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Analytical methodologies for oxidized organic compounds in the atmosphere†

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Oxidized compounds in the atmosphere can occur as emitted primary compounds or as secondary products when volatile emitted precursors react with various oxidants. Due to the presence of polar functional groups, their vapor pressures decrease, and they condense onto small particles. Thereby, they have an effect on climate change by the formation of clouds and scattering solar radiation. The particles and oxidized compounds themselves can cause serious health problems when inhaled. Therefore, it is of utmost importance to study oxidized compounds in the atmosphere. Much ongoing research is focused on the discovery of new oxidized substances and on the evaluation of their sources and factors influencing their formation. Monitoring biogenic and anthropogenic primary oxidized compounds or secondary oxidized products in chamber experiments or field campaigns is common. New discoveries have been reported, including various oxidized compounds and a new group of compounds called highly oxidized organic molecules (HOMs). Analytics of HOMs are mainly focused on chromatography and high-resolution mass spectrometry employing chemical ionization for identifying and quantifying compounds at low concentrations. Oxidized compounds can also be monitored by spectrophotometric methods in which the determinations of total amounts are based on functional groups. This review highlights recent findings on oxidized organic compounds in the atmosphere and analytical methodologies used for their detection and quantification. The discussion includes gas and liquid chromatographic methods, sampling, extraction, concentration, and derivatization procedures involved, as well as mass spectrometric and spectrophotometric methods.

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Environmental significance

Oxidized compounds in the atmosphere can occur as emitted primary compounds or as secondary products when volatile emitted precursors react with various oxidants. These are currently under intensive investigation due to environmental and health aspects. The discovery of new oxidized species and subsequent evaluation of their sources and factors influencing their formation is an area of research that has recently received considerable attention. Monitoring of biogenic and anthropogenic primary oxidized compounds or secondary oxidized products in chamber experiments or field campaigns is common. A new group of compounds called highly oxidized organic molecules (HOMs) has recently been discovered. This review highlights the latest findings on oxidized organic compounds in the atmosphere and analytical methodologies used to detect and quantify them.

1. Introduction

The atmosphere contains solid particles and liquid droplets that mix in the air we breathe and form atmospheric aerosols. Particulate matter (PM) consists of liquid and solid particles, and these can contain inorganic or organic constituents (compounds). The organic part can be divided into primary organic aerosols and secondary organic aerosols (SOAs).^{1–3}

Primary organic aerosols contain organic compounds emitted directly from original sources and they are not formed in atmospheric processes.⁴ In contrast, SOAs are formed from organic matter in the atmosphere, usually through oxidation involving ozone, anthropogenic nitrogen oxides (NO_x), and various radicals.^{1,5} Hydroxyl radicals (OH) are important oxidants during the daytime as they are formed under the influence of light.⁶ The reactions with precursors yield various reactive peroxide radicals which act as intermediates in the formation of oxidized compounds according to several proposed reaction mechanisms.^{7–11} Among the oxidation reactions occurring in the dark, the most essential are nitrate radicals (NO₃), which are formed in the reaction between ozone and nitrogen dioxide.¹² In addition to oxidants, atmospheric sulfuric acid increases SOA production by acting as an acidic

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catalyst enhancing reactions and by producing additional organosulfate compounds.¹³ Gaseous volatile organic compounds (VOCs) are typical emitted precursors that react with the above oxidants to form secondary products.

Organic aerosols (OA) have an impact on climate change because of their capability to absorb and scatter solar radiation, and they affect cloud nucleation by aerosol–cloud interactions.¹⁴ The vapor pressures of the oxidized compounds are decreased compared to their precursors, allowing them to either participate in nucleation reactions that form new particles or to condense on the surfaces of existing particles. The presence of polar organic compounds can increase the hygroscopicity of particles and improve their ability to act as condensation nuclei.^{15,16} The formation also has negative health effects by increasing PM_{2.5} and causing *e.g.* asthma symptoms and lung cancer when inhaled.¹⁷ Alpha-pinene SOA components have recently been demonstrated to potentially cause oxidative stress to lung cells.¹⁸ The oxidized forms of polycyclic aromatic hydrocarbon compounds (PAH) are considered even more toxic and bioavailable compared to their original form.^{19,20}

Modern mass spectrometric (MS) methods are the most important analytical methodologies in today's atmospheric studies. There are recent reviews discussing this topic as the technology is constantly developing and providing new information, for example, see ref. 21–23. Here, we discuss oxidized compounds commonly analyzed in the atmosphere or compounds generated in laboratory smog chamber experiments to assess emission sources and factors influencing their formation and occurrence. Several compounds have been monitored actively from the atmosphere for at least two decades,¹³ but the recently discovered new compounds are also discussed. All the most common analytical methods from sampling to the detection and identification of compounds are presented, assessing their advantages and disadvantages.

2. Oxidized compounds in the atmosphere

Oxidized compounds can be used as markers or tracers of their sources and precursors. The term tracer has a specific definition including being ideally a precursor and source specific, stable, and produced with high yield to be analyzed.^{24,25} We found only a few publications preferring and justifying the use of the term marker as the criterion for tracers are not fulfilled, especially regarding compound stability and source specificity.^{13,24} The impact of OA on climate and health is considerable, and the analysis of oxidized marker compounds is important as they provide information about the factors influencing the formation of OA, such as formation mechanisms and the origin of precursors.^{5,26} The concentrations of the markers vary locally and seasonally, which can be explained by possible different emission rates of precursors and meteorological conditions including *e.g.*, temperature, relative humidity, intensity of solar radiation, and concentration of oxidants.²⁷ In the eastern parts of the Mediterranean, the proportion of SOAs in PM has been between 65 and 78%, and sometimes more than 80%, due to

increased concentrations of precursors and ozone in the summer.^{28,29} In the urban area of Hong Kong, the secondary formation has produced 51.4% of the total organic carbon (TOC) in the atmosphere.¹² The secondary formation contributed to PM_{2.5} only 2.8% in the rural area of Nanjing (China), where emissions of primary compounds had greater contribution including sources such as biomass burning (77.4%), fungal spores (12.2%), plant debris (5.5%), and soil resuspension (2.1%).³⁰ In addition to local sources, long-distance transportation has been assessed to have a significant effect on the prevalence of compounds in the studied areas.^{3,30,31}

Here, we present specific oxidation products of the most common biogenic and anthropogenic VOC compounds, as well as other compounds emitted as primary products or less specific products oxidized from various precursors. We are aware that our selected examples provide only a brief overview as there are many publications presenting a more diverse range of different markers. In addition, we introduce highly oxidized organic molecules (HOMs), a recently discovered group of oxidized compounds. We use the term marker to align and simplify the discussion.

2.1. Isoprene markers

Isoprene (2-methyl-1,3-butadiene) is the most common biogenic VOC (BVOC) emitted especially by deciduous trees.³² Marine phytoplankton has also been found to have a possible effect on increasing isoprene emission in the spring when photosynthesis intensifies.³ Isoprene reacts with OH, first forming a peroxy radical derivative, from which the resulting reaction pathways and products depend especially on atmospheric acidity and concentrations of NO_x and sulfate.^{13,33} Ozonolysis also contributes to isoprene oxidation in which typical products are formed, albeit less compared to the OH-initiated reactions.³⁴

At low NO_x concentrations, a more favored intermediate product is isoprene-epoxydiol (IEPOX), which has an epoxy ring that can react under acidic conditions with nucleophiles (Fig. S1†).¹⁰ At higher NO_x concentrations, the peroxy radical derivative forms methacrolein that is oxidized further to methacrolein acid epoxide (MAE), which also reacts with nucleophiles under acidic conditions (Fig. S2†).¹¹ Isoprene marker concentrations have generally been found to be the lowest in winter and the highest in the summer when isoprene emission and OH concentrations are at their peak values.^{12,27,35}

The most commonly monitored IEPOX derivatives have been 2-methylerythritol and 2-methylthreitol (known as 2-methyltetrols, 2-ME),^{5,26,27,34} which are formed at low NO_x concentrations by the addition of water with acidic catalysis. Recently,^{9,36} oxidation products of 2-MEs have been detected and identified, namely 2-methyltartaric acids, 2-methylerythronic acid, 2-methylthreonic acid, and their 3-methyl isomers. These compounds have been oxidized several times, based on which (and other corresponding higher generation oxidation products in general) SOA aging could be expressed.³⁷ Other analyzed IEPOX derivatives, abundant at low NO_x concentration, are 2-methyl-2,3,4-trihydroxy-1-butene and *cis* and *trans*-2-methyl-



1,3,4-trihydroxy-1-butene (so-called C5-alkenetriols), which are formed in the same reaction path with 2-MEs.^{10,13,27,33} Aerosols containing sulfuric acid yield organosulfate derivatives of 2-ME (2-MEOS), which have been monitored frequently.^{33,38,39} These derivatives are sensitive and degraded into C5-alkenetriols and 2-ME form upon acid catalysis.^{13,38} As SOA aging indication, 2-MEOSs can react further by intermolecular rearrangement into cyclic compounds, producing tetrahydrofuran or lactone derivatives.^{13,40}

2-Methylglyceric acid (2-MGA) is also a commonly monitored isoprene marker, the concentration of which typically increases at higher NO_x concentrations.^{10,24,26} Higher relative amounts of 2-MGA to 2-MEs are due to traffic or by wood burning, which is common in the winter season.^{5,27} It has also been suggested that 2-MGA occurs in winter due to lower temperatures, as the reaction pathway forming 2-MGA is possibly more favored at lower temperatures.¹² A reaction path for the formation of 2-MEs at increased NO_x concentration has also been presented,⁴¹ as 2-MEs have been observed to be abundant in urban areas, despite increased NO_x concentrations.¹²

2.2. Terpene markers

Terpenes are significant BVOCs emitted from vegetation, especially from coniferous trees.²⁷ Alpha and beta-pinene as well as delta-3-carene and limonene are the most common monoterpenes that react with atmospheric oxidants such as ozone, OH, and NO₃.⁴² SOAs derived from terpenes have been determined to be generally more abundant in Europe than in Asia or America, where isoprene had greater contribution to SOAs.²⁴ In Alaska, isoprene has been found to be a more significant source of SOA than monoterpenes, despite the vegetation being mostly boreal coniferous forest.²⁷ Unlike isoprene, no clear seasonal variation has been observed for monoterpene markers generally.^{24,35} The highest concentrations of markers have usually been observed in the spring,^{3,27} but increased concentrations have also been observed in the fall due to long-distance transport.³ In addition to monoterpenes, a common sesquiterpene beta-caryophyllene's oxidization marker beta-caryophyllinic acid (Fig. S3†) has been monitored, however, to a lesser extent because beta-caryophyllene is reactive and less abundant in the atmosphere due to its lower volatility.⁴³ Beta-caryophyllinic acid concentrations have typically been highest in the winter, probably due to the release of beta-caryophyllene from the burning of biomass and wood.^{3,24,27}

The most common markers of monoterpenes (Fig. S3†) are the first-generation SOA markers *cis*-pinonic acid and pinic acid, and their second-generation oxidation products such as 3-hydroxyglutaric acid (3-HGA), 3-methylbutane-1,2,3-tricarboxylic acid (3-MBTCA), 3-hydroxy-4,4-dimethylglutaric acid, 3-acetylpentanedioic acid, and 3-isopropylpentanedioic acid.^{5,12,26,27,30,44} Malic acid has been proposed to be a higher-generation oxidation product of terpenes based on a high correlation with terpene markers in field studies and in chamber experiments.^{12,45} However, it is not a very specific marker as it can be formed from other precursors from anthropogenic and biogenic sources.⁴⁵

The first-generation products have been found to form under atmospheric conditions with OH or ozone and at lower NO_x concentrations, while an increase in NO_x concentrations has yielded more higher-generation products.^{34,44} In smog chamber experiments by Müller *et al.*,⁴⁴ *cis*-pinonic acid was demonstrated to react with OH radicals, as pinic acid was not observed to produce 3-MBTCA. Occurring in the gas phase, *cis*-pinonic acid was observed to have higher volatility, and its fraction increased at higher chamber temperatures leading to higher 3-MBTCA concentrations. The monoterpene SOA has been found to be less aged mostly in the spring and winter time, as the relative amount of 3-MBTCA compared to the first-generation products has been smaller.^{27,35} Correspondingly, the concentrations of 3-MBTCA or 3-HGA have been at their highest level in the summer, indicating SOA aging.^{27,35} The higher ratio of 3-MBTCA to 3-HGA has been applied to assess alpha-pinene being the dominant monoterpene rather than beta-pinene and limonene.^{26,27} Typical markers to monitor *d*-limonene SOAs include limonic acid, limonoic acid, and 3-carboxyheptanedioic acid.^{46–48}

2.3. Anthropogenic markers

Ubiquitous anthropogenic VOC and SVOC in the atmosphere include aliphatic hydrocarbons, BTEX compounds (benzene, toluene, ethylbenzene, and xylenes), other alkyl benzenes, and PAHs originating from anthropogenic sources, *e.g.*, vehicle exhaust and biomass burning.^{49,50} Aromatic compounds and unsaturated hydrocarbons containing a double bond react with ozone and OH.^{19,34,49–51} Increased concentrations of anthropogenic markers (Fig. S4†) have generally been observed in the winter when biomass burning and other anthropogenic activities are more common.^{24,31} Increased anthropogenic NO_x concentrations have also yielded oxidized nitro compounds which have been monitored and discovered recently.^{24,35,51}

2,3-Dihydroxy-4-oxopentanoic acid is a commonly analyzed marker, which is formed when benzene and other alkylbenzenes are oxidized photochemically or with ozone, regardless of NO_x concentrations.^{5,24,35,52–54} In general, this marker has been more abundant in urban areas, but increased concentrations have been observed in rural areas possibly due to biomass burning.²⁶ Another frequently analyzed oxidation product of alkylated benzenes has been benzoic acid.^{24,31,34,55} Phthalic acid and 4-methylphthalic acid have been considered to be oxidation products of naphthalene and its methylated derivatives.^{5,35,45,54,56} Sato *et al.*⁵¹ recently discovered the formation of 3,5-dinitrosalicylic acid from toluene and 4-nitrophthalic acid from naphthalene. These new products are proposed to be more specific markers.

Anthropogenic aromatic and aliphatic hydrocarbons are typical precursors of dicarboxylic acids, which are very common in the atmosphere.^{50,57} Cyclic alkenes have been reported to form mostly dicarboxylic acids in ozonolysis, and to a lesser extent hydroxy and oxo acids as well as methyl and ethyl esters.⁵⁸ The ratio between adipic acid and azelaic acid, which are formed from anthropogenic cyclohexene and biogenic oleic acid, has been applied to determine whether the acid sources are biogenic or anthropogenic.⁵



In addition to secondary formation, anthropogenic activities have a significant effect on primary emissions. There are numerous studies showing positive correlations between increased anthropogenic sources and marker concentrations. Levoglucosan is a very commonly monitored marker which concentrations have been higher in urban areas and during colder seasons due to the increased biomass burning.^{27,30,34,35} Terephthalic acid has been monitored as an oxidized marker emitted from burning plastics.³⁵ Cresols and other phenolic compounds are emitted from biomass burning, and they can react with NO_x to form methylnitrocatechols and nitroguaiacols, which have been observed at increased concentrations in the winter.^{24,54} The above-mentioned markers are common in combustion reactions; therefore, such natural processes can also be sources.

2.4. Other oxidized markers

Organic acids may not be precursor-specific since the same acids have been observed to form from different precursors or they have been primarily emitted from anthropogenic and biogenic sources.^{24,34,59} Many smaller dicarboxylic acids are formed in the oxidation of homologous larger dicarboxylic acids, which is why oxalic acid (C2) is generally the most abundant dicarboxylic acid followed by malonic acid (C3) and succinic acid (C4).^{60,61} Bikkina *et al.*⁶² demonstrated in their recent chamber experiments that the ozonolysis of isoprene yielded a great amount of oxalic acid and smaller amounts of other acids such as malonic acid, succinic acid, and glyoxylic acid. Sato *et al.*⁴⁵ recently discovered *meso*-tartaric acid to be a potential marker for toluene since *meso* diastereomer occurs rarely in nature. In addition, *DL*-isomers of tartaric acid, malic acid, succinic acid, fumaric acid, and maleic acid were produced. Saturated and multifunctional dicarboxylic acids have been observed at increased concentrations in the summer time possibly due to their formation mainly by photochemical oxidation.³¹ On the other hand, increased concentrations of dicarboxylic acids in urban areas have been observed in the autumn and winter because of increased anthropogenic activities and pollution.¹²

The atmosphere has been monitored for carbohydrates which are typically primary oxidized compounds from various sources *e.g.*, biomass burning, plant debris, and fungal spores.^{30,63} Like levoglucosan, markers such as mannosan and galactosan have been typical markers for biomass burning.³⁵ Increased concentrations of plant and fungal spore-originated markers arabinol, mannitol, and glucose have been observed in PM_{2.5} due to soil particles released into the air.³⁵ Carbonyl compounds are common in the water-soluble fraction of PM, where they can act as intermediates of oxidation reactions and have harmful effects on health.⁶⁴ Glyoxal and methylglyoxal are small dicarbonyl compounds frequently analyzed in the atmosphere,^{65–68} in which they occur as primarily emitted or being formed in oxidation reactions of several different precursors.^{49,66,69} These compounds can enhance SOA formation by oligomerization or by oxidation reactions yielding organic acids and generating more ozone and OH, increasing the

oxidation capacity of the aerosol.^{65,69,70} Short-chain aldehydes and carbonyls have been analyzed as a result of traffic emissions.⁷¹ They have possibly been formed from HOMs decomposing in the particle phase.⁷²

2.5. Highly oxidized organic molecules

Oxidation reactions in the atmosphere yield products that are rich in oxygen relative to the amount of carbon in the molecule. These compounds are called HOM compounds but the exact definition is still pending as this group of compounds is a relatively new discovery.⁷³ Ehn *et al.*⁷⁴ presented the term HOM for the first time referring to highly oxidized multifunctional compounds found in the oxidation reaction of alpha-pinene. The compounds were observed to form clusters with nitrate ions, and the oxygen-carbon ratio (O:C) of molecules was determined by high resolution mass spectrometry (HR-MS) to be between 0.7 and 1.3. The analyzed compounds were naturally charged and assumed to contain carbonyl groups and geminal diols, which is why the multifunctionality term was included in the name. The term HOM has also been referred to as highly oxidized organic compounds hereafter, where functional groups themselves are not exactly included in the definition.⁷³ Due to the high degree of oxidation, the compounds have a lowered vapor pressure which is why the group has also been denoted as extremely low-volatility organic compounds, which further emphasizes their importance in the formation of SOAs.^{72,75}

Bianchi *et al.*⁷³ recently suggested three criteria to define HOMs and to distinguish them from other oxidized species. As the first criterion, HOMs should be formed *via* an autoxidation reaction mechanism involving peroxy radicals. They preferably possess at least one peroxide group, but this requirement has been relaxed in more recent publications.^{9,76,77} The second criterion states that HOM formation occurs only under atmospherically relevant conditions. These two criteria exclude oxygen-rich carbohydrates and multifunctional organic acids, which may be primary biogenic compounds formed in the atmosphere along with other reaction pathways or their formation occurs in combustion reactions.^{44,78} Hydroperoxide groups are thought to be a reason for a high O:C ratio in HOMs,⁷⁵ and this is proposed as the third criterion for defining HOM compounds. A ratio exceeding 0.7 is considered to be the limit.⁷² However, Bianchi *et al.*⁷³ proposed as their third criterion that the HOM molecule should contain six or more oxygen atoms, but in some cases only five. The higher absolute amount of oxygen is justified by the fact that the molecule would be more likely formed by the reaction pathway suggested for HOMs.

The formation of HOMs has been found to be involved in the reaction of peroxy radicals (autoxidation).^{7,75} This reaction and its mechanism presented in Fig. S5† would explain hydroperoxide and carbonyl functionalities to occur in HOMs and in SOA constituents in general. In the reaction, the OH and molecular oxygen form a peroxy radical, which further undergoes a proton transfer (H-shift) and reacts with oxygen. A new oxidized group is added to the original molecule, which in turn further



increases the rate of autoxidation. HOMs have been found to form especially from VOC precursors with a cyclic structure and double bonds, such as monoterpenes and anthropogenic aromatic hydrocarbons.⁷⁶ Wang *et al.*⁷⁷ demonstrated in a recent study that even saturated hydrocarbons are able to form highly oxidized species by autoxidation under atmospheric and combustion engine's conditions. Also, the aldehydes employed as surrogates representing the first-generation oxidation products of alkanes were found to produce high yields of HOMs. This was a logical observation, since the initiation of autoxidation requires a polar functional group in the reacting molecule, as shown in Fig. S5.† Interestingly, the formation was also observed at higher NO_x concentrations, where the autoxidation reaction is usually terminated.

The exact structures of HOM compounds are not known so far, but structures have been proposed by applying quantum mechanical computational methods which are not covered in this review.^{76,77} HOMs are, based on experimental evidence, assumed to contain other functional groups, such as hydroxyl and carbonyl groups, in addition to peroxide groups.^{72,73,76} Nitrogen-containing compounds are likely to contain organic nitrate functionalities.^{75,79,80} In sulfate-containing aerosols, HOMs might act as precursors producing highly oxidized organosulfate compounds condensed in the particle phase.⁷² HOMs occur as monomers or form dimers with their peroxide groups.⁷⁶ The current criteria previously presented were determined by researchers specializing in HOM research, but due to the novelty of the compound group and the lack of experimental evidence, the criteria on the basis of which the classification has been made must be clearly defined for the time being.⁷³ As an example, Jaoui *et al.*⁹ have recently presented that 2-methyl-tartaric acids formed from isoprene could be classified as HOMs. It was stated that they are formed by a mechanism similar to autoxidation under normal atmospheric conditions, and the amount of oxygen in them fulfilled the presented criteria. Wang *et al.*⁷⁷ did not classify the highly oxidized species formed from saturated hydrocarbons as HOMs if they were produced at higher temperatures and/or possessed less than six oxygen atoms.

3. Analytical methods for oxidized compounds in the atmosphere

The most common analytical procedures and instrumental techniques to determine oxidized compounds in aerosol samples are presented in this section. Methods to analyze specific markers by chromatographic methods are introduced first. These offline analytical methods focus on particle phase compounds, in which sampling and sample preparation steps prior to analysis are also introduced. Capillary electromigration techniques are not covered in this review, because they have only been employed for the separation of some organic acids.^{31,61,81} Other techniques presented include HR-MS techniques to online detect gas phase HOMs and spectrophotometric applications to determine the total amount of oxidized compounds based on their functional groups.

3.1. Analysis of markers by chromatographic methods

The analysis of aerosol samples is challenging due to their complexity, which is why gas chromatography (GC) methods have often been applied to the analysis of aerosol samples.⁸² One crucial requirement is the volatility of the analytes, which does not apply to highly oxidized compounds with reduced vapor pressure.⁸³ Thereby, the compounds cannot be analyzed in their native form, but need to be derivatized to increase volatility. Some typical analyses performed with GC techniques are presented in Table 1.

Liquid chromatographic (LC) techniques are also common analytical techniques for the determination of SOA markers (Table 2). Compared to GC methods, higher molar mass compounds like humic substances, and larger dimers can be analyzed by LC methods.⁸⁴ Another advantage is that the decomposition of compounds and the formation of artifacts due to the high temperatures employed in GC can also be prevented. Cui *et al.*³⁸ demonstrated artifact formation as a result of thermal decomposition of 2-MEOS resulting in up to 188% higher concentrations of 2-MEs by GC than by LC. In addition, C5-alkene triols and 3-methyltetrahydrofuran-3,4-diols were detected in the pure standard of sulfate derivatives when analyzed by GC-MS. These were not observed upon analysis by LC-MS. It is important to consider the thermal stability of compounds as ion sources are using high temperatures to ionize analytes. For example, organic acids can decarboxylate at high temperatures, which might complicate tentative identification or decrease the sensitivity in the targeted analysis. An additional advantage of LC techniques is that by choosing a column and a detector suitable for the analyzed compounds, laborious derivatization steps can be avoided.^{2,30,85}

3.1.1. Sampling. Field samples have typically been collected with filters by employing pumps with flow rates up to about 1.0 m³ min⁻¹ and sampling times from a few hours to a few days.^{9,30,86,87} Separate collectors are employed in sampling if it is desired to analyze compounds from the gas phase and from the particle phase separately.⁸⁷ This has been typical when gas-particle distributions have been assessed for oxidized markers.⁵⁰ The advantage of a high sample volume is that a larger number of compounds end up in the collectors, enabling lower concentrations to be determined. The drawback is that the possible adsorption of gaseous analytes to the collected particle phase is increased and systematically higher concentrations will be determined.^{35,50} In addition, sampled gaseous oxidants might form additional oxidized species (*i.e.* artifacts) on filters by reacting with sampled organic compounds and matter.^{50,88}

The filter material is typically pure quartz, often baked at a high temperature (400–500 °C) for several hours prior to sampling to remove impurities.^{30,38,86} There are alternate options such as polycarbonate filters by which Kwiezinski *et al.*² obtained better recoveries for small and polar organic acids by water extraction, compared to quartz filters. Filters could be pre-weighed to determine gravimetrically the amount of collected particles after the sampling.³⁴ In this case, it is also possible to determine the amount of TOC and elemental carbon, allowing





Table 1 Selected examples on studies quantifying oxidized OA species with GC techniques

Analytes	Extraction	Derivatization	GC technique	Concentrations measured (ng m ⁻³)	Ref.
Organic acids	3 × 10 min water by UAE. Filtration and pH adjusted with potassium hydroxide to 8.5–9.0 followed by evaporation to almost dryness with rotary vapor	14% Boron trifluoride in butanol (100 °C, 1 h). Water and ACN were added and derivatives were extracted to <i>n</i> -hexane. Silylation of hydroxy groups with BSTFA (80 °C, 30 min) followed by drying with nitrogen flow and dilution into <i>n</i> -hexane	GC-EI-MS, GC-EI-TOF MS, and GC-FID. DB-5MS columns (30 m × 0.25 mm × 0.25 μm)	Hydroxycarboxylic acids: 0.1–27.3 Oxaloacetic acids: <0.005–2 Dicarboxylic acids: 5.36–42.3 Quantitated by FID	86
Carbonyl compounds	Quartz filters were extracted with 1 : 1 (v/v) ACN-DCM and concentration with rotary evaporator prior to derivatization steps	Filters impregnated with BHA (gas phase) and quartz filters (particles) were immersed in ACN containing 4% BHA (room temperature, 24 h). Solvent exchange to ethyl acetate with rotary evaporator. The concentration with N ₂ was performed followed by silylation with BSTFA (60 °C, 1 h)	GC-EI-MS DB5-column (30 m × 0.32 mm × 0.5 μm)	Glycolaldehyde: 0–826 Hydroxyacetone: 0–579 Glyoxal: 46–1200 Methylglyoxal: 88–2690 Nonanal: 0–500 Decanal: 0–230	66 and 90
Aged isoprene SOA markers	Tetracosane-d ₅₀ and <i>cis</i> -ketopinic acid were spiked on quartz-fiber filters prior to extraction. Sonication in 5 mL of MeOH for 1 h. Evaporation to dryness with nitrogen flow	Silylation in pyridine with BSTFA and 1% TMCS (70 °C, 1 h)	GC-ion trap MS EI and methane-CI RTX-5MS column (60 × 0.25 mm × 0.25 μm)	Methyltartaric acid: 1.52–6.27 2-Methylthreonic acid: 0.12–0.69 3-Methylthreonic acid: 0.04–0.22 2-Methylglyceric acid: 1.182–4.4	36
Terpene SOA and isoprene SOA markers	Quartz filter aliquots were extracted with 3 × 7 mL of DCM/MeOH (2 : 1 v/v) with sonification. Extracts were filtered and evaporated dryness with a rotary evaporator and nitrogen flow	Silylation in pyridine with BSTFA and 1% TMCS (70 °C, 3 h). Diluted in <i>n</i> -hexane and tridecane was added as ISTD prior to GC analysis	GC-EI-MS DB-5MS column (30 m × 0.25 mm × 0.25 μm)	Monoterpene markers: 0.1–25.3 Isoprene markers: 0.01–46.6 Beta-caryophyllinic acid: 0.1–4.0	27
Organic acids, terpene SOA markers, isoprene SOA markers, levoglucosan	<i>Cis</i> -ketopinic acid was spiked on quartz filters prior to extraction. Filters were extracted with 3 × 30 ml of DCM/MeOH (1 : 1 v/v) by UAE for 3 × 20 minutes. Extracts were combined, filtered, and evaporated to dryness with a rotary evaporator and nitrogen flow	Step 1: carboxyls were derivatized by adding a heated solution of MHA in ACN (70 °C, 1 h) Step 2: methylation of carboxylic groups by adding TMSD and MeOH (in an ultrasonic bath, 20 min) Step 3: silylation of hydroxyl groups with BSTFA, 1% TMCS, and pyridine addition (70 °C, 60 min)	GC-EI-MS HP-5MS column	Monocarboxylic acids: 0.82–130 Dicarboxylic acids: 0.30–88.2 Aromatic acids: 0.78–52.0 Hydroxycarboxylic acids 0.39–58.7	34
				Monoterpene markers 0.32–31.2 Isoprene markers 0.53–57.5 Levoglucosan 0.65–21.1	



Table 2 Selected examples on studies quantifying oxidized OA species with LC techniques

Analytes	Extraction and possible derivatization	LC technique and columns	Concentrations measured (ng m ⁻³)	Ref.
Glyoxal and methyl glyoxal	30 min UAE with 1 : 1 (v/v) water and MeOH. μ -SPE device with DNPH-coated C18 adsorbent was set to extracts and vortexed for 3 minutes and then dried. 150 μ L of ACN was eluted through the device by centrifuging to perform on-sorbent derivatization and extraction	LC-DAD (detection at 450 nm) C18-column (150 \times 4.6 mm, 5 μ m)	Glyoxal: 0.38 Methylglyoxal: 0.41	67
2-MEs and 2-MEOS	UAE with MeOH twice, cooling in between. Filtration and evaporation with nitrogen flow. Extract residues reconstituted to 150 μ L ACN/water (95 : 5 v/v) and diluted by a factor of 20 or 30	LC-ESI(-)-Q-TOF MS BEH amide HILIC column (100 \times 2.1 mm, 1.7 μ m)	2-MEs: 0.14–0.86 2-MEOS: 0.40–2.33	38
Organic acids and organosulfates	Extraction with 0.25 mL (13 mm quartz filters) or 3 mL (47 mm polycarbonate filters) of water with shaking for 2 h. 2.9 mL of 3 mL extract was diluted with 3.1 mL of water. All samples were filtered (0.2 μ m) prior to analysis	IC-ESI(-)-Orbitrap MS AS19-4 μ m hydroxide-selective anion-exchange column	Monoacids: 0.8–23.1 Diacids: 0.1–98.2 Citric acid: 1.8 Terpene-derived acids: 0.2–3.0	2
Carbohydrates and monoterpene SOA markers	UAE (in an ice bath) of filters with 3 \times 3 mL of MeOH for 15 min. Extracts were filtered (0.22 μ m) and evaporated with N ₂ . Reconstitution to 500 μ L of 0.01% acetic acid : ACN solution (20 : 60 v/v to analyze carbohydrates and 80 : 20 v/v for organic acids)	LC-ESI(-)-MS/MS Monoterpene SOA markers: Thermo Hypersil C ₁₈ (100 \times 2.1 mm, 3 μ m) Carbohydrates: Prevail column (150 \times 4.6 mm, 3 μ m)	Organosulfates: 0.3–17.8 Levogluconan: 32.27–201.85 Sugars: 0.47–2.63	30
Glyoxal and methyl glyoxal (snow samples)	Spike of 4-fluorobenzaldehyde (ISTD) and BHA solution (derivatization agent) to melted samples. pH adjustment to 3.5 and PDMS coated stir bar was placed for 20 h and 1000 rpm. Oxime derivatives were desorbed by stirring (500 rpm) in 1.5 mL of ACN for 3 h	LC-ESI(+)-MS/MS (ion trap analyzer) Pursuit XRS C8 column (150 \times 2.0 mm, 3 μ m)	Sugar alcohols: 0.25–3.22 Organic acids: 0.03–2.22	68
C1–C10 aldehydes and cyclohexanone	UAE of filters with 2 \times 10 mL water followed by dilution to 25 mL. 10 mL of extract were derivatized by adding acidic DNPH solution followed by mixing for 5 min	IT-SPME-cLC- μ ESI(-)-MS	Glyoxal: 0.085–16.3 ng mL ⁻¹ Methylglyoxal: 0.126–3.6 ng mL ⁻¹	64
	2 mL of derivatized extract was applied through the IT-SPME column followed by 34 μ L injection of water to clean unretained impurities	Zorbax C18-column (150 \times 0.5 mm, 5 μ m) IT-SPME: TRB-35 GC-column (43 cm \times 0.32 mm \times 3 μ m) integrated to injector	Acetaldehyde (C2): 10 C3–C4 and C6–C8 aldehydes: 1	

for the estimation of the markers' proportions in the OA.²⁷ Gas phase collectors typically contain an adsorbent like Amberlite XAD-2 (ref. 87 and 89) or Amberlite XAD-4.^{36,50}

Polar compounds are water-soluble and condense into particles, allowing them to precipitate from the atmosphere into *e.g.*, water or snow. Because of this, oxidized compounds of the atmosphere have been analyzed not only from aerosol samples but also from rainwater, snow samples, and ice core samples.^{68,90–92} There has been particular interest in snow samples to examine ice nucleation properties and the occurrence of markers in ice.^{68,91} Despite potential changes in compounds trapped in ice due to cosmic radiation,⁹³ such compounds have been analyzed to provide information about the history of climate change. For example, analysis of ice core samples taken from Alaska has revealed information about the occurrences of isoprene and terpene markers, as well as climate change-related climate oscillations from the 1660s.⁹⁴ For this, a sample was taken from a depth of 180 m, as from other upper layers depicted different eras.

3.1.2. Extraction procedures. The oxidized compounds have typically been extracted with polar solvents like water, methanol (MeOH), and their mixtures.^{2,36,38,67,86} The extraction is more challenging if a wide range of compounds with varying polarities are desired to be analyzed. In these cases, the polarity of the solvent is reduced with mid-polar solvents such as acetonitrile (ACN) and dichloromethane (DCM).^{27,34,66,87} Kowalewski and Gierczak⁹⁵ developed a multi-step derivatization method in which a solvent mixture of ACN and DCM (1 : 1 v/v) was employed for dissolving non-polar (*e.g.* C8–C10 aldehydes) and polar compounds (*e.g.* citric acid and tartaric acid). When choosing the extraction solvent, its suitability for the following analysis must also be considered. For example, anhydrides are converted into acids by water extraction, which will increase their concentration.⁸² Bateman *et al.*⁹⁶ demonstrated artifact formation as carbonyl species and organic acids were converted into hemiacetals, acetals, and methyl esters when MeOH was employed in the extraction.

Ultrasound-assisted extractions (UAE) have been very common,^{34,38,64,86} but mechanical extraction with an orbital shaker or by vortexing has also been employed providing good and quantitative recoveries.^{2,66} Albinet *et al.*⁹⁷ compared MeOH extractions using UAE and vortexing for the assessment of reference concentrations for 12 SOA markers in an urban dust standard reference material (SRM 1649b, NIST). Similar concentrations were obtained with both extraction techniques, and the reference values determined in the study made it possible to evaluate the efficiency of the extractions in a later study.²⁴ Mutzel *et al.*⁹⁸ observed the formation of peroxides when samples were extracted with water and employing UAE due to the possible formed reactive radicals. Compared to mechanically extracted samples, reduced concentrations of terpene-derived organic acids were observed in the UAE extracts, which was either caused by reactions with peroxides formed or by the decomposition of acids under ultrasound treatment. Similar formations of reactive radicals can occur not only in water but also in organic solvents and their aqueous mixtures.⁹⁹

3.1.3. Concentration procedures. Concentration is typically the next phase after the extraction step to achieve more sensitivity for the analytical method. By looking at the data in Tables 1 and 2, extracts have typically been concentrated or evaporated to near dryness using a rotary evaporator or using a gentle stream of nitrogen. These methods possess a high risk of sample loss due to the evaporation of compounds, which can be eliminated in the analysis of organic acids by ionizing them with the addition of sodium hydroxide or potassium hydroxide.^{61,86,100} Multi-step derivatization methods for GC analysis⁹⁵ have been developed specifically to avoid concentration and additional extraction steps between the derivatization steps. Derivatization is typically initiated by first converting the carbonyls into methyl oximes, followed by methylation of organic acids and silylation of hydroxyl compounds. This method has been adapted for field samples,^{34,87,101} but interfering by-product formation has been reported.⁸⁷

Microextraction techniques have been useful for concentrating the analytes and avoiding the aforementioned methods and their drawbacks. Naing and Lee⁶⁷ employed a vortex-assisted micro solid phase extraction (VA- μ SPE) technique to determine glyoxal and methylglyoxal. The μ SPE device was prepared by enclosing the C18 adsorbent in a polypropylene envelope, after which the device was immersed in a 2,4-dinitrophenylhydrazine (DNPH) solution. The device was placed in the aqueous extract from which the compounds were on-sorbent derivatized and extracted simultaneously under stirring. The compounds were detected in the extracts at concentrations of 0.14–0.15 $\mu\text{g L}^{-1}$ using LC with diode array detection (DAD).

Müller-Tautges *et al.*⁶⁸ employed stir bar sorptive extraction (SBSE) to analyze compounds deposited from the atmosphere into snow. The compounds were derivatized to phenyl oximes in melted samples from which derivatives were enriched on a polydimethylsiloxane (PDMS) coated stirring rod. The compounds were desorbed into the solvent prior to LC-MS, providing limits of detection (LOD) of 0.21–0.24 $\mu\text{g L}^{-1}$ in snow samples.

Prieto-Blanco *et al.*⁶⁴ applied in-tube solid phase microextraction (IT-SPME) after the aqueous extraction of aldehydes, and conversion of the aldehydes to DNPH derivatives. In this column switching technique, the extract was applied to a GC column integrated into the injector of an LC. Analytes were concentrated and unretained impurities were washed away, after which the eluent was directed into the capillary to desorb the analytes for subsequent LC analysis. This method using capillary LC-MS demonstrated excellent sensitivity as LODs varied between 0.9 and 2.8 ng L^{-1} . However, the required configuration was rather complex and difficult to implement.

3.1.4. Derivatization techniques. The purpose of derivatization is to convert the desired compounds into a form from which they can be analyzed. By applying reactions specific for different functional groups, the selectivity of the method can be improved. SOAs contain compounds with, *e.g.*, hydroxyl, carbonyl, and carboxyl groups due to the atmospheric oxidation reactions. These are typical functional groups that can undergo



derivatization reactions. The most employed derivatization strategies, and how they provide new characteristics for identification and detection purposes are discussed below. Reagents for functional groups and analysis techniques are summarized in Table S1.†

3.1.4.1. Carbonyl compounds. A versatile range of different derivatization reagents for carbonyl compounds are available for analyses employing both GC and LC methods. In GC analyses, multistep derivatization applying silylation is usually initiated first by converting the carbonyl groups.^{34,86,87,100,101} This avoids the direct reaction of the carbonyl group and the silylation reagent, and the possible enolization of the carbonyl groups, which all could form artifacts that complicate and interfere with the analysis.^{102,103} Fig. 1 demonstrates possible artifact formation between an aldehyde with α -hydrogen and a common silylation reagent *N*-methyl-*N*-(trimethylsilyl)trifluoroacetamide (MSTFA); the added trimethyl silane group (TMS) could lead to misidentification as hydroxyl or carboxylic species. Oxaloacetic acid is shown to represent keto–enol-tautomerization which has been observed to occur in the atmosphere when analyzed in field samples.⁸⁶ The enol form produces a conjugated system spanning the whole molecule, which is why this form occurs. An unknown aldehyde in Fig. 1 can also produce such an unpredicted (in this case, unlike oxaloacetic acid) enol form and artifact with MSTFA.¹⁰³

Ketones and aldehydes can be specifically converted to oximes, where the oxygen atoms of the group have a hydrophobic substituent depending on the reagent used. Possible reagents are *O*-methylhydroxylamine (MHA),^{34,87,89,95,101} *O*-benzylhydroxylamine (BHA),^{66,68,90} *O*-*tert*-butylhydroxylamine,^{104,105} and *O*-(2,3,4,5,6-pentafluorobenzyl)hydroxylamine (PFBHA).^{82,106} Oxime derivatives have been mainly applied for GC analyses, but they can also be employed in LC.⁶⁸ Oxime derivatization can be performed in an organic solvent (MeOH, ACN, DCM, and their mixtures) or in an aqueous matrix.^{68,82,95,101,104} On-filter derivatization has been applied on filters impregnated with BHA for sampling of gaseous carbonyl compounds in the atmosphere.⁶⁶ Reactions are slow and

a minimum of 20 hours has been required for the reactions to proceed at room temperature.^{66,68,82} The reactions are even slower with ketones, and to achieve quantitative reaction, heating the reaction mixture at 60 °C (or higher) for two hours using MHA as a reagent has been suggested.⁹⁵ Quantitative reaction has been achieved by heating at 70 °C for one hour,¹⁰¹ but recoveries have been about 70% or less for carbonyl compounds in field samples.⁸⁷ Bulky substituent containing oxime derivatives provides more specific fragmentation in mass spectra but the drawback is that they are capable of forming only mono derivatives for alpha, beta-diketones due to the steric hindrance.^{82,104} Compounds containing more carbonyl groups could be better analyzed by GC-MS applying alkyl oximes, providing smaller derivative molecular masses compared to PFBHA.¹⁰⁴

Asymmetric carbonyl compounds yield *E* and *Z* isomers of oximes which often appear in chromatograms as multiple peaks possessing similar electron impact (EI) mass spectra.¹⁰⁴ An asymmetric ketone containing one carbonyl group has been observed to produce two peaks, of which the first eluting is clearly smaller than the latter, while the case is the opposite considering aldehydes.⁹⁵ The exact number of carbonyl groups cannot necessarily be deduced from the number of peaks, as two to four different peaks have been observed for dicarbonyls.¹⁰⁵ Accurate determination would require complete separation of all isomers. Possible molecular ions appear best in the EI mass spectrum with aromatic analytes or when employing an aromatic oxime reagent, otherwise the spectral identification has been based on ion fragments and ions corresponding to the alkoxy and alkyl groups of the oxime group.^{90,95,101,104,105} The possible involvement of the nitrogen atom of the oxime group in the analogous McLafferty rearrangement has been proposed to explain other specific peaks in the mass spectrum.^{95,101} Loss of an NO group $[M-30]^+$ from the molecule derivatized with PFBHA has been observed.⁸²

In addition to oxime derivatives, acetals and ketals have also been formed from carbonyls for GC analyses. The conversion



Fig. 1 Possible artifact formation between an aldehyde and MSTFA, as well as keto–enol tautomerization of oxaloacetic acid. MSTFA artifact reproduced from ref. 103 with permission from Elsevier, copyright 1999.



has been executed in *n*-butanol solution containing 14% of boron trifluoride by heating at 100 °C for 1 hour.^{62,86,100} This strategy has the advantage that organic acids are simultaneously converted into butyl esters (discussed later). Higher incubation temperature is crucial as no dibutoxy derivatives have been observed to form at 65 °C.¹⁰² Dried solvents are recommended to avoid the formation of hemiacetals which produce an extra hydroxy moiety and misidentification of the compound.¹⁰¹ Derivatives produce fragmentations in the EI spectrum corresponding to the cleavage of butene and the butoxy group.¹⁰⁰

DNPH mentioned earlier is a common derivatization reagent specific for carbonyls. The hydrazone derivative formed by DNPH is generally separated with LC and detected by UV at 360 nm or by MS with electrospray ionization (ESI) at negative polarity.^{64,67,83} Analysis with GC is possible, but it would require optimization of temperatures to prevent thermal degradation and the isolation of derivatives from the reaction mixture containing the acid and unreacted reagent with *e.g.*, cation exchange SPE.^{71,107} Under strongly basic conditions, the hydrazone derivative forms a quinoid structure with strong absorption at 480 nm.^{83,108} Conditions have been assessed to be established only by the addition of a strong base (pH over 12), which is why this has not been applicable in LC techniques restricted by the durability of the column materials.⁸³ Aimanant and Ziemann¹⁰⁸ applied this higher pH in a spectrophotometric method (discussed later) to determine the total concentration of carbonyl compounds obtained in a chamber experiment after ozonolysis of alpha-pinene. The reaction with DNPH could be carried out like for oxime derivatization in an aqueous matrix in which the reaction could be completed in just a few minutes at room temperature.^{64,67} Compared to the oxime reaction, it is very fast, which is why DNPH derivatization should be considered for LC analyses. Girard's reagent T (trimethyl amino acetohydrazide chloride) is also a possible alternative to convert carbonyls into hydrazone derivatives. By introducing this functional group into the molecule, detection is possible either by UV absorption or by MS employing positive polarity.^{46,109}

3.1.4.2. Carboxylic acids. Alkylation of carboxylic acids (*i.e.* esterification) is a common derivatization method for GC analysis. Without using highly toxic reagents such as diazomethane,⁹⁵ esterification is often done by acid catalysis of samples dissolved in an alcohol. The most common acid is boron trifluoride, with MeOH or butanol as alkylating alcohols.^{34,86,87,100,101} Boron trichloride is a very similar catalyst that can be applied as an alternative.¹⁰² Silylation (discussed later) has also been applied to derivatize carboxyl groups, but esterification has been preferred in compound identification to distinguish between carboxyl and hydroxyl groups.^{86,87,95,100,101} For the targeted analysis, organic acids have been silylated reducing the sample pretreatment steps.^{9,27,36} These derivatives produce larger peaks compared to methyl esters, which enhances the sensitivity of the method.⁹⁵ In addition to improving the sensitivity, silylation has been shown to be more reproducible than esterification.¹¹⁰ This derivatization strategy may not be recommended for small acids, such as oxalic acid, as

sample loss may occur in the analysis when the resulting derivatives are highly volatile.⁴⁵

Butanol has been more popular in acid-catalyzed esterification reactions because butyl esters are less volatile. The methyl esters of smaller acids, such as glycolic acid and pyruvic acid, evaporate easily after esterification and in the following extractions.¹⁰² Oxalic acid converted to dibutyl esters have been determined in yields of 75% or better^{62,86} and they separate better from the solvent peak.¹⁰¹ Quantitative esterification in butanol containing 14% of boron trifluoride has typically been achieved by heating at 100 °C for 30–60 minutes.^{62,86,100,111} After the reaction, the mixture is neutralized by adding a solvent mixture containing water, the butyl esters are extracted into *n*-hexane, followed by drying. Extracts are analyzed by GC or silylation is performed first to derivatize the remaining free hydroxyl groups. Esterification alone has been found to be inadequate for acids with two or more hydroxyl groups due to their high polarity.^{101,102}

As an alternative method, trimethylsilyl diazomethane (TMSD) has been applied to methylate acids in the presence of MeOH.^{34,87,89,95,101} The reagent acts as a silylating agent if the amount of MeOH used in the reaction is too low.⁹⁵ The reaction has been carried out after the addition of the reagents in an ultrasonic bath, where a 20-minute hold was sufficient. The advantage of using TMSD is that it does not require neutralization, thereby eliminating the need for extraction and possible evaporation of methyl esters for the next derivatization step, clearly reducing the risk of sample loss.

Ion fragmentation of the esters' alkoxy groups ($[M-31]^+$ and $[M-73]^+$) and corresponding alcohols from the molecular ion have been typically observed in EI-spectra.^{95,100,102} The ions formed in these processes possibly fragment further with the loss of carbon monoxide $[-28]$. Possible cleavage of butene $[-56]$ by McLafferty rearrangement has been evaluated with butyl esters.¹⁰⁰ Another typical cleavage with esters is alpha cleavage ($[M-59]^+$ and $[M-101]^+$) and beta cleavage ($[M-73]^+$ and $[M-115]^+$). Gowda *et al.*¹⁰⁰ identified 3-hydroxyglutaric acid and 3-hydroxyadipic acid by beta cleavage, which distinguished acids from their 2-hydroxy isomers. The 74 *m/z* ion characteristic of long-chain methyl esters of mono- and dicarboxylic acids is formed by the McLafferty rearrangement.⁹⁵ With methyl esters, in addition to the protonated molecular ion, the formation of a corresponding dimer has been observed using chemical ionization (CI) with methane.¹⁰²

3.1.4.3. Hydroxyl compounds. Silylation is a derivatization method in which the active hydrogen attached to an electronegative atom is typically replaced with a TMS group. As mentioned in the section discussing the derivatization of carboxylic acids, hydroxyl groups have been derivatized by silylation when oxidized species have been analyzed in aerosol samples by GC methods. 4-(Dimethylamino)benzoyl chloride¹¹² and 4-nitrobenzoyl chloride (4-NBC)⁸³ have been proposed as alternatives to derivatize hydroxyl moieties in SOAs for LC separations. However, according to our knowledge, these have not (yet) been employed in real field studies.

Various silylation reagents are available and among these, *N,O*-bis(trimethylsilyl) trifluoroacetamide (BSTFA) is a powerful



reagent that reacts with many types of hydroxyl groups.¹¹³ BSTFA has been the most common reagent for silylation with 1% of trimethylchlorosilane (TMCS) added as a catalyst.^{5,27,36,66,86,87,101} It is important to note that BSTFA is not capable of converting sterically hindered tertiary hydroxyl groups in, *e.g.*, citric acid and tertiary alcohols.⁹⁵ Another employed reagent is MSTFA,^{24,113} which is similar to BSTFA in its silylation properties.⁹⁵ The use of *N,O*-bis(trimethylsilyl)acetamide has been avoided due to its tendency to easily produce undesired artifacts.^{95,103}

Silylation requires careful optimization of several factors including the properties and amount of the reagents and solvents, temperatures, and reaction times.^{95,103} The derivatives react with water and protic polar solvents, which is why drying and solvent exchange is always performed prior to derivatization. The derivatization involves evaporation of the sample extracts to dryness, followed by the addition of a silylation reagent and pyridine as catalyst.^{9,34,82} The reaction has been accelerated by heating at 70–80 °C for 30–60 minutes.^{27,36,82,86} Silylation is possible to perform in the gas phase, which could occur during the injection into a hot GC injector.¹⁰³

In EI spectra, the silylated compound generally has a high abundance of ion *m/z* 73, corresponding to the cleaved TMS group.^{90,101} Ion *m/z* 75 is also observed presenting the cleaved trimethyl siloxane group (TMSO) ion lacking a methyl group. If the molecular ion is not present, the molecular weight could possibly be assessed from the largest ion presenting the derivatized molecule that has lost a methyl from the TMS group [M-15]⁺.¹⁰⁰ In addition, the corresponding molecular ion [M-89]⁺ can be observed resulting from the cleavage of the entire TMSO group.⁸² The ion *m/z* 147 provides an indication of two silylated groups, which has been observed for C2–C6 dicarboxylic acids and multifunctional acids such as tartaric acid.^{82,95,100,111} The ion is assumed to be formed when the TMS groups are in close proximity to the molecule.⁹⁵ A fragment ion *m/z* 103 is a possible indication of a primary alcohol due to the loss of TMSO and a methylene group.^{90,95} If carboxylic acids were silylated, a fragment presenting alpha cleavage [M-117]⁺ can distinguish alcohols from acids.³⁶

3.1.5. Columns. A non-polar DB5 or corresponding column containing 5% phenyl and 95% PDMS has by far been the most popular column in GC configurations for compounds that have been derivatized to reduce their polarity.^{27,34,36,66,86} Flores and Doskey⁸⁹ used the more polar DB-210 column containing 50% trifluoropropyl and 50% PDMS as the first dimension column for multidimensional gas chromatography. To analyze SOA precursors and their oxidized markers, the most optimal separation was obtained with two additional columns, as 90% cyanopropyl polyphenylene column being the second column followed by polyethylene glycol-based (SolGel-WAX) column. Because the sample pretreatment involved silylation, the GC injector had an additional bed of Carbowax 20M to prevent unreacted reagent from entering the column system and reacting with the WAX material in the second-dimension column. Comprehensive two-dimensional chromatography using DB-210 and Sol-Gel WAX columns was demonstrated to provide efficient separation, which has been applied in later

studies analyzing a broad range of hydrocarbon precursors and their oxidation products at concentrations as low as pg m⁻³.^{87,101}

In LC methods, it is possible to employ a broad range of columns according to the analytes' properties. In reversed-phase liquid chromatography (RPLC), the most common column material is octadecyl (C18) or octyl silane for the separation of especially terpene-derived acids, their dimers, and DNPH-derived carbonyl compounds.^{30,67,68,114} A column containing phenyl groups has been applied for the analysis of aromatic compounds, such as nitroaromatic compounds.⁵¹ Hydrophilic interaction liquid chromatography (HILIC) columns have proven useful for more polar compounds such as saccharides, organosulfates, and organic acids including terpene derivatives, dicarboxylic acids and smaller multifunctional acids.^{39,63,85} Isomers of 2-MEs and 2-MEOS have been separated particularly well with HILIC, whereas separation was not possible with RPLC.³⁸ The smallest and highly polar organic acids, on the other hand, have been analyzed by ion chromatography (IC). Kwiezinski *et al.*² employed an IC-MS method employing an anion exchange column to determine oxalic acid, glycolic acid, malic acid, and citric acid with LODs of 2.5–10 µg L⁻¹ and more hydrophobic acids with five and ten times lower detection limits. Carbohydrates, 2-MEs and other sugar alcohols have been analyzed by high performance anion exchange chromatography (HPAEC).^{115,116}

3.1.6. Analyzers and detectors. The data in Tables 1 and 2 clearly show that the most common detector for GC and LC instrumentation is MS providing excellent sensitivity, specificity, and structural information about the analytes. Flores and Doskey¹⁰¹ observed LODs of 0.3–54 pg m⁻³ for compounds containing carbonyls, carboxyl groups, and hydroxyl groups with a sample volume of 35 m³, analyzed by multidimensional GC analysis after multi-step derivatization. With a similar derivatization procedure and GC-MS analysis, Kanellopoulos *et al.*³⁴ reported LODs between 0.16 and 0.62 ng m⁻³ for various organic acids and SOA markers of monoterpenes and isoprene, with sample collection for 24 hours at a flow rate of 2.3 m³ h⁻¹. LC-MS methods have given LOD values from extracts between 0.1 and 10 ng mL⁻¹ for organic acids and 0.075–4 ng mL⁻¹ for organosulfate compounds.^{2,38} LODs of 1–25 ng mL⁻¹ have been obtained for carbohydrates by LC-MS.³⁰

In GC, EI is the most common ionization technique that provides information about the structure of compounds through characteristic ion patterns.^{27,36,86,100,101} The advantage of the softer CI is that the molecular ions are better visible in the spectra which helps to identify the compounds better.^{36,102} The most common ionization technique for LC has been ESI, which has been applied mostly in negative mode to detect non-derivatized sugars, organic acids, and organosulfates separated with RPLC, HILIC, and IC.^{2,30,38,64} Positive mode ESI is applicable for detection of carbohydrates.³⁰ Compared to other LC works, the IC-ESI-MS configuration employed by Kwiezinski *et al.*² required a separate suppressor and post-column addition of iso-propanol to avoid non-volatile salt stacking in the ion source and to increase the sensitivity, as the eluent was an aqueous solution of potassium hydroxide. With CI employing ammonia and isobutane, it has been possible to detect DNPH



derivatives of carbonyls, hydroxyl group compounds derived with 4-nitrobenzoyl chloride, and organic acids derived with 2-nitrophenylhydrazine hydrochloride using positive polarity.⁸³

A simple quadrupole mass analyzer has been commonly employed in low resolution mass spectrometers.^{27,64,66,86} An ion trap has also been employed as another low resolution mass analyzer.^{9,36,68} For accurate identifications of compounds, HR-MS is required to determine the elemental composition or to provide more selective detection of analytes. For those purposes, the most commonly employed HR-MS analyzer has been Orbitrap^{2,36} or time of flight (TOF) which has been employed as alone or in combination with a quadrupole (Q-TOF).^{38,86,87,100,101} This tandem mass spectrometry (MS/MS) has also been established by coupling three quadrupoles.³⁰ MS/MS has also been referred for a single ion-trap analyzer that collides isolated ions with, *e.g.*, helium to provide collisional induced dissociation and multiple reaction monitoring.⁶⁸ In addition to quantitative analysis, MS/MS has been employed to identify compounds by applying neutral loss mode. For example, loss of water [M-18] and decarboxylation [M-44] have been monitored to identify carboxylic acids tentatively.^{117,118} With the same principle, other compounds with certain functional groups have been identified, such as organic hydroperoxides and peroxy acids, which could be identified by their characteristic loss of hydrogen peroxide [M-34].¹¹⁹

Detectors other than MS have been employed but to a lesser extent. Flame ionization detection (FID) in combination with GC has been employed for quantification of compounds when the qualitative information has been acquired by MS.^{86,100} FID is sensitive and has provided LODs of 5 pg m⁻³ for multifunctional organic acids using a high sample volume exceeding 7000 m³.⁸⁶ A pulsed amperometric detector has been employed in HPAEC to detect carbohydrates and 2-MEs at low LOD of 0.05 ng m⁻³.^{115,116} Ultraviolet-visible (UV-VIS) has been employed for more selective detection of DNPH-derivatized carbonyl compounds in LC.⁶⁷ The non-derivatized carbonyl or carboxyl group also acts as a chromophore, in which case UV detection can be applied also for compounds in their native form.⁶¹ However, UV detection is not so sensitive compared to MS

detection, and, *e.g.*, Prieto-Blanco *et al.*⁶⁴ obtained 30–100 times lower detection limits for DNPH-derived aldehydes compared to those obtained by IT-SPME-cLC-diode array detection (LODs ranging from 30 to 198 ng L⁻¹).¹²⁰

3.1.7. Quantitation using surrogates. Authentic standards required for accurate quantitative analysis are not always available or no synthetic methods are known, which is often the case when a wider group of compounds are analyzed. The response factors of surrogates (compounds similar to the analyzed compounds) are applied in this case to assess the concentration of the analyzed compounds. Table 3 shows different compounds and surrogates used for their quantitation. The concentrations obtained with surrogates are more likely to be semi-quantitative, as the responses produced by surrogates and detected analytes are likely to be slightly different despite their similarity.¹²¹ Kenseth *et al.*¹¹⁴ demonstrated this by determining the ionization efficiencies for synthesized *cis*-pinonic acid, pinic acid, pinolic acid, and their dimer esters employing LC(-)ESI-MS. In the comparison between the acids, their efficiencies were observed to vary by a factor of six. Compared to *cis*-pinonic acid, the ionization efficiencies of dimer esters were 19–36 times higher, which led to overestimated concentrations when *cis*-pinonic acid was applied as a surrogate. Haque *et al.*²⁷ determined varying uncertainties with the surrogates applied (shown in Table 3) when the compounds were silylated before GC-EI-MS. 2-MGA, 2-MEs, and 3-MBTCA had lower uncertainties ranging between 10 and 32%. Larger uncertainties (85 to 136%) were obtained for C5-alkene triols and beta-caryophyllinic acid.

3.2. Ion mobility mass spectrometry

Ion mobility spectrometry (IMS) is an alternative separation technique integrated with MS for the analyses of atmospheric compounds. The separation efficiency with IMS is not that great as compared to chromatography, but the selectivity of the ionization technique used can be of great help for complex samples. Generally, analytes are ionized and separated by the electric field and opposing drift gas flow based on their ion

Table 3 Selected examples of compounds separated by GC and surrogates applied for quantitation

Analytes	Applied surrogate	Ref.
Methyltheronic acids	Citramalic acid	36
2-Methyltartaric acids + 3-hydroxyglutaric acid	Tartaric acid	
2-Methylglyceric acid	Pentanedioic acid	
3-Carboxyheptanedioic acid	<i>d</i> -Arabitol	
2-MEs	<i>d</i> -Threitol	
Aliphatic monocarboxylic acids	Dodecanoic acid	87 and 101
Aliphatic dicarboxylic acids	Glutaric acid, adipic acid, azelaic acid	
Ketocarboxylic acids	<i>Cis</i> -pinonic acid	
Aromatic carboxylic acids	<i>p</i> -Toluic acid	
Carbonyls	Nopinone	
Alcohols and phenols	Hexadecanol and <i>p</i> -hydroxybenzoic acid	
Carbohydrates and glycolics (polyols)	Levoglucosan and <i>meso</i> -erythritol	
Hydroxycarboxylic acids	Malic acid	
2-MGA, 2-MEs, C5-alkene triols	<i>meso</i> -Erythritol	27
3-MBTCA, beta-caryophyllinic acid	Pinic acid	



mobility, which is highly dependent on the structure of the ion (mass, charge, and collision cross section) and its interactions with the added drift gas. In recent publications, the method has especially been applied to the separation of isomeric compounds. The methodology is applicable to online analyses of gaseous compounds and aerosol samples extracted into suitable solvents. Analysis of aerosol extracts without chromatographic separation can be challenging as other ions contained in the matrix may cause ion suppression and interfere with the separation by forming additional clusters.^{122–124}

The early application of IMS-MS in atmospheric research was reported by Krechmer *et al.*¹²² for the online analysis of a nitrate-rich gas phase with CI-IMS-TOF. The field measurements provided detection of BVOC oxidation products, agreeing well with previous results obtained with nitrate CI-MS methods (which will be discussed later in the section regarding HOM detection). The evaluation was focused on monoterpene-derived HOMs, of which monomeric species were efficiently detected. However, larger dimers were not detected, most probably due to poor ion transmission and sample losses due to interactions

with the walls of the instrument. To analyze 2-MEOS isomers, aerosol phase samples were extracted and analyzed by ESI. Isomers of 2-MEOS were separated providing information on isomer occurrence, which depended on the isomeric form of the IEPOX intermediate reacting with sulfuric acid. The separation efficiency was observed to decrease, probably due to the extracted sulfuric acid which formed clusters with the analytes in IMS.

West *et al.*¹²³ employed ESI-IMS-Q-TOF to separate alpha-pinene and limonene SOA oxidation products possessing similar mass spectra. Alpha-pinene products possessing cyclobutane rings were successfully separated from isomeric noncyclic limonene products. The collision cross sections in the employed nitrogen gas were demonstrated to be dependent on the ionization mode, showing the formation of positive sodium adducts being favorable for monomeric products and negative deprotonation for dimeric and trimeric products. Iinuma *et al.*¹²⁵ characterized alpha-pinene-derived dimer esters by LC-ESI-IMS-TOF/TOF employing collision-induced dissociation to identify their acid components. Isomeric *cis*-norpinic acid and

Table 4 Selected examples of studies analyzing HOMs and other oxidized species^a with MS techniques

Detected compounds	Information about study	Technique	Ref.
C _{8–10} H _{12–18} O _{4–9} (monomers) C _{16–20} H _{24–36} O _{8–14} (dimers)	Alpha-pinene-derived HOMs generated in chamber experiments were analyzed in both gas (online) and particle (offline) phase	Online: CIMS with iodide ionization Offline: ESI-IMS-TOF with sodium adducts [M + Na] ⁺	133
C _{8–10} H _{12–16} O _{6–11} (gas and particle phase) C _{7–10} H _{8–14} O _{7–11} S (particle phase)	Alpha-pinene ozonolysis chamber experiments and field measurements in rural areas surrounded by forest and agricultural pasture dominated by alpha-pinene emissions. The study involved the detection of HOMs in both phases including discoveries of highly oxidized organosulfate compounds and carbonyl functionalities in HOMs after DNPH derivatization	CI-API-TOF with NO ₃ [−] ionization Particle phase samples were extracted and analyzed directly or with DNPH derivatization by LC(−)ESI-TOF or UHPLC(−)IMS-QTOF technique	72
C _{3–7} H _{5–9} O _{6–7} N _{0–1} (lighter HOMs) C _{9–10} H _{14–16} O _{7–13} N _{0–2} (monomers) C _{16–20} H _{28–32} O _{9–19} N _{0–2} (dimers)	Field measurements in boreal forest area where monoterpenes were dominating BVOCs. HOMs formation and occurrence in daytime and night-time with different oxidants presents was assessed	CI-API-TOF with NO ₃ [−] ionization	79
C _{5–20} H _{6–30} O _{4–18} (monoterpenes) ^b C _{3–20} H _{5–31} N _{1–2} O _{4–14} (monoterpenes) ^b C _{4–15} H _{6–20} O _{5–22} (isoprene) ^b C _{4–15} H _{6–11} N _{1–3} O _{5–11} (isoprene) ^b	Field measurements in forest area providing possible markers for isoprene and monoterpene	CI-API-TOF with NO ₃ [−] ionization	80
C _{6–20} H _{11–38} O _{4–14} ^b C ₁₀ H ₂₁ O _{1–4} NO ₂ C ₁₀ H ₁₇ O _{1–8} NO ₂ C ₆ H ₁₁ O _{2–5} NO ₂	Oxidized carbonyl species, peroxy radicals and NO ₂ species produced from alkanes, alcohols, and carbonyls in chamber experiments representing conditions in the atmosphere and combustion engine	CI-API-TOF with NO ₃ [−] and ethylamine ionization	77

^a Wang *et al.*⁷⁷ do not classify compounds as HOMs if possessing less than six oxygen atoms and/or formed under conditions representing a combustion engine. ^b Monomers and dimers are included.



terpenylic acid did not separate in IMS using nitrogen as the buffer gas; however, the separation was successful switching into more polar carbon dioxide. Zhang *et al.*¹²⁶ studied alkyl nitrates using ESI, with a focus on the optimal addition of anions to the extracts for adduct formation. Chloride and nitrate were found to form clusters most efficiently. For identification, a specific nitrate fragment was determined by collision-induced dissociation analysis, regardless of the adduct used. The correlation of the calculated collision cross section and the mass-charge ratio was also applied to the identification, as the ratio was found to behave differently compared to organic acids or organosulfate compounds.

3.3. Mass spectrometric detection of highly oxidized organic molecules

Oxidized species have been analyzed by MS without separation prior to detection. The analysis in this case is typically for compounds present in the gas phase, but also extracts of samples taken on filters can be analyzed by MS alone. No separate sampling is required for the gas phase compounds, so the measurements have been done online in real time. Emphasis has been on the examination of HOM formation in chamber experiments or their occurrence in field samples. Selected studies focusing on HOM compounds and other oxidized species are presented in Table 4.

For the time being, HOM analytics mainly rely on MS methods where they are analyzed from the gas phase since the reactivity of their supposed peroxide groups makes them unstable and causes analytical problems.^{73,76} Peroxide compounds are challenging as they may degrade in SOAs within

a few hours.¹²⁷ Analytics are also limited because the exact structures of HOM compounds are still unknown.⁷⁶ Mutzel *et al.*⁷² have shown by LC-MS, employing DNPH derivatization, that HOMs found in the gas phase occurred also in the particle phase, and they contained carbonyl groups. Zhao *et al.*¹²⁸ applied iodometry (discussed later) to LC-ESI-MS, in which samples were qualitatively analyzed for peroxide functionalities containing species before and after iodometry. The spectra obtained were compared, in which the peaks of certain compounds had disappeared after treatment, indicating that the compound had a peroxide group. Peroxide compounds have also been successfully analyzed from the ozonolysis-generated alpha-pinene SOAs using LC, where the separated peroxides dimerized *p*-hydroxyphenylacetic acid in post-column derivatization followed by detection with a fluorescence detector.^{129,130} Despite successful peroxide compound detection with LC techniques, the use of chromatographic methods for HOM analyses is challenging, because HOMs are most likely too polar to be all separated and their original structures could change during the separation process by reacting with column materials.⁷⁶

3.3.1. Instrumentation. Atmospheric pressure interface time-of-flight (API-TOF) mass spectrometry¹³¹ has been typically employed in MS analyses of HOMs. This configuration alone has no ion source meaning that only spontaneously charged compounds can be analyzed with this technique. Early detections of HOMs were performed by this technique.^{74,132} An ion source typically employing CI has been integrated with the API-TOF instrumentation to analyze neutral species. The most common studies on HOM compounds have employed this CI-

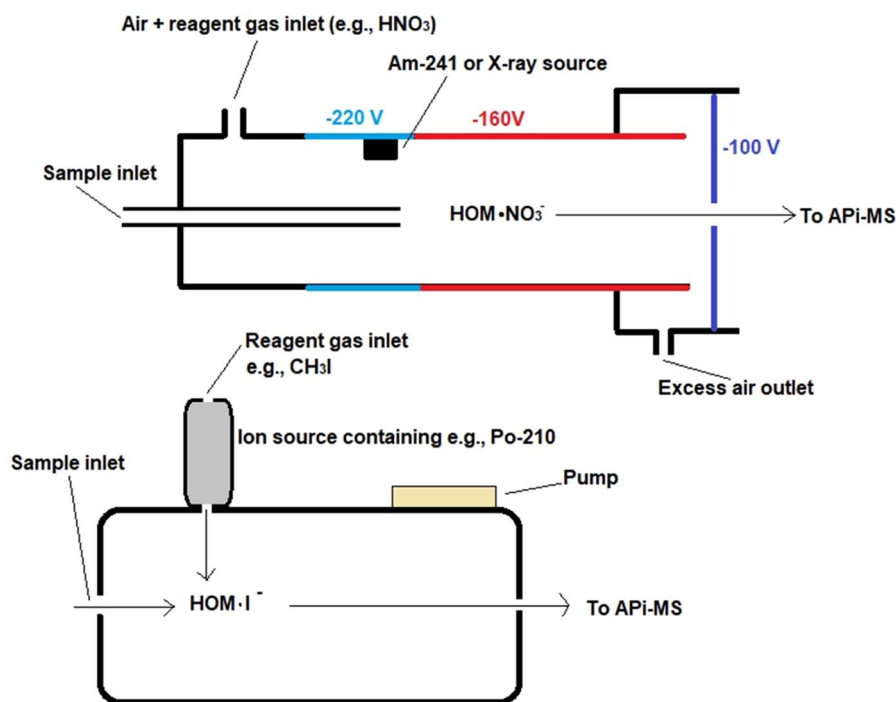


Fig. 2 Schematic structure of an Eisele-type ion source, reproduced from ref. 137 with permission from National Academy of Sciences, copyright 2014. An IMR ion source, reproduced from ref. 139 with permission from Copernicus Publications, copyright 2011.



API-TOF technique, sometimes referred to as chemical ionization mass spectrometry (CIMS or TOF-CIMS) due to the lack of consistency using uniform terms regarding different CI techniques.⁷³ Alternatively, the ESI-source alone or interfaced with IMS has also been employed for ionization of HOMs.^{72,133}

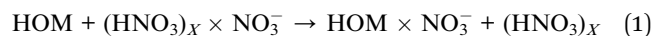
There are differences in the configurations of ion sources when employing different reagent gases, as presented in Fig. 2. The ion source of the CI-API-TOF instrumentation is typically the so-called Eisele type ion source, which is practically a flow tube model source where, in addition to the sample, pure air and reagent gas are supplied to the source.^{73,134} Reactive gas species are formed from the reagent gas with the aid of a corona needle or a radiation source, such as an alpha-emitting Am-241 or an X-ray source.^{135–137} Voltages are applied to direct ionized analytes towards the mass analyzer system. In the design of ion source, it has been important to pay particular attention to sample losses of oxidized compounds due to the adsorption on the walls and inlets as compounds have lowered vapor pressures. To eliminate this, high sample flows are applied in the ion source and the structure of the ion source is designed to practically have no walls.⁷⁵ The configuration employing iodine or acetate, in turn, has an ion-molecule reaction chamber (IMR) as an ion source, in which case instruments have been called CI-TOFMS or TOF-CIMS instead of CI-API-TOF.^{138,139} This source type also has some radioactive material to generate the reagent gas, but unlike the Eisele type ion source, a reduced pressure is applied in the ionization chamber and the supply of reagent gas to the chamber is orthogonal to the sample flow.

Proton transfer reaction mass spectrometry (PTR-MS), which can be classified as a CI technique, is optimal only for volatile compounds, as found by Riva *et al.*,¹³⁸ who compared five different CI techniques to detect about a thousand compounds formed in the ozonolysis of alpha-pinene. The comparison also included the so-called Vocus instrument, which has a discharge reagent-ion source and a focusing ion-molecule reactor compared to the traditional PTR-MS configuration.¹⁴⁰ The Vocus instrument has faster response times which enables faster analyses, decreasing the loss of SVOC on the walls and inlets. This was verified as the Vocus instrument was able to detect SVOC with an oxygen content ranging from three to five with good agreement with the iodine-CI technique included in the comparison. Li *et al.*¹⁴¹ found that the Vocus instrument is still poorly suited for the detection of peroxide compounds, most likely due to the fragmentation of peroxides in the PTR process.

In API-TOF, the ionized species entering the API interface are directed by quadrupoles and ionic lens systems before entering the mass analyzer. TOF is clearly the most commonly used mass analyzer for studies of HOMs.^{72,79,80,133} As this analyzer can provide high mass resolution, exact elemental compositions of HOMs can be assessed and compounds distinguished from each other without any separation technique. Riva *et al.*¹⁴² compared CI-Orbitrap with CI-API-TOF employing acetate and *n*-propylamine as reagent gases in both systems to detect peroxide radicals. Both mass analyzers gave results with good agreement regardless of the reagent gas, but the responses in the Orbitrap were observed to have poor linearity at the lowest concentrations.

3.3.2. Reagent gases employed in chemical ionization.

Nitric acid was the reagent gas applied in the first CI-API-TOF application.¹³⁵ After this, CI-API-TOF employing nitrate has become a very common method as HOMs and their peroxide precursors have been observed to form strong and specific clusters with nitrate ions.⁷³ This is believed to be based on the two hydroperoxide groups contained in the HOM compounds, which results in the formation of strong hydrogen bonds with nitrate.¹⁴³ Hydroxyl groups (as hydrogen bond donors) are also capable of forming similar specific nitrate cluster. The instrumentation contains generally the Eisele-type ion source in which a reactive gaseous cluster is formed with nitric acid and nitrate ions (eqn (1)).^{135,136} In addition to good selectivity towards HOMs containing hydroperoxide and hydroxyl groups, nitrate-CI-API-TOF could detect low concentrations, even less than 10^5 molecules in cm^{-3} .¹³⁸



The iodide adducts have also been studied, as iodide has been found to ionize easily oxidized and acidic compounds.¹⁴⁴ The IMR source typically utilizes methyl iodide and a Po-210 alpha radiation source to form a reactive gas cluster $\text{I}(\text{H}_2\text{O})^-$ with water.^{138,144} One of the benefits using iodide adducts is the dehydroxylation of hydroperoxides observed in the mass spectra, which could be used as a qualitative tool for identifying the compounds.¹⁴⁵ The iodide CI methodology is optimal for less oxidized semi-volatile organic compounds (SVOC), possessing oxygen atoms from three to five¹³⁸ or having the O : C ratio of about 0.6.¹⁴⁵

In addition to inorganic anions, adducts have also been formed with organic anions, such as acetate, pyruvate, and lactate.¹⁴⁶ The Eisele-type ion source employing corresponding acids as dopant gases have been employed, but the IMR ion source has also been applied to acetate.¹³⁹ Of the above anions, acetate has been the most commonly used in the studies of oxidized species. For example, Berndt *et al.*¹⁴⁷ compared nitrate and acetate ionization to study HOM compounds formed in the ozonolysis of cycloalkenes. Acetate was found to be more favorable for the analysis of compounds containing one hydroperoxide group as systematically lower concentrations were observed using nitrate. The study also demonstrated acetate's ability to form detectable ions by deprotonating analytes and forming adducts with them. Because of those two possible ionization mechanisms, acetate is not favorable for qualitative analysis as the interpretation of the mass spectrum becomes more complex.^{148,149}

The above-mentioned adducts have been detected with negative polarity, but oxidized compounds have also been analyzed with positive polarity. Methyl-, ethyl-, and *n*-propyl-amine as well as hydrazine have been employed to detect molecules containing peroxide radicals that form clusters with aminium and hydrazinium cations.¹⁵⁰ Riva *et al.*¹³⁸ compared different CI techniques and found that butylamine gave poorly correlated results to other CIMS techniques, which is thought to be due to reagent gas depletion, leading to non-linear responses. In addition to organic cationic adducts, the sodium adduct has been found to be particularly well suited for ionizing polyolic HOMs and those containing organic peroxides and ester groups.¹³³



3.3.3. Quantification of highly oxidized organic molecules.

Due to the lack of a more precise definition and structures for HOMs, it has not been possible to quantify them accurately.⁷³ Therefore, HOM total concentrations have only been determined semi-quantitatively by applying known concentrations of reagent gases employed in ionization and the calibration factor of the selected reference compound, assuming that the concentration of HOM compounds is significantly lower compared to the dopant gases.¹³⁵ This method is demonstrated in eqn (2) in which the signal of the compound detected HOM(NO_3^-) is related to the sum of the dopant gas concentrations and multiplied by the determined calibration factor C .^{72,79,135} The resulting concentration, which is typically in the unit of molecules in cm^{-3} , is not accurate and depends on the differences between the reference compound applied in the calibration factor and the analytes. The most typical reference compound in the nitrate CI method is sulfuric acid, as the clusters of HOMs and nitrates formed are assumed to be as stable as the corresponding clusters of sulfuric acid and nitrate.⁷⁵

$$[\text{HOM}] = C \times \frac{\text{HOM}(\text{NO}_3^-)}{\sum_{i=0}^2 (\text{NO}_3)_i(\text{NO}_3^-)} \quad (2)$$

3.4. Spectrophotometric applications

Spectrophotometry has been introduced as an analytical method to study SOA constituents. Compared to chromatographic and MS methods, spectrophotometric methods are far less popular, mainly due to the lower sensitivity.¹⁰⁸ Methods applied for SOA research provide only total concentrations of functional groups, not individual marker compounds. Selective functional group reactions have been applicable to LC methods, providing possible new alternatives for the derivatization reactions introduced above.⁸³ In addition to these methods, Fourier-transform infrared spectroscopy (FTIR) has also provided similar qualitative information about functional group compositions.¹⁵¹⁻¹⁵⁵

3.4.1. Determination of various functional groups.

Aimanant and Ziemann¹⁰⁸ determined spectrophotometrically oxidized species present in alpha-pinene ozonolysis products. Hydroxyl species were derivatized with 4-NBC, and carboxylic species with 2-NPH, and carboxylic species with 2-nitrophenylhydrazine (2-NPH) and N,N' -dicyclohexylcarbodiimide (DCC). 2-NPH derivatives form quinoidal structures under highly alkaline conditions as DNPH derivatives, which were employed in this application (Fig. 3). Esters were derivatized to the hydroxyamic form with hydroxylamine, followed by the addition of acidic ferric chloride solution to convert the derivative into a ferric hydroxamate form to be

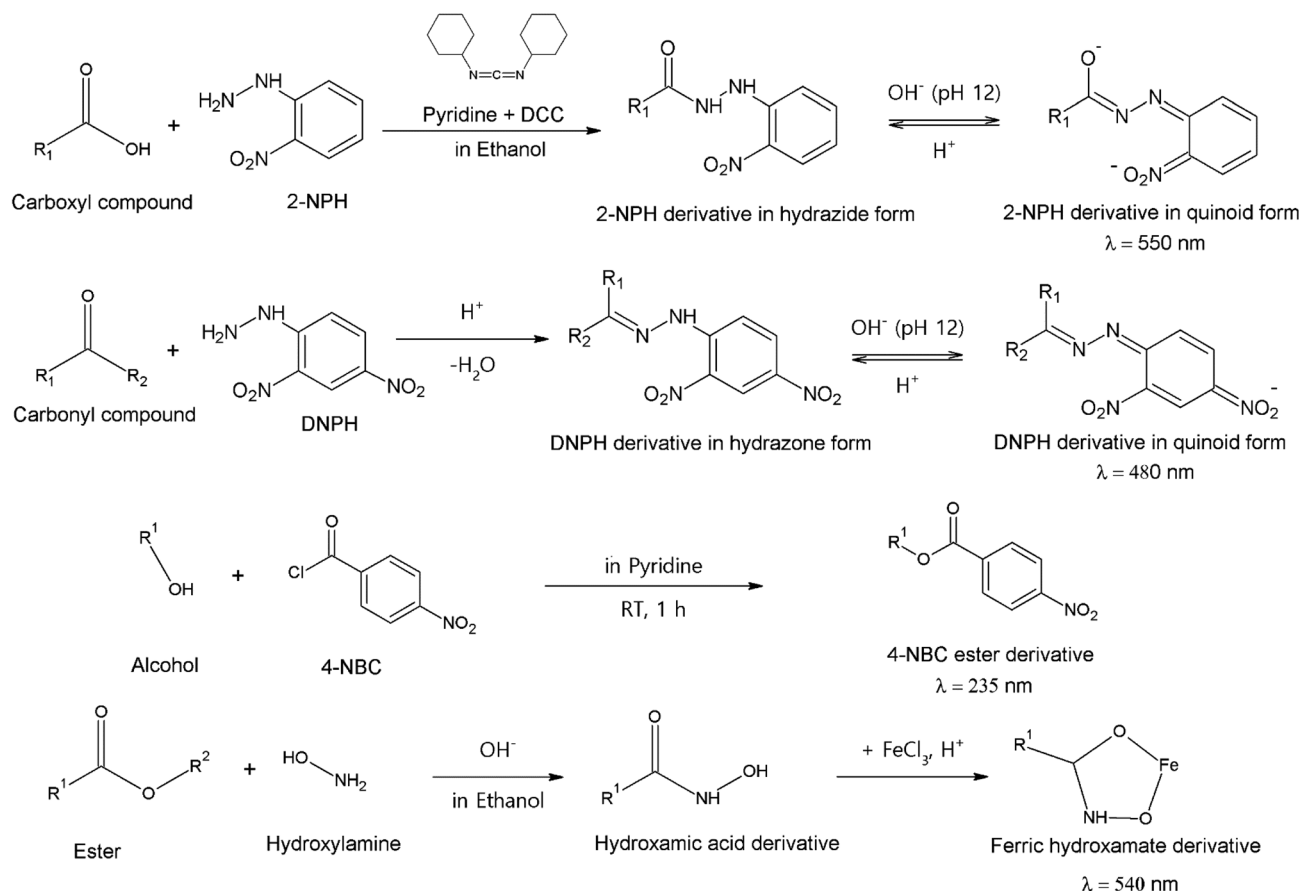


Fig. 3 Functional group-specific reactions and measurement wavelengths (λ) applied in spectrophotometric application. Reproduced from ref. 108 with permission from Taylor & Francis Group, copyright 2013.



measured. Peroxides were determined by iodometry and nitrate species were analyzed by LC-UV at 210 nm. The objective was to determine the functional group composition of laboratory-generated ozonolysis of alpha-pinene SOAs. Measured concentrations in moles were converted into masses by applying the groups' molecular weights and the total mass was subtracted from the weighed sample mass where the remaining mass represents carbon as methylene groups. Possible interferences, including increased background absorbance and decreased extraction efficiencies due to brown and black carbon, humic substances, and inorganic dust, were discussed in terms of applying this method to field samples.

This spectrophotometric method has also been applied in later studies analyzing functional group compositions obtained in chamber experiments.^{156,157} Recently, Clafin *et al.*¹⁵² have also applied this for real field samples. In that study, organonitrates were determined spectrophotometrically by measuring samples directly at 210 nm. The study also involved a comparison with the FTIR method in which functional groups were specifically detected from dried filters. The results provided justifications to determine the total concentrations of functional groups, as their abundance could indicate the origin of formed SOAs including precursors and oxidants.

Chamber test samples demonstrated the appearance of nitrates when SOAs were generated in the presence of NO_x or NO₃. Peroxide and methylene fractions increased as carboxylic, carbonyl, and ester species decreased when alpha-pinene oxidation involved OH instead of ozone. In reactions involving OH and NO_x, the ratio of methylene to nitrate of 1 indicated isoprene of being a precursor and the ratio of 2 suggested that monoterpene is the precursor. Based on observations from chamber experiments, monoterpenes and possibly isoprene were suggested precursors in field samples producing SOAs in the presence of OH and ozone at low NO_x concentration conditions. FTIR and spectrophotometry were found to provide comparable results when analyzing organonitrates, hydroxyl groups, and non-acid carbonyl groups. FTIR provided simpler and faster analyses, but spectral interferences and overlaps were the drawbacks encountered.

3.4.2. Iodometric applications. Organic peroxides are strong oxidants which have been utilized in the analysis of peroxides.¹⁵⁸ The iodometric reactions have good specificity for peroxides, and the reactions are fast enough for quantitative analysis applications, although the substituents attached to the peroxide groups have an effect on the speed of the reaction. For example, a *tert*-butyl group in the molecule has been demonstrated to slow down the reaction as the group causes steric hindrance and possibly an inductive effect that stabilizes the peroxide group.¹⁵⁹ The total amount of peroxides could be determined by spectrophotometry in which the absorption of the triiodide was measured at 420 or 470 nm.^{129,130,158,159} Mutzel *et al.*⁹⁸ optimized an iodometry-based spectrophotometric method for determining the amount of water-soluble peroxides from the alpha-pinene ozonolysis reaction mixture collected on a filter. Hydrogen peroxide was applied for calibration providing an excellent linear measurement area. The optimal pH value was 3 (adjusted with acetic acid). The amount of potassium

iodide was found to be critical, as too high or low amounts resulted in systematically higher or lower yields. By degassing the molecular oxygen for 5–15 minutes with pure nitrogen, the quantitative reaction time was 60 minutes.

Iodometry has been applied to peroxide using long path absorption photometry (LOPAP), which is a spectrophotometric technique employing iodometry introduced by Mertes *et al.*¹⁵⁹ The instrumentation is closed and pure nitrogen (5.0) is fed into the reaction chambers to eliminate the interference of oxygen. The pump inside dispenses solvents (ethanol and 0.1 M HCl in this application) into the reaction chamber, where the filter sample and solid potassium iodide have been added. After the reaction phase, the pump dispenses the sample through the filter into the long path detection chamber, called the liquid core waveguide (LCW). This is a thin capillary made of Teflon material with a refractive index lower than that of water and ethanol, which enables the light produced by the deuterium lamp to propagate in the capillary with total reflection. The absorbance is measured with a separate spectrophotometer integrated into the LOPAP instrument.

Mertes *et al.*¹⁵⁹ used a 50 cm long LCW achieving a better sensitivity according to the Lambert–Beer law. The method was assessed to be suitable for peroxides poorly soluble in water and ethanol, as the absorbances produced by benzoyl peroxide and lauroyl peroxide were assessed to be practically the same as the absorbances produced by water soluble hydrogen peroxide and peracetic acid. Analyzed smog chamber samples revealed the mass fraction of peroxides being 34% in fresh alpha-pinene ozonolysis SOAs and decreasing to 12% after remaining stable for 6 hours. Mass fractions decreased when NO_x concentration was increased, but a similar peroxide concentration time trend was observed. A molecular weight of 300 g mol⁻¹ was applied for calculations based on an estimated mixture of peroxyhemiacetal and minor hydroperoxide species forming in monoterpene SOAs generated with ozone.¹⁶⁰ Krapf *et al.*¹²⁷ have utilized this application for the determination of HOM yields from laboratory-produced alpha-pinene SOA. Assuming that HOM contains on average two hydroperoxide groups, the HOM content contributing to alpha-pinene SOA was estimated to be 5%. Also, the same above-mentioned peroxide concentration time trend was observed demonstrating high instability with half-lives shorter than an hour.

4. Conclusion and prospects

New oxidized species are constantly being discovered in the atmosphere. One of the challenges in analyzing SOA markers is the diversity and large number of compounds, which complicates the analysis. Analytical standards are not available and must be synthesized to determine concentrations more accurately for compounds and confirm their identity. In addition to using surrogates, the concentrations have been dependent on the analytical method used and there have been variations at sampling points and times. Finding new and more specific compounds would be important, especially for the assessment of aged SOAs. In the absence of specific markers, monitoring of precursor VOCs is recommended for the characterization of



SOAs. Chromatographic methods have been used to determine oxidized markers at a concentration level as low as pg m^{-3} . In GC methods, it has been possible to determine a more diverse range of compounds with a single analysis. Due to the higher separation efficiency, it has been possible to better separate the isomers of the compounds compared to LC methods and to numerically detect more compounds. The disadvantage is the need for derivatization of oxidized compounds, which makes sample processing more laborious and can create interfering artifacts without proper optimization. LC methods, on the other hand, have generally been used for the analysis of compounds directly from extracts or with fewer sample processing steps. MS, as an independent analyzer or integrated as a detector after chromatographic separation, is essential in modern research and provides good sensitivity and qualitative information of the compounds. Spectrophotometric methods are simple and can be used with cheaper equipment. Their sensitivity is lower and they do not provide information on individual compounds, which is why they have not been mainstream techniques in SOA research. Iodometric determination by spectrophotometry of peroxides from laboratory-generated SOAs has been more commonly used. HOM compounds are a relatively new discovery that has not yet been the subject of much research. There are no harmonized and unambiguous definitions, so for the time being exact criteria and definitions for HOMs should be presented in the research articles. HOMs are challenging to analyze by chromatographic techniques, and authentic standards for quantitative analysis are not yet available, so the analysis of HOMs is so far qualitative and semiquantitative using HR-MS methods and chemical ionization. The hydroperoxide and other peroxide groups in HOMs are highly reactive, which complicates the analysis of the compounds due to the poor stability of the peroxide compounds and changes in SOA samples as the peroxides react with the matrix. Currently, analysis of HOM compounds is focused on CI-API-TOF or corresponding methods, where different reagent gases with specificity to different functionalities could potentially provide new information on the structure of HOM compounds.

Author contributions

Aleksi Tiusanen: writing – original draft, conceptualization, investigation, visualization; Jose Ruiz-Jimenez: writing – review & editing, conceptualization, supervision; Kari Hartonen: writing – review & editing, conceptualization, supervision; Susanne Wiedmer: writing – review & editing, conceptualization, supervision.

Conflicts of interest

There are no conflicts of interest to declare.

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