Ozone and water vapor variability in the polar middle atmosphere observed with ground-based microwave radiometers

Guochun Shi1,2, Witali Krochin1,2, Eric Sauvageat1,2, and Gunter Stober1,2

1 Institute of Applied Physics, University of Bern, Bern, Switzerland
2 Oeschger Center for Climate Change Research, University of Bern, Bern, Switzerland

Correspondence: Guochun Shi (guochun.shi@unibe.ch)

Abstract. Leveraging continuous ozone and water vapor measurements with the two ground-based radiometers GROMOS-C and MIAWARA-C at Ny-Ålesund, Svalbard (79° N, 12° E) that started in September 2015 and combining MERRA-2 and Aura-MLS datasets, we analyze the interannual behavior and differences in ozone and water vapor and compile climatologies of both trace gases describing the annual variation of ozone and water vapor at polar latitudes. A climatological comparison of the measurements from our ground-based radiometers with reanalysis and satellite data was performed. Overall differences between GROMOS-C and Aura-MLS ozone volume mixing ratio (VMR) climatology are mainly within ±7 % throughout the middle and upper stratosphere and exceed 10 % in the lower mesosphere (1–0.1 hPa) in March and October. For the water vapor climatology, the average 5 % agreement is between MIAWARA-C and Aura-MLS water vapor VMR values throughout the stratosphere and mesosphere (100–0.01 hPa). The comparison to MERRA-2 yields an agreement that reveals discrepancies larger than 50 % above 0.2 hPa depending on the implemented radiative transfer schemes and other model physics. Furthermore, we perform a conjugate latitude comparison by defining a virtual station in the Southern Hemisphere at the geographic coordinate (79° S, 12° E) to investigate interhemispheric differences in the atmospheric compositions. Both trace gases show much more pronounced interannual and seasonal variability in the Northern Hemisphere than in the Southern Hemisphere. We estimate the effective water vapor transport vertical velocities corresponding to upwelling and downwelling periods driven by the residual circulation. In the Northern Hemisphere, the water vapor ascent rate (5 May to 20 June in 2015, 2016, 2017, 2018, and 2021 and 15 April to 31 May in 2019 and 2020) is 3.4 ± 1.9 mm s⁻¹ from MIAWARA-C and 4.6 ± 1.8 mm s⁻¹ from Aura-MLS, and the descent rate (15 September to 31 October in 2015–2021) is 5.0 ± 1.1 mm s⁻¹ from MIAWARA-C and 5.4 ± 1.5 mm s⁻¹ from Aura-MLS at the altitude range of about 50–70 km. The water vapor ascent (15 October to 30 November in 2015–2021) and descent rates (15 March to 30 April in 2015–2021) in the Southern Hemisphere are 5.2 ± 0.8 and 2.6 ± 1.4 mm s⁻¹ from Aura-MLS, respectively. The water vapor transport vertical velocities analysis further reveals a higher variability in the Northern Hemisphere and is suitable to monitor and characterize the evolution of the northern and southern polar dynamics linked to the polar vortex as a function of time and altitude.
1 Introduction

Ozone and water vapor are essential climate variables that play a key role in the radiative balance in the middle atmosphere. Their seasonal and interannual variability is closely coupled to dynamical and chemical processes, which are driven and modulated by atmospheric waves including planetary waves, gravity waves (GWs), and atmospheric tides. These atmospheric waves transport energy and momentum from their source region to the altitudes of their dissipation and thus contribute to the energy balance between different atmospheric layers.

Model results suggest that GWs drive the summer mesopause temperature up to 100 K below the radiative equilibrium (Lindzen, 1981; Smith, 2012; Becker, 2012). The extreme cold temperatures at the summer mesopause are the result of an upwelling and a corresponding adiabatic cooling of the uplifted air masses in the summer hemisphere and are accompanied by a downwelling in the winter hemisphere. The pole-to-pole circulation is often referred to as residual circulation or transformed Eulerian mean circulation (Andrews and McIntyre, 1976). Another important circulation branch at altitudes lower than the residual circulation is the Brewer–Dobson circulation (BDC) (Brewer, 1949; Dobson, 1956). BDC is a factor in stratospheric ozone and water vapor variability, but, as mentioned later in this paper, polar stratospheric clouds can have significant seasonal impacts on ozone and water vapor abundances in the lower stratosphere. The circulation is fundamentally driven by dissipating waves of tropospheric origin and broadly consists of large-scale tropical ascent and winter pole descent. BDC is much weaker during boreal summer due to the different distribution of land masses and the associated differences in the generation of planetary and GWs between both hemispheres. BDC can govern the entry and distribution of air masses and constituents from the troposphere into and within the stratosphere. The meridional transport of trace gases into the polar cap is controlled by the strength of the polar vortex during winter, which is driven by the temperature gradient between the polar cap and the mid-latitudes through the thermal wind balance in the hemispheric winter stratosphere, and the vortex forms an essential barrier separating ozone-rich air at the mid-latitudes from ozone-depleted air within the polar cap. However, planetary waves can disturb the polar vortex and even lead to its breakdown during sudden stratospheric warming events (SSWs) (Matsuno, 1971; Baldwin et al., 2021), which is accompanied by a large-scale intrusion and mixing of air masses from the mid-latitudes towards the high latitudes, helping to recover the ozone volume mixing ratio (VMR) (Schranz et al., 2020). Furthermore, the transition from the winter to the summer circulation is decisively controlled by the presence of the planetary wave activities (Matthias et al., 2021). Previous studies even concluded that dynamical forced transitions have a persistent impact on the circulation lasting several weeks (Baldwin and Dunkerton, 2001). The stratospheric quasi-biennial oscillation (QBO) modulates the Northern Hemisphere wintertime stratospheric polar vortex, resulting in its weakening and shifting (Garfinkel et al., 2012; Zhang et al., 2019). Wang et al. (2022) use reanalysis data and model simulations to demonstrate that the total column ozone and stratospheric ozone (50–10 hPa) anomalies are seasonally dependent and zonally asymmetric in the polar region, and in that work the QBO affects the polar vortex and stratospheric ozone mainly by modifying the wave number 1 activities. Tao et al. (2019) investigate an intercomparison of simulated stratospheric water vapor variations, focusing on the QBO and long-term variability and trends.

Stratospheric ozone observation results largely reflect a distribution that presents significant asymmetry in both hemispheres, with the differences reaching their maximum values in the winter and spring seasons (Shepherd, 2008). Because most ozone is found in the lower stratosphere, the differences in the column ozone distribution explain the asymmetry because of dynamic transport, as well as the interannual variability of ozone in both hemispheres (McConnell and Jin, 2008; Langematz, 2019). Long-term polar ozone observations offer better recognition and predictability of stratospheric ozone trends and an understanding of the attribution of changes. Water vapor has a chemical lifetime on the order of months in the upper stratosphere and lower mesosphere (Brasseur and Solomon, 2005); therefore, it can be used as a tracer to study a large-scale upwelling and downwelling of the air masses in the polar mesosphere. The mesosphere at the high latitudes is characterized by an annual variation with higher water vapor during local summer and lower water vapor during local winter that is mainly determined by the mean vertical transport (Forkman et al., 2005; Lee et al., 2011). Straub et al. (2010) and Schranz et al. (2019) estimate the vertical gradient of water vapor inside of the polar vortex in autumn based on microwave radiometry measurements at polar latitudes. The distribution and variability of ozone and water vapor exhibit a wealth of information on atmospheric circulation.

There are several techniques to obtain ozone and water vapor measurements in the middle atmosphere. The Aura satellite with the Microwave Limb Sounder (MLS) collects global water vapor and ozone profiles among other chemical species with coverage at a fixed local time due to its sun-synchronous orbit (Livesey et al., 2006). Ground-based observations are often performed using Brewer and Dobson instruments (Zuber et al., 2021), which provide very high-quality and high-precision ozone column densities but lack the vertically resolved information. Lidars are providing good vertical resolution to measure ozone (Brinksma et al., 1997; Bernet et al., 2021). The instruments carried with aircraft and balloon-borne instruments including ozonesondes and frost-point hygrometers perform highly vertically resolved measurements of ozone and water vapor in the upper troposphere and lower stratosphere (Zahn et al., 2014; Eckstein et al., 2017). How-
ever, there are only a few systems available and the observation time depends on tropospheric weather conditions. At tropospheric altitudes, water vapor can also be retrieved leveraging Raman lidars (Sica and Haefele, 2015, 2016). Precise water vapor measurements above the troposphere can also be collected by in situ balloon-borne sensors such as laser absorption spectrometers (Graf et al., 2021). Ground-based microwave radiometers (MWRs) allow continuous observations under all weather conditions with a time resolution of the order of hours except during rain. MWRs measuring ozone and water vapor are valuable as they complement satellite measurements, are relatively easy to maintain, have long lifetimes (which ensures long and continuous time series covering several decades), and can be operated from different locations with measurements performed autonomously on a campaign basis (Scheiben et al., 2013, 2014). Ground-based microwave radiometry is a reliable technique that performs continuous measurements to monitor the vertical profiles of ozone and water vapor VMR changes to investigate Arctic and Antarctic dynamics from diurnal to interannual timescales.

Here, we present a detailed comparison of ozone and water vapor observed by Aura-MLS at the conjugate latitude station leveraging multiyear ground-based observations from GROMOS-C and MIAWARA-C performed at Ny-Ålesund and Aura-MLS and reanalysis data. We produce and compare the polar regions’ multiyear-mean ozone and water vapor climatologies at conjugate latitude stations (Fig. 1). On the one hand, it is intended to provide a well-characterized representation of ozone and water vapor measured by the two instruments and the chemistry differences between both hemispheres concerning climatological behaviors. On the other hand, it provides a source of data for future work, including intercomparison studies and evaluation. Furthermore, we use the water vapor mixing ratio measurements from MIAWARA-C and Aura-MLS observation data to derive the ascent and descent rates. We estimate the strength of upwelling and downwelling in both hemispheres over the polar stations and discuss their interannual variability and the hemispheric differences.

We provide an overview of the datasets in Sect. 2. The time series of ozone and water vapor at conjugate latitude stations in the Northern Hemisphere (NH) and Southern Hemisphere (SH) are presented in Sect. 3. The climatologies of ozone and water vapor are discussed in Sect. 4. The transport of water vapor is discussed in Sect. 5. Sections 6 and 7 present the discussion and conclusions of this study.

2 Instruments and models

In this study, we use ozone and water vapor measurements from our two ground-based MWRs GROMOS-C and MIAWARA-C, which are only available to measure at single locations and are thus representative of a specific geographic location. Both instruments are located at Ny-Ålesund, Svalbard (79° N, 12° E), and have collected continuous data since September 2015. We extract the interannual ozone and water vapor variability between Jan 2015 and July 2022 from MERRA-2 and Aura-MLS over northern and southern polar stations. The corresponding virtual conjugate latitude station (79° S, 12° E) is shown in Fig. 1. In addition, we use temperature observations from Aura-MLS.

2.1 GROMOS-C

GROMOS-C (GRound-based Ozone MOntoring System for Campaigns) is an ozone MWR measuring the ozone emission line at 110.836 GHz at Ny-Ålesund, Svalbard (79° N, 12° E), which is described in detail in Fernández et al. (2015). It was built by the Institute of Applied Physics (IAP) at the University of Bern. GROMOS-C is very compact, meaning it can be transported and operated at remote field sites under extreme climate conditions. It further can switch the frequency of the local oscillator and measure the 115 GHz carbon monoxide (CO) emission line. The system noise temperature of the instrument is about 1080 K. The optics setup of GROMOS-C has two rotating mirrors such that observations in all four cardinal directions are possible. Therefore, GROMOS-C observes in the four cardinal directions (north, east, south, and west) under an elevation angle of 22° with a sampling time of 4 s. Ozone VMR profiles are retrieved from the ozone spectra with a temporal averaging of 2 h leveraging Atmospheric Radiative Transfer Simulator version-2 (ARTS2; Eriksson et al., 2011) and Qpack2 software (Eriksson et al., 2005) according to the optimal estimation algorithm (Rodgers, 2000). An a priori ozone profile is required for optimal estimation and is taken from an MLS climatology of the years 2004–2013. The sensitive altitude range for this instrument extends from
23 to 70 km. The vertical resolution of ozone profiles is 10–
12 km in the stratosphere and increases up to 20 km in the
mesosphere as estimated from the width of the averaging
kernels. The averaging kernels (AVKs) of GROMOS-C to-
gether with its measurement response, errors, and ozone pro-
files are shown in Appendix A (Fig. A1). In the lower strato-
sphere, the errors are below 0.3 ppmv and reach above the
stratopause values up to 0.4 ppmv. More details about the
uncertainty and the AVKs can be found in Fernández et al.

2.2 MIAWARA-C

MIAWARA-C (MIddle Atmospheric WAter vapor RA-
diometer for Campaigns) is a ground-based MWR measur-
ing the pressure-broadened rotational emission line of wa-
ter vapor at the frequency of 22 GHz. It was also built by
the University of Bern and is located at Ny-Alesund, Sval-
brand (79° N, 12° E). The MIAWARA-C front end is an un-
cooled heterodyne receiver with a system noise tempera-
ture of 150 K. The antenna is followed by a dual-polarization re-
ceiver. The incident radiation is split into vertical and hori-
zontal polarization by an orthomode transducer (OMT) lo-
cated immediately after the feedhorn. The two polarized
signals are processed in the two identical receiver chains
and separately analyzed in a fast Fourier transform (FFT)
spectrometer model Acqiris AC240. The spectrometer has a
400 MHz bandwidth and a spectral resolution of 30.5 kHz.
The standard measurement cycle of MIAWARA-C is to mea-
sure sky east, reference east, sky west, and reference west
for about 15 s each. Every 15 min the ambient load is mea-
sured for about 2 s, and the sky at 60° elevation is measured
for about 15 s. A tipping curve is performed to determine
the sky temperature at 60° elevation. The difference spec-
tra in the east and west directions and the two polarizations
are then calibrated separately with the hot and cold measure-
ments close in time. Similar to GROMOS-C, MIAWARA-
C retrieval is also performed with ARTS2 (Eriksson et al.,
2011) and QPACK software (Eriksson et al., 2005) accord-
ing to the optimal estimation algorithm (Rodgers, 2000). An
a priori measurement is taken from an MLS climatology of
the years 2004–2008. From the measured spectra we retrieve
water vapor profiles that cover an altitude range extending
from 37 to 75 km with a vertical resolution of 12–19 km and
with a time resolution of 2–4 h depending on the opacity of
the troposphere. For MIAWARA-C, retrievals are performed
with a constant time resolution and with a constant noise
of 0.014 K. The AVKs of MIAWARA-C, together with its
measurement response, errors, and water vapor profiles, are
shown in Appendix A (Fig. A2). In the upper stratosphere,
the errors are 0.5 ppmv and increase from 0.5 to 1.5 ppmv
in the mesosphere. A detailed design of the instrument and
description of the retrieval algorithm can be found in Straub
et al. (2010) and Tschanz et al. (2013).

2.3 Aura-MLS

The Microwave Limb Sounder (MLS) is one of the payloads
onboard NASA’s Earth Observing System (EOS)-Aura satel-
lite, which was launched in 2004 (Waters et al., 2006). The
satellite is in a sun-synchronous orbital altitude of 705 km
with a period of 1.7 h and 98° inclination. MLS scans the
atmospheric limb in the direction of orbital motion, which
gives almost pole-to-pole coverage (82° S to 82° N), lead-
ing to retrieved profiles at the same latitude every orbit, with
a spacing of a 1.5° great circle angle along the suborbital
track. Ozone is retrieved from the band of 240 GHz, and
water vapor is retrieved from the 183 GHz line. Tempera-
ture is derived from radiances measured from the 118 and
240 GHz channels with a vertical resolution between 3 and
6 km (Schwartz et al., 2015b). The estimated temperature
single-profile precision is 0.5–1.2 K from 100 to 0.001 hPa.
MLS provides ozone profiles (version 5) from 12 to 80 km al-
titude with a vertical resolution of 2.5–6 km (Schwartz et al.,
2015a) and water vapor profiles (version 5) from 10 to 90 km
with a vertical resolution of 3.5 km from 316 to 4.64 hPa and
15 km above 0.1 hPa (Lambert et al., 2015). The estimated
ozone single-profile precision varies from 0.2 to 0.4 ppmv
from the middle stratosphere to the lower mesosphere. For
water vapor, the estimated precision is 0.2–0.3 ppmv in most
of the stratosphere and increases to 0.7–0.8 ppmv in the mid-
dle mesosphere. It passes at Ny-Alesund twice a day at
around 04:00 and 10:00 UTC. Profiles for comparison are ex-
tracted if the location is within ±1.2° latitude and ±6° lon-
gitude of either Ny-Ålesund or the defined virtual conjugate
latitude station.

2.4 MERRA-2

The Modern-Era Retrospective Analysis for Research and
Applications, version 2 (Waters et al., 2006; Gelaro et al.,
2017, MERRA-2) is the latest global atmospheric reanaly-
ysis produced by the NASA Global Modeling and Assimila-
tion Office (GMAO) from 1980 to the present. MERRA-2
assimilates observation types not available to its predeces-
sor, MERRA, and includes updates to the Goddard Earth Ob-
serving System (GEOS) model and analysis scheme so as to
provide a viable ongoing climate analysis beyond MERRA’s
terminus. MERRA-2 provides a regularly gridded, homoge-
neous record of the global atmosphere and incorporates ad-
ditional aspects of the climate system including several im-
provements to the trace gas constituents, land surface repre-
sentation, and cryospheric processes.

In MERRA-2, methods of analysis, model uncertainties,
and observations cause uncertainties (Rienecker et al., 2011).
Davis et al. (2017) provides a comprehensive assessment of
the MERRA-2 ozone product that relatively clearly shows
the vertical distribution of ozone and water vapor in the
stratosphere and has the best agreement with stratospheric
ozone observations compared to other reanalysis products.
Wargan et al. (2017) identified that ozone in MERRA-2 data were expected to have higher uncertainties in regions of high variability, such as winter high latitudes. The uncertainties are related to the chemistry model ozone bias (Gelaro et al., 2017), which is altitude dependent in MERRA-2. Similar behavior was also found in other GEOS chemistry models (Knowland et al., 2022; Wargan et al., 2023). MERRA-2 stratospheric water vapor is also biased compared to independent observations from Aura-MLS and the Atmospheric Chemistry Experiment-Fourier Transform Spectrometer (ACE-FTS). In this study, we use the ozone and water vapor with 72 model levels from the surface up to 0.01 hPa and a horizontal resolution of 0.5° × 0.625°. The time resolution is 6 h. MERRA-2 products are accessible online through the NASA Goddard Earth Sciences Data Information Services Center (GES DISC).

3 Climatologies of ozone and water vapor

This section examines the ozone and water vapor climatologies over Ny-Ålesund, Svalbard (79° N, 12° E), and the conjugate latitude station (79° S, 12° E) generated from the GROMOS-C, MIAWARA-C, MERRA-2, and Aura-MLS datasets. It is important to evaluate how well GROMOS-C and MIAWARA-C can monitor the ozone and water vapor variability and enhance our understanding of their distribution in the Arctic middle atmosphere. The inherent variability of ozone and water vapor in the middle atmosphere can be displayed better in the resulting climatologies, as measured by the two instruments involved in this study, and provide a crucial source of data for future work including intercomparison studies and model evaluation, assessing ozone depletion, validating satellite observations, and studying climate change.

3.1 Ozone

Figure 2 shows the climatological ozone distribution as a function of pressure and time deduced from GROMOS-C, MERRA-2, and Aura-MLS in both hemispheres. Many features of ozone MWR measurement climatologies are broadly consistent with model and satellite data. Some exceptional maximum values larger than 8 ppmv in MERRA-2 data above 0.1 hPa are described in Sect. 3.

Overall, the ozone profile reveals a characteristic seasonal dependence at polar latitudes. In particular, the altitude of the maximum ozone VMR, as well as its temporal variability, exhibits a seasonality. The peak ozone VMR (approximately 6.5 ppmv) appears in the NH in the late spring, whereas autumn shows the lowest values throughout the course of the year. The maximum observed and reanalysis ozone VMR (approximately 5.5 ppmv) in the SH is 1.0 ppmv smaller than the NH maximum and occurs later in the hemispheric spring season. The primary driver of the hemispheric maximum in ozone VMR is related to circulation processes throughout the stratosphere, including those associated with the BDC, transporting ozone-rich air toward the poles in the winter–spring hemisphere. To be precise, this circulation moves the ozone-rich air from the tropical photochemical source region to high latitudes after the polar vortex broke down and essentially enables the intrusion of ozone-rich air from the mid-latitudes into the polar region and the replacement of the ozone-depleted air masses. Another essential feature is that GROMOS-C and Aura-MLS capture the tertiary ozone VMR maximum at the northern polar latitude in the early winter and in late spring at the southern polar latitude. However, the tertiary ozone maximum in GROMOS-C occurs at altitudes close to or even above the limit of 0.8 for the measurement response. MERRA-2 performs poorly when trying to capture the tertiary ozone VMR maximum in both polar latitudes. Due to the complexity of altered dynamics in the polar regions (Wargan et al., 2017) introducing extra uncertainties into numerical models and data assimilation systems, ozone VMRs exhibit dramatic variability (red shading in Fig. 2b, e) in the mesosphere.

From September to November in the Southern Hemisphere, both MERRA-2 and MLS effectively capture the presence of the ozone hole in the lower stratosphere. However, the climatology of ozone in the NH does not reflect a corresponding signature. The annual cycle of ozone at 46 hPa in both hemispheres further reveals this significant feature, as shown in Fig. 3a. MLS exhibits a greater magnitude of ozone depletion compared to MERRA-2. In Fig. 8b, the ozone levels at 3 hPa in the NH follow an annual cycle, with a peak occurring in March and a minimum in October. The SH ozone VMR maintains a constant level of 5 ppmv throughout the summer while experiencing two minimum values in February and October. Furthermore, the ozone VMR in MERRA-2 consistently remains lower than that in MLS except during autumn in the SH. In the mesosphere, there is relatively good agreement between GROMOS-C and MLS, while MERRA-2 exhibits higher variability (Fig. 3c). Both hemispheres have dramatically different seasonal variations and distribution in ozone due to differences in the stratospheric dynamics of the two hemispheres.

3.2 Water vapor

Figure 4 shows the climatology of water vapor vertical and temporal distribution from MIAWARA-C, MERRA-2, and Aura-MLS over both stations (79° S and 79° N) compiled from measurements collected between 2015 and 2021. The annual variation of water vapor with a maximum during hemispheric summer and a minimum during hemispheric winter is clearly visible. During the hemispheric winter, the middle-atmospheric water vapor maximum is shifted down to about 10 hPa, whereas it rises up to the lower mesosphere during the hemispheric summer. The characteristics of water vapor in both hemispheres depend strongly on the mesospheric pole-to-pole circulation, which is an upwelling with
moist air transporting upwards for the hemispheric summer and a corresponding downward motion with dry air into the stratosphere during the winter (Orsolini et al., 2010). In addition, due to the relatively long photochemical lifetime of water vapor, more water vapor produced by methane oxidation accumulates in summer.

The annual cycle of monthly mean water vapor VMR at three separate pressure levels (3, 0.3, and 0.02 hPa) in both hemispheres can be seen in Fig. 5. Water vapor VMR appears at a maximum in October in both hemispheres in the stratosphere (3 hPa) due to the oxidation of methane. Due to photodissociation caused by solar Lyman-alpha radiation acting as a sink and the amount of water vapor being in equilibrium between different photochemical processes and vertical transport, water vapor is more smooth, with nearly constant mixing ratios from winter to spring in the NH. In the mesosphere (0.3 and 0.02 hPa), water vapor gradient variations can be found during hemispheric spring and summer. The gradient at 0.02 hPa is steeper than at 0.3 hPa from April to July in the NH (from November to January in the SH). Furthermore, the positive gradient is weaker, but the time of increase lasts longer and shows an extreme negative gradient in the hemispheric autumn. In both hemispheres, the seasonal behavior of water vapor is almost symmetric at 0.02 hPa and has a slight asymmetry at 0.3 hPa. At 0.02 hPa the maximum value of the water vapor mixing ratio persists for 1 month, and at 0.3 hPa the decrease in water vapor is not visible until September in the NH (March in the SH), but it then appears with a steep gradient.

3.3 Relative differences

In the two ground-based radiometer measurements from GROMOS-C and MIAWARA-C, the aforementioned seasonal behavior in their ozone and water vapor distributions display very similar patterns to Aura-MLS and MERRA-2. However, there are distinct differences between the datasets. To quantitatively assess the consistency of the ozone and water vapor climatologies from GROMOS-C and MIAWARA-C, the relative differences (RDs) of ozone and water vapor VMR between the ground-based MWRs and the two other data sets are calculated using the following expression:

$$\text{RD} = \left( \frac{\langle \varphi \rangle_{\text{radiometer}} - \langle \varphi \rangle_{\text{dataset}}}{\langle \varphi \rangle_{\text{dataset}}} \right) \cdot 100\%,$$

where $\langle \varphi \rangle_{\text{radiometer}}$ is the monthly mean ozone or water vapor VMR from the ground-based radiometer measurements, and $\langle \varphi \rangle_{\text{dataset}}$ is the monthly mean ozone or water vapor VMR from the other data set.
where $\varphi$ represents either ozone or water vapor VMR. Before evaluating the differences in climatology between MWRs and MERRA-2 and Aura-MLS, each MERRA-2 and Aura-MLS profile is convolved with the averaging kernels of MWRs. The convolution is performed according to the following expression:

$$x_{\text{conv}} = x_a - A(x_a - x),$$

where $x_{\text{conv}}$ is the convolved profile, $x_a$ is the a priori profile, $A$ is the averaging kernel matrix, and $x$ is the high-resolution profile of MERRA-2 and Aura-MLS.

Figure 6 shows the relative differences of ozone and water vapor climatologies from GROMOS-C and MIAWARA-C with respect to average convolved MERRA-2 and Aura-MLS. In Fig. 6a, the largest negative RD is larger than 50% above 0.2 hPa in winter and spring because of the high bias of model ozone chemistry in the mesosphere in MERRA-2 (Wargan et al., 2023). GROMOS-C shows relatively good agreement with MERRA-2 in the lower and middle stratosphere (50–5 hPa), with RDs smaller than $\pm$5%, but includes a low RD with magnitudes greater than 10% in the upper stratosphere in autumn. GROMOS-C and Aura-MLS agree well, with RDs mainly within $\pm$7% throughout the middle and upper stratosphere, and exceed positive RD 10% in the lower mesosphere (1–0.1 hPa) in March and October (Fig. 6b).

The RD between MIAWARA-C and MERRA-2 is larger than 50% throughout the late autumn to spring months above 0.2 hPa in Fig. 6c. Simultaneously, MERRA-2 underestimates water vapor VMR at all altitudes due to the lack of assimilated observation to constrain the water vapor reanalyses in the polar mesosphere and also in part due to the methane oxidation parameterization being disabled in the GEOS Composition Forecast model (Davis et al., 2017; Knowland et al., 2022). The MIAWARA-C agrees with MERRA-2 within 7% in the stratosphere and lower mesosphere (about 100–0.3 hPa). MIAWARA-C and Aura-MLS show relatively good agreement, with RDs within 5% throughout the stratosphere and lower mesosphere (1–0.1 hPa) in March and October (Fig. 6b).

The RD between ground-based MWRs and Aura-MLS is in part the Aura-MLS ozone and water vapor profile sampling (as mentioned in Sect. 2.3) and the measurement geometry, leading to seasonal variations in the polar ozone and water vapor distribution. Furthermore, the diurnal cycle in the ozone and water vapor has not been explicitly accounted for in the GROMOS-C and MIAWARA-C measurements. Neglecting the diurnal cycle potentially contributes to positive RDs between MWR measurement and other data sets in the upper stratosphere and lower mesosphere. Overall, GROMOS-C and MIAWARA-C are valuable to monitor the distribution of stratospheric ozone and mesospheric water vapor at the polar latitudes, respectively, which gives us more details to investigate their long-term variability, sources, and trend.

4 Time series of ozone and water vapor

4.1 Ny-Ålesund, Svalbard (79$^\circ$ N, 12$^\circ$ E) in the NH

The time series of daily ozone for GROMOS-C, MERRA-2, and Aura-MLS at Ny-Ålesund, Svalbard (79$^\circ$ N, 12$^\circ$ E), extending from 2015 to 2021 are shown in Fig. 7. The ozone daily profiles measured with GROMOS-C cover a pressure range of 100–0.03 hPa, which corresponds to about 16–70 km. The horizontal upper and lower white lines indicate the bounds of the trustworthy pressure range where the mea-
surement response is larger than 0.8, meaning that the measured spectrum contributes more than 80% to the retrieved profile. The measurement data gaps are that GROMOS-C measured CO for about 2 months during winter 2017/2018 and the spectrometer had a hardware problem during winter 2016/2017 and summer 2019.

Figure 7 reveals the annual ozone cycle with higher ozone VMR in summer (about 6 ppmv) than in winter (about 4.5 ppmv) at about 5 hPa (≈35 km). The GROMOS-C ozone VMR time series together with MERRA-2 and Aura-MLS at about 5 hPa smoothed by a 30 d running mean is shown in Appendix B (Fig. B1a). We can see that all three datasets are able to capture the annual ozone variations well in the stratosphere. Stratospheric ozone largely follows the annual cycle of solar irradiation and is produced through the Chapman cycle; more specifically, ozone VMR is mainly dominated by photochemical production in the summer months in the Arctic middle atmosphere.

In late winter and spring, the stratospheric ozone’s higher variability is largely associated with the stratospheric polar vortex. Figure 7 shows the ozone VMR starting to increase up to the maximum value of about 8 ppmv for some of the years in the stratosphere when the polar vortex is disturbed or weakened by the planetary waves leading to the formation of SSWs. It shows that the planetary wave activity results in meridional transport of the ozone-rich air from the sub-tropics towards the pole and significantly perturbed the distribution of ozone. For example, the polar vortex split and shifted away from Ny-Ålesund, and ozone VMR reached about 7 ppmv during the winter 2018/2019 SSW (Schranz et al., 2020). In some years, the polar vortex is stable and strong over Ny-Ålesund, and ozone VMR sustains smaller values. At the end of the winter 2019/2020 season, the stratosphere featured an extremely strong and cold polar vortex, resulting in low stratospheric ozone in the polar regions (Lawrence et al., 2020; Inness et al., 2020). During late winter and early spring, stratospheric ozone decreases rapidly when the vortex passes over Ny-Ålesund.

Marsh et al. (2001) and Smith et al. (2009, 2018) investigate the tertiary ozone maximum in the winter middle mesosphere at high latitudes based on the models and observations. In this study, GROMOS-C and Aura-MLS present the seasonal tertiary ozone layer at 0.03–0.02 hPa (about 70–75 km) in winter months, but the tertiary ozone VMRs have higher values of 15%–20% in MERRA-2 (as shown the red shading in Fig. 7b). The red shading originates from the un-
certainties in the MERRA-2 product, which are expected to be magnified at high latitudes in winter and spring when the variability is increased compared to other seasons (Wargan et al., 2017). The anomalous atmospheric dynamics, displaced or split polar vortex, and hemispherically asymmetric conditions during SSWs may cause complexity and additional uncertainties in the estimation of ozone flux and transport terms. Figure 7 shows the consistencies of GROMOS-C with both MERRA-2 and Aura-MLS datasets in the time series of ozone below 1 hPa. A clear annual cycle in the stratosphere is well captured by all datasets, and the higher variability of ozone in winter and spring seasons is clearly visible. Remarkably, the annual variation in MERRA-2 ozone is significantly different from the observed variation throughout the mesosphere above 1 hPa.

Figure 8 shows the time series of daily water vapor VMR from MIAWARA-C, MERRA-2, and Aura-MLS at Ny-Ålesund, Svalbard (79° N, 12° E), for the period 2015–2021. MIAWARA-C continuously measures water vapor profiles which cover a pressure range from 5–0.02 hPa corresponding to about 37–75 km. The horizontal upper and lower white lines again indicate the bounds of the trustworthy pressure range where the measurement response is larger than 0.8.

The most evident feature of water vapor is its annual cycle with higher mixing ratios during local summertime and lower mixing ratios during local wintertime throughout the middle atmosphere. This seasonal behavior is mainly driven by the upward and downward branches of the mesospheric residual circulation. As shown in Appendix B (Fig. B1a), water vapor VMR has a maximum of about 7.5 ppmv in summer and a minimum of about 3.5 ppmv in winter at 0.1 hPa (approximately 60 km). Due to the air subsidence inside the polar vortex from autumn to winter, water vapor VMR reaches the maximum at 10 hPa.

In some years, such as 2020, water vapor exhibits a larger variability in late winter and spring. The variation of water vapor is mainly affected by the occurrence of a major SSW which interrupts the polar vortex; after that point, the vortex recovers, and the lower mesospheric water vapor content increases and is accompanied by a decrease in the stratosphere, corresponding to the water vapor vertical profile in the regions outside of the polar vortex (Schranz et al., 2019, 2020). In general, in MIAWARA-C the annually varying mesospheric distribution of water vapor agrees well with reanalysis data and satellite observations. However, there are notable differences at the polar latitudes in the water vapor VMR compared to MERRA-2 throughout the stratosphere and the mesosphere. While MERRA-2 shows a tendency to lower water vapor VMR compared to MIAWARA-C and MLS observations, the seasonal variations of similar amplitude do show reasonably good agreement with the observations. This tendency is likely related to the lack of assimilated observations and known deficiencies in the representation of stratospheric transport (Davis et al., 2017) in the reanalysis data.

4.2 The conjugate latitude station (79° S, 12° E) in the SH

Figures 9 and 10 show time series of ozone and water vapor VMR from MERRA-2 and Aura-MLS at the conjugate latitude station (79° S, 12° E) for the 2015–2021 period, respectively. For comparison purposes, the conjugate latitude station results are lagged by 6 months relative to those for Ny-Ålesund. Both stations exhibit annual cycles of ozone, while the conjugate latitude station shows an ozone VMR maximum of about 6 ppmv in summer and a minimum of about 4 ppmv in winter at about 5 hPa, as shown in Appendix B (Fig. B1b).

Compared to the interannual ozone variability at Ny-Ålesund, the results from the conjugate latitude station are less variable throughout the spring in the stratosphere. Compared to the NH, the planetary wave activity is much weaker in the SH, where a minimum in the ozone mixing ratios prevails over the polar latitudes during the late winter and
Figure 6. Monthly distributions of ozone and water vapor relative differences between ground-based MWRs and data sets: (a, c) MERRA-2 and (b, d) Aura-MLS.

Figure 7. Time series of daily ozone VMR as a function of pressure over Ny-Ålesund, Svalbard (79° N, 12° E), for the 2015–2021 period. Panels show ozone VMR from (a) GROMOS-C measurements, (b) MERRA-2 reanalysis data, and (c) MLS satellite observations. The vertical white lines represent the data gaps caused by the hardware and measurement problems. The horizontal upper and lower white lines indicate a measurement response of 0.8 in panel (a).

spring. There is a dominance of an isolated and stable polar vortex which inhibits the meridional ozone transport to the South Pole throughout the annual cycle, and the formation of polar stratospheric clouds promotes the production of chemically active chlorine and bromine, leading to catalytic ozone depletion in the lower stratosphere. This process is reflected well in the MLS observations and MERRA-2 data for September and October at the conjugate latitude station. Figure 10 presents the water vapor annual cycle with higher mixing ratios during local wintertime and lower mix-
Figure 8. Time series of water vapor VMR as a function of pressure over Ny-Ålesund, Svalbard (79° N, 12° E), for the 2015–2021 period. Panels show ozone VMR from (a) MIAWARA-C measurements, (b) MERRA-2 reanalysis, and (c) MLS satellite observations. The vertical white lines represent the data gaps caused by the hardware and measurement problems. The horizontal upper and lower white lines indicate a measurement response of 0.8 in panel (a).

Figure 9. The same as Fig. 7 (without GROMOS-C measurements) but for the SH at the conjugate latitude station (79° S, 12° E).

ing ratios during local summertime over the conjugate latitude station. Furthermore, polar stratospheric cloud particles can sediment with considerable velocities and irreversibly remove water and nitric acid, resulting in a substantial reduction in water vapor VMR at the lower stratosphere during the southern polar winter and early spring (Waibel et al., 1999; Tritscher et al., 2021). After September, the water vapor VMR increases again as the polar stratospheric cloud influence reduces.

While MERRA-2 generally does an excellent job at reproducing the variability of stratospheric ozone and water vapor, it fails in the mesosphere as discussed in previous sections and also underestimates the vertical extent of the ozone hole, which appears to end at lower altitudes (larger pressures) in MERRA-2 than in MLS. This is also reflected in water vapor, where MERRA-2 fails to reproduce the vertical layering (e.g., July 2015 or July 2017) and underestimates the area of dehydration. Since the reanalysis is less constrained when it
is assimilated (Davis et al., 2017; Shangguan et al., 2019), this behavior can be seen in MERRA-2 when it is compared to observations. It further demonstrates that ozone and water vapor in the NH will continue to be available from ground-based MWR observations; however, the detailed information about the SH winter as shown here for the “conjugate latitude” will be lost after the end of the last three limb sounders, such as Aura-MLS, which is still observing.

5 Dynamics and transport of water vapor

Water vapor is widely used as a tracer to investigate the dynamics of transport processes in the Arctic middle atmosphere (Lossow et al., 2009; Straub et al., 2012; Tschanz et al., 2013; Schranz et al., 2019, 2020). The chemical lifetime of water vapor is on the order of years in the lower stratosphere, months in the lower mesosphere, and weeks in the upper mesosphere (Brasseur and Solomon, 2005). Water vapor mixing ratios display different dynamical features depending on the altitude ranges because of different vertical gradients of water vapor VMR above and below its peak in the upper stratosphere. The water vapor mixing ratio is assumed to be constant for a month and a half, and the vertical velocity of air can be estimated.

The time periods of the year when water vapor VMR increases and decreases with altitude (in the upper stratosphere and lower mesosphere) measured from MIAWARA-C and Aura-MLS over both Ny-Ålesund and the conjugate latitude station are well covered in this study. We will denote the period with a stable high water vapor mixing ratio as the hemispheric summer and the positive and negative transition periods as the upwelling and downwelling branches, respectively. With the northern and southern hemispheric water vapor measurements, we calculate the effective ascent and descent rates as derived from a linear regression fit to different water vapor mixing ratio isopleths (5.5, 6.0, and 6.5 ppmv).

For instance, the ascent and descent rates from 6.5 ppmv water vapor isopleth are shown in Figs. 11 and 12).

The time period of many years of descent rate from 15 September to 31 October in the altitude range of about 50–70 km is well presented in Fig. 11 (the first and third rows). This is an estimated result and not a quantitative calculation of the water vapor descent since the water vapor dynamic and chemical reactions not directly related to descent may also affect the polar water vapor changes. However, the effect of dynamic processes such as planetary wave disturbance is relatively obvious to estimate the ascent rate. As shown in Fig. 11 (second and fourth rows), the time period of 5 years (2015, 2016, 2017, 2018, and 2021) of ascent rate starts from 5 May to 20 June, and in another 2 years (2019 and 2020) the starting time for the increase of water vapor happens earlier by approximately 20 d (from 15 April). In 2019 and 2020, MIAWARA-C and Aura-MLS observe the return of the water vapor mixing ratio to pre-winter values in mid-April, and the moist air is lifted to greater altitudes in early May. The starting time of the ascent rate in 2019 and 2020 is about 3 weeks earlier than that in other years, which is caused by several processes. The planetary waves displace the polar vortex above Ny-Ålesund, and water-vapor-rich air is transported into the upper stratosphere; furthermore, the maximum water vapor mixing ratio is re-established at an altitude of about 55 km. Wave breaking and mixing above the strongest vortex level tends to reduce the tracer gradient (Lee et al., 2011), leading to an increase in the water vapor mixing ratios. In addition, photochemical processes from solar radiation can also contribute to the accumulation of water vapor in the springtime at the stratopause altitude (Brasseur and Solomon, 2005).

In the SH, the time period of each year of the ascent rate is relatively consistent from 15 October to 30 November in the upper stratosphere and mesosphere (Fig. 12). The descent rate of water vapor from 15 March to 30 April in the SH appears to be similar for each of the 7 years,
likely due to the higher stability and strength of the southern polar vortex. Figure 13 shows the annual variability of effective ascent and descent rates in the SH and NH during a given time period. In contrast to the NH, the southern hemispheric rate exhibits less interannual variability and an approximately consistent variation with the rates corresponding to the different isopleths, illustrating the qualitative agreement in different years in the effective vertical rates for the transition periods in the SH. The uncertainties for the ascent and descent rates of airflow over northern and southern polar latitude stations are depicted by error bars in Fig. 13. Effective ascent and descent rates estimated from water vapor measurements are very significant. Table 1 gives a more quantitative perspective to compare the vertical movement of air in both hemispheres. In the NH, the average vertical velocities are $3.4 \pm 1.9 \text{ mm s}^{-1}$ from MIAWARA-C and $4.6 \pm 1.8 \text{ mm s}^{-1}$ from Aura-MLS for upwelling from spring to summer and $5.0 \pm 1.1 \text{ mm s}^{-1}$ from MIAWARA-C and $5.4 \pm 1.5 \text{ mm s}^{-1}$ from Aura-MLS for downwelling from summer to autumn. During the transition from winter to spring, the vertical velocity is $5.2 \pm 0.8 \text{ mm s}^{-1}$ for downwelling and $2.6 \pm 1.4 \text{ mm s}^{-1}$ for upwelling from autumn to winter calculated by Aura-MLS in the SH. Table 1 shows a stronger upwelling branch in the NH polar summer mesosphere as compared to the SH, and this is accompanied by a stronger downwelling branch towards the SH winter in the polar region. In general, these results assess the ability to derive middle atmospheric ascent and descent rates from water vapor measurements at polar latitudes and further provide evidence for the higher variability in the NH than in the SH.
Figure 13. Interannual variability of the effective ascent and descent rates over both Ny-Ålesund, Svalbard (79° N, 12° E), and the conjugate latitude station (79° S, 12° E) estimated from the linear regression fit to the 5.5, 6.0, and 6.5 ppmv isopleths of water vapor VMR from MIAW ARA-C and Aura-MLS. Bars indicate the uncertainties in the ascent and descent rates based on the MIAW ARA-C and Aura-MLS observations.

Table 1. Comparison of the mean values and standard deviations (SDs) of ascent and descent rates from MIAW ARA-C and Aura-MLS in both hemispheres. The mean is computed for the rates from 5.5, 6.0, and 6.5 ppmv isopleths (as shown in Fig. 13).

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<tr>
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<th>Ascent rate (mm s⁻¹)</th>
<th>Descent rate (mm s⁻¹)</th>
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<td>NH (mean ± SD)</td>
<td>SH (mean ± SD)</td>
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<tr>
<td>MIAW ARA-C</td>
<td>3.4 ± 1.9</td>
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<td>Aura-MLS</td>
<td>4.6 ± 1.8</td>
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6 Discussion

Continuous observations of essential climate variables such as ozone and water vapor are important for investigating the radiative balance of the atmosphere. Interhemispheric and interannual differences shed light on ozone and water vapor natural variability due to transport and photochemistry.

Ground-based measurements such as those performed by GROMOS-C and MIAW ARA-C provide a high-resolution and continuous data set of these trace gases at remote locations such as Ny-Ålesund. A comparison to the Aura-MLS satellite data exhibits excellent agreement throughout all altitudes in the stratosphere with GROMOS-C and the upper stratosphere and lower mesosphere with MIAW ARA-C. The climatologies of Aura-MLS and the MWRs agree within ±10% during the year. However, a climatological comparison to the reanalysis data MERRA-2 and our ground-based radiometers indicates larger discrepancies above 0.2 hPa. These increased deviations are partially due to the implemented radiative transfer schemes and other model physics used, such as interactive chemistry, which is computationally much more expensive (Gelaro et al., 2017). Furthermore, MERRA-2 includes the MLS observations of temperature and ozone in the 3DVAR data assimilation (Wargan et al., 2017). MLS observations are most important for the mesosphere and are weighted by their precision and accuracy.

The interhemispheric comparison is performed by defining a virtual conjugate latitude station (79° S, 12° E) in the Southern Hemisphere at conjugate geographic coordinates. Although the general seasonal morphology is very similar,
the Northern Hemisphere shows much more variability in the time series of ozone and water vapor. One of the most significant differences is the occurrence of the ozone hole in the Southern Hemisphere towards the end of the winter season below 10 hPa (Solomon et al., 2014). During this time the water vapor VMR measurements also exhibit a minimum due to the formation of polar stratospheric clouds (Flury et al., 2009; Bazhenov, 2019) providing even more favorable conditions for catalytic ozone destruction reactions. In Fig. 14, we present zonally averaged temperature at two conjugate latitudes and compare results between the polar and mid-latitudes from Aura-MLS observations in both hemispheres. The southern polar latitudes appear much colder, by about 20 K, than their northern counterparts, as seen in Appendix C (Fig. C1). Furthermore, the temperature gradient between the mid-latitudes and high latitudes is stronger in the Southern Hemisphere at the stratosphere and thus drives a more stable polar vortex due to the thermal wind balance, which prevents the mixing of ozone-rich air from the low latitudes and mid-latitudes into the polar cap by planetary waves as can often be observed in the Northern Hemisphere (Schranz et al., 2019). There are only a few occasions from 2015 onward where such a stable and cold polar vortex was observed in the Northern Hemisphere, i.e., in 2015/2016 and 2019/2020 in the Arctic winter (Matthias et al., 2016; Lawrence et al., 2020), which can also lead to anomalies in the middle atmospheric dynamics (Stober et al., 2017).

In addition to the strength of the polar vortex, we investigated the strength of the upwelling and downwelling in both hemispheres, which we consider a proxy of the strength of the residual circulation. Water vapor has a longer lifetime than CO at 50–70 km (Brasseur and Solomon, 2005), which makes it a more robust tracer to manifest the vertical motion of air. The effective rates of vertical transport are estimated by using the water vapor measurements following the approach presented by Straub et al. (2010). The vertical velocities are derived under the assumption that other processes, such as chemical reactions, photodissociation, or horizontal advection, are less important. Here we analyze periods around the equinoxes. During this time of the year, the horizontal gradients of temperature between the polar region and the mid-latitudes are minimal or negligible, and thus there is no strong forcing due to horizontal temperature gradients or planetary waves driving the zonal or meridional transport. Furthermore, we only investigated altitudes below typical hydroxyl (OH) layer heights, which also reduces the impact of photodissociation in our estimates (Ryan et al., 2018). We obtained vertical velocities of 3.4 ± 1.9 and 4.6 ± 1.8 mm s⁻¹ for the upwelling towards the northern hemispheric summer and vertical motions of 5.0 ± 1.1 and 5.4 ± 1.5 mm s⁻¹ downwelling during the fall transition. Furthermore, there is rather significant interannual variability in the Northern Hemisphere that is not found above the Antarctic continent. Due to the frequent occurrence of SSWs, the spring transition is more variable in the NH (Matthias et al., 2021). In the Southern Hemisphere, the spring transition towards summer is characterized by 2.6 ± 1.4 mm s⁻¹ vertical velocity, and during the transition from summer to winter a downwelling with 5.2 ± 0.8 mm s⁻¹ vertical velocity is observed. A high variability in NH winter downwelling would imply a high variability of SH summer upwelling, which is indeed observed in Fig. 13. This suggests that the interhemispheric coupling (Körnich and Becker, 2010; Orsolini et al., 2010; Smith et al., 2020) means that disturbances are transmitted from the winter stratosphere to the summer mesosphere in the other hemisphere. However, it is worth noting
that there is a lower variability in SH winter downwelling, which corresponds to a higher variability in NH summer upwelling. This observation suggests that the coupling between the winter and summer circulations within each hemisphere is influenced by various factors beyond the interhemispheric exchange. As shown in Fig. 14, with a stronger temperature gradient during SH winter, the polar vortex is relatively stable and well defined, leading to reduced variability in the downward motion of air (downwelling). For instance, the SSW events during the NH winter (Limpasuvan et al., 2016; Schranz et al., 2019, 2020) or increased stratospheric planetary wave activity (de Wit et al., 2015) lead to increased variability in the NH winter stratosphere. Simulations with a GW-resolving model response to the enhanced winter hemisphere Rossby wave activity may lead to both interhemispheric couplings through a downward shift of the GW-driven branch of the residual circulation and increased GW activity at high summer latitudes (Becker and Fritts, 2006). Other observational studies using polar mesospheric clouds and stratospheric reanalysis data exhibited an interhemispheric correlation during the summer months (Karlsson et al., 2007; Espy et al., 2011). More recent model results suggested that the strongest interhemispheric coupling signatures are found between the stratosphere and mesosphere in the opposite hemisphere (Smith et al., 2020).

7 Conclusions

Continuous ground-based measurements of ozone and water vapor remain an essential tool to understand the short and long-term evolution of the middle atmosphere, as well as for the validation and parameterization of atmospheric models. In this study, we present ozone and water vapor measurements from the two ground-based radiometers GROMOS-C and MIAWARA-C located at Ny-Ålesund, Svalbard, collected between 2015 and 2021. The data were compared to observations from MLS onboard the Aura spacecraft as well as reanalysis data MERRA-2. This comparison showed a good agreement for the climatological behavior between the ground-based radiometers and MLS to within almost ±7% for ozone at about 50–1 hPa and within ±5% for water vapor at about 100–0.02 hPa. However, we identified pronounced differences between the measurements and the reanalysis data above 0.2 hPa where MERRA-2 deviations up to 50% were visible. Ground-based observations are going to become more important within the next few years as the satellite instruments such as MLS are going to reach the end of their life and so far there are no adequate replacements in orbit.

By defining a virtual conjugate latitude station in the Southern Hemisphere, we investigated altitude-dependent interhemispheric differences. Both trace gases showed a much higher variability during the northern hemispheric winter driven by planetary wave activity. The Southern Hemisphere was characterized by a more stable polar vortex and colder temperatures in the polar cap that result in more favorable conditions to form polar stratospheric clouds and thus more efficient ozone destruction by catalytic reactions causing the well-known ozone hole. Furthermore, the polar stratospheric cloud formation was accompanied by a reduction in the water vapor VMR at the same altitudes in the lower stratosphere.

We investigated the strength of the residual circulation by estimating the upwelling and downwelling above Ny-Ålesund and the corresponding conjugate latitude station. Typical ascent rates during the summer transition reach values of 3.4–4.6 mm s\(^{-1}\), and for the downwelling in the fall transition vertical velocities of 5.0–5.4 mm s\(^{-1}\) are inferred. Correspondingly, a vertical velocity of 2.6 mm s\(^{-1}\) for the upwelling and 5.2 mm s\(^{-1}\) for the downwelling is calculated in the SH. The Northern Hemisphere also reflected a much more pronounced interannual variability compared to the southern polar latitudes. However, there is no strong correlation between upwellings and downwellings in the opposite hemispheres; this is most likely due to dynamic processes such as the QBO or weather patterns that play a role and need to be taken into account. Therefore, long-term ozone and water vapor measurements will create a deeper understanding of the mechanisms that control polar ozone and water vapor variability and predict the future evolution of middle atmospheric ozone and water vapor in climate changes.
Appendix A: Retrieved results of instruments

Figure A1. Example of GROMOS-C hourly ozone retrievals on 25 October 2021 around 01:00 UTC. Panel (a) shows the averaging kernels together with measurement response; panel (b) shows the measurement, smoothing, and observation errors; and panel (c) shows the retrieved and a priori ozone profile.

Figure A2. Example of MIAWARA-C hourly water vapor retrievals on 20 August 2021 around 23:00 UTC. Panel (a) shows the averaging kernels together with measurement response; panel (b) shows the measurement, smoothing, and observation errors; and panel (c) shows the retrieved and a priori water vapor profile.
Appendix B: Ozone and water vapor

Figure B1. GROMOS-C, MERRA-2, and Aura-MLS ozone VMR time series at about 5 hPa smoothed by a 30 d running mean over Ny-Ålesund, Svalbard (79° N, 12° E), and the conjugate latitude station (79° S, 12° E).

Figure B2. MIAWARA-C, MERRA-2, and Aura-MLS water vapor VMR time series at 0.1 hPa smoothed by a 30 d running mean over Ny-Ålesund, Svalbard (79° N, 12° E), and the conjugate latitude station (79° S, 12° E).
Appendix C: Time series of temperature at mid-latitudes and polar latitudes

Figure C1. Time series of temperature from Aura-MLS observations as a function of pressure over conjugate polar latitudes (80° N and 80° S) and conjugate mid-latitudes (60° N and 60° S) in both hemispheres.

Data availability. The GROMOS-C and MIAWARA-C level 2 data are provided by the Network for the Detection of Atmospheric Composition Change and are available at https://ndacc.larc.nasa.gov/stations/ny-alesund-norway (Schranz et al., 2020). MLS v5 data are available from the NASA Goddard Space Flight Center Earth Sciences Data and Information Services Center (GES DISC): https://doi.org/10.5067/Aura/MLS/DATA2516 (Schwartz et al., 2020). MERRA-2 data are provided by NASA at the Modeling and Assimilation Data and Information Services Center (MDISC) and are available at the model level at https://doi.org/10.5067/WWQSXQ81VFW8 (GMAO, 2015a) and the pressure level at https://doi.org/10.5067/QBZ6MG944HW0 (GMAO, 2015b).

Competing interests. The contact author has declared that none of the authors has any competing interests.

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Author contributions. GSh was responsible for the ground-based ozone measurements with GROMOS-C, performed the data analysis, and prepared the manuscript. ES provided the Aura-MLS data. WK helped with data analysis. GST designed the concept of the study and contributed to the interpretation of the results. All of the authors discussed the scientific findings and provided valuable feedback for manuscript editing.

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References


Lambert, A., Read, W., and Livesey, N.: MLS/Aura Level 2 Water Vapor (H2O) Mixing Ratio V004, Greenbelt, MD, USA, Goddard Earth Sciences Data and Information Services Center (GES DISC) [data set], https://doi.org/10.5067/Aura/MLS/DATA2009, 2015.


Schanz, F., Tschanz, B., Rüfenacht, R., Hocke, K., Palm, M., and Kämpfer, N.: Investigation of Arctic middle-atmospheric dy-


Schwartz, M., Froidevaux, L., Livesey, N., and Read, W.: MLS/Aura Level 2 Temperature V004, Greenbelt, MD, USA, Goddard Earth Sciences Data and Information Services Center (GES DISC) [data set], https://doi.org/10.5067/Aura/MLS/DATA2015b, 2015b.

Schwartz, M., Froidevaux, L., Livesey, N., and Read, W.: MLS/Aura Level 2 Ozone (O3) Mixing Ratio V005, Greenbelt, MD, USA, Goddard Earth Sciences Data and Information Services Center (GES DISC) [data set], https://doi.org/10.5067/Aura/MLS/DATA2016, 2016a.


