

Evaluating a Space-Based Indicator of Surface Ozone-NO

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2	Evaluating a space-based indicator of surface ozone-NO _x -VOC sensitivity over mid-
3	latitude source regions and application to decadal trends
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23 Abstract

24 Determining effective strategies for mitigating surface ozone (O₃) pollution requires 25 knowledge of the relative ambient concentrations of its precursors, NO_x and VOCs. The spacebased tropospheric column ratio of formaldehyde to NO₂ (FNR) has been used as an indicator to 26 27 identify NO_x-limited versus NO_x-saturated O₃ formation regimes. However, quantitative use of 28 this indicator ratio is subject to three major uncertainties: 1) the split between NO_x-limited and 29 NO_x-saturated conditions may shift; 2) the ratio of the vertically integrated column may not 30 represent the near-surface environment; 3) satellite products contain errors. We use the GEOS-31 Chem global chemical transport model to evaluate the quantitative utility of FNR observed from 32 the Ozone Monitoring Instrument over three northern mid-latitude source regions. We find that 33 FNR in the model surface layer is a robust predictor of the simulated near-surface O₃ production 34 regime. Extending this surface-based predictor to a column-based FNR requires accounting for 35 differences in the HCHO and NO₂ vertical profiles. We compare four combinations of two OMI 36 HCHO and NO_2 retrievals with modeled FNR. The spatial and temporal correlations between the 37 modeled and satellite-derived FNR vary with the choice of NO₂ product, while the mean offset 38 depends on the choice of HCHO product. Space-based FNR indicates that the spring transition to 39 NO_x-limited regimes has shifted at least a month earlier over major cities (e.g. New York, 40 London, Seoul) between 2005 and 2015. This increase in NO_x sensitivity implies that NO_x 41 emission controls will improve O_3 air quality more now than it would have a decade ago.

42 1 Introduction

43 Surface ozone (O_3) , the main component of photochemical smog, has adverse effects on 44 public health [Kampa and Castanas, 2008], agriculture [Van Dingenen et al., 2009] and ecosystems [Yue and Unger, 2014]. The global premature mortality rate due to O₃ pollution is 45 46 estimated at 0.8 million/year [Lelieveld et al., 2013]. Surface O3 formation in urban areas is non-47 linearly dependent on the availability of two classes of O_3 precursors: oxides of nitrogen (NO_x) 48 and volatile organic compounds (VOCs). That is, depending on local relative abundances of NO_x 49 to VOCs, O_3 formation can be mitigated by reducing NO_x emissions (NO_x-limited regime), or by 50 reducing VOC emissions (NO_x-saturated or VOC-limited or radical-limited regime). At regional 51 and global scales, O₃ production is largely NO_x-limited, though urban areas with high NO_x 52 emissions are frequently NO_x-saturated.

53 The non-linear dependence of surface O_3 on precursor emissions poses challenges to 54 effective mitigation of surface O_3 . Simon et al. [2015] find that U.S. summertime O_3 decreases 55 with its precursor emissions in recent decades, but wintertime O₃ increases in urban areas as NO titration declines. Urban areas with NOx-saturated O3 production chemistry should be 56 57 transitioning to NO_x-limited chemistry following the substantial nationwide NO_x emission 58 reductions implemented since the late 1990s [Pusede et al., 2015]. NO_x emissions decreased by 59 27% over Europe in the past decade, and the overall O₃ distribution narrowed [Guerreiro et al., 2014; Lefohn et al., 2017]. In China, controls on anthropogenic NO_x emissions are being 60 implemented [Gu et al., 2013; Liu et al., 2016; Souri et al., 2017], but surface O₃ may increase 61 62 due to the dominance of VOC-limited ozone formation regimes [Liu et al., 2013; Jin and 63 Holloway, 2015; Lefohn et al., 2017].

64 O₃ sensitivity to precursor emissions has been derived from models using various approaches including emission perturbation simulations [Jacob et al., 1995; Wu et al., 2009; 65 66 Tonnesen and Dennis, 2000], O₃ source apportionment [Dunker et al., 2002; Cohan et al., 2005; 67 Li et al., 2012], and adjoint modeling [Schmidt et al., 2003; Hakami et al., 2006; Zhang et al., 68 2009]. Model uncertainties, including the possibility of compensating errors, could lead to erroneous estimates of O₃ sensitivity despite accurate simulation of O₃ concentrations [Sillman, 69 70 1995; Tonnesen and Dennis, 2000]. Furthermore, the sensitivity is non-linearly dependent on the 71 magnitude of the emission perturbation [Wu et al., 2009; Fu et al., 2012].

Sillman [1995] showed that the relationships of NO_y to O₃, H₂O₂ to HNO₃, and HCHO to 72 NO_v reflect the processes that determine the non-linear sensitivity of O₃ to VOC and NO_x 73 74 precursor emissions, which has been further examined in models and measurements [Jacob et 75 al., 1995; Tonnesen and Dennis, 2000; Hammer, 2002; Stein et al., 2005]. The relative ambient 76 concentrations of HCHO and NO_v or NO₂ reflect the reactivity-weighted concentrations of VOC 77 and NO_x , respectively, and thus indicate how O_3 will respond to changes in NO_x and VOC 78 emissions [Sillman, 1995; Valin et al., 2016]. Tonnesen and Dennis [2000] suggest HCHO/NO2 79 is more useful than HCHO/NO_v because both HCHO and NO₂ have short lifetimes (~ hours), 80 and their ratio better represents the competition between OH reaction with VOC versus NO₂. 81 Martin et al. [2004a] first applied the indicator ratio to Global Ozone Monitoring 82 Experiment (GOME) retrievals of tropospheric columns of HCHO and NO₂ with a spatial

resolution of 80×40 km², and proposed that the transition from NO_x-saturated to NO_x-limited 83 84 occurs when HCHO/NO₂ equals 1, thereby diagnosing the O₃-NO_x-VOC sensitivity across the globe from space. This work has been refined and extended to Ozone Monitoring Instrument 85 86 (OMI) products to characterize O₃ sensitivity over the U.S.A. [*Choi et al.*, 2012; *Duncan et al.*, 87 2010; Chang et al., 2016] and East Asia [Jin and Holloway, 2015; Souri et al., 2017]. The finer spatial resolution of OMI (up to 13×24 km²) better captures the urban-rural gradient of O₃ 88 89 sensitivity. In addition, the OMI overpass time (~1:45 PM) is better suited to detect the 90 sensitivity of ozone production during the afternoon, when O_3 photochemical production peaks, 91 and when the boundary layer is deepest and the solar zenith angle is small, maximizing 92 instrument sensitivity to HCHO and NO₂ in the lower troposphere.

93 Table 1 summarizes previous studies that use HCHO/NO_v or HCHO/NO₂ as indicators for O₃-NO_x-VOC sensitivity. While previous studies have demonstrated the potential of the 94 95 space-based indicator ratio to identify the O_3 sensitivity to NO_x vs. VOC emission controls, the 96 quantitative application of space-based HCHO/NO₂ is subject to three major uncertainties. First, 97 different mechanisms and meteorological conditions, such as humidity, temperature and dry 98 deposition rates can affect the relationship of O₃ production to HCHO/NO₂ [Sillman, 2002; 99 Vogel et al., 2007; Liu et al., 2010]. Second, satellite observations measure the vertically 100 integrated column density, which differs from the mixing ratio near the surface, of most 101 relevance to air quality management. Variations in the vertical distribution of HCHO relative to 102 that of NO₂ also alter the relationship between the column and surface ratios [Martin et al., 103 2004b]. Third, even if column-based HCHO/NO₂ is a useful indicator of surface O₃ sensitivity, 104 satellite retrievals are subject to large uncertainties from measurement errors, surface reflectivity, 105 cloud effects, profile shape and aerosol effects [Boersma et al., 2004; Lin et al., 2014]. Duncan 106 et al. [2010] and Martin et al. [2004b] derive the ozone production regime thresholds (i.e., the 107 range of values over which the transition occurs from NO_x-saturated to NO_x-limited) from 108 modeled column densities, assuming that modeled column densities match what is retrieved from 109 space. Inter-model comparison of indicator ratios, however, shows large disagreements between 110 satellite products and models [Campbell et al., 2015]. Zhu et al. [2016] suggest HCHO satellite 111 retrievals are biased low relative to aircraft data, with the extent of this underestimate varying by 112 product. If HCHO is biased low, the extent of VOC-limited regimes will be overestimated.

113 We investigate these uncertainties by first evaluating the quantitative utility of the 114 indicator ratio HCHO/NO₂ (hereafter FNR) observed from OMI over three mid-latitude source 115 regions: North America (22 °N - 50 °N, 75 °W - 120 °W), Europe (35 °N - 60 °N, 10 °W - 30 116 °E) and East Asia (20 °N - 50 °N, 100 °E - 140 °E). Relative to a multi-year (2006 - 2012) base-117 case GEOS-Chem simulation, we conduct two perturbation simulations that separately reduce 118 NO_x and VOC emissions globally by 20% to examine the ability of FNR to detect the surface O_3 119 sensitivity to precursor emissions (Section 3.1). Using the 3-D distribution of NO₂ and HCHO 120 archived from GEOS-Chem, we examine the surface-to-column relationships of FNR and their 121 spatial and temporal variations (Section 3.2). The model-derived surface-to-column relationships 122 are then applied to determine the column-based regime threshold values. We then compare four 123 combinations of two OMI HCHO products and two OMI NO₂ products with the GEOS-Chem 124 simulations (Section 3.3). Finally, we investigate decadal trends in surface O₃ sensitivity over 125 northern mid-latitude polluted regions from 2005 to 2015 using the fine OMI products with 0.25° 126 resolution (Section 4).

127 **2** Data and Methods

128 2.1 OMI Products

OMI is on board the NASA EOS Aura satellite at ~705 km altitude in a sun-synchronous orbit with 98° inclination [*Levelt et al.*, 2006]. OMI is a nadir-viewing UV/visible spectrometer, providing daily, near global coverage with a local equator crossing time of ~1:45 PM. OMI covers two UV region (264 - 311 nm and 307 - 383 nm) and one VIS region (349 - 504 nm) with a spectral resolution between 0.42 to 0.63 nm and a spatial resolution of up to 13×24 km² at nadir [*Levelt et al.*, 2006].

135 2.1.1 OMI Tropospheric NO₂

We use two Level-2 OMI NO₂ satellite retrieval products: the Level-2 standard operational NO₂ Product (OMNO₂ SP, version 2.1) developed at NASA/Goddard Space Flight Center [*Bucsela et al.*, 2013], and the Dutch NO₂ product (DP) developed at KNMI, the Royal Netherlands Meteorological Institute (DOMINO DP, v2.0) [*Boersma et al.*, 2011]. Retrieval of tropospheric NO₂ column density involves three major steps: (1) spectral fitting to obtain a raw NO₂ slant column density; (2) separation of tropospheric and stratospheric columns; and (3) conversion from slant column to vertical column density. NASA SP and DOMINO DP differ in (2) and (3) [*Bucsela et al.*, 2013; *Boersma et al.*, 2011]. The air mass factor (AMF, the ratio of
the slant column to the vertical column density) can be expressed as the vertical integral of the
contribution of each layer to the column divided by the vertical column [*Boersma et al.*, 2011]:

$$AMF = \frac{\sum_{l} m_{l}(\mathbf{b}) \cdot x_{a,l}}{\sum_{l} x_{a,l}}$$
(1)

147 where $m_{\rm l}({\bf b})$ is the atmospheric scattering weight that is a function of satellite viewing geometry, 148 cloud pressure, cloud radiance fraction, surface pressure and reflectivity, and $x_{a,l}$ is the sub-149 column from the a priori profile for layer l. Scattering weights are included in the NASA Level-2 150 SP. DP provides averaging kernels (AK) as an alternative, which is equal to $m_l(\mathbf{b})/AMF$ [Eskes 151 and Boersma, 2003]. A recent study estimates a structural uncertainty of 42% for AMF over 152 polluted regions resulting from different priori trace gas profiles, surface albedo and cloud 153 parameters applied for AMF calculation [Lorente et al., 2017]. In this study, we calculate the 154 tropospheric AMF (AMF_{trop}) consistently for both DP and SP by using 1-hour average GEOS-155 Chem modeled NO₂ profiles sampled each day at the OMI overpass time, which enables direct 156 comparison between GEOS-Chem and the OMI products [Boersma et al., 2016]. We use the 157 stratospheric NO₂ columns and AMFs provided with the data products. We calculate the NO₂ 158 tropospheric column density at each pixel as the difference between the total and the 159 stratospheric slant column density:

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$$V_{trop} = \frac{S - S_{stra}}{AMF_{trop}} \tag{2}$$

where V_{trop} is tropospheric column density; AMF_{trop} is the tropospheric AMF; S is the de-striped 161 162 total slant column density; S_{stra} is the stratospheric slant column density. For DP, we use the TM4 assimilated stratospheric slant column density (S_{stra}) included in the product. For SP, S_{stra} is 163 164 calculated as the product of the stratospheric column density (V_{strat}) and the stratospheric AMF 165 (AMF_{strat}) included in the product. We select individual observations with cloud radiance fraction 166 lower than 30%, solar zenith angle smaller than 85° , and only those unaffected by row anomalies 167 (http://www.knmi.nl/omi/research/product/rowanomaly-background.php) [Dobber et al., 2008]. The overall uncertainty of the OMI SP and DP retrievals is on the order of $\sim 10^{15}$ molecules/cm² 168 169 over polluted areas (20% - 30% of the retrieved quantity) [Boersma et al., 2011; Bucsela et al., 170 2013]. While the effects of aerosols on satellite retrievals are not included explicitly, such effects 171 are accounted for implicitly via cloud retrievals being sensitive to the scattering effects of aerosols, though such corrections may not work well for extreme aerosol loading and highly absorbing aerosol mixtures [*Lin et al.*, 2014, 2015; *Lorente et al.*, 2017]. Evaluation of the OMI SP NO₂ (version 2.1) with ground-based and aircraft data shows that OMI products generally agree with *in situ* measurements over the U.S.A. within $\pm 20\%$ [*Lamsal et al.*, 2014]. *Marchenko et al.* [2015] suggest that the OMI retrieved slant column density is overestimated by 10 to 40%; improvements in slant column density have been made to the Version 3 NASA products, but are not yet included in DP. We use SP from Version 2.1 for the sake of consistency with DP.

179 2.1.2 OMI HCHO

180 We use two Level-2 OMI HCHO retrieval products: NASA's standard product developed 181 by the Smithsonian Astrophysical Observatory (SAO) team (OMI-SAO, v3.0) [González Abad et 182 al., 2015], and the Belgian Institute for Space Aeronomy (BIRA-IASB) retrieval (OMI-BIRA, 183 v14) [De Smedt et al., 2015]. HCHO slant columns are estimated via spectral fitting in near ultra-184 violet (UV) regions. The OMI-SAO retrieval of slant columns differs from BIRA-IASB in the 185 absorption cross sections for HCHO, BrO and NO₂ [González Abad et al., 2015; De Smedt et al., 186 2015]. We convert slant columns to vertical columns ($\Omega_{HCHO-BIRA}$ and $\Omega_{HCHO-SAO}$) via the AMF 187 (Equation 1) provided with the products. For direct comparison, we use 1-hour average GEOS-188 Chem HCHO profiles sampled each day at the OMI overpass time as the a priori vertical 189 profiles. The scattering weights are based on the scalar LIDORT radiative transfer model (v3.3) 190 for OMI-BIRA, and the VLIDORT for OMI-SAO (v2.4) [Spurr, 2008]. Latitude-dependent 191 biases due to unresolved spectral interferences are pronounced for weak absorbers such as 192 HCHO. OMI-BIRA and OMI-SAO products deal with the spectral interference differently: OMI-193 BIRA product employs a two-step across-track and zonal reference sector correction to 194 normalize the HCHO slant columns [De Smedt et al., 2015], and OMI-SAO product applies a 195 post-processing normalization for the vertical column density using a model reference sector 196 over the remote Pacific Ocean [González Abad et al., 2015]. Similar to NO₂, the effects of 197 aerosol are not accounted for explicitly for both retrievals. We select observations with cloud 198 radiance fraction less than 30%, solar zenith angle smaller than 70° , and unaffected by row 199 anomalies following the criteria suggested in De Smedt et al. [2015]. The overall error of the 200 monthly average HCHO column is about 30% for both products [de Smedt et al., 2008, 2012; 201 González Abad et al., 2015].

202 2.1.3 OMI HCHO/NO₂

203 Daily Level-2 OMI NO₂ and OMI HCHO data from 1 January 2005 to 31 December 204 2012 are re-gridded to the GEOS-Chem model grid for direct comparison with the model 205 simulations. In order to reduce the random errors in the satellite retrievals, we first calculate 206 seven-day average tropospheric NO₂ and HCHO column densities ($\Omega_{GC NO2}$ and $\Omega_{GC HCHO}$). 207 Negative columns may occur as a result of minimizing residuals during the spectral fitting below 208 the satellite detection limit and are included when constructing seven-day averages [Boeke et al., 209 2011]. We calculate four combinations of the O₃ sensitivity indicator ratio (FNR) by taking the 210 ratio of seven-day average Ω_{HCHO} to Ω_{NO2} : 1) $\Omega_{\text{HCHO}_{\text{SAO}}}/\Omega_{\text{NO2}_{\text{SP}}}$ (FNR_{OMI_SS}); 2) 211 $\Omega_{\text{HCHO BIRA}}/\Omega_{\text{NO2 SP}}$ (FNR_{OMI BS}); 3) $\Omega_{\text{HCHO BIRA}}/\Omega_{\text{NO2 DP}}$ (FNR_{OMI BD}); 4) $\Omega_{\text{HCHO SAO}}/\Omega_{\text{NO2 DP}}$ 212 (FNR_{OMI SD}). The combined relative uncertainty in FNR (σ_{FNR}/FNR) can be calculated as:

213
$$\frac{\sigma_{FNR}}{FNR} = \sqrt{\left(\frac{\sigma_{NO_2}}{\Omega_{NO_2}}\right)^2 + \left(\frac{\sigma_{HCHO}}{\Omega_{HCHO}}\right)^2 - 2\left(\frac{\sigma_{NO_2,HCHO}}{\Omega_{NO_2}\Omega_{HCHO}}\right)}$$
(3)

where σ_{NO2} and σ_{HCHO} are the estimated individual errors for OMI NO₂ and HCHO, and $\sigma_{HCHO,NO2}$ is the covariance of these errors. Assuming a 20% relative uncertainty for OMI NO₂, a 30% relative uncertainty for OMI HCHO, and that the errors of the retrieved NO₂ and HCHO products are uncorrelated (i.e. $\sigma_{HCHO,NO2}$ =0), we estimate an overall FNR uncertainty of 36%. As the effects of clouds, aerosols, and albedo on satellite retrievals may cancel out, the uncertainty in FNR is expected to be lower than 36% [*Martin et al.*, 2004a; *Duncan et al.*, 2010].

220 **2.2 GEOS-Chem**

221 We use the GEOS-Chem global 3D CTM (version 9.02; http://www.geos-chem.org) to simulate O_3 -NO_x-CO-VOC-aerosol chemistry with $2^\circ \times 2.5^\circ$ resolution for 2005 to 2012. These 222 223 simulations are driven by Modern Era-Retrospective Analysis for Research and Applications 224 (MERRA) meteorology [*Rienecker et al.*, 2011]. Base anthropogenic emissions are from the 225 Emission Database for Global Atmospheric Research (EDGAR) inventory for inorganic 226 compounds [Olivier et al., 2007] and from the Reanalysis of the Tropospheric Chemical 227 Composition (RETRO) inventory for organic compounds [Schultz et al., 2007], with regional 228 overwrites for the United States (EPA National Emissions Inventory (NEI) 2005), Canada 229 (National Pollutant Release Inventory), Mexico [Kuhns et al., 2005], Europe [Auvray and Bey, 230 2005], and South and East Asia [Streets et al., 2006; Zhang et al., 2009]. Anthropogenic NO_x 231 emissions over the U.S.A., Canada, Japan and Europe are scaled each month based on estimates 232 provided by the individual countries or regions [van Donkelaar et al., 2008]. The scale factors 233 for North American are extended to 2012, and fixed after 2006 for other regions unless 234 overwritten by regional emission inventories. No inter-annual scale factors are applied to 235 anthropogenic VOCs. Additional inventories are applied for aircraft emissions [Stettler et al., 236 2011] and shipping [Vinken et al., 2011]. Monthly biomass burning emissions are from the 237 Global Fire Emissions Database version 3 [van der Werf et al., 2010]. Biogenic VOC emissions 238 follow the Model of Emissions of Gases and Aerosols from Nature scheme version 2.1 239 [Guenther et al., 2012]. Lightning NO_x emissions are as described by Murray [2016]. Soil 240 microbial NO_x emissions are described by Hudman et al. [2012]. Monthly surface methane is 241 prescribed from the NOAA GMD global surface network as a lower boundary condition 242 [Murray, 2016]. Regional monthly average NO_x (including anthropogenic, natural and total) and 243 VOC emissions (including anthropogenic, isoprene and total) are shown in Figure S1 and S2.

244 We sample model fields as one-hour averages between 1:00 and 2:00 pm local time (LT) 245 to match the OMI overpass time. To examine the response of surface O_3 to precursor emissions, 246 we conduct two perturbation simulations in GEOS-Chem that span 2006 to 2012, following a 12-247 month initialization period beginning in January 2005. First, we decrease global NO_x emissions 248 by 20%. Second, we decrease global VOC emissions (including isoprene) by 20%. We calculate 249 FNR_{GC} using the 3-D distribution of 1-2pm LT GEOS-Chem NO₂ and HCHO. We calculate the 250 area-weighted average of all individual retrievals within each model grid cell. We sample 251 modeled HCHO and NO₂ columns for the scenes concurrent with valid OMI observations to 252 avoid sampling biases [Boersma et al., 2016]. To minimize random noise, we average both 253 modeled and observed HCHO and NO₂ columns over seven days.

3 Evaluating space-based FNR as an indicator of surface O₃ sensitivity

In this section, we first evaluate the quantitative utility of FNR from a modeling perspective (Section 3.1), by correlating modeled column and surface FNR with the surface O_3 response to NO_x versus VOC emission reductions. We then examine the vertical profiles of HCHO and NO₂ in GEOS-Chem to better understand the spatial and temporal factors affecting column FNR relative to surface FNR (Section 3.2). Section 3.3 compares seven-day average OMI FNR with that simulated from GEOS-Chem.

261 **3.1 Relating FNR to surface O₃ sensitivity**

262 Previous studies characterize the transition between NO_x-sensitive and NO_x-saturated 263 ozone production in different ways, such as the response of surface O₃ to emission perturbations 264 [e.g. Martin et al., 2004a], correlations between O_3 and NO_v or NO_z [e.g. Jacob et al., 1995], or 265 radical loss pathways [e.g. Kleinman et al., 1994; Duncan et al., 2010]. Different methods may 266 identify different threshold values marking the transition between chemical production regimes. 267 Figure 1 shows the normalized surface O₃ responses to the perturbed NO_x and VOC emissions 268 change (i.e. d[O₃]/dE) in GEOS-Chem versus the surface and column FNR averaged between 1 269 to 2 PM for all polluted model grid cells within our three regions (grid cells where multi-year average $\Omega_{\rm NO2~GC} > 2.5 \times 10^{15}$ molecules/cm²). In general, the surface O₃ response to NO_x 270 271 emission reductions increases with FNR, and the surface O₃ response to VOC emission 272 reductions decreases with surface FNR. We define negative $d[O_3]/dE_{NOx}$ as NO_x-saturated (VOC-limited) conditions. In this chemical regime, reductions in NO_x emissions increase surface 273 274 O₃ due to NO titration effects and reductions in VOC emissions decrease surface O₃. NO_x-275 limited conditions occur when the surface O_3 response to NO_x emission reductions is larger than 276 that to VOC emission reductions (i.e. $d[O_3]/dE_{NOx} > d[O_3]/dE_{VOC}$). We refer to the intermediary 277 conditions as a mixed or "transitional" regime.

278 Spatial variations in meteorological and photochemical conditions, as well as in 279 downwind transport of ozone produced in upwind grid cells, can produce a range of $d[O_3]/dE_{NOx}$ 280 sensitivities for any given FNR value (Figure 1). Despite these variations, surface FNR can 281 qualitatively distinguish between NO_x-saturated and NO_x-limited conditions (Figure 1). The 282 majority (90%) of NO_x-saturated grid cells are associated with surface FNR < 0.6; over 90% of 283 NO_x -limited conditions are associated with surface FNR > 0.9. In the model, the surface FNR 284 values thus mark a clear separation between the NO_x-saturated and NO_x-limited regimes. Figure 285 2 shows the cumulative probability of correctly identifying the NO_x-limited or the NO_x-saturated 286 regime at a given FNR value in the GEOS-Chem model. The intersection of the two lines marks 287 the point at which the probability of identifying NO_x-limited and NO_x-saturated correctly is 288 equal. This intersection occurs around 0.65 for North America, 0.5 for Europe and 0.7 for East 289 Asia (Figure 2). Below this value, the likelihood of correctly identifying NO_x-limited conditions 290 increases, while the likelihood of identifying NO_x-saturated conditions decreases.

291 Instead of defining a single cut-off value between NO_x-saturated and NO_x-limited 292 conditions, we define a range of values marking a "transitional regime" to lower the probability 293 of misclassification (i.e. incorrectly classifying NO_x-saturated as NO_x-limited or vice versa). A 294 wider transitional regime lowers the chance of misclassification but generates more grid cells 295 where ozone sensitivity is regarded as mixed or uncertain. If the regime threshold values are set 296 between 0.5 and 0.8 for North America so that the probability of misclassification is 5%, then 297 10% of NOx-saturated and 5% of NOx-limited conditions will be incorrectly considered as 298 transitional. If we widen the transitional regime to lower the probability of misclassification to 299 2%, ~50% of NO_x-saturated and ~10% of NO_x-limited conditions will instead be classified as 300 transitional (Figure 2). We define the regime threshold values as those where the cumulative 301 probability of NO_x-saturated and NO_x-limited conditions is 95% (i.e. the probability of 302 misclassification is 5%) that reflect a balance between accuracy and certainty.

303 Next, we investigate whether the above regime definition should be applied to derive the 304 regime threshold values globally, regionally or individually for each grid cell. Combining all 305 data over the polluted areas of the three regions, we find the transition regime occurs between 306 values for surface FNR of 0.4 and 0.7 (Figure 2). Separating by region, we find that the regime 307 transition occurs between smaller surface FNR values for Europe $(0.4 \sim 0.6)$ than over North 308 America and East Asia $(0.5 \sim 0.8)$ (Figure 2). Figures 3 (a) to (c) shows the classification 309 accuracy (percentage of correct classifications of NO_x-saturated or NO_x-limited conditions) when 310 we apply the regionally-derived range of values for the transition regime. The overall accuracy is 311 high (> 90%) over the majority of polluted areas in the three regions. Lower accuracy is found 312 over California (82%), England (~75%), and North East China (~80%), regions with high 313 anthropogenic emission regions. The high accuracy implies that surface FNR is a quantitatively 314 robust metric for diagnosing the response of surface O₃ to changes in VOC and NO_x emissions.

If we instead derive the regime threshold values separately in each model grid cell, we obtain spatially varying values marking the boundaries of the transitional regime, with higher threshold values over low-latitude regions. This approach, however, does not always improve the accuracy (Figure 3 (d) to (f)), which decreases over California (< 70%), Northeast U.S.A. (< 70%), England (< 70%), the Netherlands (< 60%) and North East China (40% ~ 70%). The low accuracy over these regions reflects a less pronounced correlation between FNR and d[O₃]/dE, 321 and therefore the derived regime threshold values are less stable. Our approach assumes changes 322 in $[O_3]$ in each grid cell are due to the emissions within that box, but $[O_3]$ is also influenced by 323 pollution transported from upwind regions, which could also account for the low accuracy. 324 Sillman et al. [2002] suggests the indicator ratios behave differently for rural and urban 325 environment. The global model resolution cannot fully capture these urban-rural gradients, and 326 therefore even the pixel-based derivation of values marking regime thresholds is unable to 327 characterize fine-scale variations of the photochemical environment. We thus conclude that the 328 regionally-based regime threshold values are most appropriate for surface FNR if a global model 329 is applied to derive the regime threshold. Despite spatial and temporal differences in the factors 330 affecting O₃ production (abundance of solar radiation, VOCs, NO_x and VOC speciation), surface 331 FNR can identify the large-scale variation of O₃ sensitivity.

332 While the column FNR is also able to separate the NO_x-saturated and NO_x-limited 333 conditions qualitatively, modeled column FNR correlates less significantly with surface O_3 -NO_x-334 VOC sensitivity compared to surface FNR. While $d[O_3]/dE_{NOx}$ tends to be negative at low 335 column FNR, and positive at high column FNR (Figures 1(b)), negative $d[O_3]/dE_{NOx}$ still occurs 336 for some high values of column FNR. Nevertheless, the column FNR values marking the 337 boundary between NO_x -saturated and transitional regimes are 0.9 for all three regions (Figure 2). 338 The boundary between the transitional and NO_x-limited regimes, however, varies: 1.4 for North 339 America, 1.2 for Europe and 1.6 for East Asia (Figure 2). Martin et al. [2004a] previously 340 identified a column FNR value of 1 to separate NO_x-limited and NO_x-saturated regimes (Table 1) 341 using GEOS-Chem (version 4.16), close to the threshold value of 0.9 that we find for North 342 America. Duncan et al. [2010] estimated that this regime transition occurs across a column FNR 343 range of 1 - 2 (Table 1) over California. They diagnosed this value as when the radical loss rate 344 through HO_x equals that lost through NO_x ($L_{HOx}/L_{NOx} = 1$). Using column FNR to classify the O₃ 345 sensitivity degrades accuracy across all three regions (Figure 3 (d) to (f)) by about 10% 346 compared to surface FNR. Using spatially varying regime threshold values improves the 347 classification accuracy over most area, suggesting that spatially-varying regime threshold values 348 may be more suitable for column FNR, but the accuracy is still low over those low-accuracy 349 regions identified from surface FNR.

350 **3.2** Column-to-surface relationship

351 We find that the surface and column indicators are robust, providing confidence in the 352 utility of FNR to represent photochemical conditions relevant to ozone production. We address 353 here the uncertainty as to whether the ratio of the vertically integrated column represents the 354 near-surface environment. That is, the relationship of surface to column FNR varies spatially and 355 temporally, mainly due to differences in vertical profiles of NO₂ and HCHO. As in previous 356 studies, we use a model (GEOS-Chem) to adjust column-based ratios observed from satellite 357 instruments to surface-based ratios that are more relevant to near-surface ozone formation 358 [Lamsal et al., 2008; Zhu et al., 2017]. To relate the column-based and surface-based indicator 359 ratios, we calculate the ratio of the GEOS-Chem simulated tropospheric column densities to 360 near-surface number densities of NO₂ and HCHO, defined as an effective boundary layer height 361 for each species (BLH_{eff NO2} and BLH_{eff HCHO}) [Halla et al., 2011]:

$$BLH_{\rm eff} = \frac{\Omega_x}{N} \tag{4}$$

363 where Ω_x is the model simulated tropospheric vertical column density of species x (molecules/cm²), and N is the model simulated number density of species x of the surface layer 364 (molecules/cm³). Similarly, the conversion factor (f_{c_s}) between column and surface FNR is 365 366 calculated as the ratio of column FNR to surface FNR, which is equivalent to the ratio of 367 BLH_{eff_HCHO} to BLH_{eff_NO2}. Generally, if NO₂ or HCHO is well mixed within a homogeneous 368 boundary layer, and most NO₂ or HCHO exists in the boundary layer, the effective boundary 369 layer height should approximate the meteorological boundary layer height [Halla et al., 2011]. 370 As such, it is expected that the column-to-surface relationship of trace gases depends on the PBL 371 height.

Figure 4 shows the relationships between daily meteorological PBLH and BLH_{eff_NO2} and BLH_{eff_HCHO} aggregated over polluted grid cells of the three regions from 2005 to 2012. BLH_{eff_NO2} is correlates strongly with PBLH (R = 0.85), as expected for a short-lived species emitted mainly at the surface. BLH_{eff_NO2} is higher than the simulated PBLH, implying a nonnegligible contribution of free tropospheric NO₂ to the total tropospheric column density (such as from lightning NO_x [*Travis et al.*, 2016; Figure S5]). In contrast, there is little to no relationship of the HCHO vertical profile to PBLH (R = 0.01). HCHO is a secondary photochemical product,

379 formed throughout the atmosphere, with a smaller vertical gradient NO₂, leading to smaller 380 fraction of HCHO within the boundary layer than for NO₂ (Figure 5). The vertical gradient of 381 HCHO is larger in warm season than cold season due to larger contribution of isoprene as a 382 source of HCHO, while the vertical gradient of NO_2 is smaller in warm season (Figure 5) when 383 the surface emission generally mixes through deeper boundary layer and the lightning NO_x 384 source and deep convective mixing are most active. These differences between the NO_2 and 385 HCHO vertical distributions affect the surface and column FNR, and can be accounted for by 386 adjusting the values marking the boundaries of the transitional regime to reflect seasonal and 387 spatial variations in the relationship between column FNR and surface photochemical conditions. 388 As shown in Figure 4 (c), $f_{c,s}$ is inversely correlated with PBLH (R = 0.78), largely driven by the 389 PBLH dependence of NO_2 (Figure 4 (a)).

390 The spatial variation of $f_{c s}$ implies that column-based FNR shows less spatial variability 391 than surface-based FNR (Figure S4). BLH_{eff_NO2} varies seasonally by a factor of 2 yet 392 BLH_{eff HCHO} varies little, with winter-summer differences of less than 500 m. Figure 6 shows a 393 clear seasonal cycle of f_{c_s} over polluted areas in North America and Europe, with a December 394 maximum and July minimum. The shapes of the seasonal cycles of f_{c_s} oppose those of column FNR (Figure 7), which implies that column-based FNR tends to dampen the seasonality of 395 396 surface FNR. Both Europe and North America show larger seasonal cycles than East Asia, where 397 $f_{c s}$ in January exceeds that of July by a factor of 3. The seasonal cycle of BLH_{eff HCHO} in East 398 Asia correlates with PBLH and BLH_{eff_NO2}, with maxima in spring and fall (Figure S5), yielding 399 a smaller f_c s seasonal cycle.

400 As the relationship between surface and column FNR varies spatially and temporally 401 (section 3.1), we adjust the column-based FNR values marking the transitional ozone production 402 regime by applying the modeled $f_{c s}$ to the threshold values of surface FNR. The variation in 403 column-to-surface relationships of NO₂ and HCHO is dependent on the vertical profiles, which 404 are mostly driven by meteorology. We find the column-to-surface relationship of FNR does not 405 vary much year-to-year: the standard deviation for any given month is lower than 8% for all 406 regions (Figure S6). We find no statistically significant trends in the column-to-surface 407 relationships, suggesting that the constant regime threshold values will not affect the trend 408 analysis in Section 4. Also, as we are attempting to generalize the derived regime threshold

409 values for application beyond the model simulation period, constant regime threshold values are 410 preferred. Therefore, we do not adjust the regime threshold values to include inter-annual 411 variability of the column-to-surface relationships. The pink band in Figure 7 shows the seasonal 412 cycle of these model-simulated column-based values averaged from daily data within each 413 month over the polluted regions separately within North America, Europe and East Asia. The 414 lighter band represents the 1σ deviation of these values derived from individual polluted grid 415 cells in each regional domain. Larger standard deviations occur over Europe, reflecting stronger 416 spatial variations of the column FNR values spanning the transitional regime. The coarse spatial 417 resolution of GEOS-Chem cannot capture sharp urban-rural gradients though it does resolve 418 large-scale variations in meteorology and topography. The maximum standard deviation occurs 419 in spring, when meteorological conditions range widely during the transition from winter to 420 summer and the onset of biogenic emissions.

421 Following the seasonal cycle of the column-to-surface relationship (Figure 6), the 422 transitional regime thresholds in Figure 7 are higher in the cold season than in the warm season. 423 The transition from the NO_x-saturated to the transitional regime occurs for column-based FNR 424 ranging from 0.5 in June to 1.6 in January over North America, 0.8 in June to 1.0 in January over 425 East Asia, and 0.6 in August to 1.8 in January over Europe. The thresholds between the 426 transitional and the NO_x-limited regime range from 0.8 in July to 2.5 in December for North 427 America, 1.2 in August to 1.6 in December for East Asia, and 0.9 in July to 2.7 in December for 428 Europe. East Asia shows a smaller seasonal cycle in these threshold values compared to North 429 America and Europe. The threshold from NO_x-saturated to transitional regime is generally 430 smaller than 1.0 as proposed in Duncan et al. [2010], likely due to different definitions for the 431 transitional regime (Table 1).

432 **3.3 Model and satellite comparison**

While the model demonstrates that tropospheric column ratios of HCHO to NO_2 can indicate surface O_3 sensitivity to NO_x and VOC emissions, both satellite retrievals and model simulations are subject to large uncertainties. Here we compare the OMI-derived seven-day average FNR with the GEOS-Chem base-case simulation to identify where and when the satellite products and model agree best, implying more confidence in our understanding. We restrict the comparison to polluted regions, defined as those grid cells in GEOS-Chem where annual average 439 tropospheric $\Omega_{NO2_GC} > 2.5 \times 10^{15}$ molecules/cm². Table 2 summarizes the comparison between 440 modeled and OMI FNR.

441 We find that the correlation coefficient between the model- and satellite-derived FNR 442 products depends on the choice of NO₂ product, while the mean bias depends on the choice of 443 HCHO product. FNR_{OMI} (with GEOS-Chem profiles applied) using $\Omega_{NO2 SP}$ (R: 0.44 ~ 0.74) 444 correlates better with GEOS-Chem than $\Omega_{NO2 DP}$ (R: 0.28 ~ 0.63) for all three regions. Among 445 the four combinations, FNR_{OMI SS} correlates best with FNR_{GC} over North America and East 446 Asia, while FNR_{OMI BS} correlates best over Europe. The choice of HCHO product does not influence the overall correlation, except over Europe where using Ω_{HCHO_BIRA} results in a higher 447 448 correlation coefficient compared to $\Omega_{HCHO SAO}$. The low correlation coefficient of FNR_{OMI SD} 449 and FNR_{OMI BD} is largely caused by observations with low $\Omega_{NO2 SP}$ and $\Omega_{NO2 DP}$ over clean regions ($\Omega_{NO2} < 1.5 \times 10^{15}$ molecules/cm²). We find both $\Omega_{NO2 DP}$ and $\Omega_{NO2 SP}$ match $\Omega_{NO2 GC}$ 450 over polluted regions, and the mean differences with GEOS-Chem are within 5% for both 451 452 products (Table S1). FNR_{OMI} is on average higher than FNR_{GC} by 10% to 40% if $\Omega_{\text{HCHO BIRA}}$ is 453 used, and lower than FNR_{GC} by 10% to 30% using $\Omega_{\rm HCHO}$ sao. The opposite sign of the mean 454 offset results from the large difference between two HCHO retrievals: Ω_{HCHO_SAO} is on average 455 50% lower than $\Omega_{HCHO BIRA}$ across the three regions (Table S1). Discarding observations with negative HCHO columns corrects the negative offsets of FNR_{OMI_SS} and FNR_{OMI_SD} relative to 456 the model, but increases the positive offsets of FNR_{OMI BD} and FNR_{OMI BS}. 457

458 Although the absolute values of FNR_{OMI} differ from FNR_{GC}, FNR_{OMI} is in general able to 459 capture the spatial and temporal variation of the O₃ production regime inferred from FNR_{GC}. The 460 agreement (defined as the percentage of both FNR_{GC} and FNR_{OMI} falling in the same 461 photochemical regime) is higher than 80% for warm season and 60% for cold season across three 462 regions (Table 2). The agreement also depends on the choice of HCHO product, especially in 463 warm season. Figure 7 shows the seasonal cycle of FNR_{GC} and FNR_{OMI} averaged from 2005 to 464 2012 for each region. FNR_{OMI} shows a positive offset and low correlation coefficient in cold 465 season for all four combinations, especially over Europe, reflecting the HCHO overestimate in 466 winter. Nevertheless, the products and model all agree that NO_x -saturated or transitional regimes 467 dominate in winter. FNR_{GC}, along with the four combinations of OMI observed FNR, indicate

468 NO_x-limited regimes from May to September over all three regions, but may disagree for 469 individual grid cells (Figure S7 to S10). FNR_{GC} disagrees with FNR_{OMI} more frequently in spring 470 and fall during the transitions between regimes. FNR_{OMI BD} and FNR_{OMI BS} are consistently 471 higher than FNR_{GC} over all three regions, leading to a longer NO_x-limited regime versus FNR_{GC} , 472 but they match regimes diagnosed with FNR_{GC} better in warm season (Table 2). In contrast, 473 FNR_{OMLSS} and FNR_{OMLSD} are lower than FNR_{GC}, especially in the warm season, leading to a 474 longer NO_x-saturated and transitional regime, and better match the ozone production regimes 475 indicated by FNR_{GC} in cold season (Table 2).

476 The re-gridded FNR_{OMI} at coarse resolution tends to smear spatial gradients in ozone 477 production regimes. To characterize the spatial heterogeneity of O₃ sensitivity to its precursor 478 emissions, we recommend the Level-3 OMI HCHO and NO₂ data available at $0.25^{\circ} \times 0.25^{\circ}$ for 479 general applications of the indicator ratio that do not involve comparison or interpretation with a 480 model. We compare FNR_{GC} with Level-3 FNR_{OMI BS} (FNR_{OMI BS L3}) by spatially matching 481 Level-3 data to the model grid. Using different prior profile shapes leads to minor differences in 482 AMFs for NO₂ and HCHO (Table S1). While model-satellite discrepancies are larger for 483 FNR_{OMI BS L3} vs. FNR_{OMI BS} derived with our daily GEOS-Chem profiles, the overall correlation 484 is comparable to and even better than FNR_{OMI BD} and FNR_{OMI SD} over Europe and East Asia 485 (Table 2). FNR_{OMI BS L3} is on average higher than FNR_{GC} by 20%. Evaluation with aircraft data 486 suggests GEOS-Chem underestimates HCHO concentrations by 10% over the southeast U.S.A. 487 [Zhu et al., 2016]. In order to correct this systematic model-satellite discrepancy, which likely 488 reflects the model underestimate of HCHO, we additionally correct the regime thresholds for 489 FNR_{OML BS L3} by increasing the values marking the boundaries of the transitional regimes 490 derived from section 3.2 by 20%.

491 **4 Decadal changes of O₃-NO_x-VOC sensitivity**

Here we investigate the decadal trend of surface O_3 sensitivity over polluted areas in North America, Europe, and East Asia. For this application, we use monthly average Level-3 gridded with the original standard AMFs included in the products from 2005 to 2015 [*Duncan et al.*, 2010; *Jin and Holloway*, 2015]. Satellite-derived ozone production regimes generally agree with *in-situ* ground-based studies over the three regions (Table 3), but OMI observations tend to overestimate the NO_x sensitivity. The OMI overpass time is in the afternoon, when the NO₂ level is at its daily minimum and thus ozone production is most NO_x -limited. The horizontal resolution of OMI data is likely to miss urban core NO_x -saturated regimes sampled at individual urban sites (e.g. *Pusede et al.*, [2012]).

501 Before applying the OMI L3 product to analyze decadal trends, we investigate whether 502 long-term changes in FNR are compromised by instrument degradation and data availability 503 changes. First, Marais et al. [2012] suggest an artificial increase in the background HCHO 504 column in the OMI SAO retrieval due to instrument degradation. This feature does not appear in 505 the BIRA retrieval, which applies a zonal reference sector correction (e.g., Figure S11). 506 FNR_{OML BS L3} does not show any artificial trend over remote Pacific region either (Figure S11). 507 Second, OMI data coverage has decreased over the years mostly due to growing row anomalies 508 (Figure S12), which tend to decrease monthly average HCHO column with time [De Smedt et al. 509 2015]. We find that the data coverage has declined by about 20% from 2005 to 2015 for both the 510 Level-3 OMI HCHO and NO₂ products, implying that these sampling biases may largely cancel 511 out when we take the HCHO/NO₂ ratio (Figure S12). To test the influence of this sampling bias, 512 we calculate another time series of monthly average FNR that randomly discards the 513 corresponding number of daily HCHO and NO₂ data such that the data coverage for a given 514 month is set as the minimum number of samples obtained during that month between 2005 and 515 2015 (Figure S13). There is no systematic offset due to the changing data coverage. We find that 516 sampling differences may influence the magnitude of the FNR_{OMI} trend (slope), but have little 517 impact on the diagnosed changes in the ozone production regimes (Figure S13 versus Figures 8 -518 11). Note that our definition of regimes relies on the modeled values that do not include inter-519 annual variability (pink shaded bands in Figure S13), and therefore the changes over time in the 520 ratio reflect trends in the satellite products that do not contain any information from model. It 521 should also be noted that a higher solar zenith angle cut-off was applied to the HCHO retrieval 522 mainly reflecting the lower retrieval sensitivity to HCHO as compared to NO₂. The data 523 coverage of satellite-derived HCHO is thus smaller than for NO_2 . We calculate a new time series 524 of monthly average FNR that is constructed using only days when both HCHO and NO_2 have 525 valid data. We find similar trends in FNR and in changes in the ozone production regime (Figure 526 S14) as for our original time series that includes all available data (Figures 8 - 11), but the

resampled data show larger fluctuations, due to increasing uncertainties as the number ofobservations used for temporal averaging decreases.

529 4.1 North America

530 From 2005 to 2015, NO_x sensitivity increased over populated regions of North America 531 (Figures 8 and 9). Previous studies have shown that NO₂ levels decreased by 25% to 55% over 532 the continental U.S.A. over the past decade, resulting from the implementation of nationwide 533 emission controls [Lamsal et al., 2015; Schneider et al., 2015; Tong et al., 2015; Duncan et al., 534 2016]. De Smedt et al. [2015] note a decreasing trend of HCHO (-10% to -2%) over the eastern 535 U.S.A. and California from 2005 to 2014, but the magnitude is much less significant than NO₂, 536 as the main source of HCHO from biogenic emissions fluctuates with meteorology [Guenther et 537 al., 2006; Millet et al., 2008]. The NO_x-limited regime dominates over the northeast U.S.A. in both May 2005 and 2015. The FNR_{OMI BS L3} indicates New York City was in NO_x-saturated 538 539 regime in May 2005, but shifted to NO_x-limited by 2015. The NO_x-limited regime occurred from 540 June to August in NYC in 2005, and the length of the NO_x -limited regime increased from three 541 months in 2005 to five months in 2015 (Figure 8). The average length of the NO_x -limited regime 542 in 2005 to 2009 is 3.2 months, and increases to 4.2 months for the 2011 to 2015 period. The 543 length of the NO_x-saturated regime has decreased from eight months in 2005 to five months in 544 2015. The five-year average length of the NO_x -saturated regime has decreased from 7.4 (2005 to 545 2009) to 6.0 months (2011 to 2015). Over Chicago, FNR_{OMI BS L3} varies inter-annually but 546 increases by 0.10 per year from 2005 to 2015, extending the average length of NO_x-limited 547 regime from 3.0 months between 2005 to 2009 to 4.8 months between 2011 and 2015, and 548 narrowing the NO_x-saturated regime from 7.2 to 5.2 months. Jing et al. [2014] suggest O₃ 549 formation has shifted from NO_x -limited to VOC-limited in 2008/2009, but such increasing VOC 550 sensitivity is not observed from FNR_{OMI BS L3}. We also find similar regime shifts in other cities 551 such as Detroit and Los Angeles (not shown). For other cities with pronounced emission 552 reductions [Duncan et al., 2016], such as Philadelphia, Atlanta and Phoenix, while the O_3 553 production regime remains NO_x-limited in the warm season, we find an increasing trend of 554 FNR_{OMI BS L3} and consequently enhanced NO_x sensitivity (not shown). The observed increasing 555 NO_x sensitivity over the U.S. cities implies that continued regional NO_x emission control

programs should be effective for surface O_3 mitigation, as shown in modeling studies [*Frost et al.*, 2006; *Song et al.*, 2010].

558 **4.2 Europe**

559 Similar to U.S. cities, surface O_3 production is becoming more sensitive to NO_x (NO_x -560 limited) over Europe in response to decreasing NO_x emissions. Satellite-derived NO₂ products 561 also show significant decreasing trends of -50% to -3% over Europe and Russia, driven by a 562 combination of environmental policy and reduced economic activity during recessions 563 [Castellanos and Boersma, 2012; Duncan et al., 2016]. No significant HCHO trend occurs over 564 Europe [De Smedt et al., 2015]. We find that transitional and NO_x -saturated regimes were 565 dominant over Great Britain and western Europe in July 2005. Ozone production regimes 566 transitioned to NO_x-limited regime in northern England, Germany and France in 2015 (Figure 10). Overall, an increasing trend of FNR_{OMI BS L3} occurs over London, extending the average 567 568 length of the NO_x -limited regime from 1.4 months between 2005 and 2009 to 2.4 months 569 between 2011 and 2015. The five-year average length of Surface O₃ production in London was 570 sensitive to VOC emissions most of the year between 2005 and 2014 except for July 2009 when 571 FNR_{OMI BS L3} reached the NO_x-limited regime. We find a sharp increase of FNR_{OMI BS L3} in 2015, with surface O₃ production NO_x-limited in July and August. Amsterdam also shows an 572 573 increasing trend of FNR_{OMI BS L3} that peaks in 2012, and the length of NO_x-limited regime has 574 increased from 2.0 (2005 to 2009) to 2.8 months (2011 to 2015). Note the average FNR_{OMLBS L3} 575 was relatively low in urban areas of Europe compared with cities in North America with similar 576 of NO_x emission levels. This may reflect low HCHO concentration in Europe due to lower 577 biogenic emissions [De Smedt et al., 2015]. As shown in Figure 10, monthly average 578 FNR_{OMLBS L3} over London and Amsterdam does not vary significantly with season from 2005 to 579 2015. This finding, however, does not necessarily indicate a weak seasonality of O₃ sensitivity. $f_{c,s}$ varies with season by a factor of three over Europe (Figure 6), which dampens the seasonal 580 cycle of the column-based FNR_{OMI_BS_L3}. Also, OMI observations of HCHO and NO₂ at high 581 582 latitudes are subject to large uncertainties due to signal interference of unknown species 583 [González Abad et al., 2015]. Most wintertime observations were excluded due to high solar 584 zenith angle and larger retrieval uncertainty.

585 **4.3 East Asia**

586 The trends in surface O_3 -NO_x-VOC sensitivity are uneven and mixed over East Asia, 587 where we find increasing NO_x sensitivity over Japan and Korea, and an overall increasing VOC 588 sensitivity over China (Figure 11). Changes in surface O₃ sensitivity over China have been 589 investigated in Jin and Holloway [2015] using OMI observations under the assumption of that 590 the transitional regime occurs when 1<FNR<2. This study builds upon *Jin and Holloway* [2015] 591 by incorporating the seasonality of column-to-surface relationships when defining the 592 transitional regime. We find here that the transition to the NO_x-limited regime in summer occurs 593 at FNR_{OMI BS L3} < 2, and the transition to the NO_x-saturated regime occurs at FNR_{OMI BS L3} < 1, 594 leading to a larger spatial extent of the NO_x-limited regime in summer and the NO_x-saturated 595 regime in winter compared to Jin and Holloway [2015]. Jin and Holloway [2015] show a spatial 596 and temporal expansion of the NO_x-saturated regime over East China, but the developed 597 megacities, such as Beijing, Shanghai and Guangzhou, show an increasing NO_x sensitivity due to 598 NO_x emission reduction. The duration of NO_x -limited regime extended from one month in 2005 599 to four months in 2015 over Beijing (Figure 11). The average length of the NO_x-limited regime 600 from 2005 to 2009 is 1.4 months, and increased to 2.2 months from 2011 to 2015. However, the 601 length of NO_x-saturated regime remains around 8 months throughout the entire period from 2005 602 to 2015. FNR_{OML BS L3} increased sharply in the summer of 2008, reflecting emission controls 603 during the Beijing Olympic Games [Wang et al., 2009]. For other cities over the Northern China 604 Plain such as Jinan (Figure 11(b)), O_3 production regimes in May have become NO_x-saturated 605 since 2011. FNR_{OMI BS L3} in Jinan decreased from 2005 to 2011, and remained stable since 2011, 606 likely associated with nationwide NO_x reductions from power plants [Liu et al., 2016]. The 607 length of NO_x-saturated regime has increased from eight months between 2005 and 2009 to nine 608 months between 2011 and 2015. The Pearl River Delta shows increasing NO_x sensitivity due to 609 successful NO_x emission control; O₃ sensitivity was in the transitional regime in May 2005, but 610 shifted to the NO_x-limited regime in 2015. Duncan et al. [2016] found large decreases of OMI 611 NO₂ levels over South Korea and Japan, attributed to national environmental regulations. We 612 find an increasing NO_x sensitivity over Korea and Japan accordingly. Seoul and Tokyo were in 613 the NO_x-saturated regime in May 2005, and they both shifted to the transitional regime in 2015 614 (Figure 11). The value of FNR_{OMI BS L3} was consistently below the upper boundary of 615 transitional regime in Tokyo (Figure 11(c)) and Seoul (Figure 11(d)), where surface O₃

616 production was either NO_x -saturated or transitional except for July 2015 in Seoul and August 617 2015 in Tokyo. *Duncan et al.* [2016] suggest that effective domestic control strategies may have 618 been negatively offset by increasing trans-boundary NO_x emissions from China, resulting in a 619 smaller positive trend in $FNR_{OMI_BS_L3}$ in Seoul and Tokyo than over European and U.S cities 620 with similar emission changes.

621 **5 Conclusions**

622 We use OMI observations of NO_2 and HCHO column densities, along with a global 623 chemical transport model (GEOS-Chem) to examine the sensitivities of surface O_3 pollution to 624 NO_x and VOC emissions over northern mid-latitude source regions. We use the GEOS-Chem model to determine the regime thresholds for FNR with two emission perturbation simulations. 625 626 We find that surface FNR in the model does indicate surface O₃ sensitivity, and that regionally constant FNR thresholds can separate the NOx-limited and NOx-saturated conditions to at least 627 628 90% confidence. FNR values marking the boundaries of the photochemical regimes are derived 629 from the model, and thus depend on the mechanism used to represent photochemistry. Travis et 630 al. [2016] suggest an overestimate of NO_x emissions over the eastern U.S.A. Such an 631 overestimate could lead to excessive tropospheric NO₂ columns as well as an underestimate of d[O₃]/dE_{NOx}, which may largely cancel out so that the threshold values would be less sensitive to 632 633 this error. Erroneously high NO₂ columns, however, could lead us to diagnose excessively low 634 regime threshold values over NO_x-saturated regions.

635 Column FNR shows a lower regime classification accuracy, largely due to variations in 636 column-to-surface relationships. The column-to-surface relationships for NO₂ correlate strongly 637 with PBLH, but weakly for HCHO. As a result, the column-to-surface relationship of FNR ($f_{c,s}$) 638 is inversely correlated with PBLH. Following the spatial and temporal variations of PBHL, f_{cs} 639 shows pronounced seasonal cycles with maxima in winter and minima in summer, which act to dampen the spatial and temporal variation of surface O3 sensitivity. We adjust the regime 640 641 threshold values for column-based FNR using the modeled $f_{c s}$. The derived column FNR 642 thresholds marking the boundaries between ozone production regimes vary by a factor of 3 over 643 North America and Europe. The modeled vertical profiles are also sensitive to the PBL scheme. 644 Full PBL mixing scheme is implemented in GEOS-Chem. The full-mixing scheme in GEOS-

Chem is likely to underestimate the vertical gradient of both NO₂ and HCHO [*Lin et al.*, 2010; *Zhang et al.*, 2016].

647 Even though modeled FNR can indicate surface O₃ sensitivity to NO_x vs. VOC 648 precursors, both satellite-derived and modeled FNR are subject to uncertainties. We compare 649 four combinations of two OMI HCHO products (BIRA, SAO) and OMI NO₂ (DP, SP) products 650 with GEOS-Chem simulations. The spatial and temporal correlation between the modeled and 651 observed indicator ratios depends on the choice of NO₂ product, while the mean bias depends on 652 the choice of HCHO product. We note that wintertime satellite retrievals of HCHO incur large 653 uncertainties due to diminished satellite sensitivity near the surface [De Smedt et al., 2015]. 654 Qualitatively, however, such uncertainties should not affect the conclusion that ozone production 655 is NO_x-saturated in winter over regions heavily influenced by anthropogenic emissions, as noted 656 in previous studies [Kleinman, 1991, 1994; Jacob et al., 1995]. Satellite-derived O₃ sensitivity 657 generally agree with *in situ* observations performed in previous studies. While the distinct 658 behavior of indicator ratio over urban and rural environment cannot be fully resolved from the 659 coarse resolution of global model, the finer resolution of OMI observation can explain the 660 majority of the spatial and temporal variation of O₃ sensitivity. Future work could assess the ability of the OMI indicator ratio to reveal urban fine-scale features with a higher resolution 661 662 (e.g., regional) model.

663 Combining model-derived threshold values with a decadal record of satellite 664 observations, we further investigate how O₃ production sensitivity to precursors has changed 665 over the 2005 to 2015 period. We find a general increase in FNR_{OMI} over the urban areas of 666 North America, Europe, South Korea and Japan from 2005 to 2015, driven by NO_x emission 667 reductions imposed over the past decade. The spring transition to a NO_x -limited regime has 668 shifted earlier in some megacities, and the NO_x-limited regime has become dominant in summer. 669 China shows an overall decrease in FNR_{OMI} except for the most developed areas such as Beijing, 670 Shanghai and Pearl River Delta, where emission control strategies have been implemented. In 671 our FNR analysis, HCHO serves as an indicator of reactivity-weighed VOCs, but the yield and production of HCHO from isoprene is non-linearly dependent on the NO_x level [Wolfe et al., 672 673 2016]; this non-linearity implies that FNR may underestimate increases in NO_x sensitivity as 674 NO_x emissions decline.

675 Surface O_3 sensitivity also varies throughout the day and from day to day. The suitability 676 of the FNR_{OMI} for daily variation is still limited by the uncertainties associated with the OMI 677 HCHO and NO₂ retrievals. In addition, the spatial resolution of OMI may be too coarse to reveal 678 VOC-limited chemistry in urban cores. Near-term advances in space-based observations of 679 HCHO and NO₂ from geostationary satellites as anticipated to occur over East Asia 680 (Geostationary Environment Monitoring Spectrometer), Europe (Sentinel-4) and North America 681 (Tropospheric Emissions: Monitoring of Pollution) [Lahoz et al., 2012], offer exciting opportunities to explore the potential for space-based FNR to diagnose ozone production regimes 682 683 at finer spatial and temporal scales.

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693 **Disclaimer:**

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1057 Figure 1 GEOS-Chem model estimates of the normalized ozone sensitivity to 20% decreases in global NO_x and VOC emissions $(d[O_3]/dE_{NOx}$ in orange, $d[O_3]/dE_{VOC}$ in blue) in units of 1058 molecules $cm^{-2}s^{-1}$, versus the modeled (a) surface HCHO/NO₂ and (b) tropospheric column 1059 HCHO/NO₂ aggregated over the three selected regions (North America, Europe, Asia). Each 1060 point is equal to the normalized sensitivity ratios of daily one-hour averages between 1 and 2 PM 1061 from 2006 to 2012 in a single model grid cell. We only include polluted grid cells, defined as 1062 cells with average modeled tropospheric NO₂ column densities higher than 2.5×10^{15} 1063 1064 molecules/ cm^2 .





1068 Figure 2 Cumulative probability of NO_x-saturated $(d[O_3]/dE_{NOx} < 0)$ and NO_x-limited $(d[O_3]/dE_{NOx} > d[O_3]/dE_{VOC} > 0)$ conditions, as a function of modeled (a-d) surface HCHO/NO₂ 1069 1070 and (e-h) tropospheric column HCHO/NO2 over (a, e) North America, (b, f) Europe, (c, g) East 1071 Asia and (d, h) all three regions aggregated, selecting for polluted conditions as in Figure 1. The 1072 blue line represents the cumulative probability of NO_x-saturated conditions for all HCHO/NO₂ 1073 smaller than each given value. The red line represents the cumulative probability of NOx-limited 1074 condition for HCHO/NO₂ greater than each given value. The cumulative probability indicates the likelihood of correctly identifying the NO_x-limited or the NO_x-saturated conditions at any given 1075 1076 HCHO/NO₂ as simulated by the GEOS-Chem model. The probability is calculated from the 1077 normalized sensitivity ratios of daily one-hour averages between 1 and 2 PM from 2006 to 2012 1078 (individual points in Figure 1).



Figure 3 Percentage of correct classifications based on modeled (top two rows) surface or (bottom two rows) column FNR to NO_x -saturated or NO_x -limited conditions using: (first and third rows) regionally-derived values marking the boundary of the transitional regime (second and fourth rows) pixel-based derivation of the transitional regime range over North America (first column), Europe (second column), East Asia (third column). We only include polluted grid cells, defined as cells with average modeled tropospheric NO_2 column densities higher than 2.5 × 10^{15} molecules/cm².



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1090 Figure 4 Modeled effective boundary layer height of NO_2 (a) and HCHO (b), column-to-surface 1091 conversion factor of FNR (c), versus planetary boundary layer height (PBLH) over polluted areas

1092 within the three regions (defined as in Figure 1). Each point is the GEOS-Chem daily one-hour

average from 1 to 2 PM. The black lines are the best-fit linear regression (a and b) and reciprocal

1094 regression (c).



Figure 5 Vertical profiles of HCHO and NO₂ sub-column densities averaged from daily one-hour data between 1 and 2 PM for the warm season (May to September, a) and cold season (October to April, b) from 2005 to 2012 over the polluted areas of three regions aggregated. The dash line shows the average planetary boundary layer height.

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1106 Figure 6 Seasonal cycle of the column-to-surface conversion factors of FNR (f_{c_s}) over North 1107 America, Europe and East Asia averaged from daily GEOS-Chem data for polluted areas 1108 (modeled tropospheric NO₂ column density higher than 2.5×10^{15} molecules/cm²) from 2005 to 1109 2012. The dashed error bars are 1 σ standard deviation representing spatial variations.



Figure 7 Seasonal cycle of GEOS-Chem modeled (black) and four combinations of OMI observed FNR: 1) SAO HCHO : SP NO₂ (OMI_SS); 2) BIRA HCHO : SP NO₂ (OMI_BS); 3) BIRA HCHO : DP NO₂ (OMI_BD); 4) SAO HCHO : DP NO₂ (OMI_SD), along with the seasonal cycle of column-based regime thresholds in over a) North America, b) Europe, and c) East Asia averaged for polluted areas (modeled tropospheric NO₂ column density higher than 2.5 $\times 10^{15}$ molecules/cm²) from 2005 to 2012. The dash error bars and the lighter pink band are 1 σ standard deviation representing spatial variations.



1120 Figure 8 Ozone production regimes over Northeast US in May 2005 and 2015 (top), and time 1121 series of OMI-derived FNR along with the regime threshold values (pink shading) in New York 1122 City. The regime classification uses the ratio of monthly average OMI Level-3 BIRA HCHO to 1123 Level-3 NASA SP NO₂. Solid lines indicate the warm season (May to September) and the dashed lines indicate the cold season (October to April). The transition regime threshold values 1124 1125 are adjusted based on the column-to-surface relationships (Section 3.2), and the model-satellite 1126 difference (Section 3.3). The observed FNR are monthly average OMI Level-3 BIRA HCHO to 1127 Level-3 NASA SP NO₂ for the grid cells fully covering these cities. The uncertainty (error bars) 1128 is calculated from monthly standard deviation of NO₂ and HCHO using Equation (3). The blue line shows the linear regression trend. Areas with average observed tropospheric NO₂ column 1129 densities $< 2.5 \times 10^{15}$ molecules/cm² are masked. 1130



1133 Figure 9 Same as Figure 8 but for Great Lake Region and Chicago.



Figure 10 Same as Figure 8 but for Central Europe in July 2005 and 2015 (top), and time series in London (a) and Amsterdam (b). The letters mark the approximate location of London and

1138 Amsterdam. Missing values indicate no sufficient valid observations during the month.



Figure 11 Same as Figure 8 but for East Asia in May 2005 and 2015 (top), and time series in

- Beijing (a), Jinan (b), Seoul (c), and Tokyo (d). The letters mark the approximate location of the four cities.

- 1144 Table 1 Summary of previous studies that use surface or column HCHO/NO₂ as indicators of
- 1145 surface ozone sensitivity.

Reference	Study Area	Indicator Ratio	Regime Threshold Values	Model	Observation
Tonnesen et al., [2000]	New York City Area	Surface HCHO/NO ₂	<0.8 NO _x -Saturated >1.8 NO _x -limited 0.8~1.8 Transition	RADM	Ground-based Measurements
Martin <i>et al.</i> , [2004a]	North America, East Asia and Europe	Column HCHO/NO ₂	<1.0 NO _x -Saturated >1.0 NO _x -limited	GEOS-Chem	GOME
Duncan <i>et al.</i> , [2010]	U.S.	Column HCHO/NO ₂	<1.0 NO _x -Saturated >2.0 NO _x -limited 1.0~2.0 Transition	NASA LaRC	OMI
Choi <i>et al.</i> , [2012]	U.S.	Column HCHO/NO ₂	<1.0 NO _x -Saturated >2.0 NO _x -limited 1.0~2.0 Transition	CMAQ	GOME
Chang <i>et al.</i> , [2016]	Northeast U.S.	Column HCHO/NO ₂	<1.5 NO _x -Saturated >2.3 NO _x -limited 1.5~2.3 Transition	CMAQ- DDM	OMI
Jin and Holloway [2015]	China	Column HCHO/NO ₂	<1.0 NO _x -Saturated >2.0 NO _x -limited 1.0~2.0 Transition	N/A	OMI

- 1148 Table 2 Comparison between OMI FNR and GEOS-Chem modeled FNR. Mean bias is the
- 1149 averaged difference between OMI observed minus model retrievals. Agreement is defined as the
- 1150 percentage of both FNR_{GC} and FNR_{OMI} falling in the same photochemical regime. Warm season
- 1151 includes May to September, and cold season includes October to April.

		Mean Bias	R	Agreement (warm)	Agreement (cold)
North	FNR _{GC} vs. FNR _{OMI_SS}	-25%	0.74	94%	78%
America	FNR _{GC} vs. FNR _{OMI_BS}	17%	0.74	97%	74%
	FNR _{GC} vs. FNR _{OMI_BD}	17%	0.63	97%	70%
	FNR _{GC} vs. FNR _{OMI_SD}	-26%	0.61	94%	76%
	FNR _{GC} vs. FNR _{OMI_BS_L3}	10%	0.56	96%	67%
Europe	FNR _{GC} vs. FNR _{OMI_SS}	-18%	0.44	83%	73%
	FNR _{GC} vs. FNR _{OMI_BS}	28%	0.61	90%	63%
	FNR _{GC} vs. FNR _{OMI_BD}	33%	0.44	90%	63%
	FNR _{GC} vs. FNR _{OMI_SD}	-15%	0.28	83%	71%
	FNR _{GC} vs. FNR _{OMI_BS_L3}	33%	0.44	90%	56%
East	FNR _{GC} vs. FNR _{OMI_SS}	-30%	0.68	80%	83%
Asia	FNR _{GC} vs. FNR _{OMI_BS}	10%	0.72	88%	74%
	FNR _{GC} vs. FNR _{OMI_BD}	39%	0.53	89%	70%
	FNR _{GC} vs. FNR _{OMI_SD}	-10%	0.47	85%	81%
	FNR _{GC} vs. FNR _{OMI_BS_L3}	17%	0.65	86%	74%

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Table 3 Comparison with previous *in-situ* ground-based studies over individual sites. The ozone production regime is derived from monthly average FNR using OMI Level-3 BIRA HCHO to

Level-3 NASA SP NO₂.

Period	Study Area	Ozone Sensitivity	This Study	Method and Reference	
July, 2005	Beijing	Urban: NO _x -saturated	Urban: Transitional to	CMAQ-RSM [Xing et al.,	
July, 2005	Shanghai	Rural: NO _x -Limited	NO _x -limited	2011]	
July, 2005	Guangzhou		Rural: NO _x -Limited		
Nov., 2007	Guangzhou	NO _x -saturated	NO _x -saturated	Photochemical trajectory model [<i>Cheng et al.</i> , 2010]	
June – July, 2005	Beijing	NO _x -limited or Transitional	Transitional	Ground-based Measurements [Wang et al., 2006]	
June - July, 2006	Lanzhou	NO _x -limited	NO _x -limited	Observation-based model (MCM3.2) [<i>Xue et al.</i> , 2014]	
May - June, 2005	Shanghai	NO _x -saturated	Transitional		
Aug., 2006	Beijing	Mixed s	Transitional	Observation-based photochemical box model (OBM) [<i>Lu et al.</i> , 2010]	
Nov., 2006	PRD	Mixed	Rural: NOx-Limited Urban: NO _x -saturated	Chemical Transport Model (EBM) [<i>Li et al.</i> , 2013]	
Aug., 2007	Beijing	Transitional	Transitional	1-D photochemical model [<i>Liu et al.</i> , 2012]	
Nov., 2008	PRD	NO _x -saturated	NO _x -saturated	WRF-Chem [Ye et al., 2016]	
May - June, 2010	Jiangsu	Mixed	NO _x -saturated	Observation-based model (RACM) [<i>Pan et al.</i> , 2015]	
Summer 2009 - 2011	Miyun Site (Beijing)	NO _x -saturated	Transitional to NO _x - limited	Smog Production algorithm (OBM) [<i>Wang et al.</i> , 2008]	
2009-2011	Seoul	NO _x -saturated	NO _x -saturated or Transitional	Statistical Correlation Analysis [<i>Iqbal et al.</i> , 2014]	
May - Oct., 2006	Houston	NO _x -Limited	NO _x -Limited	CAMx [Kommalapati et al., 2016]	
Summer 2008-2009	Chicago	NO _x -saturated	NO _x -Limited	Statistical trend analysis [<i>Jing et al.</i> , 2014]	
2007	Sacramento, CA	NO _x -Limited or Transitional	NO _x -Limited or Transitional	Observation-based 1-D plume model [<i>LaFranchi et al.</i> , 2011]	
2007-2010	Southern and Central San Joaquin Valley	NO _x -Limited	NO _x -Limited	Observation-based method [<i>Pusede et al.</i> , 2012]	
2007-2010	Northern San Joaquin Valley	NO _x -saturated	NO _x -Limited		
Sept., 2013	Houston	Mostly NO _x -Limited (afternoon)	NO _x -Limited	Observation-based model (CB05) [<i>Mazzuca et al.</i> , 2016]	

Figure 1.



Figure 2.



Figure 3.



Figure 4.



Figure 5.

Warm Season

Cold Season



Figure 6.



Figure 7.



Figure 8.



Figure 9.



Figure 10.



Figure 11.



2005 2006 2007 2008 2009 2010 2011 2012 2013 2014 2015

