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On the impact of the temporal variability of the collisional quenching process on the mesospheric OH emission layer: a study based on SD-WACCM4 and SABER

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Abstract. The mesospheric OH Meinel emissions are subject of many theoretical and observational studies devoted to this part of the atmosphere. Depending on the initial vibrational level of excitation the altitude of the considered OH Meinel emission is systematically shifted, which has important implications for the intercomparison of different studies considering different transition bands. Previous model studies suggest that these vertical shifts are essentially caused by the process of collisional quenching with atomic oxygen. Following this hypothesis, a recent study found experimental evidence of a coherent seasonality at tropical latitudes between vertical shifts of different OH Meinel bands and changes in atomic oxygen concentrations. Despite the consistent finding of the above mentioned hypothesis, it cannot be excluded that the actual temporal variability of the vertical shifts between different OH Meinel bands may in addition be controlled or even dominated by other processes. It remains an open question whether the observed temporal evolution is indeed mainly controlled by the modulation of the collisional quenching process with atomic oxygen. By means of a sensitivity study which employs a quenching model to simulations made with the SD-WACCM4 chemistry climate model, we aim at assessing this question. From this study we find that the observed seasonality of vertical OH Meinel shifts is only partially controlled by temporal changes in atomic oxygen concentrations, while molecular oxygen has another noticeable impact on the vertical OH Meinel shifts. This in particular becomes evident for the diurnal variability of vertical OH Meinel shifts, which reveal only a poor correlation with the atomic oxygen species. Furthermore, changes in the $H + O₃$

source gases provide another mechanism that can potentially affect the diurnal variability in addition. By comparison with limb radiance observations from the SABER/TIMED satellite this provides an explanation for the less evident diurnal response between changes in O concentrations and vertical OH Meinel shifts. On the other hand, at seasonal timescales the coherency between both quantities is again evident in SABER/TIMED but less pronounced compared to our model simulations.

1 Introduction

The hydroxyl (OH) emission layer is a prominent feature of the mesopause region. Its main production process is commonly referred to as the Bates–Nicolet mechanism [\(McDade,](#page-16-0) [1991\)](#page-16-0). This mechanism suggests the exothermic reaction between O_3 and H, which leads to rotationally, vibrationally excited OH radicals [\(Bates and Nicolet,](#page-16-1) [1950\)](#page-16-1). According to the available exothermic energy of this reaction, these radicals can have excited vibrational states up to the $\nu = 9$ quantum number. Lower vibrational states can be populated via spontaneous emission, but also through collisional quenching with ambient species. Hence, we can distinguish between different $OH(v)$ layers with respect to their vibrational excitation states.

Because different observational studies on the mesospheric OH Meinel emission rely on different transition bands, it is of general interest to understand systematic differences between the vertical profiles of the associated $OH(v)$ layers. As we know from previous rocket campaigns (e.g. see [Baker and Stair Jr.](#page-16-2) [\(1988\)](#page-16-2) for a comprehensive compilation of rocket campaigns), systematic vertical shifts exist between these layers, while further studies have shown that collisional quenching with ambient species is significantly affecting these shifts (e.g. [Dodd et al.,](#page-16-3) [1994;](#page-16-3) [Makhlouf et al.,](#page-16-4) [1995,](#page-16-4) and [Adler-Golden,](#page-16-5) [1997\)](#page-16-5). In particular atomic oxygen is an effective quencher and its impact on the vertical distribution of different $OH(v)$ layers has been recently investigated by [von Savigny et al.](#page-17-0) [\(2012\)](#page-17-0). Based on a sensitivity study, which relies on an updated version of a quenching model by [McDade](#page-16-0) [\(1991\)](#page-16-0), they suggest that quenching with atomic oxygen causes an upward shift of the individual $OH(v)$ layers with increasing vibrational state. In a follow-up study [von](#page-17-1) [Savigny and Lednyts'kyy](#page-17-1) [\(2013\)](#page-17-1) provided experimental evidence that the vertical shifts between different OH bands are indeed correlated with the amount of atomic oxygen in the altitude range of the OH emission layer. Despite the consistent findings between both studies, it should be outlined that the simulated OH profiles by [von Savigny et al.](#page-17-0) [\(2012\)](#page-17-0) were limited to a single month based on the MSIS climatology, while the effect of collisional O quenching has been considered by different scaling factors in the associated rate term. On the other hand, systematic changes in the vertical $O₃$ and H profiles will also affect the temporal variability of the vertical $OH(v)$ shifts and must be taken into account when discussing the impact of collisional quenching on the vertical structure of the $OH(v)$ layers. Thus, it remains an open question whether the temporal changes in the relative vertical $OH(v)$ shifts are mainly driven by the temporal variability of the ambient quenching species, the source gases of OH or by a combination of both.

To investigate the importance of the temporal variability of the collisional quenching on the vertical $OH(v)$ shifts, this study established an updated quenching model that is applied to simulations made with the Whole Atmosphere Community Climate Model driven with Specified Dynamical fields (SD-WACCM4).

The emphasis of this study will be on the equatorial regions, where the large amplitude of the diurnal migrating tide has a strong impact on OH airglow and ambient temperatures [\(Shepherd et al.,](#page-17-2) [2006\)](#page-17-2). Many studies have reported evidence of a semi-annual oscillation in airglow observations that is associated with the large seasonal changes in the tidal amplitude. For instance, [Marsh et al.](#page-16-6) [\(2006\)](#page-16-6) show a pronounced semi-annual oscillation in SABER OH volume-emission-rate (VER) measurements at equatorial latitudes. A similar seasonality was also recently shown for OH VER measurements from SCIAMACHY (SCanning Imaging SpectroMeter for Atmopsheric CHartographY) by [von Savigny and](#page-17-1) [Lednyts'kyy](#page-17-1) [\(2013\)](#page-17-1). In addition, a semi-annual oscillation was also reported from HRDI observations [\(Yee et al.,](#page-17-3) [1997\)](#page-17-3) and ISIS-2 observations [\(Cogger et al.,](#page-16-7) [1981\)](#page-16-7) of the $O(^1S)$ green line. Because the vertically integrated O concentration should be proportional to the integrated OH VER (see Eq. 2 in [Mlynczak et al.,](#page-16-8) [2013\)](#page-16-8), the same observed seasonal variability could also apply for the vertical $OH(v)$ shifts.

Based on the initial hypothesis that the collisional quenching with atomic oxygen is affecting the relative vertical $OH(v)$ shifts, we would therefore expect a coherent response in these shifts with the temporal evolution of the diurnal migrating tide. Accordingly, we focus on the seasonal and diurnal changes in the collisional quenching of OH with atomic oxygen. In addition, we will also consider the impact of collisional quenching with molecular oxygen, the second most efficient OH quencher after atomic oxygen [\(Adler-Golden,](#page-16-5) [1997\)](#page-16-5). The advantage of our model approach is that we can deactivate the individual collisional quenching processes to study the associated impact on the relative vertical $OH(v)$ shifts. We compare these simulations with limb radiance observations from the SABER (Sounding of the Atmosphere by Broadband Emission Radiometry) instrument onboard the TIMED (Thermosphere Ionosphere Mesosphere Energetics Dynamics) satellite and discuss the observed temporal variability of the vertical $OH(v)$ shifts with regard to our model results.

This paper is structured as follows. Section [2](#page-1-0) introduces our OH quenching model and gives a brief summary on the SD-WACCM4 and SABER data. The methodology of our analysis on the relative vertical $OH(v)$ shifts is explained in Sect. [3.](#page-4-0) This is followed by a discussion on potential sources of error in Sect. [4.](#page-4-1) Based on a case example we reexamine important systematic features of the temporal variability of the OH emission layer and O quenching species in Sect. [5](#page-5-0) to establish an expectation on the temporal evolution of vertical $OH(v)$ shifts. In Sect. [6](#page-6-0) we investigate the initial hypothesis on the role of collisional quenching on the vertical $OH(v)$ shifts by simulating the seasonal variability of the OH emission layer from the SD-WACCM4 data for different model assumptions. These simulations are then compared with experimental observations from SABER. Based on the same methods, the diurnal variability of the OH quenching process is investigated in Sect. [7.](#page-9-0) We provide a summary of our results for the seasonal and diurnal variability of the relative vertical $OH(v)$ shifts in Sect. [8](#page-14-0) and discuss their implications on the initial hypothesis.

2 Model and data description

2.1 Hydroxyl quenching model

A detailed description of the quenching model, which we use as a basis for our OH simulations, is given in [McDade and](#page-16-9) [Llewellyn](#page-16-9) [\(1988\)](#page-16-9) and [McDade](#page-16-0) [\(1991\)](#page-16-0). Here, we limit our discussion to its primary key aspects and our adjustments to simulate absolute number densities of $OH(v)$.

As mentioned in the beginning, the Bates–Nicolet mechanism suggests the principal excitation mechanism of vibrationally excited OH according to the following reaction:

$$
H + O_3 \to OH(v' \le 9) + O_2
$$

 k_1 , (R1)

where k_1 denotes the rate constant of this reaction. The released exothermic energy of this reaction leads to a preferred vibrational excitation between $v = 6$ and $v = 9$. In accordance with [von Savigny et al.](#page-17-0) [\(2012\)](#page-17-0) we assume the following processes to populate lower vibrational states:

– radiative cascade from the initially populated higher levels

$$
\text{OH}(v') \to \text{OH}(v'') + hv \qquad \qquad A(v', v'') \text{ (R2)}
$$

- **–** collisional relaxation
- $OH(v') + Q \rightarrow OH(v'') + Q$ k_3^Q $rac{Q}{3}(\nu', \nu'')$ (R3) with $Q = O_2$, N_2 .
- **–** complete OH removal

 $OH(v') + Q \rightarrow$ other products k Q

with $Q = 0, Q_2, N_2$.

Apart from these processes, the recombination of the perhydroxyl radical $(HO₂)$ with atomic oxygen as being proposed by [Krassovsky](#page-16-10) [\(1963\)](#page-16-10) could provide another mechanism to form OH with vibrational excitations below $\nu < 6$ at the mesopause. Different opinions exist on the importance of this mechanism to the general OH formation (e.g. see [Khomich et al.,](#page-16-11) [2008,](#page-16-11) for a summary of different studies), though the recent study by [Xu et al.](#page-17-4) [\(2012\)](#page-17-4) implicates that its contribution is rather negligible for vibrational states above $v = 3$. As we will discuss later, the main emphasis of our study is on vibrational states above $v = 3$, accordingly we neglect this mechanism in our following considerations.

Following [McDade](#page-16-0) [\(1991\)](#page-16-0), Eq. (3) in [von Savigny et al.](#page-17-0) [\(2012\)](#page-17-0) describes the OH concentration for steady state conditions. Here, we adjust this expression as follows:

$$
[OH(v)] = \left(A(v) + \sum_{Q} k_{L}^{Q}(v)[Q]\right)^{-1} \times
$$

$$
\left(P(v)\{k_{1}[H][O_{3}]\} + \sum_{v^{*}=v+1}^{9}[OH(v^{*})]
$$

$$
\left\{A(v^{*}, v) + \sum_{Q} k_{3}^{Q}(v^{*}, v)[Q]\right\},
$$
 (1)

where P is the nascent vibrational level distribution, $A(v)$ corresponds to the inverse radiative lifetime of OH and k_l^Q \sum_L is the total rate constant for removal of OH in vibrational level ν through Reactions [\(R3\)](#page-2-0) and [\(R4\)](#page-2-1). Accordingly, we substitute the nascent production rate p in [von Savigny et al.](#page-17-0) [\(2012\)](#page-17-0) by the $P(v)$ {k₁[H][O₃]} rate term in the nominator of Eq. [\(1\)](#page-2-2). In contrast to the work of [von Savigny et al.](#page-17-0) [\(2012\)](#page-17-0), we do not normalise Eq. [\(1\)](#page-2-2) with respect to the $\nu = 9$ vibrational state, because we aim to calculate absolute number densities of $OH(v)$ to allow for a direct comparison with the observed VER by SABER. Therefore, we have to implement absolute rate constants as well as absolute inverse radiative lifetimes in Eq. [\(1\)](#page-2-2).

For our present model simulations we use the constants listed in Table [1,](#page-3-0) assuming that multi-quantum relaxation only applies for quenching with O_2 , while the less efficient N_2 quenching is limited to single-quantum relaxation only. If we apply these assumptions to Eq. [\(1\)](#page-2-2), we get the following expression for OH as a function of vibrational state:

$$
\begin{aligned}\n\mathcal{Q}_{1}(v',v'') \quad &(\mathbb{R}3) \qquad [\text{OH}(v)] = \left(A(v) + k_{L}^{O_{2}}(v)[O_{2}] + k_{L}^{N_{2}}(v)[N_{2}] + k_{L}^{O}(v)[O]\right)^{-1} \times \\
& \left(\begin{array}{c} P(v)[k_{1}[\text{H}][O_{3}]] + \sum_{v^{*} = v+1}^{9} [OH(v^{*})][A(v^{*},v) + k_{3}^{O_{2}}(v^{*},v)[O_{2}] + k_{3}^{N_{2}}(v^{*},v)[N_{2}]\end{array}\right) \\
& \text{with } k_{3}^{N_{2}}(v^{*},v) = 0 \text{ for all } \{v^{*} > v+1\} \text{ and } k_{3}^{N_{2}}(v^{*},v) = \\
& \mathcal{Q}_{1}(v',v'') \quad &(\mathbb{R}4) \qquad k_{L}^{N_{2}}(v^{*}) \text{ for } \{v^{*} = v+1\}.\n\end{aligned}
$$

2.2 SD-WACCM4

The SD-WACCM4 simulations are based on the Whole Atmosphere Community Model, version 4 (WACCM4), which is a comprehensive free running chemistry–climate model. This model version is based on an earlier version described by [Garcia et al.](#page-16-12) [\(2007\)](#page-16-12) and has been recently extended, such that it is nudged to meteorological fields that are taken from the Global Earth Observing System Model, Version 5 (GEOS-5) of NASA's Global Modeling and Assimilation Office (GMAO).

SD-WACCM4 data were provided to us by courtesy of R. R. Garcia and D. E. Kinnison, NCAR Boulder. The same SD-WACCM4 simulations, which we consider in our study, were already applied to another study by [Hoffmann et al.](#page-16-13) [\(2012\)](#page-16-13) that investigates the dynamics of the model using mesospheric CO volume-mixing-ration (VMR) measurements. We therefore refer to this paper for a more detailed description of the model. Here, we limit our discussion to the most relevant aspects to our study.

The nudging of SD-WACCM4 with GEOS-5 meteorological fields M_{GEOS} is performed up to 50 km altitude for each time step t by replacing the model-predicted fields M_{pred} according to the following equation:

 $M_{pred}(t) = 0.99 \cdot M_{pred}(t) + 0.01 \cdot M_{GEOS}(t).$

Between 50 and 60 km altitude a linearly decreasing relaxation scheme follows until it completely switches to a freerunning mode above 60 km.

Despite the weak constraint of SD-WACCM4 by its relaxation to external meteorological fields from GEOS-5, [Hoff](#page-16-13)[mann et al.](#page-16-13) [\(2012\)](#page-16-13) show that the upper (free-running) part is

Table 1. Employed constants to Eq. [\(2\)](#page-2-3).

still strongly driven by the described nudging and closely reflecting the dynamic response, which they deduce from CO based measurements.

The horizontal resolution of the SD-WACCM4 data is $1.9^\circ \times 2.5^\circ$ in latitude and longitude. Its vertical extent reaches from the ground up to the lower thermosphere at about 137 km geopotential height (GPH) and it is divided into 66 height levels. The provided GPH values are transformed to geometric heights for our analysis. In the region from 80 km up to 95 km, which encloses the hydroxyl emission, the vertical distance between the model grid points varies from about 1.2 km to 3.6 km. The SD-WACCM4 simulations are initially performed at 0.5 h time increments, however, to save computational resources, global model results are stored as daily increments at 00:00 UTC. This limitation of our data set prevents us from studying the diurnal evolution of the OH vertical profiles at a fixed geolocation. To overcome this constraint we make the assumption that the diurnal evolution of the vertical profiles is already contained within the zonal variation of each daily model result, i.e. we convert the longitudinal information to the local solar time (LST). However, as we will discuss in Sect. [7.2,](#page-13-0) other processes exist, which can still complicate a direct comparison of the diurnal variability between SD-WACCM4 and SABER.

To simulate $OH(v)$ profiles by means of Eq. [\(2\)](#page-2-3), we convert the SD-WACCM4 chemical profiles from VMR to absolute number densities based on the provided pressure and temperature fields. In addition, we consider the SD-WACCM4 temperatures for the calculation of the temperature dependent rate constant k_1 of Reaction [\(R1\)](#page-2-4). The SD-WACCM4 data in this study cover the period between April 2010 and June 2011.

2.3 SABER

SABER is a multichannel infrared radiometer onboard the TIMED satellite. Limb profiles are taken from a circular orbit at 625 km inclined at 74° to the equator and cover a latitudinal range from 54◦ S to 82◦ N or 82◦ S to 54◦ N, depending on the phase of the yaw cycle [\(Russell III et al.,](#page-17-8) [1999\)](#page-17-8). One yaw cycle of SABER corresponds to 60 days, i.e. due to the full precession of the instrument during one cycle, this period is required to get a full coverage of local times.

SABER is equipped with two channels sensitive to OH emissions, i.e. the 1.6 μ m channel covers emissions from the $OH(5,3)/OH(4,2)$ transitions and the 2.0 µm channel covers emissions from the OH(9,7)/OH(8,6) transitions.

VER profiles from both channels are contained in the SABER Level 2a data products and will be used in our study. According to [Mertens et al.](#page-16-15) [\(2009\)](#page-16-15) the vertical resolution of the SABER VER profiles is approximately 2 km. Because the atmosphere is optically thin at altitudes above 80 km for wavelengths between 0.35 and 2.0 µm [\(Khomich](#page-16-11) [et al.,](#page-16-11) [2008\)](#page-16-11), the effect of self-absorption is negligible for the observed OH emission. Given this assumption, we can directly compare changes in our simulated OH concentrations to changes in the vertical VER profiles observed from both SABER channels.

In addition to measurements of the OH radiance, the latest SABER V2.0 data contain atomic oxygen profiles, which we use to study the impact of O quenching on the observed vertical shifts between the $1.6 \mu m$ and $2.0 \mu m$ VER profiles. As explained in [Mlynczak et al.](#page-16-8) [\(2013\)](#page-16-8), the SABER O concentrations are indirectly determined from the measured 2.0 µm VER profiles based on the following steady state assumption:

$$
\underbrace{k_1[H][O_3]}_{\alpha \text{VER}(2.0\mu\text{m})} = k_2[O][O_2],\tag{3}
$$

with k_2 denoting the reaction rate constant between O and O_2 and $k_1[H][O_3]$ being directly proportional to the observed VER. At first glance, this seems to introduce a circular reasoning in our attempt to correlate O concentrations with the vertical shifts between the $1.6 \mu m$ and $2.0 \mu m$ VER profiles. This could potentially introduce a spurious (i.e. nonphysical) correlation between both quantities, if the SABER model did not properly consider the real photochemistry and gaseous kinetics based on the steady state assumption (Eq. [3\)](#page-3-1). However, [Mlynczak et al.](#page-16-8) [\(2013\)](#page-16-8) find a close agreement between their derived day- and night-time O concentrations, which both rely on completely different methods. This indicates that the SABER model is reproducing physically meaningful O profiles. In turn, this should justify a direct comparison between SABER O concentrations and vertical shifts between both VER profiles, because we may suppose

Figure 1. Vertical OH(ν) profiles calculated from monthly averaged SD-WACCM4 model output for September 2010, 00:00 UTC at the 0° equatorial latitude. Left panel: absolute number concentrations. Right panel: normalised $OH(v)$ profiles.

that any correlation between both quantities represents a real dependency between them.

3 Methodology

For the first part of this study we have to address some general features of the vertical OH profiles to provide a basis for our analysis on the collisional quenching process. Figure [1](#page-4-2) shows vertical $OH(v)$ profiles that were simulated according to night-time conditions at equatorial latitudes based on our model approach. In general, we will limit our discussion on the night-time OH, because the relatively low abundances of daytime OH and the large Rayleigh scattering background make a comparison with OH daytime observations more difficult.

nounced vertical separation above the $OH(v)$ peak altitudes, In accordance with [von Savigny et al.](#page-17-0) [\(2012\)](#page-17-0) the vertical distribution of night-time $OH(v)$ follows single peak profiles that are shifted upwards with respect to their vibrational state. If we normalise each $OH(v)$ profile, the relative vertical shifts become clearly visible. In addition, we can notice a more prowhich according to [von Savigny et al.](#page-17-0) [\(2012\)](#page-17-0) is related to the steep vertical gradient in O concentrations and the associated more pronounced collisional deactivation of OH at the upper part of the OH emission layer. By comparison, the vertical shifts between the $OH(v)$ profiles are significantly less pronounced below the profile peak altitudes.

The systematic increase of the vertical $OH(v)$ shifts above the profile peak altitudes seems to favour this altitude region for our study on the collisional quenching with O. However, it is important to keep in mind that any changes in the vertical $OH(v)$ shifts are the convolved response to changes in the quenching and source species concentrations according to Eq. [\(2\)](#page-2-3). Therefore, finding the optimum reference points to compare the vertical shifts between two layers turns out to be less obvious than initially thought.

Another difficulty arises for the determination of relative vertical $OH(v)$ shifts from the rather coarse vertical resolu-

tion of our simulated OH profiles and observed SABER VER profiles. Despite this constraint on the vertical resolution, we can benefit from the significantly higher dynamic range of the calculated number densities and observed VERs.

To quantify the vertical $OH(v)$ shifts at the peak altitudes and above, we therefore define two different reference points, which we determine for each vertical $OH(v)$ profile:

D.1 weighted peak altitude: Zpkweighted

In analogy with [von Savigny and Lednyts'kyy](#page-17-1) [\(2013\)](#page-17-1) we weight the altitudes with the number density profile $N_{OH}(v, z)$ for each OH(v) layer:

$$
Zpk_{weighted}(v) = \frac{\int_0^\infty N_{OH}(v, z')z'dz'}{\int_0^\infty N_{OH}(v, z')dz'}
$$

D.2 shifted peak altitude: Zpk_{+HWHM}

To sense changes between the vertical $OH(v)$ shifts in the upper part of the OH layer, we interpolate the altitude above the profile peak of each $OH(v)$ layer, where $N_{OH}(v, z)$ has dropped by a factor of 0.5, i.e. the position that is shifted by the half width at half maximum (HWHM) above the profile peak.

For the SABER VER profiles, we can simply replace the number densities by the VERs in the above definitions.

4 Sources of error

While the inclusion of number densities according to D.1 and D.2 helps us to improve the vertical sensitivity of our model study, systematic departures between simulated and real number densities are a source of error for our investigation of the collisional quenching effects. The recently published study by [Smith et al.](#page-17-9) [\(2013\)](#page-17-9) indicates that WACCM tends to underestimate mesospheric ozone concentrations, which in turn will impact Reaction [\(R1\)](#page-2-4). In addition, WACCM tends to overestimate mesospheric temperatures according to [Smith](#page-17-10) [\(2012\)](#page-17-10), which will affect our calculation of the rate constant k_1 and absolute number densities from the SD-WACCM4 temperature pressure fields.

Inspection of Eq. [\(2\)](#page-2-3) shows that at least a linear departure in the $H + O_3$ source profiles from reality is not critical for our analysis, because it will cancel out in the calculation of the $OH(v)$ layer altitudes according to the above stated definitions. The situation is different for the quencher profiles, because any linear scaling of their concentrations cannot be completely factored out in Eq. [\(2\)](#page-2-3). An overestimation of temperatures should in principle lead to an underestimation of the absolute quenching gas concentrations based on the ideal gas law. In contrast, overly high temperatures will lead to an overestimation of the temperature dependent rate constant $k₁$. To get an estimate of the associated impact on the vertical shifts between different $OH(v)$ layers, we applied a constant offset of −20 K to our SD-WACCM4 temperatures.

Based on this approach, the impact appears to be minor, i.e. in the order of a few tens of metres with regard to our later analysis of profile shifts based on D.1. With respect to the O quenching species the simulated concentrations tend to be lower compared to concentrations derived from SABER as shown in [Smith et al.](#page-17-11) [\(2011\)](#page-17-11). According to the initial hypothesis on the impact of the collisional quenching with O on the vertical shifts between different $OH(v)$ layers, an underestimated rate of collisional quenching should result in less pronounced vertical shifts. Apart from the discrepancies in simulated O concentrations, the uncertainty of its collisional rate constant k_L^O will also affect our results. By comparison with the other quenchers, k_L^O has the greatest uncertainty. If we apply the upper and lower boundary of the uncertainty estimates of k_L^O from [Xu et al.](#page-17-4) [\(2012\)](#page-17-4), the changes to the vertical $OH(v)$ shifts based on D.1 range between about 100 m and 160 m with regard to our later analysis.

5 Simulated tidal signatures in OH and O: a monthly case example

Before we will address the temporal variability of vertical $OH(v)$ shifts, we have to reexamine systematic temporal changes of the entire OH emission layer and the O , O_2 quenching species for two reasons. First, we have to make sure that the temporal variability in the SD-WACCM4 data leads to a consistent evolution of the OH and O , O_2 species compared to previous studies. Second, this reexamination helps us to establish an expectation about the impact of temporal changes in the collisional quenching on the vertical OH (ν) shifts.

As motivated in the beginning, we will now consider a monthly case example around the September 2010 equinox, where the amplitude of the diurnal migrating tide maximises. For this month, a series of different model results is presented in Fig. [2.](#page-5-1) The global distribution of the integrated total column of all $OH(v = 1, 2, ..., 9)$ layers is displayed for 00:00 UTC in panel a. A general eastward decrease in the integrated OH concentrations is clearly visible. In terms of LSTs, this corresponds to a decrease of integrated OH concentrations over the course of the night. In addition, the OH concentrations are generally high at equatorial latitudes and minimise around $\pm 30^\circ$ latitude, which is consistent with the study of [Marsh et al.](#page-16-6) [\(2006\)](#page-16-6) and other observational studies stated therein. The steep decrease of integrated OH concentrations at the outer latitudinal and longitudinal margins marks the terminator between day- and night-time conditions.

Weighted OH peak altitudes Zpkweighted according to our previous definition D.1 are displayed in panel b. A systematic night-time increase in the weighted OH peak altitudes by up to 4 km is again clearly visible. Accordingly, we find a significant anti-correlation between OH peak altitudes and concentrations in comparison with panel a. Indeed, previous

Figure 2. Monthly averaged model results around September 2010 equinox. **(a)** Vertically integrated number density of simulated \sum^9 $\sum_{i=1}$ OH(v_i). **(b)** Weighted peak altitudes of simulated OH emission layer according to our definition D.1. **(c)** O concentrations weighted with vertical OH(ν = 9) profiles.

studies based on observations made with the high-resolution Doppler imager (HRDI) instrument and the Wind Imaging Interferometer (WINDII) instrument onboard the upper atmosphere research satellite (UARS) revealed the same coherent anti-correlation between OH peak altitudes and integrated concentrations. Following [Liu and Shepherd](#page-16-16) [\(2006\)](#page-16-16) and the stated referenced therein, this anti-correlation may be driven by the vertical motions associated with tides or other processes (see also [Cho and Shepherd,](#page-16-17) [2006\)](#page-16-17).

As with the determination of $OH(v)$ profile peak altitudes, several possibilities exist to quantify temporal changes in the quenching species concentrations. The simplest method is to look at the diurnal evolution of a quenching species at a constant height level. However, this method neglects any changes of the quenching species concentrations that arise from the vertical motion of the entire OH layer. To account for this, we may determine the quenching species concentration at a fixed reference point of the OH layer. Again, this method is still rather simple, because the collisional quenching is not constrained to a fixed point at the OH layer. Thus, a more sophisticated approach is to quantify the collisional quenching by weighting the vertical quencher profiles with the corresponding OH profiles (i.e. replace z' in D.1 with the number density $N_Q(z')$ of the quenching species).

For our monthly case example, we applied the latter approach for the O concentrations in Fig. [2c](#page-5-1). Accordingly, the equatorial weighted O concentrations show a pronounced maximum before midnight, which has also been confirmed by other observational studies (e.g. see [Smith](#page-17-6) [et al.,](#page-17-6) [2010\)](#page-17-6) and should therefore lead to a pronounced collisional quenching of the OH emission layer at those LSTs. Furthermore, [Smith et al.](#page-17-6) [\(2010\)](#page-17-6) report another wavenumber 1 type feature at $\pm 30^\circ$ with opposite phase, which at least seems to be reflected at 30◦ S in our simulations. Of course, we have to bear in mind that we are considering a single month only and that the temporal variability of the OH emission layer is also affecting our weighted O concentrations. Moreover, following the study of [Lu et al.](#page-16-18) [\(2012\)](#page-16-18) the magnitude of the tidal amplitude seems to be slightly underestimated by WACCM4. Despite this slight underestimation, the tidal signatures in the OH profile weighted O as well as the OH concentrations of our monthly case example show consistent characteristics with previous observations and should therefore serve as a plausible testing ground for the initial hypothesis on the collisional quenching.

If we expand our monthly case example to a full seasonal cycle, we would expect a semi-annual oscillation in the atomic oxygen concentrations, which are following the temporal changes in the amplitude of the diurnal-migrating tide as discussed before. Indeed, this oscillation is clearly visible in the simulated O concentrations, as shown in the left panel of Fig. [3.](#page-7-0) For this figure we choose an LST bin between −1 and 0 h around equatorial latitudes. Each curve represents one of the above discussed methods to quantify the O concentrations, i.e. O determined at the 0.241 Pa pressure level (green line), O interpolated at the selected $OH(v = 5)$ and $OH(v = 9)$ weighted profile peak altitude according to our definition D.1 (blue lines), O interpolated at the HWHMshifted position above the profile peak (see definition D.2 and red lines), and O weighted with either the selected $OH(v = 5)$ or $OH(v = 9)$ profile (black lines). Because we are interested in the relative temporal changes in the quenching species, each curve is subtracted by its minimum value (see legend) to allow for a better intercomparison.

In addition to atomic oxygen, we include the seasonal variability of molecular oxygen in the right panel of Fig. [3.](#page-7-0) Interestingly, we can find another semi-annual oscillation in phase with the atomic oxygen species, if we consider the curves that do not refer to the fixed 0.241 Pa level. Despite the lower quenching efficiency of O_2 compared to O , the higher absolute O_2 abundances will at least partially compensate this. Because of the increasing O_2 number density with decreasing altitude, the collisional deactivation of excited $OH(v)$ through O_2 quenching will be most pronounced at the lower part of the OH emission layer. Vice versa, the O quenching is rapidly decreasing at the lower part of the OH emission layer due to the steep vertical gradient in O number densities, thus, O_2 quenching is expected to be the dominant process of vibrational deactivation of OH at the bottom of the OH emission layer. This already seems to indicate an important role of the seasonality in the $O₂$ quenching with regard to the temporal evolution of vertical $OH(v)$ shifts.

6 Seasonal evolution of OH layer shifts

6.1 Sensitivity study

In the following, we will compare relative changes in the vertical shifts between the OH(ν = 9) and OH(ν = 5) profiles. We select these two vibrational states because each of them contributes to emissions, which can be observed by either the 1.6 or 2.0 µm SABER channel. Ideally, one must consider that each SABER channel captures a mixture of emissions that belong to two different transition bands. However, because the difference in vibrational levels between each transmission is limited to $\Delta v = 1$, we assume that we can neglect the effect of profile mixing for each channel, if we are interested in the relative vertical shift between both (mixed) OH profiles. The vertical shift between a simulated $OH(v = 9)$ and $OH(v = 5)$ profile will be calculated from the difference between either their weighted peak altitudes,

$$
\Delta Zp k_{weighted} = Zp k_{weighted} [OH(\nu = 9)] -
$$

Zp k_{weighted} [OH(\nu = 5)], (4)

or from the difference between the HWHM-shifted altitudes above the profile peaks:

$$
\Delta Zpk_{+HWHM} = Zpk_{+HWHM}[OH(\nu = 9)] -
$$

Zpk_{+HWHM}[OH(\nu = 5)]. (5)

The vertical shifts between the SABER 1.6 µm and 2.0 µm VER profiles are determined in the same way.

We will now investigate the seasonal variability of the relative vertical shifts between our simulated OH($\nu = 9$) and $OH(v = 5)$ profiles, which we denote as $OH(9;5)$ profile shifts in the following. For this task, we perform three different model runs:

- **–** all quenching species are activated
- **–** O quenching species is deactivated
- **–** O² quenching species is deactivated

Figure 3. Seasonal variability of simulated atomic and molecular oxygen concentrations from April 2010 to June 2011 according to the peak altitudes (red and blue lines), O concentration weighted with OH concentrations (black lines). From each curve the offsets listed in the legend were subtracted to allow a better intercomparison of the temporal changes. following definitions: O concentration at fixed pressure level (green line), O concentration interpolated at +HWHM-shifted and weighted

This allows us to study the impact of collisional quenching on the OH(9;5) profile shifts for both species.

the seasonal variability are much more pronounced and the run leads to a still no less clear. So far, we find the best agreement with the ini-
less clear. So far, we find the best agreement with the ini-
2011, we can find a further semi-annu weighted peak altitude definition D.1. 3 The results from our three model runs are shown in Fig. 4. Figure [4a](#page-8-0) displays the seasonal evolution of OH(9;5) profile shifts for the first model run (i.e. complete quenching considered). The left axis/solid line refer to the relative vertical shifts between weighted peak altitudes according to Eq. [\(4\)](#page-6-1). The right/dashed line refer to the relative vertical shifts at the upper part of the $OH(9;5)$ layers according to Eq. [\(5\)](#page-6-2). If we concentrate on the solid line first, we find indeed a semiannual oscillation in the $OH(9;5)$ profile shifts that is in phase with the observed changes in the O and O_2 concentrations according to Fig. [3.](#page-7-0) On the other hand, if we look at the upper part of the OH(9;5) layers (dashed line), the fluctuations in response to the seasonal changes in the quenching species is tial hypothesis on the collisional quenching process for our

the deactivated O quenching process. For both lines, we find July and October due to t Similarly to panel a, panel b shows the model run with a significant decrease in the OH(9;5) profile shifts, which (see panel c), the impact of the deaction again is consistent with the initial hypothesis. On the other is less than one half. hand, we still find a persisting semi-annual oscillation for the solid line (i.e. OH(9;5) profile shifts with respect to weighted peak altitudes) that is superimposed by another temporal maximum around mid January 2010. The seasonal response at the upper part of the $OH(9;5)$ layers (dashed line) remains less clear. If we now subtract the results from the model runs with and without O quenching, we find a clear semi-annual response in the OH(9;5) profile shifts according to the solid line in panel c. Interestingly, if we compare the increase in the OH(9;5) profile shifts between July and October between panels b and c, the contribution of the O quenching process to the temporal changes in the OH(9;5) profile shifts is just

as to stady the inpact of confisional quenching

run with deactivated O quenching. With regard to the initial

run with deactivated O quenching. With regard to the initial ts from our three model runs are shown in Fig. 4. bypothesis, this suggests that we cannot address the observed slightly above the remaining temporal changes for the model seasonality in OH(9;5) profile shifts to the modulation in the collisional O quenching only.

e collisional quenching process for our tical OH(9;5) profile shifts due to the switching between the definition D.1. deactivated and activated O_2 quenching. In comparison with We therefore repeat the same investigation of the collisional quenching process for the O_2 quencher. In analogy with panel c, panel d shows the difference in OH(9;5) profile shifts when subtracting the results from the model runs with activated and deactivated O_2 quenching. Again, the upper part of the OH(9;5) layers shows strong fluctuations (dashed lines), thus we will limit our discussion to the relative shifts between weighted peak altitudes (solid line). First of all, we find that the deactivation of the O_2 quenching in our model run leads to a still noticeable decrease in the vertical OH(9;5) profile shifts. If we neglect the maximum around January 2011, we can find a further semi-annual response in the verthe seasonal change in the $OH(9;5)$ profile shifts between July and October due to the deactivation of the O quenching (see panel c), the impact of the deactivation of O_2 quenching is less than one half.

> $\frac{1}{2}$ another temporal effect of O and O₂ quenching according to panels c and d ently, taking the sum of panels c and d leads to an improved With respect to the model run that considers all quenching species (see panel a), we notice that simply adding the still leads to considerably smaller vertical OH(9;5) profile shifts, as shown in panel e. On the other hand, the agreement in the seasonal variability between panels a and e is quite good. This becomes evident, if we determine the best scaling factor between both functions in a least-squares sense. Accordingly, the grey line in panel e denotes the ΔZp k_{weighted} profile shift values from panel a divided by 1.438. Apparagreement in the seasonal variability with respect to panel a

equatorial range between $\pm 7.5^{\circ}$ and the LST range from -1 to 0h. Solid line/left axis: OH(9;5) vertical shifts between weighted peak altitudes (see definition 3). Dashed line/right axis: OH(9;5) vertical shifts between the +HWHM-shifted peak positions (see definition D.1). on/off. (d) Difference in OH(9;5) vertical shifts when switching O₂ quenching on/off. (e) Sum of (c) and (d) (black line). In addition, ΔZ pk_{weighted} from (a) divided by 1.438 shown by grey line. (f) Full peak width of the vibrationally integrated OH(v) layer. (g)/left axis: Full peak widths of OH(9) and OH(5) layers (dashed and solid grey lines). (g)/right axis: Relative difference between the full peak widths of
the OH(0) and OH(5) layers (bleak solid line) **Figure 4. (a–d)** Seasonal variability of vertical OH(9;5) profile shifts from April 2010 to June 2011 for different model runs within the **(a)** Full quenching model run. **(b)** Deactivated O quenching model run. **(c)** Difference in OH(9;5) vertical shifts when switching O quenching the OH(9) and OH(5) layers (black solid line).

run in panel a. As noticed before, the impact of N_2 quenching in the OH rather than considering the effect of deactivating either O or $O₂$ quenching only. This again suggests the importance of the O_2 quenching to the seasonal variability. Still, the question remains, why the sum of panels c and d is smaller by a factor of 1.438 compared to the complete quenching model is an insufficient explanation, i.e. it leads to a difference between 40–50 m, if we subtract a model run with deactivated N_2 quenching from the complete quenching case in panel a (not shown). This indicates that the combined effect of O and $O₂$ quenching is larger than the sum of their individual contributions.

As discussed in the beginning, seasonal changes in the vertical $H + O_3$ profiles will affect the OH emission layer width, which in turn will also affect the OH(9;5) profile shifts. In addition to the combined effect of O and O_2 quenching, this could provide another mechanism that is driving the temporal variability. The seasonal evolution of the OH emission layer width is shown in panel f of Fig. [4.](#page-8-0) In this case, we deter-

sidering the effect of deactivating either O or mine the width of the vertical profile by the full width at half only. This again suggests the importance of maximum (FWHM) to account for changes above and below ng to the seasonal variability. Still, the ques-
the profile peak altitude. Accordingly, we find a pronounced B compared to the complete quenching model ues, which is coherent with the observed additional increase increase around the mid of January 2011 in the FWHM valin the $OH(9;5)$ profile shifts for the deactivated O quenching case (panel b). This gives an explanation, why the drop in the vertical OH(9;5) profile shifts is less pronounced after the winter solstice according to panel a. Furthermore, the larger extent of the OH profile width may also favour the rate of collisional O_2 quenching, which could explain the coherent response according to panel d of Fig. [4.](#page-8-0) On the other hand, a coherent semi-annual variability with respect to the ΔZp k_{weighted} profile shift values in panel a is not evident, which strengthens the argument of the combined effect of O and $O₂$ quenching as the dominant driving mechanism of the seasonal variability in the OH(9;5) profile shifts.

> Finally, we also consider the relative changes of the OH(9) and OH(5) peak widths, which should particularly influence

the OH(9;5) profile shifts above the profile peak altitudes. The seasonal evolution of each peak width is shown by the grey lines in panel g of Fig. [4](#page-8-0) (see caption). The difference we receive by subtracting both temporal evolutions with each other is shown by the black solid line. We find that the large relative changes in the profile widths around October 2010 and May 2011 are coherent with the observed jumps in the OH $(9;5)$ profile shifts at the upper part of the OH $(9;5)$ layers (see dashed line in Fig. [4a](#page-8-0)), i.e. the vertical shifts ΔZ pk_{+HWHM} appear to respond more sensitively to relative changes in the OH(9;5) profile widths.

6.2 Comparison with SABER

We will now focus on the seasonal variability of the vertical shifts between the SABER 1.6 and 2.0 μ m VER profiles for the period from January 2009 to December 2011. In analogy with our sensitivity study, we choose the same -1 to 0 h LST bin for the results presented in Fig. [5.](#page-10-0) Here, each point represents the mean value based on three matching yaw-cycles between 2009 to 2011. Each error bar denotes the corresponding standard deviation. Panels a and b show the seasonal variability in the VER profile shifts for two equatorial latitude bins. Again, the solid line refers to the vertical shifts between weighted peak altitudes according to Eq. [\(4\)](#page-6-1) (left axis) and the dashed line refers to the vertical shifts at the upper part of the VER profiles according to Eq. [\(5\)](#page-6-2) (right axis). The seasonal variability of derived O concentrations is displayed in panels c and d. The black line shows the O concentrations at 90 km altitude (left axis). The grey dotted and dashed lines show the VER profile weighted O concentrations with respect to the 1.6 and 2.0 μ m channel (right axis). Panels e and f show the seasonal variability of the 1.6 and $2.0 \,\mu m$ VER profile widths (dashed and dotted, left axis) as well as their relative difference (black solid line, right axis).

First of all, we notice that a semi-annual oscillation – with maxima around May and October – is also present in the SABER VER profile shifts. Indeed, we find another coherent semi-annual oscillation in the O concentrations for the 0° to $10°$ S bin. With regard to the $0°$ to $10°$ N bin a faint semiannual structure is present, but the overall change is dominated by an annual oscillation. Interestingly, in comparison with the changes in the SABER VER profile shifts, the semiannual response is more dominating in the 10◦ N rather than the 10◦ S latitude bin. Again, this indicates that the consideration of O quenching alone cannot sufficiently explain the seasonal variability of SABER VER profile shifts. In contrast to our model results, we cannot directly rule out that changes in the sources gases may significantly affect the seasonality of the observed VER profile shifts, because of the limited spectral bandwidth of SABER, which prevents us from sensing all Meinel bands of the OH emission.

If we consider the relative changes of the VER profile widths according to the black solid line in panels e and f of Fig. [5,](#page-10-0) we can find a similar coherent response in the vertical shifts at the upper part of both VER profiles (see dashed line in panels a and b), which again shows the stronger sensitivity of this profile shift definition to changes in the relative profile shapes.

7 Diurnal evolution of OH layer shifts

7.1 Sensitivity study

In analogy with our analysis of the seasonal variability of the vertical OH(9;5) profile shifts in the previous section, we perform the same three model runs where we consider the full-quenching case, the deactivation of O quenching, and the deactivation of O_2 quenching. To improve our later comparison with the observed diurnal variability from SABER, we adjust the temporal averaging period in our model runs to the same period that is required for a full SABER yaw cycle.

Accordingly, Fig. [6](#page-10-1) shows the diurnal variability of both simulated quenching species around the September 2010 equinox at equatorial latitudes, where the amplitude of the diurnal migrating tide maximises. Again, we use the same definitions to quantify changes in the O and O_2 concentrations as in Fig. [3.](#page-7-0)

While the different definitions of O concentrations only lead to a slight phase shift in the temporal evolution of the O concentrations of up to one hour, the different definitions of $O₂$ concentrations can result in quite different diurnal evolutions. With regard to the systematic increase in the nighttime OH peak altitudes (Fig. [2b](#page-5-1)), the decrease in the OH profile weighted O_2 concentrations (black lines) and interpolated $O₂$ concentrations at the weighted OH peak altitude (blue lines) appears to be the most consistent.

The results from our three model runs are shown in Fig. [7,](#page-11-0) with the solid lines (left axis) referring to the OH(9;5) profile shift according to Eq. [\(4\)](#page-6-1) and the dashed lines (right axis) referring to Eq. [\(5\)](#page-6-2) correspondingly. Panel a displays the OH(9;5) profile shifts for the first model run, which considers all quenching species. Panel b shows the OH(9;5) profile shifts for the model run with deactivated O quenching. Panels c and d show the difference in OH(9;5) profile shifts, if we subtract either the model run with deactivated O or $O₂$ quenching from the full quenching model run shown in panel a. Similarly to the previous section, we also include the sum of panels c and d in panel e.

Keeping in mind the initial hypothesis on the effect of collisional quenching, we would expect that the vertical OH(9;5) profile shifts should maximise shortly before midnight according to the maximising O concentrations. However, neither of both OH(9;5) profile shift definitions matches with this expectation according to panel a. Furthermore, we notice that the diurnal variability in the OH(9;5) profile shifts is rather opposite for both definitions. If we switch off the O quenching according to our second model run (panel b), the vertical OH(9;5) profile shifts are significantly reduced

Figure 5. SABER results based on 3 years of observation. Each point represents the mean value of three yaw cycles according to each year. Solid line/left axis: Vertical VER profile shifts between weighted peak altitudes according to Eq. [\(4\)](#page-6-1). Dashed line/right axis: Vertical VER profile shifts between the +HWHM-shifted peak positions according to Eq. (5). (c–d) O concentrations at 90 km (left axis, black solid line) peak widths of 1.6 and 2.0 µm VER profiles (dotted and dashed grey lines). (e-f)/right axis: Relative difference between the full peak widths of the 1.6 and 2.0 μ m VER profiles (black solid line). the standard deviation is denoted by the error bars. **(a–b)** Seasonal variability in the vertical shifts between the 1.6 and 2.0 µm VER profiles. and 1.6 as well as 2.0 µm VER profile weighted atomic oxygen concentrations (right axis, dotted and dashed grey lines). **(e–f)**/left axis: Full

Figure 6. Diurnal variability of simulated atomic and molecular oxygen concentrations. The same denotations apply that are used for the seasonal variability of both species in Fig. [3.](#page-7-0) The temporal averaging interval ranges from 15 September 2010 to 15 November 2010 to match the same period in our simulations that is needed for one complete SABER yaw cycle.

variability in Fig. 4b. Moreover, if we consider the impact of collisional O queno as it was also the case for our investigation of the seasonal the collisional O quenching according to panel c, a coherent response to the diurnal evolution of O is clearly visible for the ΔZp k_{weighted} values (solid line). For the HWHM-shifted positions above the profile peaks (dashed line), we can still

so the case for our investigation of the seasonal find a significant internal variability, such that the impact of collisional O quenching again remains less clear.

> As with our analysis of the seasonal variability, the collisional O_2 quenching is also significantly affecting the vertical OH(9;5) profile shifts according to Fig. [7d](#page-11-0). In comparison with the collisional O quenching the effect is still smaller with regard to the weighted peak altitudes. Furthermore,

Figure 7. Simulated diurnal evolution of vertical $OH(9,5)$ profile shifts at equatorial latitudes for the same averaging time as in Fig. [6.](#page-10-1) (a) (Eq. [4\)](#page-6-1), the dashed line refers to vertical shifts with respect to peak altitudes + HWHM (Eq. [5\)](#page-6-2). (b) Vertical shifts based on a model run with deactivated O quenching. (c) Difference between (a) and (b). (c) Difference between a full quenching model run and a model run with Full peak widths of OH(9) and OH(5) layers (dashed and solid grey lines). (g)/right axis: Relative difference between the full peak widths of the $OH(9)$ and $OH(5)$ layers (black solid line). Vertical shifts based on a model run including all quenching terms. The solid line refers to peak shifts with respect to weighted peak altitudes deactivated O_2 quenching. **(e)** Sum of **(c)** and **(d)** (black line). **(f)** Full peak width of the vibrationally integrated OH(*v*) layer. **(f)**/left axis:

consider the results based on the HWHM-shifted positions, scaling factor we receive for the seasonal the effect of the deactivation of O_2 quenching strongly exthe relative changes due to the deactivation of O_2 quenching remain rather constant after −2 h. In contrast, if we ceeds the corresponding effect for O. Moreover, the early $Zpk_{weighted}[OH(5)]$ positions are even higher than those of the $\text{Zpk}_{\text{weighted}}[\text{OH}(9)]$ positions, which leads to the negative values before −3 h.

If we consider the sum of panels c and d, as shown in panel e, the resulting OH(9;5) profile shifts are again significantly smaller compared to the model run in panel a which considers all quenching species simultaneously. Inter-

s due to the deactivation of O_2 quench-estingly, if we search for the best scaling factor between both constant after $-2h$. In contrast, if we panels, we obtain a factor of 1.430, which is very close to the scaling factor we receive for the seasonal variability. On the other hand, the agreement between the diurnal evolution according to the sum of panels c and d and the scaled panel a, which is denoted by the grey line in panel e, is less clear for the early evening hours. During the same hours, we notice a strong shrinking of the entire OH emission layer by up to 4 km due to the night-time evolution of the $H + O_3$ source gases according to Fig. [8a](#page-12-0). This may also provide an explanation for the departure between the sum of panels c and d and scaled panel a, presuming that the combined effect of O

of OH(9) and OH(5) layers (dashed and solid grey lines). (b)/right axis: relative difference between the full peak widths of the OH(9) and OH(5) layers (dashed and solid grey lines). (b)/right axis: relative difference be F_1 and F_2 and F_3 and F_4 and solid grey lines). **Figure 8. (a)** Diurnal variability of OH peak widths for the same spatial and temporal bin considered in Fig. [7.](#page-11-0) **(b)**/left axis: full peak widths OH(5) layers (black solid line).

Figure 9. Corre[la](#page-6-2)tion plots of vertical OH(9;5) profile shifts (left panels Eq. (4), right panels Eq. 5) vs. OH(9) profile weighted O concentrations. The upper panels show the correlation between OH(9;5) profile shifts and O concentrations for the full quenching model run (similarly deactivated O quenching model runs. Correlation coefficients are shown in the legend and denoted with the asterisk symbol *, if they were found to be significant according to a 90% significance level. α to be significant according to $\alpha > 0$ to significance rever. to Fig. [7a](#page-11-0)). Similarly to Fig. [7c](#page-11-0), the lower panels consider the difference in vertical OH(9;5) profile shifts between the full quenching and the

according to Fig. 7. With regard to the OH(9;5) profile shifts correlation plots in Fig. 9, we find no significant corre these are again strongly correlated with the relative changes $OH(9)$ concentrations for all seasons and both profile in the OH(9;5) profile widths according to Fig. [8b](#page-12-0).
We smooth and properly to the full user of simula and O_2 quenching can be described by its linearly scaled sum based on ΔZpk_{+HWHM} values (dashed lines), we find that

We expand our analysis to the full year of simulated $OH(v)$ populations and summarise the found correlations between vertical OH(9;5) profile shifts and quenching species con-

 σ_2 quenching can be described by its linearly scaled sum centrations in Fig. [9](#page-12-1) and Fig. [10.](#page-13-1) Following the displayed correlation plots in Fig. 9, we find no significant correlation between the vertical OH(9;5) profile shifts and weighted OH(9) concentrations for all seasons and both profile shift definitions. A weak positive correlation is only visible, if we include all data points in panel a. As with our equinoctial case example according to Fig. [7c](#page-11-0), the correlation between vertical OH(9;5) profile shifts and weighted O concentrations

Figure 10. Similar to Fig. [9](#page-12-1) but referring to the O_2 quenching species.

significantly improves, if we compare the relative changes between the model runs with activated and deactivated O quenching for weighted peak altitudes (Fig. [9c](#page-12-1)). In contrast, the correlation remains poor, if we consider OH(9;5) profile shifts at the upper part of both layers (Fig. [9d](#page-12-1)).

Figure [10](#page-13-1) shows the corresponding correlations for the $O₂$ quencher. In contrast to the O quencher, the correlations with the vertical OH(9;5) profile shifts are exceptionally high. Of course we have to bear in mind that the systematic increase in the OH night-time altitudes (see Fig. [2b](#page-5-1)) will also be reflected in the systematic decrease in OH(9) weighted O_2 concentrations. However, for the relative changes between the model runs with activated and deactivated $O₂$ quenching, we still find a significant correlation in Fig. 10c, respectively anti-correlation in Fig. [10d](#page-13-1).

In summary, the night-time evolution in the $OH(9;5)$ profile shifts can hardly be explained by the process of collisional quenching with atomic oxygen only. Again, the inclusion of molecular oxygen quenching further improves the correlation with the $OH(9;5)$ profile shifts. In addition, the simultaneous strong decrease of the entire OH layer width, driven by the $H + O_3$ source profiles, will further impact the night-time evolution of OH(9;5) profile shifts. Interestingly, the systematic changes in OH peak altitudes and associated changes in O_2 concentrations show a very strong correlation with the temporal changes in the vertical OH(9;5) profile shifts.

7.2 Observed diurnal variability from SABER

For the SABER observations we first consider the same yawcycle that was also used for the model simulations presented in Fig. [7](#page-11-0) and compare the relative shifts between the 1.6 and $2.0 \,\mu$ m VER profiles with the OH-VER weighted atomic oxygen profiles similarly to our analysis of the seasonal variability. For the observed diurnal variability, it is important to note that the temporal evolution in the observed relative OH profile shifts may significantly differ from our model results because of the existence of additional non-migrating tides as being reported by [Xu et al.](#page-17-12) [\(2010\)](#page-17-12) from SABER observations at lower latitudes. These tides would complicate a direct comparison with our model results, since we have to extract the temporal evolution from the longitudinal variability of our 00:00 UTC model outputs. However, despite the possible existence of non-migrating tides, this does not prevent us from testing the hypothesis on the impact of collisional quenching with regard to the temporal variability of the O quenching species.

In analogy with Fig. [7](#page-11-0) the SABER results are shown in Fig. [11](#page-14-1) for the same yaw-cycle and for two latitudinal bins nearby the equator. Indeed, the night-time evolution of relative OH profile shifts looks quite different compared to our modelled vertical OH(9;5) profile shifts. Furthermore, the amplification of atomic oxygen before midnight is not as evident as in our model results. Despite these discrepancies, we would expect from the systematic night-time decrease in atomic oxygen a corresponding feedback in the vertical VER

Figure 11. Diurnal variability according to SABER OH observations. $(a-b)$ Relative vertical shifts between VER(1.6 µm) and VER(2.0 µm) right axis, dotted and dashed line). (e-f) FWHM of VER(1.6 μm) profile (grey dashed line), FWHM of VER(2.0 μm) profile (grey solid line) and the difference Δ FWHM between both FWHM values (black solid line). profiles in analogy with Fig. [7.](#page-11-0) **(c–d)** O concentrations at 90 km level (left axis, solid line) and weighted with VER(1.6 μ m) and VER(2.0 μ m)

notice strong changes in the relative peak widths according profile shifts, which clearly is not the case. Again, we also to panels e and f of Fig. [7](#page-11-0) that are partially reflected in the night-time changes of VER profile shifts.

1275

12007535 142500

 $3.0e + 11$

 $4.1e+11$

 $6.4e+11$

∆Zpkweighted (m)

 \circ)

1350

1425

1500

If we expand our analysis to a full seasonal cycle, the missing correlation between VER profile shifts and O concentrations remains. Accordingly, the process of collisional O quenching appears to be insufficient to explain the night-time evolution of the OH VER profile shifts, which also agrees with our model expectations in a qualitative sense.

8 Summary and conclusion

Following the hypothesis that the process of collisional quenching is significantly affecting the vertical shifts between different $OH(v)$ layers, this study investigated the im-

h clearly is not the case. Again, we also pact of the temporal variability of the collisional quenching on the seasonal as well as the diurnal evolution of the vertical shifts between the OH(9) and OH(5) layer. This was done by establishing an updated quenching model, which uses the model output from a state-of-the-art 3D chemical climate model (SD-WACCM4) to simulate the temporal variability of both $OH(v)$ layers.

> By comparing different model runs, which consider either all quenching species or neglect the collisional quenching process by O or O_2 , we could study the actual impact of the temporal variability of O and O_2 quenching on the vertical OH(9;5) profile shifts. For the seasonal variability we find that both quenchers have a noticeable impact on the vertical OH(9;5) profile shifts, which manifests in a semi-annual variability that is following the temporal evolution of the diurnal migrating tide at the equator. Furthermore, the simultaneous quenching of both species results in about $1.4 \times$ larger

Figure 12. Correlation between relative night-time VER shifts and O concentrations from SABER observations in analogy with Fig. [9.](#page-12-1)

vertical OH(9;5) profile shifts than we would receive from the sum of their individual contributions. With regard to previous studies that were mainly focusing on the effect of collisional quenching with O, this indicates the important role of the combined effects of O and O_2 quenching on the vertical structure of the OH layer. In addition, the strong change in the OH emission layer widths around January 2011 demonstrates that temporal changes in the $H + O_3$ profiles provide another mechanism to affect the OH(9;5) profile shifts.

We found further evidence of the same seasonality in the OH(9;5) profile shifts in the SABER observations, even though the coherence with changes in the derived SABER O concentrations is not always as clearly pronounced as it is the case in our model simulations. This could reflect the stronger temporal variability in the true $H + O_3$ profiles, which is competing with the temporal changes in the collisional quenching processes.

With regard to the diurnal variability, our model study as well as our analysis of SABER VER profiles clearly show that the collisional quenching process of OH with O is insufficient to describe the temporal evolution alone. Again, the consideration of the combined effect of O and O_2 quenching is required to describe the temporal shifts in the OH(9;5) profiles. In addition, the model results suggest that the nighttime evolution of the $H + O_3$ profiles is significantly affecting the vertical OH(9;5) profile shifts for the first half of the night.

In summary, according to this study the effect of the collisional quenching does have a noticeable impact on the temporal variability of OH(9;5) profile shifts at the equator, but requires the simultaneous consideration of the O and O_2 quenching species to provide a proper description of the observed temporal evolution. While the O and O_2 quenching appears to be modulated in phase with regard to their seasonal evolution, the need of the simultaneous consideration of both quenching species becomes even more important for the diurnal evolution, where the modulation of O and O_2 can differ substantially, such that we cannot find a meaningful correlation to changes in the O quenching alone.

Critical assumptions of this study include the following:

- **–** absolute number densities derived from SD-WACCM4 temperatures (too warm),
- **–** simulated diurnal variability described by zonal variation of daily model output (conflicting with nonmigrating tides),
- **–** SABER O concentrations indirectly determined from OH Meinel emission.

Furthermore, the large uncertainty in the collisional quenching rate constant k_L^O appears to have the most significant impact on our results, showing the need for improving our understanding of the collisional quenching process.

Because of the manifold of transition bands being observed by different ground-based instruments, a thorough understanding of the driving processes of the variability of OH emission altitudes is crucial for the intercomparison and interpretation of long-term data sets. This in particular applies for studying of mesopause temperature trends by means of OH rotational temperature measurements (see [Beig et al.](#page-16-19) [\(2003\)](#page-16-19); [Beig](#page-16-20) [\(2011\)](#page-16-20) for a comprehensive review on this topic). Further improvements in the modelling of the tidal variability at the mesopause as well as the inclusion of a multi-year analysis of the features that have been discussed here would contribute to a better quantitative understanding of the systematic biases between different observational long-term studies.

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