2	tropospheric ozone variability, trends and radiative effect
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Influence of nitrogen oxides and volatile organic compounds emission changes on

Abstract

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Ozone in the troposphere is a prominent pollutant whose production is sensitive to the emissions of nitrogen oxides (NOx) and volatile organic compounds (VOC). Here, we assess the variation of tropospheric ozone levels, trends, ozone photochemical regimes, and radiative effects using the ECHAM6-HAMMOZ chemistry-climate model for the period 1998-2019 and satellite measurements. The global mean simulated trend in Tropospheric Column Ozone (TRCO) for the study period (1998–2019) is 0.89 ppb decade⁻¹. During the overlapping period with OMI/MLS observations (2005–2019), the simulated global mean TRCO trends (1.58 ppb decade⁻¹) show fair agreement with OMI/MLS estimates (1.4 ppb decade⁻¹). The simulations for doubling emissions of NOx (DoubNOx), VOCs (DoubVOC), halving of emissions NOx (HalfNOx) and VOCs (HalfVOC) show nonlinear responses to ozone trends and tropospheric ozone photochemical regimes. The DoubNOx simulations show VOC-limited regimes over Indo-Gangetic-Plains, Eastern-China, Western-Europe, and the Eastern-US, while HalfNOx simulations show NOx-limited regimes over North-America and Asia. Emissions changes in NOx (DoubNOx/HalfNOx) influence the shift in tropospheric ozone photochemical regimes compared to VOCs (DoubVOC/HalfVOC).

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The estimated global mean TO3RE during 1998-2019 from the CTL simulations is 1.21 Wm⁻². The global mean TO3RE shows enhancement by 0.36 W m⁻² in DoubNOx simulations than CTL. While TO3RE shows a reduction in other simulations compared to CTL (DoubVOC: $-0.005~\mathrm{Wm^{-2}}$; HalfNOx: $-0.12~\mathrm{Wm^{-2}}$; and HalfVOC: $-0.03~\mathrm{Wm^{-2}}$). We show that emissions

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- changes in anthropogenic NOx cause higher changes in TO3RE than anthropogenic VOCs.

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- **Key words:** Tropospheric ozone, trends, ozone photochemical regimes; ozone radiative effect.
- 58 FNR; ECHAM6-HAMMOZ model simulations.

1. Introduction

Tropospheric ozone, a major air pollutant, has been a pressing issue in recent decades due to its detrimental effect on human health and ecosystem productivity and as a short-term climate forcer (Riese et al. 2012; Gulev et al., 2021; Wang et al., 2022). Considering these harmful impacts, the assessment of tropospheric ozone levels and trends are being conducted frequently (Gaudel et al., 2018; Mills et al., 2018; Tarasick et al., 2019). Ozone trends are being assessed from surface observations, in-situ and ground-based measurements, satellite retrievals, and model simulations (Cooper et al., 2014; Cohen et al., 2018; Young et al., 2018; Tarasick et al., 2019; Archibald et al., 2020). The latest IPCC AR6 reported an enhancement in free tropospheric ozone by 2-7% decade⁻¹ in the northern mid-latitudes, 2-12% per decade in the tropics, and <5% decade⁻¹ in southern mid-latitudes (Gulev et al., 2021; Szopa et al., 2021). The Tropospheric Ozone Assessment Reports (TOAR) have documented global increases of tropospheric column ozone (TRCO) in the 20th century (Cooper et al., 2014; Lefohn et al., 2017; Schultz et al., 2017; Fleming et al., 2018; Gaudel et al., 2018; Mills et al., 2018; Young et al., 2018; Tarasick et al., 2019). Increasing tropospheric trends are explained by enhanced anthropogenic emissions (Cooper et al., 2014; Zhang et al., 2016) and modulation by climate variability (Lin et al., 2014; Lu et al., 2018). Several studies have documented increasing trends in TRCO across various regions and different time periods.

For instance, enhancement in TRCO trends globally using measurements from multiple sources such as In-service Aircraft for a Global Observing System database (IAGOS) and ozonesondes and GEOS–Chem model simulation revealed an increasing trend of 2.7 ± 1.7 and 1.9 ± 1.7 ppb decade⁻¹ between 1995 and 2017 (Wang et al., 2022). Additionally, Fiore et al. (2022) also found increasing trends ranging from 0.6 to 2.5 ppb decade⁻¹ from 1950 to 2014 globally based on the available limited surface ozone records and the Community Earth System, version-2, the Whole Atmosphere Community Climate Model, version-6 (CESM2–WACCM6) model study. Furthermore, trends in TRCO are stronger in the Northern Hemisphere (NH) than Southern Hemisphere (SH) due to larger anthropogenic emissions (Monks et al., 2015). Ozone Monitoring Instrument (OMI) and Microwave Limb Sounder (MLS) observations from 2005 until 2010 show annual TRCO burden averaged over the NH exceeds the SH by 4% at low latitudes (0° – 25°), by 12% at mid-latitudes (25° – 50°), and by 18% at high latitudes (50° – 60°) (Cooper et al., 2014). CMIP6 models also show that the

tropospheric ozone burden increased by 44 % in 2005 – 2014 compared to 1850 (Griffiths et al., 2021). Recently, studies have reported a decrease in TRCO trends globally after the Corona–Virus Disease 2019 (COVID–19) outbreak e.g., Chang et al. (2022, 2023), and Steinbrecht et al. (2021). Also, Putero et al. (2023) show widespread ozone decreases at high-elevation sites due to the COVID–19 lockdown. However, our study period (1998–2019) excludes the COVID–19 associated emission changes.

Nitrogen oxides (NOx; NO+NO₂) and volatile organic compounds (VOC) are the major precursors that define ozone photochemical regimes (Duncan et al 2010). Information on ozone photochemical regimes is of utmost importance to know ozone (O₃) levels. However, the nonlinearity in the O₃ - NOx - VOC chemistry has always posed a challenge in identifying photochemical regimes. The regime is called NOx-limited if the ozone production is directly related to a change in NOx rather than from VOC perturbations. Whereas the region where ozone production is regulated by the ambient availability of VOCs is called VOC-limited (Sillman et al., 1990; Kleinman, 1994). The ratios such as ozone to oxidized nitrogen species (O₃/(NOy-NOx), where NOy is the total reactive nitrogen), formaldehyde to total reactive nitrogen (HCHO/NO₂), formaldehyde to nitrogen dioxide (HCHO/NO₂), hydrogen peroxide to nitric acid (H₂O₂/HNO₃) are adopted to diagnose the ozone photochemical regimes (e.g., Sillman, 1995; Martin et al., 2004; Duncan et al., 2010). Among these, the most widely used indicator to identify regimes is the Formaldehyde (HCHO) to Nitrogen dioxide (NO₂) Ratio (FNR) (Martin et al., 2004; Duncan et al., 2010). In our study, we adopt FNR to identify NOxlimited or VOC-limited regimes. On par with the current effort to mitigate ozone pollution, it is important to understand how the changes in emissions of NOx and VOC affect the ozone photochemical regimes and trends (Jin et al., 2017, 2020).

Ozone is the third strongest anthropogenic greenhouse gas, also called a short-term climate forcer, producing a global average radiative forcing of 0.47 [0.24 to 0.71 W m⁻²] [5% to 95% uncertainty range] (Forster et al., 2021). Recent studies showed ozone effective radiative forcing (ERF) of 0.51 [0.25 to 0.76] W m⁻² during 1750 – 2023 (Forster et al., 2024). The knowledge of ozone radiative forcing due to changes in anthropogenic emissions of NOx and VOC will help to assess climate change. Therefore, we also show the impacts of enhanced or reduced emissions of NOx and VOC on ozone radiative effect in addition to ozone trends

124 and photochemical regimes. To achieve this, we conducted sensitivity experiments by doubling 125 and halving global NOx and VOC emissions using the state-of-the-art chemistry-climate 126 model ECHAM6–HAMMOZ for the period 1998–2019. This approach of increase/decrease of 127 emissions is important to understand the nonlinear response of ozone to emission changes. 128 These experiments are helpful for designing emission implementation strategies (e.g., Zhang et al., 2021; Wang et al., 2023). The paper is outlined as follows: satellite data and the model 129 130 experimental setup are given in section 2, and results are given in section 3, which includes a 131 comparison of simulated tropospheric column ozone with satellite data and estimated ozone 132 trends. Discussions on ozone photochemical regimes and their trends are made in sections 4 to 133 6. Estimates of ozone radiative effects are given in section 7. Conclusions are made in section 134 8.

2. Satellite data and model experiments

136 2.1. OMI Satellite Data.

- We include OMI/MLS tropospheric column ozone (TRCO) for October 2004 December 2019 and OMI NO₂, HCHO data for latitude range 60° S – 60° N (Ziemke et al., 2006; De Smedt et al., 2021; Lamsal et al., 2021). OMI/MLS TRCO is determined by subtracting MLS
- stratospheric column ozone (SCO) from OMI total column ozone each day at each grid point.
- 141 Tropopause pressure used to determine the SCO invoked the World Meteorological
- Organization (WMO) 2 K km⁻¹ lapse-rate definition from the NCEP reanalysis. The MLS data
- used to obtain SCO were derived from the MLS v4.2 ozone profiles. We estimate 1σ precision
- for the OMI/MLS monthly mean gridded TRCO product to be about 1.3 DU. Adjustments for
- drift calibration and other issues (e.g., OMI row anomaly) affecting OMI/MLS TRCO are
- discussed by Ziemke et al. (2019) and Gaudel et al. (2024).
- 147 We used OMI monthly mean Level 3 (L3) data for NO₂ and HCHO
- 148 (https://doi.org/10.18758/h2v1uo6x) that were produced in the context of the ESA CCI+
- precursors for aerosols and ozone project (De Smedt et al., 2021; Anglou et al., 2024). The
- datasets consist of the monthly mean tropospheric column densities for NO₂ and HCHO (based
- on the QA4ECV NO₂ and HCHO dataset) as measured by OMI from October 2004 to March
- 152 2019, and include minimum spatial and temporal coverage thresholds (De Smedt et al., 2018).
- OMI has an overpass time of 13.30 local time and the retrieved column densities concern clear-
- sky or partially cloudy conditions.

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2.2 IASI–SOFRID

157 The Software for a Fast Retrieval of Infrared Atmospheric Sounding Interferometer (IASI) data 158 (SOFRID) retrieves global ozone profiles from IASI radiances (Barret et al., 2011, 2021). It is 159 based on the RTTOV (Radiative Transfer for TOVS) operational radiative transfer model 160 jointly developed by ECMWF, Meteo-France, UKMO and KNMI within the NWPSAF 161 (Saunders et al., 1999; Matricardi et al., 2004). The RTTOV regression coefficients are based 162 on line-by-line computations performed using the HITRAN2004 spectroscopic database 163 (Rothman et al., 2005), and the land surface emissivity is computed with the RTTOV UW-164 IRemis module (Borbas and Ruston, 2010). The IASI-SOFRID ozone for the study period 165 (2008 to 2019) is obtained from METOP-A (2008-2018) and METOP-B (2019). 166 We use the SOFRID version 3.5 data presented and validated in Barret et al. (2021), which 167 uses dynamical a priori profiles from an O₃ profile tropopause-based climatology according to tropopause height, month, and latitude (Sofieva et al., 2014). The use of such an a priori has 168 largely improved the retrievals, especially in the SH where the previous version was biased. 169 The retrievals are performed for clear-sky conditions (cloud cover fraction < 20%). IASI-170 171 SOFRID ozone retrievals provide independent pieces of information in the troposphere, the 172 UTLS (300 – 150 hPa), and the stratosphere (150 – 25 hPa) (Barret et al., 2021). SOFRID 173 TRCO absolute biases relative to ozonesondes are lower than 8 % with root mean square error 174 (RMSE) values lower than 18 % across the six 30° latitude bands (see Barret et al. (2021)). 175 Importantly, Barret et al. (2021) have shown that relative to ozonesondes, TRCO from IASI-SOFRID display no drifts (<2.1 % decade⁻¹) for latitudes lower than 60°N and in the SH for 176 latitudes larger than 30° (<3.7 % decade⁻¹). But significant drifts are observed in the SH tropics 177 (-5.2% decade⁻¹) and in the NH at high latitudes (12.8% decade⁻¹). 178

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2.3 IASI+GOME2

IASI+GOME2 is a multispectral approach to retrieve the vertical profile of ozone and its abundance in several partial columns. It is based on the synergy of IASI and GOME2 spectral measurements in the thermal infrared and ultraviolet spectral regions, respectively, which are jointly used to improve the sensitivity of the retrieval for the lowest tropospheric ozone (below

3 km above sea level, see Cuesta et al., 2013). Studies over Europe and East Asia have shown particularly good capabilities for capturing near-surface ozone variability compared to surface in situ ozone measurements (Cuesta et al., 2018, 2022; Okamoto et al., 2023). TRCOs from IASI–GOME2 also show good agreement with several datasets of in-situ measurements for a four-year period in the tropics, with almost negligible biases and high correlations (Gaudel et al., 2024). This ozone product provides global coverage for low cloud fraction conditions (below 30%) for 12 km diameter pixels spaced 25 km apart (at nadir). The IASI–GOME2 global dataset is publicly available through the French AERIS data center, with data from 2017 to the present and covers the 90° S – 90° N latitude band. For this study, we use the monthly TRCO data between the surface and the tropopause for 2017 – 2019 for different latitude bands.

2.4 TROPOMI

The TROPOspheric Monitoring Instrument (TROPOMI) is the sole payload on the Copernicus Sentinel–5 Precursor (Sentinel–5P or S5P) satellite, which provides measurements of multiple atmospheric trace species, including NO₂ and HCHO, at high spatial and temporal resolutions (Veefkind et al., 2012). TROPOMI has a daily global coverage with a spatial resolution of 5.5 \times 3.5 km² at nadir since a long-track pixel size reduction on 6 August 2019. We have used the ESA CCI+ Level-3 gridded 1° x 1° monthly tropospheric column of NO₂ (based on L2 v2.3.1, which applies a retrieval consistent with the most recent TROPOMI L2 version) and HCHO (https://doi.org/10.18758/2imqez32) (based on L2 v2.4.1, collection 3) data from May 2018 to December 2019 for our study (De Smedt et al., 2021; Glissenaar et al., 2024). This dataset was created using the same methods as the ESA CCI+ OMI Level-3 datasets.

2.5 The ECHAM6-HAMMOZ model experiments

The ECHAM6.3–HAM2.3–MOZ1.0 aerosol chemistry–climate model (Schultz et al., 2018) used in the present study comprises the general circulation model ECHAM6 (Stevens et al., 2013), the tropospheric chemistry module, MOZ (Stevenson et al., 2006) and the aerosol module, Hamburg Aerosol Model (HAM) (Vignati et al., 2004). The gas phase chemistry is represented by the Jülich Atmospheric Mechanism (JAM) v002b mechanism (Schultz et al., 2018). This scheme is an update and an extension of terpenes and aromatics oxidation based on the MOZART–4 model (Emmons et al., 2010) chemical scheme. Tropospheric heterogeneous chemistry relevant to ozone is also included (Stadtler et al., 2018). MOZ uses

216 the same chemical preprocessor as CAM-Chem (Lamarque et al., 2012) and WACCM 217 (Kinnison et al., 2007) to generate a FORTRAN code containing the chemical solver for a 218 specific chemical mechanism. Land surface processes are modelled with JSBACH (Reick et 219 al., 2013). Biogenic VOC emissions are modelled with the MEGAN algorithm (Guenther et 220 al., 2012) which has been coupled to JSBACH (Henrot et al., 2017). The lightning NOx 221 emissions are parameterized in the ECHAM6-HAMMOZ as described by Rast et al. (2014). 222 The lightning parameterization is the same in all the simulations. The model simulations were 223 performed for the period 1998 to 2019 using the Atmospheric Chemistry and Climate Model 224 Intercomparison Project (ACCMIP) (Lamarque et al., 2010; Van Vuuren et al., 2011) emission 225 inventory. ACCMIP emission inventory includes emissions from agriculture and waste 226 burning, forest and grassland fires, aircraft, domestic fuel use, energy generation including 227 fossil fuel extraction, industry, ship traffic, solvent use, transportation, and waste management. The model was run at a T63 spectral resolution corresponding to about $1.8^{\circ} \times 1.8^{\circ}$ in the 228 horizontal dimension and 47 vertical hybrid σ – p levels from the surface up to 0.001 hPa. The 229 230 details of model parameterizations and validation are described by (Fadnavis et al., 2019b, 231 2019a, 2021b, 2021a, 2022, 2023). We performed five experiments: (1) control (CTL) and four 232 emission sensitivity experiments: (2) doubling anthropogenic emission of NOx globally 233 (DoubNOx), (3) reducing anthropogenic emissions of NOx by 50 % globally (HalfNOx), (4) 234 doubling anthropogenic emissions of all VOCs globally (DoubVOC), (5) reducing 235 anthropogenic emissions of all VOCs by 50 % globally (HalfVOC). We performed each experiment from 1998 to 2019 after a spin-up of one year. We used the Representative 236 237 Concentration Pathway (RCP) 8.5 high emission scenario (Van Vuuren et al., 2011) in all 238 model simulations. In each experiment, the monthly varying AMIP–II sea surface temperature 239 and sea ice representative of the period 1998–2019 were specified as a lower boundary 240 condition. Anthropogenic VOC emissions included in the model are listed in the supplementary 241 table S1. TRCO is computed from the satellite data and model simulations by averaging O₃ amounts 242 from the surface up to the tropopause. The partial tropospheric column is converted into a 243 244 mixing ratio assuming a constant ozone mixing ratio in the troposphere. Tropopause considered is as described by the WMO thermal tropopause definition, the lowest level at which the 245 temperature lapse rate decreases to 2 K km⁻¹ or less (WMO, 1957). The estimated tropopause 246 247 in the satellite data will show differences since the tropopause is quite variable in space and

- 248 time; its location will depend on the employed reanalysis (e.g., Hoffmann and Spang, 2022).
- 249 The vertical resolution of the satellite and the ECHAM6-HAMMOZ also affect the estimated
- 250 tropopause. For comparison of the model with satellite datasets, e.g., IASI-SOFRID,
- 251 OMI/MLS, we use model and satellite data for the same period.

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2.6 Tropospheric ozone radiative effects

- 254 The tropospheric ozone radiative effect (TO3RE) is calculated as in Pope et al. (2024). While
- 255 the radiative effect calculated in ECHAM6–HAMMOZ also includes impacts of aerosols and
- dynamical effects, here we isolate TO3RE by using the Rap et al. (2015) tropospheric ozone
- 257 radiative kernel derived from the SOCRATES offline radiative transfer model (Edwards and
- Slingo, 1996), including stratospheric temperature adjustments. To calculate the TO3RE, the
- 259 monthly averaged ECHAM6-HAMMOZ simulated ozone field is multiplied by the offline
- 260 radiative kernel (at every grid box). It is then summed from surface to the tropopause. The
- simulated ozone data are mapped onto the spatial resolution of the radiative kernel and then
- interpolated vertically onto its pressure grid. The equation for each grid box is
- 263 $TO3RE = X \text{ trop}_i = \text{surf } RK_i \times O_{3i} \times dp_i / 100$ (1)
- 264 where TO3RE is the tropospheric ozone radiative effect (W m⁻²), RK is the radiative kernel
- 265 $(W m^{-2} ppbv^{-1} 100 hPa^{-1})$, O₃ is the simulated ozone grid box value (ppbv), dp is the pressure
- 266 difference between vertical levels (hPa), and 'i' is the grid box index between the surface
- pressure level and the tropopause pressure. The tropopause pressure is identified based on the
- 268 WMO lapse rate tropopause definition. Several past studies have used this approach of using
- 269 the SOCRATES offline radiative kernel with output from model simulations to derive the
- 270 TO3RE (Rap et al., 2015; Scott et al., 2018; Rowlinson et al., 2020; Pope et al., 2024).

271 **3. Results**

272 3.1 Comparison of the simulated seasonal cycle in TRCO, NO₂ and HCHO with

- 273 satellites retrievals
- 274 In this section, we compare the estimated TRCO from the model (CTL) simulation with
- 275 OMI/MLS (2005 2019), IASI–SOFRID (2008 2019), and IASI–GOME2 (2017 2019)
- satellite retrievals. We compared simulated TRCO for the same period as individual satellite
- 277 retrievals. The comparison of monthly mean TRCO is made for 20° latitude bins in Figure 1.

278 In the northern tropics ($0^{\circ} \text{ N} - 20^{\circ} \text{ N}$) (Fig. 1a), the OMI/MLS data exhibits an annual cycle 279 with a peak in April, whereas the model indicates a peak in January. Both datasets show a 280 minimum in August. The model underestimates TRCO by 1.8 to 3.9 ppb during March to October. In the 21° N -40° N and 41° N -60° N latitude bands (Fig. 1 b–c), the model shows 281 282 a one-month lead in the peak of the annual cycle compared to OMI/MLS. In the 21° N -40° N band, the model underestimates OMI/MLS TRCO by 2.8 – 6.1 ppb during the summer 283 284 months (May–August), while it overestimates TRCO by 4.1 - 8.3 ppb from October to March. 285 The 41° N -60° N latitude band exhibits an underestimation in the model by 1.1 - 6.3 ppb during June and July, while it overestimates (0.7 - 7.5 ppb) the rest of the year. In the Southern 286 287 Hemisphere (SH), OMI/MLS and the model show a similar pattern in the seasonal cycle. The 288 model shows a one-to-two-month lead in the annual cycle. However, the model shows an 289 underestimation of TRCO for all months. The model underestimates TRCO by 0.5 to 7.1 ppb in the 0-20 °S, by 5.1-15.3 ppb in 21 °S -40 °S, and by 9.2-13.8 ppb in the 41 °S -60 °S 290 291 latitude bands. The comparison of TRCO from IASI–SOFRID with the model shows features similar to those in the OMI/MLS. In the 0° N – 20° N latitude band, the model underestimates 292 293 the TRCO by about 3.8 to 7.7 ppb from April to October and in the 21° N -60° N latitude band by 1.9 – 11.3 ppb in summer (May–August). In the SH, the model shows better agreement with 294 295 IASI–SOFRID than OMI/MLS. During the SH winter (June–August), the model overestimates TRCO by 2.8 - 6.5 ppb in the latitude range of 0° S $- 40^{\circ}$ S. Conversely, it underestimates 296 TRCO by 2.7 - 8.2 ppb in the 41° S $- 60^{\circ}$ S throughout the year, which is less compared to 297 other satellite datasets. IASI-SOFRID is known to suffer from negative drifts in the SH (Barret 298 299 et al., 2021). 300 Interestingly, the model exhibits a fair agreement with IASI-GOME2 retrieved TRCO during 301 the summer months (May-August) in the Northern Hemisphere (NH). During the winter 302 months, the estimated TRCO shows a large overestimation of 8.3 - 11.7 ppb in the NH (0° N 303 -40° N), while it is underestimated by 8.3 - 11.7 ppb in the 41° N -60° N. In the SH, a fairly 304 good agreement is observed between the model and IASI-GOME2 TRCO, especially in the 0° $S-40^{\circ}$ S latitude band. The model overestimates the TRCO by 7.4-8.8 ppb in the 0° S -20° 305 S during SH winter and underestimates by 4.7 - 6.7 ppb in the 21° S -40° S belt during SH 306 307 summer (December–January–February). An overall underestimation of about 7 – 11.2 ppb in TRCO is noted in the 41° S -60° S throughout the year. Figure 1 shows that a peak in the 308 seasonal cycle in the model is earlier than the three satellite data between 40° N and 40° S. In 309

general, the model underestimates TRCO in summer in the NH and overestimates in winter relative to OMI/MLS, and IASI–SOFRID. In the SH, the model underestimates TRCO throughout the year compared to OMI/MLS, IASI–SOFRID, and IASI–GOME2, especially in the 41° N – 60° N band. Although the model–satellite comparison is done for the same period, the differences in sampling between the model and satellite measurements may cause the observed differences. It should be noted that the spatial resolution, coverage, and diurnal sampling time differ among the satellites which also contribute to the observed differences among them.

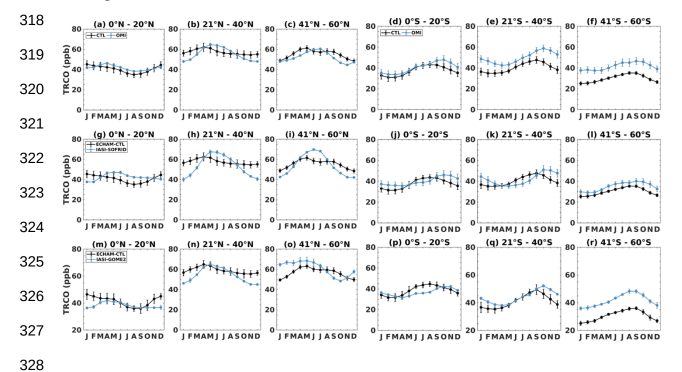


Figure 1. Time series of monthly mean TRCO (ppb) averaged for 20° wide latitude bins from (a–f) OMI/MLS (blue) and ECHAM6–HAMMOZ CTL simulations (black) for the time period October 2004 – December 2019. (g–l) same as (a–f) but for IASI–SOFRID (blue) and ECHAM6–HAMMOZ CTL simulations (black) for the period January 2008 – December 2019, and (m–r) same as (a–f) but for IASI–GOME2 (blue) and ECHAM6–HAMMOZ CTL simulations (black) for the time period January 2017 – December 2019. The vertical bars in all the figures represent 2σ standard deviation.

To evaluate our model simulations of NO_2 and HCHO, we compare the simulated tropospheric column NO_2 and HCHO with the ESA CCI+ monthly averaged TROPOMI and OMI data (Fig. 2). The simulated NO_2 reproduces the seasonal cycle but shows overestimation in the entire latitude band except 41° S -60° S in the SH. In the NH, the magnitude of overestimation in the simulated NO_2 increases with latitude. Simulated NO_2 is overestimated by 0.15 to 0.35

 $\times 10^{15}$ molecules cm⁻² in 0° N – 20° N, by 0.3 to 0.6 $\times 10^{15}$ molecules cm⁻² in 21° N – 40° N, and by 0.25 to 0.9 $\times 10^{15}$ molecules cm⁻² in 41° N – 60° N latitude bands compared to TROPOMI. Similarly, simulated NO₂ is overestimated compared to OMI by 0.16 to 0.35 $\times 10^{15}$ molecules cm⁻² in 0° N – 20° N, by 0.16 to 0.48 $\times 10^{15}$ molecules cm⁻² in 21° N – 40° N, and by 0.18 to 0.76 $\times 10^{15}$ molecules cm⁻² in 41° N – 60° N latitude belt (Fig. 2a–c and 2g–i). Although the model overestimates NO₂ in the SH, the magnitude of this overestimation is smaller compared to NH. Simulated NO₂ shows a fairly good agreement from 21° S to 60° S latitudes in the SH (Fig. 2d–f and 2j–l).

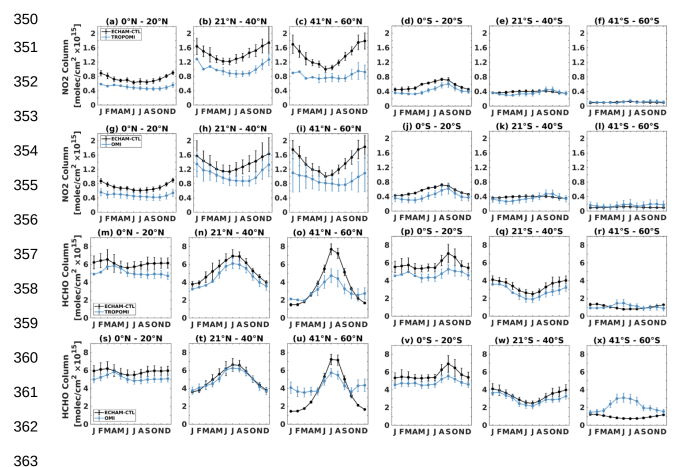


Figure 2. Time series of monthly mean tropospheric column NO_2 (molecules/cm²) averaged for 20° wide latitude bins from ECHAM6–HAMMOZ CTL simulations (black) for the time period same as (a–f) TROPOMI from May 2018 to December 2019, and (g–l) OMI from January 2005 to December 2019. (m–x) are the same as that of (a–l) but for HCHO. The vertical bars in the figures represent 2σ standard deviation.

While the simulated HCHO successfully reproduces the seasonal cycle in both hemispheres, it shows a large overestimation, particularly in the tropical region (Fig. 2m-x). The overestimation is most pronounced when compared to TROPOMI, especially in the tropics, and to a lesser extent with OMI. The model HCHO aligns reasonably well with both TROPOMI and OMI in the northern and southern mid-latitudes ($21^{\circ} \text{ N} - 40^{\circ} \text{ N}$ and $21^{\circ} \text{ S} - 40^{\circ} \text{ S}$) with a modest overestimation of $0.4 - 1.2 \times 10^{15}$ molecules cm⁻² and $0.3 - 0.5 \times 10^{15}$ molecules cm⁻² respectively in the NH and $0.4 - 1 \times 10^{15}$ molecules cm⁻² and $0.5 - 1.4 \times 10^{15}$ molecules cm⁻² respectively in the SH. However, in the 41° N -60° N band, the model overestimates HCHO compared to TROPOMI (OMI) by $0.6 - 2.9 (0.5-1.7) \times 10^{15}$ molecules cm⁻² during the NH from May to October and underestimates it by $0.08 - 1.1 (0.01 - 2.7) \times 10^{15}$ molecules cm⁻² during other months. On the contrary, the model underestimates HCHO in the 41° S -60° S during SH winter. It should be noted that TROPOMI/OMI monthly means are valid for clearsky situations, whereas the model simulations are all-day all-sky averages. In previous studies (Boersma et al. (2016) and references therein), it was shown that NO₂ is typically 15 – 20 % lower on clear-sky days than under cloudy situations due to higher photolysis rates, and faster chemical loss of NO2. Further, OMI and TROPOMI cannot sample for snowy scenes, and nighttime. There is significantly lower coverage on the NH during winter and vice versa for SH. These all can likely cause model and satellite differences. For HCHO the effect is smaller because HCHO is both produced and destroyed by OH (see Fig. 4 in Boersma et al. 2016). In addition, diurnal NO₂ observations from Geostationary Environment Monitoring Spectrometer (GEMS), Edwards et al. (2024) showed that the tropospheric column diurnal NO2 variations can be larger than 50 % of the column amount compared to once-a-day TROPOMI observations. Considering these differences, we proceed with the analysis of TRCO trends, ozone photochemical regimes, and ozone radiative effects.

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3.2. Impacts of emission changes on the spatial distribution of ozone

Figure 3 shows the spatial distribution of the simulated surface (Fig. 3a–e) and TRCO (Fig. 3f–j) concentration from ECHAM CTL simulations and the anomalies obtained from differences in DoubNOx - CTL, DoubVOC - CTL, HalfNOx - CTL, and HalfVOC - CTL simulations for the period 1998 - 2019. The CTL simulation shows high surface ozone levels (19 - 61.1 ppb) between 10° N -40° N (Fig. 3a). Doubling of NOx emission (DoubNOx) causes a global mean

enhancement of surface ozone anomalies by 4.1 [-3.8 to 13], [5th to 95th percentile] ppb. Surface ozone anomalies show an increase of 5-20 ppb across most of the globe, excluding highly urbanized regions like the Indo-Gangetic plains (IGP), Southeast China, Northeastern United States (US), and Europe. (Fig. 3b). Over these regions, a large reduction (8-20 ppb) in surface ozone anomalies is noticed, indicating ozone titration by NOx. While surface ozone anomalies from DoubVOC - CTL simulations show global mean enhancement by 0.9 [0.1 to 2.3] ppb, its magnitude is less than that of the anomalies from DoubNOx - CTL (Fig. 3c). The largest increase in surface ozone anomalies for DoubVOC is observed over IGP, Eastern China and the Eastern US (3-6 ppb). Interestingly, these are the same regions where a decrease in ozone anomalies is observed in the DoubNOx case. The decrease (increase) in ozone anomalies with an increase in NOx (VOC) emissions indicates that these regions could be NOx-saturated or VOC-limited. Reduction of NOx emissions (HalfNOx-CTL) simulations show a reduction in surface ozone anomalies (global mean by -2.5 [-7.2 to -0.7] ppb) except over North-Eastern China (Fig. 3d). Earlier, Souri et al. (2017) also reported that eastern Asia has witnessed a rise in surface ozone levels despite NOx control strategies, indicating the prevalence of VOClimited photochemistry over this region (details in section 4 to 6). However, the absence of such an increase over other VOC-limited regions point towards nonlinear ozone chemistry. While HalfVOC - CTL stimulation causes a reduction in surface ozone anomalies (global mean -0.4 [-1.4 to 0.05] ppb), an increase is observed in South America, some parts of the US, Australia, and the Indo-China peninsula (Fig. 3e). This increase could be due to a reduction in the radical destruction of ozone caused by aromatic hydrocarbons in low NOx conditions in these regions (Taraborrelli et al., 2021).

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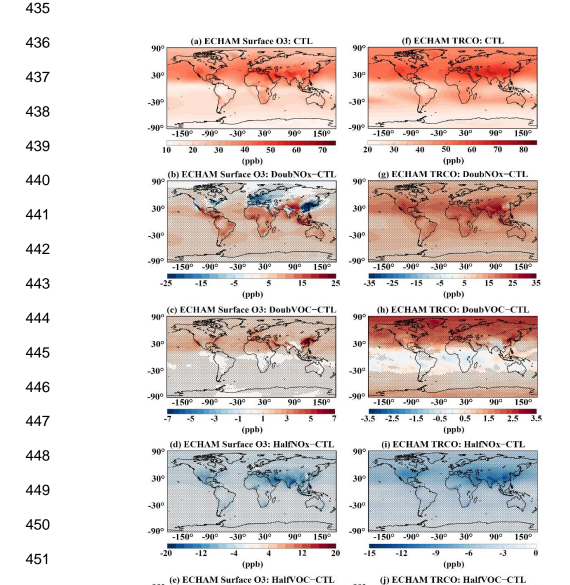
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Further, we show the impact of emission changes on the TRCO distribution (Fig. 3f–j). The estimated global mean TRCO from the CTL simulation from 1998 to 2019 is 39.4 [23.8 to 56.8] ppb (Fig. 3f). CTL simulations show higher amounts of TRCO (40.9 to 68.8 ppb) in the latitudinal band of 20° N to 40° N. These concentrations are pronounced over South and East Asia, spanning from the Mediterranean region to eastern China (Fig. 3f). TRCO anomalies from DoubNOx - CTL show enhancement by 11.7 [6.9 to 19.8] ppb (global mean) (Fig. 3g). Between 20° N – 40° N belt, the TRCO anomalies exceed by 6.1 – 29.3 ppb, particularly over South Asia. Interestingly, in highly urbanized areas such as the IGP, Southeast China, Northeast US, and Europe, there is only a marginal increase in TRCO anomalies (~5 ppb). This

suggests the existence of a distinct ozone photochemical regime in these regions. Further exploration of this aspect will be discussed in sections 4 to 6.



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-30°

-150°

(ppb)

Figure 3. Spatial distribution of surface ozone (ppb) for (a) from CTL simulations, anomalies from (b) DoubNOx - CTL, (c) DoubVOC - CTL, (d) HalfNOx - CTL, and (e) HalfVOC - CTL simulations. (f–j) are the same as that of (a–e) but for TRCO. The stippled regions in the figures indicate anomalies significant at 95% confidence based on the t-test.

-150° -90°

(ppb)

30°

The impact of the doubling of VOC emissions (anomalies from DoubVOC - CTL simulations) on TRCO is depicted in Figure 3h. An increase in global mean TRCO by 1 [-0.2 to 2.4] ppb is observed in this emission scenario. It should be noted that TRCO anomalies from DoubVOC -CTL are ten times less than that from DoubNOx - CTL (Fig. 3g and 3h). Large values of TRCO anomalies (1.5-2) are observed in the high latitudes (north of 60° N) and South and East Asia, with the largest values of more than 2.5 ppb over East China (e.g., Beijing). Interestingly, slight decreases in TRCO are seen in the tropical regions. This is consistent with the recent finding that aromatics, especially benzene, can lead to efficient ozone destruction in tropical UTLS (Rosanka et al., 2021). The TRCO anomalies in response to the reduction of NOx emission by 50% (HalfNOx - CTL) show negative TRCO anomalies all over the globe (Fig. 3i). The global mean TRCO anomalies are reduced by -3.7 [-7.9 to -1.1] ppb. Large decreases in TRCO anomalies are seen over Arabia, South and East Asian regions (2.6 – 12.8 ppb). The TRCO anomalies from HalfVOC - CTL show an overall decrease in TRCO by -0.27 [-0.97 to -0.4] ppb (Fig. 3j). Further, a small enhancement is noted in the TRCO anomalies (by 0.5 - 1 ppb) in the southern tropics and south polar region, while a decrease of -2.3 to 0.3 ppb is observed in the NH. (Fig. 3j). Figure 3 clearly portrays that the TRCO response to NOx emission change is larger than that of VOCs. There is a spatially distinct distribution in TRCO associated with the region-specific ozone photochemical regimes (more discussion on the ozone photochemical regimes will be detailed in sections 4-6).

3.3. Spatial distribution of trends in ozone

We estimate trends in TRCO from ECHAM CTL simulations (1998–2019) and OMI/MLS satellite retrievals (2005–2019). The simulated trends are compared with satellite retrieves for the period 2005–2019. Since IASI–GOME2 has a short observation period (2017–2019) and IASI–SOFRID has negative drift in the SH, only TRCO from OMI/MLS is considered for trend estimation (Fig. 4). The spatial pattern of trends from OMI/MLS shows fair agreement with model simulations (Fig 4a-b). Quantitatively, the global mean TRCO trend from OMI/MLS is slightly lower than the model (OMI/MLS:1.43 [-0.5 to 3.2] ppb decade⁻¹; ECHAM-CTL: 1.58 [0.3 to 3.3] ppb decade⁻¹). Both datasets reveal high trends, ranging from 3–4 ppb decade⁻¹, across regions such as South Asia, East Asia, and the West Pacific. OMI/MLS show negative trends over parts of Africa, South America, Australia, and the southeastern Pacific (Fig. 4b), which is not simulated in ECHAM6–HAMMOZ. Although there is fair agreement in spatial

patterns of TRCO trends between OMI/MLS and the model, the minor differences may be due to the model's tendency to underestimate ozone levels and differences in the seasonal cycle. (See Fig. 1).

TRCO trends analyzed from the Total Ozone Mapping Spectrometer (TOMS) indicate a consistent absence of trend over the tropical Pacific Ocean, with notable positive trends (4 – 5% decade⁻¹) seen in the mid-latitude Pacific regions of both hemispheres (Ziemke et al., 2005). This pattern is consistent across the ECHAM6–HAMMOZ and OMI/MLS data, although their magnitude differs (Fig. 4 a–b). TOMS data also showed trends of ~2 – 5% decade⁻¹ across broad regions of the tropical South Atlantic, India, Southeast Asia, Indonesia, and the tropical/subtropical regions of China during 1979 – 2003 (Ziemke et al., 2005; Beig and Singh, 2007) which are also simulated in the model. Further, a large positive trend of ~2.5 ppb decade⁻¹ observed near 50° S in OMI/MLS is not simulated by the model (Fig. 4a–b). The CESM2–WACCAM6 simulation from 1950 to 2014 also shows the largest trend estimate of 0.8 Tg decade⁻¹ over 20° N – 30° N (Fiore et al., 2022). Large TRCO trends over 20° N – 30° N are also seen in OMI/MLS and the model (Fig.4). Wang et al. (2021) reported TRCO trends varying between 2.55 to 5.53 ppb decade⁻¹ during 1995–2017 over South and East Asia using IAGOS, ozonesonde observations, and Goddard Earth Observing System–chemistry model (GEOS–Chem). Our model also shows similar increasing trends.

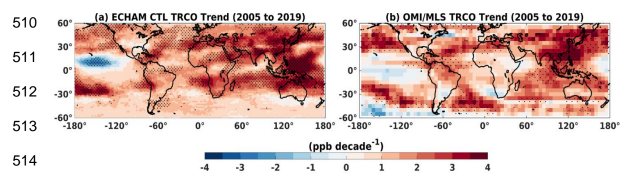


Figure 4. Trend of TRCO (ppb decade⁻¹) from (a) ECHAM CTL, and (b) OMI/MLS satellite for the period January 2005 to December 2020. Stippled regions in the figures indicate trends significant at 95% confidence based on the t-test.

Figure 5 shows the spatial distribution of estimated trends in surface ozone and TRCO from CTL simulation for the period 1998 – 2019. Changes (Doubling/halving) in the emission

of NOx and VOCs will change ozone trends. Hence, we also analyzed anomalies in ozone from DoubNOx - CTL, DoubVOC - CTL, HalfNOx - CTL, and HalfVOC - CTL. The surface ozone trend in the CTL simulation shows spatial variation with a pronounced increasing trend over South Asia and the Middle East (3 – 4 ppb decade⁻¹) (Fig. 5a). Similar pronounced increase is also seen in the TRCO trend (Fig. 5b). The estimated global mean TRCO trend from CTL is 0.89 [-0.07 to 2.1] ppb decade⁻¹. However, the negative trends in surface ozone over Mexico, certain parts of the US, and East China are barely discernible in the TRCO data. This discrepancy may stem from the interplay of mixing and transport processes, stratospheric intrusions, which are crucial when assessing ozone levels across the tropospheric column. The stratospheric ozone intrusions lead enhancement in the tropospheric ozone (Prather and Zhu, 2024).

Figure 5 c–d shows the trend in surface ozone and TRCO estimated from anomalies obtained from DoubNOx - CTL simulations. A striking feature is the large negative trend over India and China at the surface (-4.8 to -8 ppb decade⁻¹) and TRCO (-2 to -4 ppb decade⁻¹). Whereas Europe, the US, some parts of Africa and South America show positive trends at the surface (1.8 to 8 ppb decade⁻¹) and TRCO (2 to 4 8 ppb decade⁻¹). The global mean TRCO trend are 1.2 [-0.1 to 2.7] ppb decade⁻¹.

When global emissions of VOCs are doubled, the trend in ozone estimated from anomalies (DoubVOC - CTL) shows a decrease (by -0.8 to -1.9 ppb decade⁻¹) in surface ozone over Europe, Africa and some parts of the US, while strong positive trend (1.6 to 2 ppb decade⁻¹) are seen over India and China (Fig. 5e). TRCO trends show an enhancement over South Asia, Southwest Asia, China, parts of the Indian Ocean, and the western Pacific (0.8 to 1.6 ppb decade⁻¹) (Fig. 5f). A global mean TRCO trend for DoubVOC-CTL simulation is 0.5 [-0.03 to 1.04] ppb decade⁻¹. The estimated enhancement in global mean TRCO trend for DoubVOC is less than DoubNOx simulations. Figures 5 c and e also give indications of the existence of distinct ozone photochemical regimes globally. The increasing (decreasing) trend in surface ozone with an increase in VOC (NOx) over India and China indicates that these regions are in a VOC-limited regime, and vice-versa over the US and Europe indicates that these regions are in a NOx-limited regime (more discussions on sections 4 to 6).

Figure 5g—h shows the trend in surface and TRCO ozone estimated from anomalies from HalfNOx - CTL. The surface ozone trend shows a large negative trend over Europe and South Asia, while a positive trend over the US, China, and Australia (Fig. 5g). Trends in TRCO also show a large negative trend over South Asia (Fig. 5h). Although anthropogenic NOx emissions are halved compared to CTL, ozone trends are positive over large region globally. Our investigations reveal that the trend in VOC anomalies from the HalfNOx-CTL simulations is positive over the US, India, and Europe, while negative over China. Similarly, the trend in NOx anomalies from the HalfNOx-CTL simulations is positive over the US and Europe while negative over India and China (Fig. S1a-b). It should be noted that over the US and Europe the positive NOx trend estimated from anomalies (HalfNOx-CTL) suggests that NOx levels in HalfNOx are declining more slowly compared to CTL, leading to relatively higher NOx concentrations over time. Meanwhile, over India and China, the NOx level is weak in HalfNOx compared to CTL, leading to a negative trend estimated from NOx anomalies (HalfNOx-CTL).

It is known that ozone response is non-linear to NOx and VOC emission changes and depends on the local photochemical regime. In the US, estimated trends from anomalies are positive in VOC and NOx (HALFNOX-CTL). This indicates that relatively more ozone precursors in the HalfNOx contribute to the observed positive ozone trend. Over Europe, the strong positive trend estimated from NOx anomalies (compared to VOCs) enhances NOx titration effects, contributing to the observed negative ozone trend. China, being a VOC–limited regime, experiences reduced NOx titration under lower NOx conditions, resulting in positive ozone trends. Over India, the positive trend in VOC anomalies with a negative trend in NOx anomalies may enhance the radical destruction of ozone caused by aromatic hydrocarbons in low NOx conditions (Taraborrelli et al., 2021), resulting in a negative ozone trend. For HalfNOx – CTL, the global mean trend is positive 0.47 [-0.76 to 1.3] ppb decade⁻¹.

The trend in surface and TRCO ozone estimated from anomalies from HalfVOC – CTL is shown in Figure 5i-j. A large negative trend in surface ozone is noted over IGP and China, while an insignificant positive trend is noted over the US and Europe (Fig. 5i). The TRCO trend are positive over large regions in the world with pronounced high over the mid- and high latitudes although emissions of all anthropogenic VOCs are halved (Fig. 5j). The global mean trend for HalfVOC - CTL is 0.37 [-0.35 to 1.02] ppb decade -1. Our analysis shows that the

trend in NO_X anomalies and VOC anomalies from HalfVOC - CTL is decreasing over both India and China (Fig S1c-d). The negative trend in precursors might have resulted in a negative trend in ozone over these regions. The absence of strong trends in TRCO (Fig. 5 b, d, f, h, j) similar to that at the surface (Fig. 5 a, c, e, g, i) in all the simulations indicates the potential contribution of transport in the troposphere and stratospheric intrusions in TRCO. Ozone injection through stratosphere—troposphere exchange is important source of tropospheric ozone (Prather and Zhu, 2024).



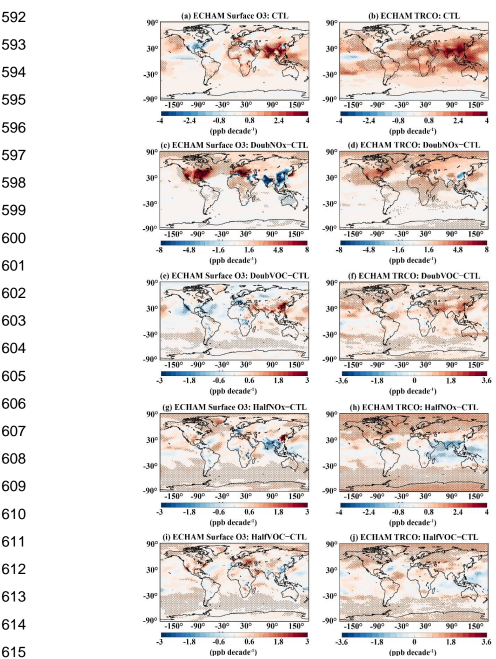


Figure 5. Trend in surface ozone (ppb decade⁻¹) for the period 1998-2019, calculated from (a) CTL, ozone anomalies in (c) DoubNOx - CTL, (e) DoubVOC - CTL, (g) HalfNOx - CTL, and (i) HalfVOC - CTL simulations. Panels (b, d, f, h, and i) represents the corresponding trend in TRCO. The stippled regions in the figures indicate significance at 95% confidence based on the t-test.

3.4. Trends in emission and tropospheric column of NO2 and HCHO

We show mean emissions of NOx (NO+NO₂) and HCHO over urban/semi-urban regions; US, Brazil, Europe, Africa, India, China, and Australia in Figure 6. High emissions of VOCs and NOx in India and China are evident in Figure 6. Furthermore, VOCs emissions are noted to be higher than NOx over all the regions. They are higher by a factor of 3.3 in the US, 11.3 in Brazil, 4.8 in Europe, 10.5 in Africa, 10.8 in India, 6.1 in China, and 6.7 in Australia.

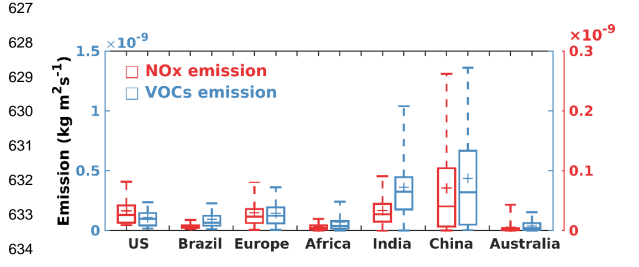


Figure 6. Box and whisker plot illustrating the NOx (NO+NO₂) and VOCs emission over the regions US (85°W - 110°W, 35°N - 44°N), Brazil (34°W - 49°W, 24°S - 3°S), European Union (9°W - 45°E, 35°N - 55°N), Central Africa (14°W - 45°E, 0° - 14°N), India (75°E - 90°E, 8°N - 30°N), China (110°E - 125°E, 30°N - 42°N), and South Australia (134°E - 154°E, 38°S - 28°S) from the ECHAM model. The box represents the 25 and 75 percentiles, and the whisker represents the 5 and 95 percentiles. The plus marker represents the mean, and the horizontal bar represents the 1 and 99 percentiles.

The trends in ozone are partly modulated by the change in the emission of its precursors and partly by meteorology (e.g., Verstraeten et al., 2015). We show trends in emissions and tropospheric column amounts of ozone precursors NO₂ and HCHO from ECHAM CTL and OMI satellite retrievals in Figure 7. NO₂ and HCHO are considered here because column densities of these will be used to identify the ozone photochemical regimes discussed later in

Sections 4–6. Emissions and tropospheric columns of HCHO and NO₂ from ECHAM–CTL show large positive trends over the South and East Asian regions (Fig. 7a–d). These regions show large positive ozone trends in both model and OMI satellite data (see Fig. 4 and 5). Over Europe and the US, the emission trend in both HCHO and NO₂ from the model is negative (Fig. 7a, c). Though a similar negative trend in tropospheric column NO₂ is seen over these regions, a marginal positive trend is noted for HCHO (Fig. 7b, d). The positive trend in column HCHO could be due to secondary production pathways from biogenic emissions or methane oxidation and transport (e.g., Anderson et al., 2017; Alvarado et al., 2020). The positive trend in ozone (Fig. 4a–b and 5a, f) along with a negative trend in NO₂ and HCHO (Fig. 7a–d) over Europe indicates that ozone production over this region has been initially controlled by VOCs (i.e., VOC-limited regime; detailed discussed in section 4). However, a large decreasing trend in NO₂ compared to that of HCHO over this region might have decreased the NOx titration effect, resulting in an increase in ozone. On the contrary, a negative trend in surface ozone (Fig. 5a) along with negative trends in NO₂ and HCHO are seen over the US (Fig. 7a-b). The decrease in both NO₂ and HCHO would have resulted in a decreasing trend in surface ozone over this region. This also indicates that the US might have been in a NOx-sensitive regime before and the large negative trend in NO₂ might have resulted in the decreasing trend in ozone (discussed further in section 4-6).

Further we compared the simulated trends in column HCHO and NO₂ with the OMI retrievals for the period 2005–2019 (Fig. 7e–h). OMI shows a positive trend in tropospheric column HCHO over South Asia $(1-1.5\times10^{15}\ \text{molecules}\ \text{cm}^{-2}\ \text{decade}^{-1})$, parts of western China $(0.75-1.25\times10^{15}\ \text{molecules}\ \text{cm}^{-2}\ \text{decade}^{-1})$, the Iranian Plateau $(0.5-1\times10^{15}\ \text{molecules}\ \text{cm}^{-2}\ \text{decade}^{-1})$, horth America $(0.5-1.5\times10^{15}\ \text{molecules}\ \text{cm}^{-2}\ \text{decade}^{-1})$, North America $(0.5-1.5\times10^{15}\ \text{molecules}\ \text{cm}^{-2}\ \text{decade}^{-1})$, and central Africa $(1-1.5\times10^{15}\ \text{molecules}\ \text{cm}^{-2}\ \text{decade}^{-1})$. The model simulated trends show reasonable agreement with OMI, except for western areas in central Africa, north Africa, southwest and southeast China, and some parts of Australia. Over these regions OMI indicates a negative trend, while the model suggests a marginal positive trend. OMI and ECHAM CTL show a good agreement in the tropospheric column NO₂ trend. Both datasets show negative trends over the eastern US and Europe, and positive trends over the Middle East, and South Asia. However, differences are seen in eastern China and central Africa, where OMI indicates a negative trend,

while the model shows a strong positive trend. The differences between simulated and OMI HCHO and NO₂ column trends may be due to sampling time and differences in seasonal cycle. Figures 4, 5, and 7 clearly indicate the impact of ozone precursors on the spatial distribution of ozone trends. This warrants a detailed discussion on the spatial distribution of ozone precursors and their impact on ozone production-sensitive regimes, which will be presented in the next section.

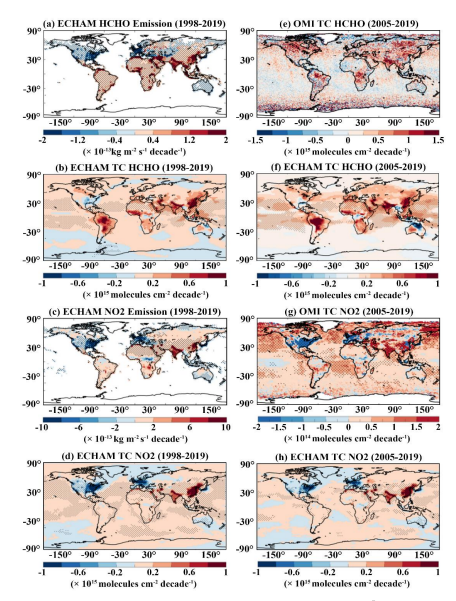


Figure 7. Trend in anthropogenic emission (kg m² s⁻¹ decade⁻¹) from ECHAM CTL simulation for the period 1998–2019 for (a–b) HCHO and NO₂ respectively. Trends in tropospheric column (TC) (molecules cm⁻² decade⁻¹) for (c–d) HCHO and NO₂ respectively. Trend in tropospheric column of HCHO from (e) OMI, and (f) ECHAM–CTL simulations for the period 2005-2019. (g–h) are the same as that of (e–f) but for tropospheric column NO₂. The stippled regions in the figures indicate significance at 95% confidence based on the t–test.

4. Influence of NOx and VOCs emissions on Formaldehyde to Nitrogen dioxide Ratio

In this section, we diagnosed the spatial distribution of tropospheric ozone production sensitivity regimes (NOx–limited/VOC–limited) associated with simulations of emission changes by using formaldehyde to nitrogen dioxide ratio (FNR). We estimate the FNR thresholds from ECHAM6–HAMMOZ model simulations adhering to the methodology outlined by Jin et al. (2017). The procedure to obtain FNR threshold from monthly mean data involves two steps: (1) obtaining the ozone response from emission sensitivity simulations (here, HalfNOx and HalfVOC simulations) by considering only the polluted cells over the study region and plotting it as a function of FNR (Fig. 8a), (2) calculating cumulative probability from this data for the conditions $d[O_3]/dE_{NOx} < 0$) (VOC limited) and $d[O_3]/dE_{NOx} > d[O_3]/dE_{VOC} > 0$) (NOx–limited) (Fig. 8b), where $d[O_3]/dE$ represents the change in ozone corresponding to a change in emission of either NOx or VOCs. This approach is applied to estimate FNR thresholds to distinctly delineate the ozone photochemical regimes as NOx or VOC–limited over major urban and semi-urban regions. The regions considered for estimating the FNR are shown in Figure 9.

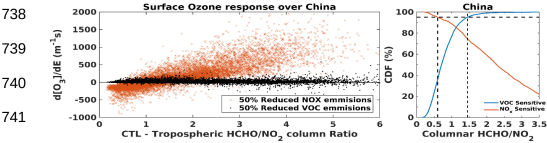


Figure 8. (a) Typical example of a normalized surface ozone sensitivity to a 50% reduction in global NOx (HalfNOx) and VOC (HalfVOC) emissions versus tropospheric column HCHO/NO $_2$ ratio derived from ECHAM6–HAMMOZ model simulation over China for the period 1998-2019, (b) Cumulative probability (CP) of VOC–sensitive (d[O $_3$]/dENOx<0) and NOx–sensitive (d[O $_3$]/dENOx>d[O $_3$]/dEVOC>0) conditions, as a function of tropospheric column HCHO/NO $_2$ as simulated by the ECHAM6–HAMMOZ model. The horizontal dashed line represents the 95% CP, and the vertical dashed lines represent the HCHO/NO $_2$ ratio corresponding to 95% CP for both the VOC–sensitive and NOx–sensitive curves demarcating the VOC–sensitive, NOx–sensitive, and transition regimes.

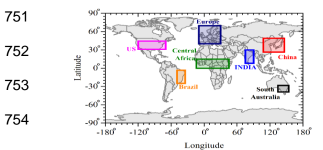


Figure 9. The rectangular box marks indicate the regions considered for estimating the HCHO/NO₂ ratio (FNR).

Further, we compared the model-estimated FNR with the OMI-derived FNR for the period 2005 – 2019. Figure 10 illustrates the comparison of FNR estimated from ECHAM6–HAMMOZ CTL simulations with OMI. The spatial map of FNR shows fairly good agreement between OMI and the model. Over the urbanized regions (e.g., South Asia, Europe, the US, and China) both the model and OMI show FNR < 4. In contrast, regions like North Canada, South America, central Africa, Australia, and Siberia exhibit high FNR values >9. There is good agreement between the model simulations and OMI, however, some minor differences are seen between the model and OMI FNR over the west coast of South America, South Africa, the Tibetan Plateau, and western Australia.



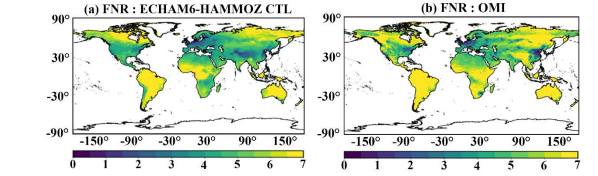


Figure 10. Spatial distribution of mean tropospheric column HCHO/NO₂ (FNR) obtained from ECHAM6–HAMMOZ CTL simulations (2005 – 2019) and OMI (2005 – 2019).

These differences could be due to the underestimation of HCHO in the model over these regions. Considering the fair performance of the model in comparison with OMI, we further analyzed the influence of changes in NOx and VOC emissions on the FNR from the model simulations, which are discussed in the subsequent sections.

Since the emission of HCHO and NO₂ varies with the seasons across the globe (e.g., De Smedt et al., 2015; Wang et al., 2017; Surl et al., 2018; Kumar et al., 2020; Goldberg et al., 2021; Guan et al., 2021), understanding the seasonal changes in FNR is crucial for comprehending shifts in ozone photochemical regimes. In this regard, we extracted the seasonal changes in

transition limits for the major urban and semi-urban regions shown in Figure 9 and summarized in Table 3. Figure 11 illustrates the seasonal variation of estimated FNR from both OMI data and model simulations across these key urban regions. In general, all regions exhibit distinct seasonal variations in transition limits (Table 3). Previously reported transition limits over the US (2-5): Johnson et al., 2024; 1.1-4: Schroeder et al., 2017) and China 0.6-1.5/1.25-2.39 (Chen et al., 2023) during the summer season are also compared with our model estimates. The estimated FNR values from the ECHAM6–HAMOZ simulations show fair agreement over both locations (0.4-4.6) in the US and 0.58-2.56 in China) with some minor differences. These minor discrepancies in the estimated FNR could be due to differences in the chosen location, time period and dataset used. Chen et al. (2023) have also reported that the transition limits depend on the region considered for the analysis.

Based on the threshold values depicted in Table 3 and the mean FNR in Figure 11, the seasonal change in ozone photochemical regimes over the key regions associated with the different emission scenarios are assessed. In the CTL simulation (Fig. 11e – h), the US, Europe, and China are found to be in the transition regime, while all other regions are NOx–limited during winter. In spring, every region except India remains NOx–limited, with India transitioning into the transition regime. During summer and autumn, all regions shift to a NOx–limited condition. We further compared the model-estimated regional FNR from the CTL simulation with the OMI-derived FNR shown in Figure 11a – d. The ozone photochemical regimes inferred from both OMI and the model show consistent results except during winter. During winter, the US, Europe and China are NOx limited in OMI, while our model shows them in the transition regimes.

Doubling NOx (DoubNOx) leads to a shift to a VOC-limited regime in all regions except Africa and Australia during winter, spring, and autumn (Fig 11i-1). The relatively high VOC contributions in Africa and Australia likely keep these regions in the transition regime. During summer, the US, Europe, Africa and Australia transform to the transition regimes, while all other regions remain VOC-limited. In both the DoubVOC and HalfNOx scenarios (Fig 11m-t), ozone photochemical regimes show no seasonality. All regions consistently exhibit a NOx-limited regime throughout all seasons. In the HalfVOC simulation (Figure 11u-x), the US, Europe, and China are in transition regimes, while all other regions become NOx-limited

during winter. India remains in a transition regime during all other seasons, whereas other regions consistently exhibit NOx–limited conditions. Figure 11 also depicts that DoubNOx and HalfNOx simulations greatly impact the shift in ozone photochemical regimes compared with DoubVOC and HalfVOC simulations. This indicates that ozone photochemistry is highly sensitive to changes in NOx emissions globally.

Table 3. Seasonal mean estimated values of the tropospheric HCHO/NO₂ columns threshold ratios from ECHAM6–HAMMOZ model control simulation to identify the NOx and VOC sensitive regimes across regions mentioned in Figure 9. The FNR less than the lower limit indicates VOC–limited, and that higher than the upper limit indicates NOx–limited regimes.

Sr. No.	Regions	Transition limits of FNR							
		DJF		MAM		JJA		SON	
1	US	0.48	1.04	0.49	1.15	0.49	4.69	0.45	1.39
2	Brazil	2.93	7.79	2.93	6.66	2.93	6.02	3.12	8.44
3	European Union	0.33	1.13	0.33	1.17	0.33	3.32	0.3	1.45
4	Central Africa	2.95	7.26	2.92	5.66	2.93	6.56	3.14	7.06
5	India	2.23	3.91	2.22	9.19	2.22	5.76	2.27	5.29
6	China	0.56	1.85	0.57	1.86	0.58	2.56	1.14	2.01
7	South Australia	1.1	5.54	1.09	2.3	1.09	1.82	1.12	3.93

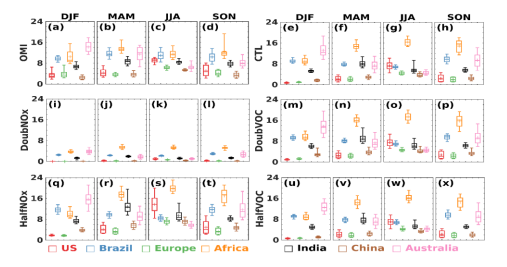


Figure 11. Box and whisker plot illustrating the long-term seasonal average FNR over the regions depicted in Fig.7 from (a-d) OMI observations, and model (e-h) CTL, (i-l) DoubNOx, (m-p) DoubVOC, (q-t) HalfNOx, and (u-x) HalfVOC simulations. Box represents 25 and 75 percentile and whisker represents 5 and 95 percentiles. The plus marker represents the mean, and the horizontal bar represents the 1 and 99 percentiles.

5. Influence of NOx and VOCs emissions on trends of Formaldehyde to Nitrogen dioxide Ratio

Trend analysis is carried out on FNR to understand the temporal evolution of ozone photochemical regimes associated with different emission scenarios. Figure 12 illustrates trends of FNR during the period 1998 – 2019 from CTL, DoubNOx, DoubVOC, HalfNOx, and HalfVOC simulations. In CTL simulation, decreasing (negative) trends in FNR are seen over the Asian region (-0.4 to -1.2 decade⁻¹) and Australia (-0.8 to -1.6 decade⁻¹), and an increasing (positive) trend in Europe (0.2 decade⁻¹) and the US (0.8 – 1.4 decade⁻¹) (Fig. 12a). These observed trends in FNR are mainly driven by the region-specific trends in HCHO and NO₂ (Fig. 7). Figure 7 shows a higher positive trend in NO₂ than in HCHO in the Asia region, causing an overall decreasing trend in FNR, indicating a tendency towards VOC-limited regimes. Whereas, over the US and Europe, there is a higher negative trend in NO₂ than HCHO, causing a positive trend in FNR, indicating a tendency towards a NOx–limited regime. A recent study by Elshorbany et al. (2024) also reported a significant positive trend over Europe and the US and a negative trend over Asia using the OMI-based tropospheric column HCHO/NO₂ ratio. Further, long-term column measurements of HCHO and NO₂ from OMI over India and China have revealed an increasing trend in NO₂ compared to that of HCHO, causing a decreasing trend in FNR over these regions (Jin and Holloway, 2015; Mahajan et al., 2015).

DoubNOx simulation (Fig. 12b) shows a similar spatial trend pattern to that of CTL simulation (Fig. 12a). However, the magnitude of this trend is less than that of the CTL. For example, a weak positive trend is noted in the US and Europe (0.2 – 0.4 decade⁻¹), while trends over India, and China are less negative (-0.2 to -0.4 decade⁻¹) in DoubNOx than CTL. (Fig. 12b). On the contrary, the magnitude of the positive trend over Canada and the negative trend over central Africa increased in DoubNOx emission, while the negative trend over Australia became nominal and insignificant. This indicates that Canada and central Africa have a tendency to become NOx–limited and VOC–limited respectively.

In DoubVOC simulations, trends are marginally increasing over the US, Canada, and Europe compared to the CTL (Fig. 12a and 12c). A notable change is observed over the Middle East and Amazon, where trends become more negative and positive respectively compared to CTL.

The negative trends over Australia in the CTL become nominal and insignificant in the

DoubVOC simulation. In HalfNOx simulations (Fig. 12d), the positive trends are higher over

the US, Europe and Amazon, while negative trends prevail over India, China and northeast Australia. Meanwhile, in HalfVOC simulation, marginal changes are noted globally compared to CTL. The most pronounced change in the FNR trend is observed over West Australia, where the negative trend in CTL becomes positive in HalfVOC (Fig. 12e). Figure 12f clearly shows that the trend in FNR is always negative over India and China for all the simulations, indicating that these regions have a tendency to become VOC–limited, while the positive trends over Europe and US show a tendency to become more NOx–limited. Further, from Figures 5, 11 and 12, we can infer that the relation between trends in FNR and ozone exhibits a nonlinearity. For example, even though FNR shows a negative trend over India and China for all the simulations, the TRCO trend depends on the specific emission scenario.

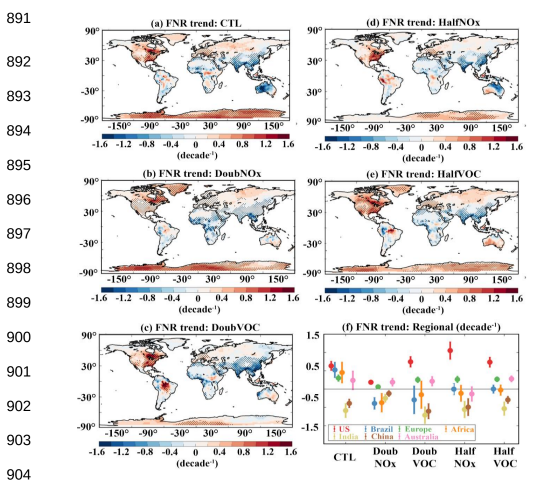


Figure 12. Trends in the tropospheric column HCHO/NO₂ ratio during 1998 – 2019 from ECHAM6–HAMMOZ simulations for (a) for CTL, (b) DoubNOx, (c) DoubVOC, (d) HalfNOx, (e) HalfVOC simulations. The stippled region indicates the trend significant at 95% confidence based on the t-test. (f) scatter plot illustrating the long-term trend and standard deviation over the regions depicted in Fig.9.

6. Tropospheric ozone radiative effects

The impact of emission changes on the tropospheric ozone radiative effect (TO3RE) is 912 913 estimated using the ECHAM6 model output and a radiative kernel method (see data and model 914 experiments). The estimated TO3RE for different model simulations are shown in Figure 13. 915 In the CTL simulations (Fig. 13a), the estimated global mean area-weighted average TO3RE for the period 1998 to 2019 is 1.21 [1.1 to 1.3] W m⁻². High TO3RE is noted over North Africa 916 and the Middle East region in NH (~2.2 W m⁻²), while in SH, it is over Australia and South 917 Africa (~1.2 W m⁻²). TO3RE estimates from TES measurements (2005 – 2009) also show a 918 peak of 1.0 W m⁻² in northern Africa, the Mediterranean, and the Middle East in June–July– 919 August (Bowman et al. 2013). Recently, Pope et al. (2024) reported TO3RE estimates from 920 IASI-SOFID, IASI-FORLI, and IASI-IMS for the period 2008 – 2017. The values reported 921 by Pope et al. (2024) are comparable with our CTL simulation (e.g. IASI–FORLI: 1.23 W m⁻², 922 IASI-SOFRID: 1.21 W m⁻², IASI-IMS: 1.21 W m⁻², ECHAM6: 1.22 W m⁻²). 923

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The anomalies of TO3RE from DoubNOx-CTL simulations are shown in Figure 13b. Doubling of NOx emission causes an enhancement in TO3RE by 0.36 [0.23 to 0.5] W m⁻² compared to the CTL simulation. It shows a peak over the Middle East and adjacent North Africa (0.7 W m⁻²). A similar peak over this region is also seen in the CTL simulation. Doubling of VOC emissions causes a marginal decrease in global mean TO3RE by -0.005 [-0.05 to 0.04] W m⁻². TRCO enhancement for doubling NOx is also higher than doubling VOC (see Fig.3). DoubVOC-CTL simulations (Fig. 13 c) show a peak over the Arctic (0.02 W m⁻²). The TO3RE anomalies are negative between $30^{\circ} \text{ N} - 30^{\circ} \text{ S}$. These negative anomalies in TO3RE between $30^{\circ} \text{ S} - 30^{\circ} \text{ N}$ (Fig. 13c) can be attributed to negative anomalies of TRCO (Fig. 3h). The reduction of NOx emission by 50% reduced global mean TO3RE by -0.12 [-0.2 to -0.05] W ${\rm m}^{-2}$ than CTL. The anomalies in TO3RE from HalfNOx-CTL simulations (Fig. 13d) show negative anomalies all over the globe, with a strong decrease over the Middle East and adjacent North Africa (-0.25 W m⁻²). Figures 13b and 13d show that the effect of enhancement/reduction of NOx emission is high over the Middle East and adjacent North Africa. The reduction of VOC emission by 50% reduced global mean TO3RE by -0.03 [-0.07] to 0.021 W m⁻² than CTL simulations (Fig. 13e). HalfVOC - CTL simulations show negative

anomalies of TO3RE between 40° S -40° N and positive 0.015 W m⁻² (low confidence) over mid-high latitudes in NH and SH. From Figure 13, it is interesting to note that the magnitude of TO3RE and its response to emission change is pronounced over the Middle East compared to all other regions. Further, Figure 13f depicts the latitude variation of zonal mean TO3RE for different sensitivity simulations. It is clear from Figure 13f that the TO3RE response to emission change is large at the northern and southern mid-latitudes, around $\pm 30^{\circ}$. Also, Figure 13f clearly indicates that the impacts of NOx emission changes are larger than VOCs throughout the latitude band.

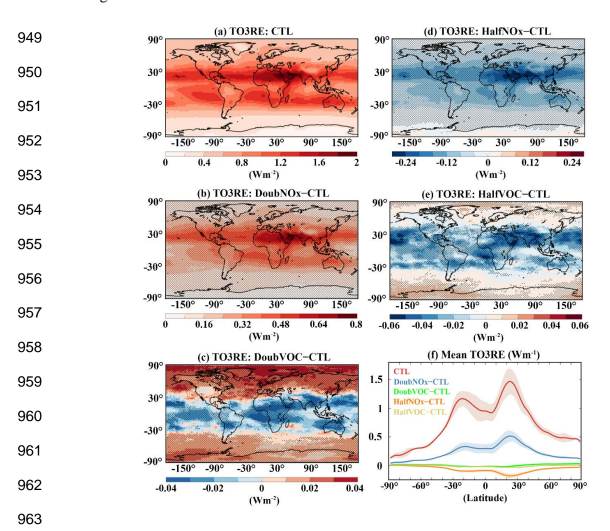


Figure 13. Tropospheric Ozone radiative effects (TO3RE) (W m⁻²) for (a) CTL, and anomalies from (b) DoubNOx - CTL, (c) DoubVOC - CTL, (d) HalfNOx - CTL, (e) HalfVOC - CTL simulations. Stippled regions in Figures (b–e) indicate TO3RE significant at 95 % confidence level based on the t-test, (f) line plot for zonal mean TO3RE (W m⁻²) from CTL, DoubNOx - CTL, DoubVOC - CTL, HalfNOx - CTL, HalfVOC - CTL, shades indicate standard deviation.

7 Summary and Conclusions

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- 971 In this study, we report variation of tropospheric ozone levels, trends, photochemical regimes
- and radiative effects using the state-of-the-art ECHAM6–HAMMOZ chemistry-climate model
- 973 simulations from 1998 to 2019. The model simulations are validated against multiple satellite
- 974 observations. Our analysis shows that
- 975 1. The estimated global mean trend in TRCO from CTL simulations for the period 1998 -
- 976 2019 is 0.89 [-0.07 to 2.1] ppb decade⁻¹. Trend estimates from OMI/MLS (1.43 [-0.5 to
- 977 3.2] ppb decade⁻¹) for the period January 2005 to December 2019 show good agreement
- with CTL $(1.58 [0.3 \text{ to } 3.3] \text{ ppb decade}^{-1})$ for the same period.
- 979 2. TRCO anomalies from DoubNOx CTL simulations show positive trends over Europe, the
- US, Africa, and South America, with a global mean trend of 1.2 [-0.1 to 2.7] ppb decade⁻¹.
- However, India and China show decreasing TRCO trend -2 to -4 ppb decade⁻¹. Surface
- ozone anomalies over these regions show strong negative trends -4.8 to -8 ppb decade⁻¹.
- 983 3. Global mean TRCO trend anomalies from DOUBNOX CTL simulation is 1.2 [-0.1 to
- 984 2.7] ppb decade⁻¹, while for DoubVOC CTL is 0.5 [-0.03 to 1.04] ppb decade⁻¹. Global
- mean TRCO trend anomalies from HalfNOx CTL is 0.47 [-0.76 to 1.3] ppb decade⁻¹ and
- 986 for HalfVOC CTL is 0.37 [-0.35 to 1.02] ppb decade⁻¹.
- 987 4. The spatial distribution of TRCO anomalies shows that enhancement is nearly 12 times
- 988 higher in DoubNOx CTL than in DoubVOC CTL simulations. The largest increase in
- 989 surface ozone anomalies from DoubVOC CTL is observed over Indo-Gangetic Plains,
- Eastern China and the eastern United States (4-6 ppb), where a decrease in surface ozone
- anomalies is observed in the DoubNOx CTL simulation. This decrease (increase) in ozone
- with an increase in NOx (VOC) indicates that these regions are VOC-limited.
- 994 5. The FNR transition thresholds exhibit pronounced seasonal variability. In the CTL
- simulation, the US, Europe, and China remain in the transition regime during winter, while
- other regions are predominantly NOx-limited. In spring, NOx-limited conditions persist
- 997 across all regions except India. A widespread shift to NOx-limited regimes is observed
- 998 during summer and autumn.

The DoubNOx simulation indicates a consistent shift toward VOC-limited regimes across
 most regions, except Africa and Australia, during winter, spring, and autumn. In summer,
 VOC-limited conditions prevail in most regions, whereas the US, Europe, Africa, and
 Australia are in a transition regime.

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- 7. In both the DoubVOC and HalfNOx scenarios all regions consistently exhibit a NOx–limited regime throughout all seasons. In the HalfVOC simulation the US, Europe, and China are in transition regimes, while all other regions become NOx–limited during winter.

 India remains in a transition regime during all other seasons, whereas other regions consistently exhibit NOx–limited conditions.
- 1010 8. Comparison of all the emission simulations, DoubNOx and HalfNOx simulations influence 1011 the shift in tropospheric ozone photochemical regimes compared to DoubVOC and 1012 HalfVOC simulations, highlighting the global sensitivity of ozone photochemistry to NOx 1013 emissions changes.
- 9. Trends estimated from modelled FNR are negative over India (-0.6 decade⁻¹) and China (-0.4 decade⁻¹) in all the simulations, indicating that these regions have a tendency to become VOC–limited, while the positive trends over Europe (0.3 decade⁻¹), US (0.63 decade⁻¹), and Africa (0.45 decade⁻¹), indicating a tendency to become more NOx–limited.
- 1019 1.3] W m⁻² which is increased by the doubling of NO_X emissions (DoubNO_X CTL) by
 1020 0.36 [0.23 to 0.5] W m⁻² and VOCs by -0.005 [-0.05 to 0.04] W m⁻² (DoubVOC CTL).
 1021 However, halving NO_X (HalfNO_X CTL) emissions shows a reduction in the global mean

10. The estimated global mean tropospheric ozone radiative effect (TO3RE) is 1.21 [1.1 to

- TO3RE by -0.12 [-0.2 to -0.05] W m⁻² and VOC (HalfVOC CTL) by -0.03 [-0.07 to 0.02]
- 1023 W m^{-2} .

- 1024 11. We show that anthropogenic NO_X emissions have a higher impact on tropospheric ozone levels, trends, and radiative effects than VOC emissions globally.
- Our study highlights the dominant role of anthropogenic NOx emissions in shaping tropospheric ozone trends, photochemical regimes, and radiative forcing over the past two decades. Sensitivity experiments reveal that NOx emission changes have a significantly larger impact on ozone levels and radiative forcing than VOCs, with regions transitioning from VOC-limited to NOx-limited regimes as NOx reductions continue. With current emission trends, air quality regulations, and industrial growth continuing to evolve, our study reinforces the need for region-specific mitigation strategies to

1032	effectively manage ozone pollution. This study emphasizes the importance of carefully balancing NOx
1033	and VOC controls to optimize air quality management and climate mitigation efforts. While climate
1034	change affects ozone chemistry, its impact over our study period (1998-2019) is minimal, given the
1035	relatively small global rate of temperature increase (\sim 0.3–0.4 $^{\circ}$ C) (IPCC AR6). Studies suggest that an
1036	ozone climate penalty emerges only after a 2-3°C temperature increase (Zanis et al., 2022), primarily
1037	in high-emission regions. Furthermore, rising temperatures increase water vapor, which reduces the
1038	ozone lifetime in remote areas. Thus, while climate change will play a growing role in future ozone
1039	variability, our findings confirm that anthropogenic emissions remain the dominant driver of recent
1040	ozone trends. As mitigation strategies evolve, targeted NOx and VOC controls must account for
1041	regional photochemical regimes, while future research should explore the potential interactions between
1042	climate change, natural VOCs, and ozone formation in a warming world.

- 1043 **Author's contribution**: SF and YE initiated the manuscript. SF made the model simulations.
- VS and SC did the analysis of model simulations. Satellite datasets are provided by JZ, BB,
- 1045 EF, IG, ID, MR, IS. All authors contributed to the writing of the manuscript.

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1047 **Competing interests:** At least one of the (co-)authors is a member of the editorial board of 1048 Atmospheric Chemistry and Physics.

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- 1056 scheme (award reference 4000137140).

1057 Data availability

- Available from the TOAR FTP server (ftpshare.al.noaa.gov).
- 1059 Code availability
- 1060 Available from the corresponding author upon reasonable request.

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