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Retrieval of Martian ozone and dust from SPICAM spectrometer for MY27–MY28

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Ozone (O₃) is important in the stabilisation of CO₂ in the Martian atmosphere and thus it is important to study the spatio-temporal variability of O₃. We retrieve two years of total columnar O₃ from raw spectral data provided by the SPectroscopy for the Investigation of the Characteristics of the Atmosphere of Mars (SPICAM) instrument aboard the *Mars Express*. The seasonal variability is studied in tropical, mid- and high latitudes and is compared with simulations by a photochemical coupled general circulation model (GCM). The high latitudes exhibit the largest seasonal variations in O₃, with a winter high and a summer low and a comparison with GCM results is good in general. We have studied the correlation of O₃ with dust, retrieved simultaneously from SPICAM observations. In southern tropical latitudes, the columnar O₃ is seen to increase during a global dust storm year (Martian year (MY) 28) compared to the O₃ column values during a year without global dust storm (MY27), although the water vapour column between these years remains unchanged. This indicates the radiative impact of dust on O₃ and its retrieval. We also study the ozone–carbon monoxide correlation as a tracer of dynamics. The dynamical contribution to the O₃ column is found to be the highest during winter over the southern polar region.

Keywords. Ozone–dust correlation; retrieval of ozone; O₃–CO correlation; SPICAM; Martian ozone.

1. Introduction

Stability and composition of the Martian atmosphere are controlled by photochemical processes (Krasnopolsky 2006), of which ozone (O₃) is one of the most active species. Hydroxyl ions (OH), which are a sink for O₃ and carbon monoxide (CO), have not been extensively observed on Mars. Thus, O₃ and CO and their correlation have become important options for use as tracers for OH. O₃ is also important in determining the habitability of Mars, as it modulates the surface ultraviolet (UV) flux. It was detected for the first time by the absorptions of the Hartley band in the UV and the IR features by the UV spectrometer aboard

Mariner 7 flyby (Barth and Hord 1971). Since then, there have been many Earth- and space-based remote-sensing observations which have explored O₃ in the Martian atmosphere. To mention a few: The UV spectrometer aboard Mariner 9 (Barth *et al.* 1973), spectroscopy for the investigation of the characteristics of the atmosphere of Mars (SPICAM) aboard the *Mars Express* (Perrier *et al.* 2006) and MARs Colour Imager (MARCI) (Clancy *et al.* 2016) aboard the *Mars Reconnaissance Orbiter* (MRO). Ground-based studies such as the heterodyne instrument for planetary wind and composition (HIPWAC) (Fast *et al.* 2009) of the NASA Goddard Space Flight Center InfraRed Heterodyne Spectrometer (IRHS) (Fast *et al.* 2006),

the hubble space telescope faint object spectrograph (Clancy *et al.* 1996) and a high-resolution infrared telescope (CSHELL) at the NASA infrared telescope facility (Novak *et al.* 2002) provided significant insight into Martian O₃. These observations have improved our knowledge about the seasonal and temporal variations of O₃.

The global distribution of O₃ (longitudinal averaged) is reported from remote-sensing observations by Perrier *et al.* (2006) and Clancy *et al.* (2016). In recent years, vertical profiles of O₃ measured using a SPICAM UV spectrometer have shown the presence of two distinct O₃ layers at low-to-mid latitudes (Lebonnois *et al.* 2006). The first peak occurs near the surface and the second nocturnal peak between 25 and 60 km was detected for the first time by solar occultation observations from Phobos (Blamont and Chassefiere 1993). Based on the stellar occultation measurements by SPICAM, Montmessin and Lefèvre (2013) reported a third layer of O₃ over the winter south pole between 35 and 70 km with a peak at about 50 km. This layer has been attributed to the cross equatorial Hadley cell in which warmer air rises and travels poleward before cooling and sinking at higher latitudes. The deep downwelling air is rich in oxygen molecules, which recombine in the polar night to form O₃. O₃ can also be affected by the topography of Mars, through the effect of gravity waves. An increase in O₃ column over Hellas basin is attributed to a topographically induced transport of the polar night air, enriched in O₃ (Clancy *et al.* 2016).

These studies lead us to believe that there should be an anti-correlation of O₃ with water vapour (Lefèvre *et al.* 2004; Fast *et al.* 2006; Fedorova *et al.* 2006; Clancy *et al.* 2016; Willame *et al.* 2017). This anti-correlation was inferred for the first time from Mariner 9 observations (Barth *et al.* 1973). The photolysis of water vapour provides odd hydrogen radicals (HO_x) which are responsible for the catalytic loss of O₃ (Nair *et al.* 1994). Both polar caps and the regolith trap water vapour when the temperature is low. The northern polar cap is a greater source of water vapour compared to the southern polar cap. The atmosphere over other regions of Mars contains comparatively less water vapour. O₃ is mainly produced in the sunlit latitudes where oxygen atoms are abundant due to the dissociation of CO₂ and O₂ and combine with O₂ to produce O₃. During winter over polar-regions, the contribution to the O₃ column comes mainly from the transport of O₃ from sunlit

latitudes. During summer, polar caps release water vapour and destroy the O₃ column. However, it is believed that the degree of anti-correlation varies from region to region (Lefèvre *et al.* 2004). Latitudinal and seasonal variations in the temperature of the Martian atmosphere control the water vapour saturation level (hygropause) and thus the O₃ distribution. Thus, e.g., during the spring and fall when the temperatures are low, high O₃ is observed in the mid-to-high latitudes. As a result of the anti-correlation with water vapour, O₃ is at the maximum during winter and at the minimum during summer in general for all latitudes (Barth *et al.* 1973). It is our aim to study the seasonal and meridional variation of O₃ using observations from remote sensing and simulations from a model.

Dust aerosol is the most important component of Martian meteorology. Dust affects the incoming and outgoing solar radiation by its radiative properties. It heats up by absorbing in the visible region of the solar spectrum and cools down by radiating in the infrared region, thus modifying the temperature profile (Gierasch and Goody 1972; Zurek 1978). Daytime absorption of solar radiation lifts dust to a height 20–30 km above the surface (Guzewich *et al.* 2013). The Martian atmosphere is characterised by global dust storms, regional dust storms and local dust storms (Martin and Zurek 1993; Smith 2009; Sheel and Haider 2016). These dust events change the vertical profile of dust (Heavens *et al.* 2011). Dust optical depth varies seasonally and also with the global and regional storms (Sheel and Haider 2016). The annual variation of these storms is not fully known. As the optical depth increases, solar radiation cannot penetrate deeper. This reduces the availability of solar radiation at many layers of the atmosphere and thus affects the photochemistry of Mars. In this study, we will retrieve the optical depths of dust along with the O₃ column for various latitude regions and for various seasons.

O₃ in the atmosphere of Mars is closely linked to CO (produced by the photodissociation of CO₂ and is mainly destroyed by oxidation with OH radicals). CO can be used as a dynamical tracer in the Martian atmosphere because of its longer lifetime (González-Galindo *et al.* 2005), and as shown by Clancy *et al.* (2016) has been used to determine the increase of O₃ in the Hellas basin. In this study, we will, therefore, also examine the variability of the CO–O₃ correlation in the Martian atmosphere.

2. Objective

The observations of O₃ on Mars are important not only because of their importance in photochemistry, but also to test photochemistry-coupled general circulation models (GCMs) which simulate the spatio-temporal variation of O₃ and related species. Also, the variations in the UV environment of Mars are mainly due to O₃ and dust. O₃ strongly absorbs UV in the 255-nm region and hence a continuous monitoring and modelling of O₃ is essential to characterise the UV radiation reaching the surface of Mars. Dust not only affects the retrieval of accurate O₃ abundances, especially in regions with high O₃ (Clancy *et al.* 2016), but can also directly affect O₃ abundances due to its radiative impact on photochemistry. In this paper, we have retrieved two Martian years (MY27 and MY28) of total O₃ abundances (O₃ column) from SPICAM to study the seasonal variability of O₃ and its correlation with dust and CO. The objectives of this paper comprise (i) a study of the seasonal variation of total O₃ in different latitude regions from retrieved and modelled abundances; (ii) a study of the retrieval of Martian O₃ in the presence and absence of dust storms and (iii) a study of the correlation of O₃ with CO. The model used here is the Laboratoire de Météorologie Dynamique (LMD) photochemistry coupled GCM (LMD-GCM) described later in section 4.

3. O₃ retrieval

The observations from SPICAM UV spectrometer aboard the *Mars Express* are used to retrieve columnar O₃. These observations are available in analogue to digital units (ADU) from the radiance recorded by the spectrometer in the nadir mode (Bertaux *et al.* 2006; Perrier *et al.* 2006). The spectral resolution of the spectrometer is about 1 nm/pixel, and the resolving power is 120–300. SPICAM records the UV radiation in the dayside, backscattered by the atmospheric gases and dust in the Martian atmosphere. Observations are recorded in 110–320 nm range, allowing the measurement of O₃, which has a characteristic Hartley band absorption spectrum in the wavelength range of 210–300 nm, with a peak at 255 nm. This characteristic absorption gets imprinted in the backscattered solar radiation received by SPICAM. In this wavelength region, other species such as CO₂, N₂, O₂ and H₂O are

less absorptive in comparison with O₃, which make the O₃ column determination easier. For the current retrieval, we have used measurements of O₃ absorption cross sections from Malicet *et al.* (1995), which are consistent with the recent measurements of Gorshchev *et al.* (2014) and Serdyuchenko *et al.* (2014). However, Rayleigh scattering and absorption and scattering by dust have to be accounted for while retrieving O₃. In order to retrieve O₃ values from the SPICAM observations, theoretical spectra need to be derived and fitted. The radiative transfer model and the model atmosphere used to do so are described in section 3.2.

3.1 Raw data and retrieval method

The *Mars Express* which is in an elliptical polar orbit with a 7-h period, started observations in January 2004 at solar longitude (L_s) = 331°. We have retrieved observations from MY27 and MY28. This corresponds to approximately 4149 orbits from orbit number 61 (MY27, L_s = 7°; 19 March 2004) to orbit number 5055 (MY28, L_s = 359.74°; 9 December 2007). The numbers of orbits in individual years MY27 and MY28 were 2331 and 1818, respectively. The analysis resulted in around 51,721 high-S/N spectra (MY_27: 28,273 and MY_28: 23,448).

We have taken level 1A observation spectra from the Geosciences Node of ESA's Planetary Science Archive (<ftp://psa.esac.esa.int/pub/>). The Level 1A data are corrected for dark charge, electronic noise, erroneous data and damages by cosmic rays and saturations. Each nadir observation is typically 30-min long and records almost 1000 spectra, spectral radiance being recorded each second. SPICAM records spectra in five contiguous bins each second. Each bin is characterised by the number of charge-coupled device (CCD) lines grouped together (Bertaux *et al.* 2006). Spectra recorded in each bin are averaged to obtain a mean spectrum from which O₃ is extracted. We then averaged 50 consecutive spectra to improve the signal-to-noise ratio. This results in 30–40 individual spectra. Each recorded spectra (I) is divided by the flux at the top of the atmosphere (F), yielding the radiance factor I/F.

Perrier *et al.* (2006) have reported that the spectra suffer from the internal stray light by 8–10% of its total intensity. Accordingly, we describe here the internal stray light correction that we have applied to each spectrum. Stray light contributions to the data can be taken as 10% of the

recorded intensity (Bertaux *et al.* 2006). Therefore, we integrated each spectrum over the operating range of the SPICAM-UV spectrometer ($\lambda = 118\text{--}320$ nm):

$$I = \int I_\lambda d\lambda. \quad (1)$$

Bertaux *et al.* (2006) provided spectra of stray light for each band. Each of those spectra has been integrated over the wavelength range 118–320 nm. The stray light associated with any particular spectrum recorded by SPICAM at wavelength λ is given as

$$I_{\text{stray}}(\lambda) = 0.1 \times I \times \left(I_b(\lambda) / \int I_b(\lambda) d\lambda \right), \quad (2)$$

where $I_b(\lambda)$ is the intensity (in ADU) associated with the stray light spectrum. Therefore, the corrected spectra are given as

$$I_{\text{corrected}}(\lambda) = I_{\text{SPICAM}}(\lambda) - I_{\text{stray}}(\lambda). \quad (3)$$

The averaging over 50 consecutive spectra results in 30–40 individual spectra. This has been done in order to increase the signal-to-noise ratio. The final intensity is expressed in terms of kilo Rayleigh (kR) per nanometre, and consequently converted into the radiance factor (I/F), where πF is the solar flux received by SPICAM.

3.2 Forward model

To simulate a theoretical spectrum corresponding to each observational spectrum, we developed a forward model. We solve the following radiative transfer equation in the nadir direction assuming a plane parallel atmosphere:

$$\frac{dI_\nu(z)}{d\tau_\nu(z)} = S_\nu(z) - I_\nu(z), \quad (4)$$

where $I_\nu(z)$ is the spectral radiance at height z in units of power per unit area, per unit solid angle, per unit frequency interval in the neighbourhood of the frequency ν . $S_\nu(z)$ is the source function at height z . The optical depth of the atmosphere at height z is defined as

$$\tau_\nu(z) = \int_0^z k_\nu(z') \rho_i(z') dz', \quad (5)$$

where k_ν is the coefficient of extinction due to absorption and scattering by a species i whose density is ρ_i . The integral form of the above radiative transfer equation is written as (Evans 1998)

$$I_\nu(z) = \int_0^z S_\nu(z') k_\nu(z') \rho_i(z') \times \exp \left[- \int_{z'}^z k_\nu(t) \rho_i(t) dt \right] dz' + I_\nu(0) \exp(-\tau_\nu(z)), \quad (6)$$

where the variable $t = z - z'$. The first term on the right-hand side represents the scattering contribution of the diffuse radiations and the second term represents the contribution of the direct radiation in the direction of $I_\nu(z)$. Numerically, we perform the above integral using the discrete ordinate method and representing the angular part of the source functions S_ν and I_ν in terms of spherical harmonics expansion. The method is called the simple harmonic discrete ordinate method (SHDOM) (Pincus and Evans 2009). This method is computationally more efficient than standard methods. The source function includes contribution due to the scattering and absorption from diffused radiation.

3.2.1 Model atmosphere for the radiative transfer model

To retrieve the O_3 column from the SPICAM observed radiance, a model atmosphere is required for the geometry of each spectrum. This atmosphere is constructed with data from the Mars Climate Database (MCD) version 5.2 (Millour *et al.* 2014) and is divided into 12 equal layers up to 60 km. For the retrieval of O_3 , surface pressure, temperature and dust vertical profile are important inputs and appropriate calculations of dust extinction and Rayleigh scattering are required. The MCD is a result of the three-dimensional (3D) LMD GCM which is a well-tested dynamical model for Mars (Encrenaz *et al.* 2012; Clancy *et al.* 2013), hence constructing the model atmosphere from MCD is suitable for our purpose. Among the other inputs, we have provided dust-effective radius and O_3 vertical profiles to the radiative transfer model, suitable for the geometry of the corresponding spectra to be fitted.

3.2.2 Rayleigh scattering

The Martian atmosphere is composed of 95.7% CO_2 , 2.07% Ar and 2.03% N_2 (Franz *et al.* 2015), hence being a CO_2 -dominated atmosphere, it is 2.5 times more scattering than Earth's atmosphere (Forget and Pierrehumbert 1997). Thus, the calculation of CO_2 Rayleigh scattering cross section

becomes important. In earlier studies, theoretical calculation of Rayleigh scattering for CO₂ does not agree with measurements (Sneep and Ubachs 2005). We have calculated the cross section from the fitted equation $\sigma = \bar{\sigma} \times v^{4+\epsilon}$ with appropriate values of the parameters (Willame *et al.* 2017). As for the cross sections of N₂ and Ar, we have calculated from theory based on the refractive index with data from Thalman *et al.* (2014). The parameter values are better than 1% and the experiment was made for the wavelength range between 408 and 532 nm, which we have extrapolated to the UV region that is important for O₃ retrieval.

3.2.3 Dust optical depth and scattering

Dust scattering and absorption are calculated from the Mie theory considering spherical dust particles. The phase function for dust scattering is calculated from the Legendre series expansion, the coefficients of which are given as inputs to the SHDOM solver. The refractive index in the UV wavelength range of consideration has been taken from Wolff *et al.* (2010). In figure 1, the phase function calculated from the Mie theory for a particle of radius 0.1 μm and its comparison with the Henyey–Greenstein phase function is shown. In our calculation, we have used the full series of the phase function, a truncation being used only when the values of the coefficients are less than 0.00001. The scattering and extinction efficiencies (S_{ext} and Q_{ext}) are shown in figure 2 for wavelength 220 nm. These are calculated for the refractive index, $m = 1.486 - i \cdot 0.0155$. The real part is a slowly varying function of wavelength and this is

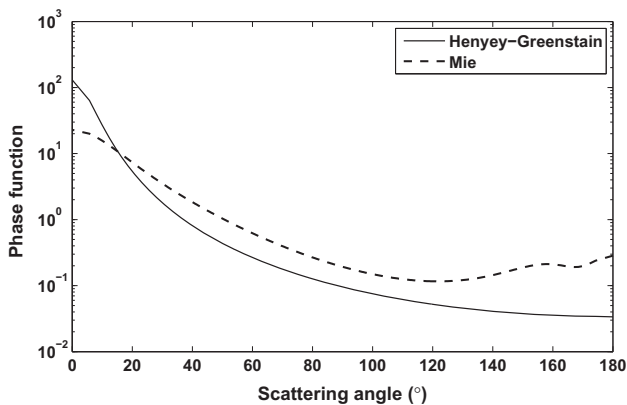


Figure 1. Phase function for particles of radius 0.1 μm calculated from Mie theory and compared with the Henyey–Greenstein phase function.

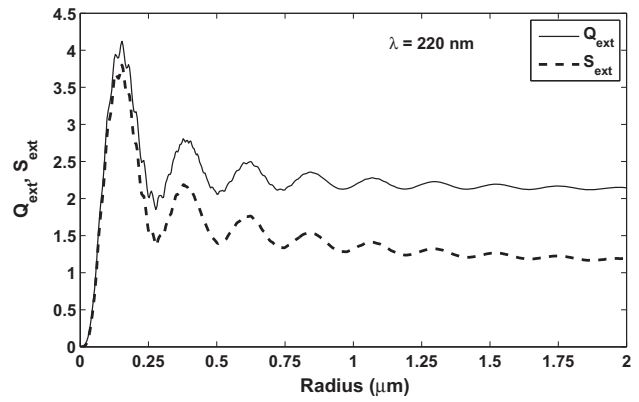


Figure 2. Scattering and extinction efficiencies (S_{ext} and Q_{ext}) for $m = 1.486 - i \cdot 0.0155$ and $\lambda = 220$ nm.

why we have taken a constant value 1.486 for the present retrieval (Wolff *et al.* 2009). As for the imaginary part, the proper behaviour with wavelength in the region 200–300 nm is not known; hence, we have used a value suggested by Wolff *et al.* (2009) for $\lambda = 668$ nm. The real part represents the dust scattering while the imaginary part represents absorption by dust. As can be seen from figure 2, the single scattering albedo defined as the ratio $S_{\text{ext}}/Q_{\text{ext}}$ is >0.9 for smaller particles. For particles with a higher radius, the single scattering albedo is reduced to values approaching 0.6.

3.3 Fitting method

Now that we have both theoretical and observational spectra, we compare them by using a best fit to determine columnar O₃. For this, we select nine points from the observation spectra in the range 210–300 nm. Corresponding to the observed radiance at each point, a theoretical radiance has been calculated using the radiative transfer model discussed in section 3.2, with a prior value of O₃ column, surface albedo and dust opacity. Mars surface albedo varies from place to place and therefore albedo at 210 and 300 nm are also retrieved simultaneously with O₃. However, the values of albedo are forced to be <1 . All parameters are kept positive throughout the retrieval. In order to determine total column O₃, the sum of the square of weighted differences (χ^2) has to be minimised. For this purpose, we have used the Levenberg–Marquardt minimisation method (LM method) which is a nonlinear chi-square minimisation method where multiple parameters can be fitted. The LM method which updates the parameter values adaptively, works better than the gradient

descent and Gauss–Newton methods. The convergence criterion for the method is that χ^2 defined as follows, is <1 :

$$\chi^2 = (y - \hat{y}(p))^T W (y - \hat{y}(p)), \quad (7)$$

where $y(\lambda_i)$ are the observations which are functions of wavelength λ_i and $\hat{y}(\lambda_i; p)$ is the fitting function for a vector of parameter p and W is the inverse of the error covariance matrix. We estimate the uncertainties on the O_3 column due to our retrieval method. For a known O_3 column, we generate synthetic spectra with the forward model. O_3 columns are then retrieved assuming that these synthetic spectra were observed by SPICAM. For high dust and moderate O_3 ($\sim 10 \mu\text{m atm}$), the errors range between 10% and 30%. The errors are in the same range for low dust and low O_3 ($\sim 0.5 \mu\text{m atm}$). For low dust and high O_3 ($\sim 20 \mu\text{m atm}$), errors in the retrieved O_3 values are within 10% (Perrier *et al.* 2006). An example of a fitted spectrum for the orbit number 902 (latitude = 31.92°N and $L_s = 95.31^\circ$) is shown in figure 3, where the radiance factor (as defined in section 3.1) is plotted against the wavelength. An O_3 columnar value of $0.23 \mu\text{m atm}$ is retrieved for this spectrum. The retrieved values of albedo are commensurate with a poor reflector of the UV radiation, and are lower compared to those retrieved by Perrier *et al.* (2006) in the polar region and also during some seasons in the equatorial region. We have observed that the retrieved O_3 values are not very sensitive to the values of albedo especially when there are clouds present.

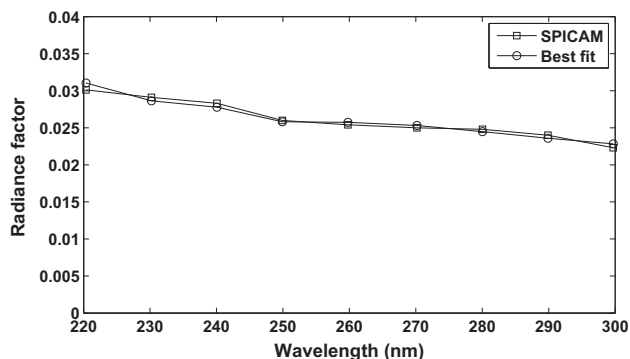


Figure 3. Fitted spectrum for orbit number 902 (latitude = 31.92°N , $L_s = 95.31^\circ$). An O_3 columnar value of $0.23 \mu\text{m atm}$ is retrieved for this spectrum.

4. General circulation model

In this work, we use the LMD-GCM, developed in collaboration with the Laboratoire de Météorologie Dynamique, the Open University, the University of Oxford and the Instituto de Astrofísica de Andalucía. This model uses physical parameterisations such as effects due to CO_2 and dust adsorption and emission, thermal conduction in the soil, CO_2 condensation and sublimation, sub-grid scale dynamics, vertical diffusion and convection (Forget *et al.* 1999). These are coupled to a spectral dynamical core and van-Leer advection scheme with mass conservation to transport tracers.

The dust distribution is prescribed horizontally using an interpolation of numerous sets of observations from orbiters and landers using the Kriging method (Montabone *et al.* 2015) and vertically distributed treating it at equal footing same as the tracers (Forget *et al.* 1999). We run the model on a grid of 3.75° latitude $\times 5.675^\circ$ longitude, with 32 vertical hybrid sigma levels extending to an altitude of ~ 122 km.

O_3 , water vapour, CO and other species are simulated by a photochemical module (Lefèvre *et al.* 2004) coupled with the GCM. The module consists of 36 chemical and 14 photochemical reactions. Chemical species are updated every 10 min of cycling over each atmospheric column, computing the tendencies for each chemical species at each level in the column. The chemical species are then transported by the van-Leer advection scheme. The advection scheme uses wind fields updated by the dynamical core to determine the O_3 concentration at each model grid point every 30 min. The chemical fields are exchanged between the GCM and photochemical model each time step for a fully interactive coupling between dynamics, physical parameterisations and chemical species. The model runs with appropriate boundary conditions. We run the model on the PRL 100 Tf high-performance computing (HPC) Linux cluster with 100 nodes at the Physical Research Laboratory Ahmedabad, India.

5. Results and discussions

Longitudinally averaged columnar values of O_3 retrieved from SPICAM for MY27 are shown as a function of latitude and season (L_s) in figure 4. For the current retrieval, the highest O_3 columnar values are within $25 \mu\text{m atm}$ over the northern polar

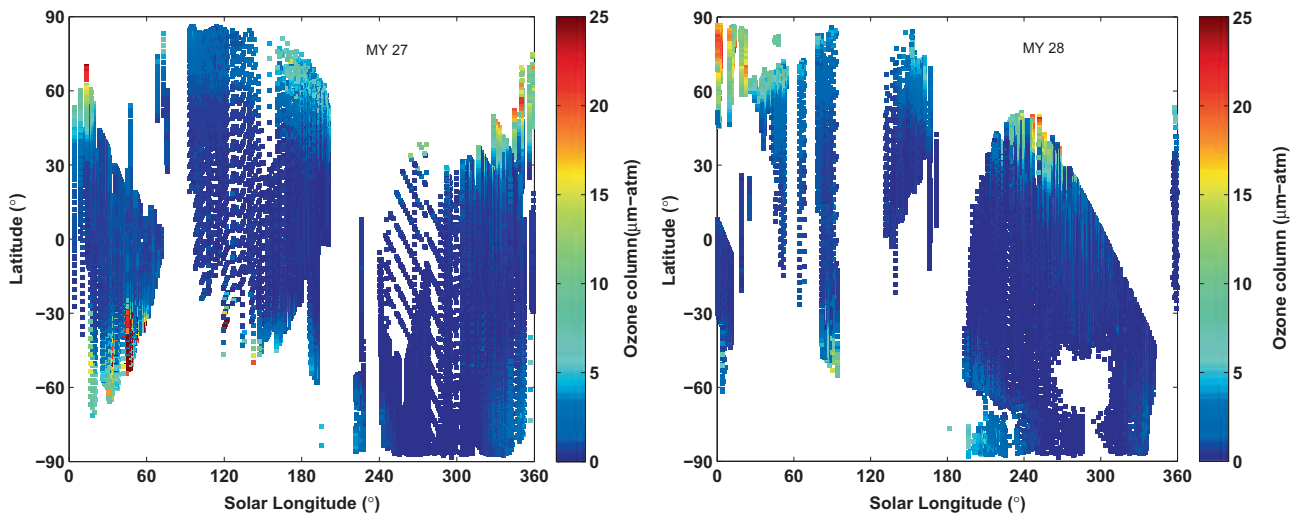


Figure 4. O₃ columnar values retrieved from SPICAM for MY27 as a function of latitude and season.

region during spring and winter. The O₃ retrieved for this work matches well with those retrieved by Willame *et al.* (2017) and Clancy *et al.* (2016), who have reported similar maximum values. However, Perrier *et al.* (2006) reported high columnar O₃ amounts of about 40 µm atm. The seasonal and regional features of the O₃ column can be seen in figure 4. In the Martian atmosphere, O₃ is anti-correlated with the water vapour (Krasnopolsky 1993; Nair *et al.* 1994; Lefèvre *et al.* 2004), since O₃ is destroyed by the odd H radicals produced from water vapour. This results in higher column values for O₃ during winter and almost negligible values during summer. During winter, water vapour condenses on the poles leading to a lesser loss of O₃ and thus higher column amounts. Over the high northern latitudes in spring, high O₃ values in the range 12–21 µm atm are retrieved. During the polar winter, night observations are not possible. During summer, around $L_s = 120^\circ$, O₃ values are of the order of 1 µm atm while they increase at the end of summer to 5 µm atm. During the southern summer, the columnar values of O₃ are <0.1 µm atm, which is a direct effect of hygro-pause elevation.

The O₃ data retrieved and discussed above are averaged over latitude intervals of 30° and L_s intervals of 2° and their comparison with simulations from the LMD-GCM is shown in figure 5 for MY27 and MY28 and for the six latitude regions: tropical (0° to ±30°), mid (±30° to ±60°) and high (±60° to ±90°) latitudes. Certain features are clearly visible in the GCM-simulated O₃ variability. Firstly, maximum O₃ columns (~20 µm atm) are seen at high latitudes and the polar region in the southern fall/winter due to low water vapour. This model

simulated peak occurs at $L_s = 100^\circ$ for polar latitudes and between $L_s = 50^\circ$ and 100° for high latitudes (30°S and 60°S). The O₃ columns retrieved from SPICAM observations also show high latitude O₃ maxima but for polar winter, data were not available. In contrast, very low O₃ due to the release of water vapour from the polar cap can be seen from $L_s = 240-360^\circ$ (southern summer) in the southern polar region. O₃ retrievals for the southern polar region for $L_s = 30-190^\circ$ are not available, as SPICAM cannot observe the winter poles due to the absence of solar irradiance. But from the comparison with GCM results, a reasonable match is seen with the retrieved O₃ data in different regions. In the equatorial regions, the seasonal variation of O₃ is not as pronounced as in the polar region.

The global map of dust optical depth retrieved from SPICAM is shown in figure 6 as a function of season for MY28. The figure indicates a background aerosol loading with optical depths <0.5 between $L_s = 0^\circ$ and 160° . Optical depths measured by a thermal emission imaging system (THEMIS) at 9.3 µm start to increase after $L_s = 220^\circ$ during MY28, reaching about 1.2 at $L_s = 280^\circ$ during the global dust storm period of that year (Sheel and Haider 2016). Similarly, the global dust storm can be seen from the current retrieval of dust optical depth (figure 6), when optical depths start increasing only after $L_s = 220^\circ$ and during $L_s = 240-330^\circ$, the values increase to >1.0. The retrieval is thus able to capture the global dust storm and thus compare qualitatively with the THEMIS observations (Smith 2009). After the major storm, it took about a few months for the

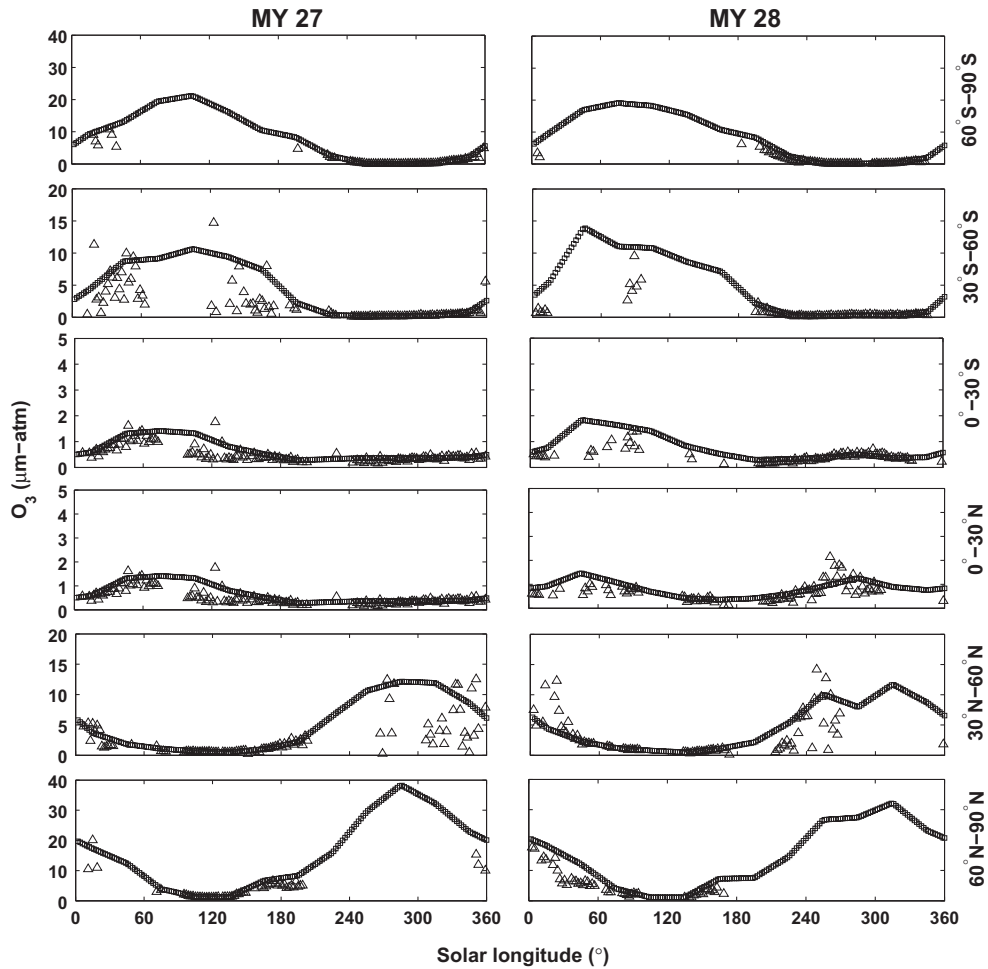


Figure 5. Zonally averaged O₃ retrieved from SPICAM and its comparison with simulations from the LMD-GCM for MY27 and MY28, averaged over the six latitude regions: tropical (0 to ±30), mid (±30 to ±60) and high (±60 to ±90).

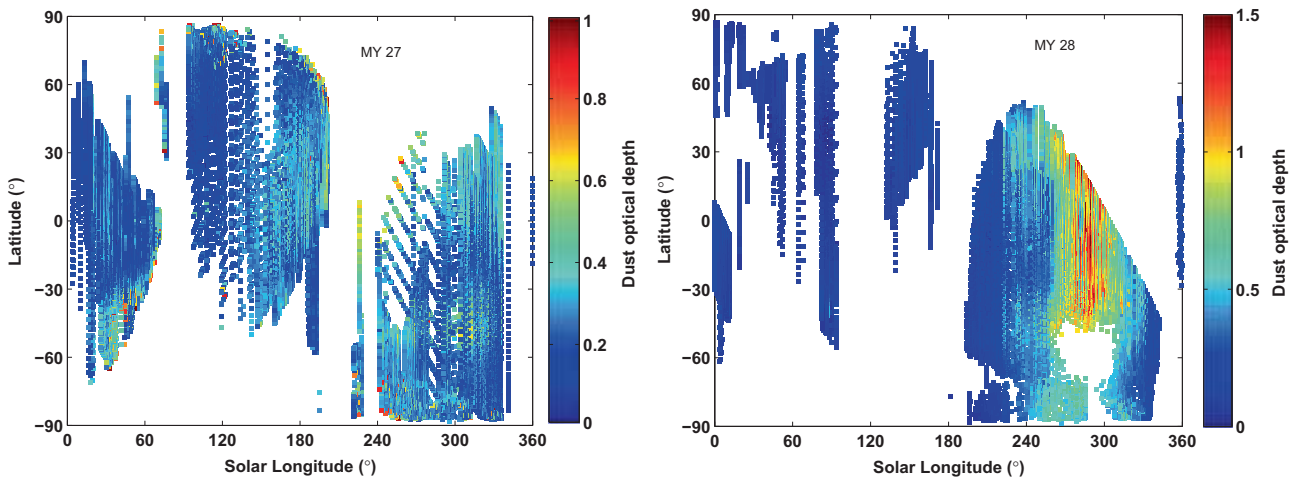


Figure 6. Dust optical depth retrieved from SPICAM at $\lambda = 300$ nm for MY28 as a function of latitude and season.

dust to settle down to the background dust loading. Many of these major dust storms begin in the subtropical region in the southern summer season after perihelion, when Mars receives a

higher amount of radiative energy from the Sun compared to other seasons. The increased solar insolation increases the temperature and wind speeds lift dust from the surface into the

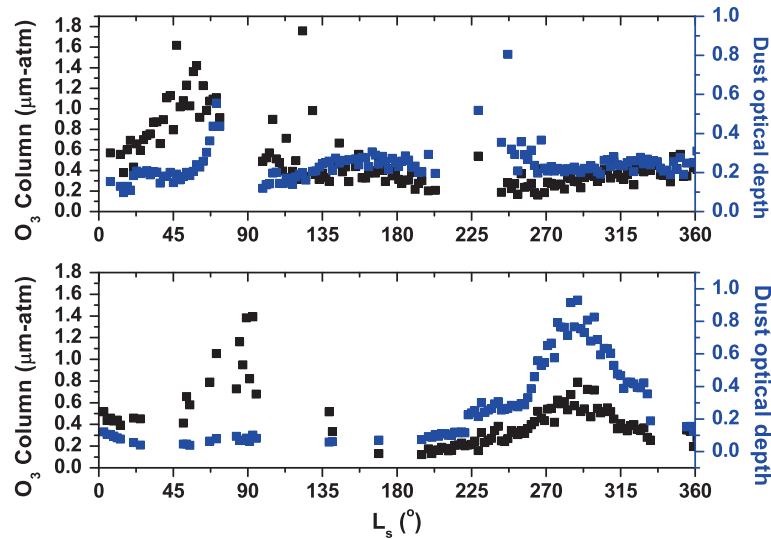


Figure 7. Correlation of O_3 and dust retrieved from observations by SPICAM, for the latitude region $0\text{--}30^\circ\text{S}$. The upper panel is for MY27 (no global dust storm) and the lower panel is for MY28 (global dust storm).

atmosphere of Mars (Lemmon *et al.* 2015). The retrieval of dust optical depths can be affected by the presence of clouds, as the retrieval methods yield artificially high dust optical depths, possibly due to the presence of water-ice clouds (which we do not retrieve). In general, clouds occur in the cold aphelion seasons. For example, dust optical depths are found higher in the southern polar region during $L_s = 0\text{--}50^\circ$ in MY27. Due to a lack of data in the southern polar region, those high values of dust optical depth affected by clouds could not be seen during MY28.

5.1 O_3 and dust

O_3 is very reactive, and apart from being sensitive to changes in water vapour in the atmosphere, it is also affected by the dust loading. Dust, which is known to affect various other parameters in the atmosphere, including the ionosphere (Sheel and Haider 2016), is highly variable. Many dust storms originate in the southern hemisphere and some of these grow into global dust storms in certain years, while regional or local dust storms occur almost every year (Sheel and Haider 2016). A climatology of 9 Martian years of dust loading has revealed that there is always a background aerosol loading of optical depth of about 0.1 between $L_s = 0^\circ$ and 160° , corresponding to a period when dust storms are absent, while the optical depth increases every year (e.g., by a factor of 5 at the southern summer tropics), to values in the range 0.3–0.5 corresponding to regional dust storms which typically occur at $L_s = 220^\circ$ (Sheel

and Haider 2016). For global dust storms, the peak optical depth can reach up to, e.g., 1.7 in the southern tropics during the MY25 global dust storm at $L_s = 210^\circ$. During the regional and global dust storms, detached dust layers have also been observed and modelled (cf. Sheel and Haider 2016), which could lift dust to altitudes up to 50 km, between $L_s = 150^\circ$ and 280° . Here, we consider two cases: (i) MY28 which witnessed a global dust storm and (ii) the preceding year MY27, in which a regional storm did occur, though it could not eventually turn into a global storm. In figure 7, we have shown the seasonal behaviour of O_3 and dust (total optical depth) for MY27 and MY28 for southern low latitudes ($0\text{--}30^\circ\text{S}$).

A comparison of the panels of figure 7 indicates an increase of O_3 with the increase of dust. As discussed above (figures 4 and 5), not much variation in O_3 is seen in the tropical latitudes. However, in figure 7, we see a striking feature of O_3 increasing with dust during the global dust storm year of MY28 from $L_s = 210^\circ$ to $L_s = 330^\circ$. The O_3 columnar values continue to increase after $L_s = 225^\circ$ in MY28 (figure 7), for dust optical depths more than 0.5. The water vapour column shown in figure 8 indicates that for the 2 years, the amount of water vapour available in the atmosphere is constant (Navarro *et al.* 2014); therefore, the increase in O_3 can be attributed to the increased dust loading and its radiative impact. This behaviour of the O_3 column during the global dust storm can be validated with the LMD-GCM simulation for MY28 with high dust (see figure 9). Other model simulations show that the photodissociation rates

are drastically reduced during dust storms due to the radiative impact of dust, thereby increasing O_3 abundances which may be typically increased by 10–50% (Lindner 1988).

5.2 CO– O_3 correlation

CO in the Martian atmosphere has been observed in the 1.6 and 2.35 μm region by ground-based and spacecraft spectroscopy by OMEGA and PFS on the *Mars Express*, CRISM on *Mars Reconnaissance Orbiter*, and the heterodyne instrument for the far infrared (HIFI) aboard the Herschel space observatory (cf. Hartogh *et al.* 2010). CO and O_3 are closely related via their common source (photolysis of CO_2) and their common sink (loss with HO_x radicals). The chemical lifetime of CO in the Martian atmosphere is in order of years while the O_3 lifetime is of few minutes, therefore the

CO– O_3 correlation cannot be expected based on their chemistry. However, it has been shown that the dynamical transport of O atoms from sunlit latitudes increases O_3 over the southern polar region (Lefèvre *et al.* 2008). CO being a non-condensable and stable gas is also used as a dynamical tracer. Therefore, in order to investigate the dynamical effect on the O_3 columnar amount, we evaluate the correlation of GCM simulated O_3 column (as SPICAM observations for polar nights are not available) with the GCM-simulated CO column for each season and over six latitude regions as shown in figure 10. Modelled CO concentrations qualitatively compare well with observations (Smith *et al.* 2009). As can be seen from the figure, CO does not show much variation with O_3 over the latitude region 60°S – 60°N compared to over the polar region, indicating insignificant dynamical contributions to the O_3 column. However, during winter over the polar region, the CO column increases with the O_3 column, the correlation being greater over the southern polar region compared to the northern region. This indicates a higher dynamical contribution towards the increase in the O_3 columnar value over the southern polar winter.

To derive the percentage contribution of the dynamics, we compare the O_3 vertical profile from the 3D and the one-dimensional (1D) version of the LMD-GCM, as the 3D version includes the large-scale Martian dynamics. Apart from this, all other physical processes such as CO_2 condensation, radiative interaction within the atmosphere, water cycle, dust condition and solar insolation are kept the same. Both the models were initiated at $L_s = 0$ and ran with the full photochemistry module of

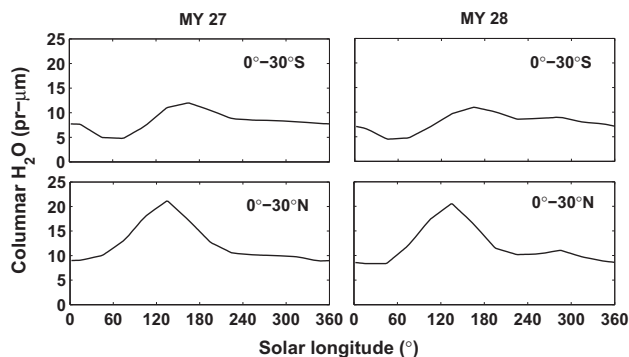


Figure 8. Seasonal variation of zonally averaged water vapour columnar abundance simulated by the LMD-GCM for MY27 and MY28 and for the latitude regions 30 – 0°S and 0 – 30°N .

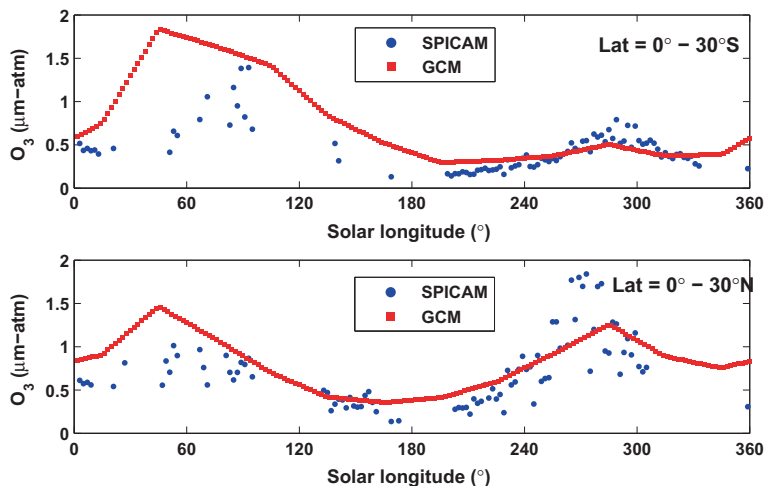


Figure 9. Comparison of the retrieved O_3 column from SPICAM observation with the GCM simulated values for MY28 and the latitude regions 30 – 0°S and 0 – 30°N .

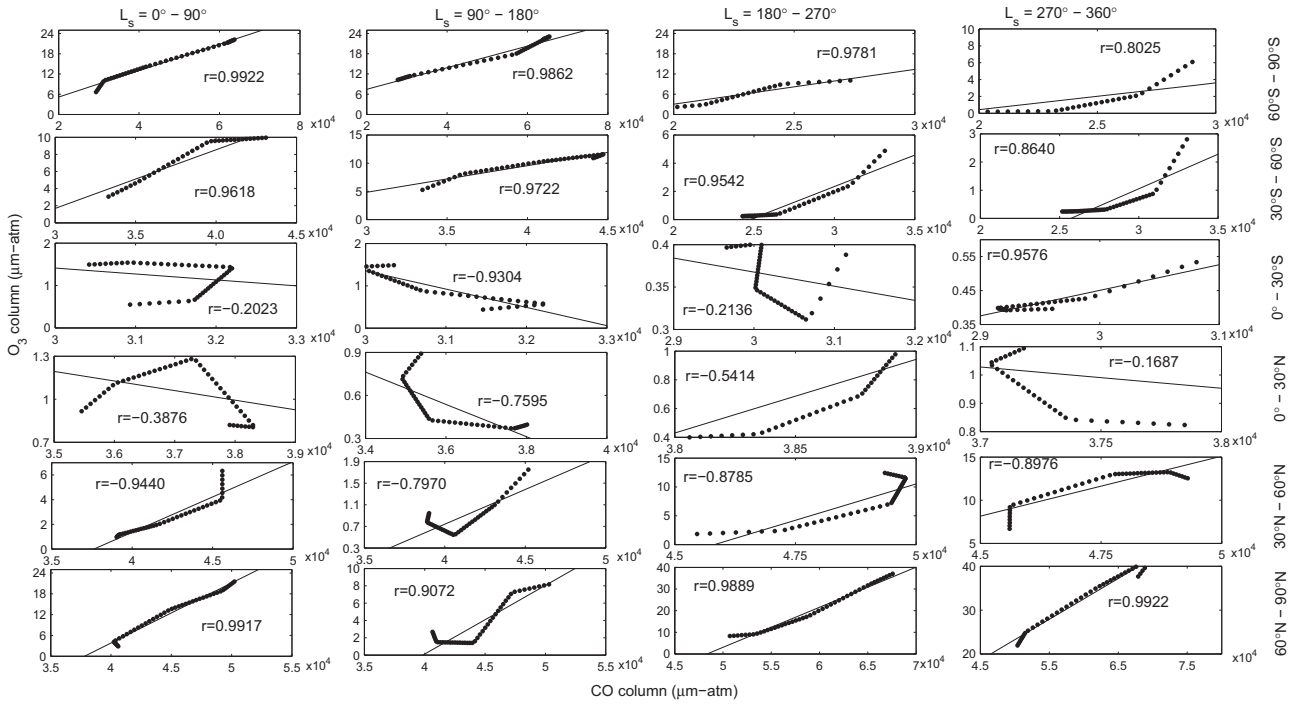


Figure 10. Correlation between O₃ and CO from the LMD-GCM, for the four seasons: spring, summer, autumn and winter, and the six latitude regions as in figure 5.

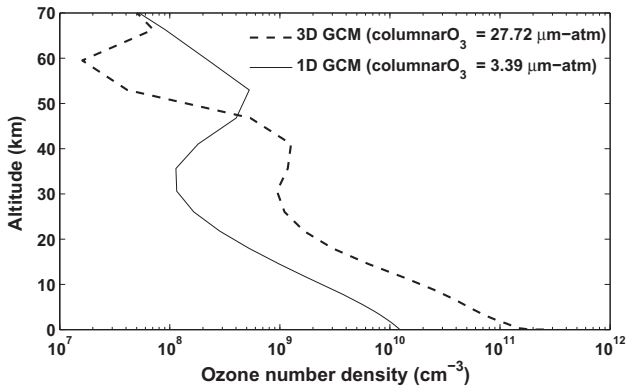


Figure 11. O₃ vertical profiles simulated for the 3D and 1D version of LMD-GCM. The 3D version includes dynamics whereas the 1D version does not include dynamics.

Lefèvre *et al.* (2004). For the 1D model, we input a reference surface pressure of 610 Pa and a temperature profile for the polar region. In figure 11, we show the number density of O₃ derived from both model calculations over 70°S during $L_s = 45^\circ$. In polar nights, the concentration of O atoms reduces drastically due to the absence of its major source (photolysis of CO₂ and O₂) as well as the absence of transport. This leads to a decrease in O₃ concentrations too, reflecting a decrease in the O₃ column of about $\sim 24 \mu\text{m atm}$ compared to the case when dynamics is present (figure 11). The shape of the O₃ vertical profile is decided by the H₂O profile

and the hygropause altitude. In the case of 3D simulation, hygropause occurs at a lower altitude, and hence, the second peak in O₃ occurs at a lower altitude compared to the 1D simulation. The exact cause of the increase in water vapour saturation level in 1D simulation could not be pointed out.

6. Summary and conclusions

We have retrieved O₃ columnar abundances and dust for MY27–MY28 from raw spectral radiances recorded by SPICAM aboard *Mars Express* and by using a forward radiative transfer model. We have also run the photochemistry coupled LMD-GCM on our HPC cluster, to simulate O₃, CO and water vapour.

The seasonal variability of total columnar O₃ was studied for 2 Martian years at tropical, mid and high latitudes. Seasonal variations in O₃ are not prominent in tropical latitudes, with O₃ column values observed in this region below 10 $\mu\text{m atm}$. For mid-latitudes, O₃ exhibits a peak in both northern and southern winters. The GCM simulations compare reasonably well with observations in these regions. A pronounced seasonal variability of O₃ is seen in the GCM results in high latitudes of both hemispheres, with a maximum in winter and minimum in summer.

Unfortunately, SPICAM observations are not available very frequently in the polar region. An O_3 –water vapour anti-correlation is seen at high latitudes, as reported by earlier studies.

We have also retrieved dust optical depths from SPICAM along with O_3 , and find that during dust storms, O_3 columnar amounts increase with the increase in dust optical depth due to the radiative impact of dust. The increase of O_3 can be as high as 50% of the O_3 without global dust storm. A similar result can be seen in O_3 simulated by the LMD-GCM for the dark dust storm of MY28.

To study the effect of dynamics on O_3 columnar abundance, the correlation between GCM-simulated CO and O_3 has been analysed. We find that the southern polar region exhibits a higher increase in the O_3 column due to the transport of oxygen atoms. During $L_s = 45^\circ$, the O_3 column over the southern polar region can increase from 3 to 27 $\mu\text{m atm}$ due to dynamical effect. On the other hand, in the northern polar region, the dynamical contribution is much less than the chemical contribution, while over the equator and tropical region, the dynamical effect is insignificant.

We have studied the seasonal variation of O_3 retrieved from SPICAM observations and simulated by the LMD-GCM, for different latitude regions. The seasonal variability is more prominent for higher latitudes compared to lower latitudes. The gross features are similar to that reported by previous studies. The retrieved O_3 values during dust storms are high, which we attribute to the radiative impact of dust. We also identify the effect of dynamics on the columnar O_3 , made possible by the 3D GCM simulations.

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