Direct detection and quantification of malondialdehyde vapour in humid air using selected ion mass spectrometry supported by gas chromatography mass spectrometry

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Abstract

RATIONALE:
It has been proposed that malondialdehyde (MDA) reflects free oxygen-radical lipid peroxidation and can be useful as a biomarker to track this process. For the analysis of MDA molecules in humid air by selected ion flow tube mass spectrometry (SIFT-MS), the rate coefficients and the ion product distributions for the reactions of the SIFT-MS reagent ions with volatile MDA in the presence of water vapour are required.

METHODS:
The SIFT technique has been used to determine the rate coefficients and ion product distributions for the reaction of H3O+, NO+ and O2•− with gas phase MDA. Then, in support of SIFT-MS analysis of MDA, solid phase microextraction, SPME, coupled with gas chromatography mass spectrometry, GC/MS, has been used to confirm MDA identification.

RESULTS:
The primary product ions have been identified for the reactions of H3O+, NO+ and O2•− with MDA and the formation of their hydrates formed in humid samples is described. Thus, the following combinations of reagent and the analyte ions (given as m/z values) are adopted for SIFT-MS analyses of MDA in the gas phase: H3O+: 109; NO+: 89, 102; O2•−: 72, 90, 108, 126. The detection and quantification of MDA released by a cell culture by SIFT-MS is demonstrated.

CONCLUSIONS:
This detailed study has provided the kinetics data required for the SIFT-MS analysis of MDA in humid air, including exhaled breath and the headspace of liquid phase biogenic media. Consequently, the quantification of MDA in humid gaseous media can now be used to assess oxidative stress in biogenic systems.

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Malondialdehyde (MDA) is a frequently used indicator of free oxygen-radical-induced lipid peroxidation. Thus, MDA monitoring is of interest in all diseases in which the molecular aetiology increases ‘oxidative stress’, including (but by no means limited to) atherosclerosis, diabetes, liver disorders, cancer, autoimmune and neurogenerative disease [1, 2] [3-6]. Most usually, MDA is assayed in liquid state serum, plasma, urine, saliva and other biological samples by colorimetric and fluorimetric techniques [7]. As far as we are aware, the direct quantification of MDA molecules in the gas phase has not been achieved, but should this be possible, it could be a valuable target compound and a non-invasive biomarker via headspace studies of biogenic fluids and in breath analysis for bacterial lung infection or even lung cancer. However, the high reactivity and instability of MDA [6] might inhibit its use as a gas phase biomarker. Given our recent focus on the exploitation of SIFT-MS analysis of exhaled breath [8-11] and headspace of mammalian and bacterial cell cultures [12-15] when MDA has often been mentioned, we now feel constrained to investigate the potentially rewarding prospect of gas phase MDA analysis. Thus, in the present study we describe a method for the accurate real-time quantification of gas phase MDA using selected ion flow tube mass spectrometry (SIFT-MS) supported by solid phase microextraction coupled with gas chromatography mass spectrometry (SPME/GC/MS) for compound confirmation. This study follows similar focused kinetics studies of the ion chemistry of H2CO [16], H2S [17], HCN [18], CH4 [19] and C5H12 [10] as reported previously in Rapid Comm. Mass Spec., which were required to achieve analysis of these compounds in humid air by SIFT-MS.

**Background MDA chemistry**

MDA (see Scheme 1) is one of many aldehydes that are produced during the peroxidative decomposition of unsaturated fatty acids [20]. Compared to free radicals, most aldehydes are quite stable, diffuse from cells and attack targets far from their site of their production. About 32 aldehydes have been identified as products of lipid peroxidation [21] including saturated aldehydes (propanal, butanal, hexanal, octanal and decanal), 2,3-trans-unsaturated-aldehydes (hexenal, octenal, nonenal, decenal and undecenal) and a series of 4-hydroxy-2,3-trans-unsaturated aldehydes (4-hydroxyundecenal and 4-hydroxynonenal). MDA has been considered for some time to be the most important lipid peroxidation metabolite [21].

![Scheme 1](image)

Scheme 1. Two structural forms of malondialdehyde with molecular formula C₃H₄O₂ and relative molar mass 72.07 g/mol.

Pure MDA is a solid with melting point of 72-74°C. It is known to be relatively stable in pH neutral solutions but unstable under acidic conditions. Under physiological conditions (pH 5-7) it is presumed to be present in the form of a much less reactive conjugate base [−O−CH=CH−CHO] [22]. MDA can interact with many biological molecules including lipoproteins, peptides and DNA, altering their chemical behaviour [21], which has attracted much attention in different
research areas. Its high reactivity is based on its electrophilicity that makes it strongly reactive toward cellular compartments and lipid cell membranes. It has been detected in many animal and vegetable foodstuffs and plants and is found in human and animal tissue as an end-product of lipid peroxidation.

**Current methods used for MDA quantification**

Several methods are currently used to identify and quantify MDA in biological fluids, including spectrophotometry and fluorimetric detection, high performance liquid chromatography (HPLC), gas chromatography (GC) and immunological techniques [5]. The most common method for MDA analyses is based on the reaction with 2-thiobarbituric acid (TBA) in acidic media at 100 °C forming the MDA-TBA complex that can be quantified by spectrophotometry or fluorimetry [2]. In spite of their simplicity and low cost, such analyses have received wide criticism over the years. The main problem is the lack of sensitivity and specificity, since TBA reacts with a variety of compounds (so called TBA reactive species, TBARS), including other aldehydes, sugars, amino acids, bilirubin and albumin [21]. The results of TBARS analyses, even though often quoted in the units of MDA concentration (for example μmol/L), do not necessarily indicate the presence of MDA. Selective extraction of MDA molecules from biological samples is difficult because they are small, polar, relatively unstable and highly water-soluble [23]. Therefore, analytical derivatization is commonly employed to stabilise the MDA molecules and also increase their volatility. Both HPLC [24] and GC have been used [25-27] that demonstrated this approach to be more specific than the widely used TBARS [2]. The use of TBARS and HPLC in combination has been also reported [28]. Especially interesting to the subject of biomarker detection are the recent HPLC studies of MDA in exhaled breath condensate [29, 30]. Also relevant are the GC analyses of derivatised MDA in blood headspace using solid phase microextraction (SPME) [31].

**Rationale for the present study**

There is a growing interest in non-invasive monitoring of lipid peroxidation processes by analyses of volatile organic compounds in the headspace of biological samples or in exhaled breath. Volatile hydrocarbons resulting from polyunsaturated fatty acids oxidation have been studied previously in this context [10, 32]. Several aldehydes resulting from peroxidative processes have also been studied in the gas phase using SIFT-MS and GC/MS [33, 34]. A detailed SIFT-MS investigation of head space aldehydes as biomarkers of peroxide oxidation processes has been carried out by B. Ross and co-authors [35]. This study involved SIFT-MS quantification of 18 saturated volatile aldehydes however, MDA was not included. The promising volatile biomarker of lipid peroxidation MDA has so far only been quantified in headspace after its derivatization [21]. For the analysis of MDA molecules in humid air by selected ion flow tube mass spectrometry (SIFT-MS), the rate coefficients and the product ion distributions for the reactions of the SIFT-MS reagent ions with gas phase MDA molecules in the presence of water vapour are required and this need is the prime motivation for the present study.
Experimental

As a first step, a selected ion flow tube, SIFT, study was performed of the reaction of H₃O⁺, NO⁺ and O₂²⁺ precursor ions with MDA molecules in order to establish the primary product ions of the reactions and hence to identify appropriate analyte ions for the analysis of MDA by SIFT-MS. To achieve this, the rate coefficients for the analytical reactions are also required, but, as we will explain, it was not possible to determine these experimentally, so theory has to be invoked. Inevitably, secondary reactions of the primary product ions of the reactions with water molecules occur when analysing humid mixtures, so these reactions were also studied in order to identify which hydrate ions must be included as additional analyte ions in the SIFT-MS kinetics library in order to achieve accurate analyses of MDA in humid air [36-39]. The collected kinetic data were then used to formulate the appropriate kinetics library entries [36, 40]. Additional to these SIFT studies it was considered important to perform the GC/MS experiments described below in order to corroborate the SIFT mass spectrometric identification of the synthesised MDA that was used in all these experiments. Finally, pilot real-time analyses of MDA formed via lipid peroxidation were carried out to test the efficacy of detection and analysis of MDA by SIFT-MS.

Selected ion flow tube, SIFT, determination of product ion branching ratios

The SIFT technique has been described in detail for the determination of the rate constants and ion product distributions of ion-molecule reactions, including many reactions of H₃O⁺, NO⁺ and O₂²⁺ with many classes of organic compounds [10,16-19,38,41,42]. Ion swarms [43] of H₃O⁺, NO⁺ and O₂²⁺ precursors are formed by injecting mass filtered ions into He carrier gas. Then, in the present experiments, the headspace of an aqueous solution containing a trace amount (typically less than 10 ppmv) of MDA vapour, prepared as described below, was introduced into the SIFT instrument (Profile 3 SIFT-MS manufactured by Instrument Science Limited, Crewe, UK) via a heated calibrated capillary and full scan (FS) mass spectra comprising the precursor ions and the product ions and any of their hydrates were acquired. The m/z mass spectral range covered was chosen as 10–180 that was wide enough to encompass all the primary product ions and their hydrates. For each precursor ion species, five full mass spectra (FS) were obtained with a total integration time of 60 s, as exemplified later. The major ion products were identified and their count rates were precisely determined in separate experiments using the multi-ion monitoring (MIM) mode of SIFT and SIFT-MS [42]. In order to determine the primary product branching ratios of the reactions, the percentages of the individual product ions were plotted on a linear scale as a function of the sample mixture flow rate, as described previously [38, 41] and exemplified later.

Preparation of the MDA solution

Pure MDA is an unstable aldehyde that cannot be easily stored and shipped. Thus, for the purposes of these experiments, it was necessary to synthesize the MDA, and this was done by diluting 10 µL of 1,1,3,3-Tetraethoxypropane (TEP) with 10 mL of 0.1 M HCl solution, this mixture then being incubated at 100°C for 10 min to hydrolyse the TEP to MDA [27]. Thus, a standard solution of MDA at a concentration of 300 mg L⁻¹ was obtained. Ethanol is a by-product of this synthesis and appears in the solution headspace at concentrations that can
interfere with SIFT and SIFT-MS measurements. Fortunately, its concentration can be
minimised by judicious choice of the TEP/HCl ratio and by partial venting to reduce the
solution/headsapce ethanol concentration. As can be seen later in the SIFT-MS spectra, the
ethanol headspace concentration is reduced to acceptably low levels by these procedures. The
chemicals and reagents used were purchased from Sigma Aldrich spol. s r.o. (Sokolovská 100/94
Praha 8).

The headspace of this solution was used both as the sample air/vapour mixture for the SIFT
studies and for the SPME/GC/MS extraction/pre-concentration studies. Additional GC/MS
studies involved the commonly used derivatization method. Thus, the MDA was derivatized by
pentaphluorophenylhydrazine (PFPH) during 10 min of incubation time at 70°C in 15 mL vials
closed with septa in which 5 mL of MDA solution or sample was mixed with 1 mg of PFPH
(powder 99%) [22]. Then the volatile MDA derivative was adsorbed from the headspace using an
SPME fibre and injected into the GC/MS instrument in the usual way.

Cancer cell culture

Lung adenocarcinoma cancer cells (A549) in culture are used as a model for the pilot test of the
developed real-time SIFT-MS quantification method. STR-validated A549 cells were cultured
according to ATCC (Manassas, Virginia) recommendations in the RPMI-1640 medium
(developed by Moore et. al. at Roswell Park Memorial Institute, Buffalo NY [44]) supplemented
with 10% heat-inactivated foetal calf serum (FCS), 2mM glutamine, and 1% penicillin/streptomycin. The confluence (fraction of surface area covered by cells) was 80%.

SPME/GC/MS sampling and analysis protocol

The volatile organic compounds (VOCs) were extracted from the headspace of the standard
MDA mixture along with the MDA-PFPH derivative using carboxene polydimethylsiloxane
(CAR/PDMS)-coated SPME fibres (Supelco, Bellefonte, PA, USA) for a period of 15 min at a
temperature of 37°C. Then the fibres were directly inserted into an injector of the GC/MS
instrument (FOCUS GC with split/splitless injector (SSL), ITQ 700 ion trap mass spectrometer
using electron ionisation, Thermo Fisher Scientific Inc., USA ) held at 220°C. A GC/MS capillary
column TG-624 (fused 100% cyanopropylphenyl polysiloxane, 30m x 0.25mm ID x 1.0um film.,
Thermo Fisher Scientific Inc., USA) was used under the following conditions: splitless injection,
helium carrier gas flow of 1 mL/min, temperature program 38°C (hold 3 min) 20°C/min ramp
up to 220°C (hold 3min). The total run time was 15 min. Electrons at energy of 70eV were used
to generate ions which were analysed by the ion trap operating in the scan mode (m/z 15–400,
scan rate 1 scan/s).

Results and discussion

The structure of this section is as follows: firstly, the results of the ion chemistry studies i.e. the
experimentally determined ion products and the calculated rate coefficients for the reactions of
H3O+, NO+ and O2•• with MDA molecules in the presence of water molecules are given. Then a
short section indicates the appropriate entries to the SIFT-MS kinetics library that are required
for MDA analysis. This is followed by a brief section describing the SPME/GC/MS experiments
that verifies the identification of MDA. Finally, an example is given of the production and detection of MDA by an *in vitro* cell culture subjected to peroxidative stress, which shows that SIFT-MS can track the production of MDA in biological systems.

**Reaction of H₃O⁺ with MDA.**

A SIFT spectrum obtained using H₃O⁺ precursor ions when the headspace of the MDA solution (1,1,3,3-Tetraethoxypropane (TEP) with 0.1 M HCl, Sigma Aldrich spol. s r.o. Sokolovská 100/94 186 00 Praha 8) is introduced into the helium carrier gas is shown in Fig. 1, plotted as the log of the ion count rate, c/s, against the mass-to-charge ratio, m/z, of the ions. Immediately obvious are the majority H₃O⁺ (m/z 19) and its hydrates H₃O⁺(H₂O)₁,₂,₃ at m/z values of 37, 55, 73 and much smaller ion peaks at m/z 91 and 109 that could be designated as the higher-order H₃O⁺(H₂O)₄,₅ hydrates. However, very significantly, the nominal molecular mass of MDA (C₃H₄O₂) is 72 Daltons and because of the high proton affinity of aldehydes [39] it is confidently expected that the MDA molecule will accept a proton from H₃O⁺ on every H₃O⁺/MDA collision to produce MDAH⁺ ions also at m/z 73. These MDAH⁺ ions will efficiently hydrate to produce MDAH⁺(H₂O)₁,₂ ions at m/z 91 and 109. So the overlap of the m/z 73 ions precludes the use of MDAH⁺ as an analyte ion for MDA analysis by SIFT-MS as does the ions at m/z 91, because the H₃O⁺(H₂O)₄ ions are relatively abundant product ions when humid samples are being analysed. Fortunately, H₃O⁺(H₂O)₅ production is inefficient and thus m/z 109 ions can be used as an analyte ion for MDA analysis. However, the relative fraction of MDAH⁺(H₂O)₂ in the total analyte ions needs to be known and, even then, only an approximate quantification of MDA can be obtained. Because of these problems, further consideration of H₃O⁺ as a reagent ion for SIFT-MS analysis of MDA is not given here and attention is now given to the more suitable NO⁺ and O₂⁺ reagent ions. Note in the spectrum of Fig. 1 that the peak levels of the characteristic product ions of the H₃O⁺/ethanol reaction at m/z values of 47, 65 and 83 are relatively small.

**Reaction of NO⁺ with MDA**

A SIFT spectrum obtained using NO⁺ precursor ions when the headspace of the MDA solution is introduced into the helium carrier gas is shown in Fig. 2, as usual plotted as the log c/s, against m/z. Immediately obvious are the majority NO⁺ ions at m/z 30 and its hydrates NO⁺(H₂O)₁,₂ at m/z 48 and 66. The primary product ion at m/z of 71 is the result of the hydride ion extraction reaction between NO⁺ and C₃H₄O₂ (MDA) resulting in C₃H₃O₂⁺; this ion partially hydrates to its monohydrate C₃H₃O₂⁺H₂O at m/z 89. The product ion at m/z 102 is the adduct ion NO⁺C₃H₄O₂ also formed in primary reaction and that at m/z 120 is its monohydrate NO⁺C₃H₄O₂.H₂O. Parallel hydride ion transfer and adduct formation is not uncommon in the reactions of NO⁺ with aldehydes, especially so for unsaturated aldehydes [34, 39, 45].

It is clear from these data that NO⁺ reagent ions are a much better prospect for the quantification of MDA by SIFT-MS. So in order to determine how the relative signal levels of the four product ions at m/z 71, 89, 102 and 120 vary with sample humidity the standard approach to branching ratio variations has been taken in which the signal levels (c/s) of these ions are measured whilst introducing the humid headspace of the standard solutions of MDA at varying flow rate into the
helium carrier gas of the instrument. The humidity of the sample headspace was varied by raising its temperature in a controlled way, and since the sample flow is only a small fraction of the helium carrier gas flow, the increase in temperature of the helium/sample mixture is insignificant. The humidity is determined from the relative signal levels of the precursor ion hydrates, as is the routine procedure in SIFT-MS \cite{35,46}. Sample data are shown in Fig.3 revealing that the count rate of the major primary product ion at \( m/z \) 71 decreases rapidly with increasing humidity until at 6% humidity (corresponding to the saturated water vapour pressure at body temperature of 37°C). The hydrate at \( m/z \) 89 becomes comparable in magnitude to the ion at \( m/z \) 71 under these SIFT (and SIFT-MS) conditions. The adduct ion at \( m/z \) 102 is never greater than about 20% of the total product ion signal, and its hydrate at \( m/z \) 120 is not more than about 5%.

Reaction of O\(_2\)\(^{+}\) with MDA

A SIFT spectrum obtained using O\(_2\)\(^{+}\)precursor ions when the headspace of the MDA solution is introduced into the helium carrier gas is shown in Fig. 4, again plotted as the log c/s, against \( m/z \). Immediately obvious are the majority O\(_2\)\(^{+}\) ions (\( m/z \) 32) and the H\(_3\)O\(^+(\text{H}_2\text{O})_{0,1,2,3} \) ions at \( m/z \) 19, 37, 55 and 73 that are always seen and are formed in the complex ion chemistry of O\(_2\)\(^{+}\) with water molecules \cite{47} (in contrast, see the relative signal levels of these ions in Fig. 1). The very small signal at \( m/z \) 50 is the monohydrate of the O\(_2\)\(^{+}\). The clearly identifiable product ion originating from MDA are those at \( m/z \) 72, 90, 108 and 126, but this ion chemistry is complicated and multiple smaller product ions are seen at \( m/z \) 43 and 71 that are the result of fragmentation in the relatively energetic charge transfer occurring in the reaction of O\(_2\)\(^{+}\) with MDA molecules; this fragmentation process is quite common in the reactions of O\(_2\)\(^{+}\) with polyatomic molecules \cite{38,42}. For the moment, we choose to exclude these minor fragment ions and, as for the NO\(^{+}\) reaction, the signal levels of the aforementioned four product ions were measured as a function of the sample gas humidity. Sample data are shown in Figure 5 revealing that the count rate of the major parent cation MDA\(^{+}\) at \( m/z \) 72 decreases rapidly with increasing humidity until at 6% humidity its hydrates MDA\(^{+}(\text{H}_2\text{O})_{2,3} \) at \( m/z \) 108 and 126 becomes comparable to its magnitude under these SIFT (and SIFT-MS) conditions. Curiously, the monohydrate at \( m/z \) 90 is always a small fraction of the total ion signal; what is remarkable and unusual is the rapidity of the hydration to the dihydrate and trihydrate ions, which may be a feature of molecules containing both –OH and –CHO groups, but the gas phase reactions of such molecules with ions have rarely been studied.

Construction of kinetics library entries for the analysis of MDA by SIFT-MS

The prime objective of the above ion chemistry study was to identify the most suitable reagent ion species with which to quantify MDA in the gas phase and to detect and quantify MDA in the humid air that is the headspace above biological fluids. The following are the \( m/z \) values of the major product ions for the reactions of H\(_3\)O\(^{+}\), NO\(^{+}\) and O\(_2\)\(^{+}\) with MDA molecules, those given in bold being potential analyte ions for SIFT-MS analysis of MDA:

\[ \text{H}_3\text{O}^{+}; \, 73, \, 91, \, \textbf{109} : \, \text{NO}^{+}; \, 71, \, 89, \, 102, \, \textbf{120} : \, \text{O}_2^{+}\text{O}^{+}; \, 43, \, 71, \, 72, \, 90, \, 108, \, 126 \]
The rationale behind the inclusions/exclusions as analyte ions of any of these product ions for each reagent ion is given in the previous sections. Excluding any of the product ions requires that their fractions of the total product ions must be accounted for when constructing the kinetics library entries for SIFT-MS analyses [36, 40]. This is done by applying a multiplication factor (greater than unity) to those product ions included as analyte ions, as can be seen in the constructed kinetics library entry shown in Table 1; this is elaborated upon below. These multiplication factors are obtained from data such as those shown in Figures 3 and 5 by optimising the results of calculations of MDA concentrations so that they are constant across the entire range of humidity.

For accurate SIFT-MS analyses, rate coefficients for the analytical reactions must also be entered into the kinetics library. Experimental difficulties often preclude their measurement, usually because the preparation of gaseous mixtures of volatile organic compounds at accurately known concentrations is difficult. Such is especially so for the unstable MDA. Under such circumstances, the only approach is to rely on theory to calculate the collisional rate coefficients. Fortunately, it has often been shown that when proton transfer from an ion to a molecule is exothermic (by more than 25 kJ/mol) then it occurs on every collision, i.e. at the collisional or gas kinetic rate [48]. Because the proton affinity of all aldehyde molecules exceeds that of the water molecule [49] then proton transfer is facile from H₃O⁺ to MDA and the collisional rate coefficient can be calculated by the theory of Su and Chesnavich [50]. The collisional rate coefficients are calculated ab-initio using the DFT B3LYP method (NWChem software version 6.5, Pacific Northwest National Laboratory, Richland, Washington, US) using a dipole moment of 4.8 D, and a polarizability of 6 Å² calculated according Miller and Savchik [51] are given in Table 1. However, it cannot be tacitly assumed that the reactions of NO⁺ and O₂⁺ with MDA proceed at their collisional rates, but it is often possible to perform SIFT experiments in which the loss rates of these two ion species can be compared to that of H₃O⁺ from which data the rate coefficients for the NO⁺ and O₂⁺ can be deduced. But such experiments require that sufficient flows of the organic vapour, in this case MDA, can be introduced into the helium carrier gas to significantly reduce the signals of the reactant precursor ions [41]. Unfortunately, this could not be achieved because the production process for MDA adopted in these experiments could not realise high enough concentrations of pure MDA in the liquid headspace. Nevertheless, many studies in many laboratories have shown that the reactions of NO⁺ and O₂⁺ with aldehydes and most other polyatomic molecules do proceed at or very close to their respective collisional rates and so it is acceptable to use the aforementioned collisional theory to calculate appropriate rate coefficients for their reactions with MDA. Thus, the calculated rate coefficients for the reactions of H₂O⁺, NO⁺ and O₂⁺ with MDA molecules and those of the hydrates H₂O⁺(H₂O)₁,₂,₃ and NO⁺(H₂O)₁,₂, which must be included as reagent ions for accurate SIFT-MS analyses [36, 37, 40], are adopted in the kinetics library entries. These are given in Table 1 for the analysis by SIFT-MS of MDA in samples at 6% absolute humidity using the three reagent ions. Further discussion of the construction of these kinetics library entries in the format used in Profile 3 SIFT-MS software is given in recent papers [36, 38].

GC/MS verification of MDA synthesis
The ion chemistry studies described above, in particular the ion products of the NO+ reaction that conform to the expectations for the reaction of NO+ with aldehydes [34, 39, 45, 52], confirmed the identity of the synthesized molecule as MDA. To our knowledge, direct gas phase detection and analysis of MDA molecules has not been carried out previously; thus, we felt constrained to reinforce and verify the presence of MDA.

The detection of MDA by preparing a derivative using pentafluorophenylhydrazine (PFPH) to form the stable volatile adduct N-pentafluorophenylpyrazole followed by extraction of the derivative from the headspace using SPME followed by GC/MS analysis is well established [25-27]. The structure of MDA-PFPH derivative is shown in Fig. 6. Its electron ionisation (EI) fragmentation pattern is known [27], the major ion being its molecular ion at m/z 234 with other minor fragment ions at m/z 117, 167, 180, 194 and 207. So, following this precedent, we used PFPH to prepare the derivative of the assumed MDA in the standard water solution, (300mg/L) as used for the ion chemistry studies, and used SPME/GC/MS for its analysis.

Fig. 6 shows the limited SPME/GC–MS chromatogram indicating a retention time for the MDA–PFPH derivative of 12.17 minutes in the GC column used (TG-624 fused 100% cyanopropylphenyl polysiloxane). The EI mass spectrum obtained for this eluted compound is also shown in Fig. 6 where it can be seen that the m/z values of the peaks (m/z 234 for the molecular radical cation and fragments at 117, 167, 180, 194 and 207) conform to those given above and thus correspond to the known mass spectrum from previous works[27]. This we take as confirmation that the synthesized molecule is indeed MDA. It is interesting to note that a clear peak appeared in the wider GC chromatogram at a retention time of 5.60 minutes. This we postulate to be underivatized MDA molecules, but the EI spectrum for MDA is not included in the GC/MS database and is presumably unknown. So we performed this SPME/GC/MS experiment again, confirmed an elution time of 5.60 minutes and thus obtained the 70 eV MS fragmentation spectrum for MDA, as is shown in Figure 7. As far as we know, the EI mass spectrum of MDA has not been given in any NIST database and has not been published previously. The parent molecular ion MDA+ at m/z 72 is obvious, but the major fragment ion is at m/z 71 that results from loss of an H atom from the nascent excited (MDA+)*. The minor ion at m/z 54 may results from loss of a water molecule [MDA-H2O]+ and the ion at m/z 43 probably result from the loss of an HCO moiety as is typical of EI fragmentation of aldehydes. This study clearly demonstrates that the direct detection of MDA in the gas phase has been achieved and it can be quantified using SIFT-M

**SIFT-MS quantification of MDA in the headspace of a model biological fluid**

To test the claim stated in the last sentence of the previous section, we carried out the following experiment. Lung adenocarcinoma cancer cells (A549) (of particular interest to a member of the team (S.S.A) and the Imperial College London Oncology Group [53]) were cultured in serum medium. Hydrogen peroxide (H2O2) was then added at a lethal concentration (20mM) to initiate peroxidation stress, which is expected to generate MDA [54-56]. Direct and continuous SIFT-MS analysis of the headspace was performed for a few minutes, beginning immediately after the
addition of the H$_2$O$_2$, using NO$^+$ reagent ions and the analyte ions at $m/z$ values of 71, 89, 102 and 120. Due to overlaps with other headspace compounds only $m/z$ 89 and 102 were included for quantification (see Table 1). The results obtained are given in Fig. 8 which show that the count rates of all four analyte ions increased to peak values after a time of about 8 minutes (Fig. 8a), thus reflecting the MDA concentration which increased for about 8 minutes and then began to decrease. The limit of detection, as calculated from the count rates of the product ions$^{[40]}$, was 0.3 parts-per-billion by volume, ppbv, and the precision of quantification was ±3 ppbv. These data indicate that either MDA production slows down due to cell membrane destruction and death or that the MDA initially produced, which resides in the culture, was depleted as the headspace was continuously sampled into the SIFT-MS instrument. This speculation is not the major point of this experiment; rather, it is to demonstrate that MDA can now be detected and quantified in the gas phase in real time by SIFT-MS obviating derivatisation that can disturb the biological system.

**Concluding remarks**

The SIFT-MS analytical method has often been shown to be a valuable and accurate method for ambient analysis of trace compounds in humid air, including exhaled breath. This requires that the ion chemistry underpinning the analyses is well understood, and in this regard, special attention needs to be given to some compounds because of the peculiar aspects of their reactions with ions in the gas phase. This is especially so when attempting to analyse unstable molecules such as the biologically important MDA, the focus of the present study. Thus, following a detailed study of the relevant ion chemistry, we have shown that MDA can now be directly detected and quantified by SIFT-MS in real time in humid air like the headspace of biological fluids such as cell and tissue cultures, avoiding the complications associated with molecular derivatisation of the MDA. An additional contribution made by this study is the provision of the EI mass spectra of MDA, which facilitates the direct GC/MS analysis of MDA molecules following SPME collection. These advances open the way to detailed investigations of peroxidation processes in biology.

**Acknowledgements**

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Table 1. SIFT-MS kinetics library entries\(^a\) for the determination of the concentrations of MDA in humid samples using H\(_3\)O\(^+\), NO\(^+\) and O\(_2\)\(^{+}\) reagent ions at 300K

<table>
<thead>
<tr>
<th>MDA (H(_3)O(^+))</th>
<th>MDA(O(_2)(^{+}))</th>
<th>MDA(NO(^+))</th>
</tr>
</thead>
<tbody>
<tr>
<td>4 precursors</td>
<td>1 precursor</td>
<td>3 precursors</td>
</tr>
<tr>
<td>19 5.7e-9 1.0</td>
<td>32 4.7e-9 1.0</td>
<td>30 4.8e-9 1.0</td>
</tr>
<tr>
<td>37 4.5e-9 1.0</td>
<td>4 products</td>
<td>48 4.1e-9 1.0</td>
</tr>
<tr>
<td>55 4.0e-9 1.0</td>
<td>72 1</td>
<td>66 3.8e-9 1.0</td>
</tr>
<tr>
<td>73 3.7e-9 1.0</td>
<td>90 1</td>
<td>4 products</td>
</tr>
<tr>
<td>1 product</td>
<td>108 1</td>
<td>71 1</td>
</tr>
<tr>
<td>109 5.0</td>
<td>126 1</td>
<td>89 1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>102 2.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>120 1</td>
</tr>
</tbody>
</table>

\(^a\) Each entry contains a list of precursor ion m/z values each followed by the rate coefficient in units of cm\(^3\)s\(^{-1}\). Then a list of product ion m/z values is given. At the end of each row a coefficient is given that is used to multiply the count rate measured at the given m/z which accounts for any excluded analyte ions; see the text for further explanation.
References.


Fig. 1 Full scan spectrum obtained when the humid headspace of the MDA solution is flowed into the helium carrier gas/H3O+ ion swarm of the SIFT instrument showing the product ions at m/z 91 that comprise H3O+(H2O)4 and MDAH+(H2O) ions, and MDAH+(H2O)2 ions and at m/z 109; see text for further explanation.
Fig. 2 Full scan spectrum obtained when the humid headspace of the MDA solution is flowed into the helium carrier gas/NO$^+$ ion swarm of the SIFT instrument showing four characteristic product ions at $m/z$ 71, 89, 102, 120 due to MDA; see text for further explanation.
Fig. 3 Relative intensities of the product ions MDA-H⁺ (C₃H₅O₂⁺), NO⁺MDA (C₃H₄O₂NO⁺) and their monohydrates as a function on the sample humidity within the range 1% to 6% observed by SIFT using NO⁺ precursor ions.
Fig. 4 Full scan spectrum obtained when the humid headspace of the MDA solution is flowed into the helium carrier gas/O2• ion swarm of the SIFT instrument showing product ions at m/z 43, 71, 72, 90, 108 and 126; see text for further explanation.
**Fig. 5** Relative intensities of the product parent cation $\text{MDA}^+ (\text{C}_3\text{H}_4\text{O}_2^+)$ and its hydrates as a function on the sample humidity within the range 1% to 5% observed by SIFT using $\text{O}_2^{+•}$ precursor ions.
Fig. 6 a) SPME/GC–MS chromatogram showing the peak at an elution time of 12.17 min due to the derivative MDA-PFPH (structure shown) and b) and the EI mass spectrum obtained for the MDA-PFPH showing the molecular ion an \( m/z \) of 234 and several fragments.
Fig. 7. SPME/GC–MS chromatogram (a) and EI mass spectrum (b) obtained for the extraction and analysis of the headspace of the standard aqueous solution of MDA (300mg/L). The peak at 5.60 min is due to free MDA.
Fig. 8 SIFT-MS real time quantification of MDA in the headspace of a culture of lung adenocarcinoma cancer cells (A549) after 20 mM of H₂O₂ has been added at time 0 at a culture temperature of 38°C. a) shows the product ion count rates using NO⁺ precursor ions (count rate 500000 c/s) and b) shows the gas phase quantification of MDA in parts-per-billion by volume, ppbv as calculated from the count rate of product ions at m/z 71, 89 and 102.