Technical Note: Constraining the hydroxyl (OH) radical in the tropics with satellite observations of its drivers: First steps toward assessing the feasibility of a global observation strategy

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Daniel C. Anderson^{1,2}, Bryan N. Duncan², Julie M. Nicely^{2,3}, Junhua Liu^{2,4}, Sarah A. Strode^{2,4}, Melanie B. Follette-Cook⁵

- 5 Follett 6
- 7 1. GESTAR II, University of Maryland Baltimore County, Baltimore, MD, USA
- Atmospheric Chemistry and Dynamics Laboratory, NASA Goddard Space Flight Center, Greenbelt,
 MD, USA
- 10 3. Earth System Science Interdisciplinary Center, University of Maryland, College Park, MD, USA
- 11 4. GESTAR II, Morgan State University, Baltimore, MD, USA
- Mesoscale Atmospheric Processes Laboratory, NASA Goddard Space Flight Center, Greenbelt, MD,
 USA
- 14

15 Correspondence to: Daniel C. Anderson (daniel.c.anderson@nasa.gov)

16 17 Abstract

- 18 Despite its importance in controlling the abundance of methane (CH₄) and a myriad of other
- 19 tropospheric species, the hydroxyl radical (OH) is poorly constrained due to its large spatial
- 20 heterogeneity and the inability to measure tropospheric OH with satellites. Here, we present a
- 21 methodology to infer tropospheric column OH (TCOH) in the tropics over the open oceans using a
- 22 combination of a machine learning model, output from a simulation of the GEOS model, and satellite
- 23 observations. Our overall goals are to assess the feasibility of our methodology, to identify potential
- 24 limitations, and to suggest areas of improvement in the current observational network. The
- 25 methodology reproduces the variability of TCOH from independent 3D model output and of
- 26 observations from the Atmospheric Tomography mission (ATom). While the methodology also
- 27 reproduces the magnitude of the 3D model validation set, the accuracy of the magnitude when applied
- to observations is uncertain because current observations are insufficient to fully evaluate the machine
- learning model. Despite large uncertainties in some of the satellite retrievals necessary to infer OH,
 particularly for NO₂ and HCHO, current satellite observations are of sufficient guality to apply the
- 31 machine learning methodology, resulting in an error comparable to that of *in situ* OH observations.
- 32 Finally, the methodology is not limited to a specific suite of satellite retrievals. Comparison of TCOH
- determined from two sets of retrievals does show, however, that systematic biases in NO₂, resulting
- 34 both from retrieval algorithm and instrumental differences, lead to relative biases in the calculated
- 35 TCOH. Further evaluation of NO₂ retrievals in the remote atmosphere is needed to determine their
- 36 accuracy. With slight modifications, a similar methodology could likely be expanded to the extra-tropics
- and over land, with the benefits of increasing our understanding of the atmospheric oxidation capacity
- $\label{eq:and_standard} 38 \qquad \text{and, for instance, informing understanding of recent CH}_4 \, \text{trends.}$
- 39

40 **1** Introduction

- 41 The hydroxyl radical (OH) dictates the lifetime of many tropospheric species, including carbon monoxide
- 42 (CO), methane (CH₄), and numerous volatile organic compounds (VOCs). Knowledge of OH is therefore
- 43 necessary to understand the abundance, distribution, and variability of these species. For instance,
- 44 Rigby et al. (2017) and Laughner et al. (2021) attribute recent trends and increases in CH₄ at least
- 45 partially to changes in OH abundance. Current constraints on OH are insufficient, however, to assess its
- 46 relative importance in controlling these trends (Turner et al., 2017).
- 47

48 Differences in OH distributions among chemistry transport (CTM) and chemistry climate models (CCM)

- 49 suggest that these models are insufficient to inform understanding of OH abundance and variability
- 50 without further observational constraints. OH abundance can differ by up to 80% among models
- 51 constrained with identical emissions in intercomparison projects (Voulgarakis et al., 2013;Nicely et 52 al.; Zhao et al., 2019; Murray et al., 2021), with modeled trends disagreeing with those derived from
- 53
- observationally constrained methods (Stevenson et al., 2020). Variables such as the photolysis 54 frequency of O_3 (JO¹D) (Nicely et al., 2020), the NO_x lifetime (NO_x = NO + NO₂), and the oxidation
- 55 efficiency of VOCs (Murray et al., 2021) contribute to these inter-model variations in OH. Using
- 56 Gaussian emulation, Wild et al. (2020) found that the relative importance of drivers of OH variability
- 57 differed widely among three CTMs. Likewise, the response of OH to the El Niño Southern Oscillation
- 58 (ENSO), the dominant mode of OH variability on monthly and seasonal timescales (e.g. Anderson et al.,
- 59 2021; Turner et al., 2018), and other modes of internal climate variability can vary widely among models 60 (Anderson et al., 2021).
- 61

62 Despite this need for better constraints, observations of tropospheric OH are limited. The hydroxyl

- 63 radical has a lifetime of approximately 1s (Mao et al., 2009), resulting in large spatial heterogeneity in
- 64 both the horizontal and vertical. This spatial heterogeneity is further caused by the large variation in the
- 65 relative importance of drivers of OH loss and production in different regions of the atmosphere (e.g.
- 66 Spivakovsky et al., 1990;Lelieveld et al., 2016). A strategic, representative in situ observational network
- 67 is therefore unfeasible. As a result, observations of OH are generally limited to intensive field campaigns
- 68 (Miller and Brune, 2022) that have narrow spatial and temporal coverage. While remotely-sensed OH
- 69 observations are available, those from satellites are limited to the stratosphere (e.g., Pickett et al.,
- 70 2008), while ground-based observations of total column OH are dominated by the stratospheric
- 71 contribution (e.g., Burnett and Minschwaner, 1998).
- 72

73 Reference gases with well-characterized sources and an OH sink, such as methyl chloroform (MCF), can 74 be used to infer OH abundance (Lovelock, 1977). This methodology, however, generally yields no 75 information on spatial heterogeneity beyond the hemispheric scale (e.g., Montzka et al., 2011; Rigby et 76 al., 2017; Naus et al., 2019), although there has been recent success when using three dimensional 77 inversion techniques (Naus et al., 2021). For MCF in particular, recent declines in tropospheric 78 abundance will soon dictate the need for a new reference species (Liang et al., 2017).

79

80 Multiple studies have attempted to constrain OH through the creation of proxies and the application of 81 satellite retrievals of OH drivers. Murray et al. (2014) showed that global OH strongly correlated with a

- 82 combination of JO¹D, water vapor ($H_2O_{(v)}$), and the tropospheric sources of reactive nitrogen and carbon
- 83 in the GEOS-Chem model. Murray et al. (2021) demonstrated that OH correlated with this proxy in
- 84 multiple CTMs, although the relationship differs strongly among models. Miyazaki et al. (2020) created
- 85 a data assimilation framework that ingested satellite observations of CO, NO₂, O₃, and HNO₃ (nitric acid)
- 86 into multiple CTMs. The data assimilation reduced the spread in average OH among the models and
- 87 brought the interhemispheric ratio closer to unity, in line with values suggested by MCF observations
- 88 (e.g. Patra et al., 2014). These results demonstrate that the incorporation of satellite observations into a
- 89 modeling framework can improve the representation of OH. Wolfe et al. (2019) developed a proxy for
- 90 OH based on formaldehyde (HCHO) production and loss rates. They applied that proxy to satellite HCHO
- 91 observations to estimate OH columns in the remote troposphere, a region where HCHO abundance is
- 92 low and the satellite retrievals are reflective of the *a priori* (Zhu et al., 2016). Using machine learning,
- 93 chemical transport model output, and retrievals of NO₂ and HCHO, Zhu et al. (2022b) developed a
- 94 method to estimate surface OH in North American urban areas. Finally, Pimlott et al. (2022) used a
- 95 steady state approximation of OH, including primary production from H_2O and O_3 and loss from CO, CH₄,

- 96 and O₃, to estimate OH between 600 and 700 hPa using observations from IASI (Infrared Atmospheric
- 97 Sounding Interferometer). A logical next step, building on the results of these studies, is the
- 98 development of a methodology to constrain OH that ingests multiple satellite retrievals, encompasses
- 99 the breadth of OH chemical and dynamical drivers, and spans a significant enough portion of the globe
- to inform variability and trends in CH₄ and CO loss.
- 101

102 Combining machine learning, chemical transport model (CTM) output, and satellite data has the 103 potential to constrain tropospheric column OH (TCOH). A variety of machine learning techniques, such 104 as neural networks (Nicely et al., 2017; Nicely et al., 2020; Kelp et al., 2020), self-organizing maps 105 (Stauffer et al., 2016), random forest regression (Keller and Evans, 2019), and gradient boosted 106 regression trees (GBRTs) (Ivatt and Evans, 2020; Zhu et al., 2022b; Anderson et al., 2022) show promise in 107 helping to solve problems in atmospheric chemistry. In particular, Zhu et al. (2022b) and Anderson et al. 108 (2022) demonstrated the ability of GBRTs to predict OH from a chemical transport model with 109 reasonable accuracy. GBRT models (Elith et al., 2008; Chen and Guestrin, 2016) use an ensemble of 110 decision trees to predict the value of a target based on multiple inputs, even for targets with highly non-111 linear dependencies on the inputs.

112

113 Here, we present a methodology to infer clear sky TCOH in the tropics from space-based observations of

- 114 its chemical and dynamical drivers with the goal of assessing the feasibility of our methodology,
- identifying potential limitations, and suggesting areas of improvement in the current observational network. We train a GBRT model using output from a simulation of the NASA GEOS (Goddard Earth
- 117 Observing System) model, and then estimate TCOH in the actual atmosphere at the satellite overpass
- 118 time using inputs from a suite of satellite retrievals. In Section 2, we describe the methodology for
- 119 generating the machine learning model as well as the satellite retrievals used to constrain TCOH. We
- 120 then evaluate the suitability of MERRA2 GMI as a training dataset (Sect. 3) and, in Section 4, present a
- satellite-constrained OH product for one month from each season. Finally, in Section 5, we explore
- 122 potential methodological limitations and benefits, including lack of validation data, the impacts of
- observational uncertainties, and the ability to use different satellites and retrievals as inputs to the GBRTmodel.
- 125

2 Description of the methodology to generate the GBRT model and of the associated datasets

Our overall aim is to demonstrate the feasibility of our approach to constrain TCOH with satellite-based observations over broad regional scales. As a first step, we restrict our analysis to latitudes equatorward of 25° and regions over water. We chose to focus initially on this domain as it has appreciable OH concentrations and simplified chemistry, as compared to regions with large biogenic and anthropogenic VOC emissions. Nevertheless, this portion of the atmosphere accounts for 50 – 60% of global CO and CH₄ loss. In this section, we describe the creation of the machine learning model used to predict TCOH

- (Sect. 2.1) for this region as well as the satellite products used as inputs to the machine learning model(Sect. 2.2).
- 135

136 **2.1 Creation of the TCOH model**

137 2.1.1 Creation of the GBRT training dataset

- 138 For the machine learning model training dataset, we use a subset of output from the MERRA2 GMI
- 139 simulation (https://acd-ext.gsfc.nasa.gov/Projects/GEOSCCM/MERRA2GMI/). MERRA2 GMI is a 40 year
- 140 (1980 2019) simulation of the NASA GEOS model run in replay mode (Orbe et al., 2017) with MERRA2
- 141 (Modern Era Retrospective analysis for Research and Applications, version 2) meteorology (Gelaro et al.,
- 142 2017). The simulation has a resolution of c180 on the cubed sphere (approximately 0.625° longitude by
- 143 0.5° latitude) with 72 vertical layers and uses the Global Modeling Initiative (GMI) chemical mechanism

- 144 (Duncan et al., 2007; Strahan et al., 2007). Output is available at daily- and monthly-averaged resolution,
- as well as instantaneous values at 10:00 and 14:00 LST. These times are within approximately 30
- 146 minutes of the overpass times of the satellites described in Section 2.2. Anderson et al. (2021) and
- 147 Strode et al. (2019) provide detailed information about the simulation, including emissions.
- 148

149 The training target for the machine learning model is TCOH. In Anderson et al. (2022), we developed a 150 GBRT parameterization trained on MERRA2 GMI output to predict in situ OH concentrations using 27 151 inputs, only a small fraction of which are observable from space. That parameterization, designed to be 152 integrated into the GEOS modeling framework, performed better when there was a separate model for 153 each month as opposed to one model for all months. While that GBRT model is not appropriate for the 154 application described here, we employ a similar approach, creating a separate set of TCOH training 155 targets for each month. We use instantaneous OH output from MERRA2 GMI at 14:00 local time for 156 each day of a given month across the years 2005 to 2019, a timeframe that maximizes overlap between 157 the operational lifetime of the satellites listed in Table 1 and the period of the MERRA2 GMI simulation. 158 We omitted data from 2017 to evaluate model performance. For a given month and year, we calculate 159 daily tropospheric column values across the grid, filtering out columns where the maximum cloud 160 fraction in that column was greater than 30% in order to align the training targets more closely with 161 satellite data, where retrievals of some species are often filtered for cloud cover. This yields 162 approximately 43,000 valid grid boxes per day. For each year, we then average these values to monthly 163 resolution. This results in approximately 600,000 total training targets for each month over the 15-year period.

- 164 165
- Table 1: Input variables to the machine learning model and the corresponding satellite retrieval used to create the
 satellite OH product. Overpass times are ~13:30 LST for all satellites except MOPITT, which has a 10:30 LST
- 168 overpass.

Variable	Satellite retrieval	Original horizontal and temporal resolution	Reference
Total O₃ column	OMI TOMS-Like L3 version 3	$0.25^{\circ} \times 0.25^{\circ}$, daily	McPeters et al. (2015)
Tropospheric NO ₂ column	OMI GSFC L3 version 4	$0.25^{\circ} \times 0.25^{\circ}$, daily	Lamsal et al. (2021)
CO column	MOPITT L3 version 8	$1.0^{\circ} imes 1.0^{\circ}$, monthly	Deeter et al. (2019)
HCHO column	OMI SAO L3 version 3	$0.1^\circ \times 0.1^\circ$, daily	González Abad et al. (2015)
$H_2O_{(v)}$ column	AIRS L3 version 6	$1.0^\circ imes 1.0^\circ$, monthly	Susskind et al. (2014)
Sea surface temperature	MUR L4 version 4.2	$0.25^{\circ} imes 0.25^{\circ}$, daily	Chin et al. (2017)
Aerosol optical depth at 550 nm	MODIS Aqua L3 collection 6	$0.5^{\circ} \times 0.5^{\circ}$, daily	Levy et al. (2013)
H ₂ O _(v) layers: 925 – 850 hPa, 850 – 700 hPa, 700 – 600 hPa, 600 – 500 hPa, 500 – 400 hPa, 400 – 300 hPa, and 300 - 250 hPa	AIRS L3 version 6	$1.0^\circ imes 1.0^\circ$, monthly	Susskind et al. (2014)
Solar zenith angle	N/A		
Latitude	N/A		

169

170 We selected the input variables for the machine learning model (Table 1) based on their relevance to OH

171 chemistry and variability as well as our current ability to observe the variable with satellites.

- 172 Performance was similar for a model including total column ozone only and for a model also including
- 173 the tropospheric column. We therefore use total column ozone because of the uncertainties inherent in

- 174 separating the column into two parts in the satellite retrieval. We chose the water vapor layers to
- 175 correspond with the Atmospheric Infrared Sounder (AIRS) layers product. Layers are averages over the
- indicated pressure range, and we denote the layer names by the highest pressure in that range. We
- include sea surface temperatures (SST) as a proxy for the Indian Ocean Dipole and ENSO, which has a
 strong impact on OH variability in the tropics (Anderson et al., 2021;Turner et al., 2018;Naus et al.,
- strong impact on OH variability in the tropics (Anderson et al., 2021;Turner et al., 2018;Naus et al.,
 2021). In addition, we include latitude and solar zenith angle as previous work has shown that these
- variables can explain a large fraction of the spatial OH variability (Duncan et al., 2000;Anderson et al.,
- 181 2022).
- 182
- 183 We sampled the MERRA2 GMI output to create the training dataset in the same manner as for the TCOH 184 targets. The inputs to the machine learning model each correspond to the same model column as the
- 185 OH target. All column values are instantaneous and taken from 14:00 to correspond with satellite
- 186 overpass times, except for CO, which is for 10:00, near the Measurement of Pollution in the
- 187 Troposphere (MOPITT) overpass time. Model performance was similar when using CO output at 14:00
- and 10:00, likely because of limited diurnal variability in CO column in the study region. SSTs are
- 189 monthly averages of 24-hour averaged values, and we calculated solar zenith angle at the surface for
- 190 noon on the 15th of a given month.
- 191

192 **2.1.2 Creation and tuning of the GBRT model**

We used the XGBoost package (Chen and Guestrin, 2016) version 0.81 in Python version 3.6 to create a
GBRT model of TCOH for each month using the training datasets from MERRA2 GMI. For each month,
we used 90% of the dataset for model training and the remainder for model validation. As mentioned in
Section 2.1.1, we also used MERRA2 GMI output from 2017, which was omitted from the training
dataset, as further validation.

198

199 To maximize parameterization performance while also balancing the potential of overfitting, we tuned 200 hyperparameters, including the learning rate, the maximum tree depth, and the number of trees. We 201 chose hyperparameter values that minimized the parameterization root mean square error (RMSE) of 202 the training dataset. We set the learning rate, which controls the magnitude of change when adding a 203 new tree, to 0.1, while we varied the maximum tree depth and number of trees from 6 to 22 and from 204 10 to 150, respectively. For both maximum tree depth and number of trees, RMSE initially dropped 205 significantly with increasing value, representing sharp improvement in parameterization performance. 206 RMSE values eventually plateaued, increasing parameterization runtime without noticeably improving 207 performance. A combination of a maximum tree depth of 18 and 100 trees balanced performance with 208 model training and run time.

209

To determine whether the inputs to the machine learning model improved or hindered performance, we performed a "leave one out" analysis. Using 5-fold cross validation, we retrained the model, individually omitting each of the inputs, to determine the percent difference between the mean RMSE of the 5 folds for the model without a specific input and one including all inputs. Omitting the inputs listed in Table 1 lead to increases in the RMSE, suggesting that each is necessary for improved model performance. As a result of this analysis, we do not use water vapor layers for pressures less than 300 hPa because these decreased model performance.

217

Finally, we found that it was not necessary to apply satellite averaging kernels and shape factors to the

- training dataset. Of the satellite retrievals used in this work (discussed in Sect. 2.2 and listed in Table 1),
- only CO, HCHO, and NO₂ could require convolving the model with the averaging kernel. Shape factors
- $221 \qquad \mbox{for the OMI NO}_2 \ \mbox{retrieval are determined from a similar setup of the GEOS model, also employing the}$

- 222 GMI chemical mechanism and MERRA2 meteorology. Applying the satellite shape factors to the
- 223 simulation discussed here would therefore not result in significant changes in the modeled NO₂
- 224 (Anderson et al., 2021). To test whether it is necessary to apply the averaging kernels for CO and HCHO,
- we created a separate training dataset, where we convolved the daily MERRA2 GMI output with the
- averaging kernel and a priori from the level 2 data for both species for February 2005 2019. All other
- inputs were kept the same. We then retrained the model with these adjusted CO and HCHO variables.
- When we applied the satellite data to the model for February 2017, as described in Section 4, the resulting TCOH differed by less than 1% on average from the model that did not include averaging kerne
- resulting TCOH differed by less than 1% on average from the model that did not include averaging kernel information. This level of uncertainty is significantly smaller than the other uncertainties discussed in
- 231 Section 5, so we do not include averaging kernels in our analysis.
- 232

233 2.2 Description of satellite products

To create the observationally-constrained OH product, we use multiple satellite retrievals, listed in Table 1 and briefly described here. Each instrument is located onboard a polar orbiting satellite that provides near global coverage daily. For each satellite retrieval, we use the level 3 gridded product, with the

exception of SST which is level 4. Where necessary, we regridded the retrieval to a common horizontal

- 238 grid with a resolution of $1.0^{\circ} \times 1.0^{\circ}$ and averaged to the monthly scale.
- 239

240 We use these resolutions because, in the study domain, individual pixel retrievals, particularly of NO₂

and HCHO, are frequently at or below detection limits (González Abad et al., 2015;Lamsal et al., 2021),

242 necessitating averaging to relatively coarse temporal and spatial scales. Missing data due to cloud cover

243 and the Ozone Monitoring Instrument (OMI) row anomaly further increase the need for monthly-scale

averaging. While other satellites, such as OMPS (Ozone Mapping and Profiler Suite) and TROPOMI

245 (Tropospheric Monitoring Instrument), provide retrievals with increased signal to noise ratios and more

complete data coverage, the satellites used here cover a far longer time period. The $1.0^{\circ} \times 1.0^{\circ}$ and

- 247 monthly resolutions, in combination with the long data record, are sufficient to understand regional
- trends in TCOH and some aspects of TCOH temporal and spatial variability.
- 249

We use retrievals of three species – HCHO, O₃, and NO₂ – from OMI, an ultraviolet-visible spectrometer located onboard the Aura satellite, which has an overpass of approximately 13:30 local solar time (LST). We use the Smithsonian Astrophysical Observatory (SAO) version 3 HCHO retrieval (González Abad et al., 2015). Wolfe et al. (2019) found that this retrieval captured the variability of the HCHO columns in the remote atmosphere observed during the Atmospheric Tomography (ATom) campaign with little bias. For total column O₃, we use the TOMS-like (Total Ozone Mapping Spectrometer) retrieval version 3

(McPeters et al., 2015), which agrees with ground-based and other satellite observations within
 approximately 1% (Labow et al., 2013). Finally, we use the Goddard Space Flight Center version 4 NO2
 transcribering adjump ratriaval (Lampa) at al. 2021). While provides studies have theroughly avaluated

tropospheric column retrieval (Lamsal et al., 2021). While previous studies have thoroughly evaluated
 this retrieval in more polluted atmospheres (e.g., Lamsal et al., 2014;Choi et al., 2020), evaluation in the

- 260 remote tropical atmosphere, as defined in this study, is limited.
- 261

262 For water vapor and aerosol optical depth (AOD) at 550 nm, we use retrievals from AIRS and the

263 Moderate Resolution Imaging Spectroradiometer (MODIS) instruments, respectively, both located

onboard the Aqua satellite with an overpass of approximately 13:30 LST. We use the total column water

vapor standard physical retrieval as well as the 7 water vapor layers listed in Table 1 (Susskind et al.,

- 266 2014). Multiple studies have evaluated the accuracy of the AIRS $H_2O_{(v)}$ column and layers retrievals in
- the remote tropical atmosphere, finding bias of 5% or less and high correlation against both remote and
- 268 in situ observations (Bedka et al., 2010;Anderson et al., 2016;Pérez-Ramírez et al., 2019). We use

- 269 collection 6 of the dark target MODIS AOD retrieval at 550 nm, which is highly correlated with
- 270 observations from the AERONET network over the ocean (Levy et al., 2013).
- 271

272 We also use retrievals of CO from MOPITT, which is onboard the Terra satellite with an overpass of 273 10:30 LST. We use the version 8 retrieval that includes both near and thermal infrared radiances (Deeter

- 274 et al., 2019). CO retrievals from MOPITT in the remote tropics generally agree with ground-based
- 275 remotely-sensed observations within 10% (Hedelius et al., 2019; Buchholz et al., 2017).
- 276
- 277 Finally, we use SSTs from the Multi-scale Ultra-high Resolution (MUR) analysis, which combines
- 278 nighttime SST observations from multiple satellite platforms, including MODIS, as well as in situ
- 279 observations and agrees with other SST analyses within 0.36° C (Chin et al., 2017).
- 280

281 3 Evaluating the Suitability of the MERRA2 GMI Simulation as a Training Dataset

282 Before generating the GBRT model to predict TCOH, we first demonstrate that the MERRA2 GMI 283 simulation is suitable to use as a training dataset. Because of the paucity of in situ observations of OH 284 over most of the globe, we necessarily use output from an atmospheric chemistry model to train the 285 machine learning model. The atmospheric chemistry model output must reasonably capture the 286 distribution, magnitude, and ENSO-related variability of OH and the drivers listed in Table 1, as GBRT 287 models are unable to extrapolate beyond the photochemical environments on which they are trained 288 (Anderson et al., 2022).

289

290 3.1 Comparison of the Distribution and Magnitude of Simulated OH Drivers to Observations

291 Simulated OH from MERRA2 GMI agrees with observations over the remote ocean within the

- 292 instrumental uncertainty. Anderson et al. (2021) compared MERRA2 GMI output to in situ observations
- 293 from the first two deployments of ATom, finding modest correlation (r² values between 0.3 and 0.78
- 294 depending on the hemisphere and season) between observations and the model. The average
- 295 normalized mean bias was on the order of 20%, a slight high bias but within the 2σ observational
- 296 uncertainty of 35%. Agreement was highest in the remote atmosphere, whereas the largest error was in
- 297 regions of fresh, continental outflow off the coasts of South America and New Zealand.
- 298







302 use $H_2O_{(v)}$ at 700 hPa as an example for all $H_2O_{(v)}$ layers. Distributions of the other $H_2O_{(v)}$ layers are shown in Figure

S1. We also indicate the r^2 of the correlation between MERRA2 GMI output for February 2017 and the

304 corresponding satellite retrieval as well as the normalized mean bias of that output.

305 The simulation captures both the observed variability and the magnitude of the majority of GBRT model 306 inputs with reasonable fidelity, suggesting that the satellite retrievals highlighted in Section 2.2 are 307 suitable inputs for a machine learning model trained on MERRA2 GMI output (Fig.1). Figure 1 compares 308 the distribution of the February training dataset created from the MERRA2 GMI simulation for 2005 – 309 2019 to the satellite observations of the indicated species for February 2017, a month omitted from the 310 training dataset. Distributions of the remaining water vapor layers are shown in Figure S1. In addition, 311 correlations between observations and MERRA2 GMI output for February 2017 are shown, as an 312 example, in Figures S2 and S3. With the exception of HCHO, distributions of the species are similar 313 between the observations and MERRA2 GMI, with the training dataset encompassing the full range of 314 almost all species. A GBRT model trained on MERRA2 GMI will therefore likely not have to extrapolate 315 to photochemical environments on which it was not trained when applied to the satellite data. Further, 316 MERRA2 GMI total column O₃, H₂O_(v) column, AOD, CO, and SSTs are all highly correlated (r^2 of 0.65 or 317 higher) with their respective satellite observations, and biases are within 10%, on average. Anderson et 318 al. (2021) did show that MERRA2 GMI CO columns demonstrate biases of opposite sign in the Northern 319 and Southern Hemispheres, however.

320

321 Agreement between MERRA2 GMI and satellite observations for NO₂, HCHO, and the H₂O_(v) layers is

more variable than for the other species. While modeled NO_2 is moderately correlated with

323 observations ($r^2 = 0.68$) with relatively similar distributions, MERRA2 GMI has a NMB of 63%. This

disagreement is most pronounced at low column values, however, where observational uncertainty is
 large. Further, Anderson et al. (2021) demonstrated distinct regions of bias in NO₂ related to biomass

burning and lightning emissions. Modeled HCHO, on the other hand, is not correlated with observations
 and is biased low by -77%. Modeled water vapor layers are all modestly correlated with observations (r²

of 0.64 or greater) but vary in their bias, with the 925, 850, 700, and 300 hPa layers biased within 30%

- and the remaining layers biased up to 71%.
- 330

331 The satellite product is insensitive to the differences between the HCHO distribution of the satellite and 332 training dataset highlighted in Figure 1. To determine the effects of the difference in HCHO distribution, 333 we extended the training dataset to cover the full time period of the MERRA2 GMI simulation (1980 – 334 2019) and then subsampled the resultant data to match the satellite HCHO distribution. Extending the 335 training dataset to 1980 allows for the subsampled training dataset to have a similar size (~600,000 336 points) as the original training set. We then created a new machine learning model using this sub-337 sampled dataset and calculated OH fields for Feb. 2017 using the satellite inputs from Table 1. We 338 compared this to the TCOH field calculated from a model using the original training dataset, finding 339 agreement within 5%. Similarly, the satellite-constrained TCOH product discussed in Section 4.2 differs 340 by only 3% on average from one determined with a GBRT model that excludes HCHO as an input, 341 suggesting the limited impact of potential errors in the MERRA2 GMI HCHO distribution on model

342 performance. These uncertainties are small in comparison to that resulting from uncertainties in the

343 NO₂ and HCHO satellite retrievals discussed in Section 5.2. If the uncertainty of the satellite inputs

decreases, as retrievals and instruments improve, then it will become necessary to more closely align

345 the training and observed HCHO distributions.

346

Finally, because NO₂ and HCHO have the largest differences between satellite observations and the training dataset, we trained a separate machine learning model to predict TCOH, omitting these two

- 349 species as inputs. When this model was evaluated using the independent MERRA2 GMI output
- described in Section 4.1, the NRMSE was 10.1%, a more than factor of 2 degradation in performance as
- 351 compared to the baseline model. This suggests that omitting these species from the machine learning
- model would result in a greater uncertainty in the final TCOH product than that which results from the
- retrieval uncertainties and the potential discrepancies between observations and the training dataset.
- 354

355 **3.2 Evaluation of the simulated ENSO-related variability of OH drivers**

356 Because ENSO is the dominant mode of OH variability (Anderson et al., 2021; Turner et al., 2018), the 357 training dataset must also capture the ENSO-related variability of the GBRT model inputs. Anderson et 358 al. (2021) demonstrated that the correlation of columns of CO, $H_2O_{(v)}$, and to a lesser extent NO₂, from the MERRA2 GMI simulation with the Multivariate ENSO Index (MEI) (Wolter and Timlin, 2011) agreed 359 360 closely with correlations of the corresponding species for observations from MOPITT, AIRS, and OMI. 361 Unsurprisingly, based on the strong correlation and low bias of MERRA2 GMI SSTs with observations, the 362 simulation also captures the relationship between SSTs and ENSO. The simulation therefore sufficiently 363 captures the ENSO-related variability of these species to act as training data for the GBRT model. We 364 now evaluate this relationship for the remaining GBRT model inputs.

365

366 The MERRA2 GMI-simulated ENSO-related variability of AOD and the various water vapor layers also 367 agrees well with observations. Figures 2 and S4 show the correlation of AOD, HCHO, and the various 368 $H_2O_{(v)}$ layers with the MEI for the satellite retrievals and MERRA2 GMI. MERRA2 GMI captures the 369 general distribution and magnitude of correlation between AOD and ENSO, despite the low optical 370 depths over much of the domain. There are some regional differences, however, particularly in the 371 eastern Southern Hemispheric Pacific. For the H₂O_(v) layers, the simulation underestimates the 372 magnitude of the correlation in some areas, but in general, there is excellent agreement for all layers 373 throughout the troposphere. This suggests that, despite the high bias discussed above, including the 374 $H_2O_{(v)}$ layers could provide important, vertically-resolved information to the machine learning model. 375



376 Regression Coefficient
 377 Figure 2: Distribution of the regression coefficient of a linear least squares fit of the indicated variable against the
 378 MEI for the respective satellite retrieval (a, c, and e) and MERRA2 GMI (b,d, and f) for February. Regressions of
 379 AOD are for 2010 to 2019, the years for which we have a one-degree, gridded satellite product, while HCHO and

water vapor 700 hPa are for 2005 to 2019. Satellite data are on a 1°×1° grid while model output is at the native
 model resolution.

382 Modeled accuracy of the HCHO-ENSO relationship is more difficult to assess. While both the OMI

383 retrieval and MERRA2 GMI demonstrate broad regions of anti-correlation between HCHO and ENSO, the

384 correlations with OMI HCHO are weaker and noisier than for the other satellite retrievals. Over much of

- the domain, HCHO abundance is low, often at or below the retrieval detection limit, suggesting that the
- 386 HCHO retrieval might not be of sufficient quality to capture ENSO-related variability. We investigate the
- 387 impacts of the HCHO observational uncertainty in Section 5.
- 388

 $\label{eq:stars} 389 \qquad \mbox{Finally, because we use total column O}_3 \mbox{ as an input to the GBRT model, we do not evaluate the} \\$

- $\label{eq:solution} 390 \qquad \mbox{relationship between ENSO and O_3, as the stratosphere dominates the O_3 column and the ENSO-related}$
- variability is mostly confined to the troposphere. Oman et al. (2013) found that a GEOS CCM simulation
- and a combination of O_3 retrievals from the Microwave Limb Sounder (MLS) and the Tropospheric
- 393 Emission Spectrometer (TES) exhibited similar ENSO-related variability in the middle and upper
- troposphere, demonstrating that simulations in the GEOS framework can capture this relationship. If a
- 395 TES-like satellite retrieval were currently available, it could be a valuable contributor to the GBRT model
- described here, as it would provide vertically-resolved information about one of the primary drivers ofOH production.
- 398

399 4 Tropical tropospheric column OH constrained with observations of its drivers

We now demonstrate the ability of the GBRT model to determine TCOH. First, we show that the GBRT model can reproduce MERRA2 GMI modeled TCOH from a year independent of the training dataset, a
so-called "hold out set" (Sect. 4.1). We then input satellite data from one month from each season into
the GBRT model to evaluate the realism of the calculated TCOH fields (Sect 4.2).

404

405 **4.1 Evaluation with an independent year from MERRA2 GMI**

406 The machine learning model is able to capture both the magnitude and the variability of TCOH across 407 each season when applied to MERRA2 GMI output from 2017, a year independent of the training 408 dataset. For August 2017 (Fig. 3b), the predicted TCOH is highly correlated with MERRA2 GMI (r² of 409 0.98). TCOH from the machine learning model agrees with the CTM simulation within 4.8% on average. 410 The overall normalized mean bias (NMB) is negligible (-0.1%), although there are some regions of 411 coherent bias (Fig. 3a). Results are similar for February, May, and October 2017 (Fig. S5). The 412 normalized root mean square error for each of these months is comparable to that found for a GBRT 413 parameterization of OH created with a similar methodology that included 27 inputs (Anderson et al., 414

- 414 2022). This suggests that limiting inputs to model variables observable from space does not degrade the 415 ability of the machine learning model to predict TCOH. The low bias and high correlation between the
- ability of the machine learning model to predict TCOH. The low bias and high correlation between the
- 416 GBRT and MERRA2 GMI TCOH for all four months examined here also suggests that any potential
- 417 overfitting by the GBRT model is minimal.





421 against MERRA2 GMI for the same month **(b)**. The r² of a linear, least squares regression, along with the normalized 422 mean bias (NMB) and normalized root mean square error (NRMSE), are also indicated.

423 **4.2 TCOH from satellite observations of its drivers**

- 424 We now apply satellite data from the four months corresponding to the ATom campaign (Aug. 2016,
- 425 Feb. 2017, Oct. 2017, and May 2018) to the GBRT model to determine TCOH fields across the tropics.
- 426 More details about ATom as well as evaluation of the GBRT model with ATom observations are in
- 427 Section 5. We use the satellite observations listed in Table 1, all of which have been averaged to the
- 428 monthly scale and to a $1^{\circ} \times 1^{\circ}$ horizontal resolution. We include only grid boxes with observations for
- all GBRT model inputs and where those observations are within the range of the corresponding inputs
- 430 from the training dataset. Because the satellite inputs for most species exclude grid boxes with a cloud
- 431 fraction greater than approximately 30%, the product presented here represents predominantly clear432 sky conditions.
- 432 433
- 434 The GBRT model and multi-satellite inputs yield TCOH fields that are geophysically credible based on our
- 435 current understanding of OH photochemistry. Although the domain-wide average changes little with
- 436 season, with a minimum of 5.84×10^{12} molecules/cm² in May 2018 and a maximum of 6.35×10^{12}
- 437 molecules/cm² in August 2016, the spatial distribution varies widely among the four months (Fig. 4). In
- 438 both Feb. 2017 and Aug. 2016, TCOH minimizes in the winter hemisphere, consistent with lower OH
- 439 production due to low insolation. The reverse is true for the summer hemisphere. In addition, TCOH
- 440 maximizes in regions with strong continental outflow and along coastlines, regions likely to be impacted
- 441 by anthropogenic and biomass burning emissions of OH drivers.
- 442



- 443 Mean: 6.35x
- **Figure 4:** TCOH calculated with the machine learning model using satellite inputs for the months of each ATom
- deployment: Feb. 2017 (a), May 2018 (b), Aug. 2016 (c), and Oct. 2017 (d). The mean, domain-wide TCOH value in molecules/cm² for each month is also indicated.
- 447 In general, TCOH from the multi-satellite product differs in both magnitude and distribution from the
- 448 MERRA2 GMI simulation. For example, for Feb. 2017, mean MERRA2 GMI TCOH is 6.96×10^{12}
- 449 molecules/cm², 12% higher than the satellite product (Fig. S6). This is consistent with the comparison to 450 *in situ* observations discussed in Section 3.1 where MERRA2 GMI overestimates ATom observations by
- 450 and underestimates CH₄ lifetime, suggesting that the satellite product is again of reasonable
- 451 and underestimates Ch₄ metine, suggesting that the satellite product is again of reasonable 452 magnitude. While understanding the satellite/model differences in TCOH is beyond the scope of this
- 452 magnitude. While understanding the satellite/model differences in FCOH's beyond the scope of the 453 work, we consider the variety in TCOH spatial distributions generated by the GBRT model to be
- 454 promising. The difference between the satellite-constrained product and MERRA2 GMI lends some
- 455 confidence that the GBRT model is not overfit or "tied" to geographic determiners in the training
- 456 dataset, but rather, is sensitive to variations in the chemical and dynamical drivers of OH. These results
- 457 all suggest that the methodology presented here can produce a reasonable satellite TCOH product in the

458 tropics, with values and distributions independent of the chemistry model used to create the GBRT 459 model.

460

461 5 Understanding and mitigating potential challenges in using this methodology to constrain TCOH

462 In this section, we outline possible limitations of the machine learning methodology and the current 463 observational network of the GBRT model inputs and provide potential means to mitigate these 464 limitations where necessary. In section 5.1, we discuss the current lack of sufficient in situ observations 465 to thoroughly evaluate the methodology, highlighting this point by validating the GBRT model with data 466 from the ATom campaign. In section 5.2, we investigate the impacts of random retrieval errors in 467 satellite retrievals on the TCOH product, while in section 5.3, we evaluate the impacts on TCOH when 468 using different satellite retrievals as inputs.

469

470 5.1 Insufficient in situ observations for thorough independent evaluation

471 While we demonstrated in Section 4.1 that TCOH calculated with the GBRT model agrees closely with a 472 hold-out set from MERRA2 GMI, it is also important to demonstrate that the GBRT model can replicate 473

observed TCOH from the actual atmosphere. Because the satellite TCOH product shown in Figure 4 is 474

monthly and at a $1^{\circ} \times 1^{\circ}$ resolution, however, there are no observations with which to evaluate the 475

product. We can test the ability of the GBRT model to reproduce observed TCOH from field campaigns,

476 however, assuming there are concomitant observations of the input species listed in Table 1. The 477 additional need for tropospheric column values of many of these species severely limits the datasets

478 available for validation. To our knowledge, the ATom campaign is the only source of the required inputs

479 with enough observations to attempt a limited validation.

480

481 During ATom (Thompson et al., 2022), scientists measured a suite of air quality and climate relevant 482 trace gases and aerosols throughout the atmosphere above the remote Pacific and Atlantic. ATom took 483 place in four parts: ATom 1 (July – August 2016), ATom 2 (January – February 2017), ATom 3 (September 484 October 2017), and ATom 4 (April – May 2018). During each deployment, flights consisted of a series 485 of ascents and descents across all tropical latitudes over the Pacific and Atlantic Oceans. This allows for 486 the calculation of tropospheric column content of the observed species and evaluation of the machine 487 learning model across most latitudes of our study domain and across all seasons.

488

489 To evaluate the GBRT model performance, we calculated TCOH using a modified GBRT model and

490 observations from the ATom deployments as inputs. We then compared the values to the observed OH

491 columns. To calculate the column values from the observations, we averaged data into 25 hPa pressure

bins for each ATom profile. We filled in missing data using a log-linear interpolation and then integrated 492

493 the column. Our analysis here includes only profiles with observations of all necessary species, that

494 spanned at least 700 hPa, and where less than 25% of the pressure bin values were interpolated. We

495 also omitted any profiles that had pressure bins with negative OH values. In addition, we restrict our

496 analysis to latitudes within 25° of the equator and profiles conducted between 12:00 and 15:00 LST.

497 Values for total column O₃, AOD, and SSTs, for which there were no observations during ATom, were 498 taken from the MERRA2 GMI simulation from the grid box closest to the center of the respective profile.

499 Because ATom profiles did not span the entire tropospheric column, we trained a separate GBRT model

500 where OH and all tropospheric column input variables were substituted with columns spanning 990 -

501 250 hPa, the median range of ATom profiles. This allows for a more direct comparison between

502 observed and modeled TCOH. The spatial distribution of the valid ATom columns and the corresponding

503 columns calculated with the GBRT model are shown in Figure S7.

504



505 506

 $(x10^{12} \text{ molecules cm}^{-2})$

Figure 5: Regression of TCOH observed from the ATom deployments against that predicted from the GBRT model. 507 Error bars represent the 2σ observational uncertainty as reported in Brune et al. (2020) and the GBRT uncertainty 508 described in Section 5.2. The r^2 of a linear least squares fit and the mean bias are also shown.

509 The GBRT model captures the variability of the observed TCOH, and, while there is a modest overall high

510 bias, the median normalized absolute error of 28.3% is within observational uncertainty. When applied

511 to all ATom deployments, predicted TCOH is correlated with the observations with an r^2 of 0.67 and a

512 mean bias of 1.14×10^{12} molecules/cm² (Fig.5). Many of the data points agree within the combined

513 modeled and observational uncertainty. The r^2 values for individual deployments are 0.88 for ATom 1, 514 0.73 for ATom2, and 0.78 for ATom 3 and 4. The level of agreement between observed and predicted

515 OH is comparable or better than that of other methods to infer OH from space. For example, Pimlott et

516 al. (2022) found an r of 0.78 ($r^2 = 0.61$) when estimating ATom OH using a steady state approach, with r

517 values ranging from 0.51 to 0.85 (r^2 of 0.26 to 0.72) for the different deployments. The level of

518 agreement we show here therefore demonstrates the validity of the machine learning method to

519 capture the variability of OH.

520

521 The source of the model/measurement disagreement, with over- and underprediction at low and high

522 column content respectively, is unclear, although there are multiple potential error sources. For 523

example, a typical profile taken during ATom spanned 300 – 400 km in latitude, disconnecting the top 524 and bottom of the profile in space. This is in contrast to the data used to train the model, which were

525 vertical columns over one location. This could lead to a degradation in model performance when

526 applied to ATom, since the columns are not directly analogous to the training dataset. These effects are

527 likely limited because ATom observations are in the remote atmosphere, where the spatial distribution

528 of relevant species is likely to be more homogeneous than over land.

529

530 Further, there is a known interference with the ATom NO₂ observations, suggesting another possible

531 contributor to disagreement between measured and modeled OH. Because of thermal degradation of

532 NO_2 reservoir species, such as organic nitrates and peroxyacetyl nitrate, in the instrument inlet, ATom

- 533 NO₂ observations are likely biased high (Silvern et al., 2018;Shah et al., 2023;Nault et al., 2015). To test
- 534 the potential impact of NO_2 on the predicted OH columns, we applied the ATom observations to a model
- 535 that omits NO₂ as an input. Removing NO₂ increases the r² to 0.74, decreases the mean bias to 0.82 \times
- 536 10¹² molecules/cm², and decreases the median normalized absolute error slightly to 25.7% (Fig. S8).
- 537 These improvements in performance suggest that errors in NO₂ could be contributing to the
- 538 measurement/model differences. Omitting NO₂ does, however, likely introduce additional errors as NO_x
- 539 compounds are essential to OH production in some regions of the atmosphere. When we apply the hold

540 out set from MERRA2 GMI to this model, for example, the NRMSE increases by approximately 50%,

- 541 highlighting the importance of keeping NO₂ as an input variable.
- 542

For more certain evaluation of the GBRT model with observations, greater certainty in the in situ NO2
 observations is needed. Although the in situ observations are insufficient to evaluate the absolute

accuracy of the product, the results presented here demonstrate that a machine learning model trained
 on data from a CTM simulation can capture TCOH variability in the actual atmosphere and suggest that

- 547 predicted OH columns agree with observations within instrumental uncertainty.
- 548

549 **5.2** Impacts of uncertainties in the satellite retrievals on TCOH

In the remote atmosphere where HCHO and NO₂ abundances are low, retrieval uncertainty of an
individual pixel for both species can be on the order of 100% and is often reflective of the *a priori*(González Abad et al., 2015;Lamsal et al., 2021). Given the importance of these species to the GBRT
model as well as to OH chemistry, it is necessary to determine how the propagation of the retrieval
uncertainties from these and other model inputs impacts the predicted TCOH.

555

556 We determined the total uncertainty in TCOH from all inputs as well as the resultant uncertainty from 557 each individual input for Feb. 2017. First, we estimated an average retrieval uncertainty for each input 558 based on reported values in the retrieval files or from the literature (Table S1). We note that for NO_2 559 and HCHO we use a fit uncertainty for a single retrieval. Because we are using monthly-averaged data at 560 $1^{\circ} \times 1^{\circ}$ horizontal resolution, this likely significantly overestimates the actual uncertainty in these 561 retrievals as the random error from individual pixels will tend to cancel when averaged over such large 562 spatial and temporal scales. Our results are therefore an upper bound on the estimated TCOH 563 uncertainty.

564

Next, for each grid box and model input, we created a Gaussian distribution of 2000 values with the modeled value for Feb. 2017 as the mean and the estimated uncertainty as the standard deviation. For each input, we then ran the GBRT model 2000 times to create a distribution of predicted TCOH values for each grid box. The normalized uncertainty in TCOH attributable to a given input is the ratio of the standard deviation of the resultant distribution divided by the mean value. We repeated this process individually for all inputs. In addition, to estimate a total uncertainty in TCOH, we varied all inputs simultaneously with the same Gaussian distributions described above.

572

573 Uncertainty from the NO₂ retrieval, and to a lesser extent HCHO, dominates the total uncertainty in the 574 TCOH product but is of a magnitude comparable to that of in situ OH observations. Median TCOH 1σ 575 uncertainty resulting from NO₂ is 16.5%, with maxima in the remote atmosphere in regions where NO₂ 576 columns are low. Median uncertainty in TCOH resulting from HCHO is 7%, averaged over the study 577 domain, despite the large uncertainty in the HCHO retrieval itself. In contrast to NO₂, uncertainties in 578 TCOH resulting from HCHO maximize in regions with higher HCHO columns (Fig. 6). The magnitude of 579 that uncertainty is likely an overestimate as the actual retrieval uncertainty for HCHO in these regions is 580 significantly lower than the value assumed for the error analysis. In comparison, median TCOH 581 uncertainties resulting from other inputs are 2.9% or less (Figs. S9 and S10). Total TCOH uncertainty is 582 16.6% and is dominated by the NO₂ uncertainty. This uncertainty analysis is in general agreement with 583 the model feature importance (Supplementary Fig. 11), a measure of the relative importance of GBRT 584 model inputs, where HCHO and NO₂ consistently have the largest values of the satellite inputs. 585



586 Median: 16.6% Median: 16.

589 These results demonstrate that the satellite retrieval inputs to the machine learning model are of

590 sufficient quality to produce a meaningful TCOH data product when averaged over large spatial and

591 temporal scales. The 2σ uncertainty in TCOH resulting from the uncertainties in these retrievals is on

the order of that reported for in situ OH observations (Brune et al., 2020). As discussed earlier, this is also likely an upper bound on the uncertainty from random retrieval errors, and uncertainties could be

also likely an upper bound on the uncertainty from random retrieval errors, and uncertainties could be reduced through further averaging, although at the expense of reduced spatial and temporal resolution.

595 Improving the satellite retrievals of NO₂ and HCHO in the remote atmosphere, using retrievals with less

596 noise over the remote atmosphere such as HCHO from OMPS (González Abad et al., 2016), or

incorporating data from satellites with higher resolution, such as TROPOMI, could also reduce the
 uncertainty in their retrievals and thus in TCOH. As discussed in the next section, however, systematic

biases between satellite retrievals can also lead to uncertainties in the TCOH.

600

601 5.3 Sensitivity of TCOH to different satellite retrievals of GBRT inputs

602 The satellite retrievals listed in Table 1 provide the benefit of a long record, with data from most 603 retrievals available from at least 2005 to the present. Such a rich dataset would allow for long-term 604 trend analysis of TCOH. These instruments are near the end of their life cycle, however, so it is 605 instructive to see how retrievals from newer satellites impact the predicted TCOH from the GBRT model. 606 In addition, although these newer satellites, such as TROPOMI, have a significantly shorter observational 607 record than those in Table 1, TROPOMI also has finer spatial resolution and the added advantage of providing retrievals for CO, NO₂, O₃, HCHO, and H₂O_(v). Using retrievals of multiple species from the 608 609 same instrument could negate errors resulting from differences in viewing geometry as well as from

610 overpass time. Here, we investigate the effects of applying retrievals from TROPOMI to the machine

- 611 learning model and compare them to the results from the product described in Section 4, highlighting
- 612 potential impacts resulting from instrumental differences as well as those resulting from differences in 613 retrieval algorithms. The results emphasize the need for thorough retrieval validation in the remote
- 614 atmosphere, particularly of NO₂.
- 615

616 5.3.1 Description of TROPOMI and a modified GBRT model

617 TROPOMI, a successor instrument to OMI, is a spectrometer covering portions of the ultraviolet, visible,

- and infrared spectrum (Veefkind et al., 2012). It is located onboard the Sentinel 5 Precursor satellite,
- 619 which is polar orbiting and has a local overpass time of approximately 13:30. Horizontal resolution for

620 the month examined here (May 2018) is as high as 7 km \times 3.5 km at nadir. We have gridded the Level 2 621 product for each species to a $1^{\circ} \times 1^{\circ}$ resolution and averaged the data to the monthly scale, applying the 622 recommended quality flags and filtering for cloud fraction greater than 30%.

623

624 We use two different retrievals of TROPOMI NO₂ for this analysis. First, we use the KNMI (Royal

- 625 Netherlands Meteorological Institute) NO₂ retrieval (van Geffen et al., 2020), which is based on the
- 626 DOMINO (Dutch OMI NO₂ product) retrieval developed for the OMI instrument. Wang et al. (2020)
- 627 found that this retrieval was biased high when compared to ship-based observations from a MAX-DOAS
- 628 instrument over the remote oceans, while Verhoelst et al. (2021) found good agreement between the
- 629 retrieval and ground-based observations in Reunion. In addition, we use the MINDS (Multi-Decadal
- 630 Nitrogen Dioxide and Derived Products from Satellites) retrieval, which uses the same algorithm as for
- 631 the OMI product described in Section 2 (Lamsal et al., 2022). This retrieval has not been evaluated in 632 the remote tropics.
- 633

634 We also use TROPOMI retrievals of HCHO, $H_2O_{(v)}$ column, total column O_3 , and CO. The HCHO retrieval

- 635 (De Smedt et al., 2018) was found to have a 30% low bias with respect to an OMI retrieval using the 636
- same algorithm due to differences in cloud processing (De Smedt et al., 2021). While evaluation in the 637
- remote tropics is limited, the TROPOMI retrieval does overestimate HCHO in polluted regions (De Smedt
- et al., 2021) when compared to ground-based observations. The TROPOMI H₂O_(v) (Chan et al., 2022) 638 639 retrieval has a slight dry bias with comparison to other satellite products, while the total column O₃
- 640 retrieval (Garane et al., 2019) agrees within 0 - 1.5% with ground-based observations. Finally, the CO
- 641 retrieval (Borsdorff et al., 2019) agrees with MOPITT over the oceans within 3% on average (Martínez-
- 642 Alonso et al., 2020). TROPOMI does not have an equivalent retrieval of the AIRS $H_2O_{(v)}$ layers.
- 643

644 To calculate TCOH using TROPOMI data, we trained a separate machine learning model using all inputs 645 from Table 1 except the water vapor layers, for which there are no TROPOMI retrievals. Removal of the 646 layers from the machine learning model does not significantly degrade performance. For example, for 647 May 2017, removing the $H_2O_{(v)}$ layers from the model, increases the NRMSE from 5.34% to 5.73% when 648 applying the GBRT model to the hold out set. For this new model, we then calculate TCOH using 649 TROPOMI data, including the KNMI NO₂ retrieval. For SSTs and AOD, we use the MUR and MODIS 650 products respectively. While TROPOMI does have an aerosol product, the UV aerosol index, the 651 corresponding output from the MERRA2 GMI simulation is unavailable. We refer to this TCOH as the 652 TROPOMI-KNMI product. We have also calculated TCOH using the satellite retrievals in Table 1, except 653 for the water vapor layers, using this GBRT model, and refer to that as the OMI/MOPITT/AIRS product. 654 We restrict our analysis to May 2018, the only month for which we have TROPOMI water vapor data.

655

656 5.3.2 TROPOMI data applied to the GBRT model

657 TCOH from the TROPOMI-KNMI product is higher than that from the OMI/MOPITT/AIRS product for May 658 2018. Figure 7 shows TCOH calculated from the TROPOMI-KNMI product as well as the percent 659 difference between the two products. While there is modest correlation between the two ($r^2 = 0.63$), 660 the TROPOMI product is 27.6% higher than the OMI/MOPITT/AIRS product, with higher values across 661 almost the entire domain. Differences between the products are most pronounced in the Indian Ocean 662 and off the coasts of Indonesia and the Philippines.

663

664 In general, observations from TROPOMI agree with those from the satellites in Table 1, with the 665 exception of NO₂ and HCHO. Ozone, $H_2O_{(v)}$, and CO from TROPOMI are highly correlated (r² of 0.85 or 666 higher) and agree within 10% on average (Fig. S12) with their respective retrievals from OMI, MOPITT, 667 and AIRS. On the other hand, TROPOMI KNMI-NO₂ is systematically higher (145% on average), and

- **668** TROPOMI HCHO is 20% lower than their corresponding OMI retrievals. The higher TCOH from the
- 669 TROPOMI product is consistent with the increase in NO₂, which would lead to higher secondary
- 670 production of OH. Further, while TROPOMI KNMI-NO₂ is modestly correlated with OMI NO₂ ($r^2 = 0.61$),
- TROPOMI and OMI HCHO are not correlated ($r^2 = 0.23$), highlighting the difficulty of the HCHO retrieval.
- 672 Note that we are not seeking to determine which retrieval, if any, is more accurate. We are highlighting
- 673 the differences to emphasize the impact that systematic differences in retrieval magnitudes of GBRT
- 674 model inputs can have on the resultant TCOH.
- 675



676 677

Figure 7: TCOH for May 2018 determined using TROPOMI inputs, including the KNMI NO₂ retrieval (a). The

difference between the TROPOMI and multi-satellite product is shown in (b). Panel (c) shows the regression of
 TCOH calculated from TROPOMI against that calculated from retrievals from MOPITT, OMI, and AIRS as well as the
 percent difference between the two TCOH products.

681 NO₂ drives the differences between the two TCOH products. To determine the impacts of the different

682 TROPOMI inputs on the TCOH product, we individually swapped each TROPOMI input into the

683 OMI/MOPITT/AIRS product, replacing the corresponding input from Table 1. We then determined the

- difference in TCOH from the OMI/MOPITT/AIRS product that does not include TROPOMI. While this
- 685 method will not yield the exact contribution from a particular retrieval because of the non-linear nature 686 of OH chemistry, it does yield information about the relative importance of each species. Swapping in
- 687 TROPOMI CO, H₂O_(v), and O₃ changed TCOH by less than 2%, while using TROPOMI HCHO increased
- TCOH by 3%. In contrast, TROPOMI NO₂ increased TCOH by 29%, showing that the higher TCOH in the TROPOMI product is driven by differences in NO₂.
- 690

The increased TCOH in the TROPOMI product likely results from a combination of differences in the NO₂
 retrieval algorithm as well as instrumental differences. Comparison of the KNMI and MINDS retrievals

- 693 illustrate this point. When compared to OMI, the MINDS NO₂ retrieval is 58% higher for May 2018, as
- 694 compared to 145% higher for the KNMI retrieval. The closer agreement is unsurprising since the MINDS
- 695 NO₂ uses the same retrieval algorithm as for OMI. Substituting the MINDS NO₂ as an input to the
- 696 TROPOMI product (TROPOMI-MINDS product) reduces the difference with respect to the
- 697 OMI/MOPITT/AIRS product to 18% (Fig. S13). While this is an improvement in agreement, the
- 698 differences in TCOH as well as the lack of change in r² value still suggest that differences between OMI
- and TROPOMI unrelated to the retrieval algorithm account for some of the discrepancy. In addition, the
- training dataset does not take TROPOMI averaging kernels and shape factors into account, which could
- 701 also contribute to the observed differences.

702

The results here demonstrate the sensitivity of the methodology to any systematic bias in the input

retrievals. As with the random error analysis, the level of uncertainty introduced by these biases is low

ros enough to allow for a meaningful OH product. Despite these differences, the methodology to determine

TCOH using machine learning that we have presented here still captures the variability in TCOH,

consistent with the ATom evaluation outlined in Section 5.1. To reduce the uncertainty of TCOH, better

- evaluation of NO₂ in the remote atmosphere is needed to determine which retrievals, if any, are
 accurate.
- 710

711 6 Discussion and recommendations for future observations

The method of estimating clear-sky TCOH presented here has the potential to increase our understanding of the atmospheric oxidation capacity. Because of the long record of observations from MOPITT, OMI, AIRS, and MODIS, we can calculate tropical TCOH from 2005 to the present, and since the methodology is not constrained to a particular satellite, newer satellite missions could extend the dataset beyond the end of these instruments' lifetimes. In addition, this methodology will provide subhemispheric information on OH variability, supplementing information available from MCF inversions.

718

The methodology could be expanded to the extra-tropics and over land, allowing for global constraints on OH. Expansion over land will likely require additional satellite retrievals, like that of isoprene (Wells

et al., 2020), in regions with more complex VOC chemistry than in the remote atmosphere. Expanding

this product beyond the tropics could increase understanding of global CH₄, CO, and VOC trends and

variability and allow for a wider range of satellite retrievals as inputs. For example, current and

vpcoming geostationary air quality satellites such as Sentinel 4, TEMPO (Tropospheric Emissions:

725 Monitoring of Pollution), and GEMS (Geostationary Environment Monitoring Spectrometer) could

726 provide retrievals of most of the necessary inputs to the machine learning model, allowing for the 727 understanding of diurnal variability in TCOH and potentially in the diurnal variability of ozone production

- 728 (Zhu et al., 2022a).
- 729

A similar methodology could likely be used to determine OH at different layers of the atmosphere.

731 Because CH₄ loss is not evenly distributed throughout the tropospheric column, vertically resolved OH

would better help inform this process. Vertically-resolved OH could also help understand differences in

733 OH drivers in the upper and lower troposphere (Spivakovsky et al., 1990;Lelieveld et al., 2016), which

734 can often be decoupled from the column. While column inputs, such as those discussed here, could be

used, the inclusion of vertically resolved satellite retrievals, such as the AIRS $H_2O_{(v)}$ layers, would provide

additional information. Tropospheric O₃ at different atmospheric layers, such as that previously
 provided by the TES satellite, could also be invaluable here, as O₃ is a large driver of primary OH

- 738 production.
- 739

540 Satellite-derived OH would also a provide a much-needed, observational constraint on OH variability in

global chemistry models. Because the methodology can capture variability in TCOH of both

observations and 3-dimensional model output, TCOH trends from a satellite-constrained product could
 be used to evaluate modeled trends and as well as the spatial variability resulting from events like ENSO.

745 While the satellite-derived OH could not explicitly indicate the cause of differences, the spatial

744 While the satellite-derived OF could not explicitly indicate the cause of differences, the spatial 745 distribution of the differences as well as differences in observed and modeled machine learning model

746 inputs could indicate potential dynamical or emission sources of error in the 3D model.

747

Further, the combination of the satellite-derived OH and the machine learning model could help identify
 the impacts of any diagnosed errors in emissions inventories as well as the impacts of unexpected

- 750 events, such as COVID-19-related shutdowns, on TCOH. For example, if there are significant
- discrepancies between observed and modeled NO₂ in a specific region of the atmosphere, the satellite
- NO₂ could be scaled to more closely match the 3D model values and then be input into the machine
- 753 learning model. The difference in TCOH would then indicate the relative impact of the model error. This
- vould serve as a computationally efficient complement to other methodologies constraining models
- with observations (e.g. Miyazaki et al., 2020; Miyazaki et al., 2021) to identify the impacts of these errors
- on the atmospheric oxidation capacity. A similar methodology could be used for unexpected events that
 significantly impact emissions of OH drivers, allowing for quick determination of their potential impacts
- 758 on the atmospheric oxidation capacity before emissions inventories could be revised.
- 759
- 760 While we have shown that the methodology captures the variability of observed OH and generally
- agrees with observations within measurement uncertainty, it is unclear whether differences result from
- 762 GBRT model deficiencies or structural differences between the *in situ* observations and the training
- dataset. Additional field campaigns with observations of OH and the GBRT model inputs would allow for
- a more thorough evaluation of both the OH product and the methodology itself. Such a field campaign
- vould need to provide complete tropospheric columns of all species and cover less horizontal distance
- than the ATom profiles (e.g. from spiral flight patterns). In situ observations of NO₂ without significant
- interference from NO_X reservoir species are also needed to reduce uncertainty. Alternatively, NO_2 and
- other species could be measured through aircraft-based remote sensing. Finally, repeated sampling over
- the same locations for multiple days within a defined area would allow for meaningful statistical analysis
 while also allowing for the comparison of TCOH columns calculated from satellite observations.
- 771
- Finally, accuracy of the TCOH product is dependent on the accuracy of the satellite retrievals input into
 the machine learning model, with the NO₂ retrieval having the largest effect. To reduce the uncertainty
- of the TCOH product, more information about the accuracy of individual NO₂ retrievals is required.
- 775 Currently, there is little validation of OMI and TROPOMI NO₂ retrievals in the remote, tropical
- atmosphere, so it is difficult to assess which retrievals, if any, are correct. Recent efforts, such as the
- 777 QA4ECV (Quality Assurance for the Essential Climate Variables), to improve NO₂ retrieval algorithms
- have reduced uncertainty, particularly over land (Boersma et al., 2018), although it is unclear how the
- accuracy of these retrievals translates to the remote tropics as validation data are still extremely limited.
- 780 Even retrievals of TROPOMI and OMI made with the same algorithm show differences, suggesting that
- 781 instrumental differences could also affect the results. Future satellite missions should focus on trying to
- reduce the uncertainty in NO₂ retrievals, particularly in the remote atmosphere, both through
- 783 improvements in instrument design and algorithm development.

784785 7 Data Availability

- 786 Output from the MERRA2 GMI simulation are publicly available at <u>https://acd-</u>
- 787 <u>ext.gsfc.nasa.gov/Projects/GEOSCCM/MERRA2GMI/</u> (NASA Goddard Space Flight Center, 2023). All
- 788 satellite products, except for TROPOMI water vapor, are available at <u>https://disc.gsfc.nasa.gov (GES</u>
- 789 <u>DISC, 2023</u>). Data from the ATom campaign are located at https://daac.ornl.gov (Wofsy et al., 2021).
- 790

791 8 Author contributions

- 792 DCA wrote the manuscript, performed the data analysis, and created the GBRT model. DCA, BND, JMN,
- and MBFC developed the idea for the methodology. SAS performed three-dimensional modeling for the
- work. JMN provided advice on machine learning. JL helped perform data analysis. All authors helped
- 795 develop ideas for the analysis and contributed to the manuscript.
- 796
- 797 9 Competing Interests

- BND is a member of the editorial board of Atmospheric Chemistry and Physics. The peer-review process
 was guided by an independent editor, and the authors also have no other competing interests to
- 799 was guid800 declare.
- 800 801

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810 10. References

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