). However, it is observed
 539 that after a certain reaction time, the concentrations of 22DMpropanal and HCHO decrease. This decrease could be
 540 attributed to loss processes involving reactions with Cl atoms and/or photolysis. Additionally, the concentrations of
 541 acetone and nitrated compounds are observed to increase more than expected (profile with an upward trend, Fig.
 542 14Sb), likely due to contributions from secondary chemistry, such as the previously discussed reactions, of

22DMpropanal with Cl or its photolysis. For the nitrated compounds, a change in the trend is observed around 2 minutes likely due to the formation of nitrated peroxy carbonyl compounds because of the presence of large amounts of NO₂ in the reaction mixture. The profile of nitrated compounds from the reaction of 33DMbutanal with NO₃ radical (Fig. 14Sd), shows an increase from the start of the reaction, attributed to the presence of NO₂ in the reaction medium from the beginning.

The concentration plots of the products formed against the variation 33DMbutanal, used to obtain the yields, are shown in Fig. 15S. For of HCHO and 22DMpropanal, where the concentration decreases by react with the main oxidant, the yield has been recalculated using the formalism by Tuazon et al. 1986. In the reaction of 33DMbutanal with the OH radical, the yield of HCHO and nitrated compounds has not been determined for the same reason as in the 33DMbutanone reactions. A summary of the estimated yields of reaction products, identified and quantified through FTIR analysis is presented in Table 4 for the reaction of 33DMbutanal with the three oxidants.

SPME/GC-TOFMS experiments

The SPME/GC-TOFMS chromatograms in EI mode show peaks at different retention times that increase with reaction time (see Fig. 16S), indicating that they correspond to reaction products. The number of significant peaks is higher in the reaction of 33DMbutanal with Cl atoms and NO₃ radicals compared to the reaction with OH radicals, where only 3 peaks are observed (see Fig. 17S-20S). This difference is likely due to the different attack positions (channels) of the oxidant: 3 for Cl atoms, 2 for NO₃ radicals, and 1 for OH radicals, as estimated by the SAR method (see Table 1S). Some of these peaks have similar retention times and mass spectra (see Table 3S), indicating that they are the same products. Some of these products are also identified in the FTIR analysis (see Table 4) such as acetone ($t_r = 2.2$ min) and 22DMpropanal ($t_r = 3.08/3.43$ min, confirmation with the injection of a real sample of 22DMpropanal).

It is interesting to note the presence of a common peak for all reactions that appears at 9.56 min (8.95 min in the experiment with a new chromatographic column). The mass spectrum of this peak is assigned to 3,3-dimethylbutanoic acid (33DMbutanoic acid) by the NIST database software, with a similarity index of approximately 87%. The formation of this compound cannot be explained by the general scheme proposed (Scheme 2S), which is based on principles of atmospheric reactivity and bibliographic studies of similar compounds (Aschmann et al., 2010; Atkinson, 2007). To explain the formation of this peak, other pathway was initially proposed, where the oxy-3,3-dimethylbutyryl radical (channel I), undergoes isomerization (see Scheme S4). This pathway leads to the formation of 4-oxo-3,3-dimethylbutanoic acid, whose mass spectrum shows a fragmentation pattern very similar to the peak with a t_r of 9.56 min. Taking into account that 33DMbutanoic acid is a commercial compound, a sample was injected into the SPME/GC-TOFMS system. The chromatogram showed a peak at approximately 9 minutes, with a mass spectrum (see Fig. 21S) identical to that of the peaks (9.56/8.95 min, see Table 3S) which positively confirms the formation of 33DMbutanoic acid in the reactions of 33DMbutanal with Cl atoms and \cdot OH and NO₃ radicals. The characteristic IR bands of 33DMbutanoic acid seem to be present in the residual FTIR spectra obtained for the reaction of 33DMbutanal with Cl atoms (Fig. 22S). Recent studies have also detected organic acids from reactions of saturated aldehydes (Asensio et al., 2022; Bo et al., 2022).

The remaining chromatographic peaks shown in Fig. 17S-20S, have been assigned to reaction products depicted in Scheme 2S (except the peaks at 16.2 and 21 min). Table 3S contains the mass spectra and their product assignments.

581 In general, the complete interpretation of mass spectra is complex due to the similar structures of the products
582 formed, leading to similar fragmentation patterns. The EI mass spectrum of each peak was analyzed using the NIST
583 database of GC-MS. The similitude index for most reaction products is low except for acetone (98%) and of peak at
584 5.47 min in the NO₃ reaction, assigned to 2,2-dimethylpropanol (22DMpropanol) with a similarity index of 73.5%.
585 The mass spectrum of peaks around 8 min (7.88, 8.23 and 8.26 min) could correspond to several structurally similar
586 products for example, 2,2-dimethyl-butanodial (22DMbutanodial), 3,3-dimethyl-oxobutanal (33DMObutanal) and
587 2,2-dimethylpropanoic acid (22DMpropanoic acid). A commercial sample of 22DMpropanoic acid injected into the
588 SPME/GC-TOFMS system showed a chromatogram with a peak at 8 minutes, matching the mass spectrum of the
589 peak at 8.23 min observed in the reaction of 33DMbutanal with Cl atoms with NO. Fig. 22S, shows an IR reference
590 spectrum of 22DMpropanoic acid. 22DMbutanodial and 33DMObutanal could not be positively confirmed due to
591 the lack of commercial samples.

592 As can be seen in Table 3S, some of the proposed reaction products are dicarbonyl and/or hydroxycarbonyl
593 compounds with IR bands also present in the residual spectra (Fig. 12S and Fig. 13S). Hydroxycarbonyl compounds
594 tend to cyclize to dihydrofurans via acid-catalyzed heterogeneous reactions (Atkinson et al., 2008). For example, 4-
595 hydroxy-3,3-dimethylbutanal (4H33DMbutanal, t_r = 15.78 min) can cyclize to form 2,3-dihydro-4,4-dimethylfuran
596 (23DH44DMfuran). Another cyclization process can occur from an alkoxy carbonyl radical, such as the 4-formyl-
597 2,2-dimethylbutan-1-yloxy radical, leading to the formation of 2,2-dimethyltetrahydrofuran-2-one
598 (22DMTHfuranone, t_r = 13.25 min). A similar cyclization could explain the formation of 3,4-dimethyldihydrofuran-
599 2,3-dione (34DMDHfuran23dione, t_r = 16.2 min) observed in the reaction with Cl atoms and assigned by the NIST
600 database with a similarity index of 62%. The peak at 6.97/6.86 min, observed in the Cl + NO and NO₃ radical
601 reactions, has been assigned to a nitrated compound, possibly peroxy-3,3-dimethylbutyryl nitrate also detected in
602 the FTIR analysis. The intensity of this peak is low, likely due to thermal decomposition in the chromatograph
603 injector. Another small peak at 7.15 min, observed in the NO₃ and OH radical reactions, could correspond to peroxy
604 nitrite (in the case of the OH reaction) and peroxy nitrate (for the NO₃ radical reaction), based on IR bands of peroxy
605 compounds at 793 cm⁻¹ observed in the FTIR spectra (see Fig. 13S). Fig. 13S also shows an IR band at 810 cm⁻¹ for
606 the Cl + NO and OH reactions, characteristic of alkoxy nitrated compounds. This common product was not detected
607 in SPME/GC-TOFMS, possibly due to adsorption onto the fiber. Generally, SPME/GC-TOFMS is not effective for
608 sampling for nitrated compounds.

609 The products assigned in the qualitative analysis of the SPME/GC-TOFMS experiments, along with the products
610 identified and quantified using FTIR are shown in Table 4. The total carbon yield in % is calculated with Eq. (II).

611

Table 4. Estimated yields and total carbon (%) for reaction products formed in the reactions of 33DMbutanal with the atmospheric oxidants using FTIR and the products identified in the qualitative analysis using GC-TOFMS.

FTIR				
Reaction	Yield 22DMpropanal ^{a,b} (%)±2σ	Yield Acetone (%)±2σ	Yield HCHO ^{b,c} (%)±2σ	Yield Nitrated comp. (%)±2σ
33DMbutanal + Cl	28.4±0.4	31.1±0.6	37.4±0.6	-
	29.6±0.3	26.8±0.4	38.5±0.7	-
	29.9±2.0	27.5±0.3	40.5±2.1	-
Average	29.3±1.4	28.5±4.8	38.8±3.1	-
Total carbon (%)	45			
33DMbutanal + Cl + NO	7.1±0.5	20.3±0.6	83.3±3.8	52.1±3.1
	7.1±0.2	20.8±0.3	89.2±2.3	50.3±3.2
	6.7±0.2	22.4±1.0	95.3±6.2	52.5±2.0
Average	7.0±0.4	21.1±2.1	89.3±12.0	51.6±2.3
Total carbon ^d (%)	31			
33DMbutanal + 2 OH + NO	Not observed	24.9±0.8	-	-
		34.6±1.0	-	-
		37.1±0.7	-	-
Average		32.2±13.0	-	-
Total carbon (%)	16			
33DMbutanal + NO ₃	Not observed	Not observed	Not observed	102±2.3
				105±1.0
				108±1.0
Average				107±1.0
				106±5.0
GC-TOFMS ^e				
Product (Retention time/min)	Cl	Cl + NO	OH + NO	NO ₃
Acetone (2.2)	X	X	X	X
Hydroxyacetone (2.7)	X			
22DMpropanal (3.4)	X	X		X
22DMpropanol (5.8)	X	X		X
Peroxy-3,3-dimethylbutyryl nitrate (7)		X		X
Peroxy nitrated compound (7.2)			X	X
22DMbutanodial/33DMObutanal/ 22DMpropanoic acid (8-8.2)	X	X		X
33DMbutanoic acid (9.6)	X	X	X	X
22DMTHfuran-2-one (13.3)	X	X		
4H33DMbutanal/ 2 3DH44DMfuran (15.8)	X			

^aThe rate coefficient used to correct the concentration of 22DMpropanal has been $k_{Cl}=1.42\times10^{-10}$ cm³ molecule⁻¹ s⁻¹ from Calvert et al., (2011). ^b Yields have been estimated using the reference IR spectra from the Eurochamp database (Rodenas et al., 2017). ^cThe rate coefficient used to correct the concentration of formaldehyde has been $k_{Cl}=7.2\times10^{-11}$ cm³ molecule⁻¹ s⁻¹ from IUPAC (2017). ^dNitrated compounds have not been accounted for. ^eThe positive identification and quantification were not possible to scarce of commercial compounds. Only 22DMpropanal, 33DMbutanoic acid and 33DMbutanoic acid were confirmed with a commercial sample. The quoted error in the individual yield (2σ) is two times the statistical errors from the regression analysis (2×σ_{slope}). The quoted error in the average yield (2σ) is two times the Standard deviation (2×σ).

The analysis of these quantified compounds provides insights into the percentage of each reaction channel or favored pathway. Acetone is a reaction product formed from all channels (see Scheme 2S). According to Aschmann et al.

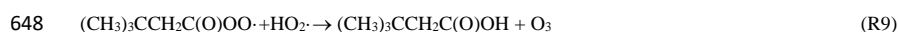
Con formato: Fuente: Negrita

Con formato: Fuente: Negrita

(2010) about reaction of 33DMbutanone with the OH radical the percentage of channel I is 94%, with acetone, tert-butyl nitrite and tert-butyl nitrate as the main reaction products. Therefore, considering the results of Aschmann et al. (2010), the acetone quantifies in our work for the reaction of 33DMbutanone with $\cdot\text{OH}$ (32%) would be formed mainly through channel I. In the reaction with Cl without NO, the acetone yield (28%) is close to the percentage predicted by the SAR method for channel I (30%). HCHO is also formed through all three channels. However, the highest yield of formaldehyde, along with the significant decrease of 22DMpropanal obtained in the reaction of Cl + NO compared to the yields of Cl reaction without NO, suggest that, in the presence of NO, the reaction of the peroxy species generated in channel I or channel II to form nitrated compounds is favored over the expense of the self-reaction of RO_2 .

The total carbon calculated for the Cl reaction in the absence of NO, based on the yields of HCHO, acetone, and 22DMpropanal, accounts for only 45%. The remaining carbon can be explained by the formation of other compounds identified through SPME/GC-TOFMS. Lower yields in the Cl reaction with NO (33%) and $\cdot\text{OH}$ (16%) could be attributed to the significant formation of nitrated compounds, not included in the total carbon calculation. For the NO_3 radical reaction, the linear trend observed in Fig. 15S (d) indicates that 100% of the reacted 33DMbutanal forms nitrated compounds. If only peroxy-3,3-dimethylbutyryl nitrate was generated, the total carbon would be 100%, indicating that no other compounds were formed in the reactions with NO_3 radical. However, as in the case of reactions with 33DMbutanone, the yield of nitrated compounds should be considered with caution. The identification of other compounds in the SPME/GC-TOFMS analysis could be due to the rapid decomposition of peroxy-nitrated compounds in the injection port or overestimation of nitrated compound yields. Quantification is necessary to determine if these products are generated in significant amounts, but this is not possible due to the lack of commercial compounds or the characteristics of the SPME sampling method.

No exclusive compounds for each reaction channel (I, II, or III) have been quantified for the 33DMbutanal reaction, making it impossible to determine the percentage of each channel. The formation of 33DMbutanoic acid in all reactions suggest another reaction pathway not considered in Scheme 2S, as $\text{RO}_2\cdot + \text{HO}_2\cdot$ -reactions are less likely in the presence of NO/ NO_2 . A proposed reaction pathway explains the formation of the 33DMbutanoic acid:



The IR bands of ozone are not observed in the residual spectra of Fig. 13S, likely due to the low amount of O_3 and overlapping IR bands. A similar reaction was proposed for the $\text{CH}_3\text{C}(\text{O})\text{OO}$ radical by other authors (Groß et al., 2014; Dillon and Crowley, 2008; Tomas et al., 2001). On the other hand, 22DMpropanoic acid (observed in the 33DMbutanal with Cl atoms) could be a secondary product from the degradation of 22DMpropanal. The yields of organic acids from the corresponding aldehyde must be very low (Bo et al., 2022), making pathway (R9) a minority reaction.

Based on the discussion about the reaction products done above, previous studies of 33DMbutanal with the OH radical (Aschmann et al., 2010) and estimated percentages for each channel using the SAR method, the scheme 2S initially proposed by the reaction mechanism for 33DMbutanal with Cl atoms, OH and NO_3 radicals has been simplified in Fig. 8.

$$\tau_{ox} = \frac{1}{k_{ox}[Ox]} \quad (III)$$

where τ_{ox} represents the lifetime for the considered reaction, k_{ox} is the rate coefficient, and $[Ox]$ refers to the typical atmospheric concentration of the oxidant. The atmospheric lifetimes of 33DMbutanone and 33DMbutanal were calculated using the rate coefficients and the concentration of oxidant from Table 2. For 33DMbutanal the k_{OH} and k_{NO_3} have been the averaged from the two rate coefficients available in the literature.

The ketones and aldehydes can significantly absorb light in the tropospheric actinic region $\lambda > 290$ nm, and the photolysis could play an important degradation process (Mellouki et al., 2015; Atkinson, 2003). There are not any experimental data about UV-Vis absorption cross-sections of 33DMbutanal that could allow the calculation of the photolysis rate. The value of $J_{(0, -\theta)} = 3 \times 10^{-5} \text{ s}^{-1}$ calculated for 3-methyl-butanal (similar compound to 33DMbutanal) by Lanza et al. (2008), has been used to estimate the photolysis lifetime. For 33DMbutanone a photolysis rate of $J(z, \theta) = 2.4 \times 10^{-6} \text{ s}^{-1}$ was estimated by Mapelli et al., (2023).

Regarding the deposition process, the lifetime associated with wet deposition could be estimated with the Eq. (IV) by Chen et al. 2003:

$$\tau_{wet} = \frac{H_{atm}}{k_H v_{pm} RT} \quad (IV)$$

where k_H is the Henry's law constant; H_{atm} is the height in the troposphere ($H_{atm} = 630$ m), v_{pm} is the average precipitation rate (536 mm yr^{-1} for Spain, (<http://www.aemet.es>, last access: 4 October 2024), R is the gas constant ($8.14 \text{ Pa m}^3/\text{mol K}$); and T is the temperature, considered to be constant and equal to 298 K . In the literature, there is only one data of Henry's constant for 33DMbutanone of $4.3 \times 10^{-2} \text{ mol/m}^3\text{Pa}$ (Sander, 2023, Hovorka et al., 2019) which has been used for 33DMbutanal as well.

Taking into account all those degradation processes, a global lifetime (τ_{global}) has also been calculated for 33DMbutanone and 33DMbutanal with the the Eq. (V):

$$\tau_{global} = \left[\frac{1}{\tau_{Cl}} + \frac{1}{\tau_{OH}} + \frac{1}{\tau_{NO_3}} + \frac{1}{\tau_{phot}} + \frac{1}{\tau_{wet}} \right]^{-1} \quad (V)$$

Table 5. Atmospheric lifetimes of 33DMbutanone and 33DMbutanal.

	^a τ_{Cl} (days)	τ_{Cl}^* (days)	^b τ_{OH} (days)	^c τ_{NO_3} (days)	τ_{phot} (days)	τ_{wet} (years)	τ_{global} (days)	[*] τ_{global} (days)
33DMbutanal	91	0.7	0.5	1.2	0.4	11	0.2	0.14
33DMbutanone	274	2.1	9.2	-	5	11	2.5	1.14

^aDetermined using the 24 h average $[Cl] = 1 \times 10^3 \text{ atoms cm}^{-3}$ (global average) (Platt and Jansen, 1995). ^{*}Determined using the peak of $[Cl]$ in coastal and industrial areas at $1.3 \times 10^5 \text{ atoms cm}^{-3}$ (Spicer et al., 1998). ^b Determined using the 12 h average of $[OH] = 1 \times 10^6 \text{ radicals cm}^{-3}$ (12-hour average) (Prinn et al., 2001). ^c $[NO_3] = 5 \times 10^8 \text{ radicals cm}^{-3}$ (Atkinson, 2000).

It can be observed that the dominant tropospheric loss processes of 33DMbutanone and 33DMbutanal are their reactions with OH radicals and photolysis process (note that photolysis lifetime depends on the atmospheric conditions considered), followed by their reaction with NO_3 radicals at night. However, in places where there is a

699 peak concentration of Cl atoms (coastal areas), the reaction with Cl atoms may compete with photolysis and reaction
700 with OH radicals as their main degradation process. The calculated wet lifetime of 11 years indicate that the wet
701 deposition can be considered negligible.

702 The shorter global lifetimes of ~4 hours for 33DMbutanal and ~2 days for 33DMbutanone indicate that these
703 compounds are degraded near their generation sources. The products created in the degradation reactions of
704 33DMbutanone and 33DMbutanal may also have environmental implications. Thus, formaldehyde is classified as
705 potentially carcinogenic to humans (NTP, 2021). Acetone, 22DMpropanal and organic nitrates (PAN and
706 peroxybutyrylnitrate) are also key components in photochemical smog episodes, a major contemporary
707 environmental issue. The multifunctional compounds such as oxocarboxyls, dicarboxyls, hydroxycarboxyls and
708 organic acids are products with polar groups characterized by low volatility, which could facilitate the formation of
709 secondary organic aerosols (SOA) (Asensio et al., 2022, Calvert et al., 2011). Moreover, the nitrated compounds
710 generated can act as NO_x reservoir species, especially during the night (Altshuler, 1993) and could have an
711 influence on global scale.

712 The potential for ozone formation of 33DMbutanone and 33DMbutanal has been evaluated calculating their
713 Photochemical Ozone Creation Potential (POCP) according to the method of Jenkin et al., (2017). The
714 Photochemical Ozone Creation Potential estimated (POCP_E), were 68 and 58 for conditions in NW Europe and
715 urban areas of the USA, respectively for 33DMbutanal and 26 and 15 for conditions in NW Europe and urban areas
716 of the USA, respectively for 33DMbutanone. Comparing with other series of organic compounds (Jenkin et al.,
717 2017), the values of POCP_E for 33DMbutanal indicate that it is an important contributor to tropospheric ozone
718 generation.

719 Regarding the calculation of the GWP (global warming potential) parameter, the method of Hodnebrog et al., (2020)
720 and the lifetime calculated above have been used to estimate the GWP of 33DMbutanone. A value of 0.13 for a time
721 horizon of 20 years has been obtained, and therefore, the direct contribution to the radiative forcing of climate can
722 be considered negligible. The GWP for 33DMbutanal has not been calculated because its lifetime is shorter than
723 that of 33DMbutanone, and thus, its expected GWP is likely to be lower.

724 5 Conclusions

725 In this work, the rate coefficient for the reaction of 33DMbutanal with Cl atoms has been determined for the first
726 time. Additionally, the rate coefficients for 33DMbutanone with Cl atoms and the OH radicals have been measured,
727 aligning with existing literature data. The kinetic findings, along with previous studies on other carbonyl compounds,
728 confirm that reactivity is influenced by the type of carbonyl group (aldehyde vs. ketone) and the number and position
729 of methyl groups. This research has expanded the database on these compounds, especially regarding their reactions
730 with Cl atoms.

731 The study of reaction products using FTIR and GC-TOFMS allows to identify and quantify acetone, formaldehyde,
732 and 22DMpropanal, alongside multifunctional products like hydroxycarboxyl, oxocarboxyl, and nitrated compounds
733 such as PAN and peroxybutyryl nitrate. The differing acetone yields from 33DMbutanone with Cl atoms and OH
734 radical suggest the importance of the RO₂· + ·OH reaction in unpolluted atmospheres. The proposed mechanism for
735 33DMbutanone indicates that hydrogen abstraction from the tert-butyl methyl group is the primary pathway for Cl
736 and ·OH, confirming SAR predictions. In the 33DMbutanal reaction, hydrogen abstraction occurs from various

functional groups depending on the reacting species (Cl atoms, OH and NO₃ radicals), also aligning with SAR predictions. The positive identification of 33DMbutanoic acid implies a pathway in the reaction mechanisms of 33DMbutanal, that initially have not been considered.

The atmospheric conditions determine the reaction products obtained in the atmospheric degradation of 33DMbutanal and 33DMbutanone. Thus, in polluted environments with high concentrations of NO_x, nitrated organic compounds (RONO₂) are formed. Moreover, when the concentration of NO₂ is higher than that of NO, ozone is formed. In a clean atmosphere, as in the case of the experiments with Cl atoms in the absence of NO_x, the reaction products are hydroxy/oxo carbonyl compounds.

Atmospherically, both 33DMbutanal and 33DMbutanone degrade within a few hours and 2 days respectively during the day, implying that degradation happens close to the emission sources. Their direct contribution to radiative forcing is minimal. However, their estimated POCP values suggest a potential role in tropospheric ozone formation, especially for 33DMbutanal. The multifunctional products formed may contribute to secondary organic aerosol formation, and their further oxidation in the troposphere could enhance photochemical smog, impacting air quality and human health.

Data availability. The underlying research data are available upon email request from the contact author of this work.

Supplement. The electronic Supplement includes additional tables and figures.

Author contributions.

Inmaculada Aranda: Formal analysis, validation, investigation, methodology, writing-original draft. **Pilar Martín:** Conceptualization, supervision, methodology, writing-original draft. **Sagrario Salgado** Conceptualization, supervision, methodology, writing-original draft. **Florentina Villanueva:** Supervision, methodology. **Beatriz Cabañas:** Conceptualization, supervision, funding acquisition.

Acknowledgments

The authors thank for financial support. I. Aranda thanks UCLM for funding her research contract (Plan Propio de I+D+i) cofinanced by FSE. The authors would like to thank Dr. Diana Rodriguez for her help in carrying out some of the experiments.

Financial support. This research has been supported by the Ministry of Science, Innovation and Universities (Project RTI 2018-099503-B-I00) and the Junta de Comunidades de Castilla-La Mancha (Project SBPLY/21/180501/000283).

References

-Altshuller, A. P.: PANs in the Atmosphere, Air & Waste, 43, 1221–1230, <https://doi.org/10.1080/1073161X.1993.10467199>, 1993.

-Aranda, I., Salgado, S., Martín, P., Villanueva, F., Martinez, E., Cabañas, B: Atmospheric degradation of 3-ethoxy-1-propanol by reactions with Cl, OH and NO₃, Chemosphere, 281, 130755-130764, <https://doi.org/10.1016/j.chemosphere.2021.130755>, 2021.

Código de campo cambiado

Código de campo cambiado

772 -Aschmann, S. M., Arey J., Atkinson, R: Kinetics and Products of the Reactions of OH Radicals with 4,4-Dimethyl-
 773 1-pentene and 3,3-Dimethylbutanal at 296 ± 2 K, J. Phys. Chem. A, 114, 18, 5810–5816,
 774 <https://doi.org/10.1021/jp101893g>, 2010.

775 -Asensio, M., Antiñolo, M., Blázquez, S., Albaladejo, J., and Jiménez, E.: Evaluation of the daytime tropospheric
 776 loss of 2-methylbutanal, Atmos. Chem. Phys., 22, 2689–2701, <https://doi.org/10.5194/acp-22-2689-2022>, 2022.

777 -Atkinson, R.: Rate constants for the atmospheric reactions of alkoxy radicals: An updated estimation method,
 778 Atmos. Environ., 41, Issue 38, 8468–8485, <https://doi.org/10.1016/j.atmosenv.2007.07.002>, 2007.

779 -Atkinson, R.: Atmospheric Degradation of Volatile Organic Compounds. Chem. Rev.103, 12, 4605–4638,
 780 <https://doi.org/10.1021/cr0206420>, 2003.

781 -Atkinson, R.: Atmospheric chemistry of VOCs and NO(x). Atmos. Environ. 34, 2063–2101,
 782 [https://doi.org/10.1016/S1352-2310\(99\)00460-4](https://doi.org/10.1016/S1352-2310(99)00460-4), 2000.

783 -Atkinson, R., Arey J., Aschmann S. M: Atmospheric chemistry of alkanes: Review and recent developments,
 784 Atmos. Environ. 42(23),5859–5871, <https://doi.org/10.1016/j.atmosenv.2007.08.040>, 2008.

785 -Bao, J., Li, H., Wu, Z., Zhang, X., Zhang, H., Li, Y., Qian, J., Chen, J., Deng, L: Atmospheric carbonyls in a heavy
 786 ozone pollution episode at a metropolis in Southwest China: Characteristics, health risk assessment, sources analysis,
 787 J. Environ. Sci (China), 113, 40–54, <https://doi.org/10.1016/j.jes.2021.05.029>, 2022.

788 -Berndt, T., Scholz, W., Mentler, B., Fischer, L., Hermann, H., Kulmala, M., Hansel, A.: Accretion Product
 789 Formation from Self- and Cross-Reactions of RO₂ Radicals in the Atmosphere, Angew. Chem. Int. Ed.
 790 26;57(14):3820–3824, <https://doi.org/10.1002/anie.201710989>, 2018.

791 -Bo, S., Weigang, W., Cici, F., Yuchan, Z., Zheng, S., Yanli, Z., Maofa, G.: Study on the reaction of 3-methyl-2-
 792 butenal and 3-methylbutanal with Cl atoms: kinetics and reaction mechanism, J. Environ. Sci., 116, 2022, 25–33,
 793 <https://doi.org/10.1016/j.jes.2021.03.032>, 2022.

794 -Bottorff, B., Lew, M. M., Woo, Y., Rickly, P., Rollings, M. D., Deming, B., Anderson, D. C., Wood, E., Alwe, H.
 795 D., Millet, D. B., Weinheimer, A., Tyndall, G., Ortega, J., Dusanter, S., Leonardis, T., Flynn, J., Erickson, M.,
 796 Alvarez, S., Rivera-Rios, J. C., Shutter, J. D., Keutsch, F., Helmig, D., Wang, W., Allen, H. M., Slade, J. H., Shepson,
 797 P. B., Bertman, S., and Stevens, P. S.: OH, HO₂, and RO₂ radical chemistry in a rural forest environment:
 798 measurements, model comparisons, and evidence of a missing radical sink, Atmos. Chem. Phys., 23, 10287–10311,
 799 <https://doi.org/10.5194/acp-23-10287-2023>, 2023.

800 -Byrne, F. P., Forier, B., Bossaert, G., Hoebbers, C., Farmer, T. J., and Hunt, A. J.: A methodical selection process
 801 for the development of ketones and esters as bio-based replacements for traditional hydrocarbon solvents, Green
 802 Chem., 20, 4003–4011, <https://doi.org/10.1039/C8GC01132J>, 2018.

803 -Calvert, J. G., Mellouki, A., Orlando, J. J., Pilling, M. J., Wallington, T. J.: The Mechanisms of Atmospheric
 804 Oxidation of the Oxygenates. Oxford University Press, New York, 2011.

805 -Carter, W. P. L.: Estimation of Rate Constants for Reactions of Organic Compounds under Atmospheric Conditions,
 806 Atmosphere, 12, 1250, <https://doi.org/10.3390/atmos12101250>, 2021.

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

807 -Chen, L., Takenaka, N., Bandow, H., Maeda, Y.: Henry's law constants for C2–C3 fluorinated alcohols and their
 808 wet deposition in the atmosphere, *Atmos. Environ.*, 37, 34, 4817–4822,
 809 <https://doi.org/10.1016/j.atmosenv.2003.08.002>, 2003.

810 -Colmenar, I., Martín, P., Cabanas, B., Salgado, S., Tapia, A., Aranda, I.: Atmospheric fate of a series of saturated
 811 alcohols: kinetic and mechanistic study, *Atmos. Chem. Phys.* 20, 699–720. [https://doi.org/10.5194/acp-20-699-](https://doi.org/10.5194/acp-20-699-2020)
 812 [2020](https://doi.org/10.5194/acp-20-699-2020), 2020.

813 -Colmenar, I., Martin, P., Cabanas, B., Salgado, S., Martinez, E.: Analysis of reaction products formed in the gas
 814 phase reaction of E,E-2,4-hexadienal with atmospheric oxidants: reaction mechanisms and atmospheric
 815 implications, *Atmos. Environ.* 176, 188–200. <https://doi.org/10.1016/j.atmosenv.2017.12.027>, 2018.

816 -D'Anna, B., Andresen, W., Gefen, Z., and Nielsen, C. J.: Kinetic study of OH and NO₃ radical reactions with 14
 817 aliphatic aldehydes, *Phys. Chem. Chem. Phys.*, 3, 3057–3063, doi.org/10.1039/B103623H, 2001.

818 -D'Anna, B. and Nielsen C.J.: Kinetic study of the vapour-phase reaction between aliphatic aldehydes and the nitrate
 819 radical, *J. Chem. Soc., Faraday Trans*, 93(19), 3479–3483, <https://doi.org/10.1039/A702719B>, 1997.

820 -Dillon, T. J. and Crowley, J. N.: Direct detection of OH formation in the reactions of HO₂ with CH₃C(O)O₂ and
 821 other substituted peroxy radicals, *Atmos. Chem. Phys.*, 8, 4877–4889, <https://doi.org/10.5194/acp-8-4877-2008>,
 822 2008.

823 -EUROCHAMP, 2020 <https://data.eurochamp.org/data-access/ir-spectra/#/>

824 -Farrugia, L. N., Bejan, I., Smith, S. C., Medeiros, D. J., and Seakins, P. W.: Revised structure activity parameters
 825 derived from new rate coefficient determinations for the reactions of chlorine atoms with a series of seven ketones
 826 at 290 K and 1 atm, *Chem. Phys. Lett.*, 640 87–93, <https://doi.org/10.1016/j.cplett.2015.09.055>, 2015.

827 -Finlayson-Pitts. B. J., Pitts, Jr. J. N.: *Chemistry of the Upper and Lower Atmosphere: Theory, Experiments, and*
 828 *Applications*. Academic Press, 2000.

829 -Fittschen, C.: The reaction of peroxy radicals with OH radicals, *Chem. Phys. Lett.*, 725, 102–108,
 830 <https://doi.org/10.1016/j.cplett.2019.04.002>, 2019.

831 -Groß, C. B. M., Dillon, T. J., Schuster, G., Lelieveld, J., and Crowley, J. N.: Direct kinetic study of OH and O₃
 832 formation in the reaction of CH₃C(O)O₂ with HO₂, *The Journal of Physical Chemistry A*, 118, 974–985,
 833 <https://doi.org/10.1021/jp412380z>, 2014.

834 -Glasius, M., Calogirou, A., Jensen, N., Hjorth, J., Nielsen, C.: Kinetic Study of Gas Phase Reactions of
 835 Pinonaldehyde and Structurally Related Compounds, *Int. J. Chem. Kinet.* 7; 527–533,
 836 [https://doi.org/10.1002/\(SICI\)1097-4601\(1997\)29:7<527::AID-KIN7>3.0.CO;2-W](https://doi.org/10.1002/(SICI)1097-4601(1997)29:7<527::AID-KIN7>3.0.CO;2-W), 1997.

837 -Heald, C. L.; Kroll, J. H. *The Fuel of Atmospheric Chemistry: Toward a Complete Description of Reactive Organic*
 838 *Carbon*, *Sci. Adv.*, 6, 8967, <https://doi.org/10.1126/sciadv.aay8967>, 2020.

839 -Hodnebrog, Ø., Aamaas, B., Fuglestad, J. S., Marston, G., Myhre, G., Nielsen, C. J., et al.: Updated global warming

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Con formato: Francés (Francia)

Con formato: Francés (Francia)

Con formato: Francés (Francia)

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

840 potentials and radiative efficiencies of halocarbons and other weak atmospheric absorbers, *Reviews of Geophysics*,
841 58, e2019RG000691, <https://doi.org/10.1029/2019RG000691>, 2020.

842 -Hovorka, Š., Vrbka, P., Bermúdez-Salguero, C., Böhme, A., & Dohnal, V.: Air–water partitioning of C5 and C6
843 alkanones: measurement, critical compilation, correlation, and recommended data, *J. Chem. Eng. Data*, 64, 5765–
844 5774, <https://doi.org/10.1021/ACS.JCED.9B00726>, 2019.

845 -IUPAC. Task Group on Atmospheric Chemical Kinetic. <http://iupac.pole-ether.fr>, 2007.

846 -Jenkin, M. E., Valorso, R., Aumont, B., and Rickard, A. R.: Estimation of rate coefficients and branching ratios for
847 reactions of organic peroxy radicals for use in automated mechanism construction, *Atmos. Chem. Phys.*, 19, 7691–
848 7717, <https://doi.org/10.5194/acp-19-7691-2019>, 2019.

849 -Jenkin, M. E., Valorso, R., Aumont, B., Rickard, A. R., and Wallington, T. J.: Estimation of rate coefficients and
850 branching ratios for gas-phase reactions of OH with aliphatic organic compounds for use in automated mechanism
851 construction, *Atmos. Chem. Phys.*, 18, 9297–9328, <https://doi.org/10.5194/acp-18-9297-2018>, 2018.

852 -Jenkin, M. E., Derwent, R. G., Wallington, T. J.: Photochemical ozone creation potentials for volatile organic
853 compounds: Rationalization and estimation, *Atmos. Environ.* 163, 128–137.
854 <https://doi.org/10.1016/j.atmosenv.2017.05.024>, 2017.

855 -Jenkin, M. E., Saunders, S. M., and Pilling, M. J.: The tropospheric degradation of volatile organic compounds: a
856 protocol for mechanism development, *Atmos. Environ.*, 31, 81–104, [https://doi.org/10.1016/S1352-2310\(96\)00105-](https://doi.org/10.1016/S1352-2310(96)00105-7)
857 [7](https://doi.org/10.1016/S1352-2310(96)00105-7), 1997.

858 -Kerdouci, J., Picquet-Varrault, B., Doussin, J. F.: Structure–activity relationship for the gas-phase reactions of NO3
859 radical with organic compounds: Update and extension to aldehydes, *Atmos. Environ.* 2014, 84, 363–372,
860 <https://doi.org/10.1016/j.atmosenv.2013.11.024>, 2014.

861 -Kwok, E.S.C., Atkinson, R.: Estimation of hydroxyl radical reaction rate constants for gas-phase organic
862 compounds using a structure-reactivity relationship: An update, *Atmos. Environ.* 29, 1685–1695,
863 [https://doi.org/10.1016/1352-2310\(95\)00069-B](https://doi.org/10.1016/1352-2310(95)00069-B), 1995.

864 -Lanza, B., Jiménez, E., Ballesteros, B., Albaladejo, J.: Absorption cross section determination of biogenic C5-
865 aldehydes in the actinic region, *Chem. Phys. Lett.*, 454, 184–189, <https://doi.org/10.1016/j.cplett.2008.02.020>, 2008.

866 -Liu, Q., Gao, Y., Huang, W., Ling, Z., Wang, Z., Wang, X.: Carbonyl compounds in the atmosphere: A review of
867 abundance, source and their contributions to O3 and SOA formation, *Atmos. Res.*, 274, 106184,
868 <https://doi.org/10.1016/j.atmosres.2022.106184>, 2022.

869 -Mapelli, C., Donnelly, J. K., Hogan, Ú. E., Rickard, A. R., Robinson, A. T., Byrne, F., McElroy, C. R., Curchod,
870 B. F. E., Hollas, D., and Dillon, T. J.: Atmospheric oxidation of new 'green' solvents part II: methyl pivalate and
871 pinacolone, *Atmos. Chem. Phys.*, 23, 7767–7779, <https://doi.org/10.5194/acp-23-7767-2023>, 2023.

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

872 -McGillen, M. R., Carter, W. P. L., Mellouki, A., Orlando, J. J., Picquet-Varrault, B. and Wallington T. J.: Database
873 for the kinetics of the gas-phase atmospheric reactions of organic compounds, *Earth Syst. Sci. Data*, 12, 1203–1216,
874 <https://doi.org/10.5194/essd-12-1203-2020>, 2020.

Código de campo cambiado

875 -Mellouki, A., Wallington, T. J., and Chen, J.: Atmospheric chemistry of oxygenated volatile organic compounds:
876 impacts on air quality and climate, *Chem. Rev.*, 115, 3984–4014, <https://doi.org/10.1021/cr500549n>, 2015.

877 -NTP (National Toxicology Program). Report on Carcinogens, Fifteenth Edition. Research Triangle Park, NC: U.S.
878 Department of Health and Human Services, Public Health Service, doi.org/10.22427/NTP-OTHER-1003, 2021.

Código de campo cambiado

879 -Pate C. T., Sprung J. L. and Pitts J. N.: Chemical ionization mass spectrum of peroxyacetylnitrate, *J. Mass*
880 *Spectrom.*, 11, 552–555, <https://api.semanticscholar.org/CorpusID:95293138>, 1976.

Código de campo cambiado

881 -Phillips, G. J., Pouvesle, N., Thieser, J., Schuster, G., Axinte, R., Fischer, H., Williams, J., Lelieveld, J., and
882 Crowley, J. N.: Peroxyacetyl nitrate (PAN) and peroxyacetic acid (PAA) measurements by iodide chemical
883 ionisation mass spectrometry: first analysis of results in the boreal forest and implications for the measurement of
884 PAN fluxes, *Atmos. Chem. Phys.*, 13, 1129–1139, <https://doi.org/10.5194/acp-13-1129-2013>, 2013.

885 -Platt, U., Jansen, C.: Observation and role of the free radicals NO₃, ClO, BrO and IO in the troposphere, *Faraday*
886 *Discuss* 100, 175–198, <https://doi.org/10.1039/FD9950000175>, 1995.

Código de campo cambiado

887 -Prinn, R.G., Huang, J., Weiss, R.F., Cunnold, D.M., Fraser, P.J., Simmonds, P.G., McCulloch, A., Harth, C.,
888 Salameh, P., O'Doherty, S., Wang, R.H.J., Porter, L., Miller, B.R.: Evidence for Substantial Variations of
889 Atmospheric Hydroxyl Radicals in the Past Two Decades, *Science*. 292, 1882–1888.
890 doi.org/10.1126/science.1058673, 2001.

Código de campo cambiado

891 -Ren, Y., Wang, J., Grosselin, B., Daele, V., and Mellouki, A.: Kinetic and product studies of Cl atoms reactions
892 with a series of branched ketones, *J. Environ. Sci.*, 71, 271–282, <https://doi.org/10.1016/j.jes.2018.03.036>, 2018.

Código de campo cambiado

893 -Ródenas, M. IR spectrum: FORMALDEHYDE HCHO (Version 1.0) [Data set]. AERIS. [https://doi.org](https://doi.org/10.25326/K17C-0762)
894 [/10.25326/K17C-0762](https://doi.org/10.25326/K17C-0762), 2017.

Código de campo cambiado

895 -Sander, R.: Compilation of Henry's law constants (version 5.0.0) for water as solvent, *Atmos. Chem. Phys.*, 23,
896 10901–12440, <https://doi.org/10.5194/acp-23-10901-2023>, 2023.

Código de campo cambiado

897 -Saunders, S. M., Jenkin, M. E., Derwent, R. G., and Pilling, M. J.: Protocol for the development of the Master
898 Chemical Mechanism, MCM v3 (Part A): tropospheric degradation of non-aromatic volatile organic compounds,
899 *Atmos. Chem. Phys.*, 3, 161–180, <https://doi.org/10.5194/acp-3-161-2003>, 2003

Código de campo cambiado

900 -Schott, G., Davidson, N.: Shock Waves in Chemical Kinetics: The Decomposition of N₂O₅ at High Temperatures,
901 *J. Am. Chem. Soc.* 80, 1841–1853. <https://doi.org/10.1021/ja01541a019>, 1958.

902 -Spicer, C.W., Chapman, E.G., Finlayson-Pitts, B.J., Plastringe, R.A., Hubbe, J.M., Fast, J.D., Berkowitz, C.M.:
903 Unexpected high concentrations of molecular chlorine in coastal air, *Nature* 394, 353–356,
904 <https://doi.org/10.1038/28584>, 1998.

Código de campo cambiado

905 -Tadic, J. M., Moortgat, G. K., Bera, P. P., Loewenstein, M., Yates, E. L., and Lee, T. J.: Photochemistry and
 906 photophysics of n-butanal, 3-methylbutanal, and 3,3 dimethylbutanal: Experimental and theoretical study, J. Phys.
 907 Chem. A, 116, 5830–5839, <https://doi.org/10.1021/jp208665v>, 2012.

908 -Tanielyan, S.K. and Augustine, R. L.: Synthesis of 3,3-dimethylbutanol and 3,3-dimethylbutanal, important
 909 intermediates in the synthesis of Neotame, Top Catal., 55, 625-630, <https://doi.org/10.1007/s11244-012-9841-z>,
 910 2012.

911 -Taylor, W.D., Allston, T.D., Moscato, M.J., Fazekas, G.B., Kozlowski, R., Takacs, G.A.: Atmospheric
 912 photodissociation lifetimes for Nitromethane, Methyl Nitrite and Methyl Nitrate, Int. J. Chem. Kinet. 12, 4, 231–
 913 240, <https://doi.org/10.1002/kin.550120404>, 1980.

914 -Tomas, A., Villenave, E., and Lesclaux, R.: Reactions of the HO₂ radical with CH₃CHO and CH₃C(O)O₂ in the
 915 gas phase, J. Phys. Chem. A, 105, 3505–3514, <https://doi.org/10.1021/jp003762p>, 2001.

916 -Tuazon, E.C., Atkinson, R.: A product study of the gas-phase reaction of Isoprene with the OH radical in the
 917 presence of Nox, Int. J. Chem. Kinet. 22, 1221–1236, <https://doi.org/10.1002/kin.550221202>, 1990.

918 -Tuazon, E. C., Leod, H. M., Atkinson, R., and Carter, W. P. L.: α -Dicarbonyl Yields from the NO_x Air
 919 Photooxidations of a Series of Aromatic Hydrocarbons in Air, Environ. Sci. Technol., 20, 383–387,
 920 <https://doi.org/10.1021/es00146a010>, 1986.

921 -US EPA. [2024]. Estimation Programs Interface Suite™ for Microsoft® Windows, v 4.11 or insert version used].
 922 United States Environmental Protection Agency, Washington, DC, USA.

923 -Vereecken, L., and Peeters, J.: A structure–activity relationship for the rate coefficient of H-migration in substituted
 924 alkoxy radicals, Phys. Chem. Chem. Phys., 12, 12608–12620, <https://doi.org/10.1039/C0CP00387E>, 2010.

925 -Vereecken, L., and Peeters, J.: Decomposition of substituted alkoxy radicals—part I: a generalized structure–
 926 activity relationship for reaction barrier heights, Phys. Chem. Chem. Phys., 11, 9062–9074,
 927 <https://doi.org/10.1039/B909712K>, 2009.

928 -Vila, J. A., Argüello, G. A., Malanca, F. E.: Kinetics studies of 5-methyl-2-hexanol, 2,2-dimethyl-3-hexanol, and
 929 2,4,4-trimethyl-1-pentanol with chlorine atoms: Photooxidation mechanism of 2,4,4-trimethyl-1-pentanol, Int. J.
 930 Chem. Kinet., 52(1), 29-34, <https://doi.org/10.1002/kin.21327>, 2020.

931 -Wallington, T. J. and Kurylo, M. J.: Flash Photolysis Resonance Fluorescence Investigation of the Gas-Phase
 932 Reactions of OH Radicals with a Series of Aliphatic Ketones over the Temperature Range 240–440 K, J. Phys.
 933 Chem., 91, 5050–5054, <https://doi.org/10.1021/j100303a033>, 1987.

934 -Winiberg, F. A. F., Dillon, T. J., Orr, S. C., Groß, C. B. M., Bejan, I., Brumby, C. A., Evans, M. J., Smith, S. C.,
 935 Heard, D. E., and Seakins, P. W.: Direct measurements of OH and other product yields from the HO₂ + CH₃C(O)O₂
 936 reaction, Atmos. Chem. Phys., 16, 4023–4042, <https://doi.org/10.5194/acp-16-4023-2016>, 2016.

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

Código de campo cambiado

937 -Xiong, K., Zheng, X., Jiang, M., Gao, D., Wang, F., and Chen Y.: Phase Equilibrium on Extraction Methylphenols
938 from Aqueous Solution with 3,3-Dimethyl-2-butanone at 333.2 K and 353.2 K, J. Chem. Eng. Data, 63, 7, 2376–
939 2383, <https://doi.org/10.1021/acs.jced.7b00932>, 2018.

940 -Zhou, X., Zhou, X., Wang, C., Zhou, H.: Environmental and human health impacts of volatile organic compounds:
941 A perspective review, Chemosphere, 313, <https://doi.org/10.1016/j.chemosphere.2022.137489>, 2023.

Código de campo cambiado