# Tomographic retrieval algorithm of OH concentration profiles using Double Spatial Heterodyne Spectrometers

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Abstract. The hydroxyl radical (OH) determines the capability of atmospheric self-cleansing and is one of the significant oxidants in atmospheric photochemistry reactions. The global OH has been monitored by satellites with the traditional limb mode in the past decades. This observed mode can achieve high-resolution vertical OH data, but cannot obtain the enough horizontal OH data for inverting high-precision OH concentrations because OH has the high reactivity that makes OH concentrations extremely low and distributions complicated. The Double Spatial Heterodyne Spectrometers (DSHS) is designed to achieve more high-resolution and detailed OH data. This sensor can measure the OH by the three-dimensional limb mode to obtain the comprehensive OH data in the atmosphere. We propose a new tomographic retrieval algorithm here based on the simulated observation data because the DSHS will work officially on the orbit in the future.

- We build up an accurate forward model. The main part of it is the SCIATRAN radiative transfer model which is modified according to the radiation transmission theory. The error of results obtained by the forward model is  $\pm 44.30\%$  in the lower atmosphere such as 21 km height and decreases gradually until the limit of observation altitude . We also construct the tomographic retrieval algorithm that the core is a lookup table method. A tomographic observed database is built up
- through the atmospheric model, the spatial information (the position of the target area and satellite position), the date parameters, the observation geometries, OH concentrations, and simulated observation data. The OH concentrations can be found from it directly. If there are no corresponding query conditions in the tomographic observed database, the cubic spline interpolation is used to obtain the OH concentrations. The inversion results are given and the errors of them increase as the altitudes rise until about 41 km height then start to decrease. The errors of the inversion results reach the maximum about ±25.03% at the 41 km and decrease to ±8.09% at the limited observation height. They are also small in the lower atmosphere which are ±12.96% at the 21 km. In summary, the tomographic retrieval algorithm can obtain
- the more accurate OH concentrations even in the lower atmosphere where the OH data is not well and avoids the setting of initial guess values for solving the iteration problems. Our research not only provides support of scientific theory for the construction of DSHS, but also gives a new retrieval algorithm idea for other radicals.

## 35 **1 Introduction**

The OH plays an important and initial role in atmospheric photochemistry reactions because of its strong oxidation. It can remove many natural and anthropogenic compositions which are significant for the air quality, the ozone distributions even the climate change from the atmosphere (Lu et al., 2019;Stevens et al., 1994). OH mainly derives from the reactions of  $O(^{1}D)$ , a photolytic product of ozone in the ultraviolet band, with water vapor in the middle and upper atmosphere.

- 40 The removal of OH is affected by various compounds like nitrogen oxides, sulfur dioxide, carbon monoxide, methane, and other volatile organic compounds (Lelieveld et al., 2016;Wolfe et al., 2019;Zhang et al., 2018). Therefore, on the one hand, the OH affects the photochemical and kinetic processes in the atmosphere, reflects the short-term and longterm climatic evolution processes in some respects. On the other hand, it is of great significance to enhance the understanding of the atmospheric physical and chemical processes.
- 45 It is a great challenge to monitor the OH in the atmosphere because of its low concentrations and strong activity. The Fluorescence Assay by Gas Expansion (FAGE) has been used to excite the OH radical continuously by the 308 nm excitation mechanism for generating the fluorescent signal. It obtains the OH concentrations through the relationship between the OH concentrations and the fluorescent signal (Hard et al., 1984). The Differential Optical Absorption Spectroscopy (DOAS) has been used to obtain the OH concentrations because the absorption of OH follows the Lambert-
- 50 Beer absorption law (Perner et al., 1976). The Chemical Ionization Mass Spectrometry (CIMS) has been used to collect ions of OH instead of the photons based on OH oxidation for obtaining the OH concentrations (Mauldin et al., 1998). These methods are commonly used to measure the OH concentrations in the actual limited environments and laboratories. Besides, the <sup>14</sup>CO oxidation method depends on the <sup>14</sup>CO<sub>2</sub> concentration, enrichment coefficient, reaction rate constant, and reaction time to get the OH concentrations due to the OH oxidability (Felton et al., 1990). The Scrubbing using the
- 55 salicylic acid Technique has been used to obtain OH concentrations by an acid and its production rate (Salmon et al., 2004). The Spin Trapping method has been used to obtain the OH concentrations by electron spin trap and 4-OH-POBN in the laboratory for some theoretical researches (Watanabe et al., 1982). Apart from the six physical and chemical methods mentioned above, many researchers used the high-precision spectrum data from the ground-based instruments especially the Fourier Transform Ultraviolet Spectrometer (FTUVS) in the Table Mountain: The OH P<sub>1</sub>(1) absorption
- 60 spectrums which are measured by FTUVS have been used to invert the OH concentrations (Cageao et al., 2001). The OH Q<sub>1</sub>(2) absorption spectrums have been used as an auxiliary spectrum to improve the accuracy of OH concentrations (Mills et al., 2002). An improved retrieval method is described that uses an average method based on spectral fits to multiple lines weighted by line strength and fitting precision to obtain the OH concentrations. (Cheung et al., 2008). These methods are restricted by the features of the instruments which can provide accurate OH data in the limited lower
- 65 atmosphere but are difficult to provide enough data for inverting OH concentrations from mesopause to tropopause. The development of satellite technology carried out the chance of global wide-scale detection of OH. Conway retrieved OH concentrations by the least square fitting method until the success of the Middle Atmosphere High-Resolution Spectrometer Investigation (MAHRSI) which provided the first observed data of OH in the mesosphere with diffractions grating technology from space (Conway et al., 1999). A new interferometric technology called spatial heterodyne
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- 70 spectroscopy has been applied to the Spatial Heterodyne Imager for Mesospheric Radicals (SHIMMER) for reaching the higher spectral resolution on the mid-deck of the space shuttle with small size and no mobile optical components. These satellite sensors measured the OH solar resonance fluorescence by the limb mode from the low Earth orbit to obtain more accurate OH data. Englert used the same method as the MAHRSI to invert OH concentrations (Englert et al., 2008). Besides these sensors mentioned above, the 2.5-THz radiometer on the Microwave Limb Sounder (MLS) has been
- 75 designed to measure the thermal emission signal of OH from stratosphere to mesosphere because the spectral region around the pair of strong OH lines at 2.51 and 2.514 THz is clean relatively. The sensitive thermal emission data is used to invert the volume mixture ratio of OH concentrations. However, the results have a poor signal to noise ratio in some atmospheric regions. Livesey used the standard optimal estimation method to obtain the OH concentrations from the calibrated MLS observed data (Level 1B) with the radiative transfer equation. He also corrected the noisy products by a
- 80 separate task using the full radiance dataset as the relevant band after the main process (Livesey et al., 2006). Wang used the average orthogonal fitting method to handle with the MLS data for obtaining the OH concentrations and compared with results of the ground-based FTUVS in some areas. They matched very well (Wang et al., 2008). Damiani handled the MLS data of nighttime winter in the northern hemisphere from the latitude of 75° to 82° by the regional mean method to research the change of OH concentrations in the short-term (day) and long-term (weeks) dimension (Damiani et al., 85 2010).
  - The MAHRSI and SHIMMER measured the OH solar resonance fluorescence in the  $A^2\Sigma^+ X^2\Pi(0,0)$  band at around 309 nm, proved that is the more possible and effective way to monitor atmospheric OH from space load at present. The traditional limb mode can obtain high-resolution vertical OH radiance profiles which contain the information of OH concentrations at different tangent planes because of the benefit of spectral technology. However, the OH radiance which
- 90 is measured at one field of view is the sum of the overall contributions in the corresponding fields of view, but the OH is inhomogeneous in fact. These will lead to some precision errors (Englert et al., 2010). Anhui Institute of Optics and Fine Mechanics, Chinese Academy of Sciences, has done some researches on the DSHS in advance to improve the horizontal resolution of OH data and achieve the better structure of OH profiles. The DSHS, will work on the orbit at 500 km, is designed to measure the OH solar resonance fluorescence from 15 to 85 km by using the double spatial
- 95 heterodyne spectrometers in orthogonal layout. The field of view angle is designed as 2 degree and the spectral resolution will reach 0.02 nm at least. This paper gives the OH interferogram which is emulated by a forward model and describes a new method called the tomographic retrieval algorithm which uses simulated DSHS data and theory of the lookup table method to invert OH concentrations. The OH concentrations in the target area will be found out in the tomographic observed database which constructs by the atmospheric model, the spatial information, the date parameters, the
- 100 observation geometries, OH concentrations productions, and the simulated observation data. The retrieval algorithm can obtain more accurate OH concentrations in the middle and upper atmosphere especially in the lower atmosphere than the results which are obtained by traditional limb retrieval algorithms and can raise the inversion speed compared with the traditional limb retrieval algorithms.

## 2 Measurement strategy

### **2.1 The principle of spatial heterodyne spectroscopy**

The DSHS is designed based on the principle of the spatial heterodyne spectrometer which the optical structure is shown in Fig. 1. The two planar mirrors in the traditional Michelson interferometer are replaced by two diffraction gratings (grating 1, grating 2). The light, enters through aperture A, is collimated by the lens L1. It is split into two beams of coherent light of equal intensity by the beam splitter. One of them is reflected by the beam splitter and incidents on the grating 1. The light is diffracted by the grating 1 and returns to the beam splitter. The other light incidents on the grating 110 2 through the beam splitter as well. It is diffracted by the grating 2 and reflects back to the beam splitter. The gratings are fixed which are placed at a Littrow angle  $\theta$  with the orthogonal plane of the optical axis in the spatial heterodyne spectrometer system. The light incidents on the gratings by  $\theta$  angle and diffracts back with  $\theta$  angle at some wavenumbers. It is called the light of Littrow wavenumber  $\sigma_0$ . The two exiting wavefronts of light of  $\sigma_0$  are diffracted 115 by the gratings and perpendicular to the optical axis. The phase difference is zero and the interference fringe spatial frequency is zero that cannot form the interferogram. However, the light of non-Littrow wavenumber has an  $\pm \gamma$  angle with the optical axis. The light forms interferogram which will image on the imaging detector by the optical imaging systems L2 and L3. So, the overlapping two wave surfaces have an angle of  $2\gamma$  which are calculated by Eq. (1) (Dohi and Suzuki, 1971):

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$$\sigma[\sin\theta + \sin(\theta - \gamma)] = \frac{m}{d} \tag{1}$$

where the  $\sigma$  is the wavenumber of light, m is the order of diffraction, 1/d is the grating groove density. The angle between the light of arbitrary wavenumber and the exiting light of Littrow wavenumber is  $\gamma$ . The spatial frequency of two  $\sigma$  light is given by Eq. (2):

$$f_x = 2\sigma \sin \gamma \approx 4(\sigma - \sigma_0) \tan \theta \tag{2}$$

For the input spectrum  $B(\sigma)$ , the intensity on the imaging detector is given by Eq. (3):

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$$I(x) = \int_0^\infty B(\sigma) \left( 1 + \cos\left(2\pi (4(\sigma - \sigma_0)x\tan\theta)\right) \right) d\sigma$$
(3)

The spectral curve  $B(\sigma)$  can recover from interferogram I(x) by Fourier transform algorithm.



Figure 1. Schematic diagram of spatial heterodyne spectrometer.

Furthermore, the spatial heterodyne spectroscopy can acquire spatial distribution of one-dimensional spectral information through the two-dimensional detection technology. The scene in the field of view is divided into multiple field of view slices which are less or equal to the number of rows in the focal plane array by adding a cylindrical mirror to the front or rear optical system. The interferograms of each field of view slices are imaged on the corresponding detector rows respectively. The several rows on the detector correspond to a layered spectral information in a field of view. The corresponding spatial resolution unit on the detector can indicate the spatial information when the target signal appears in a certain range of line. These mean that the spatial heterodyne spectroscopy can obtain the radiance of atmospheric information at different altitudes simultaneously without scanning the atmosphere from a height to other height respectively. It is important for monitoring the OH to reduce the time of data acquisition because the OH is

### 2.2 Instrument innovation

variable fast in the spatial and temporal dimension.

- 150 The spatial heterodyne spectroscopy technology, an ultra-high-resolution spectroscopy technology used in the DSHS, has the characteristics of high flux, no mobile part and small size. The DSHS which is shown in Fig. 2 mainly consists of two spatial heterodyne spectrometers in the orthogonal layout to obtain the enough data for inverting the OH concentrations. It includes the telescope cylinder system, the collimating system, the information processing system, the imaging system, and some optical components else. The field of view angle is designed as 2 degree. One of the spatial heterodyne spectrometers which scans along the satellite working orbit is defined as SHS1, the other which scans across
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the satellite working orbit is defined as SHS2. The scanning direction of SHS2 is orthogonal to SHS1s' due to the special design of the orthogonal layout. The hierarchical detection of a series of observed radiance in the same target area comes true when the satellite flies along its orbital path. It means the sensor can obtain the data of each target height like the multi-angle method.



**Figure 2.** Illustration of the Double Spatial Heterodyne Spectrometers. the left part (a) shows the vertical view of the DSHS and the right part (b) shows the overall mold of the DSHS.

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The DSHS will detect the altitudes from 15 to 85 km by the three-dimensional limb mode. It can obtain the observed data in the target area along the satellite working orbit and across the satellite working orbit respectively at the same time because the SHS1 and SHS2 are orthogonal to each other. Figure 3 shows how the tomographic data is obtained by the three-dimensional limb mode.

- 170 As Fig. 3 (a) shows that there are no intersection regions between the field of view slices of SHS1 and SHS2s' at the time of T1. The tomographic data cannot be obtained in this situation. The three-dimensional structure of OH data is unable to be reconstructed. The data of SHS1 and SHS2 at different time will be used to get an intersection region for solving this issue. The different time which meet the requirements of three-dimensional limb mode of the DSHS (we define one time as the time of T1 and the other time as the time of T2) are shown in Fig. 3 (b). At the time of T1, the
- 175 SHS1 and SHS2 finish a limb scanning in the vertical dimension (along the satellite working orbit) and the horizontal dimension (across the satellite working orbit). Then the satellite platform will need to reorient for getting the appropriate vector for the measurement at the time of T2 and finishes the observation project. The spectrometers are on during this process. The DSHS has the function of hierarchical imaging in the spatial dimensional which means it can obtain the data without scanning the target area. So, The DSHS records at T1 and T2 is very brief. The line of sight at the time of
- 180 T1 will intersect with the line of sight at the time of T2. The intersection region (target area) will form. However, the composition of tomographic data does not need all data from the SHS1 and SHS2 at the time of T1 and T2. Some data should be omitted. As Fig. 3 (c) shows that the field of view slices of T2-SHS2 (a horizontal dimension spectrometer at the time of T2) are orthogonal to the field of view slices of T1-SHS1 (a vertical dimension spectrometer at the time of T1) in an intersection region. The data obtained by the SHS2 at the time of T1 and the SHS1 at the time of T2 will be

185 omitted in this measurement process. However, these data may be used for different measurements in other target areas to obtain the tomographic data when they meet the requirements of three-dimensional limb mode. Thus, the threedimensional segmentation of the target area will complete. The three-dimensional division of the target atmosphere which consists of the intersection regions of SHS1 and SHS2 at different time is accomplished finally with the satellite platform moving. Different combinations of data for different intersection regions from two spatial heterodyne 190 spectrometers are used to invert OH concentrations at different time.



Figure 3. Three-dimensional limb mode of DSHS at the time of T1 and T2 (a). The DSHS consists of two spatial
heterodyne spectrometers (SHS1 and SHS2). The data of target area cannot be obtained by DSHS at the time of T1 and T2 respectively (b). DSHS monitors the OH in the target area using the data of SHS1 at the time of T1 and SHS2 at the time of T2 together (c). The arrow indicates the direction of the flight of the spacecraft.

The scanning direction will not be completely vertical or parallel to the surface when the DSHS will be on the orbit but

- will have an angle  $\eta$ . So, the satellite positions at the time of T2 where T2-SHS2 is orthogonal to T1-SHS1 in the intersection region are different with the  $\eta$  changing. In order to display and descript the satellite position of T2 conveniently, the  $\eta$  is set to 0° temporarily. However, a unified set of  $\eta$  is 30° in the relevant calculation during the pre-research phase.
- The satellite positions and observation geometries are given at the time of T1. The satellite positions at the time of T2 need to calculate for realizing the three-dimensional segmentation of the target atmosphere. It indicates that there are two satellite positions where the scanning directions of T2-SHS2 and T1-SHS1 are orthogonal to each other in the intersection region at the time of T2 based on the theory mentioned above. Figure 4 shows that there is a tangent point which is defined as P1 when the DSHS detects the atmosphere by the three-dimensional limb mode. The intersection region along the lines of SHS1 (the blue line in Fig. 4) and SHS2 are the observed data at the time of T1. There
- 210 will be two lines of sight (the red lines in Fig. 4) where the scanning direction of SHS2 is orthogonal to the SHS1s' under the premise that the working altitude of the satellite is constant. The satellite position on the same side to the time of T1 is defined as the position at the time of T2-1, and on the opposite side to the time of T1 is defined as the position at the time of T2-2. The two lines of sight at the time of T2 (T2-1 and T2-2) and a single line of sight at the time of T1 form two intersection regions R1 and R2 which are symmetrical about P1 along the line of sight at the time of T1. Figure 5
- 215 shows the relationship of the satellite positions at the time of T1 and T2 in the space. These two intersection regions are on the two sides of the tangent point and are at the altitude of 500 km. The distance of the satellite position at the time of T2-2 from the time of T1 is farther than the satellite position at the time of T2-1 that means the satellite needs more time to work from the position of the time of T1 to the position at the time of T2-2. The observed data will change a lot in this process and make bigger difference in the target area at the time of T1. The satellite position at the time of T2-2
- 220 should be omitted. Meanwhile, the satellite position at the time of T2-1 is near the position at the time of T1, the observed data at the position of the time of T2-1 can reflect the OH distribution in the target area better because the OH information does not change too much in the short working time of satellite. The distances between the intersection region and the tangent point determine the spatial resolution of the three-dimensional limb mode. That means the smaller distance between the P1 and R1 (or R2) is, the higher spatial resolution will be.

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**Figure 4.** Limb geometry diagram for the three-dimensional limb mode at the time of T1 and T2. The blue line is the line of sight of SHS1. The red lines are the lines of sight of SHS2.



Figure 5. Relative location of DSHS by three-dimensional limb mode. There are two locations of the satellite at the time of T2 which meet the requirement of three-dimensional limb mode. Map data: © Google Earth, Image IBCAO and Landsat/Copernicus.

So, the position of the time of T2-1 is picked up as the other satellite location where the field of view slices are orthogonal to each other at the intersection region. The OH concentrations at different heights are obtained by the combinations of observed data from the SHS1 at the time of T1 and SHS2 at the time of T2 in the intersection region. Although the observed data are different because the geometric position and the viewing angle are of difference between the time of T1 and T2 the OH concentrations in the target area will also be constant. The data forms and the structures of the observed data of the SHS1 and SHS2 are the same in three-dimensional limb mode. The distance between the intersection point R1 and tangent point P1 is presumed as 50 km which is used in the subsequent research.

## 245 **3 Forward modeling**

The forward model is used to express a physical process of how the atmospheric state parameter is obtained by the sensors. A precise forward model is significant for the design of the instrument and the retrieval algorithm. The observed data received by the DSHS with an ultra-high spectrum resolution makes up by two parts in this research: the atmospheric background radiance and the OH fluorescence emission radiance. The atmospheric background radiance is the result that the solar radiance is subjected to Rayleigh scattering and trace gas absorption. The OH fluorescence emission radiance is subjected to the OH fluorescence emission mechanism. These data can be simulated by a forward model which is constructed by the flow chart in Fig. 6.



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Figure 6. Flow chart of the forward model.

The forward model is mainly constituted by a radiative transfer model called the SCIATARN which is developed for the Scanning Imaging Absorption spectrometer for Atmospheric CHartographY (SCIAMACHY) (Rozanov et al., 2014). However, some parts of it need to modify for meeting the characters of the DSHS and OH.

First, the radiance source, the solar radiation, is considered in this forward model for the radiance transmission of OH in

the middle and upper atmosphere because (1) OH fluorescence emission radiance will be generated from the excited state to the ground state when the OH is excited by the solar radiation. A high-precision calculation of the OH fluorescence emission radiance requires the high-resolution solar radiation, (2) the DSHS has a spectral resolution of 0.02 nm at least. The higher spectral resolution solar radiation will be used to get more realistic values. So, the National

- Solar Observatory (NSO) solar spectrum with the resolution of  $8.6 \times 10^{-4}$  nm is used for this forward model. In addition to the NSO, the precision of the OH spectrum is also significant. SCIATRAN just takes the Rayleigh scattered radiance function, the ozone absorption function, the OH self-absorption function into account, but ignores the effect of OH fluorescence emission radiance. The Lifbase software is used to calculate the OH emission spectrum in a temperature
- 270 ranges according to the Bremen atmospheric model (Luque et al., 1999; Sinnhuber et al., 2009). The OH emission spectrum database is built up based on its. The database is put into the SCIATRAN as a source function. The corresponding OH spectral emission data can be got from the database when the observed radiance is simulated at some conditions.

Next, the OH absorption spectrum database in the SCIATRAN bases on the HITRAN 2012 database by default. However,

- 275 the HITRAN 2012 database does not include the OH absorption profiles in the ultraviolet band. The OH absorption profiles in the HITRAN 2014 are used to solve this problem. The HITRAN 2012 is still retained but the OH UV absorption lines data from the HITRAN 2014 are added into it.
  - Finally, the precise atmospheric composition parameters are the assurance to simulate the observed radiance received by the sensors. That means the high-quality OH concentrations database is needed when the radiance transfer is simulated
- 280 accurately. For one thing, the MLS on the Aura has monitored the global OH for many years and the data can be acquired publicly at present. These data are used to build up the OH concentrations database. MLS was launched in 15 July 2004 aboard on Aura and kept offering consequent OH data until 2009. It was stopped monitoring from the November 2009 to August 2011 to extend the service life of the sensor. It was restarted to monitor OH on the August and the September each year continually to obtain annual change trends of OH data. For another thing, the solar activity was relatively
- stable and maintained at a low level in the whole solar activity cycle from 2005 to 2009. The solar activity from 2010 to 285 2016 had just ended an active cycle, and the level of solar activity in the next few years would be similar to the situation from 2005 to 2009. Therefore, the OH concentrations data from 2005 to 2009 is a great reference for the research of atmospheric OH concentrations by the DSHS in the next few years. The MLS OH concentrations Level 2 geophysical production is used and averaged within a certain range for reducing the random errors to build up the OH concentrations
- database. The N32 Gaussian grid from European Centre for Medium-range Weather Forecasts is selected as the averaged 290 method due to the random errors of MLS OH concentrations in high latitude are much larger than them in low latitude. It is a lattice-level coordinate system for the scientific model of spheres in earth science. The latitude and longitude zones in the grid are divided unconventionally. The latitude band intervals of the northern and southern hemispheres are symmetrical about the equator. The latitude band intervals and the number of longitude bands on a latitude gradually
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- decrease with the increase of the latitude to ensure that each grid area is approximately equal. In addition, the four seasons definition is used, the first quarter (March/April/May), the second quarter (June/July/August), the third quarter (September/October/November), the fourth quarter (December/January/February), as the temporal resolution for the OH concentrations database. The season is the first quarter and the OH concentrations database is N32-seasonly database.
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The simulated atmospheric background radiance is mainly subjected by the function of Rayleigh scattering, ozone absorption, and OH self-absorption. It is calculated based on some parameters such as the spatial, temporal and observation geometries. An observed radiance profile is given as an example in the (27 °N,106 °E) area in Fig. 7. The intensity of simulated observation radiance and atmospheric background radiance at some tangent heights are also given in Table 1.



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Figure 7. An observed radiance profile obtained by the modified SCIATRAN.

	Table 1	. The intensi	y of	simulate	d observa	tion rac	diance and	l atmosr	oheric	backgro	ound ra	adiance a	t some t	angent	heig	zhts
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Altitude (km)	Intensity of simulated observation radiance (phot <sup>-1</sup> s <sup>-1</sup> cm <sup>-2</sup> nm <sup>-1</sup> sr <sup>-1</sup> )	Intensity of simulated atmospheric background radiance (phot <sup>-1</sup> s <sup>-1</sup> cm <sup>-2</sup> nm <sup>-1</sup> sr <sup>-1</sup> )
21	8.933×10 <sup>13</sup>	8.885×10 <sup>13</sup>
31	9.436×10 <sup>13</sup>	9.379×10 <sup>13</sup>
41	$1.043 \times 10^{14}$	$1.034 \times 10^{14}$
51	5.184×10 <sup>13</sup>	5.086×10 <sup>13</sup>
61	$1.498 \times 10^{13}$	$1.429 \times 10^{13}$
71	$3.682 \times 10^{12}$	3.195×10 <sup>12</sup>
81	6.030×10 <sup>11</sup>	5.866×10 <sup>11</sup>

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The results of modified SCIATRAN will be imaged in the imaging system which collects complete data from all pixels on one line of CCD. The digital number (DN) values of each pixel in each row are generated by Eq. (4) at different tangent heights:

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$$S_{CCD}(i,j) = \int_{y(j)}^{y(j+1)} \int_{x(i)}^{x(i+1)} \int_{0}^{\infty} B(\sigma, y) R(\sigma) \left(1 + \cos\{2\pi[4(\sigma - \sigma_0)x\tan\theta]\}\right) d\sigma dx dy$$
(4)

where *i* is the number of rows along the line, *j* is the tangent height from 15 to 85 km. The x(i) and y(j) is the areas where grating imaging image in the CCD,  $B(\sigma, y)$  is the input radiance spectrum,  $R(\sigma)$  is the instrument function,  $\sigma$  is the wavelength number and the  $\theta$  is the Littrow angle. The interferogram will be got based on the instrument parameters in Table 2.

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#### Table 2. Instrument parameters for emulation

Parameters	value
Littrow length (nm)	306
Grating groove density (lines/mm)	1000
Numbers of interferogram samples	1024
Grating width (line/mm)	15
Littrow angle (°)	8.80

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An emulated interferogram is shown in Fig. 8. It is the two-dimensional observed radiance interferogram constructing by 1024 columns which indicate numbers of interferogram samples and 36 rows which indicate the observed tangent heights. The observed radiance gradually increases to reach the peak value which is the bright of interferogram with the altitude increases until the altitude around 40 km. The bright decreases gradually until the upper limit of the detection height.





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The errors of this forward model mainly depend on three factors based on the modified parts above: the atmospheric model, the Doppler effect, and the instrument calibration errors. First, the ozone in the atmospheric model is the influential factor that needs to be considered because its absorption cannot be ignored in the ultraviolet band. Second,



the satellite has different heliocentric speeds when it flies over the area where the local solar time is noon or is around the morning faint line. The speed will lead to the shift of observed radiance at the ranges of wavelength. Third, there are many sources of errors in the process of converting electrical signals into radiance values. The instrument calibration errors are assumed as  $\pm 5\%$  because the DSHS has not been officially working. The total errors of observed radiance are also given in Table 3. Zhang had made the detailed analysis for errors and the reasons (Zhang et al., 2017).

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Table 3. Total errors of observed radiance caused by different factors at some tangent heights

Tangent height (km)	Errors caused by the atmospheric model (%)	Errors caused by Doppler effect (%)	Errors caused by instrument calibration errors (%)	Total errors (%)
81	From -2.65 to 5.67	From -0.87 to 0.85	From -5 to 5	From -9.52 to 9.52
71	From -2.61 to 5.62	From -1.18 to 0.95	From -5 to 5	From -9.52 to 9.52
61	From -2.81 to 5.83	From -0.98 to 0.83	From -5 to 5	From -9.67 to 9.67
51	From -4.90 to 8.17	From -0.91 to 0.78	From -5 to 5	From -11.93 to 11.93
41	From -16.49 to 26.17	From -0.82 to 0.67	From -5 to 5	From -31.75 to 31.75
31	From -18.92 to 36.64	From -0.77 to 0.65	From -5 to 5	From -41.85 to 41.85
21	From -19.56 to 39.10	From -0.76 to 0.65	From -5 to 5	From -44.30 to 44.30

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The DSHS is constituted by double spatial heterodyne spectrometers in the orthogonal layout. The distance between the intersection point and the tangent point is defined as 50 km and the angle between the spatial heterodyne spectrometer scanning direction and the vertical/horizontal direction is defined as 30°. The SHS2 positions at the time of T2 can be calculated according to the SHS1 observation geometry parameters at the time of T1 in the target area based on the detection theory of three-dimensional limb mode. The simulated observation radiance of SHS2 and SHS1 can be obtained by the modified SCIATRAN above. The partial associated geometric parameters of the time of T1 and the calculated geometric parameters of the time of T2 are given in Table 4.

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**Table 4.** Locations and geometry parameters of SHS1 at the time of T1 at some tangent height and the corresponding parameters of SHS2 at the time of T2

Altitudo		The time of T1			The time of T2	
(lum)	Latitude	Longitude	Azimuth	Latitude	Longitude	Azimuth
(KIII)	(°)	(°)	angle (°)	(°)	(°)	angle (°)
21	54.501	121.483	321.98	55.071	121.534	321.041
31	54.501	121.483	321.98	55.117	121.303	321.045
41	54.501	121.483	321.98	55.163	121.073	321.050
51	54.501	121.483	321.98	55.210	120.844	321.056
61	54.501	121.483	321.98	55.255	120.616	321.062
71	54.501	121.483	321.98	55.31	120.389	321.070
81	54.501	121.483	321.98	55.346	120.163	321.078

The intersection region of the time of T1 and T2 mentioned above is (67.46 °N,86.618 °E). The observed radiance received by SHS1 at the time of T1 and received by SHS2 at the time of T2 is composed by a group of different satellite positions at tangent heights which is shown in Fig. 9.



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**Figure 9.** The simulated observation data in the intersection region at the time of T1 and T2. The dash line is the simulated observation data received by SHS1 at the time of T1 and the solid line is the simulated observation data received by the SHS2 at the time of T2.

375 Although the observation geometries and satellite positions are different at different time, both spatial heterodyne spectrometers scan the atmosphere by the same limb mode. The observed radiance profile still maintains the distribution

in the range from 308 to 310 nm band. There are four global observed radiance databases for this research. The first quarter observed radiance is given as an example: the NSO solar spectrum is used as the radiance source, the Bremen global atmospheric model, the first quarter OH concentrations data of N32-seasonly OH concentrations database and the geometric parameters of MLS OH production are also used to simulate the observed radiance in the range from 308 to 310 nm band at the time of T1 and T2. The results are shown in Fig. 10 which the observed radiance received by SHS1 at the time of T1 and Fig. 11 which the observed radiance received by SHS2 at the time of T2.



Figure 10. The simulated observation radiance received by SHS1 at the time of T1 under the different conditions



390 Figure 11. The simulated observation radiance received by SHS2 at the time of T2 under the different conditions

There are some differences between the Fig. 10 and the Fig. 11 because of the observed geometries and satellite positions. However, the whole observed radiance trends are same. The observed radiance increases from  $8 \times 10^{13}$  phot<sup>-1</sup> s<sup>-1</sup> cm<sup>-2</sup> nm<sup>-1</sup> sr<sup>-1</sup> to  $1.4 \times 10^{14}$  phot<sup>-1</sup> s<sup>-1</sup> cm<sup>-2</sup> nm<sup>-1</sup> sr<sup>-1</sup> until the altitude around 45 km and decreases to about  $4 \times 10^{13}$  phot<sup>-1</sup> s<sup>-1</sup> cm<sup>-2</sup> nm<sup>-1</sup> sr<sup>-1</sup> with the altitudes increase. The lowest observed radiance is in the region around the 30 °N and gradually increases towards the two poles at about 41 km. The reason for this trend is that the solar zenith angle is the minimum value around the 30 °N and then increases towards the two poles. At the 71 km, the observed geometries are not the only influencing factor and the distributions of OH concentrations are another one. The Rayleigh scattered radiance is weak and OH fluorescence emission radiance becomes strong. In addition to this one, the OH concentrations in the first quarter are low in the region around 60 °N that make the high radiance values appear in the southern hemisphere.

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## **4 Inversion modeling**

## 4.1 Interferogram pretreatments

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The method of forming the interferogram by the spatial heterodyne spectroscopy is the spatial heterodyne modulation. The signal-to-noise ratio is susceptible to many factors, and the fine spectral lines of the noisy signal can also be detected simultaneously. The interferogram needs to be pretreated for solving these problems and transformed to obtain the OH fluorescence emission radiance. The whole pretreatments are shown in Fig. 12.



#### 410 **Figure 12.** Flow chart of the interferogram pretreatments.

First, the interference data with low-frequency baseline will make low-frequency spurious signals in the process of the Fourier spectrum transform. The methods of removing the baseline mainly include the polynomial linear fitting to baseline, the first-order differential de-baseline at present. The function of the first-order differential de-baseline method is the high-pass filtering. It is the most useful way to remove the baseline (Ye et al., 2009). The interferogram obtained 415 by the spatial heterodyne spectrometer is the interference data in the interval of finite optical path difference from -L to +L. That means an interferometric function forces a sudden drop to zero outside of this range. It will cause the interferogram to appear sharp discontinuities in the edge interval. The spectral profile has side lobes in which the positive side lobes are the source of the false signal, and the negative side lobes will lead to the adjacent weak spectral signals to be submerged. The apodization method is used to mitigate the discontinuity of the interferogram edge through 420 multiplying the interferogram function by a progressive weight function. The interferogram after apodization is subjected to the Fourier spectrum transform for obtaining the corresponding spectrum. However, the high-spectral resolution observed radiance data which is simulated by the forward model mentioned above includes the atmospheric background radiance. It must be identified and removed from the observed radiance data before the OH fluorescence emission 425 radiance is obtained because: (1) the atmospheric background radiance accounts for more than 95% of observed radiance

- that will make OH fluorescence emission radiance low, (2) some errors of atmospheric background radiance will transfer
  - 19

into the OH fluorescence emission radiance when the observed radiance of the interferogram form is recovered to the spectral form. The intensity of simulated observation radiance and atmospheric background radiance at some tangent heights are given in Table 1. The OH fluorescence emission radiance can be calculated by subtracting the atmospheric background radiance from the observed radiance. The OH fluorescence emission radiance graph which is shown in Fig.

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13 can be obtained after these steps and the Fourier spectrum transform can be done.



#### 435 **Figure 13.** Graph of OH fluorescence emission radiance.

The fifth step is the wavelength calibration. The zero position of the spectral data points correspond to the Littrow wavelength number in the Fourier transform spectrum, so the wavelength calibration equation is given by the Eq. (5):

440 
$$\delta_i = -1.3166574i + 32679.738$$
 (5)

where *i* is from 0 to 1024. The  $\delta_i$  is the symbol of intensity and the unit of it is phot<sup>-1</sup>s<sup>-1</sup>cm<sup>-2</sup>nm<sup>-1</sup>sr<sup>-1</sup>. The sixth step is the radiometric calibration. The purpose of the radiometric calibration is to determine the quantitative relationship between the output signal of instrument and the spectral radiance because the spectrometer obtains the DN values directly. A liner fit method is used to simulate the different sets of known radiance values at each spectral data point for establishing a fitting relationship in the calibration process by Eq. (6):

$$S_{\delta} = B_{\delta} \cdot K_{\delta} + \varepsilon_{\delta} \tag{6}$$

450 where  $S_{\delta}$  is the DN values at the wavelength number  $\delta$ .  $B_{\delta}$  is the assumed incident spectrum radiance.  $K_{\delta}$  is the calibration factor and  $\varepsilon_{\delta}$  is the deviation caused by other factors. A spectrum of OH fluorescence emission radiance is shown in Fig. 14.





Figure 14. OH fluorescence emission radiance spectrum extracted from observed radiance.

## 4.2 Constructions of the retrieval algorithm

simulated through the forward model in this research. However, it cannot be done to obtain OH concentrations in an
 inverse way directly. The atmospheric inversion problems are not an inversion of the forward model but use some related algorithms to estimate the atmospheric parameters for finding the best state parameters through the observed radiance data. The DSHS consists of two spatial heterodyne spectrometers in the orthogonal layout for monitoring the OH from 15 to 85 km. It will provide large and complicated data. The traditional iterative retrieval algorithm will take more time to obtain the OH concentrations in this case because of the character of iteration. A new retrieval algorithm that is suitable for the DSHS needs to be constructed for inverting the accurate OH concentrations.

465 for the DSHS needs to be constructed for inverting the accurate OH concentrations. The precondition of the three-dimensional limb mode for monitoring the OH is that the time interval must be relatively short. Otherwise, the OH concentrations will change. The observed data cannot reflect the accurate OH concentrations. The distance between the satellite positions at the time of T1 and T2 is approximately 66 km when the intersection point is assumed as 50 km away from the tangent point mentioned in the Sect. 2. So, the intervals between those points are

It is an inversion process to obtain the OH concentrations by the DSHS observed radiance. The observed radiance is

- 470 very small. Additionally, the OH concentrations do not change much in a very short time according to MLS products. The short interval time between the T1 and T2 and small change of OH concentrations in a very short time support the theoretical premise of the new retrieval algorithm and make DSHS possible to detect the atmosphere by the threedimensional limb mode. An algorithm called tomographic retrieval algorithm is constructed to invert OH concentrations from 15 to 85 km based on the theory above. The core of this retrieval algorithm is the lookup table method. The most
- 475 important step for it is the construction of the tomographic observed database. The main influence factors for making up

the tomographic observed database contain the atmospheric model, the observation geometries (the solar zenith angle and the relative azimuth angle), the spatial information (position of the target area and satellite position), date parameters, the OH concentrations, and the simulated observation data. The solar zenith angle and the relative azimuth angle have a greater influence on the observed radiance. The spatial information and date parameters will affect the OH concentrations

- 480 and the atmospheric model. The solar zenith angle is changed between  $0^{\circ}$  to  $100^{\circ}$  and the relative azimuth angle is changed between  $0^{\circ}$  to  $180^{\circ}$ . The corresponding OH concentrations are found out according to the date parameters and spatial information such as the season, latitude, and longitude of the target area from the OH concentrations database which has been built up in the Sect.3. The OH concentrations are changed in the ranges between 50% to 150% times amount of the original corresponding one at different tangent height for more possible situations. Many combinations of
- 485 various state parameters can be generated according to these changes. The observed data received by DSHS in these combinations is simulated through the forward model mentioned above. The tomographic observed database can be built up for the tomographic retrieval algorithm. Each parameter that constitutes the database is accurate and the errors of them are analyzed in Table 3. The method of constructing the database is reasonable which is shown in Fig. 15. Table 5 gives an example about subset of rows of the part tomographic observed database. The SZA is the solar zenith angle, the 490 AA is the relative azimuth angle, the MultiND height is the tangent height.



Figure 15. Construction of the tomographic observed database.

N	Latitude	Longitude	SZA	AA	MultiND_height	Secon
IN	(°)	(°)	(°)	(°)	(km)	Season
1	54.240	125	0	0	15	1
2	54.240	125	0	0	15	2
3	54.240	125	0	0	15	3
4	54.240	125	0	0	15	4
5	54.240	125	0	0	17	1
6	54.240	125	0	0	17	2
7	54.240	125	0	0	17	3

Based on the satellite positions, the observation geometries, the date parameters, the observed radiance at all wavelengths

of SHS1 at the time of T1 and the corresponding observed radiance at all wavelengths of SHS2 at the time of T2, the OH concentrations in the target area can be found in the tomographic observed database directly. The tomographic observed database is effective and reasonable enough. However, it is impossible to include all combinations in the tomographic observed database because 1) the size of tomographic observed database is limited and 2) all situations cannot be considered when the sensor will work in the orbit. The interpolation methods and the threshold judgment methods are always used to solve such problems. A scientific and precise threshold in the threshold judgment method can be obtained according to a lot of OH concentrations data and experiments. The OH concentrations data are little now. So, if there is no corresponding query condition in the database, the cubic spline interpolation is used to calculate the OH concentrations. The cubic spline interpolation not only has higher stability than other interpolation algorithms but

also can maintain the continuity and smoothness of the interpolation function under the premise of convergence. Figure

16 shows the flow chart that the OH concentrations obtain by the tomographic retrieval algorithm.

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**Figure 16.** Flow chart of the tomographic retrieval algorithm. The OH concentrations can be obtained from the tomographic observed database or by the cubic spline interpolation algorithm.

## **5** Results and discussion

## 515 **5.1 Inversion results**

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There are four tomographic observed databases based on the four seasons. The profile data in the database are averaged to obtain the monthly average concentration profile data. So, it will not introduce discontinuities to the retrieved OH time series when an abrupt switch happens. Here, the spring season tomographic observed database is used as an example to prove the feasibility and superiority of the tomographic retrieval algorithm. The OH concentrations are obtained in the (67.46 °N, 86.168 °E) through satellite positions, observation geometries, date parameters which are given in Table 5. If there is a corresponding query condition, the OH concentrations can be given directly from the tomographic observed database. Otherwise, the OH concentrations will be calculated by the cubic spline interpolation. The inversion



Figure 17. Inversion results in the target area using the tomographic retrieval algorithm.

530

around 45 km which closely relates to the ozone concentrations in the stratopause as the heights increase. A valley value of OH concentrations about  $5.0 \times 10^6$  molecules cm<sup>-3</sup> appears around 65 km. As the height continues to rise, the second maximum peak value of OH concentrations about 1.0×107 molecules cm<sup>-3</sup> appears at around 70 km which is mainly affected by the water vapor concentrations in the mesosphere. The OH concentrations above the 75 km and below the 25 km is very low (about 2.5×10<sup>6</sup> molecules cm<sup>-3</sup>). In general, the OH concentrations increase with the altitudes rise until around 40 km and reach the valley value at around 65 km. The second maximum value peak which is affected by 535 the water vapor concentrations appears at around 75 km, then the OH concentrations continue to decrease until the limited altitudes of detection.

The results indicate that a maximum peak value of OH concentrations is about  $2.0 \times 10^7$  molecules cm<sup>-3</sup>. It appears at

### 5.2 Discussion

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We also constructed an iterative retrieval algorithm to compare with the tomographic retrieval algorithm here for reflecting the outstanding advantages of the tomographic retrieval algorithm. The schematic diagram of iterative inversion is shown in Fig. 18. The simulated radiance can be obtained under the condition which an initial OH concentrations guess can get from the OH concentrations database. The residuals of the simulated and observed data will be calculated to judge the convergence. The initial OH concentrations guess values are the inversion results if the residual meets the accurate requirements of research. Otherwise, the retrieval algorithm will be used to obtain the inversion results.

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Figure 18. Schematic diagram of iterative inversion.

550 The LSUV (Limb scan of Scattered UV radiation) retrieval algorithm (Aruga and Heath, 1988;Aruga and Igarashi, 1976) is modified to invert OH concentrations at each tangent height. The tangent height is defined as *m* and the target area is divided into *n* atmospheric layers from the bottom to the top. One-order approximate Taylor expansion is used to establish a linear relationship between the OH concentrations and observed radiance at each atmospheric layer by Eq. (7):

555 
$$F_i = \left[\sum_{j=1}^n D_{ij} \cdot y_j\right] \varepsilon_i \quad \begin{cases} i = 1, \dots, n \\ j = 1, \dots, m \end{cases}$$
(7)

where  $\varepsilon_i$ , the inversion coefficient, is approximately 1 when the OH concentrations are closed to the guess OH concentrations. The  $F_i$  is the difference between the observed data and simulated data by Eq. (8) at the *i* layer:

560 
$$F_i = I(h_i) - I_s(h_i), i = 1, 2, ..., m$$
 (8)

The  $D_{ij}$  which is given by Eq. (9) is the partial derivative of radiance data at the *i* tangent height:

$$D_{ij} = \left[\frac{\partial F_i(y_j)}{\partial y_j}\right]_{y_j=0} = \frac{I_s^* \left(h_i, \frac{\Delta y_j}{2}\right) - I_s^* \left(h_i, -\frac{\Delta y_j}{2}\right)}{\Delta y_j}$$
(9)

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The  $y_j$  in Eq. (10) is the relative error between the OH concentrations N(j) which can get through the  $y_j$  and  $N_g(j)$  when the converges meet the accurate requirement or the number of iterations exceeds the threshold of iteration number and the guess concentrations  $N_g(j)$ :

570 
$$y_j = \frac{[N(j) - N_g(j)]}{N_g(j)}$$
  $j = 1, 2, ..., n$  (10)

The approximate OH concentrations at each tangent height  $(Q)_i$  can be got based on Eq. (10) through Eq. (11):

$$(Q)_i = \frac{F_i}{\sum_{j=1}^n D_{ij}} \tag{11}$$

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The approximate OH concentrations  $(Q)_i$  at *i* tangent height is determined by the OH concentrations on several atmospheres near the *i*. The weight function  $P_{ij}$  is established from the  $D_{ij}$  by Eq. (12):

$$P_{ij} = \frac{D_{ij}}{\sum_{j=1}^{n} D_{ij}}$$
(12)

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where  $P_{ij}$  is the proportion of the contribution of OH concentrations on the radiance of each tangent height in each layer. The accurate relative errors of the OH concentrations and guess OH concentrations in each layer  $y_j$  can be obtained from  $P_{ij}$  by Eq. (13):

585 
$$y_j = \frac{\sum_{i=1}^m Q_j \cdot P_{ij}}{\sum_{i=1}^m \cdot P_{ij}}$$
 (13)

However, the difference between the OH concentrations and guess OH concentrations is large in the actual satellite detection, the  $\varepsilon_i$  is not 1, the  $\varepsilon_i^*$  is used to replace it by Eq. (14):

590	$\varepsilon_{i}^{*} = \frac{F_{i}(y_{1}^{*}, y_{2}^{*}, \dots, y_{n}^{*})}{\sum_{j=1}^{n} D_{ij} y_{j}^{*}}$	(1	4)
	,		

Finally, the precise relative error  $y_j$  between the OH concentrations and the guess OH concentrations can be obtained. The OH concentrations N(j) can get through the  $y_j$  and  $N_g(j)$ . The convergence error  $R_i^l$  between the observed data  $I(h_i)$  and simulated data  $I_s(h_i)$  at *i* layer which based on the guess OH concentrations  $N_g(j)$  is calculated by Eq. (15):

 $1(v_i)$  and  $2minimized a minimized by <math>2q_i(1)$ 

$$R_{i}^{l} = \frac{I(h_{i}) - I_{s}^{(l)}(h_{i})}{I_{s}^{(l)}(h_{i})} = \frac{F_{i}^{(l)}}{I_{s}^{(l)}(h_{i})}$$
(15)

where *l* is the number of iterations. The average residual  $R^l$  based on the  $R_i^l$  at different tangent heights can be calculated by Eq. (16):

600

$$R^{l} = \sqrt{\frac{\sum_{i=1}^{m} \left[R_{i}^{l}\right]^{2}}{m}}$$
(16)

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The threshold of average residual and the numbers of iteration are set based on the inversion experience. The iteration result which meets the precision requirement is the final result, otherwise the prior  $N_g(j)$  will be replaced by the N(j) which is calculated by the inversion algorithm and the iteration process will continue until the residual converges meet the accurate requirement or the number of iterations exceeds the threshold of iteration number. The iteration stops when the average residual decreases below 0.005 or the number of iterations reaches 15 times in this research.

The observed radiance which is simulated by the forward model are used by the LSUV retrieval algorithm to invert OH concentrations. When the iteration results meet the accurate requirements of research, the OH concentrations can be obtained. The OH concentrations obtained by the LSUV retrieval algorithm are credible in the upper atmosphere which are the same as the results of MAHSRI and SHIMMER (Conway et al., 1999;Englert et al., 2010). However, the OH concentrations in the lower atmosphere such as below 30 km are unsuitable for scientific research because the interference factors like water vapor and ozone are too much. A single spatial heterodyne spectrometer with traditional limb mode cannot also obtain enough and high-quality data to invert the OH concentrations that will make the inversion

615 results unscientific especially in the lower atmosphere. That is the reason why there are no OH concentrations in these regions from the MAHRSI and SHIMMER (Harlander et al., 2002). The tomographic retrieval algorithm and the LSUV are two independent algorithms. The tomographic retrieval algorithm

effectively avoids the constraints of the initial guess values and does not take a lot of time to iterate for obtaining the results in the inversion progress. The errors of this retrieval algorithm just include the errors caused by the interpolation method and the observed radiance errors which are used in this algorithm. The observed radiance errors are affected by

- 620 method and the observed radiance errors which are used in this algorithm. The observed radiance errors are affected by the atmospheric model, the Doppler effect and the instrument calibration errors. The OH fluorescence emission radiance which is separated from the observed radiance is the most important part for the OH concentrations, so the contributions of each part to the OH fluorescence emission radiance is analyzed by the following method. Some certain amounts as the disturbances are applied to the three parts above. The OH fluorescence emission radiance obtained based on these
- 625 parts is considered as the incorrect results. The original parts are taken as the actual and undisturbed parameters, the results based on these parts are the true OH fluorescence emission radiance. The relative errors between the true and

incorrect OH fluorescence emission radiance are calculated by Eq. (17):

$$Err = \frac{[I]_{err} - [I]_{act}}{[I]_{act}} \times 100\%$$

$$(17)$$

630

where the  $[I]_{err}$  is the incorrect OH fluorescence emission radiance and the  $[I]_{act}$  is the true OH fluorescence emission radiance.

### 5.2.1 Influence of the atmospheric model

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The atmosphere environment is complicated. Although, the atmospheric model used in this research is scientific and reasonable, it cannot reflect the accurate atmospheric condition which leads to some errors between true OH fluorescence emission radiance and incorrect OH fluorescence emission radiance. These will cause an error when the OH concentrations are looked up or calculated. Many factors can affect the OH fluorescence emission radiance in the atmospheric model such as some physical parameters like temperature, pressure and some atmospheric composition like nitrogen dioxide and formaldehyde. Here, the main composition we consider is ozone which is a substance that plays a 640 major role in the process of radiation transmission as solar energy in the ultraviolet band and is the main source and sink of OH radicals. The other relevant source of uncertainty will be considered when the DSHS works on the orbit in the following research. The original ozone profiles are assumed as the actual atmospheric state parameter and the amount of ozone profile for the disturbances is set as  $\pm 30\%$  (Conway et al., 2000). The relative errors of the OH fluorescence emission radiance caused by the atmospheric model are calculated by Eq. (17) and are shown in Fig. 19. The relative 645 errors of the OH fluorescence emission radiance at some tangent height are also given in Table 6.



Figure 19. Relative errors of OH fluorescence emission radiance caused by the atmospheric model in the tomographic retrieval algorithm. The red error bars indicate the relative errors caused by the atmospheric model.

Tangent height (km)	Relative errors caused by the atmospheric model (%)
81	From -2.71 to 5.72
71	From -2.72 to 5.75
61	From -2.96 to 5.92
51	From -4.92 to 8.19
41	From -12.57 to 20.92
31	From -6.78 to 10.98
21	From -6.00 to 9.51

The relative errors caused by the atmospheric model indicate that the atmospheric model has a strong impact on the OH fluorescence emission radiance especially in the lower atmosphere where the ozone concentrations are high. The errors caused by  $\pm 30\%$  uncertainty of the atmospheric model have a peak value from -12.57 % to 20.92 % at the 41 km. It also appears by the same calculation method which is from -12.19 % to 20.28 % in the LSUV retrieval algorithm. The concentrations of ozone gradually decrease with the relative errors decreasing to about -3% to 5% when the heights increase. The effect of atmospheric model is smaller in the upper atmosphere. The distribution of errors has the same tendency in the LSUV retrieval algorithm that the relative errors increase and then gradually decrease after reaching the 660 peak value of altitude. These mean the relative errors caused by the atmospheric model are inevitable. If the more actual atmospheric model is used, the more accurate OH fluorescence emission radiance may be obtained which is the key for the retrieval algorithm.

### **5.2.2 Influence of the Doppler effect**

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The satellite has different heliocentric speeds on its working orbit around the earth at different times. It will cause a pseudo-random offset of observed data within  $\pm 5 \times 10^{-4}$  nm at the range of wavelength. The observed radiance is calculated within  $\pm 5 \times 10^{-4}$  nm disturbance at the wavelength. The OH fluorescence emission radiance which is separated from the observed radiance is too weak.

A little Doppler effect will cause a great influence on the weak OH fluorescence emission radiance. The magnitude of wavelength shift caused by the Doppler effect is much smaller than the spectral resolution. "Finding peak" is used to

find a peak value which needs to be larger than the values on its sides. It is impossible to correct the offset by "Finding 670 Peak" method. However, "Taking peak" is defined to find the highest peak value in the OH fluorescence emission

radiance. The "Taking Peak" is used for reducing the impact of the Doppler effect. Therefore, the OH fluorescence

emission radiance at the peak position is used instead of the wavelength from 308 nm to 310 nm when calculating the relative errors caused by the Doppler effect at a single tangent height. The relative errors of the OH fluorescence emission

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radiance caused by the Doppler effect are calculated by Eq. (17) and are shown in Fig. 20. The relative errors of OH fluorescence emission radiance caused by the Doppler effect at some tangent heights are also given in Table 7.



680 Figure 20. Relative errors of OH fluorescence emission radiance caused by the Doppler effect in tomographic retrieval algorithm. The red error bars indicate the relative errors caused by the Doppler effect.

Tangent height	Relative errors caused by the
(km)	(%)
81	From -0.39 to 0.35
71	From -0.20 to 0.31
61	From -0.11 to 0.34
51	From -0.88 to 0.41
41	From -2.16 to 0.93
31	From -3.14 to 1.44
21	From -3.71 to 1.62

Table 7. Relative errors of OH fluorescence emission radiance caused by the Doppler effect at some tangent heights

The relative errors of OH fluorescence emission radiance caused by the Doppler effect decrease when the altitudes rise. 685 The tendency of relative errors is same, but the relative errors of the tomographic retrieval algorithm are smaller than

the LSUV retrieval algorithm. These indicate that the tomographic retrieval algorithm can obtain more accurate OH concentrations than the LSUV retrieval algorithm although the errors caused by the Doppler effect cannot avoid when the spectral resolution of DSHS is extremely high.

### **5.2.3 Influence of other factors**

Some other errors exist apart from the errors above. For one thing, there are various sources make some errors between the radiance received by the sensors and the actual radiance in the instrument calibration process. The instrument calibration errors will cause an error of the inversion result. We assumed the instrument calibration errors are  $\pm 5\%$  in this research after referring to the detector parameters due to the DSHS do not officially work (Ye et al., 2009). For 695 another, the error caused by the interpolation method is the main factor we need to calculate. The cubic spline interpolation method not only has higher stability but also can ensure the continuity and smoothness of the interpolation function under the premise of ensuring the convergence compared with other interpolation methods like the linear interpolation method and some others. The error estimation of the interpolation results is as accurate as possible. The OH concentrations can be obtained by the tomographic retrieval algorithm according to the atmospheric model, spatial 700 information, date parameters, observation geometries and observed radiance. The interpolation errors are also the other error factors which affect the accuracy of OH concentrations. We give a certain disturbance to different parameters for calculating the effect of the interpolation algorithm on the OH concentrations. The solar zenith angle is changed between  $0^{\circ}$  to  $100^{\circ}$  and the magnitude perturbation is  $10^{\circ}$ . The relative azimuth angle is changed between  $0^{\circ}$  to  $180^{\circ}$  and the magnitude perturbation is also 10°. The OH concentrations are changed in the ranges between 50% to 150% times amount 705 of the original corresponding one at different tangent height for more possible situations and the magnitude perturbation is 10%. The date parameters, longitude, and latitude can determine the atmospheric model. So, the influence of date parameters, longitude, and latitude is calculated through disturbance of the ozone concentrations and the magnitude perturbation is the 30% times amount of the original corresponding one. It is about 0.32% for the six-dimensional cubic spline interpolation of OH concentrations, date parameters, longitude, latitude, solar zenith angle, and azimuth angle 710 after research and related experiment. Therefore, 0.32% is taken as the error of the interpolation algorithm in the tomographic retrieval algorithm.

## 5.2.4 Analysis of total errors in the OH concentrations

The errors of retrieved results must be considered after the discussions about the errors of OH radiance. The theoretical basis for calculating the total errors of inversion results is the error transfer formula which is always used to calculate the indirect measurement error (Gao et al., 2009). The total errors based on a single error factor often have an asymmetric distribution. The corresponding standard error according to the single each error factor should be calculated if the error transfer formula is used. It is calculated according to the B-class standard uncertainty evaluation method. The relationship

between input parameters and results can be expressed by Eq. (18):

720 
$$[I] = f(x_1, x_2, x_3, \dots, x_n)$$
 (18)

where  $x_1, x_2, x_3, ..., x_n$  are the input parameters and [I] indicates the inversion results. Equation (19) can be got when the error  $\Delta x_i$  is considered and then does a Taylor expansion:

725 
$$[I] + \Delta[I] = f(x_1, x_2, x_3, ..., x_n) + \sum_{i=1}^n \frac{\partial f}{\partial x_i} \Delta x_i$$
 (19)

The maximum value of relative error can be calculated by Eq. (20):

$$\frac{\Delta[I]}{[I]} = \sum_{i=1}^{n} \left| \frac{\partial f}{\partial x_i} \right| \frac{\Delta x_i}{[I]}$$
(20)

730

where the right part of equal sign means the errors caused by  $x_1, x_2, x_3, ..., x_n$ . The error transfer formula can be derived from Eq. (20) when the errors are squared. The total errors RSS can be calculated by the "square root" method with Eq. (21):

735 
$$RSS = \sqrt{\sum_{i=1}^{n} \left\{ \frac{\partial[I]}{x_i} \frac{\Delta x_i}{[I]} \right\}^2}$$
(21)

The total errors of the results caused by each factor can be obtained by Eq. (21). The total errors of the inversion results caused by each error factor are calculated which are shown in Fig. 21 and the total errors of inversion results at some tangent heights are given in Table 8.



Figure 21. Total errors of inversion results caused by some factors mentioned above. The red error bars indicate the relative errors caused by some factors mentioned above.

Tangent height (km)	Total errors of inversion results (%)
81	From -8.09 to 8.09
71	From -8.11 to 8.11
61	From -8.31 to 8.31
51	From -10.83 to 10.83
41	From -25.03 to 25.03
31	From -14.27 to 14.27
21	From -12.96 to 12.96

745 Table 8. Total errors of inversion results at some tangent heights

As Table 8 indicates, the total errors of OH concentrations initially increase with the heights rising. It reaches the maximum value range from -25.03 % to 25.03 % at the 41 km and then decreases from -8.09 % to 8.09 % as the heights continue to rise until the limited height. The total errors of the inversion results in the lower atmosphere are small 750 especially. However, the OH concentrations obtained by the LSUV algorithm in the same area are unsuitable for scientific research mentioned in the Sect. 5.2. It is a great improvement to compare with the results of the LSUV retrieval algorithm which has a characteristic of iteration. The relative errors of the OH concentrations obtained by the LSUV algorithm are calculated by Eq (17) and the error transfer formula mentioned above. The initial guess values have a great influence on the iterative speed and the inversion results for the iterative retrieval algorithm. A realistic initial guess value can achieve higher iteration efficiency and more accurate inversion results. The initial guess values used in the LSUV retrieval

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algorithm are obtained from the OH concentrations database through the date parameters and spatial information such as the season, latitude, and longitude of the target area. The corresponding accuracy ranges and useful height ranges of MLS OH concentrations products are also given by the MLS data quality and description document (Livesey et al., 2015). Therefore, the data accuracy is assigned at the upper limited height to an altitude greater than the upper limit of

- 760 MLS data altitude and at the lower limited height to the altitude smaller than the lower limit of MLS data altitude. The initial guess values can be obtained by using the data accuracy to perform the positive and negative turbulence for the initial OH concentrations values. A minimum positive value is given to the data less than zero for the physical meaning when a negative value may occur in the negative direction. The two initial guess values are used in the LSUV retrieval algorithm separately. Although, the initial guess values above are chosen scientifically and reasonably, the inversion
- 765 results are unsuitable for scientific research in the lower atmosphere that the relative errors caused by the initial guess values reach from -95.39 % to 545.44 %. Besides the initial guess values, the inversion results are also affected by the errors caused by the atmospheric model, the Doppler effect and the instrument calibration error. The relative errors of inversion results obtained by the LSUV retrieval algorithm are small from 30 km to 85 km. The total errors of inversion results are from -24.83 % to 24.83 % at the 31 km and then decrease gradually until the altitude of 51 km. A little increase
- occurs from the altitude of 60 km to 70 km and reaches from -37.01 % to 37.01 % at the 81 km. However, the inversion results are bad in the lower atmosphere. The total errors of inversion results are from-553.16 % to 553.16 % at the 21 km. The total errors of inversion results are 2 times larger than them by tomographic retrieval algorithm at some heights like 71 and 61 km. However, the total errors of inversion results are 4.625 times larger than the errors of the tomographic retrieval algorithm at the 21 km especially. The application altitudes of the LSUV retrieval algorithm are limited. The
- 775 main reasons lead to these unscientific inversion results are two parts. The OH fluorescence emission radiance, the main factor for the retrieval algorithm, is not strong enough due to the ozone optical depth becomes large in the lower atmosphere. That will lead to the observed radiance does not come from the tangent area but the near field. The lower atmospheric environment is complicated. The human activity will produce a lot of substances which will join in the atmospheric environment by the atmospheric convection motion. The aerosol also distributes here. These will lead to a
- 780 bad accuracy of detection. So, the useful altitudes of the LSUV retrieval algorithm based on the data of single spatial heterodyne spectrometer are limited. The DSHS can counteract the errors by the two spatial heterodyne spectrometers because of its special optical design which the inversion results of the retrieval algorithm have proved. These analyses above also indicate that the tomographic retrieval algorithm is a feasible retrieval algorithm for the atmosphere from 15
- to 85 km. The lookup table method avoids the intervention of initial guess values effectively by establishing a
  tomographic observed database of multi-dimensional variables and is understood easily. It also avoids the complicated iterative optimization process, and the OH concentrations can be obtained directly from the tomographic observed database. These make the speed of inversion process fast compared with the LSUV retrieval algorithm. The tomographic retrieval algorithm is a suitable retrieval algorithm for the three-dimensional limb mode of DSHS as well. The three-dimensional limb mode provides big and numerous observed data. The speed of traditional iterative retrieval algorithms is slow and will cost a lot of time. However, it is difficult to quantify how much fast the tomographic algorithm is
- 790 is slow and will cost a lot of time. However, it is difficult to quantify how much fast the tomographic algorithm is absolutely. The traditional iterative retrieval algorithm will take several or even tens of hours to obtain OH concentrations due to the complex iterative process. The tomographic algorithm just needs minutes for obtaining the OH concentration
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due to the usage of the lookup table method. The results obtained by the traditional iterative retrieval algorithm in the lower atmosphere are unscientific but the OH concentrations obtained by the tomographic algorithm in the same area

- are accurate. The tomographic retrieval algorithm solves these problems above and improves the efficiency of retrieval algorithm which is important for the OH concentrations. The LSUV retrieval algorithm can invert OH concentrations well under the conditions that the OH fluorescence emission radiance is strong and the interference factors like water vapor and ozone are little. However, a single spatial heterodyne spectrometer with traditional limb mode cannot obtain enough and high-quality data to invert the OH concentrations that will make the inversion results unscientific especially
- 800 in the lower atmosphere. The initial guess values play a significant role in the LSUV retrieval algorithm due to the feature of the iteration. The determination of the initial guess values involves many factors and there is no best way to determine the initial guess values at present. This feature is particularly pronounced in the lower atmosphere where OH fluorescence emission radiance is less sensitive to OH concentrations.
- In summary, the LSUV retrieval algorithm based on the iterative method can invert the OH concentrations well in the 805 higher atmosphere like mesosphere but cannot obtain the accurate OH concentrations in the lower atmosphere like the bottom of stratosphere. Many factors lead to this result like the atmospheric model, the limit of initial guess values and something else. The tomographic retrieval algorithm using the lookup table method can obtain the accurate OH concentrations from stratosphere to mesopause through the feature of the three-dimensional limb mode. This retrieval algorithm not only saves the time of the inversion process but also solves the problem that the OH concentrations are unsuitable for scientific research in the lower atmosphere.

## **6** Conclusions

The OH is the key oxidant in the atmosphere and has a great influence on the atmospheric photochemistry process. The DSHS based on the spatial heterodyne spectroscopy will monitor the OH with the three-dimensional limb mode in the future. A forward model is constructed to simulate the observed data of DSHS accurately. A new retrieval algorithm for

- 815 obtaining the OH concentrations is also proposed based on the simulations of the forward model. The distinctive feature of this algorithm is the usage of a lookup table method. The MLS OH concentrations products and N32 Gaussian grid are used to construct the OH concentrations database of four seasons. The observed radiance is obtained by the forward model. The other factors like spatial information and observation geometries are also simulated according to the characteristic of DSHS for the tomographic retrieval algorithm. The tomographic observed database is established based
- 820 on the parameters above. The OH concentrations in the target area are obtained through finding in the tomographic observed database directly. The cubic spline interpolation method is also used to obtain the OH concentrations without the corresponding query conditions. The errors are also analyzed caused by the atmospheric model, the instrument calibration error, the Doppler effect, and the interpolation algorithm. The results show that the tomographic retrieval algorithm is not only faster compared with the LSUV retrieval algorithm but also improves the inversion precision of the OH concentrations especially in the lower atmosphere.

There are still some problems to be solved when the DSHS will officially work in the future. The performances of two

spatial heterodyne spectrometers are inevitably different because they are affected by the temperature, humidity, and electromagnetic environment during manufacturing although the bi-orthogonal structure is theoretically identical in the design parameters. The instrument calibration errors of the instrument are only considered to be 5% at present. This part

830 will be optimized according to the actual working condition after the instrument officially works. The actual observed data is the key point for this research. Some aircraft flight experiments have been done and the data is being analyzed.

Code/Data availability. All code and data can be obtained from the corresponding author upon request.

835 **Author contribution.** YA and JM designed the study. YA and GB performed the simulations and carried out the data analysis. JM, WX, and XW provided useful comments on the paper. YA prepared the manuscript with contributions from all co-authors.

Competing interests. The authors declare that they have no conflict of interest.

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