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VUV photoionization aerosol mass spectrometric study on the iodine oxide particles formed from O_3 -initiated photooxidation of diiodomethane (CH_2I_2)

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lodine oxide particles (IOPs) formed from O_3 -initiated photooxidation of diiodomethane have been investigated based on the combination of a thermal desorption/tunable vacuum ultraviolet time-of-flight photoionization aerosol mass spectrometer (TD-VUV-TOF-PIAMS) with a flow reactor for the first time. Characterization of the home-made flow reactor was performed, which indicates the applicability of its combination with TD-VUV-TOF-PIAMS. Based on that, aerosol mass spectra of IOP formation from photooxidation of CH_2I_2/O_3 were studied on-line taking full advantage of both the virtues of the flow reactor and TD-VUV-TOF-PIAMS. The main chemical components of IOPs, including atomic and molecular iodine (I, I_2), iodine oxides (IO, OIO, I_2O and I_2O_3) and hydrogen-containing iodine species (HI, HIO and HIO $_3$), were observed and identified based on the corresponding photoionization energy (PIE) curves, and the probable chemical composition and formation mechanism of IOPs were proposed. The work has not only improved the understanding of the formation mechanism of IOPs, but also demonstrated the capability of TD-VUV-TOF-PIAMS for direct molecular characterization of aerosols in flow reactor experiments, whose potential application in mass spectrometric studies of atmospheric aerosols is anticipated.

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Introduction

Atmospheric iodine chemistry has received increasing attention during the last few decades¹ owing to its important role in the catalytic destruction of ozone²,³ and new particle formation (NPF) in the marine boundary layer (MBL).⁴ Although having been known for more than one century,⁵ NPF in the costal atmosphere was postulated to be related with classical binary (H₂SO₄/H₂O) or ternary (H₂SO₄/NH₃/H₂O) nucleation for a long time. Only recently, has it been identified that iodine compounds (I₂ or alkyl iodides) play a key role in ultrafine particle bursts at the coastal zones, and NPF is formed mainly during formation of iodine oxide particles (IOPs).⁶-9

$$I_2$$
, CH_3I , CH_2I_2 , $etc. \xrightarrow{h\nu} I + products$ (R1)

$$I + O_3 \rightarrow IO + O_2 \tag{R2}$$

Then OIO will be formed from the IO self-reaction:17,18

$$IO + IO \rightarrow OIO + I$$
 (R3)

The following recombination of OIO with IO or another OIO, followed by oxidation by O₃, will result in the formation of condensable iodine oxides (I_xO_y , x = 2, 3, 4; y = 3, 4, 5):^{11,18,19}

$$OIO + IO \rightarrow I_2O_3$$
 (R4)

$$OIO + OIO \rightarrow I_2O_4$$
 (R5)

$$I_2O_3 + O_3 \rightarrow I_2O_4 + O_2$$
 (R6)

$$I_2O_4 + O_3 \rightarrow I_2O_5 + O_2$$
 (R7)

Nucleation and growth of IOPs, originating from iodine photochemistry, has been investigated in numerous laboratory studies. $^{10-16}$ Generally, it is believed that I atoms produced from the photolysis of iodocarbons or I_2 will be oxidized by O_3 leading to the production of IO: 1

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RSC Advances

At last, condensation or further polymerization of I_xO_y will lead to formation of IOPs. So IOPs formation was traditionally believed to occur only when the precursor of iodine, O₃, and UV radiation are present simultaneously. Although a number of studies have tried to rationalize the mechanism of IOPs formation, it is still controversial about which is the most probable IOPs chemical composition, I₂O₄ or I₂O₅. While both of transmission electron microscope analysis and modelling calculations of IOPs generated from photochemical reaction of I2 and O3 in dry conditions indicated that IOPs are essentially I₂O₅, ¹¹ mass spectrometric study complemented with ab initio quantum calculations showed that I2O4 may be the most plausible candidate to initiate nucleation of IOPs,14 and experimental studies on IOPs formation from the two different IO generating ways (I + O₃ and O + I₂) based on flow tube also showed that polymerization of I₂O₄ play a pivotal role in IOPs formation.13 Therefore unambiguous evidence of chemical compositions of IOPs is still a major research challenge in

understanding the mechanism of IOPs formation. The application of mass spectrometric techniques to the measurement and characterization of aerosols represents a significant advance in the field of atmospheric science. Over the past decades, aerosol mass spectrometry techniques have been developed considerably, offering off-line or on-line chemical analysis for individual particles or on ensembles. 20,21 Although their application in laboratory and field has resulted in many important improvements in understanding of atmospheric aerosols and their processing, 22-24 these traditional aerosol mass spectrometry techniques suffered for extensive fragmentation of organic constituents due to "hard ionization" (for example, electron ionization or multiphoton ionization), which prevents specification of chemical components of atmospheric aerosols. Inspiringly, aerosol mass spectrometry coupled with vacuum ultraviolet (VUV) photoionization based on synchrotron radiation (SR) sources has been proved to be an effective "soft" ionization method for real-time, molecular component analysis of organic particles.25-27 SR light sources is featured with high photon flux and tunability. High photon flux makes it has better detection sensitivity than traditional VUV lamp for ultrafine particle analysis. Tunability makes it capable of analyzing broad range of constituents of atmospheric aerosols in two ways. One is it can offer fragment-free or less fragment mass spectra by single-photon ionization. The second is it can identify different chemical composition ambiguously by their ionization energies (IE) or appearance energies (AE) while not only by m/z values through tuning ionization energy. Making the advantage of these features, AMS coupled with a SR source has been successfully applied to laboratory studies of secondary organic aerosols (SOAs) formation and reaction kinetics.28-32 However, in the laboratory studies based on the combination of smog chamber with SR-VUV photoionization AMS, the feature of identifying compounds by IE or AE cannot be made the full advantage. 30-32 The reason is that the acquisition of photoionization efficiency (PIE) curves (eventually to obtain IE or AE) needs tens of minutes to several hours depending on the scanning range and scanning step of SR light, while at the same time, the concentrations of the compounds

(aerosol particles or gaseous reagents and products) in smog chamber is always changing according to the reaction progress. So it is inevitable that the observed PIE curves are the superimposition of the evolution curve of the compounds to be measured with the inherent PIE curves just depending on the energy of the exciting SR light, which inhabits the accurate identification of the compounds especially for the weak signals of mass spectra. In contrast to that in smog chamber, the concentration of the compounds to be measured in a fixed sampling site only depends on the total flow rate for a fixed flow reactor, ^{33,34} which is particularly suitable for PIE curve measurement as aforementioned.

In this work, our motivation was to make full advantages of both the virtues of flow reactor and thermal desorption/tunable vacuum ultraviolet time-of-flight photoionization aerosol mass spectrometry (TD-VUV-TOF-PIAMS) to study the formation mechanism of IOPs. As CH2I2 is the most abundant iodinecontaining compounds from biogenic emissions35 and plays an important role in the formation of iodine oxides in marine boundary layer, 36 photooxidation of CH₂I₂/O₃ was chosen as the reaction system to produce IOPs. Firstly, characterization of home-made flow reactor was performed. Then aerosol mass spectra of IOPs forming from photooxidation of CH₂I₂/O₃ were investigated based on the combination of the flow reactor and the TD-VUV-TOF-PIAMS, and the main chemical components of IOPs were identified based on the PIE curves. Finally, the probable chemical composition and formation mechanism of IOPs, as well as the potential application of the combination of the flow reactor and the TD-VUV-TOF-PIAMS based on SR were discussed.

2. Experimental methods

Chemical analysis of IOPs formation from photooxidation of CH_2I_2 at the presence of O_3 was performed based on the combination of the TD-VUV-TOF-PIAMS and the home-made flow reactor (Fig. 1) at the Atomic and Molecular Physics Beamline of the National Synchrotron Radiation Laboratory (NSRL) in Hefei (China).³⁷

As shown in Fig. 1, the flow reactor used in this study is similar in spirit to the setup in Kamens's group in University of North Carolina.33 Briefly, it consisted of two 1 m straight sections of a 2.5 cm diameter quartz glass tube and one Ushaped connection tube. While there are 16 sampling ports distributed every 10 cm along the straight sections, only the exit of the tube acted as the sampling ports in this study. Generally, the total flow rates through the tube were in the range of 1.2-2.0 l min⁻¹ corresponding to a Reynolds number of 146-243, which indicates the laminar flow conditions in the flow reactor. Similar to that in our smog chamber experiments, 38,39 the flow reactor was continuously flushed with dry zero air until the background particle concentration less than 20 cm⁻³ prior to each run. The zero air also acted as precursor to produce O₃ as well as the bath gas of CH2I2. The flow rates of CH2I2 and O3 were controlled respectively with mass flow meters, and their concentration in the flow tube were monitored separately with a gas chromatograph-electron capture detector (GC-180 ECD,

Paper

Zero Air

O3 generator

Exhaust

Annular diffusion denuder

SMPS

Detection System

TD-VUV-TOF-PIAMS

Fig. 1 Schematic diagram of the flow reactor and the TD-VUV-TOF-PIAMS used for chemical analysis of IOPs.

Agilent 7820) and O_3 analyzer (TEI model 49i). A sintered glass mixing plate filled with glass beads at the entry of the reactor ensured immediate mixing of CH_2I_2 and ozone. As UV radiation was always kept on during the experiments and the photolysis of CH_2I_2 is a fast process, the time of the mixing of CH_2I_2 and ozone is taken as the zero point of the reaction of $I+O_3$. At the exit of the flow reactor, an annular diffusion denuder was used to remove the residual O_3 and CH_2I_2 to make sure the end of the gaseous reaction of $CH_2I_2+O_3$. The concentrations and size distributions of IOPs formed from the photooxidation of CH_2I_2/O_3 were monitored on-line with a Scanning Mobility Particle Sizer (SMPS, TSI, model 3936, TSI, USA) consisting of a differential mobility analyzer (DMA, TSI model 3080) and a condensation particle counter (CPC, TSI model 3775).

Real-time chemical analysis of IOPs were performed with the TD-VUV-TOF-PIAMS developed at the Atomic and Molecular Physics Beamline of the NSRL. 30 IOPs formed in the flow reactor were tightly focused and sampled into the mass spectrometer through a typical aerodynamic lens assembly 41-43 with a 200 µmdiameter orifice. After passing through a three-stage differential pumping system, the particles hit on an 8 mm diameter temperature-controlled copper tip located at the ionization region of the main chamber and were flash thermally vaporized at an appropriate temperature (in the range of 293-873 K). In this work, the same vaporization temperature (453 K) was chosen in all tests. The first reason is for high quality of mass spectrum of IOPs. The second one is this temperature is less than the decomposition temperature of I₂O₄ (460 K), ^{44,45} so this choice can keep the high intensity of the signal of mass spectrum while avoid the decomposition of I₂O₄ assumed they exist in IOPs. And the last one, keeping the constant vaporization temperature will make us comparing the IE values from different tests without considering the temperature effect on IE. The plume of the vaporized particles was then photoionized

with the tunable VUV beam from synchrotron radiation. The undulator-based spherical grating monochromator (SGM) beamline was operated with the lowest-energy grating, generating photons with energy from 7.5 to 22.5 eV. Generally, a photon flux of 10^{14} photons per second can be produced and the resolving power of 2700 ($E/\Delta E$) at 15.9 eV can be achieved with adjusting the entrance- and exit-slit width of the grazing-incidence monochromator. Ions formed in the photoionization region were then detected with a reflectron TOFMS. When the undulator is fixed, photoionization mass spectra of IOPs can be investigated at a fixed ionization energy. While scanning the undulator continuously, PIE curves can be extracted resulting in the characterization of the ions at molecular level, which is crucial to learn the chemical component of IOPs and the formation mechanism of IOPs.

Results and discussion

3.1 Characterization of the flow reactor

As aforementioned, the evolution of the components in smog chamber is always going on following the initiation of the reaction, which impedes the effective utilization of PIE curves when smog chamber is combined with SR photoionization AMS. Different from the smog chamber, flow reactor is characterized by the fixed reaction situation if the sampling port and the total flow rate are fixed, no matter how long the sampling duration time. So prior to the combination with the TD-VUV-TOF-PIAMS at NSRL, the stabilization of the home-made flow reactor was tested.

Three test experiments were performed according to the stabilization of the particle number concentration, the size distributions and the mass concentration of IOPs formed from photooxidation of $\mathrm{CH_2I_2/O_3}$ system in the flow reactor under different reaction conditions. As shown in Table 1 and Fig. 2,

RSC Advances

Table 1 List of the reaction conditions and the parameters of IOPs in test experiments

Test no.	Flow rate ^a (L min ⁻¹)	$[CH_2I_2]_0$ (ppm)	[O ₃] ₀ (ppm)	Mean number conc. (# 10 ⁶ /cm ⁻³)	Mode diameter (nm)	Mean mass conc. ^c (10 ³ μg m ⁻³)
#1	1.8	$\textbf{0.56} \pm \textbf{0.03}$	2.32 ± 0.12	8.53 ± 0.39	57.3 ± 0.9	8.62 ± 0.41
#2	1.3	0.52 ± 0.03	3.22 ± 0.16	$\textbf{7.62} \pm \textbf{0.80}$	66.1 ± 0.2	11.97 ± 0.26
#3 ^b	4.0	1.26 ± 0.25	0.50 ± 0.1	$\textbf{45.37} \pm \textbf{5.35}$	14.5 ± 0.3	$\textbf{0.69} \pm \textbf{0.03}$

^a Flow rate = total inlet flow rate (CH₂I₂ + O₃) = total outlet flow rate (SMPS sampling rate + pump rate). ^b Nano DMA was used in test #3. ^c All particles are presumed to be compact and spherical, and the particle density was set at 5.0 g cm⁻³ for all sizes.¹³

given the initial CH_2I_2 and O_3 concentrations ($[CH_2I_2]_0 \sim$ 0.5 ppm, $[O_3]_0$ 2.3-3.2 ppm), when total flow rates in flow reactor were kept less than 2L min⁻¹, the main diameter

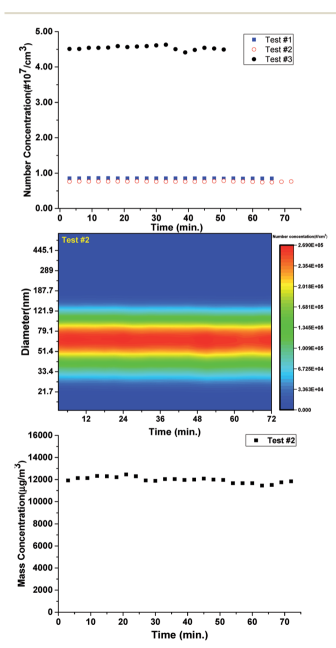


Fig. 2 The variation of the particle number concentration (a), the size distributions (b), and the mass concentration (c) of IOPs in the flow reactor according to the duration time.

distribution of IOPs formed was accumulation mode.46 For example, in test #2, the mode diameter of IOPs was 66.1 nm. And it is obvious that the production of IOPs was always kept stable given the constant precursor, oxidant and flow rate (in other words, fixed reaction time). In test #1 and #2, the variation of mean number concentration was less than 10%, the variation of central diameter was less than 2% and the variation of mean mass concentration, which is the key parameter to the bulk measurement with TD-VUV-TOF-PIAMS, was less than 5% during one hour test time. In test #3, the total flow rate in flow reactor was kept at 4L min⁻¹, corresponding to 16.5 seconds reaction time, and the concentration of O3 was around one sixth of that in test #2. In this case, Aitken mode⁴⁶ of IOPs produced in flow reactor, central diameter of which is about 14.5 nm, can be observed. Even for Aitken mode particles in test #3, the variation of mean number concentration is only 12% and the variation of mean mass concentration is 4% during about one hour test.

All of the tests show the perfect stabilization of the flow reactor, which indicates its applicability to be combined with TD-VUV-TOF-PIAMS based on SR to make full use of PIE curves. In this work, however, only IOPs with accumulation mode have been investigated with aerosol photoionization mass spectrum.

3.2 Mass spectrometric study on IOPs

3.2.1 Photoionization mass spectra of IOPs. Based on the combination of the flow reactor and the TD-VUV-TOF-PIAMS, photoionization mass spectra of IOPs formed from photooxidation of CH₂I₂/O₃ can be measured on-line when the radiation lights are on. In order to make sure the mass spectra are from IOPs, the background spectra were measured in two different ways. One was for the vacuum chamber of mass spectrometer, when the inlet of mass spectrometer was blocked. The other was for the compounds in flow reactor when the radiation lights were off while CH₂I₂/O₃ were still input. It is strange to find that the two background spectra showed the similar patterns especially in the range of m/z = 100 to m/z = 300 (only the later is shown in Fig. 3 as a representative), which indicates that the background are mainly resulted from the vacuum chamber of mass spectrometer. However, the background cannot be completely eliminated even the chamber had been baked and the vaporizer had been added with higher temperature (\sim 600 K) than the following normal experiments in order to desorb the potential pollution.

Paper

Fig. 3 The mass spectra of IOPs formed from photooxidation of CH_2I_2/O_3 (black line) and the background spectra (red line) at the photon energy of 11.0 eV. (a.u., arbitrary units).

A typical thermal desorption photoionization mass spectrum of IOPs, as well as the background, at photon energy of 11.0 eV is also shown in Fig. 3, where the vaporization temperature of heater is 453 K. As the major components of air, it is not strange to observe the signals of H_2O (m/z 18), N_2 (m/z 28), NO (m/z 30) and O_2 (m/z 32) in the background spectrum regarding to their high photoionization efficiencies and cross-section.⁴⁷ The peaks of ethanol (m/z 46), which was generally used as solvent to clean the orifice of the aerodynamic lens in this work, and its photofragment (m/z) 45) can also be identified in the background spectrum. In the range of m/z = 100 to m/z = 300, there are some unidentified while always existed features, which are labeled with stars as shown in Fig. 3. Compared with the background spectra in the range of m/z = 100 to m/z = 300, there are many new features resulted from the thermal desorption and photoionization of IOPs, corresponding to m/z = 127, 128, 143, 144, 159, 176, 254, 270 and 302.

As a "footprint" of a molecule or a radical, ionization energy or appearance energy can be used to identify molecular composition unambiguously. It is well known that photoionization mass spectrometry based on SR is characteristic of obtaining PIE curves by scanning photon energy, and IE or AE can be determined from PIE curves. In order to determined the corresponding molecular composition of the peaks observed in mass spectra of IOPs in Fig. 3, the PIE curves of these components were obtained by gating and integrating the corresponding peaks while scanning the photon energy in the range of 8.5-11.0 eV. To minimize the influence of the blocking of the orifice of aerodynamic lens by IOPs, the larger energy interval (0.1 eV) was applied in full range scanning in this work, while small energy interval (0.02 or 0.03 eV) was adopted around the threshold energy regions of IE or AE to obtain more precise value. Several typical PIE curves, corresponding to m/z = 127, 143, 159, 254 and 302, are illustrated in Fig. 4, where Fig. 4ad were acquired at 0.1 eV energy interval and Fig. 4e and f at 0.02 or 0.03 eV energy interval, and all the data points have been normalized by the corresponding photon intensities monitored simultaneously with a silicon photodiode (SXUV100, International Radiation Detector, Inc.). Both of Fig. 4a and f correspond

to m/z = 143 component in mass spectra of IOPs in Fig. 3, which shows how the more precise value of IE can be gotten.

All of the IEs derived from the PIE curves just like those shown in Fig. 4 are summarized in Table 2 with the references' data. And based on the IEs and the corresponding molecular weight, the assignments of these main components of IOPs were performed. There are three species in the photoionization mass spectra of IOPs. The first are only iodinecontaining, such as I^+ (m/z 127) and I_2^+ (m/z 254) with the IEs of 10.38 and 9.16 eV respectively. The second are iodine oxides, such as $IO^{+}(m/z 143)$, $OIO^{+}(m/z 159)$, $I_{2}O^{+}(m/z 270)$ and $I_2O_3^+$ (m/z 302). While the IEs of IO, OIO and I_2O_3 were measured as 9.49, 9.80 and 9.90 eV respectively, only the upper limit of IE of I2O could be obtained (<9.30 eV) for the weak signal in low photon energy. The third one are hydrogencontaining iodine species, such as HI^+ (m/z 128), HIO^+ (m/z 144) and HIO_3^+ (m/z 176), in which only the IE of HIO were measured as 9.70 eV and the peaks for HI⁺ and HIO₃ + were too weak to acquire PIE curves.

Although some of the chemical components observed here had been observed before in gaseous or particle phase, only a few IEs of these species had been reported. The slow dark reaction of I2 and O3 has been studied experimentally using 118 nm VUV light photoionization TOF-MS by Gómez Martín et al.,14 where atomic and molecular iodine, hydrogen iodide and iodine oxides have been observed in the gas phase products and the corresponding IEs has been estimated based on quantum chemistry calculations. The chemical composition of IOPs formed from photodissociation of CH₂I₂ has been studied by Hoffmann et al. using on-line atmospheric pressure chemical ionization mass spectrometry,15 and I-, I2-, IO-, IO2- and IO₃ have been detected in their works. Based on electron ionization aerosol mass spectrometry, Jimenez et al. have also studied new particle formation from photooxidation of CH2I2, and most of the species observed in this work have also been investigated in their work.16 Regarding to their technique of ionization, assignment of the chemical composition of IOPs were mainly depended on molecular weights not on IEs in these studies. As far as we know, only the IEs of I, I2, IO and HIO in gas phase have been investigated experimentally, 48,50,51 and it is the first time to directly investigate the IEs of iodine oxides from IOPs experimentally in this work. Although the IE values obtained in this work show good agreement with literature values, it seems that out values are systematically a little lower than the literature values obtained experimentally in gas phase. For example, IE of I2 form IOPs obtained in this work is 9.16 eV, while that obtained in gas phase is 9.31 eV (see Table 2). Regarding to the temperature of thermal desorption (453 K) used here, the population of excited vibrational states may be the partial reason resulting in the lower IEs in this work. For example, the fraction of I_2 in the second vibrational state (v_2) at

453 K is
$$f_2 = \left(1 - \mathrm{e}^{-\frac{\Theta_{\mathrm{vib}}}{T}}\right) \mathrm{e}^{-\frac{2\Theta_{\mathrm{vib}}}{T}} = 0.13$$
 (where Θ_{vib} , named vibrational temperature, is 308 K for $\mathrm{I_2}$), ⁵² and the ionization of the second vibrational state of $\mathrm{I_2}$ will result in about 0.06 eV

lower of ionization energy as $\tilde{\nu_e}$ of I_2 is 214.52 cm⁻¹.

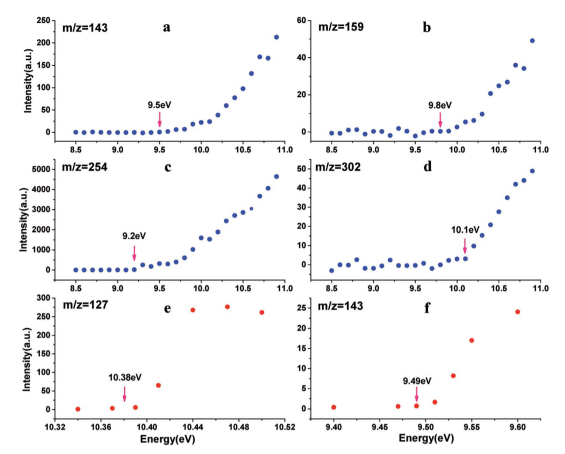


Fig. 4 PIE curves for m/z 143 (panel a and f), m/z 159 (panel b), m/z 254 (panel c), m/z 302 (panel d), m/z 127 (panel e) ((a-d), larger energy interval scanning; (e and f), small energy interval scanning). (a.u., arbitrary units).

Table 2 The assignment of the chemical components of IOPs based on their ionization energy value (this work and the references)

Ion	m/z	IE (eV) (this work)	IE (eV) (theo.) (ref.) ^a	IE (eV) (exp.) (ref.) ^b
I^+	127	10.38 ± 0.02	10.35	10.451
HI^{+}	128	N/A^c	10.50	10.386
IO^{+}	143	9.49 ± 0.02	9.56	$9.66^d, 9.745^e$
HIO^{+}	144	9.70 ± 0.03	N/A	9.811
OIO^{+}	159	9.80 ± 0.10	$9.72, 9.793^f$	N/A
HIO_3	176	N/A	N/A	N/A
I_2^+	254	9.16 ± 0.05	9.47	9.307
I_2O^+	270	<9.30	9.02	N/A
$I_2O_3^+$	302	$\textbf{9.90} \pm \textbf{0.03}$	9.97	N/A

^a ref. 14. ^b Ref. 48. ^c N/A not available. ^d Ref. 49. ^e Ref. 50. ^f Ref. 51.

3.2.2 Interpretation of IOPs chemistry. Based on the photoionization mass spectra of IOPs and the PIE curves, the main chemical composition of IOPs have been identified, which confirmed the presence of iodine oxides, iodine oxyacids as well as atomic and molecular iodine in IOPs observed before.16

Regarding to the facts that no IOPs were formed and no signal were observed in background mass spectra when the photooxidation of CH₂I₂/O₃ was not initiated, it is presumed that the observed I+ and I2+ may result from the thermal

decomposition or fragmentation of iodine oxides and/or hydrogen-containing iodine species in the IOPs.15,16 The main components of iodine oxides in IOPs are still controversial and the knowledge about them is still far deficient. Although highly oxidized iodine oxides, such as I_2O_v (y = 1, ..., 5) and I_3O_v (y = 1,..., 7) have been observed in gaseous products generated from the reaction of I2 and O3,14 the largest iodine oxide observed in IOPs is only I₂O₃ in this work and the work of Jimenez et al.¹⁶ Traditionally, I₂O₄ and I₂O₅ are believed to be the two main candidates via gas-phase collisions to nucleate the new IOPs. The crystal structure of I2O4 was approximated as a onedimensional solid comprising infinite -I-O-IO2-O- chains and weaker interchain I-O bands.53 While in solid I2O5, a I2O5 unit is composed of two IO3 pyramids which have one O atom in common, simply as O2I-O-IO2.54 However, both of the two iodine oxides are known to decompose under thermal induction:

$$5I_2O_4(150-200 \,^{\circ}C) \rightarrow 4I_2O_5 + I_2$$
 (R8)

$$2I_2O_5(\sim 400 \,^{\circ}C) \rightarrow 2I_2 + 5O_2$$
 (R9)

As the IOPs were vaporized at 180 °C in this work, the thermal decomposition of I₂O₄ may be the one potential source of the detected I⁺ and I₂⁺.

Paper

Secondly, although single-photon ionization in TD-VUV-TOF-PIAMS instead of 70 eV electron impact ionization used in traditional AMS was applied in this work, there was no guarantee to avoid the photodissociation of I₂O₄ or I₂O₅ during photoionization. Thus the photodissociation of I₂O₄ and I₂O₅ may be another reason for no detection of $I_2O_4^+$ and $I_2O_5^+$, as well as the detection of I⁺, IO⁺, OIO⁺, I₂O⁺ and I₂O₃⁺. A mass spectroscopic study on I₂O₅ aerosol particles generated from atomizing standard I2O5 sample will be performed in the near future, which is presumed to offer some useful information to verify this possibility. Thirdly, regarding to the weak peak of I₂O⁺ and even weaker signal of I₂O₃⁺, it is presumed that the photoionization cross sections of high iodine oxides, such as I₂O₄ and I₂O₅, are too small, thus the concentrations of I₂O₄⁺ and I₂O₅⁺ were too low that were beyond the detection limit of the aero mass spectrometer used in this work.

As for no detection of I_2O_4 and I_2O_5 , that whether I_2O_4 or I_2O_5 is the condensable unit of IOPs is still unclear and cannot be identified directly just depending on mass spectra. However, by combining the results of this work and the other studies, ^{10–18} the identity of I, IO, OIO, I_2 , I_2O and I_2O_3 , that is to say the reaction mechanism of (R1)–(R4) aforementioned in Introduction, has been consolidated.

It is interesting to observe ions of iodine acids, such as HI⁺, HIO⁺ and HIO₃⁺, in the mass spectra of IOPs. Similar ion signals of iodine acids were also observed before in AMS of aerosol particles formation from photooxidation of CH₂I₂ (ref. 16) and from I₂O₅ dissolved in water⁵⁵ in laboratory studies, as well as from new particle formation in an iodine-rich, coastal atmospheric environment in recent field research.⁵⁶ The detection of HI⁺, HIO⁺ and HIO₃⁺ indicates that either there are larger oxyacids in IOPs or there are new production of oxyacids on AMS vaporizer. 16 It is presumed in one way that when the IOPs impact on AMS vaporizer, I2O4 will thermally decomposes to I₂O₅ via (R9), and then I₂O₅ will react with H₂O to generate HIO₃. It is noteworthy that the peak of H₂O are hardly to be observed in the mass spectra of IOPs being compared with the background spectrum (see Fig. 3). However, more deep work are needed to identify whether the disappearance of H2O mainly results from its reaction with I₂O₅. While in dry conditions, the iodine oxyacids in IOPs may result from some unknown gasphase chemistry or from the inhomogeneous reaction between OH, HO2 or H2O with iodine oxides in IOPs.16 Regarding to the relative humidity (RH $\sim 23\%$) and the temperature of the vaporizer (T = 453 K) in this work, both of the aforementioned formation passages of HIO₃ are plausible, whichever indicates the presence of the higher iodine oxides.

4. Conclusions

In this work, chemical analysis of IOPs formed from the photooxidation of CH_2I_2 has been performed on-line based on the combination of the flow reactor system with the TD-VUV-TOF-PIAMS. Atomic and molecular iodine (I, I_2), iodine oxides (IO, OIO, I_2O and I_2O_3) and hydrogen-containing iodine species (HI, HIO and HIO₃) were observed in aerosol mass spectra, and their corresponding IEs were identified directly in aerosol phase

for the first time. The formation mechanism of IO, OIO, I2O and I₂O₃, as well as the potential source of I and I₂ has been consolidated. I2O4 and I2O5 haven't been detected directly in IOPs mass spectra in this work, the detection of HIO and HIO₃, however, indicates the presence of the higher iodine oxides in IOPs. Although more work are needed to be carried out for the CH₂I₂/O₃ reaction system itself, the study on chemical composition of IOPs carried in this work has shown that the combination of the flow reactor system with the SR-based TD-VUV-TOF-PIAMS is to be a promising approach for on-line study of chemical components and formation mechanism of atmospheric aerosol by making the full advantage of flow tube (sampling-duration-time-independent) and SR photoionization (identify the chemical composition ambiguously on IE or AE). As for experimental technique, more work is needed to resolve the block of the orifice of aerodynamic lens and to improve the detect limit of the TD-VUV-TOF-PIAMS.

Conflicts of interest

There are no conflicts to declare.

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