Insights into the long-term (2005-2021) spatiotemporal evolution of summer ozone production sensitivity in the Northern Hemisphere derived with OMI

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13 Abstract. Tropospheric ozone (O_3) formation depends on the relative abundance of precursor species, nitrogen oxides (NO_x) and 14 volatile organic compounds (VOCs). Advancements in satellite retrievals of formaldehyde (HCHO) and nitrogen dioxide (NO₂) 15 vertical column densities (VCDs), and the corresponding HCHO/NO₂ ratios (FNRs), provide the opportunity to diagnose the 16 spatiotemporal evolution of O_3 production sensitivity regimes. This study investigates trends of summertime VCD HCHO, NO₂, 17 and Ozone Monitoring Instrument (OMI)-derived FNRs in the Northern Hemisphere from 2005 to 2021. FNR trends were analysed 18 for polluted regions, specifically for 46 highly populated cities, over the entire 17-year period and in 2020 when global 19 anthropogenic emissions were reduced due to COVID-19 lockdown restrictions. It was determined that OMI-derived FNRs have 20 increased on average ~65% across cities in the Northern Hemisphere. Increasing OMI-derived FNRs indicates a general transition 21 from radical-limited to NO_x -limited regimes. The increasing trend is driven by reduced NO_2 concentrations because of emission 22 control strategies of NO_x. OMI FNR trends were compared to ground-based in situ measurements in US cities and determined they 23 can capture the trends in increasing FNRs (R = 0.91) and decreasing NO₂ (R = 0.98) occurring at the surface. OMI FNRs in urban 24 areas were higher (~20%) in 2020 for most cities studied here compared to 2019 and 2021. In addition to studying the longest 25 period of OMI FNRs across the Northern Hemisphere to-date, the capabilities and challenges of using satellite VCD FNRs to study 26 surface-level O₃ production sensitivity regimes are discussed.

Short Summary. Satellites, such as the Ozone Monitoring Instrument (OMI), retrieve proxy species of ozone (O₃) formation (formaldehyde and nitrogen dioxide) and the ratios (FNRs) which can define O₃ production sensitivity regimes. Here we investigate trends of OMI FNRs from 2005 to 2021 and they have increased in major cities suggesting a transition from radical- to nitrogen oxide-limited regimes. OMI also observed the impact of reduced emissions during the 2020 COVID-lockdown resulting in increased FNRs.

32 1 Introduction

33 Tropospheric ozone (O₃) is a harmful pollutant which has detrimental impacts on air quality leading to adverse human health and

34 premature mortality, and negative impacts on vegetation and agriculture (US EPA, 2006; Tai et al., 2014; GMD, 2020). A myriad

35 of volatile organic compounds (VOCs) can be photochemically oxidized through a complex series of chemical reactions involving

36 nitrogen oxides (NO_x = nitric oxide [NO] + nitrogen dioxide [NO₂]) which leads to tropospheric O₃ formation (Haagen-Smit, 1952;

37 Monks et al., 2015; Seinfeld and Pandis, 2016). The complex O_3 -NO_x-VOC chemical relationship results in local nonlinear O_3

38 formation which is sensitivity to the relative abundances of its precursor species (NO_x and VOCs), generally categorized as "NO_x-39 limited" versus "radical-limited" photochemical regimes (Sillman et al., 1990; Kleinman, 1994). In a NO_x-limited regime, local 40 O_3 production increases/decreases with increased/reduced NO_x emissions and concentrations, with no impact from VOC 41 perturbations. Whereas in a radical-limited regime (also known as "VOC-limited", "hydrocarbon-limited", or "NOx-saturated") 42 the formation of local O₃ increases/decreases with increased/reduced VOC emissions and concentrations; however, can also 43 slightly be impacted by NO_x emission and concentration changes. The accurate knowledge of regional and local O_3 photochemical 44 regimes is critical for developing emission control strategies to reduce surface O₃ concentrations. Overall, studying the 45 spatiotemporal evolution of the nonlinear O₃-NO_x-VOC chemistry is critical to policy decision making (National Research Council, 46 1991) and important as a fundamental scientific problem (Sillman, 1999).

47 Diagnosing regional and local O₃ photochemical regimes has always been recognized as a challenging task. Measurements 48 of proxy or indicator species (e.g., total reactive nitrogen, HCHO, hydrogen peroxide, nitric acid), and estimating the correlations 49 of such species, are the observation-based or model-observation synthesis approaches to detect O_3 sensitivity regimes (Sillman, 50 1995; Jacob et al., 1995; Tonnesen and Dennis, 2000). These measurements and associated studies are typically limited to field 51 campaign time periods and locations which hinders the spatiotemporal coverage of such data posing an obstacle for investigating 52 global and regional O_3 production sensitivity over multi-year time periods. The response of O_3 formation to changes in precursor 53 emissions can also be assessed through modeling approaches such as source apportionment studies (Li et al., 2012), forward model 54 sensitivity simulations (Wu et al., 2009), and simulations using adjoint model capabilities (Zhang et al., 2009). However, 55 uncertainties inherent in model predictions of O_3 physicochemical processes are a critical issue. Milford et al. (1994) and Sillman 56 (1995) first introduced the concept of detecting photochemical regimes using the ratio of ambient concentrations of two "indicator 57 species" HCHO to NO₂ (hereafter indicator species refers to HCHO and NO₂; the ratio of these two species as FNR), which can 58 be used to represent VOCs and NOx which are directly involved in O3-NOx-VOC chemistry and are readily measured via in situ 59 sampling and satellite remote-sensing techniques. These two indicator species are the most suitable candidates for tropospheric 60 column and planetary boundary layer (PBL) O_3 sensitivity analysis using satellites due to: 1) the sensitivity of nadir-looking 61 satellites to boundary layer FNRs, 2) most other indicator species (e.g., hydrogen peroxide, VOCs other than HCHO) cannot be 62 readily measured via satellites and the retrievals of those species having less sensitivity to surface conditions (e.g., limb-scanning 63 satellites), and 3) the short atmospheric lifetime of HCHO and NO₂ allowing these species to be suitable for proxies of surface 64 emissions of NO_x and VOCs.

65 Surface and PBL O₃ production sensitivity diagnosed with the in situ measurements of FNRs (although sparse in spatial 66 and temporal coverage) should be more accurate compared to satellite-based approaches of retrieving column-integrated 67 concentrations (Schroeder et al., 2017); however, the spatiotemporal coverage of polar orbiting satellites is a clear advantage over 68 in situ techniques. The advancements in satellite remote-sensing over the last two decades, to retrieve HCHO and NO₂ vertical 69 column density (VCD) data (Burrows et al., 1999; González Abad et al., 2019), have emerged as a new observation-based tool to 70 detect the spatiotemporal evolution of O_3 sensitivity from a global- to local-scale (Martin et al., 2004; Jin et al., 2020). Martin et 71 al. (2004) first demonstrated the capability of FNR VCDs from the Global Ozone Monitoring Experiment (GOME) satellite to 72 detect photochemical regimes. Subsequently, this technique was adopted by more studies using other satellite instruments such as 73 Ozone Monitoring Instrument (OMI), GOME-2, and TROPOspheric Monitoring Instrument (TROPOMI) (Duncan et al., 2010; 74 Witte et al., 2011; Choi et al., 2012; Choi and Souri, 2015; Chang et al., 2016; Jin and Holloway 2015; Souri et al., 2017; Jin et 75 al., 2017; 2020; Wang et al., 2021; Tao et al., 2022; Johnson et al., 2023; Acdan et al., 2023) up to a point that the results have 76 been suggested to potentially be used to inform State-Implementation Plans (SIP) in the United States (US) (Jin et al., 2018). 77 However, the accurate diagnosis of surface O_3 sensitivity regimes is impeded by numerous uncertainty components which can be

78 broadly classified into two major categories: 1) inherent uncertainties associated with the approach of relating indicator species to 79 diagnose local O_3 sensitivity at a location/time period, and 2) uncertainties associated with satellite-retrieved column-integrated 80 concentrations of indicator species to infer surface O₃ sensitivity. The former uncertainty arises from numerous factors: difficulties 81 in identifying accurate FNR "threshold" values (hereafter, "threshold" refers to threshold ratio values) separating different O_3 82 sensitivity regimes over a location and time period (Schroeder et al., 2017; Jin et al., 2017), dependency of ambient O_3 and its 83 formation to factors other than precursor species such as water vapor, meteorology, deposition, transport, and aerosol interaction 84 (e.g., Kleinman et al., 2005; Camalier and Dolwick, 2007), varying sensitivity of HCHO VCD data to anthropogenic VOCs (Jin et 85 al., 2020), and dependence of NO₂ in the production of ambient HCHO concentrations (Souri et al., 2020). These inherent 86 uncertainty sources limit the utility of satellite-based data for diagnosing O_3 sensitivity regimes. Fortunately, recent studies have 87 investigated these discrepancies in the methodology of using satellite-derived FNRs to infer O₃ sensitivity regimes using data from 88 airborne campaign data and 0-D photochemical box models (e.g., Schroeder et al., 2017; Souri et al., 2020; Souri et al., 2023a).

89 This study investigates 17 years (2005-2021) of OMI satellite sensor data which provides consistent near daily global 90 coverage of VCD retrievals of HCHO and NO₂ (Levelt et al., 2018) that are well suited to investigate the long-term spatiotemporal 91 evolution of O_3 sensitivity regimes. Numerous studies have used OMI VCD data up to the year 2016 to assess the trends in FNR 92 values over specific regions, mostly over the US and East Asia (Mahajan et al., 2015; Jin and Holloway, 2015; Souri et al., 2017; 93 Jin et al., 2017, 2020). Extending the OMI data set out to 2021 is novel and allows for the investigation of COVID-19 lockdown 94 restrictions on FNRs throughout the Northern Hemisphere. In this study we investigate the capability of VCD HCHO, NO₂, and 95 FNR data from OMI to reflect the trends in PBL and surface level O₃ production sensitivity regimes. We do not calculate actual 96 magnitudes of surface HCHO, NO₂, and FNRs derived with OMI VCD data as these proxy products are heavily reliant on chemical 97 transport models (CTMs) and spatiotemporally sparse ancillary information, both sources which have large uncertainties (discussed 98 in Sect. 4). Satellite retrieval errors can be reduced by averaging satellite data temporally (seasonal, annual, or multi-year means) 99 and spatially (by averaging individual satellite pixels across 10's to 100's of kilometers), although such averaging approaches 100 preclude the analysis of O_3 sensitivity regimes at high spatiotemporal scales. Several studies have therefore focused on assessing 101 O₃ production sensitivity using spatially-averaged satellite data aggregated to monthly, seasonal, or multi-year means over large 102 areas (e.g., Jin et al., 2020). In this study, we investigate the long-term changes in summer-mean (June, July, and August [JJA]) 103 VCD FNRs across numerous polluted cities (cities with high NO₂ VCDs) in the Northern Hemisphere. This manuscript is structured 104 in the following way. Section 2 describes the OMI retrievals, surface concentration measurements, "bottom-up" emission 105 inventories, and the approach to conduct spatiotemporal variability and trend analysis. In Sect. 3, we describe the comparison of 106 satellite VCD FNRs to surface measurements and the analysis of OMI-derived FNR values over Northern Hemisphere cities. 107 Section 4 discusses the capabilities and issues with applying satellite-derived FNRs for studying O₃ production sensitivity and 108 concluding remarks are provided in Sect. 5.

109 2. Materials and methods

110 2.1 OMI satellite sensor

The OMI sensor is a Dutch-Finnish built payload on the NASA Earth Observing System Aura satellite. The Aura platform flies as part of the Afternoon-Train satellite constellation along a sun-synchronous polar Low Earth Orbit (Schoeberl et al., 2006). Aura passes through the sunlit part of the Earth 14 times a day with a local overpass time of ~1:45 p.m. at the equator with near-complete daily global coverage (Levelt et al., 2006). OMI is a nadir-viewing solar backscatter grating spectrograph which takes retrievals in the ultraviolet (264–311 nm [UV1] and 307–383 nm [UV2]) and visible (349–504 nm) wavelengths (Levelt et al., 2006, 2018; Schenkeveld et al., 2017). The OMI instrument has a swath width of 2,600 km (60 pixels across track) with near-nadir spatial

- $117 \qquad \text{resolution of 13 km (along-track)} \times 24 \text{ km (cross-track) and near-swath edge pixel size of 40 km} \times 250 \text{ km}. \text{ OMI has been widely}$
- used by the atmospheric science, air quality, and health impact assessment communities since its launch on July 15, 2004 (e.g.,
 Levelt et al., 2018). The "row anomaly" appeared starting in May 2007 affecting the data quality of certain rows of OMI pixels
- 120 (Dobber et al., 2008, Schenkeveld et al., 2017) and is avoided in the data products used in this study.

121 2.1.1 OMI HCHO

122 This study applies the NASA-released operational OMI HCHO version 3 collection 3 (OMHCHO) gridded level 3 (L3) VCD data 123 at a spatial resolution of $0.1^{\circ} \times 0.1^{\circ}$ latitude \times longitude using the Smithsonian Astrophysical Observatory (SAO) retrieval 124 algorithm (González Abad et al., 2015). The OMHCHO retrieval applies a nonlinear fitting to the OMI-measured backscattered 125 radiances in the UV2 spectral window following the Basic Optical Absorption Spectroscopy method (Chance, 1998) to get slant 126 column densities (SCDs). The SCDs are then converted to VCDs by applying the air mass factor (AMF) formulation of Palmer et 127 al. (2001), with scattering weights calculated using the Linearized Discrete Ordinate Radiative Transfer version 2.4RT (VLIDORT) 128 Radiative Transfer Model (RTM) (Spurr, 2006) and HCHO shape factors simulated using the GEOS-Chem global CTM at a spatial 129 resolution of $2^{\circ} \times 2.5^{\circ}$ latitude \times longitude. The OMHCHO VCD product has a postprocessing bias correction (De Smedt et al., 130 2008) applied by comparing daily HCHO VCDs with background VCDs simulated with GEOS-Chem over a clean region in the 131 Pacific Ocean (González Abad et al., 2015). González Abad et al. (2015) estimated the uncertainty of the OMHCHO product 132 ranging from 45% to 105%, with relative contributions from the slant column retrievals (45% - 100%) and AMF calculations 133 (~35%). Model evaluation studies have shown that CTMs have errors and uncertainties in their predictions of HCHO in the clean 134 regions of the Pacific Ocean which could also contribute to overall OMI HCHO bias/errors (Anderson et al., 2017). Validation of 135 OMHCHO with aircraft-based observations indicates a high bias (66.1% - 112.1%) for HCHO-poor environments and low bias (-136 44.5% - -21.7%) for HCHO-rich environments (Zhu et al., 2016, 2020; Johnson et al., 2023). The OMHCHO product has been 137 used widely for estimating trends of VOC emissions (e.g., Marais et al., 2012; Shen et al., 2019) and inferring surface HCHO 138 concentrations (Zhu et al., 2017a).

139 2.1.2 OMI NO₂

140 The NASA-released standard OMI NO₂ (OMNO2) version 4 collection 3 gridded L3 high resolution VCD data at the spatial 141 resolution of $0.1^{\circ} \times 0.1^{\circ}$ latitude \times longitude was applied in this study (Lamsal et al., 2015; Krotkov et al., 2017). The OMNO2 142 retrieval uses the Differential Optical Absorption Spectroscopy method (Marchenko et al., 2015) to derive tropospheric SCDs by 143 spectrally fitting OMI-detected backscattered radiance in the visible wavelength window with a pseudo reference spectrum 144 (Chance and Spurr, 1997). The stratospheric contribution of the SCD is then subtracted and the residual tropospheric SCDs are 145 then converted to tropospheric VCDs by applying an AMF based on scattering weights calculated using the Total Ozone Mapping 146 Spectrometer (TOMS) radiative transfer model (TOMRAD) (Dave, 1964) and shape factor profiles simulated using the Global 147 Modeling Initiative (GMI) CTM at a spatial resolution of $1^{\circ} \times 1.25^{\circ}$ latitude \times longitude (Krotkov et al., 2017). The uncertainty of 148 the OMNO2 VCD product varies with cloudiness and pollution levels but is in the range of $\sim 20\% - 60\%$ (Bucsela et al., 2013), 149 with relative contributions from the spectral fitting (~10% over polluted regions, Boersma et al., 2011), stratospheric correction 150 (<5%), and AMF calculations (10% – 20%). The OMNO2 product has been used for a wide range of applications including the 151 estimation of spatiotemporal variability and trends of NO_x emissions (e.g., Krotkov et al., 2016), NO₂ surface concentrations (e.g., 152 Kharol et al., 2015; Lamsal et al., 2015), information about atmospheric particulate organic matter (Philip et al., 2014), and surface 153 O₃ sensitivity regime detections (e.g., Duncan et al., 2010; Jin et al., 2017).

154 2.1.3 Calculation of VCD FNR values

The daily L3 OMHCHO and OMNO2 products were filtered and processed for calculating FNR values. Both the operational products were already filtered for daily VCDs with an effective cloud fraction >30%, solar zenith angle >70% (for HCHO) and >85% (for NO₂), removing pixels affected by row anomalies, and Level 2 data quality flags not designated as good (see more details in the user's guides for OMNO2 [https://disc.gsfc.nasa.gov/datasets/OMNO2d_003/summary] and OMHCHO [https://acdisc.gesdisc.eosdis.nasa.gov/data/Aura_OMI_Level3/OMHCHOd.003/doc/README_OMHCHOd_v003.pdf]).

- 160 During this study, we avoided unrealistically large positive and negative values for both indicator species which occur due to
- 161 uncertainties in slant column retrievals and the calculation of tropospheric VCDs. We followed Zhu et al. (2020) to filter out HCHO
- $\label{eq:linear} \begin{array}{ll} \mbox{daily VCDs outside the range of -8.0 \times 10^{15}$ molecules/cm^2$ to 7.6×10^{16} molecules/cm^2$. The OMNO2 L3 product already included 1.5×10^{16} molecules/cm^2$ to 7.6×10^{16} molecules/cm^2$. The OMNO2 L3 product already included 1.5×10^{16} molecules/cm^2$ to 7.6×10^{16} molecules/cm^2$. The OMNO2 L3 product already included 1.5×10^{16} molecules/cm^2$ to 7.6×10^{16} molecules/cm^2$. The OMNO2 L3 product already included 1.5×10^{16} molecules/cm^2$ to 7.6×10^{16} molecules/cm^2$. The OMNO2 L3 product already included 1.5×10^{16} molecules/cm^2$ to 7.6×10^{16} molecules/cm^2$. The OMNO2 L3 product already included 1.5×10^{16} molecules/cm^2$ to 7.6×10^{16} molecules/cm^2$. The OMNO2 L3 product already included 1.5×10^{16} molecules/cm^2$ to 7.6×10^{16} molecules/cm^2$. The OMNO2 L3 product already included 1.5×10^{16} molecules/cm^2$ to 7.6×10^{16} molecules/c$
- an upper limit of 1×10^{17} molecules/cm² and we applied a lower limit of -1×10^{15} molecules/cm² below which NO₂ VCD are
- assumed in this study to be unrealistic. After data filtering, OMI VCD FNRs are calculated by taking the ratio of HCHO:NO₂ for
- each grid of the summer-mean products.

166 2.2 Surface measurement data

167 To determine if OMI VCD FNRs can replicate the trends of PBL and surface-level FNRs, long-term trends in OMI-derived VCD 168 HCHO and NO₂ are compared to in situ measurement data from the United States Environmental Protection Agency's Air Quality 169 System (US EPA AQS; https://www.epa.gov/aqs). We focus this evaluation on the US due to the much denser in situ measurement 170 networks compared to other global regions. Hourly data from the EPA AQS NO₂ data were averaged daily from 1 pm to 3 pm 171 local time to be consistent with the OMI overpass time. Since there is insufficient hourly data for HCHO from the EPA AQS 172 network, we use 24-hour average data for the HCHO evaluation which is provided by the EPA. AQS data for HCHO and NO₂ 173 from each site are only used for days in which both species are measured. Valid and continuous data points were then averaged to 174 obtain seasonal summertime mean (JJA) values from 2005 to 2019 to be intercompared with corresponding OMI VCD values.

The AQS NO₂ data suffers from potential interference of reactive nitrogen species while measuring NO converted from NO₂ in molybdenum catalytic converters, since other reactive species also get converted to NO. We attempted to account for this interference by applying a model simulated correction factor (CF; Eq. 1) to the raw AQS data, following the approach of previous studies (Lamsal et al., 2008; 2010; Cooper et al., 2020).

179
$$CF = \frac{NO_2}{NO_2 + (0.15 \times HNO_3) + (0.95 \times PAN) + Alkyl Nitrates}$$
(1)

180 The CF is calculated using the MERRA2-GMI simulated concentrations of NO₂, HNO₃, peroxyacetyl nitrate (PAN), and alkyl 181 nitrates, and by applying an assumed effective conversion efficiency of 15% for HNO₃ and 95% for PAN (see Lamsal et al., 2010). 182 The AQS's method to measure HCHO is affected by interference from species such as O₃ and NO₂ and since there is insufficient 183 information to correct those interferences, here we use uncorrected AQS HCHO data.

184 2.3 Surface emissions of NO_x

To compare the long-term evolution of FNRs with human-induced changes in precursor emissions (anthropogenic emissions of NO_x), we used the most recent Community Emissions Data System (CEDS v_2021_04_21) NO_x bottom-up emission data set (McDuffie et al., 2020). As explained in the results section of this manuscript, we focus our analysis on trends in NO_x emission instead of HCHO as it was determined that trends in NO₂ concentrations clearly drive the global trends in FNRs. The CEDS data provides monthly anthropogenic NO_x emissions at $0.5^{\circ} \times 0.5^{\circ}$ horizontal spatial scales from 1750 – 2019. For this study we analyse trends in anthropogenic NO_x emissions (source sectors: Agriculture; 1: Energy; 2: Industrial; 3: Transportation; 4: Residential,

- 191 Commercial, Other; 5: Solvents production and application; 6: Waste; 7: International Shipping) between 2005 2019 to overlap
- 192 with OMI observations. We used mean emissions for summer months (JJA) for each year to intercompare with OMI derived NO₂
- and FNR trends. Since our focus in this study was to assess the overall relationship of long-term changes in OMI-derived FNR
- values and corresponding changes in the anthropogenic NO_x over a city/region, we do not consider other natural sources (e.g.,
 biomass burning) contributing to ambient concentrations of urban NO₂.

196 2.4 Spatiotemporal analysis of FNRs

- 197 The spatiotemporal analysis of OMI-derived VCD NO₂ and HCHO values was conducted as follows. First, summer-mean trends 198 from 2005 to 2021 of HCHO and NO₂ VCDs and FNR values were calculated at the native spatial resolution $(0.1^{\circ} \times 0.1^{\circ})$. Long-199 term trends were calculated for each grid of HCHO, NO₂, and FNRs with ordinary least-squares linear regression (at various 200 confidence levels calculated with the Mann-Kendall Test) similar to past studies (e.g., Boys, et al., 2014; Kharol et al., 2015; 201 Geddes et al., 2016). To reduce retrieval random errors and improve precision, we focus on summer-mean data for each year and 202 multi-year means (three multi-year means: 2005 - 2010, 2011 - 2015 and 2016 - 2021) around 46 cities across the Northern 203 Hemisphere. The focus on the summer season was also chosen to utilize HCHO VCD retrievals with significantly better signal to 204 noise ratios compared to winter, spring, and fall months. High levels of surface HCHO concentrations over source regions form 205 due to the higher oxidant availability in summer (González Abad et al., 2015; Zhu et al., 2014; 2017a; 2017b) which leads to better 206 retrievals of HCHO VCDs. We restrict our analysis to the Northern Hemisphere as most continental polluted regions exist there. 207 We assessed the evolution of FNRs over urban and rural/suburban (hereinafter referred to just as rural) areas around cities. To 208 define urban city regions, we used the hybrid dataset, CGLC-MODIS-LCZ (Demuzere et al., 2023), which is based on the 209 Copernicus Global Land Service Land Cover (CGLC) product resampled to MODIS IGBP classes (CGLC-MODIS) and the global 210 map of Local Climate Zones (LCZ) (Stewart and Oke., 2012) that describe the heterogeneous urban land surface. This data set was 211 interpolated onto a $0.1^{\circ} \times 0.1^{\circ}$ grid to match the resolution of the L3 OMI satellite data. Urban classification is defined by the 212 CGLC-MODIS-LCZ land use categories 51 - 60, which includes a range of urban land use from sparely built to compact high-rise 213 including the heavy industry category. These land use categories capture both urban and suburban landscapes. Rural grids are those 214 not defined as urban and within 7×7 grid boxes of the city center. The CGLC-MODIS-LCZ urban and rural maps derived for this 215 study are static and will not capture urban expansion which has occurred over the last two decades. However, since our urban 216 classification includes both urban and suburban landscapes (including sparsely built-up areas), the transition from suburban to 217 urban landscapes will already be included in our urban map. The only thing not captured would be the transition from completely 218 vegetative areas to more built-up landscapes which are expected to have a minor impact on the results of this study.
- 219 In this study, we focus only on the spatiotemporal variability of indicator ratios, rather than the exact ozone sensitivity 220 regimes which can be inferred from these ratios. Although several previous studies assigned ratio values to certain O₃ regimes 221 (e.g., Jin et al., 2017; Souri et al., 2017) based on previous modelling and limited-observational studies, large uncertainty exists in 222 the classification of O₃ regimes using FNR values (Schroeder et al., 2017; Jin et al., 2020; Souri et al., 2021). Nevertheless, 223 whenever the ratio values were assessed over a city/region, we also presented the threshold ratio values (separating O₃ regimes) 224 suggested by Jin et al. (2020) for some cities in the US (Los Angeles, New York, Chicago, Washington DC, Pittsburgh, Atlanta 225 and Houston), Wang et al. (2021) for cities in China, and Duncan et al. (2010) for all other cities/regions. Note that the threshold 226 FNR values (< 1 as radical-limited versus > 2 as NO_x-limited) suggested by Duncan et al. (2010) is a crude approximation as 227 opposed to more recent and observationally-constrained threshold ratio values suggested by Jin et al. (2020) and Wang et al. (2021). 228 We believe that an accurate classification of O₃ regimes is still an ongoing research topic (Schroeder et al., 2017; Jin et al., 2020;
- 229 Souri et al., 2021) which should be addressed in future studies.

230 **3. Results**

231 3.1 Long-term mean OMI data

232 Figure 1 shows the long-term mean (2005-2021) maps of OMI-derived VCDs of HCHO and NO₂ and the corresponding column 233 FNR values. Formaldehyde enhancements reflect surface emissions of anthropogenic VOC (densely populated regions in China; 234 Shen et al., 2019), biogenic isoprene (southeast US; Millet et al., 2008), and biomass burning (South Asia; Mahajan et al., 2015). 235 OMI VCD NO₂ is abundant over urban areas primarily due to fossil fuel combustion emissions from traffic (Duncan et al., 2015) 236 and over regions with large industrial activities (Krotkov et al., 2016). The column FNRs clearly reveal lower values over cities 237 (FNR < 2), marginal values over rural/suburban regions surrounding large cities (FNR in the range of 2 - 5), and higher values 238 elsewhere (FNR > 5). The lower FNRs over cities suggest radical-limited conditions, and larger FNR values in non-polluted 239 background regions reflect NO_x-limited conditions (Martin et al., 2004; Duncan et al., 2010; Jin et al., 2017; 2020; Wang et al., 240 2021). Lower FNR values retrieved by OMI are most noticeable in the highly populated regions of the US (e.g., Los Angeles, New 241 York, Chicago), Europe (e.g., London, Amsterdam, Paris), East Asia (e.g., Beijing, Shanghai, Jinan), and Middle East (e.g., Dubai, 242 Tehran, Riyadh) where tropospheric column NO₂ abundances are enhanced. The highest FNR values are observed in regions of 243 the southeast US and south Asia (e.g., Malaysia) where there are no large cities and enhanced tropospheric column HCHO 244 abundances, primarily from biogenic emissions, are observed.

245 3.2 Capability of OMI VCD data to observe surface-level FNR trends

246 Before assessing VCD FNR trends, we compared trends in OMI NO₂ and HCHO VCD data, and corresponding tropospheric 247 column FNRs, to surface in situ measurements from EPA AQS in the US in order to determine whether OMI VCD information 248 tracks trends occurring at the surface. Figure 2 shows the 15-year time-series (2005-2019) comparison between normalized time 249 series of OMI VCD indicator species abundances and FNRs and AQS data over select cities (US cities with continuous AQS data), 250 and over all cities averaged across the continental US (USA, 373 separate sites). Table 1 shows the correlation between OMI VCD 251 and AQS in situ NO₂, HCHO, and FNR summer mean values in addition to the simple linear regression slope of normalized trends 252 from OMI and AQS for both indicator species and FNRs. Figure 2 shows that both OMI VCD and in situ AQS data have relatively 253 neutral trends in HCHO between 2005 and 2019 for most of the large urban cities of the US. While there is large interannual 254 variability in HCHO concentrations, the long-term trends are relatively flat. On average, the normalized linear trends in surface 255 HCHO in urban regions of the US was -0.05 yr⁻¹ and OMI VCDs was +0.15 yr⁻¹. OMI VCD HCHO data is unable to replicate the 256 interannual variability and long-term trends of surface data displayed by the low correlation values and opposing trends in multiple 257 large cities in the US. The inability of OMI to reflect the variability in HCHO observed at the surface is likely due to the coarse 258 spatial resolution of the OMI footprint, large noise in OMI HCHO retrievals (e.g., Johnson et al., 2023, Souri et al., 2023a), and 259 complex vertical distributions of HCHO complicating satellite retrievals and representation of surface values (e.g., Souri et al., 260 2023b).

OMI VCD and in situ AQS data of NO₂ display a different story where statistically significant reductions at a 95% confidence level in NO₂ concentrations are observed by both data sources. The normalized trends in NO₂ from both measurement platforms are in strong agreement (see Fig. 2 and Table 1). Correlation between OMI VCD and AQS NO₂ was near 1.0 (R=0.98) and both data sources had normalized linear regression slopes of ~-0.20. This suggests that OMI is able to observe the strong reduction in NO₂ concentrations at the surface measured by AQS across the US. Both data sources suggest that NO₂ reduced between 2005-2019 and the strongest negative trends were in the large cities of the US such as New York, Chicago, and Los Angeles. The near neutral trend in HCHO and large decreases in NO₂ results in both OMI VCD and in situ data sources observing an increasing trend in FNR data in all major cities of the US shown in Fig. 2. The normalized linear regression trend slopes of FNRs (0.21 from both OMI and AQS data) are all statistically significant reductions at a 95% confidence level and are nearly equal and opposite to NO₂, suggesting the reduction of NO₂ is the primary driver of FNR trends over time. It is encouraging that OMI VCD data is able to accurately reproduce the normalized trends in surface FNRs in the US. This agrees with the recent studies from Jin et al. (2017) and Souri et al. (2023a) which show that ratios of mid-day tropospheric VCD FNRs to PBL and surfacelevel concentrations are near unity. Since OMI VCD FNRs appear to be able to replicate the trends in surface FNRs, the rest of this study focuses on the trends of FNRs from OMI VCD data for the Northern Hemisphere.

275 3.3 17-year trend in OMI observations

276 This study investigates the 17-year trend of OMI VCD of HCHO, NO₂ and FNR values between 2005 and 2021. Figure 3 shows 277 the long-term trend in OMI VCD HCHO, NO₂ and FNR values at an 85% confidence level ($p \le 0.15$) (Fig. S1 shows the same 278 trend values at a 99% [$p \le 0.01$] confidence level and for all grid cells with OMI retrievals). The same information shown in Fig. 279 3 is displayed individually for North America, Europe, and Asia in the supplemental information (see Fig. S2). Formaldehyde VCDs increased by $\sim 0.5 \times 10^{14}$ molecules cm⁻² yr⁻¹ over most of the Northern Hemisphere, with reductions up to $\sim -0.5 \times 10^{14}$ 280 281 molecules cm⁻² yr⁻¹ over the southeast US. This trend is consistent with previous studies documenting increases in multi-satellite 282 (including OMI) summer mean HCHO over Northern China (during 2005-2016 [Shen et al., 2019] and 2005-2014 [Souri et al., 283 2017]) and increases in most regions in the US (during 2005-2014 [Zhu et al., 2017b]) due to increasing anthropogenic VOC 284 emissions. Some near neutral trends, to small decreases, are seen in Fig. 3 in eastern China. The decrease in HCHO over Southern 285 China could be due to reductions in anthropogenic VOCs (Souri et al., 2017; Itahashi et al., 2022) or biogenic VOC emissions as 286 noted by Jin and Holloway (2015). The decreases in summer mean OMI HCHO over the southeast US were also documented in 287 earlier studies (De Smedt et al. 2015; Zhu et al., 2017b). Note that the trend in HCHO calculated in this study is influenced by 288 yearly variations in temperature, in contrast to Shen et al. (2019) and Zhu et al. (2017b) that corrected for the impact of varying 289 temperature on HCHO VCDs.

290 It should be noted that the NASA-released operational OMI HCHO version 3 collection 3 data product used in this study 291 has been shown to have a positive drift due to instrument aging (e.g., Marais et al., 2012; Zhu et al., 2014, 2017b). This positive 292 trend in OMI HCHO data displayed in Fig. 3 is likely largely impacted by the artificial positive drift in the collection 3 OMI data. 293 A new NASA OMI HCHO version 3 collection 4 product is in development using the SAO algorithm which has removed this 294 positive drift in HCHO (Ayazpour et al., 2024; personal communication with the SAO HCHO algorithm team). This new HCHO 295 retrieval product shows that HCHO has a near-neutral trend across most of the populated cities in the Northern Hemisphere. This 296 new collection 4 retrieval data is not vet peer-reviewed or available to the public therefore is not used here and the remaining 297 results in this study use OMI HCHO version 3 collection 3 data. However, to test the potential impact on the results of this study 298 using an OMI VCD product with this average positive drift eliminated, we removed the mean annual Northern Hemispheric HCHO 299 trend (~0.004 DU yr⁻¹) from the collection 3 data and evaluate the resulting FNR trends over 18 selected large cities in the Northern 300 Hemisphere which is discussed in Sect. 3.4.

The negative trend in NO₂ OMI VCDs over populated regions of the US, Europe, and eastern China, and increases in the South Asia and Middle East regions, seen in Fig. 3 are consistent with several previous studies (e.g., Hilboll et al., 2013; Jin et al., 2017). The decreases in eastern China and Europe are as large as -2.0×10^{14} molecules cm⁻² yr⁻¹ while reductions in NO₂ in the US are between -0.1 to -1.0×10^{14} molecules cm⁻² yr⁻¹. The decreasing NO₂ trend in eastern China could be due to recent reductions in anthropogenic NO_x emissions after year 2011 (e.g., Fan et al., 2021). It is well demonstrated that the tropospheric NO₂ decreases in most Northern Hemisphere regions, particularly in urban regions, is due to reductions in anthropogenic NO_x emissions implemented through national governmental policies (e.g., Duncan et al., 2016; Koplitz et al., 2021). Figure S3 shows the trends in CEDS anthropogenic NO_x emissions between 2005-2019 which have nearly identical regions of reduction as those retrieved by OMI NO₂. Overall, the summer-mean trend in VCD NO₂ estimated in this study is generally consistent with the reported satellitebased annual mean surface NO₂ trend estimated on a global-scale (Geddes et al., 2015) and over the US (Kharol et al., 2015; Lamsal et al., 2015).

The most notable feature in Fig. 3 is the general increasing trend in OMI VCD FNR values over most of the polluted regions in the Northern Hemisphere. The increasing OMI column FNR values suggest a trend towards more NO_x -limited regimes around cities in recent years which has been noted by some previous studies (Jin et al., 2017; 2020; Souri et al., 2017). Increases in FNRs in the populated regions of China, Europe, and US reach values between 0.1 and 0.2 yr⁻¹. The increases in FNRs are driven mostly by the reductions in NO_2 rather than the small variations in HCHO, as evident in Fig. 2 and 3. The following sections focus on the assessment of the evolution of summer mean OMI-derived VCD FNRs over numerous selected cities in the Northern Hemisphere.

319 3.4 Evolution of OMI FNRs around populated cities in the Northern Hemisphere

320 Figure 4 shows the time series of summer mean OMI VCD FNRs from 2005 to 2021 over 18 selected large cities. The 321 corresponding normalized time series trends of OMI-derived NO₂ abundances and FNRs, and CEDS anthrophonic emissions of 322 NO_x over these cities, are displayed in Fig. 5. From Fig. 4 it can be seen that the largest positive trends in OMI FNRs during the 323 2005-2021 time period occurred over three mega-cities in the US: Los Angeles, New York and Chicago. Time series of the actual 324 magnitudes of OMI VCD NO₂ and HCHO abundances over the selected 18 large cities are shown in Fig. S4. In addition to increases 325 in FNRs in US cities, relatively large increases in FNRs are also evident in European (e.g., London) and Asian (e.g., Guangzhuo) 326 cities. To test whether the positive drift in the NASA OMI HCHO collection 3 data significantly impacted the results of the FNR 327 trends over the 18 selected large cities in the Northern Hemisphere we present these same results in Fig. S5 with the OMI data 328 which has the annual average Northern Hemispheric HCHO trend remove (more representative of OMI HCHO version 3 collection 329 4 data) and Fig. S6 shows the spatial trends of HCHO, NO₂, and FNRs over the Northern Hemisphere using this detrended HCHO 330 data. Comparing Figs. S5 and 4, it is seen that while some of the FNR values are slightly lower in magnitude the positive trends 331 are very similar using collection 3 HCHO retrievals and a data product with the positive drift removed. Throughout the Northern 332 Hemisphere HCHO trends now display both positive and negative values (see Fig S6) instead of the constant positive trends from 333 the OMI HCHO collection 3 product. Using the detrended OMI HCHO data does result in more negative FNR trends in remote 334 regions outside of large urban regions; however, over urban areas, and rural regions surrounding large cities, the FNR trends are 335 still positive as displayed in Fig. S5 and S6. Overall, using the OMI HCHO version 3 collection 3 data product does not significantly 336 impact the FNR results in large cities in the Northern Hemisphere focused on in this study. Future studies investigating FNRs 337 conducted when the NASA OMI HCHO version 3 collection 4 data is available to the public should however use this new product 338 to present more accurate results compared to those shown here using the NASA OMI HCHO version 3 collection 3 product.

Figure 5 reveals that the increases in OMI FNR values over the selected 18 mega-cities are linked with decreases in NO_2 abundances due to reductions in anthropogenic NO_x emissions. The spatial map of trends in CEDS anthrophonic emissions of NO_x across the Northern Hemisphere between 2005-2019 are shown in Fig. S3 (timeseries of CEDS NO_x emission magnitudes for the selected 18 mega-cities shown in Fig. S7). Based on the O_3 production sensitivity regime thresholds suggested by Jin et al. (2020) (note that these thresholds are applicable for VCD data), all the US cities shown in Fig. 4 that were VOC-limited in the early 2000's show clear transition towards NO_x -limited and transitional regimes in recent years. Major cities in Europe such as London and Amsterdam have also experienced increasing FNRs moving from VOC-limited regimes to transitional, or even NO_x -limited,

346 regimes in recent years (based on thresholds from Duncan et al. [2010]). Increases in the magnitudes of FNRs were generally 347 smaller in large cities of Asia; however, only Neijiang doesn't display some noticeable increases in FNRs in recent years. In 348 Neijiang, CEDS anthropogenic NO_x emissions are decreasing after 2012; however, OMI does not retrieve decreasing NO_2 349 abundances leading to the near-neutral trend in FNR values. Based on the O_3 production sensitivity regimes thresholds defined by 350 Wang et al. (2021) and Duncan et al. (2010), major cities in Asia have FNR values which are in the transitional or NO_x -limited 351 regimes in recent years besides Beijing, Shanghai, Jinan, and Riyadh (surrounding rural region is in the transitional regimes) (see 352 Fig. 4). Figure 5 shows these large Asian cities, besides Riyadh, implemented NO_x emission control strategies in ~2012 and have 353 recent negative trends in OMI NO₂; however, based on Wang et al. (2021) these urban regions have O₃ production which is still 354 limited by VOCs. Overall, it is difficult to conclude if these major cities in the Northern Hemisphere have in fact transitioned to 355 NO_x -limited and transitional regimes due to the large uncertainties in the exact threshold FNR values which separate these chemical 356 regimes.

357 In the vast majority of cities between 2005-2021 OMI retrieved larger FNR values in the rural regions surrounding urban 358 regions in the Northern Hemisphere compared to the urban city centers. Figure 6 shows spatial maps of OMI-derived VCD FNRs 359 around the selected 18 cities discussed above for two 6-year averages: 2005-2010 and 2015-2021 reflecting the earliest and most 360 recent years of OMI data studied here. The spatial maps of OMI-derived HCHO and NO₂ VCD values for these same time periods 361 over the 18 cities are displayed in Fig. S8 and S9, respectively. Figure 6 shows that OMI is able to retrieve the differences in FNRs 362 in urban and rural regions surrounding large cities in the Northern Hemisphere (Fig. S10 shows the same information in Fig. 6 363 except with the CGLC-MODIS-LCZ urban grids used to separate urban and rural values). In urban areas of cities, where emission 364 sources of NO_x are largest, OMI FNRs tend to be lower indicating more tendency towards VOC-sensitive O₃ production regimes 365 compared to the surrounding rural regions. This figure also displays the decadal-scale changes (2016–2021 versus 2005–2010) in 366 the OMI FNRs over the Northern Hemisphere urban regions and surrounding rural regions. In all 18 cities FNR values increase in 367 both rural and urban areas with noticeable increases in the spatial coverage of potentially NO_x -limited O_3 production regimes. 368 These spatial distributions of increasing FNR values retrieved by OMI are clearly correlated with decreasing tropospheric NO_2 369 over the vast majority of cities displayed in Fig. S9. Large cities in the US show the clearest increase in the spatial coverage of 370 potentially NO_x-limited O₃ production regimes; however, European and Asian cities also follow a similar pattern with less increase 371 in FNR magnitude overall. Recent studies have also noted that NO_x-limited regimes have expanded spatially into the city centers, 372 on a decadal-scale, throughout the Northern Hemisphere (Jin et al., 2017) and in the US (Jin et al., 2020). This has large implications 373 for O₃ sensitivity analysis and development of future emission control strategies for improving air quality.

374 Figure 7 shows the changes in OMI FNRs (multi-year averaged values: 2005-2010, 2011-2015, and 2016-2021) over 46 375 cities in the Northern Hemisphere. The vast majority of urban regions in the Northern Hemisphere (44 of the 46 selected cities) 376 experienced increasing FNRs between 2005-2010 and 2016-2021. OMI FNRs for Tehran, Iran and Neijiang, China were two 377 selected cities which did not display increasing values. On average, FNRs in urban areas of the selected cities increased by ~65% 378 between 2005-2010 and 2016-2021. Similar to urban regions, the vast majority of rural regions surrounding urban areas in the 379 Northern Hemisphere (44 of the 46 selected cities) experienced increasing FNRs between 2005-2010 and 2016-2021. The average 380 increase of FNRs in the rural regions increased slightly less (~38%) compared to urban areas. In agreement with results discussed 381 above, FNR values in rural regions are larger compared to city centers. However, OMI VCD FNR differences between rural and 382 urban regions were reduced by ~15% on average over the 17-year time period. This suggests that the urban/rural interface of FNRs 383 is becoming less drastic and NO_x -limited O_3 production regimes that in the past were predominantly observed in rural regions have 384 expanded into the urban regions of larger cities. More accurate assessment of the actual threshold ratio values separating the 385 different O₃ production regimes would allow for the determination of exactly what extent of each city has in fact transitioned to 386 NO_x-limited regimes. Overall, Fig. 7 demonstrates that the long-term record of OMI observations can observe the impact of global

emissions reduction strategies on air quality and O₃ sensitivity regimes throughout the Northern Hemisphere.

388 3.5 Impact of the COVID-lockdown on FNRs in the Northern Hemisphere

389 The global impacts of the COVID-lockdown in 2020 on atmospheric pollution, such as the reduction of tropospheric NO₂, has 390 been well documented to have impacted O_3 sensitivity regimes in the PBL and mid- to upper-troposphere (e.g., Goldberg et al., 391 2020; He et al., 2020, Cooper et al., 2022; Nussbaumer et al., 2022). Here we studied, for the first time, OMI derived VCD FNRs 392 to evaluate the impact of the COVID-lockdown on summer-mean FNRs in 2020 throughout the Northern Hemisphere compared 393 to the year before (2019) and after (2021). Figure 8 shows the changes in OMI FNRs before, during, and after (2019, 2020, 2021) 394 the COVID-lockdown over the selected 46 cities discussed in this study. Out of the 46 selected cities, 32 of the urban regions 395 (~70%) experienced higher FNRs in 2020 compared to 2019. On average, the cities that experienced increased FNRs in 2020 had 396 values which were ~19% higher compared to 2019. Similarly, 26 of the urban regions (~57%) experienced higher FNRs in 2020 397 compared to 2021 and these city centers had FNR values ~18% larger. OMI also retrieved increased FNR values in rural regions 398 surrounding city centers throughout the Northern Hemisphere during the COVID-lockdown period of 2020 (see Fig. 8). A similar 399 number of city's rural areas experienced increased FNRs in 2020 compared to 2019 and 2021 as what was observed for urban 400 areas. The increases in FNRs for rural regions in 2020 compared to 2019 and 2021 were 16% and 13%. The OMI data evaluated 401 here suggest that the majority of cities in the Northern Hemisphere, and surrounding rural regions, tended to have O₃ production 402 which was more sensitivity to NO_x emissions/concentrations in 2020 compared to the year before and after. Cooper et al. (2022) 403 demonstrated that in 2020 NO₂ concentrations were on average ~30% lower during COVID-lockdown periods and these reductions 404 were from decreased anthropogenic emissions and can't be explained by meteorological differences. The degree of reduction in 405 NO₂ determined in Cooper et al. (2022) agrees well with the OMI VCD FNR increases determined during our study of ~20%.

406 3.6 Comparison of OMI and TROPOMI FNR spatiotemporal variability in US cities

407 To expand upon previous studies which investigated OMI FNR trends published prior to the availability of TROPOMI retrievals 408 (e.g., Mahajan et al., 2015; Jin and Holloway, 2015; Souri et al., 2017; Jin et al., 2017, 2020), here we compare the ability of OMI 409 and TROPOMI to reproduced inter-city and interannual FNR variability in the US measured by EPA AQS sites in the 7 major US 410 cities illustrated in Fig. 2. For this purpose, we applied TROPOMI L2 HCHO version 2.1 and NO₂ version 3.6.2 interpolated to a 411 standardized $0.1^{\circ} \times 0.1^{\circ}$ grid format. Figure 9 shows the normalized summer mean FNRs (city-specific annual FNR values 412 normalized by the 7-city FNR mean) for the 7 selected US cities for 2018 and 2019. For both years, TROPOMI was able to 413 reproduce the inter-city variability in normalized FNRs more accurately compared to OMI for 5 of the 7 US cities. This is further 414 emphasized by the fact that TROPOMI reproduced 48% and 93% of the inter-city FNR variance (R²) measured by AQS data for 415 2018 and 2019, respectively, while OMI only reproduced ~30% of the FNR variability measured in both years. Furthermore, 416 TROPOMI accurately reproduced the direction of change in AQS measured FNRs between 2018 and 2019 for 6 of the 7 cities 417 (85%) while OMI was only able to reproduce the FNR differences for 3 of the 7 cities (43%). The improved capability of 418 TROPOMI to capture spatiotemporal FNR variability compared to OMI is to be expected as recent studies have demonstrated 419 improved HCHO and NO₂ retrievals from the newer and higher spatial resolution sensor (e.g., Souri et al., 2023a; Johnson et al., 420 2023) and OMI is far past the expected lifetime of the sensor. Future studies should intercompare the two sensor's retrievals of 421 FNRs for the entire lifetime of TROPOMI which overlaps with OMI (2017-present) to fully understand the improvements when 422 applying TROPOMI.

423 4. Discussing the use of OMI data to assess O₃ production sensitivity regimes

424 The OMI satellite sensor offers a continuous data record across the globe with sufficient spatiotemporal resolution to assess 425 tropospheric O₃ production sensitivity which cannot be achieved with in situ observations. The analysis in this study demonstrated 426 that the 17-year record of OMI-retrieved HCHO and NO₂ data offered an unprecedented opportunity to assess the long-term 427 evolution of VCD, and likely surface-level, FNR values, with potential future applications in linking these ratio changes with 428 changes in surface O₃ regimes. Here we show that OMI VCD data of FNRs replicate the trends observed with surface in situ 429 information. In order to produce actual satellite-derived surface values of HCHO, NO₂, and FNRs using VCD retrievals requires 430 algorithms which largely depend on CTM-predicted vertical distributions of these trace gases (e.g. Zhu et al., 2017a; Jin et al., 431 2017; Cooper et al., 2020). Surface-based and aircraft in situ observations are also used for this purpose; however, these 432 observations have minimal observational coverage due to being very spatiotemporally limited (e.g., Souri et al., 2023a). Using 433 model simulations or in situ data to convert satellite VCD information to surface-level HCHO, NO₂, and FNRs concentrations are 434 both inhibited by errors. In situ observations are too sparse and CTMs have system-specific errors/biases and differ largely in their 435 prediction of HCHO and NO₂ vertical distributions (Lamsal et al., 2008; Geddes et al., 2016; Souri et al., 2023b). This results in 436 large uncertainties in surface-level FNRs when convolving satellite VCDs. Our study does not address the conversion of OMI 437 VCDs to surface-level values, but clearly shows that this spaceborne sensor can capture the trends in surface-level FNRs. The 438 ability of VCD information from low earth orbit satellites to capture mid-day surface-level FNR trends might be due to studies 439 showing ratios of VCD FNRs to PBL/surface values are near unity (Jin et al., 2017; Souri et al., 2023a). However, during times 440 where HCHO and NO₂ vertical profiles in the troposphere are not similar to climatological averages, models and satellites may be 441 challenged to accurately assess conversion ratios of VCD FNRs to PBL/surface values (e.g., Souri et al., 2023b). Overall, if 442 methods are improved to reduce the uncertainty in the conversion of satellite VCD retrievals of HCHO and NO₂ to surface-level 443 concentrations, then satellite-derived surface-level FNRs will be able to be applied in higher confidence for air quality research 444 and potentially policy decisions.

445 OMI VCD FNR and NO₂ retrievals display high correlation with surface in situ data trends (see Table 1). However, this 446 satellite demonstrated less capability to replicate trends of surface-level observations of HCHO. The vertical structure of HCHO 447 can be complex which complicates the relationship between VCDs and surface-level values (Souri et al., 2023b); however, OMI 448 has also been shown to have large systematic and random biases in HCHO retrievals which drive the overall errors in OMI-derived 449 VCD FNRs (Johnson et al., 2023; Souri et al., 2023a). However, since decreasing NO₂ emissions/concentrations driven by NO_x 450 emission control strategies is the primary reason for the increasing trends of FNRs at the surface, while HCHO has near-neutral 451 trends, and OMI NO₂ VCDs have much lower errors compared to HCHO (Johnson et al., 2023; Souri et al., 2023a), this study 452 shows that OMI VCD data of indicator species can still replicate surface-level trends of FNRs. Emission control strategies for 453 VOCs have also been shown to have caused regional reductions in the concentrations of these compounds; however, it is 454 challenging to derive and assess the impact of VOC emission control strategies as there are thousands of different VOC compounds 455 all with different chemical reactivity (Pei et al., 2022). Furthermore, a large fraction of VOCs is emitted from biogenic sources 456 which cannot be controlled through changes in human activities (Guenther et al., 1995). In addition to retrieval errors, the coarse 457 spatial resolution of OMI and other spaceborne sensors results in representation errors when compared to point-source surface 458 observations (Souri et al., 2022). This also likely contributes to the challenge of satellite-derived HCHO, NO₂, and FNRs to 459 replicate trends and variability determined from in situ measurements located in city centers.

Besides the southeast US and some small areas of eastern China, OMI HCHO version 3 collection 3 data results in mostly
positive trends between 2005-2021. A portion of this positive trend could be due to the OMI instrument drift which has been
identified in past studies (e.g., Marais et al., 2012; Zhu et al., 2016, 2017b). Currently, there is an OMI HCHO version 3 collection

463 4 product in development using the SAO algorithm; however, has yet to be published and therefore could not be applied in this 464 study. A major difference between the collection 3 and 4 data products is how level 1b (L1b) data is produced as described in 465 Kleipool et al. (2022). Furthermore, changes in molecular absorption cross-sections and input parameters for SCD calculations, 466 AMF calculations, and latitudinal bias corrections are applied in collection 4 OMI HCHO data. Preliminary analysis of the 467 differences in OMI VCD HCHO using the collection 3 and 4 data demonstrates changes in the magnitudes and trends of this 468 species (personal communication with the SAO OMI HCHO algorithm team). These differences could have an impact on the 469 analysis of trends in global FNRs such as that conducted in this study. Once the OMI HCHO version 3 collection 4 product becomes 470 available to the public the analysis in this study should be redone with the updated HCHO data. Overall, we don't expect that the 471 results in FNR trends will be impacted much due to observed trends in OMI VCD and surface-level NO₂ being much larger than 472 that in HCHO (see Sect. 3.2).

473 Due to the inadequacy of our current quantitative understanding of exact threshold FNR values marking the O_3 474 photochemical regime transitions (e.g., Schroeder et al., 2017), this study avoided explicitly linking OMI FNRs with exact chemical 475 regimes. An accurate diagnosis of surface O_3 sensitivity requires more in-depth analysis of satellite FNRs at a higher spatiotemporal 476 resolution, and accurately relating these FNRs to O_3 regimes by estimating the threshold ratio values applicable to specific regions 477 and time periods. Global CTM simulations (Jin et al., 2017), photochemical box modeling utilizing measurement data from 478 airborne field campaigns (Schroeder et al., 2017; Souri et al., 2020), and observation-based methods linking ratio values with 479 surface O_3 concentrations (Jin et al., 2020; Wang et al., 2021) could lead to the derivation of more accurate regime threshold ratio 480 values. Nevertheless, the OMI-derived VCD FNR values investigated in this study revealed many aspects of O_3 sensitivity to NO_x 481 versus VOCs. In general, OMI-derived summer FNR values indicated radical limited regimes within many cities in the Northern 482 Hemisphere (FNRs < 2) and NO_x-limited regimes over the rural regions around those cities (FNRs > 3). The analysis of multi-year 483 summer mean OMI HCHO and NO₂ values revealed a positive trend in FNRs indicating a transition from radical-limited to NO_x-484 limited regimes, especially during more recent years. The positive trend in OMI FNRs over most cities are mainly due to decreases 485 in NO₂ resulting from the decrease in anthropogenic NO_x emissions and mixed-variations in biogenic VOC sources. This study 486 evaluated FNR trends for 46 large Northern Hemisphere cities which expands on other recent studies which evaluated O_3 487 production sensitivity regimes around smaller numbers of cities (e.g., Jin et al., 2017). Due to the majority of highly-populated 488 Northern Hemisphere cities, outside of the Tropics, residing in developed nations, the increasing trend in FNRs due to 489 anthropogenic emissions reductions holds true for the vast majority of the 46 cities studied here.

490 In the earliest years of studying satellite-derived FNRs it was hoped this data source could potentially be used for policy 491 decisions and developing emission control strategies (Martin et al., 2004). However, more recent studies starting with Duncan et 492 al. (2010) suggested that satellite retrievals may have errors too large for applying FNRs for air quality regulations. Furthermore, 493 satellite data typically has to be temporally averaged to reduce noise in the retrievals which may mask out important O₃ exceedance 494 events and the indicator species characteristics on these days (Schroeder et al., 2017). The recent study by Souri et al. (2023a) 495 compiled a comprehensive error budget for using satellite retrievals to assess surface-level FNRs. This study showed that total 496 relative error in satellite FNRs over large cities tend to be ~50% whereas over rural regions there are much larger errors (>100%). 497 The majority of this error comes from noise in satellite retrievals (40%-90%), especially from HCHO retrievals, and the rest of 498 this error is associated with the ability of indicator species to accurately describe complex O_3 chemistry (~20%), VCD to surface 499 translation (~19%), and spatial representation (~13%). These total relative errors are likely too large to apply satellite FNRs for air 500 quality regulation purposes; however, still provide a useful scientific research product for investigating long-term, and short-term 501 events (e.g., meteorological variations, droughts/floods, wildfires, socioeconomic events, etc.), impacts of emissions on O_3 502 production sensitivity regimes.

503 4 Conclusions

504 This study applied the 17-year data record of OMI satellite sensor's summer mean VCD HCHO, NO₂, and FNRs between 2005 505 and 2021 over the Northern Hemisphere to understand the long-term evolution of O₃ photochemical regimes. This expands the 506 global OMI record of VCD FNRs out to 2021 further than previous studies (e.g., Jin et al., 2020). The long-term trends for 46 507 highly populated cities in the Northern Hemisphere agree with past work which have shown that FNRs are primarily increasing 508 due to reductions in emissions/concentrations of NO_x (Duncan et al., 2010; Jin et al., 2017, 2020). OMI VCD NO₂ data are 509 decreasing in most urban regions of the Northern Hemisphere, while HCHO data is near-neutral or slightly increasing, resulting in 510 the increasing FNR trends. The extension of OMI FNR data out to 2021 suggests a continuing trend towards more NO_x-limited O₃ 511 production sensitivity regimes within and around cities throughout the Northern Hemisphere.

512 Another unique finding in our study is the extension of OMI FNR data out to 2021 covering the impact of the COVID-513 19 lockdown of 2020. Out of the 46 selected cities, ~70% of urban regions experienced higher FNRs in 2020 compared to 2019 514 and ~57% had higher FNRs in 2020 compared to 2021. OMI FNRs were 18%-19% higher in 2020 compared to the year before 515 and after the COVID-lockdown in 2020. We studied summer-mean FNRs in this study; however, COVID-lockdown restrictions 516 were largest in the spring of 2020, thus the full impact of COVID-lockdown restrictions on VCD FNRs was likely larger than that 517 experienced in the summer. A similar percentage of rural areas around the 46 selected cities experience higher FNRs in 2020 518 compared to 2019 and 2021; however, the increases in FNRs were smaller (13%-16%) compared to urban areas. The OMI data 519 evaluated here suggests that the majority of cities in the Northern Hemisphere, and surrounding rural regions, tended to have O_3 520 production which was more sensitive to NO_x emissions/concentrations in 2020 compared to the year before and after.

521 Past studies have shown that mid-day FNR VCDs are similar to those observed at the surface (Jin et al., 2017; Souri et 522 al., 2023a). However, during times where HCHO and NO₂ vertical profiles in the troposphere are complex, models and satellites 523 may be challenged to accurately assess conversion ratios of VCD FNRs to PBL/surface values (e.g., Souri et al., 2023b). This 524 study shows that on average the long-term trends of OMI VCD FNRs agree well with in observations at the surface in cities 525 distributed around the US, suggesting that this satellite is capable to assess the long-term trends of surface-level O₃ production 526 sensitivity regimes. However, the magnitudes of both indicator species calculated with satellite VCDs using scaling factors derived 527 with CTMs and/or in situ observations are highly uncertain. Both OMI VCD and surface in situ data of HCHO, NO₂, and FNRs 528 emphasize that the increasing trend in FNRs is driven by reduced emissions/concentrations of NO2 while HCHO has a near-neutral 529 trend. While OMI VCD HCHO trends and variability do not agree well with surface in situ observations; OMI does replicate the 530 strong decreasing trend of NO₂ observed at the surface resulting in the agreement between OMI and surface data of FNR trends.

531 Higher spatiotemporal retrievals from newer low earth orbit (e.g., TROPOMI) and geostationary (e.g., Tropospheric 532 Emissions: Monitoring of Pollution [TEMPO], Geostationary Environment Monitoring Spectrometer [GEMS]) satellite sensors 533 provide more insight into the short-term (daily, diurnal) recent (2017-present) evolution of O₃ photochemical regimes. Compared 534 to OMI, TROPOMI was shown to retrieve VCD FNR values with more accuracy and better precision compared to OMI 535 observations primarily due to improvements in HCHO product performance (e.g., Johnson et al., 2023). Johnson et al. (2023) 536 demonstrated that TROPOMI can retrieve spatiotemporal HCHO variability with uncertainties low enough to capture FNR 537 variability on a daily basis, while OMI was not. The current study compared the capability of OMI and TROPOMI to capture inter-538 city and interannual FNR variability measured by ground-based AQS data (using normalized FNR values) for 7 select US cities. 539 Here we quantitatively showed that TROPOMI was able to reproduce the spatiotemporal variability in observed FNRs more 540 accurately compared to OMI for the reasons mentioned above. Future improvements in satellite HCHO retrievals will allow for 541 more accurate retrievals of FNRs on a daily- to monthly-scale. TEMPO and GEMS provide HCHO and NO₂ VCD information at 542 1- to 3-hour temporal resolution and higher spatial resolution compared to both OMI and TROPOMI which will allow for the

543 assessment of diurnal FNR variability on a regional scale. This has not yet been possible as TEMPO and GEMS are the first 544 UV/VIS spectrometers on geostationary platforms with spatial resolution high enough to retrieve air quality relevant HCHO and 545 NO₂ VCD data. The new diurnal and high spatial resolution information from these geostationary satellites are expected to greatly 546 improve the understanding of FNRs. As OMI is set to be decommissioned in the coming years, it is critical to merge TROPOMI 547 HCHO and NO₂ VCD data with OMI in order to continue the long-term data set from 2005-present. Furthermore, combining 548 retrieved information from geostationary satellites with once-a-day low earth orbit data will provide a vast wealth of information 549 about global daily to hourly variability in FNRs.

550 While recent studies have shown that OMI FNR retrieval errors are likely too large to apply in air quality regulation and 551 for deriving emission control strategies to reduce surface-level O₃ concentrations, this long-term satellite product provides a useful 552 scientific research product for investigating atmospheric O₃ chemistry and investigating qualitative impacts of emission changes 553 on O₃ production sensitivity regimes. This is especially true for regions of the globe outside of the US and Europe that have limited 554 long-term surface in situ observation networks able to measure HCHO and NO₂ concentrations. Improvements in HCHO and NO₂ 555 VCD retrieval algorithms and methods to derive VCD to surface/PBL conversion factors for these indicator species would greatly 556 improve the ability to apply OMI and other satellite products to study surface air quality. Data assimilation and inverse models 557 have been combined with satellite retrievals of HCHO and NO₂ data to constrain predictions of NO_x and VOC emissions and 558 resulting O_3 chemistry (e.g., Souri et al., 2020). These satellite data-constrained models can then be used to assess trends and 559 variability in FNRs, indicator species emissions, and O_3 photochemistry regimes. These improvements in satellite retrieval 560 algorithms, CTMs, data assimilation and inverse modeling techniques, along with studies to better define the actual O₃ production 561 sensitivity regime thresholds, will allow for a more confident investigation of long-term air quality and the impacts of NO_x and 562 VOC emission changes on O₃ production sensitivity.

563 Code and Data Availability. The OMI HCHO L3 data used in this paper is publicly available at 564 https://acdisc.gesdisc.eosdis.nasa.gov/data//Aura_OMI_Level3/OMHCHOd.003/ (last access: 24 October 2021). The OMI NO2 565 High Resolution L3 data is also available at а public data repository: 566 https://avdc.gsfc.nasa.gov/pub/data/satellite/Aura/OMI/V03/L3/OMNO2d_HR/OMNO2d_HRD/ (last access: 3 December 2021). 567 The CEDS emission inventory used in this work is also publicly available (https://data.pnnl.gov/group/nodes/dataset/13488) (last 568 access: July 22, 2023). EPA AQS data of HCHO and NO₂ can be downloaded from: https://www.epa.gov/outdoor-air-quality-569 data/download-daily-data (last access: August 3, 2023).

570 *Supplement*. The supplement related to this article is available.

571 *Author Contributions.* MJ and SP obtained the funding for this project. MJ, SP, and RK were fundamental in developing the 572 investigation strategy of this study. SJ, SP, SM, S M-H, and YS conducted the analysis which produce the results presented in this 573 manuscript. Finally, MJ and SP were the primary authors which wrote the text of the manuscript.

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

Acknowledgements. Computational resources were provided by the NASA High-End Computing Program through the NASA
 Advanced Supercomputing Division at NASA Ames Research Center. We acknowledge the United States Environmental

- 577 Protection Agency for the free availability of in situ data. The views, opinions and findings of this paper are those of the authors
- 578 and should not be construed as an official NASA or United States Government position, policy, or decision.
- 579 Financial support. The MJ, SP, SM, RK, S M-H, and YS acknowledge funding support from the NASA Earth Science Division's
- 580 Aura Science Team (NNH19ZDA001N-AURAST) as part of the Upper Atmosphere Research Program (UARP). A part of this
- 581 material is based upon work supported by the NSF National Center for Atmospheric Research, which is a major facility sponsored
- by the U.S. National Science Foundation under Cooperative Agreement No. 1852977.

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860 Tables

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Table 1. Statistics of the correlation of OMI and AQS ("Obs.") normalized trends for HCHO, NO₂, and FNRs for major cities in the US and the average of all cities in the US (USA urban areas) between 2005-2019. Slopes of the trends for each species (units of yr⁻¹) are also provided. The values in italic font are linear regression slopes which are statistically significant at a 95% confidence level ($p \le 0.05$).

Cities	Corr. HCHO	Slope HCHO Obs.	Slope HCHO OMI	Corr. NO ₂	Slope NO ₂ Obs.	Slope NO ₂ OMI	Corr. FNR	Slope FNR Obs.	Slope FNR OMI
New York	0.01	0.10	0.05	0.91	-0.21	-0.20	0.78	0.20	0.20
Los Angeles	-0.01	-0.14	0.14	0.98	-0.21	-0.22	0.84	0.17	0.22
Chicago	NaN	NaN	0.07	0.91	-0.22	-0.21	NaN	NaN	0.18
Washington DC	0.38	0.11	0.05	0.89	-0.19	-0.20	0.77	0.18	0.21
Pittsburgh	0.51	0.06	0.10	0.64	-0.21	-0.16	0.55	0.22	0.16
Atlanta	NaN	NaN	-0.07	0.33	-0.01	-0.17	NaN	NaN	0.18
Houston	0.40	-0.18	-0.06	0.79	-0.17	-0.17	-0.27	-0.12	0.13
USA urban areas	0.24	-0.05	0.15	0.98	-0.21	-0.20	0.91	0.21	0.21

866 Correlation values are the correlation coefficient (R).

867 NaN values indicate cities where particular species data is not available for all years between 2005-2019.



870 871 Figure 1: OMI-derived multi-year (2005-2021) summer mean (June-August) HCHO VCDs (top row), NO2 VCDs (middle row), and resulting VCD FNRs at 0.1° × 0.1° latitude × longitude grid cells. Values of FNRs are displayed only for polluted 872 873 regions (NO₂ VCD > 1.2×10^{15} molecule cm⁻²). The white color indicates data gaps or oceanic grid cells.



Figure 2: Normalized time series (values from 2005 to 2019 normalized to 2005-2019 mean) of summer mean OMI HCHO
and NO₂ VCDs and column FNRs (solid lines). The same information is shown for surface concentrations from the EPAAQS in situ observations (dashed lines) over select cities and over all urban monitoring sites in the United States (bottom
right panel).



Figure 3: OMI-derived trends in summer mean (June-August) time series of HCHO (top row) and NO₂ (middle row) VCDs (units in molecule cm⁻² yr⁻¹), and corresponding FNR values (bottom row; unitless yr⁻¹) at $0.1^{\circ} \times 0.1^{\circ}$ latitude × longitude grid cells between 2005 and 2021. Values in the bottom row are displayed only for polluted regions (OMI NO₂ VCD > 1.2 × 10¹⁵ molecule cm⁻²). The white color indicates data gaps or oceanic grid cells. All trend values that are displayed are at an 85% confidence level (p ≤ 0.15) for better visualization of spatial trend variability. Figure S1 shows the trend values at 99% confidence level and for all grid cells.



Figure 4: Time series of OMI-derived summer mean (June-August) FNR VCD values for 18 selected cities across the North
Hemisphere from 2005 to 2021. The different colors illustrate mean FNR values for urban (red) and rural areas around
each city (blue). Grey shaded areas represent the transition zone of ozone production sensitivity regime threshold values
as suggested by Jin et al. (2020) (cities in United States: Los Angeles, New York, Chicago, Washington DC, Pittsburgh,
Atlanta and Houston), Wang et al. (2021) (cities in China: Beijing, Shanghai, Guangzhou, Neijiang and Jinan), and Duncan
et al. (2010) (other cities). For interpretation, FNR values that are less than the transition zone have O₃ production which
is VOC-limited and FNR values larger than the transition zone have O₃ production which is NO_x-limited.



899Figure 5: Time series of normalized OMI-derived summer mean (June-August) VCD NO2 and FNR trend values and900corresponding trends in anthropogenic emission of NOx from the CEDS bottom-up inventory over the selected 18 cities901across the Northern Hemisphere from 2005 to 2021. CEDS emissions data is only displayed until 2019 due to this being the902most recent year of availability.



Figure 6: OMI-derived summer mean (June-August) FNR VCD values for 18 selected cities across the Northern Hemisphere during 2005-2010 and 2016-2021. The black circle represents each city center. CGLC-MODIS-LCZ urban grids for each city are displayed in Fig. S10. Grey color indicates data gaps or oceanic grid cells.



909HCHO/NO2HCHO/NO2910Figure 7: OMI-derived summer-mean (June-August) FNR VCD values for a select 46 cities across the Northern911Hemisphere during 2005-2010 (blue), 2011-2015 (red) and 2016-2021 (orange). Each column represents mean ratio values912for urban city (left column) and the surrounding rural areas (right column).



914HCHO/NO2HCHO/NO2915Figure 8: OMI-derived summer-mean (June-August) FNR VCD values for a select 46 cities across the Northern916Hemisphere during 2019 (blue), 2020 (red) and 2021 (orange). Each column represents mean FNR values for urban city

917 areas (left column) and the surrounding rural regions (right column).



Figure 9: Normalized OMI and TROPOMI summer mean FNRs (each city FNR normalized by the 7-city mean) for each of the 7
 selected US cities for 2018 (top) and 2019 (bottom). The same information is shown for surface concentrations from the EPA-

921 selected US cities for 2018 (top) and 2019 (botted922 AQS in situ observations over the select cities.