Acid-yield measurements of the gas-phase ozonolysis of ethene as a function of humidity using Chemical Ionisation Mass Spectrometry (CIMS)

K. E. Leather\textsuperscript{1}, M. R. McGillen\textsuperscript{1,*}, M. C. Cooke\textsuperscript{2}, S. R. Utembe\textsuperscript{1,2}, A. T. Archibald\textsuperscript{3}, M. E. Jenkin\textsuperscript{2,4}, R. G. Derwent\textsuperscript{5}, D. E. Shallcross\textsuperscript{2}, and C. J. Percival\textsuperscript{1}

\textsuperscript{1}The Centre for Atmospheric Science, The School of Earth, Atmospheric and Environmental Science, The University of Manchester, Simon Building, Brunswick Street, Manchester, M13 9PL, UK
\textsuperscript{2}Biogeochemistry Research Centre, School of Chemistry, The University of Bristol, Cantock’s Close BS8 1TS, UK
\textsuperscript{3}Centre for Atmospheric Science, Department of Chemistry, University of Cambridge, Lensfield Road, Cambridge, CB2 1EW, UK
\textsuperscript{4}Atmospheric Chemistry Services, Okehampton, Devon, EX20 1FB, UK
\textsuperscript{5}rdscientific, Newbury, Berkshire, UK
\textsuperscript{*}current address: Chemical Sciences Division, Earth System Research Laboratory, National Oceanic and Atmospheric Administration (NOAA), 325 Broadway, Boulder, CO 80305, USA

Correspondence to: C. J. Percival (c.percival@manchester.ac.uk)

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Abstract. Gas-phase ethene ozonolysis experiments were conducted at room temperature to determine formic acid yields as a function of relative humidity (RH) using the integrated EXTreme RAnge chamber-Chemical Ionisation Mass Spectrometry technique, employing a CH\textsubscript{3}I ionisation scheme. RHs studied were <1, 11, 21, 27, 30 % and formic acid yields of (0.07 ± 0.01) and (0.41 ± 0.07) were determined at <1 % RH and 30 % RH respectively, showing a strong water dependence. It has been possible to estimate the ratio of the rate coefficient for the reaction of the Criegee biradical, CH\textsubscript{2}OO with water compared with decomposition. This analysis suggests that the rate of reaction with water ranges between 1\times 10^{-12} – 1\times 10^{-15} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} and will therefore dominate its loss with respect to bimolecular processes in the atmosphere. Global model integrations suggest that this reaction between CH\textsubscript{2}OO and water may dominate the production of HC(O)OH in the atmosphere.

1 Introduction

Organic acids are ubiquitous in the gas and aerosol phase, and are common constituents of global precipitation (Keene and Galloway, 1983). Organic acids have been measured in urban, rural, marine and remote areas (Talbot et al., 1988; Chebbi and Carlier, 1996; Kawamura et al., 2001). The contribution of organic acids to the acidity of precipitation and subsequent effects on aquatic and terrestrial ecosystems has been documented by Keene and Galloway (1986). Formic and acetic acid can dominate free acidity of precipitation thereby having an influence on pH-dependent chemical reactions and even OH cloud chemistry (Jacob et al., 1986). Low molecular weight organic salts – presumably the product of organic acid dissolution – are present in the fine fraction of aerosols, whose physical properties, namely hygroscopicity, possess relatively low critical supersaturations, allowing the activation of cloud droplets and subsequently affecting total indirect radiative forcing (Yu, 2000).

Sources of carboxylic acids include biogenic and anthropogenic primary emissions, biomass burning and
hydrocarbon oxidation, though their relative fluxes are poorly constrained (Chebbi and Carlier, 1996; Paulot et al., 2011). The major sinks of carboxylic acids are dry and wet deposition as a result of their low reactivity towards OH and NO3. However, the chemical loss via reaction with OH is poorly constrained resulting from the uncertainty in the reported rate coefficient (Atkinson et al., 2006). The modelled atmospheric lifetime of formic acid has been calculated to be 3.2 days (Paulot et al., 2011).

Global models under predict formic acid concentrations (von Kuhlmann et al., 2003; Rinsland et al., 2004; Paulot et al., 2011) especially in the marine boundary layer where [HC(O)OH] can be underestimated by a factor of 10–50, this discrepancy has been attributed to missing sources such as higher biogenic emissions during the growing season (Rinsland et al., 2004) and ageing of organic aerosols (Paulot et al., 2011). Also, the oxidation of VOC precursors leading to the production of formic acid has been suggested to be a significant source (Arlander et al., 1990), for instance the ozonolysis of ethene. Ethene emissions have been estimated to be about 15 Tg yr−1 (EDGAR, 1996) with about 162 Gmol yr−1 from the oceans (Paulot et al., 2011), and the presence of a major formic acid-producing reaction channel would therefore be of major importance to atmospheric chemical modelling.

This study focuses on the production of formic acid from ethene ozonolysis. Intuitively, monitoring the products of this reaction ought to be easier than many ozonolysis reactions since the first-generation products possess a carbon number of one and are likely to be of maximum volatility. However, there still remain considerable inconsistencies in formic acid yields reported in the literature (Neeb et al., 1997; Wolff et al., 1997; Orzechowska and Paulson, 2005). Ozonolysis proceeds via a 1,3-cycloaddition across the olefinic bond to produce a primary ozonide, the decomposition of which forms a carbonyl moiety and a Criegee biradical each with unit yield (Scheme 1).

It is the fate of the Criegee biradical that determines the end product yield and this has provoked much attention in the atmospheric chemistry community (Johnson and Marston, 2008 and references therein). Here the mechanisms highlighted shall focus on acid production pathways. It was first suggested by O’Neal and Blumstein (1973) that the Criegee biradical may isomerise to form a dioxirane intermediate, leading to the formation of carboxylic acids, as detailed by Orzechowska and Paulson (2005), this hypothesis is supported by the theoretical calculations of Cremer et al. (1998) (Scheme 2).

Formic acid may also be produced from bimolecular reactions. Calvert et al. (1978) suggested that in the presence of water, acid production can be significantly enhanced via reaction of the stabilised Criegee radical with water (Scheme 3). The formation of HC(O)OH via Scheme 3 has been further supported by the theoretical results of Hatakeyama et al. (1981), Crehuet et al. (2001) and Anglada et al. (2002). Minor pathways such as cross reactions of reactive intermediates can form secondary ozonides, for instance reaction between Criegee biradicals and carbonyls, which have been suggested to lead to the formation of acids (Neeb et al., 1996).

Despite the importance of these formic acid-producing channels, there have been relatively few experimental determinations of HC(O)OH yields from the ozonolysis of ethene. Wolff et al. (1997) and Orzechowska and Paulson (2005) report high formic acid yields (0.36) for ethene ozonolysis conducted in dry conditions compared with that of Neeb et al. (1997) (0.01) (see Table 1). Discrepancies in yields reported under humid conditions also exist as Neeb and co-workers obtain 0.42 at 65 % RH yet Orzechowska and Paulson report 0.33 at 65 % RH. Wolff et al. (1997) and Orzechowska and Paulson (2005) both use indirect analytical techniques to quantify acid yields, whereas Neeb et al. (1997) use FTIR. All the techniques have the potential for significant errors as a result of sampling efficiency, spectral overlap and low sensitivity. This study aims to resolve the discrepancy by quantifying acid yields using the highly selective and sensitive technique, Chemical Ionisation Mass Spectrometry (CIMS).

2 Experimental

Experiments were conducted in the dark in the 123 L Teflon®-coated EXTRA chamber, described in detail elsewhere (Leather et al., 2010, 2011 and McGillen et al., 2011), and shown in Fig. 1. FVMQ o-ring seals within the end flanges ensured that the chamber was leak tight, whilst reinforcement ribs afford maximum pressures of 3750 Torr. Seven sample ports are incorporated into the end flanges, to enable simultaneous sampling by a range of analytical techniques. The EXTRA chamber can be operated over
Figures

Scheme 1

![Diagram](image)

A schematic diagram of the formation of HC(O)OH under wet conditions.

Scheme 3

![Diagram](image)

A schematic diagram of the formation of HC(O)OH under wet conditions.

**Table 1.** Formic acid yields previously reported, to the best of our knowledge.

<table>
<thead>
<tr>
<th>RH %</th>
<th>HC(O)OH yield</th>
</tr>
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<tbody>
<tr>
<td>0</td>
<td>0.36±0.05 Wolff et al. (1997)</td>
</tr>
<tr>
<td>0.36±0.07 Orzechowska and Paulson (2005)</td>
<td></td>
</tr>
<tr>
<td>0.01</td>
<td>Neeb et al. (1997)</td>
</tr>
<tr>
<td>20</td>
<td>0.4±0.13 Wolff et al. (1997)</td>
</tr>
<tr>
<td>65</td>
<td>0.42 Neeb et al. (1997)</td>
</tr>
<tr>
<td>65</td>
<td>0.33±0.06 Orzechowska and Paulson (2005)</td>
</tr>
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</table>

HC(O)OH was detected using $\Gamma^-$ as the reagent ion. $\Gamma^-$ was generated by combining a 1.5 STP1 min$^{-1}$ flow of N$_2$ and a 1 scm flow of 0.5% CH$_3$I/H$_2$O/N$_2$ gas mixture and passing it through a Po(210) Nuclecel ionizer (NRD Inc.). HC(O)OH was ionised by $\Gamma^-$ via an adduct reaction,

$$\Gamma^- \cdot \text{H}_2\text{O}_n + \text{HC(O)OH} \rightarrow \text{HC(O)OH} \cdot \Gamma^- \cdot \text{H}_2\text{O}_n$$

which enabled formic acid to be detected selectively at $m/z = 171.65$ (Slusher et al., 2004).

Ions were detected with a quadrupole mass spectrometer in a three-stage differentially pumped vacuum chamber, as shown in Fig. 2. A sample of the ion molecule gas flow containing reactant ions is drawn into the collision dissociation chamber through a 0.38 mm aperture which was held at a potential of $-0.17$ V to focus charged reactant molecules. The collision dissociation chamber was pumped by a molecular drag pump (Alcatel MDP-5011) backed by a scroll pump (ULVAC DISL-100) and held at approximately 20 Torr. The ions were further focused by an octopole ion guide, stainless steel with a 1.00 mm aperture held at $-0.36$ V and passed into a second chamber containing the further octopole ion guide and passed into the rear chamber via a stainless steel plate with a 1.00 mm aperture held at $-0.48$ V which contained the quadrupole mass filter (ABB Extrel, Merlin). This second and rear chamber were each pumped by a turbo-molecular pump (Varian 81-M) backed by the molecular drag pump (Alcatel MDP-5011). Under typical operating conditions the rear chamber was held at a pressure of approximately $9 \times 10^{-6}$ Torr. Ions were detected using a channeltron (Dtech 402A-H) via negative ion counting.

Gaseous reagents were added to the chamber at a known flow rate and duration using calibrated 1179 MKS mass flow controllers. The chamber contained an atmosphere of nitrogen preceding the addition of reagent gases. Ethene was introduced from a dilute ethene/nitrogen gas mixture. Ozone was produced by flowing oxygen through a UVP ozone generator (97-0067-02).

For the addition of water or formic acid, known volumes were injected into a Pyrex impinger, becoming volatilised by exposure to the evacuated chamber, assisted by an additional nitrogen carrier gas purge. The initial reactant concentrations were varied as follows; $[\text{O}_3] = 2.46 \times 10^{12} - 9.84 \times 10^{13}$ molecule cm$^{-3}$, $\text{H}_2\text{O}\leq 1-30\%$ RH and $\text{C}_2\text{H}_4 = 4.92 \times 10^{13} - 2.23 \times 10^{15}$ molecule cm$^{-3}$.

3 Materials

Ethene (Sigma Aldrich, 99.54 %) was purified by repeated freeze-pump-thaw cycles. N$_2$, O$_2$ (99.6 %) were used as supplied. Formic acid (Fisher Scientific UK, 98/100 %) was used without further purification. Purified water ($\geq 15.0$ MΩ cm) was obtained from a PURELAB Option-S 7/15 (ELGA).
3.1 Global model description

The Global Chemistry Transport model CRI-STOCHEM has been used to assess the mass of products formed in the atmosphere using data from this study. CRI-STOCHEM is described in detail in (Utembe et al., 2010 and Archibald et al., 2010). The model used is an updated version of the UK Meteorological Office tropospheric chemistry transport model (STOCHEM) described by Collins et al. (1997), with updates reported in detail in the recent paper of Utembe et al. (2010). STOCHEM is a global 3-dimensional CTM which uses a Lagrangian approach to advect 50,000 air parcels using a 4th-order Runge-Kutta scheme with advection time steps of 3 hours. The transport and radiation models are driven by archived meteorological data, generated by the Met office numerical weather prediction models as analysis fields with a resolution of 1.25° longitude and 0.83° latitude and on 12 vertical levels extending to 100 hPa. Full details of the model version employed are given in Derwent et al. (2008).

The common representative intermediates mechanism (CRIv2-R5) (Jenkin et al., 2008; Watson et al., 2008; Utembe et al., 2009), which represents the chemistry of methane and 22 emitted non-methane hydrocarbons was employed in the model. Each parcel contains the concentrations of 219 species involved in 618 photolytic, gas-phase and heterogeneous chemical reactions, with a 5 min time step. The formation of secondary organic aerosol (SOA) is represented using 14 species, which are derived from the oxidation of aromatic hydrocarbons, monoterpenes, and isoprene (see Utembe et al., 2011).

The surface emissions (man-made, biomass burning, vegetation, oceans, soil and “other” surface emissions) are
distributed using two-dimensional source maps. Emissions
totals for the base case run for CO, NO$_x$ and non-methane hy-
drocarbons are taken from the Precursor of Ozone and their
Effects in the Troposphere (POET) inventory (Granier et al.,
2005) for the year 1998. The emission of aromatic species
ortho-xylene, benzene and toluene were taken from Henze et
al. (2008). Biomass burning emission of ethyne, formalde-
hyde and acetic acid are produced using scaling factors from
Andreae and Merlet (2001) per mole of CO emitted. NASA
inventories are used for aircraft NO$_x$ emissions for 1992
taken from Penner et al. (1999). The lightning and aircraft
NO$_x$ emissions are monthly averages and are 3-dimensional
in distribution.

4 Results and discussion

4.1 Assessment of instrument sensitivity.

Dilute mixtures of HC(O)OH in deionized water were in-
jected into the Chamber with no other gases present and the
HC(O)OH$^{-}$ signal was monitored. From a linear plot of
[HC(O)OH] vs. HC(O)OH$^{-}$ signal it is estimated that the
sensitivity for HC(O)OH was $2.39 \times 10^4$ molecule cm$^{-3}$
for a signal to noise ratio of one and a time constant of 1 s.

4.2 Rate coefficient determination

The gas-phase rate coefficient of the reaction of ethene with
ozone was determined using the absolute method. The rate
equation is shown in Eq. (2);

$$-d\ln[O_3]/dt = k'[\text{ethene}]$$

where $k'$ is the pseudo-first order rate coefficient given by
$k' = k[O_3]$. For each experiment, the slope $k'$ was obtained
using the linear regression of $\ln[O_3]$ vs. time for a broad
range of alkene concentrations. First-order plots exhibited
linear decays, (Fig. 3) having typical $R^2$ of 0.99, indicat-
ing first-order kinetic behaviour. The plot of $k'$ vs. ini-
tial [ethene] also exhibited a strong linear relationship, from
which the gradient $k$, the bimolecular rate coefficient for
the reaction was determined (Fig. 4). The rate was found to
be $(1.62 \pm 0.14) \times 10^{-18}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, in excellent
agreement with the literature recommendation (Atkinson et
al., 2000).

4.3 Product yields

Product yields were determined in excess ethene conditions,
in excess typically by a factor of 300–400. Initial compar-
ison of HC(O)OH signal shows that [HC(O)OH] at RH 30 %
exceeds that of RH < 1 % by more than a factor of 7, in the
initial stages during ethene ozonolysis, which suggests that
reaction Scheme 3 dominates in the presence of water. Fig-
ure 5 shows two temporal profiles of the formic acid pro-
duced. The curve passing through the [HC(O)OH] exper-
imentally determined values utilises the literature retrieved
rate coefficient of $1.58 \times 10^{-18}$ cm$^3$ molecule$^{-1}$ s$^{-1}$
(Atkinson et al., 2000) and the line of best fit is obtained by vary-
ng the branching ratio to HC(O)OH formation. [HC(O)OH]
yields were quantified as a function of relative humidity (RH),
as summarised in Table 2.

In Fig. 6 it is clear to see that the formic acid yield in-
creases from RH < 1–30 %, in keeping with the work of Neeb
et al. (1997) but in disagreement with the work of Orze-
chowska and Paulson (2005). Between 20–30 % RH the in-
crease is less pronounced but still apparent. There are several
possible explanations for the observed behaviour and each
will be discussed in the next section. Previous studies of the
ozonolysis of alkenes have chosen to refer to Criegee birad-
cals formed during ozonolysis as either stabilised or unsta-
bilised (e.g. Kroll et al., 2001a, b; Johnson and Marston,
2008). In the ozonolysis reaction, ozone reacts with the
olefinic bond of an alkene through a 1,3-cycloaddition forming a primary ozonide, which decomposes to form a Criegee biradical and a carbonyl coproduct. This nascent Criegee biradical may possess a range of (vibrational) energies and depending on energy barriers, a certain fraction will be able to isomerise or decompose (unstabilised), with the remainder being below these energy barriers (stabilised), affording a long enough lifetime to be able to undergo bimolecular reactions. As pressure is increased more Criegee biradicals will be ‘stabilised’ as a result of collisional quenching. The results of the HC(O)OH yield as a function of RH can be analysed using two possible scenarios:

4.4 Scenario 1

In a scenario where all Criegee biradicals are in a stabilised form in this system then it is possible to model the HC(O)OH yield as a function of RH. It is assumed that the Criegee radical has one of two fates, reaction with H₂O to form HC(O)OH (Reaction R3) or decomposition (possibly wall

 Table 2. A summary of the experimentally determined HC(O)OH yield obtained in this study, errors quoted are at the 1σ level of sensitivity calibrations.

<table>
<thead>
<tr>
<th>RH %</th>
<th>HC(O)OH Yield</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;1</td>
<td>0.07±0.01</td>
</tr>
<tr>
<td>11</td>
<td>0.18±0.03</td>
</tr>
<tr>
<td>21</td>
<td>0.36±0.05</td>
</tr>
<tr>
<td>27</td>
<td>0.40±0.06</td>
</tr>
<tr>
<td>30</td>
<td>0.41±0.06</td>
</tr>
</tbody>
</table>

Clearly it is not possible to obtain a unique fit to the experimental data as there are no direct measurements of the rate of reaction of the Criegee radical with water (k₃). Indeed, estimates for the reaction rate of the Criegee radical with water range over three orders of magnitude (Calvert et al., 2000). However, a ratio between k₃ and k₄ emerges, where k₄/k₃ is 3.3×10¹⁷ molecule cm⁻³ to obtain an excellent fit to the measurement data. Assuming that k₃ has a maximum value of around 1.5×10⁻¹⁰ cm³ molecule⁻¹ s⁻¹ (gas kinetic limit) this puts an upper limit on the decomposition rate of the stabilised Criegee biradical of 5×10⁻⁷ s⁻¹, similarly, if k₃ is around 1.5×10⁻¹⁷ cm³ molecule⁻¹ s⁻¹ as suggested by indirect measurements then k₄ is only 5 s⁻¹, much lower than theoretical (e.g. Ryzhkov and Ariya, 2004) estimates. Indeed, the work of Ryzhkov and Ariya (2004) suggest a value of k₄ between 5×10⁵ s⁻¹ and 5×10² s⁻¹, which provides a range for k₃ of 1.5×10⁻¹⁵ cm³ molecule⁻¹ s⁻¹–1.5×10⁻¹⁵ cm³ molecule⁻¹ s⁻¹.

4.5 Scenario 2

Alternatively, in a scenario where all Criegee biradicals are in an unstabilised form in this system then it is also possible to model the HC(O)OH yield as a function of RH. Following the work of Kroll et al. (2001a, b), we note that for unstabilised CH₂OO biradicals, the dominant fate is to isomerise to dioxirane which can then rearrange to form HC(O)OH in a vibrationally excited state, the so called “hot acid” channel. This hot acid can either be stabilised (here by water as well as the bath gas) or decompose to form OH. A similar model
which provides a value for the decomposition rate of the un-stabilised Criegee bi-radical of approximately $5 \times 10^{7}$ s$^{-1}$.

A variety of experiments and theoretical investigations suggest that between 35 %–54 % (Alam et al., 2011) of the CH$_2$OO formed are stabilised and it is assumed that the rest must be un-stabilised. The production of OH, believed to arise predominantly (exclusively) from un-stabilised CH$_2$OO is reported to be between 0–22 % of all Criegee biradicals formed (Alam et al., 2011). Data from this work suggests that the yield of HC(O)OH is at least 40 % and indeed is still increasing at the highest water vapour investigated. Hence, if the lower estimates for the fraction of stabilised CH$_2$OO are correct (∼35–40 %), then HC(O)OH must also be formed from un-stabilised CH$_2$OO, presumably via quenching of the “hot acid”.

Hence, current research suggests that scenario 1 could represent up to about 54 % of CH$_2$OO formed. If it is assumed that all the observed HC(O)OH is from this channel only and that the maximum yield is 0.54, then $k_4/k_3$ is $\sim 1 \times 10^{17}$ molecule cm$^{-3}$ and the value for $k_3$ is larger still than estimated in scenario 1, although this study cannot provide a unique value. Can scenario 2 exist? If the lowest evaluations of the amount of stabilised CH$_2$OO are correct, then it would appear that stabilisation is occurring at atmospheric pressure with the increasing relative humidity. Indeed, at high pressures stabilisation must occur, although there is considerable debate as to the value required to approach this high pressure limit. This is further supported by the results of Alam et al. (2011) who report a decrease in radical yield with increased RH, i.e. the stabilised Criegee radical is being removed via reaction with water. In an analogous system, the ozonolysis of alpha pinene, Tillmann et al. (2010) report an increase in the yield of pinonaldehyde with RH as a result of the increased titration of the SCI with water. The result of Tillmann et al. (2010) would suggest the reaction of SCI with water is a ubiquitous phenomenon with far reaching implications for tropospheric chemistry.

In order to further investigate this system and obtain experimental evidence to validate the production of hydroxymethylhydroperoxide (HMHP) during ethene ozonolysis, HMHP was synthesized according to accepted synthesis described by Marklund et al. (1971) and Neeb et al. (1997). The existence of HMHP was confirmed by the appearance of an absorption band at 950–1075 cm$^{-1}$ (Bauerle, 1999; Minkoff, 1954). However, using I$^+$ chemistry, the CIMS instrument was not sensitive to the detection of HMHP, though this does not rule out HMHP production and the detection of HMHP could be achieved using an alternative ionisation scheme or an additional analytical technique. Wolff et al. (1997) did not observe enhancement of HMHP in humid conditions and so do not accept scheme 3 to be responsible for acid production as a result of alkene ozonolysis. However, Neeb et al. (1997) detect high HMHP yields during ethene ozonolysis though they suggest that secondary chemistry through heterogeneous processes led to acid formation.
Recent theoretical work (Anglada et al., 2002) suggests that HC(O)OH is produced via the formation of HMHP through a Criegee intermediate water complex, and HC(O)OH yields increase as a result of increasing relative humidity, which supports the observation of this study.

The formic acid product yields obtained in this study are in good agreement with Neeb et al. (1997) across the range of RH studied. However, this work disagrees with dry yields reported by Wolff et al. (1997) and by Orzechowska and Paulson (2005). Both studies utilise an indirect method of detection of HC(O)OH, which involves a sampling step. Orzechowska and Paulson suggest that formic acid is not a major product of ethene ozonolysis and attribute acid production to the decomposition of HMHP on the solid-phase microextraction (SPME) fibre sampling system. This explanation is somewhat paradoxical, since if HMHP decomposition caused spuriously high acid yields in dry conditions, it is uncertain why HMHP was present in the system in the first place given that its formation is dependent on the presence of water (see Scheme 3). Wolff et al. (1997) also observe large formic acid yields at low RH. However, the formic acid yield that they observe is the sum of primary formic, formic anhydride and HPMF, which could explain the discrepancy under dry conditions.

CIMS is the most sensitive technique to date used to probe the production of HC(O)OH in the ethene + O3 system. Whilst CIMS is selective to HC(O)OH there still remains the possibility that formic acid production is enhanced by heterogeneous processes during ethene ozonolysis. Temperature and pressure control allow this system to be baked out during cleanout procedures, producing a small measured \( k_w \) (wall loss rate coefficient) with respect to ozone and HC(O)OH and so one can expect little impact on HC(O)OH yields from heterogeneous losses. The first-order decay rate of ozone and HC(O)OH with respect to walls were determined to be 6.94 \( \times 10^{-6} \) s\(^{-1}\) and 5.46 \( \times 10^{-7} \) s\(^{-1}\) respectively. Although studies by Neeb et al. (1997) report a time lag between \( d[\text{HC(O)OH}]/dt \) and \( -d[O_3]/dt \) indicating secondary heterogeneous HC(O)OH production, this is not apparent here and so is not concordant with this study.

### 4.6 Loss of CH\(_2\)OO

The dominant loss process for the reaction of the simplest Criegee bi-radical, CH\(_2\)OO (e.g. Taatjes et al., 2008), in the atmosphere on the one hand is not straightforward because of the lack of definitive rate coefficient data. However, it emerges from global model yields that with a rate coefficient of around 1 \( \times 10^{-17} \) cm\(^3\) molecule\(^{-1}\) s\(^{-1}\) reaction with H\(_2\)O should dominate its loss globally. Reaction with NO, NO\(_2\) and SO\(_2\) all compete with water at around the 5 ppbv level (urban environment) if one assumes a rate coefficient of 1 \( \times 10^{-12} \) cm\(^3\) molecule\(^{-1}\) s\(^{-1}\) for these species with CH\(_2\)OO in each case. However, if our estimate is correct, a value of 1 \( \times 10^{-17} \) cm\(^3\) molecule\(^{-1}\) s\(^{-1}\) is probably too small, leading to the conclusion that reaction with water dominates non-decompositional loss in the background atmosphere. Taking the initial analysis at face value and assuming that the production curve (be it considered scenario 1, scenario 2, or some combination) is correct, this study places an upper limit of about 65 % for the yield of HC(O)OH from the decomposition of CH\(_2\)OO formed in the atmosphere (from ethene ozonolysis). The 65 % corresponds to a typical water vapour levels encountered where ethene oxidation occurs in the model. If it is assumed that HC(O)OH can only be formed from stabilised CH\(_2\)OO then the maximum is 54 % (Alam et al., 2011), which is reached assuming a water concentration of around 6 \( \times 10^{17} \) molecule cm\(^{-3}\).

### 5 Model results

Data from this study for the ratio of decomposition of CH\(_2\)OO with reaction with water to produce HC(O)OH has been used in the base case global model integration. In the model there are two photochemical sources of CH\(_2\)OO, ozonolysis of ethene and ozonolysis of isoprene. The base case integration produces 1.0 or 0.8 Tg yr\(^{-1}\) HC(O)OH assuming either 0.65 or 0.52 as the yield from the ozonolysis of ethene. For the ozonolysis of isoprene, it is assumed that the yield of CH\(_2\)OO is \( \sim 0.6 \), where the biradical is a product partner on formation of either methylvinyl ketone or methacrolein. If we assume that HC(O)OH can only be formed from stabilised CH\(_2\)OO and that this is limited to 0.52, the formation of HC(O)OH from CH\(_2\)OO formed from the ozonolysis of isoprene we arrive at a yield of 9.5 Tg yr\(^{-1}\). Effectively this a total yield of about 0.3 and is consistent with Neeb et al. (1996). However, if we assume that the yield of formic acid increases with water vapour according to relationship established in Fig. 6, the temperatures and relative humidities encountered where isoprene is present in the model (i.e. tropics) produces yields of between 0.65 to 0.8 leading to a yield of 11.5 to 14.5 Tg yr\(^{-1}\). In addition reaction of OH with acetylene (ethyne) yields 3.7 Tg yr\(^{-1}\) and from direct emissions, 5.5 Tg yr\(^{-1}\) biomass burning and 1.8 Tg yr\(^{-1}\) anthropogenic sources (combustion) giving a total source of 21.3 to 26.5 Tg yr\(^{-1}\). Hence the base case produces 14 to 19.2 Tg yr\(^{-1}\) from photochemical and 7.3 Tg yr\(^{-1}\) from direct emissions compared with a recent estimate of 48.6 Tg yr\(^{-1}\) from photochemical and 8.1 Tg/year from direct emissions (Paulot et al., 2011). It is clear that the combined production of HC(O)OH from the ozonolysis of ethene and isoprene is very important in the model studies here and all of this arises from the reaction of CH\(_2\)OO (formed from ozonolysis) with H\(_2\)O. Such an assertion is in agreement with other studies such as von Kuhlmann et al. (2003). Loss processes include reaction with OH (9 %), wet deposition (50 %) and dry deposition (41 %), balancing the production processes. Figure 8 shows the surface level yearly average HC(O)OH from the base case integration.
The CRI-STOCHEM model has one of the most detailed chemistry schemes for a global model, but there will be other sources of CH$_2$OO that are not included in this model (e.g. the multitude of short-lived alkenes that are not included) and therefore the reaction of CH$_2$OO with H$_2$O would appear to dominate the in situ formation of HC(O)OH (as suggested by Paulot et al. (2011)). Models underestimate HC(O)OH measurements, especially over the oceans, where in-situ production following the reaction of CH$_2$OO with water will be at its peak.

### 5.1 Possible sources of CH$_2$OO missing from the global model

Stable products from isoprene oxidation, methyl vinyl ketone and methacrolein are included but in the simplified chemical scheme, ozonolysis does not yield HC(O)OH. Using the yields of CH$_2$OO from the work of Aschmann et al. (1996) and Grosjean et al. (1993), the yield of CH$_2$OO is between 0.85–0.95 and even assuming that the fraction that produces HC(O)OH is 0.52 yields around 4.3 Tg yr$^{-1}$, and assuming a fraction of up to 0.8 yields around 6.7 Tg yr$^{-1}$.

Monoterpenes are included in the model but assumed to react as either $\alpha$-pinene or $\beta$-pinene and in the simplified mechanism used do not form CH$_2$OO. Lee et al. (2006) have measured the yield of HC(O)OH from ozonolysis of a series of monoterpenes and found that for $\alpha$-pinene (RH = 4.1 %) the yield was 7.5 % and for $\beta$-pinene (RH = 6.3 %) the yield was 4 %. We recognise that the structure of the monoter- pene will of course dictate whether CH$_2$OO is formed from ozonolysis and that only $\beta$-pinene has a structure that can produce CH$_2$OO. If we assume that $\beta$-pinene represents all the monoterpene emission ($\sim$127 Tg yr$^{-1}$) and that the yield of nopinone (the co-product to CH$_2$OO formation) from $\beta$-pinene ozonolysis is around 20 % (Lee et al., 2006) we can invoke the 0.52 yield to produce an estimate of about 1 Tg yr$^{-1}$, if the yield is 0.8 then this rises to about 1.5 Tg yr$^{-1}$. Larsen et al. (2001) report HC(O)OH yields from ozonolysis of $\beta$-pinene (38 %), using these data produces 3.5 Tg yr$^{-1}$. Adding the monoterpenes, methyl vinyl ketone and methacrolein yields (up to $\sim$10 Tg yr$^{-1}$) with the base case estimate produces a photochemical yield from $\sim$25–30 Tg yr$^{-1}$, closer to the biogenic estimate of Paulot et al. (2011).

Furthermore, all 1-alkenes (Johnson and Marston, 2008) can undergo ozonolysis to yield CH$_2$OO and subsequently HC(O)OH. Hence there are myriad small sources of HC(O)OH that will contribute to global HC(O)OH.

### 6 Conclusions

This study has confirmed that the yield of HC(O)OH from the ozonolysis of ethene has a strong water dependence, rising rapidly with additional water. Assuming a simple two channel model for the fate of the CH$_2$OO radical it has been possible to estimate the ratio of the rate coefficient for the reaction with water compared ($k_3$) with decomposition ($k_4$). Such an analysis suggests that $k_3$ probably ranges between $1 \times 10^{-12}–1 \times 10^{-15}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ and as such will indeed be the dominant loss process, other than decomposition, for this radical in the atmosphere. Global model integrations confirm that this reaction between CH$_2$OO with water is responsible for over half the production of HC(O)OH. However, HC(O)OH is still underestimated by the model. Unless there are missing biological sources, one is tempted to conclude that the myriad missing short-lived alkenes that could all contribute to CH$_2$OO production could provide the missing source, particularly in the marine boundary layer where Reaction (R3) will be at its highest rate. Further analysis shows that monoterpene oxidation and the ozonolysis of methyl vinyl ketone and methacrolein could contribute up to 10 Tg yr$^{-1}$ to the HC(O)OH budget.

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