2 Enhancing Long-Term Trend Simulation of Global 3 Tropospheric OH and Its Drivers from 2005-2019: A 4 Synergistic Integration of Model Simulations and Satellite 5 Observations

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Amir H. Souri^{1,2*}, Bryan N. Duncan¹, Sarah A. Strode^{1,2}, Daniel C. Anderson^{1,3}, Michael E. Manyin^{1,4}, Junhua Liu^{1,2}, Luke D. Oman¹, Zhen Zhang^{5,6}, and Brad Weir^{2,7}

- 9
- ¹Atmospheric Chemistry and Dynamics Laboratory, NASA Goddard Space Flight Center (GSFC),
- 11 Greenbelt, MD, USA
- 12 ²GESTAR II, Morgan State University, Baltimore, MD, USA
- 13 ³GESTAR II, University of Maryland Baltimore County, Baltimore, MD, USA
- ⁴Science Systems and Applications, Inc., Lanham, MD, USA
- 15 ⁵National Tibetan Plateau Data Center (TPDC), State Key Laboratory of Tibetan Plateau Earth System,
- 16 Environment and Resource (TPESER), Institute of Tibetan Plateau Research, Chinese Academy of17 Sciences, Beijing, China
- 18 ⁶Earth System Science Interdisciplinary Center, University of Maryland, College Park, MD, USA
- ⁷NASA Global Modeling and Assimilation Office (GMAO), Goddard Space Flight Center, Greenbelt,
 MD, USA
- 20
- 22 * Corresponding author: <u>a.souri@nasa.gov</u>
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24 Abstract

25 The tropospheric hydroxyl radical (TOH) is a key player in regulating oxidation of various compounds in Earth's atmosphere. Despite its pivotal role, the spatiotemporal distributions of OH are poorly constrained. 26 27 Past modeling studies suggest that the main drivers of OH, including NO₂, tropospheric ozone (TO₃), and 28 $H_2O(v)$, have increased TOH globally. However, these findings often offer a global average and may not 29 include more recent changes in diverse compounds emitted on various spatiotemporal scales. Here, we aim 30 to deepen our understanding of global TOH trends for more recent years (2005-2019) at 1×1 degrees. To 31 achieve this, we use satellite observations of HCHO and NO₂ to constrain simulated TOH using a technique 32 based on a Bayesian data fusion method, alongside a machine learning module named ECCOH, which is 33 integrated into NASA's GEOS global model. This innovative module helps efficiently predict the 34 convoluted response of TOH to its drivers/proxies in a statistical way. Aura Ozone Monitoring Instrument 35 (OMI) NO₂ observations suggest that the simulation has high biases over biomass burning activities in 36 Africa and Eastern Europe, resulting in overestimation of up to 20% in TOH, regionally. OMI HCHO primarily impacts oceans where TOH linearly correlates with this proxy. Five key parameters including 37 38 TO_3 , $H_2O(v)$, NO_2 , HCHO, and stratospheric ozone can collectively explain 65% of variance in TOH trends. 39 The overall trend of TOH influenced by NO_2 remains positive, but it varies greatly because of the 40 differences in the signs of anthropogenic emissions. Over oceans, TOH trends are primarily positive in the 41 northern hemisphere, resulting from the upward trends in HCHO, TO_3 , and $H_2O(v)$. Using the present 42 framework, we can tap the power of satellites to quickly gain a deeper understanding of simulated TOH 43 trends and biases.

44 **1. Introduction**

45 The hydroxyl radical (OH) regulates the lifetimes of a vast number of key atmospheric compounds, 46 such as sulfur dioxide (SO_2) , nitrogen dioxide (NO_2) , volatile organic compounds (VOCs), carbon 47 monoxide (CO), and methane (CH₄). Despite its outsized importance for atmospheric chemistry and 48 climate, our knowledge on both the abundance and long-term trends of OH is limited due to its sparse 49 observations, manifesting in large discrepancies between simulated OH among global models (e.g., Naik 50 et al., 2013; Zhao et al., 2019; Murray et al., 2021; Fiore et al., 2024). Particularly, these discrepancies can 51 introduce large uncertainties when it comes to precisely representing methane (Holmes et al., 2013; Nguyen 52 et al., 2020), a potent greenhouse gas. Consequently, to understand the potential impact of this warming 53 agent on climate shifts and extreme weather events, it is essential to accurately simulate methane 54 concentration within a coupled climate model, such as the NASA's Goddard Earth Observing System 55 (GEOS) model (Molod et al., 2015; Nielsen et al., 2017), which requires reasonable representation of its 56 major sink – reaction with OH.

57 Despite the challenges posed by OH's short lifespan of less than two seconds, low-pressure laser-58 induced fluorescence spectroscopy has proven invaluable in measuring OH for over twenty airborne field 59 campaigns (Miller and Brune, 2020). These datasets have been instrumental in verifying the efficacy of 60 chemical mechanisms involving varying reaction rate coefficients and aerosol heterogeneous chemistry 61 (Brune et al., 2019; Miller and Brune, 2020; Brune et al., 2022), understanding urban air quality (Brune et 62 al., 2022; Souri et al., 2023), as well as identifying potential sources of HO_x (OH+HO₂) that may have been 63 hampered due to instrument detection limits and/or unmeasured compounds (e.g., Ren et al., 2008). 64 However, while these observations offer valuable insights, they are limited in time and space and cannot 65 provide a full picture of tropospheric OH abundance.

There are several approaches that have been employed to constrain OH needed for replicating
observed values of a tracer whose primary sink is OH and its sources are relatively well known. One notable
method is methyl chloroform (MCF) inversion (Patra et al., 2014; Turner et al., 2017; Rigby et al., 2017;

Naus et al., 2019). However, this method only provides hemispheric-average OH and is thus insufficient toresolve the spatial distribution of OH.

71 A more sophisticated approach to constraining OH is to incorporate well-characterized satellite 72 observations of factors known to influence OH, such as NO₂, CO, ozone, and formaldehyde (HCHO), into a chemical transport model using inverse modeling and/or chemical data assimilation methods (Sandu and 73 74 Chai, 2011; Bocquet et al., 2015). This method offers a crucial advantage in that it accounts for the 75 interconnectedness of various chemical and physical processes within model increments. For example, 76 adjustments to NO_x levels will impact nitrate and ozone concentrations, which in turn affect the HO₂ uptake 77 through aerosols, OH, and radiation, reciprocally leading to a more accurate representation of NO_x. Several 78 studies have used subsets of satellite observations to improve HO_x and ozone chemistry, with Miyazaki et 79 al. (2020) using a diverse range of observations, including CO, NO₂, O₃, and nitric acid (HNO₃), to improve 80 model predictions using a local ensemble Kalman filter. The incorporation of these observations led to a 81 reduction in the asymmetric OH ratio between the northern and southern hemispheres, aligning better with 82 MCF results (Patra et al., 2014). Similarly, Souri et al. (2020a) leveraged well-characterized observations of HCHO and NO₂ to improve ozone chemistry over East Asia using non-linear analytical Bayesian 83 84 inversion, observing significant changes in OH levels after adjusting biogenic VOC in southeast Asia. 85 While incorporating these observations into atmospheric models offers a comprehensive way to gain 86 insights into spatiotemporal OH variability, it is complicated by several layers of complexity, such as 87 unidentified satellite biases, unresolved scales in satellite observations, and errors in models including 88 transport, chemical mechanisms, vertical diffusion, and depositions rates. Understanding how these errors 89 could cloud the realistic determination of OH requires running constrained models under various 90 realizations, which is computationally prohibitive.

91 Researchers have developed OH predictors based on a set of key parameters, offering reasonable 92 spatial and temporal coverage without compromising computational efficiency (Spivakovsky et al., 2000; Duncan et al., 2000; Elshorbany et al., 2016; Nicely et al., 2018; Wolfe et al., 2019; Nicely et al., 2020; 93 94 Anderson et al., 2022, Zhu et al., 2022; Anderson et al., 2023; Baublitz et al., 2023). These studies fall into 95 four categories, the first of which uses box model photochemical simulations to predict OH levels under a 96 steady-state assumption, using a blend of pre-modeled fields and various observations influencing OH (Spivakovsky et al., 2000; Nicely et al., 2018). The second group uses proxy observations (e.g., HCHO or 97 water, H₂O) of OH in remote areas (Wolfe et al., 2019; Baublitz et al., 2023). The third group employs 98 99 high-order polynomials to establish an empirical relationship between OH and different parameters, 100 avoiding the need to solve numerous differential equations in chemical mechanisms (Duncan et al., 2000; Elshorbany et al., 2016). Finally, the fourth group leverages powerful machine learning algorithms to 101 102 encapsulate the complexities between OH and its key influencers to efficiently predict OH using a 103 comprehensive dataset which is easily exchangeable between models (Nicely et al., 2020; Anderson et al., 104 2022; Zhu et al., 2022; Anderson et al., 2023).

105 In this work, we demonstrate the potential of a new approach to constrain simulated OH that uses 106 satellite observations to adjust the input parameters to an improved parameterization of OH (Anderson et 107 al., 2022), within the Efficient CH₄-CO-OH (ECCOH) (pronounced "echo") configuration (Elshorbany et 108 al., 2016) of NASA's GEOS model. We use the Modern-Era Retrospective analysis for Research and 109 Applications, Version 2 (MERRA2) reanalysis data (Molod et al., 2015) to constrain meteorology and 110 adjust two critical OH inputs using the latest Aura Ozone Monitoring Instrument (OMI) NO2 and HCHO 111 retrievals (Lamsal et al., 2021; Nowlan et al., 2023) from 2005-2019 worldwide. Through conducting a 112 range of experiments, we determine the extent to which leveraging OMI NO₂ and HCHO observations can 113 enhance current representations of these two species derived from a global model simulation, MERRA2-114 GMI (hereafter M2GMI) (Strode et al., 2019), so that we can achieve more accurate portrayals of OH 115 abundance and its long-term trends. Ultimately, we deconvolve the intricate OH trend maps into five critical 116 parameters using various modeling experiments, including tropospheric ozone, stratospheric ozone, NO₂, 117 HCHO, and H₂O.

118 Our paper is structured into several sections. In sections 2.1 to 2.3, we discuss the model 119 configurations, Bayesian data fusion algorithm, and satellite observations used. In section 2.4, we outline 120 our modeling experiments, which aim to uncover the impact of various key OH inputs on its trends and 121 assess the effect of OMI adjustments. In section 3.1, we examine the discrepancies between our prior knowledge from M2GMI and OMI observations and demonstrate how the data fusion can mitigate these 122 123 differences. In section 3.2, we delve into the effect of OMI adjustments to NO₂ and HCHO on tropospheric 124 OH (TOH) magnitudes across the globe. In section 3.3, we focus on understanding the long-term effect of 125 a set of key inputs on OH and how well they can replicate our most dynamic representation of TOH. In 126 Section 4, we summarize the potential of using satellite observations in conjunction with well-characterized 127 models to identify biases and long-term trends in TOH and discuss the limitations of our current analysis

128 and potential paths forward.

129 2. Models, Methods, and Measurements

130 2.1. Models

131 *2.1.1. GEOS*

132 The GEOS model (Molod et al., 2015; Nielsen et al., 2017) simulates global weather with 133 1° longitude $\times 1^{\circ}$ latitude spatial resolution. The model follows 72 hybrid sigma values ranging 134 from the surface to 0.01 hPa. We employ a cumulus parameterization to consider deep convection 135 (Moorthi and Suarez, 1992). Cloud microphysics is determined by a single-moment 136 parameterization based on Bacmeister et al. (2006). We activate the "replay" option (Orbe et al., 137 2017) to constrain several meteorological variables using the MERRA-2. Sea surface temperatures 138 and ice content are pre-described from various observations (Nielsen et al., 2017; Reynolds et al., 139 2007). Speciated aerosol concentrations and their optical properties are simulated by the GOCART 140 model (Chin et al., 2002) within GEOS. The rapid radiative transfer model for GCMs (RRTMG) resolves the long- and short-wave radiation imposed by GOCART-simulated aerosols, allowing for 141 142 the direct impact of aerosol on meteorology to be taken into consideration (Nielsen et al., 2017). 143 The period of simulation starts in 2005 and ends in 2020. Ten years before 2005 are considered for 144 the spin-up of meteorological, CO, and CH₄ fields.

145 *2.1.2. ECCOH*

146 A computationally-efficient module, named ECCOH was developed to simulate the 147 chemistry of the CH₄-CO-OH cycle in the GEOS-5 model framework (Elshorbany et al., 2016). 148 CO and CH₄ tracers are explicitly simulated and their emissions are discussed in Sections 2.1.2.1 149 and 2.1.2.2. A key component of ECCOH is the parameterization of tropospheric OH, which was 150 developed using a gradient boosted regression tree machine learning algorithm (Anderson et al., 151 2022) and is a function of chemical, solar irradiance, and meteorological variables. The training 152 dataset of chemical and meteorological variables was a 40-year daily M2GMI model simulation 153 (Strode et al. 2019), which includes tropospheric chemistry involving 120 species and 400 reactions 154 with the GMI mechanism (Duncan et al., 2007a and the references therein) and uses MERRA-2 155 reanalysis to constrain transport and meteorology at 0.625×0.5 degrees.

156 We present the variables used as inputs to the parameterization of OH for this study in 157 Table 1. The daily archived chemical inputs are from the M2GMI simulation with several variables 158 being constrained with observations. For instance, both NO₂ and HCHO fields are corrected 159 whenever satellite observations are available as described in Section 2.2.1. We chose NO₂, an 160 observable compound from satellites and a reasonable proxy for NO_x that has been shown to affect 161 OH (e.g., Zhao et al., 2020; Anderson et al., 2022). HCHO is used as a proxy for VOC oxidation 162 via OH in remote oceanic regions (Wolfe et al., 2019). 163 There are also long-term satellite data records of other OH drivers, including water vapor 164 (e.g., Aqua AIRS) and total ozone column (e.g., Aura OMI), that we could also consider. However, 165 the GEOS MERRA-2 system already assimilates satellite datasets of water vapor and the M2GMI 166 simulation simulates well (i.e., <4%) the total ozone column as compared to observations (Figure S1). The integrated water vapor columns from MERRA2 and microwave-based satellite 167 168 observations over-ocean also agree well (<5%), especially after 2000 when many satellite 169 observations have been used in the reanalysis data (Figure 3 in Bosilovich et al., 2017). Therefore, 170 the application of the "replay" mode constrains various meteorological fields, providing a more 171 realistic reconstruction of OH studied here.

Tropospheric ozone is another critical input to the parameterization of OH. Although we will compare M2GMI tropospheric ozone with satellite observations to locate any differences, reliable measurements of tropospheric ozone from satellites are lacking due to the limited sensitivity of the retrievals to ozone in low altitudes. Therefore, our study refrains from imposing any observational constraint on tropospheric ozone.

Throughout the paper, TOH is determined based on the methane-reaction-weighted OH suggested by Lawrence et al. (2001).

179 2.1.2.1. Monthly CO emissions

We use a modified version of EDGAR (Emissions Database for Global Atmospheric Research) v5.0 (Crippa et al., 2019), which is a comprehensive database that provides estimates of sector-based CO emissions from human activities (i.e., anthropogenic) on a global scale. Previous studies (e.g., Zheng et al., 2019) suggested a large underestimation of EDGAR CO emissions for India and China. Accordingly, we scale up the residential and transportation emissions from China by a factor of 1.6, and the residential emissions from India by a factor of 1.2 based on Zheng et al. (2019). The emissions spanned the entirety of the study period, from 2005 until 2020, and were prepared monthly at a spatial resolution of $0.1^{\circ} \times 0.1^{\circ}$. The daily biomass burning emissions are CMIP6 emissions, which derived from on the Global Fire Emissions Database version 4 with small fires (GFED4s) (van Marle et al., 2017). To account for the chemical production of CO from the oxidation of non-methane VOCs, we adopt the CO yield estimates from Duncan et al. (2007b) (i.e., a molar yield of 20% from isoprene, 20% from monoterpenes, 100% from methanol, 67% from acetone, 19% from anthropogenic VOC emissions, and 11% from biomass burning VOC sources) and released these CO emissions in the first vertical level of the model. With regards to the biogenic VOC emissions used for the above CO production estimates, we use offline MEGAN calculations using a GEOS-Chem (v13.2.0) run. CO production from CH₄ oxidation is calculated online for each model box.

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2.1.2.2. Monthly CH₄ emissions

In this study, several bottom-up CH₄ emissions related to anthropogenic, wetland, natural, and biomass burning sources are used to simulate CH₄. The monthly-basis anthropogenic sources are derived from EDGARv6 (Ferrario et al., 2021). The biomass burning emissions come from the GFED4s. Because EDGARv6 accounts for agricultural waste burning, we exclude this specific source from the GFED4. Following Strode et al. (2020), we use modified monthly-basis natural emissions from ocean, termite, and mud volcano emissions. Wetland emissions are derived from an improved dynamic wetland emission framework at $0.5^{\circ} \times 0.5^{\circ}$ based on the TOPography-based hydrological model (TOPMODEL) (Zhang et al., 2016; Zhang et al., 2023). A climatological sink of CH₄ from soil uptake is subtracted from the total CH₄ emissions.

Input Group	Variables (Directly Constrained)	Source	Temporal Resolution
Offline Chemical Species	NO₂, HCHO , O ₃ , isoprene, acetone, H_2O_2 , propene, propane, methyl hydroperoxide, ethane, C4 and C5 alkanes, and stratospheric O ₃ columns	M2GMI (offline) (Strode et al. 2019)	Daily- averaged
Online Chemical Species	CO and CH ₄	GEOS (online)	Daily- averaged
Meteorologic al Fields	T, P, Qv, and cloud fraction	GEOS (online)	Daily- averaged
Optical Properties	Aerosol optical depth; ice crystal cloud optical depth; and water droplet cloud optical depth at above and below of a given model vertical layer.	GEOS (online)	Daily- averaged
Geographic Information	Latitude and solar zenith angle (SZA)	Calculated	Fixed for latitude, bu daily for SZA based on local noontime
Surface Properties	Surface UV albedo	OMI LER climatology (Qin et al., 2019; Fasnacht et al., 2019)	Monthly (climatolo y)

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214 2.2. Methods

215 2.2.1. Bayesian data fusion for NO₂ and HCHO fields using OMI retrievals

To improve the representation of M2GMI NO₂ and HCHO concentrations and their long-216 217 term trends, which are used as input to the parameterization of OH in ECCOH, we scale their 218 columnar mass using Aura OMI observations of NO₂ and HCHO columns (described in Sections 219 2.3.1 and 2.3.2) using an offline version of the optimal interpolation (OI) method (Parish and 220 Derber, 1992; Jung et al., 2019) with an appropriate regularization. If we assume that the error covariances of M2GMI columns and OMI ones follow a Gaussian distribution with zero means and 221 their relationships are linear, we can estimate new columns using Bayes' theorem (Rodgers, 2000): 222 $\mathbf{X}_{a} = \mathbf{X}_{b} + \gamma \mathbf{B} \mathbf{H}^{\mathrm{T}} (\gamma \mathbf{H} \mathbf{B} \mathbf{H}^{\mathrm{T}} + \mathbf{E})^{-1} (\mathbf{Y} - \mathbf{H} \mathbf{X}_{b})$ where \mathbf{X}_{b} is the prior M2GMI columns (i.e., background), \mathbf{X}_{a} is the posterior M2GMI columns 223 224 (i.e., analysis), **B** is the error covariance matrix of the a priori, **E** is the sum squares of error 225 covariance matrix of the observations and the representation errors, Y is the observations, and H is 226 the observational operator which is equivalent to the identity matrix in our case. The instrument 227 error part of E is populated by the average sum of precision error squares the satellite product 228 provides. We interpolate both E and Y into the M2GMI grid box using a mass-conserved linear 229 barycentric interpolation method. In this method, both OMI observations and errors in the L2 230 granules provided at their irregular grid have been projected into a common grid of 0.25×0.25 231 degrees using Delaunay triangulation bi-linear interpolation. Subsequently, we convolve these re-232 gridded maps with a box filter whose kernel size is equivalent to the rounded fraction of M2GMI 233 grid box size to the re-grided OMI pixel size based on Souri et al. (2022). This interpolation method 234 removes the spatial representation error resulting from the unresolved scales in M2GMI columns. 235 Nonetheless, we did not take into account the errors of unresolved processes in M2GMI to augment 236 to E. The National Meteorological Center's (NMC) approach is a common technique for calculating 237 **B** in atmospheric models (Parish and Derber 1992; Souri et al., 2020b); however, due to computing 238 constraints, rerunning the M2GMI model to create the 24-hour prediction segments needed in the 239 NMC method was not possible. Instead, we initialize **B** by setting it to 50% errors for NO₂ and 240 HCHO, both of which are subject to regularization. γ is the regularization factor designed for 241 achieving the best fit (minimum residuals between Y and HX_b) while minimizing the effect of the 242 noise in the observations (minimum variance in X_a). To this end, we seek an optimal regularization 243 factor based on finding the "knee point" in the curve of the incremental regularization factors 244 (ranging from 0.1 to 10) and the degrees of freedom obtained from the optimization. The γ value 245 is determined based on the average of all data points in a month and does not vary from pixel to 246 pixel to ease the interpretation of the result. We did not account for the non-diagonal spatial 247 correlations of **B**, as it requires us to carry out the NMC method. We use the ratio of X_b/X_a to 248 uniformly scale the three-dimensional concentrations of the target gas (i.e., NO₂ or HCHO). The 249 error associated with the constrained M2GMI columns can be obtained via

 $\mathbf{S}_{\mathbf{a}} = (\mathbf{I} - \gamma \mathbf{B} \mathbf{H}^{\mathrm{T}} (\gamma \mathbf{H} \mathbf{B} \mathbf{H}^{\mathrm{T}} + \mathbf{E})^{-1} \mathbf{H}) \times \gamma \mathbf{B}$

(2)

The averaging kernels (AK) describe the amount of information gained from the observations are represented by

$$AK = I - \frac{S_a}{B}$$
(3)

where **I** is the identity matrix.

In our research, we have created an open-source Python package called OI-SAT-GMI (Souri, 2024), which possesses the ability to download and process OMI level 2 products, perform air mass factor (AMF) recalculation, and conduct mass-conserved interpolation, while also executing the OI algorithm.

In our approach, the adjustments are implemented to the M2GMI output (i.e., a data fusion approach instead of data assimilation one), thereby restricting the full use of improved NO₂ and HCHO representation for more accurate simulation of other chemical compounds impacted by NO₂ and HCHO, including ozone (e.g., Souri et al., 2020a, 2021). Nevertheless, as the accuracy of NO₂ concentrations can significantly impact OH and HCHO is strongly tied to VOC oxidation through OH in remote ocean areas (Wolfe et al., 2019), the adjustments are expected to be beneficial in achieving a more robust representation of OH.

264 2.2.2. Trend analysis

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We determine a linear trend in a time series based on fitting the following equation accounting for a seasonal cycle and shorter frequencies in the observations:

$$\mathbf{y} = a_0 + a_1 \mathbf{t} + \sum_{i=1}^3 a_{i+1} \cos 2\pi \omega_i (\mathbf{t} - \varphi_i)$$
(4)

The equation comprises several variables, including y (data points) on monthly-basis, a_0 as the mean, a_1 as the linear trend, t as time (fractional year), a_{i+1} , ω_i , and φ_i are the amplitude, frequency, and phase, respectively. We consider three harmonics ($\omega_i = 1,2,3$) to account for seasonal cycle (ω =1) and higher frequencies. To assess the statistical significance of a trend, we employ the Mann-Kendall test and consider a trend to be significant if the linear trend passes the test at a 95% confidence level.

In the context of trend analysis, the careful examination of errors in observations (y) is a critical aspect often overlooked. However, when the errors of observations are obtainable, such as those obtained from satellites or constrained M2GMI fields, we determine the parameters by applying a weighted estimation. This estimation is optimized using the Levenberg–Marquardt algorithm (Marquardt et al., 1996) using *SciPy* open-source package. Considering the errors in the observational data deemphasizes more uncertain data, resulting in a more realistic determination of the linear trend.

280 2.2.3. OH response calculations

281 To elucidate the response of OH to different input parametrizations, such as NO_2 , HCHO, 282 and O_3 , we determine the semi-normalized sensitivities through a traditional finite difference 283 method:

$$SOH_i = \frac{[OH]_i^{110\%} - [OH]_i^{90\%}}{0.2}$$
(5)

284 where $[OH]_i^{110\%}$ and $[OH]_i^{90\%}$ are OH concentrations from perturbing input parameters (*i*) by 1.1 285 and 0.9 scaling factors in the ECCOH offline framework (Anderson et al., 2022). These calculations 286 are solely used to better understand why OH changes in a particular way relative to the changes in 287 its drivers. In our online modeling framework, OH is simultaneously affected by the dynamic 288 changes of various variables considered in the parametrization of OH.

289It is crucial to acknowledge that ECCOH has established an implicit relationship between290OH and various input parameters statistically. These perturbations could involve a range of291physiochemical processes that are challenging to fully decipher. For example, the perturbation of292NO2, acting as a surrogate of reactive nitrogen, involves chemical reactions that include reactive293nitrogen like NO+HO2 and NO2+OH, ozone formation, aerosol HOx uptake, and radiation.294Nonetheless, it may not be feasible to understand the extent to which ECCOH has been considered.295Therefore, the presented perturbations in this work should be viewed qualitatively.

296 2.3. Measurements

297 2.3.1. OMI MINDS tropospheric NO₂ columns

To improve the representation of NO₂ fields used as input to the parameterization of OH, we constrain the archived monthly fields with the most updated NASA standard tropospheric NO₂ product (v4.0; Lamsal et al., 2021) from Aura OMI. Aura has a local equatorial overpass time of 13:45 and nearly daily global coverage. This new OMI product version is improved in multiple aspects as compared to the former products, including surface reflectance and cloud retrieval (Lamsal et al., 2021).

304 The validation of OMI tropospheric NO₂ columns from the comparison to integrated 305 aircraft spirals obtained from diverse air quality campaigns revealed a good level of correlation 306 (r>0.7) (Choi et al., 2020). However, large mean biases, approximately 40%, were observed. These 307 biases come from various sources, including systematic biases in prognostic data utilized in the 308 retrieval, biases inherent in the aircraft data, spatial representation errors (Judd et al., 2020; Souri 309 et al., 2022), and temporal representation errors. The spatial representation errors have been 310 recognized to notoriously drift the slopes from the unity line in validation studies (Souri et al., 311 2022). Notably, Choi et al. (2020) achieved a substantial reduction in mean biases, decreasing from 312 40% to 16%, through the downscaling of OMI data into a finer resolution domain using a regional 313 chemical transport model. Likewise, Pinardi et al. (2020) reduced the biases between MAX-DOAS and OMI NO₂ observations by considering a radial dilution factor to account for the mismatch scales between the satellite footprint and the pointwise observations. These studies showed that the true statistics describing OMI biases are unknown, but they tended to be milder than those derived from directly comparing large pixels with pointwise measurements. It is important to highlight that discrepancies between M2GMI and OMI NO₂ will surpass the reported biases, thereby underscoring the product's reliability over diverse geographical regions.

The long-term trends of tropospheric NO₂ columns have undergone extensive comparative
analyses with in-situ observations (Lamsal et al., 2015; Pinardi et al., 2020), regulatory inputs, and
assessments of human and biomass burning activities (Duncan et al., 2016; Choi and Souri,
2015a,b; Krotkov et al., 2016; Jin and Holloway, 2015; Souri et al., 2017; Rueter et al., 2014; de
Foy et al., 2016; Hickman et al., 2021).
We prefer level 2 over level 3 products to enable the recalculation of AMFs with time-

We prefer level 2 over level 3 products to enable the recalculation of AMFs with timevarying shape factors derived from the M2GMI simulation. We removed low-quality pixels using the main quality flag, cloud fraction >30%, terrain reflectivity >20%, and those pixels affected by the "row anomaly" complication. The data product, which has a spatial resolution ranging from ~13 km × 24 km (at nadir) to ~24 km × 160 km (at extremities of the scanline), were then regridded to the M2GMI grid (0.625°×0.5° degrees) using a mass-conserved linear barycentric interpolation method. The AMF recalculation was performed via:

$$VCD_{new}^{\square} = \frac{VCD_{old}^{\square}AMF_{old}^{\square}}{AMF_{new}^{\square}}$$
(6)

332 where VCD_{old} and AMF_{old} are the default states of tropospheric vertical columns and air mass 333 factors. AMF_{new} is determined by summing the product of scattering weights and the M2GMI partial 334 columns from the surface to the tropopause level prescribed in the OMI level 2 data.

335 2.3.2. OMI SAO total HCHO columns

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336 For the same reason as OMI NO₂, we use OMI SAO total columns based on a newly-337 developed algorithm framework by Nowlan et al. (2023). The new retrieval represents a major step 338 forward in the surface albedo treatment including the bidirectional reflectance distribution function 339 for land (BRDF) from the MODIS product (MCD43C1 Version 6.1) extended to the UV 340 wavelengths using a principal component algorithm. Since there are no MODIS BRDF data 341 available over water, the algorithm uses the Cox-Munk slope distribution to estimate the surface 342 reflectance of water bodies (Cox and Munk, 1954). An important issue with the long-term record 343 of OMI HCHO measurements is the artificial increasing trend brought on by sensor degradation 344 (Choi and Souri, 2015a,b, Gonzalez Abad et al., 2015). The algorithm uses an earthshine spectrum 345 over the Pacific Ocean with a latitudinal and solar zenith-dependent correction factor described in 346 Nowlan et al. (2023) to mitigate this artifact.

347The new SAO algorithm has been validated with Ozone Mapping and Profiler Suite348(OMPS) data radiance with respect to Fourier-transform Infrared Spectroscopy (FTIR) in-situ349measurements in 2012-2020, showing a relative bias of 30% based on monthly-averaged data350(Kwon et al., 2023). While the validation results based on the OMI radiance have not been released351yet, it is likely for the biases to stay at roughly the same range of errors at monthly-gridded OMI352data onto the M2GMI grid which is comparable to the OMPS footprint (50 km).

353Once again, we used Eq.6 to recalculate OMI HCHO total columns with dynamical shape354profiles produced during the M2GMI simulation. We remove unwanted pixels using the following355criteria: the main quality flag, cloud fraction >40%, and flag for pixels affected by the row anomaly.356We then regridded the data to the MERRA-2 GMI grid using the same approach used for OMI357NO2.

358 2.4. Experiments

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We perform a series of experiments to investigate the sensitivity of OH to geophysical variables known to influence or to be tied with OH. Table 2 lists all sensitivity experiments along with their purposes and differences from an analysis (i.e., constrained) experiment. The pillar of all experiments is the analysis experiment (*Sanalysis*) which uses i) chemical variables from a full-chemistry simulation as input to the parameterization of OH in ECCOH (Section 2.1.2; Table 1); ii) transport and metrological fields constrained by MERRA2 reanalysis data (Section 2.1.1); iii) long-term estimates of monthly CO and CH₄ emissions (Section 2.1.2.1 and 2.1.2.2); iv) optical depths of clouds and aerosols along with observed climatology of OMI UV surface albedo; and v) the NO₂ and HCHO fields constrained by the Bayesian data fusion method (Section 2.2.1).

To examine the importance of having NO₂ and HCHO fields constrained with OMI data, we design three experiments imitating *Sanalysis*, but withholding the OI scaling factors one at a time. We then subtract these model outputs from those of *Sanalysis* and name them as *SOMInitro*, *SOMIform*, and *SOMInitroform*.

372 The other experiments are intended to systematically isolate the chemical effect of a 373 specific driver/proxy of OH trends. Due to the significant impact of NO₂, tropospheric ozone, 374 stratospheric ozone column, and water vapor on the primary or secondary pathways of OH 375 loss/production (Naik et al., 2013; Murray et al., 2013; Strode et al., 2015; Nicely et al., 2018; Zhao 376 et al., 2020; Anderson et al., 2021), we include four experiments (SOHwv, SOHnitro, 377 SOHtropozone, and SOHstratozone) to single out each effect on OH trends. Additionally, we 378 include HCHO (SOHform), a robust proxy for VOC oxidation via OH in remote ocean regions 379 (Wolfe et al., 2019) to understand how those chemical pathways have changed over time. In these 380 experiments, we set the target driver constant to the monthly values in the first year of simulation, 381 and subsequently subtract these model outputs from Sanalysis. Amongst various OH 382 drivers/proxies studied here, water vapor is simulated online based on the GEOS simulation; to 383 conduct SOHwy which aims at isolating the water vapor effect on OH without affecting 384 meteorology, we set water vapor fields fed to the parametrization of OH to the offline MERRA2 385 based on the monthly-varying 2005 simulations. Simultaneously, GEOS is allowed to simulate 386 water vapor online to address meteorology. This ensures that the meteorology remains consistent 387 across both SOHwv and Sanalysis.

Using ambient gas concentrations in the ECCOH model poses a challenge in distinguishing
 the respective factors contributing to their variations. For instance, it is difficult to discern the
 distinct influences of lightning-produced NO₂ versus anthropogenic NO₂ on the abundance of OH.
 However, an advantageous feature of our approach is that various observational sources constrain
 the data fields used via the Bayesian data fusion method or MERRA2 reanalysis data.

Table 2. The experiments designed to assess the effect of various OH drivers/proxies and OMI constraints on TOH trends and magnitudes.

Model Scenario	Term	Difference from the analysis run	Purpose
Analysis (constrained)	Sanalysis		The "best effort" to simulate the evolution of the CH ₄ -CO-OH cycle from 2005-2019.
Analysis - ^a defaulting to NO ₂ M2GMI	SOMInitro	Uses archived M2GMI monthly- averaged NO ₂ concentration fields.	Isolate the importance of constraining M2GMI NO ₂ concentration fields with OMI observations.
Analysis - defaulting to HCHO M2GMI	SOMIform	Uses archived M2GMI monthly- averaged HCHO concentration fields.	Isolate the importance of constraining M2GMI HCHO concentration fields with OMI observations.
Analysis - defaulting to NO ₂ and HCHO M2GMI	SOMInitroform	Uses archived M2GMI monthly- averaged NO ₂ and HCHO concentration fields.	Isolate the importance of constraining M2GMI NO ₂ and HCHO concentration fields with OMI observations.
Analysis - fixed H ₂ O vapor	SOHwv	The dynamical water vapor fields fed to the parameterization of OH are fixed to the monthly-varying 2005.	Isolate the impact of the long-term trend of water vapor on OH.
Analysis - fixed tropospheric ozone	SOHtropozone	M2GMI ozone fields are set to the monthly-varying 2005.	Isolate the impact of the long-term trend of tropospheric ozone burden on OH.
Analysis - fixed NO ₂	SOHnitro	M2GMI NO ₂ fields are set to the monthly-varying 2005.	Isolate the impact of the long-term trend of NO ₂ on OH.
Analysis - fixed HCHO	SOHform	M2GMI HCHO fields are set to the monthly-varying 2005.	Understand the long-term trend of HCHO strongly tied with VOC oxidation via OH in remote regions.
Analysis - fixed stratospheric ozone column	SOHstratozone	M2GMI stratospheric ozone field fed to the parameterization of OH is set to the monthly-varying 2005.	Isolate the impact of the long-term trend of stratospheric ozone columns on OH.

395 ^a "-" denotes the subtraction operator.

396 3. Results and Discussion

397 3.1. Spatial distributions and trends analysis of several inputs to the parameterization of OH

We begin our analysis with an examination of the long-term trends and magnitudes of two key inputs
 (HCHO and NO₂) to the parameterization of OH. Some other key parameters, such as total ozone columns,
 tropospheric ozone columns, and water vapor are also shown in Figure S1-3, Figure S7-8, and Text S1.

401 *3.1.1. Tropospheric NO*₂ columns

We performed two sets of comparisons; the first comparison involves examining the differences in the
 tropospheric NO₂ columns in the M2GMI relative to those of OMI before and after applying the OI correction.
 The second comparison focuses on the global 2-D maps of long-term linear trends of OMI, M2GMI prior to
 and after the Bayesian data fusion correction synched at the satellite viewing condition.

Figure 1 demonstrates the absolute difference in M2GMI tropospheric NO_2 columns with respect to those of OMI before (the a priori) and after (the a posteriori) the data fusion application along with AK in 2005-2019. In-land regions show positive biases over several regions, including central Africa (box A), the Midwest U.S. (box B), and Europe (box C). The same tendency was observed in Anderson et al. (2021). The largest contributor to NO_2 in box A and box C is biomass burning activities (Jaeglé et al., 2005; Giglio et al., 2012), suggesting that either the emission factors and/or the total dry mass burnt were possibly too high in these regions.

413 M2GMI overestimates NO_2 concentrations in non-urban areas in box B which tend to be more severe 414 during summertime. Although soil NO_x emissions could be the first explanation for this phenomenon, 415 accounting for about 30% of tropospheric NO_2 columns in the region according to Vinken et al. (2014), the 416 soil NO_x parameterization used in M2GMI relies on Yienger and Levy (1995), which is known to have a low 417 bias (Jaeglé et al., 2005; Hudman et al., 2012; Vinken et al., 2014; Souri et al., 2016). Therefore, there may be 418 other uncertainties in the model concerning chemistry (e.g., Canty et al., 2015) or area anthropogenic NO_x 419 emissions (Hassler et al., 2016) causing the bias.

A large portion of metropolitan areas in the Middle East, Europe, and the U.S. shows an 420 421 underestimation of NO₂ in M2GMI. Moreover, OMI observations reveal large positive biases over the North 422 China Plain (NCP), a region exhibiting exceptionally high NO₂ levels (e.g., Duncan et al., 2016; Krotkov et 423 al., 2016; Souri et al., 2017). This is primarily because of not accounting for the recent aggressive emissions 424 mitigation in China in the bottom-up emission inventory used in the model. We observe several regions over 425 China and Yellow Sea underestimating NO₂ with respect to OMI observations that do not improve 426 considerably after the adjustments. This tendency is a result of the use of a fractional error for populating the 427 error covariance matrix of the a priori, rendering the prior error too low. Although we used a regularization 428 factor to battle this problem, it did not vary from region to region. A regionally-adaptive regularization factor 429 could be a possible remedy for this problem but at a cost of overcomplicating the interpretation of the results.

430 Expectedly, the Bayesian fusion greatly mitigates the regional biases, with notable reductions 431 observed over central Africa, China, the U.S., Amazon, and Europe. The regional biases (>80%) well exceed the reported biases associated with OMI tropospheric NO_2 product (<40%), suggesting that the adjustments 432 433 should be considered as improvement. Nonetheless, it is important to acquire an abundance of long-term 434 records from surface spectrometers such as MAX-DOAS and Pandora to comprehensively evaluate the degree 435 of enhancement of M2GMI constrained by OMI within the troposphere, which is currently unavailable for the 436 period of 2005-2019 to our best knowledge. The reduction in the biases over remote areas in the tropics is less 437 noteworthy due to large errors in the observations. In other words, it is difficult to have high confidence in the 438 degree of deficiency the model can have in simulating NO_2 over pristine areas by comparing it to OMI. This 439 notion mathematically manifests in low AK in remote areas showing that rich information from OMI tropospheric NO₂ gravitates more towards polluted regions. This finding assumes that the regularized 440 441 covariance matrix of the prior error does not substantially vary between land and ocean and is isotropic.

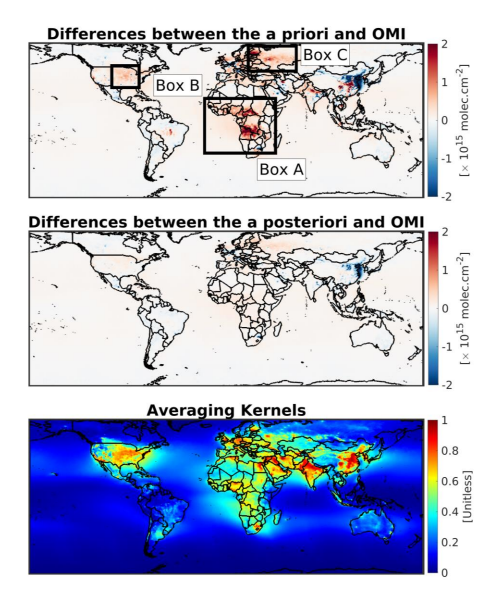


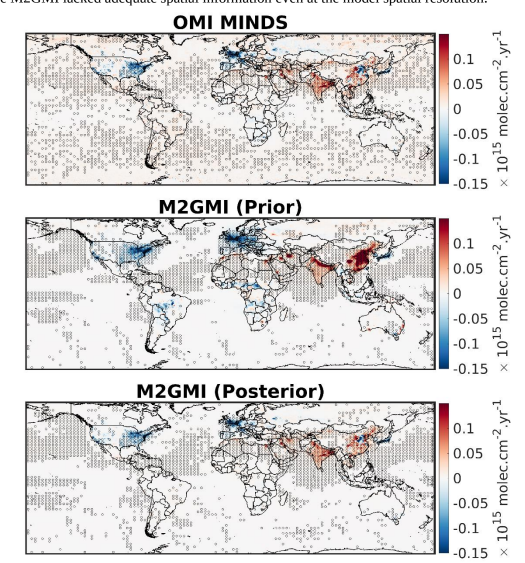
Figure 1. The global maps of M2GMI tropospheric NO₂ annual difference with respect to those of OMI before applying the Bayesian data fusion correction factors (top) and after (middle) in 2005-2019; the mean of averaging kernels describing the gained information from OMI (bottom). Grids in high latitudes are removed from the figure due to too few numbers of samples OMI provided.

447

448 Figure 2 illustrates the linear trends of tropospheric NO₂ between 2005 and 2019 observed by OMI 449 and simulated by the M2GMI before and after using the OI algorithm. The errors in OMI observations and the 450 constrained M2GMI are considered while calculating the trends. Focusing on the trends by OMI, we observe 451 a consistent picture compared to former studies (Duncan et al., 2016; Choi and Souri, 2015a,b; Krotkov et al., 2016; Jin and Holloway, 2015; Souri et al., 2017). High income countries, such as the U.S., those located in 452 453 the western Europe, and major cities in Russia, undergo a significant reduction of NO₂ concentrations due to 454 the implementation of emission mitigation regulations. Additionally, low and moderate income countries, such 455 as those in the Middle East, northern Africa, and India, have seen upward trends in NO₂. Various signs of 456 trends are observed in East Asia. Due to recent effective regulations in China (Zhang et al., 2012), we observe downward trends in the NCP region (Rueter et al., 2014; de Foy et al., 2016; Souri et al., 2017). The downward 457 458 trend predominantly starting from 2011-2012 counteracts the upward trend in prior years resulting in

459 statistically insignificant linear trends. Both Japan and South Korea show downward trends during the period460 of 2005-2019 (Duncan et al., 2016; Souri et al., 2017).

461 Encouragingly, the model prior simulation of the tropospheric NO_2 trend is consistent with OMI over 462 most of the polluted regions except for China, where the bottom-emission inventories used in the M2GMI fail 463 to reflect recent mitigation efforts occurring in NCP region. The posterior estimation is in a higher degree of 464 agreement compared to OMI (Text S2). An encouraging observation arising from the comparison of the 465 M2GMI prior with the posterior NO_2 trends is the achievement of a higher spatial variance (information) in 466 low and medium income countries (e.g., India and Iran). This finding suggests that the emission inventories 467 used in the M2GMI lacked adequate spatial information even at the model spatial resolution.



468

Figure 2. The global maps of linear trends of annual tropospheric NO₂ columns observed by OMI and
simulated by M2GMI before and after using the Bayesian fusion. The model simulations are sampled at the
exact time and location of OMI, and masked if OMI observations were unavailable due to data quality criteria
used. The dots indicate statistically significant trends at 95% confidence interval.

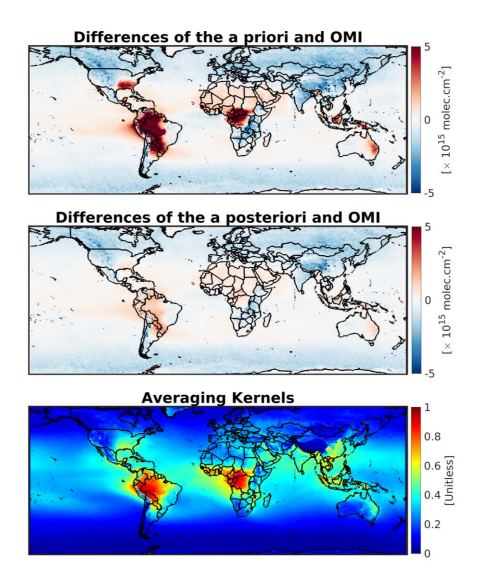
473

474 *3.1.2. Total HCHO columns*

475 We validate the simulated HCHO concentrations, drawing inspiration from the NO₂ comparison 476 framework. Figure 3 illustrates the absolute differences in simulated HCHO total columns with respect to OMI 477 before and after the Bayesian data fusion application, in addition to AK. The prior model simulation has 478 considerable skill in capturing the HCHO total columns over several areas, such as the Middle East, Europe, 479 India, and East Asia. However, marked positive biases are discernible in regions with abundant isoprene 480 emissions, such as the Amazon, southeast Asia, southeast U.S., and central Africa. This outcome is most likely 481 due to an overestimation of biogenic emissions; various investigations have reported a predominantly positive 482 bias (between a factor of 2 to 3) linked to isoprene emissions estimated by the Model of Emissions of Gases 483 and Aerosols from Nature (MEGAN) using satellite measurements in isoprene-rich regions (e.g., Millet et al., 484 2008; Stavrakou et al., 2009; Marais et al., 2012; Bauwens et al., 2016; Souri et al., 2020a).

485 The simulated HCHO concentrations are relatively too low over pristine areas, such as high latitudes and over mountains. This may be attributed to an underestimation of CH₄ in M2GMI because of assigning its 486 487 values as background conditions (Strode et al. 2019). The integration of OMI satellite data has proven effective 488 at reducing the biases in areas where HCHO concentrations are large because the signal-to-noise ratio tends to 489 be large resulting in high AKs. Nonetheless, there are some adjustments over remote areas. In fact, OMI HCHO columns provide more information than OMI NO₂ in remote areas because background HCHO concentrations 490 are not extremely low due to evenly distributed methane and methanol concentrations. It is worth noting that 491 the biases in M2GMI well exceed the expected OMI HCHO column biases, suggesting that the adjustments to 492

493 HCHO improve the model.



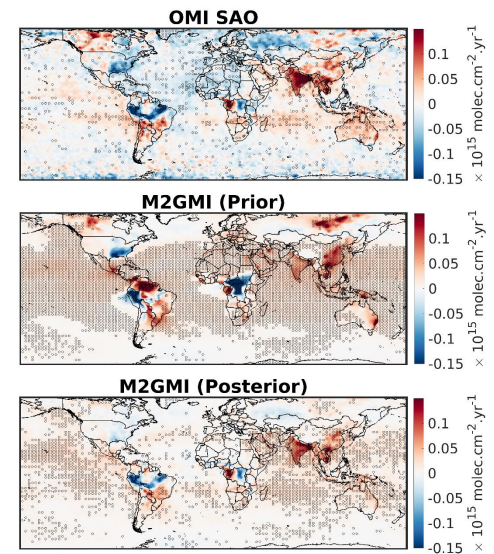
495 Figure 3. Same as Figure 1 but for HCHO total columns.

496 Figure 4 shows the global maps of HCHO total column trends derived from OMI, the prior M2GMI, 497 and the posterior M2GMI. The widespread upward trends in HCHO over India are evident due to lack of 498 effective efforts on cutting emissions related to volatile organic compounds (e.g., De Smedt et al., 2015; 499 Kuttippurath et al., 2022; Bauwens et al., 2022). We observe HCHO columns going up in the northwestern US 500 and over oil sands in Canada, possibly due to increased evergreen needleleaf forest and an increase in crude oil production (Zhu et al., 2017), respectively. The downward trends over the southeast US could be due to a 501 502 decrease in drought events (Figure S5), which significantly affect isoprene emissions and the oxidation of VOCs (Duncan et al. 2009; Naimark et al., 2021; Wang et al., 2022). Alternatively, this downward trend could 503 504 be partially due to the dampened HCHO production from VOC oxidation due to reduced NO_x emissions 505 (Marais et al., 2014; Wolfe et al., 2016; Souri et al., 2020c). In agreement with previous studies (Stavrakou et al. 2017, Souri et al., 2017, Shen et al., 2019, Souri et al., 2020a), HCHO columns increase over the NCP. 506 507 HCHO columns tend to decrease over parts of central Africa (e.g., Democratic Republic of the Congo) and the 508 Amazon basin potentially due to reduced deforestation rates (De Smedt et al., 2015; Jones et al., 2022). 509 However, a large variability in the sign of HCHO trends over these regions is seen; Congo shows an opposite 510 trend in comparison to that of Democratic Republic of the Congo; the northern portion of the Amazon basin 511 increased. Encouragingly, the prior knowledge captures the upward trends over India and China along with

downward trends over central Africa. However, the magnitudes and spatial features of these trends are notentirely in line with respect to OMI.

We do not fully understand HCHO trends over oceans. Part of these patterns might be caused by transport from nearby sources. For instance, areas around south Asia, South America, and Gulf of Mexico can be affected by the trends over the land in their proximity. However, trends over several areas, such as the southern part of the Indian Ocean, Australia, and Sahara, are not fully explainable by nearby sources. It is possible that certain patterns can be linked to climate variability or OH (Wolfe et al., 2019) affecting the oxidation of background VOCs; an in-depth understanding of HCHO trends over oceans certainly deserves a separate follow-up study.

The posterior estimates better line up with the OMI trends, especially over the Amazon, India, and Central Africa (Text S3). The correction factors, however, worsen the trends over the southeast US and Canada. One possible explanation for this may be the varying errors from the data fusion algorithm, which tend to be reduced more in summertime than in wintertime due to the larger OMI HCHO signal. This results in some degree of inconsistencies of the linear trend over these regions with larger interannual and interdecadal variabilities.



527

Figure 4. Same as Figure 2 but for total HCHO columns. The linear trends in OMI SAO are smoothed by amedian filter for better visualization.

530

531 In sum, we saw that M2GMI NO₂ and HCHO, both inputs to the parameterization of OH, were broadly 532 better presented through the integration of OMI observations. Consequently, the improvement is expected to elevate the level of reliability in the experimental outcomes, particular in the context of SOHnitro and 533 534 SOHform simulations. As for other important compounds, such as stratospheric columns, tropospheric O₃, and 535 water vapor, the comparison of the model with OMI total ozone columns shows a strong degree of agreement (<4% biases) with no significant trend in low-mid latitudes (Figure S1 and S2). The well-documented upward 536 trend in tropospheric ozone in the northern hemisphere is well reproduced by M2GMI (Figure S3). We did not 537 538 validate GEOS water vapor simulations, because of the use of MERRA2 reanalysis, which is thoroughly 539 validated in Bosilovich et al. (2017). Furthermore, the comparison of integrated water vapor linear trends from 540 our GEOS-5 run (2005-2019) with satellite data presented in Borger et al. (2022) shows a remarkable 541 agreement (Figure S7-8).

542 3.2. Added value of OMI on simulated tropospheric OH

543 Here, we present the results from three OMI-related experiments (*SOMInitro*, *SOMIform*, 544 *SOMInitroform*) to understand the effect of OMI adjustments made to M2GMI on TOH. Moreover, we 545 calculate the response of TOH to NO_2 and HCHO using Eq.5.

546 Figure 5 consists of three columns, illustrating the percentage adjustments made by OMI NO₂ using 547 OI, the response of TOH to NO₂ concentrations, and the simulated TOH derived from the SOMInitro 548 experiment. The observed pattern of increments aligns with the improvements seen in Figure 1, with positive (negative) values indicating underestimation (overestimation) of M2GMI. Broadly, the overestimates 549 550 dominate over underestimates resulting in the global tropospheric NO₂ reduction by $\sim 4\%$. Upon segregating 551 the increments into four distinct seasons, it becomes evident that the adjustments do not uniformly apply to 552 every season. This non-uniformity is primarily attributed to biases in M2GMI, influenced by biomass burning 553 (box A, C) (Section 3.1.1), both of which exhibit strong seasonality.

Deciphering the precise chemical processes influencing the response of OH to NO₂ using a machine-554 learning approach is challenging. However, it is widely recognized that reactive nitrogen has positive feedback 555 on tropospheric OH through increased NO+HO₂ and ozone (Murray et al., 2021; Zhao et al., 2020; He et al., 556 2021). Considering NO₂ as a surrogate for reactive nitrogen, similar tendencies are expected, as evident from 557 558 the positive numbers from the sensitivity results obtained from offline calculations. The response of TOH to NO₂ displays a pronounced seasonal cycle stemming mainly from photochemistry. It is believed to have some 559 negative values for the sensitivity of OH to NO₂ for extremely polluted regions due to radical termination 560 through NO₂+OH or ozone titration (Nicely et al., 2018). While we have not identified any negative values in 561 562 the tropospheric domain, we have observed significant negative values of OH when perturbing NO_2 at the model surface layer (Figure S26). This tendency highlights the ECCOH's ability to account for non-linearities. 563

The impact of adjustments made by OMI NO₂ on TOH is most substantial over regions where both the adjustments and TOH responses to NO₂ are significant. For instance, the large adjustments made over Europe in DJF do not substantially affect TOH because the response value is low due to reduced photochemistry.

568 On a global scale, changes to TOH are much milder (1% reduction) than those occurring regionally. 569 For instance, we see substantial regional impacts (up to 20%) over many areas such as Central Africa, the 570 Midwest US, the Middle East, and Eastern Europe. In light of the global reduction in OH, we observe global 571 column average methane mixing ratios (XCH₄) to increase by 10 ppbv on average (Text S4). This 572 augmentation happens monotonically with an increase of 0.9 ppbv per year, ultimately resulting in ~15 ppbv 573 difference at the end of the simulation (Figure S13). This is essentially due to the long lifetime of CH₄. 574 Likewise, the TOH reduction results in column average CO mixing ratio (XCO) enhancements which transpire 575 more locally than XCH₄ does due to the shorter XCO lifetime. The XCO enhancements reach above 10 ppbv

576 in Africa (Text S5).

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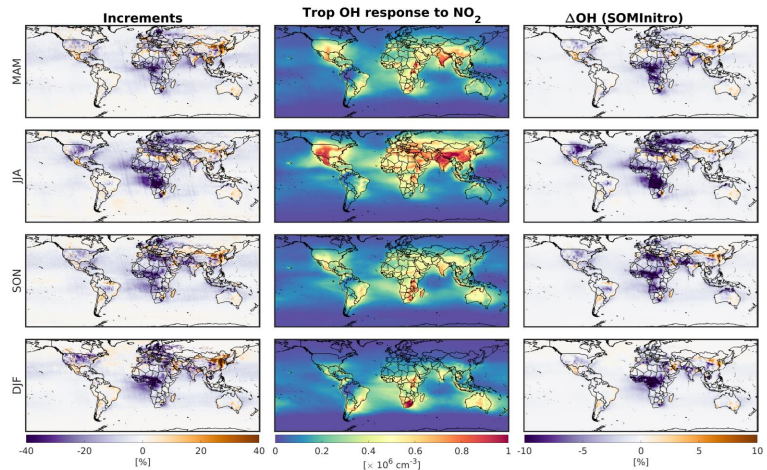


Figure 5. (first column) the percentage of adjustments applied to M2GMI NO₂ fields within the troposphere suggested by OMI tropospheric NO₂ columns for four different seasons, (second column) the semi-normalized response of tropospheric OH to tropospheric NO₂ changes based on ECCOH offline calculations, and (third column) the resulting effect of the adjustments on tropospheric OH derived from the online simulation (*SOMInitro*). MAM, JJA, SON, and DJF are acronyms for March-April-May, June-July-August, September-October-November, and December-January-February.

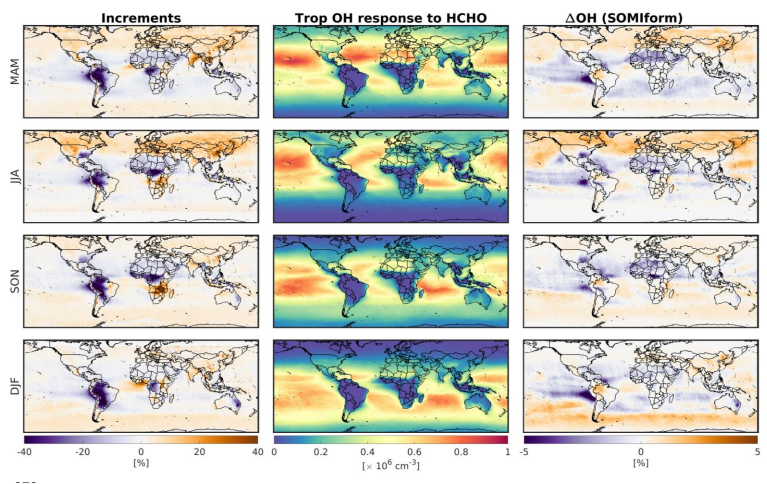
Figure 6 demonstrates the same scheme as Figure 5 but with a focus on the *SOMIform*. Marked negative increments are found in regions characterized by elevated isoprene concentrations because of the overestimations of M2GMI biogenic isoprene emissions. Positive increments are mostly confined to high latitudes and certain areas of East Asia (Section 3.1.2).

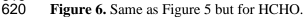
The interplay between HCHO and OH is contingent on the intricate dynamics governing HCHO production from the oxidation of VOCs and methane and HCHO loss from various chemical pathways (Valin et al., 2016; Wolfe et al., 2019). In remote areas where HO_x is low, the prevailing sink of HCHO is through photolysis. Conversely, in more polluted areas, the reaction of HCHO+OH emerges as a competing loss pathway. Assuming a steady-state approximation, which is a reasonable assumption for pristine areas, the photolysis loss of HCHO dominates over the reaction with OH, resulting in a linear relationship between HCHO and OH. In other words, high (low) HCHO concentrations are indicative of high (low) TOH. It is because of this that we use HCHO as a proxy of TOH in remote oceans regions. In regions characterized by heightened HO_x levels, OH and HCHO become decoupled. Encouragingly, our implicit parametrization of OH has considerable skill at elucidating these intricate chemical tendencies; specifically, it reveals muted responses in regions with relatively tangible pollution levels, whereas positive responses are evident in oceanic regions. Like results obtained for NO₂, the response map has a seasonal cycle due to photochemistry.

Because of the muted response of TOH to HCHO over land, a substantial portion of geographical
regions undergoing significant adjustments made by OMI becomes less important. TOH primarily changes
over oceanic areas in a way that it decreases in low latitudes but increases in high latitudes. The largest
reduction occurs in Amazon downwind where both increments and responses display large magnitudes. As a
result of these changes, we see a marginal increase in XCH₄ over tropics where OMI increments reduced TOH.
The HCHO adjustment did not noticeably affect XCO either (Text S5).

Modifications on HCHO by OMI do not signal substantial changes in background VOC oxidation
through OH. In fact, TOH changes by this proxy are of an order of magnitude less than those by OMI NO₂.
This tendency is a result of two key factors: i) the adjustments wield their major influence over oceans where
M2GMI has a fair performance, and ii) the amount of information obtained from OMI HCHO (i.e., AK)
remains somewhat limited in remote areas due to low signal-to-noise ratios.

614 Due to the rather independent nature of the TOH responses to NO_2 and HCHO, where the former 615 prevails over land and the latter over ocean, the concurrent adjustments of HCHO and NO_2 using OMI (i.e., 616 *SOMInitroform*) results in a rather linear combination of outcomes derived from *SOMIform* and *SOMInitro* 617 (Figure S21). This linear outcome is characterized by a large decrease in TOH in low latitudes and a moderate 618 increase in high latitudes resulting in a decrease of global TOH by ~1%.





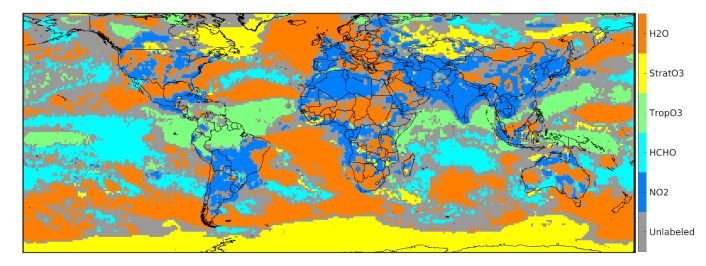
621 3.3. Synergy of the model and satellite observations to explain TOH long-term trends

622 3.3.1. The dominant contributor to TOH trends

Here, we take advantage of the wealth of information from satellites and our well-characterized model used for the inputs to the parameterization of OH to rank the dominant contributor to TOH linear trends. By assuming that TOH follows a linear combination of each individual experiment designed to isolate OH driver/proxy (i.e., *SOHnitro, SOHform, SOHtropozone, SOHstratozone*, and *SOHwv*), wherein second (or higher) chemical feedback is disregarded, we can determine the biggest contributor to the TOH trend for each model grid box by finding which driver/proxy holds the largest absolute amount. We only label a grid if the absolute linear trend of the dominant driver/proxy surpasses the second most dominant one by 30%.

630 Figure 7 illustrates the dominant factor explaining TOH trends. Several patterns can be found from 631 this result: i) NO₂ plays a significant role in TOH trends in various polluted areas, such as Asia and the Middle 632 East; ii) the upward trend of TOH over the western Pacific Ocean is primarily attributed to increased 633 tropospheric ozone from Asia (e.g., Lin et al., 2017); also, we observe a significant fraction of TOH over the 634 tropical Atlantic Ocean increasing because of rising tropospheric ozone from Africa and Central/South 635 America (Edwards et al., 2003); iii) HCHO is convolved with TOH trends over tropical oceans); iv) water 636 vapor plays a pivotal role in shaping TOH trends over oceans across the globe; iv) stratospheric ozone columns 637 are mostly significant over the South Pole due to the ozone healing process (Figure S2). The next sections will 638 focus on the magnitude of these trends and the degree to which they can collectively explain the variance in 639 TOH trends compared to Sanalysis.

It is important to recognize that the analysis presented here should be interpreted as a relative
assessment of a limited number of TOH drivers/proxies, rather than an exhaustive evaluation of all the physical
and chemical processes that are tied to TOH. Nonetheless, the data presented offers valuable insights into the
TOH trends and can be used as a basis for further research.



645

Figure 7. The major contributor to TOH trends based on the largest absolute trends of TOH drivers/proxiesabove 30% of the second most dominant factor.

648 3.3.2. Magnitudes of linear trends of TOH key inputs

Figure 8 shows the linear TOH trends influenced by NO₂ (*SOHnitro*), HCHO (*SOHform*), water
 vapor (*SOHwv*), tropospheric ozone (*SOHtropozone*), and stratospheric ozone (*SOHstratozone*). A

651 discussion on each parameter will follow:

652 SOHnitro – The trends in TOH driven by NO₂ show a strong correlation with the a posteriori trend 653 discussed in Section 3.1.1, with low- and medium-income countries experiencing an increase in TOH due to 654 rising NO_2 levels, while high-income countries see a reduction in TOH due to the opposite trend. The most significant increase in TOH is observed over India, where both the NO₂ trend and TOH sensitivity to NO₂ are 655 prominent. The most rapid regional decline in TOH seems to be over the NCP, because of NO_x reductions that 656 657 began after 2011. This finding is particularly noteworthy since M2GMI did not reproduce this trend without 658 OMI as a constraint. The trend in TOH resulting from NO_2 is predominantly anthropogenic in nature. This 659 aligns with the findings of Chua et al. (2023), who observed that the impact of lightning NO_x emissions on TOH trends was relatively minor. The global trend in TOH driven by NO₂ is positive, but with considerable 660 661 variation due to the significant disparities in how anthropogenic NO_x emissions have changed.

662 *SOHform* – We saw that HCHO was a reasonable proxy for TOH over oceans. Accordingly, the TOH 663 trends primarily are observed over oceans, especially over the Pacific and the Indian Oceans. This lines up 664 with the information gathered from the analysis of M2GMI and OMI HCHO observations (Figure 4). These 665 upward HCHO trends, as discussed in Section 3.1.2, may be influenced by transport and dynamics. It is worth 666 noting that the increase in TOH tied to this proxy (HCHO) is a global tendency, attributable to the relatively 667 uniform rise in HCHO levels across oceans.

668 SOHwv – Water vapor is a primary source of OH. The offline sensitivity of ECCOH captures this 669 tendency (Figure S22). Accordingly, the TOH linear trends mirror those of IWV (Figure S8) with major increases over oceans. Similar to other experiments, the global TOH increases because of rising water vapor 670 671 in the atmosphere. We acknowledge that understanding the reasons for changes in water vapor, which our 672 model shows to agree with Broger et al. (2022), is a complex subject that goes beyond the scope of our research. It requires an in-depth understanding of the water cycle, evapotranspiration and precipitation rates, and the 673 674 effect of temperature on the air's capacity to hold moisture, known as the Clausius Clapeyron relationship. 675 However, a great deal of effort has been made to demonstrate that global water vapor levels have increased significantly in recent decades. This is based on reanalysis data, microwave satellites, and in-situ 676 measurements (Trenberth et al., 2005; Chen and Liu, 2016; Wang and Liu, 2020; Allan et al., 2023), which is 677 678 consistent with what our model shows, as it is well-constrained by MERRA2 reanalysis data.

679 SOHtropozone - The impact of tropospheric ozone on OH formation is widely acknowledged 680 (Lelieveld et al., 2016). Likewise, our ECCOH offline sensitivity tests have revealed a largely positive correlation between tropospheric ozone and OH (Figure S23). Consequently, the linear trends observed in 681 682 TOH closely mirror those of tropospheric ozone in M2GMI (Figure S3). This tendency is especially noticeable 683 in the Atlantic Ocean, East and Southeast Asia, as well as the northern region of the Pacific Ocean, where 684 rising ozone levels have increased TOH. M2GMI suggests that tropospheric ozone levels in the southern 685 hemisphere have decreased (Text S1), potentially leading to a downward trend in TOH, an observation that has yet to be fully confirmed (e.g., Thompson et al., 2021). This finding is especially important given past 686 687 research indicating that models tend to exaggerate TOH asymmetry between the northern-southern hemispheres (Strode et al., 2015; Naik et al., 2013). The decrease in the simulated tropospheric ozone may 688 689 offer a plausible explanation for this tendency, but further verification is deemed necessary. Like the previous 690 experiments, tropospheric ozone on average leads to a global increase in TOH in 2005-2019.

SOHstratozone –Stratospheric ozone columns reduce UV actinic fluxes leading to a reduction in
 tropospheric JO¹D and thus OH, a tendency well reproduced by ECCOH (Figure S24). Nonetheless,
 stratospheric columns did not change noticeably in the tropics and mid-latitudes where OH production is
 important; consequently, the linear trends are close to zero or faintly negative due to a slight upward trend in
 the column. This tendency results in a rather uniform decrease of TOH globally.

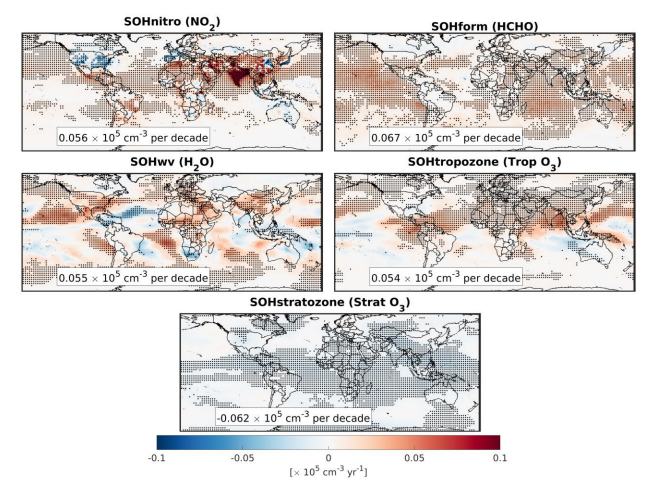


Figure 8. The contribution of each TOH key input (addressed in this study) to TOH in 2005-2019. HCHO, NO₂, and water vapor results are observationally constrained. Stratospheric ozone columns yielded comparable results compared to total ozone columns observed by OMI, however a large portion of tropospheric ozone trend has remained unverified in the southern hemisphere. ENSO affects the variability of TOH (Anderson et al., 2021), so we add a linear term to Eq.4 that is a function of the Niño 3.4 Index. This helps prevent ENSO from affecting the subsequent results.

703 3.3.3. OMI contributions to TOH trends

704 It is attractive to gauge the additional information gained from OMI on better representing the linear 705 trends of TOH. To achieve this, we need to analyze three sets of model output: one with OMI scaling factors, 706 one without OMI scaling factors, and one with the NO₂ and HCHO drivers (i.e., SOHnitro and SOHform). The 707 linear trends from these sets of model results are shown in Figure 9. The trends in the first column illustrate 708 the overall effect of NO_2 and HCHO on TOH trends, while the two other subplots isolate the effect of OMI 709 from the prior information based on M2GMI. M2GMI plays a significant role in shaping the trends in SOHnitro, possibly due to the small discrepancy between the trends in OMI and M2GMI columns over regions 710 711 where TOH is responsive to the driver. The most significant impact of OMI on NO_2 is visible over NCP. 712 Concerning HCHO, OMI slows down the upward trends in TOH over oceans which was suggested by M2GMI. 713 In general, M2GMI largely dictates the overall shape of TOH trends driven by NO₂ and HCHO possibly due 714 to small difference between the model and OMI observations and/or limited informational content in OMI.

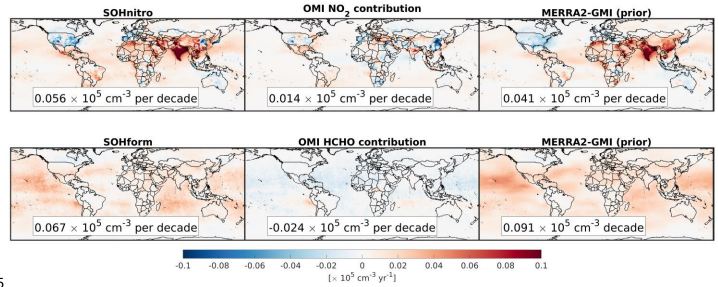
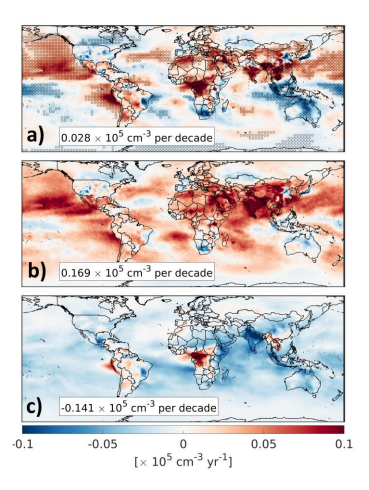


Figure 9. The resulting effect of tropospheric NO₂ and HCHO on TOH linear trends during 2005-2019 (first column); the contributions of OMI information added on top of the prior knowledge (M2GMI) (middle column); the effect of the prior knowledge on shaping TOH linear trends (last column).

719 *3.3.4.* How well can these experiments explain the simulated trends collectively?

720 We find that there is a good degree of correlation between the sum of the linear trends and those of 721 Sanalysis (R²=0.65) indicating that a good portion of variability in TOH trend can be well explained by these 722 experiments (Figure S25). Figure 10a shows the linear trend of TOH from Sanalysis in 2005-2019, and Figure 723 10b shows the sum of the linear trends of the five OH key inputs. These maps are one of the most recent and 724 detailed TOH trends available, relative to newer studies (Nicely et al., 2018; Zhao et al., 2020; Chua et al., 2023). The TOH trend from Sanalysis varies greatly, where positive values are prevalent over northern parts 725 of the Pacific Ocean, the Middle East, central Africa, and several regions over East Asia. Negative trends are 726 727 found over the US, southeast Asia, and the southern part of the Pacific Ocean. The linear sum of the 728 experiments strongly aligns with Sanalysis, particularly over the northern hemisphere, reinforcing that the 729 selected parameters are sensible choices to reproduce a large portion of variance in TOH trend.



730

Figure 10. (a) The linear trends derived from *Sanalysis* experiment, the "best effort" to simulate the evolution of the CH₄-CO-OH cycle, from 2005-2019. The statistically significant trends are superimposed by dots. (b) The linear summation of the five selected TOH influencers including water vapor, NO₂, HCHO, stratospheric and tropospheric ozone, showing a strong degree of correspondence to the top panel, particularly in the northern hemisphere. (c) The unexplained portion of the TOH trends, which was not explainable by five experiments addressed in this research.

738 Revealing the unexplained portion of TOH trends, which cannot be attributed to the selected TOH 739 experiments, is necessary. Within the model, various physiochemical factors such as CO, CH₄, dynamics, 740 aerosols, and clouds can impact the TOH trends. Although we will not delve into these drivers in this study, 741 we can identify unexplained parts of TOH trends by subtracting the sum of trends derived from the five primary 742 TOH key inputs from those of Sanalysis, which discounts second (or higher) chemical feedback. Figure 10c 743 displays the unexplained TOH trends between 2005 and 2019. It is readily apparent that there are uniform and 744 significant downward trends in TOH in the tropics and subtropics where photochemistry is strong. This is most 745 likely triggered by increasing concentrations of CH₄, which is demonstrated in Figure S10, causing OH levels 746 to decrease over time. It is very probable that the extent of these downward trends in TOH has been exaggerated 747 in our model because of the simulated CH₄ increasing too rapidly compared to in-situ observations. The 748 overestimation of the upward trend in CH₄ in our model compared to in-situ observations could be caused by 749 the biases (~3%) in sources minus sinks and/or the initial condition. Consequently, the globally-averaged TOH 750 trend derived from Sanalysis may be slower than it should be. Lastly, an unexplained strong upward trend in 751 TOH over central Africa lingers.

752 **4.** Conclusion

753 While a comprehensive multi-sensor/multi-species data assimilation and inverse modeling approach, such as Souri et al. (2020a), Miyazaki et al. (2020), and Souri et al. (2021), would be ideal for fully harnessing 754 the potential of satellite information on improving multiple aspects of a model representing OH, it will be 755 prohibitively expensive. Therefore, our simplified approach serves the purpose of understanding the first-order 756 effects of observational adjustments to TOH drivers/proxies before committing substantial resources to the 757 implementation/execution of an observationally-constrained, full-chemistry model. Here, we implemented the 758 759 newest version of the parameterization of OH, following Anderson et al. (2022), within NASA's GEOS model, 760 presenting an opportunity to understand and mitigate TOH biases caused by misrepresentation of HCHO and 761 NO₂ concentrations with respect to the state-of-the-art OMI NO₂ and HCHO retrievals using Bayesian data fusion, as well as to unravel the intricacies of TOH to its key inputs such as tropospheric and stratospheric 762 763 ozone and water vapor.

764 We found large positive biases in tropospheric NO_2 columns in M2GMI, the archived model used as 765 an input to the parameterization of OH, compared to OMI over Africa, Eastern Europe, and the Midwest US. 766 Because of a large positive effect of NO₂ (a surrogate for NOx) on TOH, a tendency well captured by our 767 implicit parameterization, these overestimations introduced significant regional biases in TOH up to 20%, and 768 a global overestimation of TOH by 1%. Consistent with former work, we saw distinct disparities in the sign 769 of linear trends of tropospheric NO₂ over high- and medium-income countries (i.e., negative) and low-income countries (i.e., positive). While M2GMI generally replicated these trends, notable deviations were identified 770 771 over China leading to an erroneous trend of TOH.

Pronounced inaccuracies with regards to both the simulated HCHO magnitude and trend in M2GMI were revealed by OMI over land. However, this proxy for OH was loosely connected to TOH in areas where photolysis was not the major sink of HCHO (Wolfe et al., 2019), especially over land. Over oceans, where HCHO and TOH were highly correlated, adjustments to M2GMI by OMI HCHO were relatively mild resulting in small alterations to TOH which was by an order of magnitude lower than those of NO₂. These mild alterations speak to either an insufficient amount of information in OMI or the reasonable accuracy of M2GMI over pristine areas.

779 In general, five variables including NO₂, HCHO, water vapor, tropospheric ozone, and stratospheric 780 ozone, could collectively account for 65% of the variance in TOH trends globally. To estimate this, we 781 executed various modeling experiments to isolate the effect of NO₂, HCHO, water vapor, tropospheric ozone, and stratospheric ozone on long-term trends of TOH in 2005-2019 at 1°×1° resolution. Except for tropospheric 782 783 ozone, these variables were either constrained by observations or aligned with independent observations, 784 boosting confidence in our trend results. Given the robust positive correlation between OH and NO₂, HCHO, 785 water vapor, and tropospheric ozone over regions where photochemistry was active, TOH trends influenced 786 by these variables closely mirrored the trends in their respective drivers/proxies. For instance, high- and 787 medium-income countries exhibited negative TOH trends driven by NO₂. Rising tropospheric ozone over east and south Asia, heavily vetted by various observations (Guadel et al., 2018), led to an upswing in TOH over 788 789 the Pacific Ocean. The trend of water vapor, greatly in agreement with independent observation (Broger et al., 790 2022), was dominantly positive over oceans leading to further enhancement of TOH. Rising HCHO over 791 Pacific and Indian Ocean suggested by constrained M2GMI was associated with increased TOH. The effect 792 of stratospheric ozone on TOH was marginal in low and mid latitudes due to negligible changes in stratospheric ozone columns in M2GMI reconfirmed by OMI total ozone column observations. 793

A large offset between our analysis experiment with varying CO and CH₄ concentrations was observed after removing the sum of the linear trends derived from these five key experiments from the analysis experiment, indicating that our future research using ECCOH should include new experiments isolating the effects of CO, CH₄, and transport (e.g., Gaubert et al., 2017; Zhao et al., 2020). Those experiments will refine the investigation of the unexplained portion of the TOH trend.

The development of an effective parameterization of OH, that is capable of integrating advanced satellite-based gas retrievals and improved weather forecast models enabled us to unravel the convoluted response of TOH to various parameters. Nonetheless, it is important to recognize some of the limitations 802 associated with our work: first, the offline nature of the Bayesian data fusion algorithm makes the entire 803 experiment blind to the interconnected responses of various compounds, such as ozone or aerosols, to 804 adjustments to NO₂ and HCHO. Despite this limitation, our work has provided valuable insights into the firstorder effects of adjustments on TOH key inputs. This can help quickly identify areas where our prior 805 knowledge is least reliable to simulate TOH. Second, the machine learning algorithm employed for 806 807 parameterizing OH is implicit and its response to drivers/proxies is complex, making it difficult to 808 quantitatively verify against full chemistry models. However, by including a vast number of parameters in the 809 parameterization, Anderson et al. (2022) boosted its ability to understand the convoluted chemistry of OH. This has allowed for reproducing OH for events not included in the training dataset (Anderson et al., 2022, 810 811 2023, 2024).

812 The longevity and stability of Aura's record of observations have played a significant role in 813 constraining/assessing several important variables pertaining to TOH on a global scale. This is exemplified by 814 the wealth of information obtained from OMI NO₂, HCHO, water vapor, total ozone columns, and Microwave Limb Sounding (MLS) temperature and ozone, that are used directly or indirectly in our analysis. However, 815 as Aura's mission comes to an end, there will be a gap in the monitoring of these variables. TROPOMI, OMI's 816 817 successor, can help fill this gap, but its record of observation is still short; therefore, it is important to invest 818 in research to harmonize data from multiple satellite observations such as OMI and TROPOMI (e.g., Hilboll 819 et al., 2013). This is because each sensor can have different biases and spatial representativity, which can lead 820 to inconsistencies and potentially conflicting values if they are used together.

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824 Data availability

- 825 Satellite data can be accessed for Level 2 OMI tropospheric NO₂ at
- 826 <u>https://doi.org/10.5067/MEASURES/MINDS/DATA204</u> (Lamsal et al., 2022), Level 2 OMI total ozone
- 827 columns at https://disc.gsfc.nasa.gov/datasets/OMTO3_003/summary (Bhartia, 2005), OMI SAO HCHO at
- 828 <u>https://waps.cfa.harvard.edu/sao_atmos/data/omi_hcho/OMI-HCHO-L2/</u> (Gonzalez Abad, 2023), MOPITT
- 829 CO (https://doi.org/10.5067/TERRA/MOPITT/MOP03JM_L3.008) (NASA LARC, 2000), OMI/MLS TO₃
 830 at https://acd-ext.gsfc.nasa.gov/Data_services/cloud_slice/data/tco_omimls.nc (Ziemke, 2023).
- 831 In-situ CO and CH₄ observations can be obtained from
- 832 <u>https://gml.noaa.gov/dv/data/index.php?category=Greenhouse%2BGases</u> (Helmig et al., 2021; Lan et al., 2021).
- 834 MERRA2-GMI model outputs can be downloaded from https://acd-
- 835 ext.gsfc.nasa.gov/Projects/GEOSCCM/MERRA2GMI/ (NASA Goddard Space Flight Center, 2023).

836 Code availability

- 837 OI-SAT-GMI python package developed for this research can be found from
- 838 <u>https://doi.org/10.5281/zenodo.10520136</u> (Souri, 2024).
- 6EOS-Quickchem used to run the modeling experiments encompassing ECCOH can be found from
 https://github.com/GEOS-ESM/QuickChem.git.
- 841 GEOS model can be obtained from <u>https://github.com/GEOS-ESM/GEOSgcm.git</u>.
- 842 Offline ECCOH calculations to derive the sensitivity of TOH to different drivers/proxies can be obtained
 843 from https://doi.org/10.5281/zenodo.10685100

844 Authors contributions

- 845 A.H.S and B.N.D designed the research. A.H.S analyzed the data, conducted the simulations, made all the
- figures, and wrote the original manuscript. B.N.D helped with conceptualization, fund raising, and writing.
- 847 S.A.S helped configuring the model and interpreting the results. M.E.M and D.C.A implemented the improved
- 848 ECCOH module into GEOS-5 Quickchem. J.L. thoroughly validated the model with respect to CO and CH₄
- 849 observations. B.W. provided an improved CO emission inventory. L.D.O provided M2GMI and helped

- 850 interpret it. Z.Z. provided improved wetland CH₄ emissions. All the authors contributed to the discussion and
- edited the paper.
- 852 **Competing interests**
- 853 B.N.D is a member of the editorial board of Atmospheric Chemistry and Physics.

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