1 Unraveling Pathways of Elevated Ozone Induced by the 2020

2 Lockdown in Europe by an Observationally Constrained Regional

3 Model using TROPOMI

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15 Abstract. Questions about how emissions are changing during the COVID-19 lockdown 16 periods cannot be answered by observations of atmospheric trace gas concentrations alone, in part 17 due to simultaneous changes in atmospheric transport, emissions, dynamics, photochemistry, and 18 chemical feedback. A chemical transport model simulation benefiting from a multi-species 19 inversion framework using well-characterized observations should differentiate those influences 20 enabling to closely examine changes in emissions. Accordingly, we jointly constrain NO_x and 21 VOC emissions using well-characterized TROPOMI HCHO and NO₂ columns during the months 22 of March, April, and May 2020 (lockdown) and 2019 (baseline). We observe a noticeable decline 23 in the magnitude of NO_x emissions in March 2020 (14-31%) in several major cities including Paris, 24 London, Madrid, and Milan expanding further to Rome, Brussels, Frankfurt, Warsaw, Belgrade, 25 Kyiv, and Moscow (34-51%) in April. However, NO_x emissions remain at somewhat similar 26 values or even higher in some portions of the UK, Poland, and Moscow in March 2020 compared 27 to the baseline possibly due to the timeline of restrictions. Comparisons against surface monitoring 28 stations indicate that the constrained model underrepresents the reduction in surface NO_2 . This 29 underrepresentation correlates with the TROPOMI frequency impacted by cloudiness. During the 30 month of April, when ample TROPOMI samples are present, the surface NO₂ reductions occurring 31 in polluted areas are described fairly well by the model (model: $-21\pm17\%$, observation: $-29\pm21\%$). 32 The observational constraint on VOC emissions is found to be generally weak except for lower

33 latitudes. Results support an increase in surface ozone during the lockdown. In April, the 34 constrained model features a reasonable agreement with maximum daily 8 h average (MDA8) 35 ozone changes observed at the surface (r=0.43), specifically over central Europe where ozone 36 enhancements prevail (model: +3.73±3.94%, +1.79 ppbv, observation: +7.35±11.27%, +3.76 37 ppbv). The model suggests that physical processes (dry deposition, advection, and diffusion) 38 decrease MDA8 surface ozone in the same month on average by -4.83 ppbv, while ozone 39 production rates dampened by largely negative $J_{NO2}[NO_2]-k_{NO+O3}[NO][O_3]$ become less negative, 40 leading ozone to increase by +5.89 ppby. Experiments involving fixed anthropogenic emissions 41 suggest that meteorology contributes to 42% enhancement in MDA8 surface ozone over the same 42 region with the remaining part (58%) coming from changes in anthropogenic emissions. Results 43 illustrate the capability of satellite data of major ozone precursors to help atmospheric models 44 capture ozone changes induced by abrupt emission anomalies.

45 **1. Introduction**

46 Continuous monitoring of air pollution by satellites can help our understanding of both 47 anthropogenic and biogenic variability and change caused by rapid economic recession 48 [Castellanos and Boersma, 2012] and regulations [Krotkov et al., 2016; Souri et al., 2020a]. Earth's 49 atmosphere has substantially become more polluted since the industrial era in comparison to its 50 original environmental condition [Li and Lin, 2015], thus any abrupt hiatus in anthropogenic (man-51 made) emissions should result in an immediate impact on relatively short lifetime pollutants such 52 as nitrogen dioxide (NO₂), formaldehyde (HCHO), and tropospheric ozone (O₃). The beginning of 53 the global COVID-19 pandemic in early 2020 [Fauci et al., 2020] provided such an abrupt change 54 in human activities [Le Quéré et al., 2020]. A first step to fully understand how much of these 55 impacts are related to the pandemic lockdowns is to disentangle the physiochemical processes 56 determining their ambient concentrations. Unraveling those processes require precise, continuous 57 observations of physical and chemical states and emission rates, which are not routinely available 58 on global, continental and regional scales. Therefore, we resort to using a model realization 59 attempting to reproduce such an intricate system. Models without observational guidance are 60 incapable of numerically representing the real world [Lorenz, 1963], so our best option to improve 61 a model is to constrain some of its prognostic inputs using well-characterized observations. 62 Accordingly, the framework of this study is centered around inverse modeling and data 63 assimilation.

64 Significant attention has been given to documenting the lockdown-related changes in 65 atmospheric composition around the world using both in-situ and satellite observations [e.g., 66 Sicard et al., 2020; Shi and Brasseur, 2020; Lee et al., 2020; Salma et al., 2020; Le Quéré et al., 67 2020; He et al., 2020; Le et al., 2020; Miyazaki et al., 2020; Liu et al., 2020; Barré et al., 2020; 68 Goldberg et al., 2020; Ordóñez et al., 2020; Wyche et al., 2021; Bekbulat et al., 2020; Gaubert et 69 al., 2021; Sun et al., 2021]. The broad picture is consistent among these studies; the lockdown 70 drastically reduced the concentrations of NO_x , CO, and SO_2 and some types of particulate matter, 71 whereas the concentrations of several secondarily formed compounds such as ozone behaved in 72 non-linear ways due to emissions and/or meteorology.

The motivations of this study are to determine the capability of a regional model constrained by satellite HCHO and NO₂ columns to capture near-surface pollution, and if local ozone production rates are the driving factors for heightening ozone pollution during the 2020 lockdown. In other words, what physiochemical processes are associated with the elevated ozone? How representative are satellite observations at capturing surface air quality through an inversion context? Is meteorology the primary factor in shaping elevated ozone as suggested by Ordóñez et al. [2020]?

80 To address these pivotal questions, it is desirable to constrain models using multi-species 81 observations because relationships between the atmospheric compounds such as HCHO and NO₂ 82 are importantly intertwined [Marais et al., 2012; Valin et al., 2016; Wolfe et al., 2016; Souri et al., 83 2020a,b]. Accordingly we build our inversion framework upon a non-linear joint analytical 84 inversion of NO_x and VOCs proposed in Souri et al. [2020a] using the TROPOspheric Monitoring 85 Instrument (TROPOMI) HCHO and NO₂ observations in Europe. Performing this type of 86 inversion not only enables us to precisely quantify the changes in emissions (along with its 87 uncertainty, as the inversion framework is analytical) but also paves the way for estimating the 88 resulting changes on different pathways of ozone.

89 2. Measurements, Modeling, and Methods

90

2.1. Satellite Observations

91 2.1.1. TROPOMI NO₂

92We use daily offline S5P TROPOMI tropospheric NO2 slant columns93[Copernicus Sentinel data processed by ESA and Koninklijk Nederlands94Meteorologisch Instituut (KNMI), 2019] derived from a two-step framework

95 involving DOAS spectral fitting in conjunction with a stratosphere/troposphere
96 decoupler [Boersma et al., 2018]. The time periods of this study are March, April,
97 and May 2020 and 2019. The data provide Jacobians of light intensity with respect
98 to optical thickness (i.e., vertically-resolved scattering weights) which are
99 dependent on scene surface reflectivity, the cloudiness of the assumed Lambertian
100 clouds, and the sensor viewing geometry.

101Aerosol effects on the scattering weights are not taken into consideration.102Based on radiative transfer calculations and satellite-based aerosol products, Jung103Y. et al [2019] and Cooper et al. [2019] observed small changes (<10%) in AMFs</td>104with and without considering the aerosol impacts in Europe in springtime. This105tendency likely results from a low aerosol optical depth.

106 The 2019 TROPOMI observations used in this study have a spatial resolution of 7×3.5 km², whereas those in 2020 have a spatial resolution of 5.5×3.5 107 108 km². The NO₂ products for the study time period were produced by processor 109 versions v01.02.02 (1 March 2019 - 20 March 2019) and v01.03.02 (20 March 110 2019 onward). The v01.03.02 processor includes an update to the FRESCO-S cloud 111 algorithm and improvements to a quality flag variable. NO₂ validation from 112 processors v01.02.02 and v01.03.02 shows similar biases and dispersion [Lambert 113 et al., 2020], as do comparisons from before and after the pixel spatial resolution 114 change [Verhoelst et al., 2021]. We extract good quality pixels based on the main 115 quality flag (qa flag) > 0.75, which removes retrievals flagged as bad and pixels 116 over snow/ice or with cloud radiance fractions > 0.5, and resample them to our 15-117 km regional model (discussed later) using the bilinear interpolation. Since vertical 118 column densities (VCDs) depend on assumed gas profile shape (i.e., they are quasi-119 observations), we recalculate those shape factors using profiles from our 120 constrained chemical transport model. Shape factors are re-estimated by calculating 121 the ratio of the vertical column of total air to the simulated vertical column of NO_2 122 multiplied by the mixing ratios of NO₂ profile from the regional model [Martin et 123 al., 2002].

124Satellite remote sensing observations are usually far more stable than they125are accurate. This can make the data practical for measuring relative changes in

126 emissions, but may necessitate the use of a bias correction for absolute emissions 127 estimates. Moreover, the systematic and random errors associated with satellite 128 retrievals may differ markedly from location to location. It is therefore crucial to 129 thoroughly validate columns against independent observations. To this end, we 130 compile statistics reported in several validation studies focusing on the TROPOMI 131 tropospheric NO₂ product and summarize their findings in Table 1. The most 132 comprehensive global study to date is a comparison of TROPOMI tropospheric 133 NO₂ with that derived from 19 MAX-DOAS instruments [Verhoelst et al., 2021]. 134 This study indicates there is a low bias in TROPOMI tropospheric NO₂ of -23 to -135 37% relative to MAX-DOAS at clean to moderately polluted sites, and as large as 136 -51% at highly polluted sites. When considering all sites, the overall median bias in this study was found to be -37%, with a dispersion of 3.5×10^{15} molec/cm² 137 (defined as half of the 68% interpercentile). No obvious seasonal patterns were 138 139 found in the biases. These results are consistent with other validation studies which 140 have observed a low bias in TROPOMI tropospheric NO₂ [Chan et al., 2020; Griffin 141 et al., 2019; Judd et al., 2020]. A potential significant source of bias in polluted 142 regions is the relatively low spatial resolution $(1 \times 1^{\circ})$ TM5-MP prior profiles used 143 in the TROPOMI air mass factor calculation. Several validation studies have shown 144 the low bias in TROPOMI NO₂ can be reduced in polluted regions by 5-17% 145 through the use of higher spatial resolution model a priori profiles or other 146 improvements in the AMF calculation [Chan et al., 2020; Griffin et al., 2019; Judd 147 et al., 2020; Zhao et al., 2020].

Directly incorporating these numbers into an inversion model is challenging, mainly because of spatiotemporal variability in the satellite errors. Ideally, the relationship between errors and retrieval inputs (e.g., albedo, scene radiance, profiles, etc.) would be used as an additional cost function in the inversion, commonly known as variational bias correction [e.g., Auligné et al., 2007]. In the absence of such relationships, we use the biases reported in the validation studies.

155In the case of NO2, we uniformly scale up the satellite tropospheric columns156by 25%. This bias estimate is derived by first assuming a 37% low bias in the

157 columns over polluted regions as reported by Verhoelst et al. [2021]. In turn, this 158 low bias can be mitigated somewhat by the application of high spatial resolution 159 profiles in the air mass factor calculation, such as the ones used in this study. Table 160 1 summarizes the results from several TROPOMI validation studies at specific 161 locations that calculated NO₂ using model profiles with higher spatial resolution 162 than the operational TROPOMI $(1^{\circ} \times 1^{\circ})$ profiles (see Table 1 columns 163 "Modification" and "Modified Bias"). In these studies, modified columns show 164 increases ranging from 0 - 25%. Based on these results, we assume a low bias of 165 37% can be mitigated by $\sim 12\%$ through the use of high spatial resolution profiles. 166 for a resulting total low bias of 25%. This bias is likely not valid over pristine areas, 167 where validation studies show lower biases in TROPOMI NO₂ [Verhoelst et al., 168 2021, Wang et al., 2020, Zhao et al., 2020]; nonetheless, we previously observed 169 in Souri et al. [2020a] that the low signal-to-noise ratios of those column amounts 170 resulted in small changes in the top-down emissions. We assume the errors of 171 observations originate from two main sources: i) the precision error provided with 172 the data (e_{precision}) and ii) a fixed error estimated from comparisons to in-situ 173 measurements (e_{const}). Mathematically, the final error is:

$$e_0^2 = e_{const}^2 + \frac{1}{n^2} \sum_{i=1}^n e_{precision,i}^2$$
(1)

174where *n* is the number of samples for a given grid and e_{const} equals to 1.1×10^{15} 175molec/cm² (<6×10¹⁵ molec/cm²) in clean regions and 3.5×10^{15} molec/cm²176(>=6×10¹⁵ molec/cm²) in moderately to highly polluted regions. These regions are177defined based on the wide ranges reported in Verhoelst et al. [2021] (3-14×10¹⁵178molec/cm² for moderately to highly polluted regions).

179 *2.1.2. MODIS AOD*

180To improve the simulation of total aerosol mass, we use the collection 6181MODIS aerosol optical depth (AOD) from both Aqua (~ 13:30 LT) and Terra (~18210:30 LT) platforms over both land and ocean [Levy et al., 2013] (available at183<u>https://ladsweb.modaps.eosdis.nasa.gov</u>, access May 2020). We independently184validate all three major products, namely the deep blue, the dark target and a185combined dark blue products by comparing to AOD values measured by

- 186 AERONET over Europe at the same time period of this study. Only good and very 187 good (qa>=2) pixels are selected for the comparison. The AERONET AOD data 188 are computed based on the values at 500 nm and Angstrom Exponent in the 440-189 675 nm range. We collocate two datasets if they are within 10 km radius and less 190 than 30 mins apart. The dark blue product results in the best agreement (r>0.87) 191 with a high bias of <0.05 (Figure S3, and S4). This product is therefore chosen for 192 the data assimilation. We remove the bias and assign the value of the covariance 193 matrix of observations to the RMSE values obtained from the comparison.
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2.2. Surface Measurements

195 UV photometry and chemiluminescence surface ozone and NO₂ measurements all 196 over continental Europe are used to investigate possible changes in their concentrations 197 induced by the lockdown (https://discomap.eea.europa.eu/map/fme/AirQualityExport.htm, 198 access June 2020). The NO₂ chemiluminescence measurements are usually overestimated 199 due to interferences from the NO_z family (PAN, organic nitrate, HNO₃, etc.). We assume 200 that the interferences are not significantly different between the baseline and lockdown 201 mainly due to relatively low photochemistry in early spring [Lamsal et al., 2008] compared 202 to summertime. Additionally, the correction needs a careful evaluation of the model with 203 regards to the NO_z family whose measurements are not available in this case study.

More than 6450 meteorological stations archived on NOAA's integrated surface database (<u>https://www.ncei.noaa.gov/data/global-hourly/</u>, access April 2020) are used to validate the performance of our weather model in terms of several prognostic inputs including ambient air temperature, air humidity, and U and V wind components.

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2.3. WRF-CMAQ Modeling

209 The regional air quality simulations at 15×15 km² are carried out with the widely 210 used CMAQ v5.2.1 (https://doi.org/10.5281/zenodo.1212601) in conjunction with WRF 211 v3.9.1 [Skamarock et al. 2008] models. The models overlap and cover continental Europe 212 and some portions of Africa and Middle East. The domain consists of 483 east-west, 383 213 north-south grids, and 37 unevenly spaced eta levels (Figure 1). The simulation time period 214 is from March to May 2019 and 2020 (six months). Since IC/BC are taken from already 215 spun-up National Centers for Environmental Prediction (NCEP) FNL (final) reanalysis and 216 GEOS-Chem v12.9.3 (10.5281/zenodo.3974569) runs, we only spin up the models for the 217 month of February. The chemistry configuration of the CMAO model mainly consists of 218 CB05 with chlorine chemistry (gases) and AERO6 (aerosol). Hourly-basis biogenic 219 emissions are processed by the offline standalone Model of Emissions of Gases and 220 Aerosols from Nature (MEGAN) v2.1 model [Guenther et al., 2012] based on high-221 resolution plant functional maps made by Ke et al. [2012]. The biogenic emission factors 222 are estimated based on the PFT-specific information provided in Guenther et al. [2012]. 223 The biogenic VOCs include a wide range of compounds including isoprene, monoterpenes, 224 aromatic VOCs, and methanol. Soil NO_x emissions are estimated by Yienger and Levy, 225 [1999]. Lightning NO_x emissions are based on in-line calculations involving convective 226 precipitation rates and cloud vertical distributions. Lightning NO_x emissions are not 227 constrained in the model. Anthropogenic emissions are based on the Community Emissions 228 Data System (CEDS) inventory in 2014 [Hoesly et al., 2018]. Diurnal scales are not 229 considered for the anthropogenic emissions. We also output the CMAQ integrated process 230 analysis quantifying the contribution of each process to the amount of compounds. The 231 physical setting of WRF includes the Lin microphysics scheme [Lin et al., 1983], the Grell 232 3-D ensemble cumulus scheme [Grell and Dévényi, 2002], the RRTMG radiation scheme, 233 ACM2 planetary boundary layer parametrization [Pleim, 2007], and Pleim-Xu land-234 surface scheme [Xiu and Pleim, 2001]. To minimize the deviation of the model from the 235 reanalysis data, we turn on the grid nudging option with respect to wind, moisture, and 236 temperature only outside of the PBL region. The inclusion of this option only outside of 237 the PBL region is because we do not want the coarse reanalysis data wash out the relatively 238 high-resolution dynamics. Moreover, leaf area index and the sea surface temperature are 239 updated every 6 hours based on satellite measurements included in the reanalysis data. 240 Extensive model evaluations based upon surface observations show a striking 241 correspondence (Table S1, S2) which is indicative of reasonable energy budget and 242 transport in our model.

243

2.4. Inverse Modeling and Data Assimilation

To adjust the bottom-up emission inventories, we follow a non-linear joint inversion method proposed in Souri et al. [2020a]. Briefly, a Gauss-Newton algorithm is utilized to incrementally solve the Bayes' quadratic function in analytical fashion. The posterior emissions are then derived by

$$\mathbf{x}_{i+1} = \mathbf{x}_a + \mathbf{G}[\mathbf{y} - F(\mathbf{x}_i) + K_i(\mathbf{x}_i - \mathbf{x}_a)]$$
⁽²⁾

248 where *y* is bias-corrected monthly-averaged TROPOMI NO₂ and HCHO observations (see 249 S.A1), \mathbf{x}_a (or \mathbf{x}_0) is the prior emissions, \mathbf{x}_i is the posterior emission at the *i*th increment, *F* 250 is the forward model (here WRF-CMAQ) to project the emissions onto columns space, **G** 251 is the Kalman gain,

$$\mathbf{G} = \mathbf{S}_e \ K_i^T \left(K_i \mathbf{S}_e \ K_i^T + \mathbf{S}_o \right)^{-1}$$
(3)

252 and K_i (= $K(\mathbf{x}_i)$) is the Jacobian matrix calculated explicitly from the model using the finite 253 difference method by perturbing separately NO_x and VOC emissions by 20%. The 254 perturbations are applied for each iteration. The model outputs along with Jacobians and 255 emissions are spatiotemporally co-registered with the observations. S_0 and S_e are the error 256 covariance matrices of the observations and emissions. Similar to Souri et al. [2020a], the 257 prior errors in anthropogenic NO_x and VOCs emissions are set to 50% and 150%. 258 respectively. In terms of the biogenic emissions, the errors are set to 200% for both NO_x 259 and VOCs. The instrument covariance matrices are populated with squared-sum of the 260 aforementioned errors based on the compilation of the validation studies and precision 261 errors provided with the data (Eq.1). Both error matrices are assumed diagonal. The 262 inversion window is monthly meaning we have three separate correction factors in months 263 of March, April, and May. The covariance matrix of the a posteriori is calculated by:

$$\hat{\mathbf{S}}_e = (\mathbf{I} - \mathbf{G}\hat{K} \)\mathbf{S}_e \tag{4}$$

264 where \hat{K} is the Jacobian from the *ith* iteration. Here we iterate Eq.2 three times. The 265 averaging kernels (A) are given by:

$$\mathbf{A} = \mathbf{I} - \hat{\mathbf{S}}_e \mathbf{S}_e^{-1} \tag{5}$$

- 266Not only does this method considers non-linear chemical feedback among NO2-267HCHO-NOx-VOC by simultaneously incorporating the HCHO and NO2 in the inversion268framework, it also permits quantification of A that explicitly explains the amount of269information obtained from the observation. Low A indicates low G making the a posteriori270to be rather independent of the observational constraint.
- An important caveat with this inversion system is that we do not take the model parameter error (such as errors in chemistry, cloud microphysics, and PBL) into account. To properly estimate the forward model parameter errors, one needs to calculate the

274 sensitivity matrix of the columns to the model parameters combined with the sensitivity 275 matrix of the columns to the emissions (*K*) [Rodgers, 2000]. The former calculation is 276 computationally expensive. Moreover, the spatiotemporal varying model parameter errors 277 may not be known in detail. The consequence of disregarding the model parameter errors 278 is the overconfidence in the top-down estimates (i.e., overestimations of AKs).

We also correct total aerosol mass by daily assimilating the MODIS dark blue AOD observations following the algorithm discussed in Jung et al. [2019]. Briefly, the assimilation framework uses a modified optimal interpolation method adjusting uniformly all relevant aerosol masses in a column as a function of a weighted-distance and appropriate errors.

284 **3. Results and Discussion**

285

3.1. Variability of NO₂ columns seen by TROPOMI

286 We assess difference maps of NO₂ columns (and HCHO in S1) in 2020 with respect to 287 those in 2019 during the months of March, April and May. The difference maps along with the 288 absolute values of the tropospheric NO₂ columns are shown in Figure 2. Regardless of the year, 289 we observe a noticeable reduction in NO₂ as we approach warmer months which can be explained 290 by increases in OH concentrations (higher water vapor content, solar radiation, and O₃ levels), 291 faster vertical mixing due to larger sensible fluxes (more diluted columns for a given receptor due 292 having a greater chance of experiencing stronger winds in higher altitudes), and a reduction in 293 temperature-dependent light-duty diesel NO_x emissions [Grange et al., 2019]. This sequential 294 decline of NO₂ obscures the quantitative interpretation of the satellite observations in two ways: 295 first, as noted by Silvern et al. [2019], the free tropospheric background NO₂ levels, which are 296 highly uncertain, becomes comparable to those located at near-surface, and second, the relatively 297 lower signal-to-noise ratios reduce the amount of information that we can obtain for inverting NO_x 298 emissions (discussed later).

The anomaly map (2020 vs 2019) in March indicates pronounced decreases in tropospheric NO₂ columns over several countries including France, Spain, Italy, and Germany (box A). In contrast, we see increases in the magnitude of the NO₂ columns over some portions of the UK excluding London (box B), northeastern Germany (box C), and Moscow, Russia (box D). A recent study [Barré et al. 2020] observed roughly the same tendency which was attributable to meteorological changes. While those changes are indeed an important piece of information, we

305 should recognize that the degree of the enforced restrictions varies temporally; moreover changes 306 in emission heavily rely on the dominant emission sector (e.g., mobile or industry). For instance, 307 according to TASS press [https://tass.com/society/1144123, accessed Sep 2020], Russian 308 governments did not take significant measures to control the virus before April 15, immediately 309 evident in the large NO₂ enhancement over Moscow in March (box D). During the next two months 310 (April and May), we observe a major turnaround over this city (box F and H). In May, the anomaly 311 of the tropospheric NO₂ suggests that the reduction in NO_x emissions abruptly experiences a hiatus 312 in central Europe (box G). However it is crucial to note that these maps are based upon sporadic 313 clear-sky pixels that might obscure the full portrayal of emissions changes happening throughout 314 the period (discussed later).

315

3.2. Top-Down estimates of NO_x emissions

Following the inversion and the data assimilation frameworks, we adjust the total amounts 316 317 of VOC, NO_x emissions, and aerosols mass using TROPOMI HCHO, NO₂ and MODIS AOD 318 observations. We focus on the topic of gas phase chemistry (i.e., ozone and its precursors) implying 319 that the aerosol data assimilation is carried out to partially remove errors associated with radiation 320 [e.g., Jung et al., 2019] or heterogenous chemistry [Jacob, 2000]; therefore, the aspect of aerosol 321 changes induced by the lockdown will be examined in a separate study. Furthermore, we observe 322 a relatively weak observational constraint from TROPOMI HCHO on VOC emissions, especially 323 in higher latitudes; accordingly, the relevant discussion on this subject is presented in S2.

The spatial distributions of magnitude of the top-down NO_x and their corresponding changes and averaging kernels are shown in Figure 3. Moreover, the monthly values of the a posteriori and the a priori are shown in Figure S5 and S6. It is worth emphasizing that we use identical prior values in terms of anthropogenic emissions in both years.

328 According to Figure 3, large averaging kernels associated with NO_x emissions are confined 329 in high-emitting regions suggesting that the most valid estimates can be found in areas undergoing 330 strong TROPOMI NO₂ signals. We observe an improvement in the statistics associated with 331 simulated surface NO₂ using the posterior emissions compared to the surface measurements in 332 many places around Europe with an exception to northeastern Germany where TROPOMI NO₂ 333 observations deviates the model from the measurements (Figs S9-S12; Tables S3, S4). The large 334 underestimation of the model in terms of surface NO₂ concentrations is most likely due to the 335 underestimation of the CEDS inventory [e.g., Figure 11 in Sun et al., 2021]. However, it is worth

336 noting that the disagreements between the model and the surface measurements do not solely 337 reflect the uncertainty in the emissions. A major complication arises from the fact that the point 338 measurements represent concentrations locally, whereas the model grids $(15 \times 15 \text{ km}^2)$ are (at best) 339 the average of infinitesimal points integrated over the grid space. Essentially, no one should expect 340 that these quantities will completely line up, unless one transforms the point measurements to the 341 grids (i.e., rasterization) by carefully modeling the spatial auto-correlation (or semivariograms) of 342 the point data [Souri et al., 2021]. Additionally, there is uncertainty about the chemical mechanism 343 utilized in the model. In particular, Souri et al. [2017] observed a large overestimation (~ factor 4) 344 of daily-averaged total nitrate (HNO₃ + NO₃⁻) in the CB05/AERO6 mechanism despite moderately 345 reasonable nitrate (NO_3^-) simulations. This was attributed to a large overestimation of N_2O_5 346 hydrolysis rate [Bertram and Thornton, 2009] which is the primary loss pathway of NO_x in low 347 photochemically active regions [Shah et al., 2020]. The interferences from the NO_z family on the 348 surface measurements might be still present in springtime in midlatitudes ($\sim 10-30\%$) [Lamsal et 349 al., 2008]. Last but not the least, the PBL parameterization controlling the level of vertical mixing 350 rates has errors primarily due to soil moisture not being observationally constrained in the model 351 [Huang et al., 2021].

352 The discrepancies between the simulated tropospheric NO₂ columns versus TROPOMI are 353 mitigated by the inversion (Figure S13 and S14). Immediately apparent in Figure 3 is a strong 354 correlation between anomaly maps of TROPOMI tropospheric NO₂ (Figure 2) and those of top-355 down emissions. We observe reductions in NO_x emissions in March (14-31%) in several major 356 cities including Paris, London, Madrid, and Milan; the reductions further expand to Rome, 357 Brussels, Frankfurt, Warsaw, Kyiv, Moscow, and Belgrade with higher magnitudes (34-51%) in 358 April. In general, the level of NO_x reduction is somewhat higher in April relative to months of 359 March and May possibly due to temporal variabilities associated with the restrictions; for example, 360 UK and Poland governments enforced the restrictions starting in the last week of March to the 361 middle of April (see Figure S1 in Okruszek et al. [2020]; https://www.bbc.com/news/uk-362 51981653, accessed in March 2020). The decreased anthropogenic NO_x emissions in the strait of 363 Gibraltar and Alboran Sea reveal reportedly reduced ship activities [United Nations Conference 364 on Trade and Development Report, Accessed Dec 2020]. The numbers in May indicate that several 365 countries in central and eastern Europe (shown in box G in Figure 2) likely eased coronavirus lockdown restrictions, a picture that has yet to be verified by surface measurements (discussedlater).

368 3.3. Disparities in near-surface concentrations suggested by the constrained model versus 369 those by in-situ measurements

370 **3.3.1**. **NO**₂

371 It is necessary to examine whether the constrained model can precisely represent the 372 changes observed by surface measurements. Several factors can complicate this analysis: i) having 373 overconfidence in the constrained model where the satellite observations used were uncertain; this 374 problem can be addressed by considering grid cells whose averaging kernels are above a threshold 375 (here 0.5), ii) not accounting for spatial representativity function when it comes to directly 376 comparing two datasets at different scales (i.e., point measurements vs the model grids); a 377 statistical construction of the spatial representivity function [Janic et al., 2016; Souri et al., 2021] 378 requires a dense observational network so that we can build a semivariogram; instead, we only 379 consider model grid cells having more than two stations; those observations then are then averaged, 380 iii) interferences of the NO_z family on NO₂ chemiluminescence measurements [Dickerson et al., 381 2019] which can be partly discounted when calculating differences, iv) model uncertainties, 382 especially with respect to turbulent and convective fluxes that are heavily determined by 383 representing local heterogeneity of forces and non-hydrostatic dynamics [Emanuel, 1994], all of 384 which are challenging to fully resolve in a 15-km resolution.

385 With these caveats in mind, we plot the daily-averaged changes of surface NO₂ 386 concentrations in 2020 relative to 2019 derived by the model and the European air quality network 387 for the months of March, April, and May (Figure 4). Large gaps in Figure 4 are caused by 388 considering grid cells with averaging kernels>0.5 and number of samples>2. The constrained 389 model correlates reasonably well with the changes observed by the surface measurements in April, 390 but it fails to fully reflect those in March and May. The surface measurements in March reinforce 391 increases (or negligible changes) in NO_2 in northeastern Germany and UK, although the 392 magnitudes are not as large as those suggested by the model (and TROPOMI NO₂ columns). A 393 number of factors can contribute to these large discrepancies: i) the surface measurements were 394 present throughout the month of March, whereas TROPOMI data were frequently absent due to 395 cloudiness resulting in some degree of temporal representativity issues; ii) the statistics used for 396 the TROPOMI bias-correction may not always hold true, since each individual pixel can deviate 397 from the norm of the reported biases; iii) the shape of NO₂ profiles simulated by the WRF-CMAQ 398 can sometimes be uncertain due to errors in the PBL parameterization or the difficulties with 399 resolving the non-hydrostatic components (where vertical motions are comparable to horizontal 400 ones) [e.g., Pouvaei et al., 2021]; this complication can result in unrealistic changes in the columns. 401 The constrained model tends to consistently underrepresent the decline in NO₂ in March (model: 402 $-11\pm21\%$, observation: $-19\pm16\%$), April (model: $-21\pm17\%$, observation: $-29\pm21\%$), and May 403 (model: $-12\pm18\%$, observation: $-25\pm20\%$). The frequency of TROPOMI data heavily impacted by 404 cloudiness is an important factor effectively leading to the underrepresentation of the model in a 405 course of a month. Figure 5 depicts the average number of days that TROPOMI was able to sample 406 on in both years (individual years are shown in Figure S18 and S19). There is a strong degree of 407 correlation between the frequency of the data and the discrepancy between the model versus the 408 surface observations. This is especially the case for May when we see too few days to be able to 409 realistically reproduce NO₂ changes.

Given the reasonable performance of our model at reproducing the changes observed over the surface in April, a result of abundant samples from TROPOMI, we only focus on this month for the subsequent analysis.

413

3.3.2. Ozone

414 Figure 6 depicts the changes in maximum daily 8 h average (MDA8) surface ozone 415 concentrations suggested by the measurements and the constrained model in April 2020 with 416 respect to 2019. Immediately obvious from the observations is the elevated surface ozone 417 concentrations up to 32% in places where NO_x emissions drastically decreased such as Germany, 418 Italy, France, UK, Switzerland, and Belgium (shown as box L). This tendency potentially driven 419 by ozone chemistry [Sicard et al., 2020a; Shi and Brasseur, 2020; Grange et al. 2020; Salma et al., 420 2020; Lee et al., 2020] and/or meteorology [Lee et al., 2020; Wyche et al., 2021; Ordóñez et al., 421 2020] has drawn much attention. The challenge is to set up a model that is the characteristic of 422 such a complex tendency [e.g., Parrish et al., 2014]. Encouragingly, our constrained model does 423 have skill in describing the ozone enhancements over the whole domain (r=0.43). In the proximity 424 of central Europe (shown as box L), the enhanced MDA8 ozone concentration observed by the 425 observations is 7.35±11.27% (+3.76 ppbv) which is nearly a factor of two larger than that of the model (3.73±3.94%, +1.79 ppbv). 426

427 We plot the simulated MDA8 surface ozone concentrations in April 2020 (lockdown), 428 April 2019 (baseline), and their differences in Figure 7. Surface ozone concentrations show a 429 strong latitudinal gradient with lower values in higher latitudes, underscoring the importance role 430 of solar radiation in the formation of ozone. Meanwhile, the Mediterranean basin is prone to 431 elevated concentrations of ozone resulting from different factors including calm weather, the 432 transport from neighboring countries, atmospheric recirculation in coastal environments, and local 433 emissions [Lelieveld et al., 2002]. While we observe a strong variability in the difference map, 434 signaling various sources and sinks (discussed later), three distinctive features in 2020 in 435 comparison to 2019 are evident: i) higher concentrations over the central Europe (up to 5 ppbv), 436 ii) lower concentrations in eastern Europe (-2.67 ± 1.65 ppbv) due to the 2019 biomass burning 437 activities (see S1 and S2) and larger snow cover fraction accelerating photolysis [e.g., 438 Rappenglück et al., 2014], and iii) lower values in the Iberian Peninsula (-0.51±1.41 ppbv) 439 [Ordóñez et al., 2020].

440 While the remaining model uncertainty could be either improved or characterized by 441 including more observations (if available), reconfiguring the physiochemical mechanisms used, 442 and constraining chemical boundary conditions, it is imperative to gauge the contribution of each 443 process (i.e., transport, chemistry, etc.) in forming ozone changes. Here we mainly make use of 444 the CMAQ process analysis. A direct use of the process analysis output (in unit of ppbv hr⁻¹) can 445 be confusing as both physiochemical processes and underlying concentrations are inextricably 446 linked together. To be able to isolate each process (in unit of hr⁻¹), we normalize the outputs by 447 ozone concentrations. We average each process at the same hours used in calculating MDA8. 448 Figure 8 shows the major model processes, namely horizontal transport (horizontal advection plus 449 diffusion), vertical transport (vertical advection plus diffusion), dry deposition, and chemistry in 450 2020, 2019, and their differences. Positive (negative) values indicate a source (sink) for ozone. 451 Regarding the horizontal transport, the values mostly follow the transport pattern and are 452 dependent on whether the advected air mass is more or less polluted. The vertical transport 453 correlates with the PBLH which is an indicator of the atmospheric stability and turbulence, 454 although we should not rule out the impact of the subgrid convective transport that can occur 455 sporadically. Low PBLHs are usually associated with more stable (or sometimes capping 456 inversion) and weaker vertical mixing [e.g., Nevius and Evans, 2018]. Vertical transport which is 457 majorly dictated by the vertical diffusion is by far the most influential factor in the magnitude of 458 ozone [e.g., Cuchiara et al., 2014]. In contrast to NO₂ and HCHO, a stronger vertical diffusion 459 increases surface ozone due to positive gradients of ozone with respect to altitude. However, the 460 aerodynamic resistance controlling dry deposition velocity [Seinfield and Pandis, 2006] is also a 461 function of turbulent transport. For example, during daytime, intensified turbulence exposes more 462 pollution to surface deposition. It is because of this reason that we see the dry deposition process 463 largely counteracting vertical transport. This will leave the chemistry process the major driver of 464 the ozone changes.

465 We separately sum the quantities of the physical processes and PO₃ contributing to MDA8 466 surface ozone changes binned to box L. The physical processes lead to -4.83 ppbv changes in the 467 MDA8 ozone mainly due to a relatively larger dry deposition in 2020, whereas $P(O_3)$ contributes 468 to +5.89 ppby. The net effect is +1.06 ppby which is slightly smaller than the simulated changes 469 in MDA8 ozone in this region (+1.79 ppbv). This apparent discrepancy is caused by the differences 470 in boundary and initial conditions which are not quantifiable by the process analysis and would 471 require additional sensitivity test. Nonetheless, we believe these numbers should provide 472 convincing evidence on the fact that chemistry has promoted the enhancements of surface ozone 473 during the lockdown.

474 Chemistry is also a function of meteorology, specifically solar radiation and temperature. 475 A typical scenario to isolate emissions from meteorology is by running the model with fixed 476 anthropogenic emissions (and boundary conditions) and subtracting the outputs from the variable 477 emission output. Figure 9 shows the contribution of anthropogenic emissions (VOCs and NO_x) to 478 the changes seen over the surface. The anthropogenic emissions make up roughly 58% of the 479 changes. The map is strongly in line with the changes in NO_x emissions constrained by TROPOMI. 480 The impact of meteorology plus biogenic changes (the former is dominant) highly correlates with 481 anomalies in both surface air temperature and photolysis rates dictated by synoptic conditions 482 (Figure S17). We observe negligible ozone changes due to emissions over Iberian Peninsula 483 reinforcing the significance of the meteorological impacts [Ordóñez et al., 2020].

484

3.4.Ozone chemistry

Figure 10 shows the numerically-solved ozone production rates (PO₃) simulated by the constrained model during the MDA8 hours period. We observe positive PO₃ in less polluted areas and eastern Europe where biomass burning activities occurred in 2019 (see S1 and S2), while negative PO₃ in major cities. Negative values in PO₃ are indicative of either loss in O₃ or O₃-NO- 489 NO₂ partitioning. The difference in PO₃ between the two years suggests that the ozone 490 enhancement in box L is caused by a reduction in negative PO₃ in 2020 over major cities compared 491 to 2019. To examine which pathways are contributing to this pattern, we attempt to analytically 492 reproduce the numerically-solved PO₃ (Figure 10) through two different equations: the first 493 equation widely applied in photochemically active environments follows [Kleinman et al., 2002]:

$$P(O_{3}) = k_{HO_{2}+NO}[HO_{2}][NO] + \sum k_{RO_{2i}+NO}[RO_{2i}][NO] - k_{OH+NO_{2}+M}[OH][NO_{2}][M] - k_{HO_{2}+O_{3}}[HO_{2}][O_{3}] - k_{OH+O_{3}}[OH][O_{3}] - k_{O(^{1}D)+H_{2}O}[O(^{1}D)][H_{2}O] - L(O_{3} + VOCs)$$
(6)

This equation yields negative values only if the O₃ loss pathways including NO₂+OH, HO_x+O₃, O¹D+H₂O and O₃+VOCs dominate over the first two terms. The second equation which is independent of RO₂ and HO₂ concentrations [Thornton et al., 2002], is:

$$P(O_3) = jNO_2[NO_2] - k_{NO+O_3}[O_3][NO]$$
(7)

- In summer, this equation tends to be positive during early afternoon, almost zero during afternoon (steady-state), and negative in early morning (or night) in which the second term (O₃ titration) is leading. Any abrupt changes in NO_x and VOC, and photolysis can directly affect Eq.7 moving PO₃ out of the diel steady-state. The assumption of the steady-state (PO₃ from Eq.7 equals to zero) is also not valid if an air parcel is in the vicinity of high-emitting NO_x sources [Thornton et al., 2002].
- 502 Figure 11 displays the reactions rates of each individual component involved in Eq.6 503 averaged during the MDA8 hours. HO₂+NO is the dominant chemical source of ozone correlating 504 well with the changes in NO_x and prevailing chemical conditions regimes (NO_x-sensitive vs VOC-505 sensitive). Souri et al. [2020a] found the reaction of RO₂+NO to be primarily dependent on VOCs. 506 Likewise, we observe a strong degree of correlation between the anomaly of RO₂+NO and that of 507 VOCs (Figure S1 and S2). Figure 11 indicates that the chemical pathways of ozone loss are rather 508 constant between the two years; therefore the largely negative PO₃ over urban areas shown 509 previously in Figure 10 is not reproducible using this equation. Figure 12 shows the reactions rates 510 of $J_{NO2}[NO_2]$, $k_{NO+O3}[NO][O_3]$, and the difference during the MDA8 hours. The difference maps 511 replicate the largely negative PO₃ over cities suggesting that we are not in the diel steady-state, 512 and O₃ titration is prevailing due to relatively low photochemistry in the springtime. Table 2 lists 513 the averaged reactions rates involved in Eq.6 and 7 along with the numerically-solved PO₃ shown

in Figure 10 over box L. These numbers suggest that the major chemical pathways of enhanced ozone are through $J_{NO2}[NO_2]$ and $k_{NO+O3}[NO][O_3]$, implying that O_3 -NO-NO₂ partitioning is more consequential than other chemical pathways. This analysis strongly coincides with Lee et al.

- 517 [2020] and Wyche et al. [2021] who observed roughly constant O_3 +NO₂ concentrations over the
- 518 UK before and during the lockdown 2020.

4. Summary

520 The slowdown in human activities due to the COVID-19 pandemic had a large impact on 521 air pollution over Europe [Barré et al. 2020; Siccard et al., 2020; Sun et al., 2021]. Satellite 522 monitoring systems with large spatial coverage help shed light on the spatial and temporal extent 523 of those impacts. The relationships between satellite-derived columns and near-surface emissions 524 have proven difficult to fully establish without using realistic models, capable of providing insights 525 on the convoluted processes involving chemistry, dynamics, transport, and photochemistry and 526 therefore help with deciphering what anomaly maps of satellite concentrations are suggesting [e.g., Goldberg et al., 2020]. To address these challenges, we jointly constrained NO_x and VOC 527 528 emissions using TROPOMI HCHO and NO₂ columns following a non-linear Gauss Newton 529 method developed in Souri et al. [2020a], in addition to assimilating MODIS AOD observations 530 based on Jung et al. [2019]. The constrained emissions also permitted investigating the 531 simultaneous effects of physiochemical processes contributing to ozone formation, illuminating 532 the complexities associated with non-linear chemistry.

533 Several implications of the derived emissions for the months of March, April, and May 534 2020 (lockdown) relative to those in 2019 (baseline) were investigated. First, as previously 535 reported [Sicard et al., 2020; Barré et al. 2020], we observed a significant reduction in NO_x in 536 March (14-31%) in several major polluted regions including Paris, London, Madrid, and Milan. 537 The reductions were further seen in other cities such as Rome, Brussels, Frankfurt, Warsaw, 538 Belgrade, Kyiv, and Moscow (34-51%) in April. Second, NO_x emissions decreased drastically in 539 April rather than March in UK, Moscow, and Poland due to the timeline of restrictions. Third, the 540 changes in NO_x suggested by TROPOMI NO₂ and the constrained model over northeastern 541 Germany in March and Eastern Europe in May were unrealistic, possibly due to observations or 542 the model issues. Fourth, we observed a weak observational constraint on VOC emissions from 543 TROPOMI HCHO except for lower latitudes.

The constrained model calculations gave good representations of near-surface NO₂ changes in April (model: $-21\pm17\%$, observation: $-29\pm21\%$) in places where the top-down estimates are strongly constrained by TROPOMI (averaging kernels > 0.5), but inferior representations in other months, especially in May (model: $-12\pm18\%$, observation: $-25\pm20\%$). This tendency mainly arose from TROPOMI observation frequencies; too few days (10-26% out of a month) in May due to cloudiness precluded the determination of realistic NO_x emission changes.

550 We observed surface MDA8 ozone increase from both model and measurements in April 551 2020 with respect to the baseline. Comparisons of calculation by the constrained model in terms 552 of MDA8 surface ozone found a reasonable agreement with observations in the proximity of 553 central Europe in April (model: +3.73±3.94%, +1.79 ppbv, observation: +7.35±11.27%, +3.76 554 ppby). These comparisons indicate that the performance of the constrained model to reproduce the 555 ozone enhancement feature is promising, suggesting fruitful information in TROPOMI, although 556 reasons behind the underestimation of the enhancement remained unexplained. It was clear that 557 the dominantly negative ozone production rates dictated by O₃-NO-NO₂ partitioning (J_{NO2}[NO₂]-558 $k_{NO+O3}[NO][O_3]$ became less negative primarily due to the reduced NO_x emissions in urban areas 559 where O₃ titration occurred. This tendency was in agreement with studies of Lee et al. [2020] and 560 Wyche et al. [2021]. We found negligible differences in ozone production from [HO₂+RO₂][NO] 561 and ozone loss from O^1D+H_2O and O_3+HO_x between the two years suggesting photochemistry 562 was rather low in the springtime over Europe.

563 We further quantified the contributions of physical processes (transport, diffusion and dry 564 deposition) and chemistry to the formation/loss of ozone using the integrated process rates. The 565 physical processes decreased MDA8 ozone by -4.83 ppbv resulting from relatively larger dry deposition in 2020, whereas chemistry (ozone production) augmented ozone levels by +5.89 ppby, 566 567 indicating that rising ozone was primarily impacted by changes in chemistry. Enhanced air 568 temperature and photolysis in 2020, both of which were well captured in our model, also affected 569 chemistry. Experiments with fixed anthropogenic emissions underwent significant enhancement 570 in surface MDA8 ozone over central Europe, but those only contribute to 42% of the total 571 enhancement indicating that anthropogenic emissions were the major factor.

572 The results shown here reveal previously unquantified characteristics of ozone and its 573 precursors emission changes during the lockdown 2020 in Europe. We have been able to measure 574 the amount of changes along with the level of confidence in NO_x (and partly VOC emissions)

- 575 using a state-of-the-art inversion technique by leveraging satellite observations, which in turn,
- allowed us to unravel the physiochemical processes contributing to increased ozone in Europe.
- 577 Unless a comprehensive air quality campaign targeting COVID-19 related lockdown is available,
- 578 we recommend that the impact of lockdown on air pollution should be examined through the lens
- 579 of well-established models constrained by publicly available data, especially those from space in
- 580 less cloudy environments.

581 Author contributions

- 582 AHS designed the research, analyzed the data, conducted the inverse modeling and atmospheric
- 583 modeling (for CMAQ, GEOS-Chem, WRF, and MEGAN), made all figures, and wrote the paper.
- 584 JB validated WRF-CMAQ model and reformatted the surface observation files. CRN and GGA
- 585 did literature review regarding the TROPOMI validation. YJ validated MODIS AOD. DW helped
- 586 with implementing the AOD assimilation framework. KC, JM, and XL guided the discussion. All
- 587 authors contributed to discussion and edited the paper.

588 Data availability

- 589 The atmospheric inversion data are publicly available from Souri et al. [2021]. The model outputs
- 590 are available upon the request from absouri@cfa.harvard.edu. The links on where to download
- 591 surface and satellite observations that are used in this study are already provided in the text.

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- 601

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- 937 938 **Table 1.** Statistics reported in several validations studies comparing TROPOMItropospheric NO2 against independent observations.

Study	Location	Time	Benchmark	Bias (low)	Dispersio	Modificatio	Modified
		Period	Instrument		n	n	Bias (low)
Chan et al. 2020	Munich	May 2018-Apr 2019	MAX-DOAS	30%	N/A	In-situ MAX- DOAS profiles	17%
Griffin et al. 2019	Canadian Oil Sands	Mar-May 2018 (v1.01)	Pandora (direct Sun)	15-30%	N/A	Higher resolution profiles (10 km) and albedo	0-25%
Judd et al. 2020	New York	Jun-Sep 2018	GeoTASO	19-33%	N/A	Higher resolution profiles (12 km)	7-19%
Verhoelst et al. 2020	Global	Apr 2018- Feb 2020	MAX-DOAS	37% (average), 23-51% (range)	3.5×10 ¹⁵ molec/cm ²	N/A	N/A
Wang P. et al. 2020	Atlantic and Pacific Oceans	4 campaigns during Dec 2018- Jul 2019	MAX-DOAS	Negligible	N/A	N/A	N/A
Zhao et al. 2020	Greater Toronto Area	Mar 2018- Mar 2019	Pandora (direct Sun)	24-28% (suburban/ur ban) +4-10% (rural)		Higher resolution profiles (10 km) and albedo	13-24% (suburban/ urban) +14-15% (rural)

Reactions	Production (P) or loss (L)	April 2020 (ppbv/hr)	April 2019 (ppbv/hr)	Net diff ^a (ppbv/hr)
HO ₂ +NO	Р	0.85	0.91	-0.06
RO ₂ +NO	Р	0.44	0.41	+0.03
NO ₂ +OH	L	0.10	0.14	+0.04
O^1D+H_2O	L	0.07	0.08	+0.01
O ₃ +VOCs	L	0.01	0.01	0.00
O ₃ +HO _x	L	0.09	0.08	-0.01
J _{NO2} [NO ₂]	Р	14.61	27.28	-12.67
k _{NO+O3} [NO][O ₃]	L	15.11	28.52	+13.40
J _{NO2} [NO ₂]- k _{NO+O3} [NO][O ₃]	N/A	-0.50	-1.24	+0.74
Numerically solved PO ₃	N/A	-0.79	-1.53	+0.74

949 Table 2. Reaction rates relating to the chemical pathways of ozone formation and loss over box L950 (proximity of central Europe).

951 ^a A positive net difference indicates higher (lower) production (loss) in 2020 with respect to 2019.
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CMAQ 15-km Domain

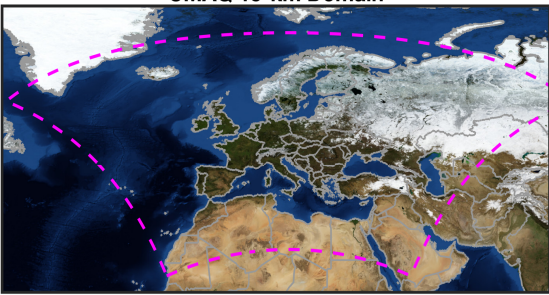


Figure 1. The WRF-CMAQ 15 km domain covering Europe. The background picture is based on the publicly available NASA Blue Marble (© NASA). 957



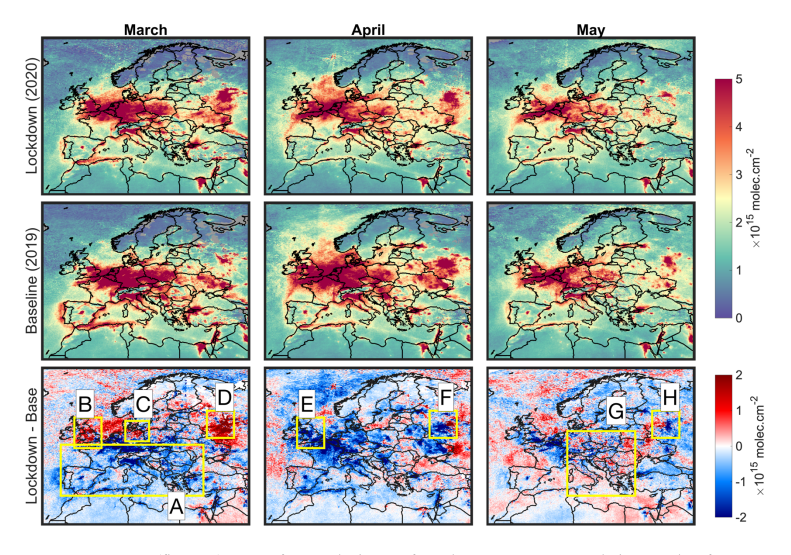


Figure 2. (first row) Maps of tropospheric NO₂ from the TROPOMI sensor during months of March, April, and May in 2020 (lockdown). (second row) Same as the first row but for the baseline year (2019). (last row), Difference of the columns in 2020 with respect to those of 2019. All columns are corrected for the bias and their AMFs are recalculated iteratively based on the posterior profiles derived from our inverse modeling practice. The satellite-derived columns are subject to errors, so a direct interpretation of their magnitudes cannot be performed in a robust manner.

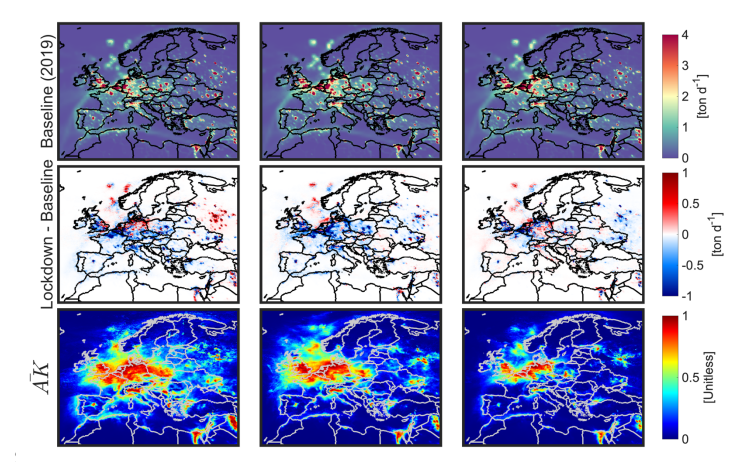
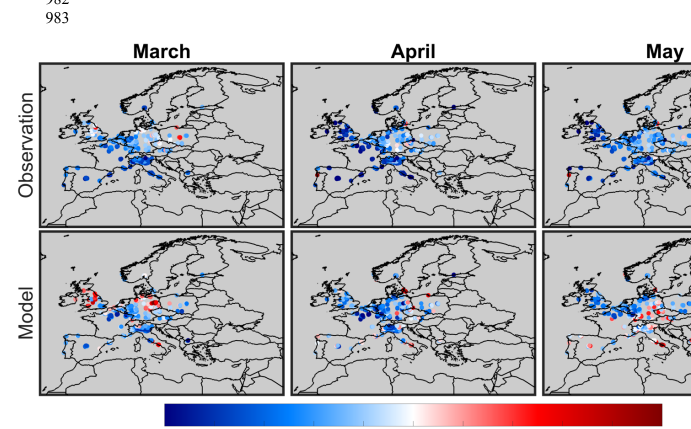


Figure 3. Top-down estimates of total NO_x during months of March, April and May in 2019 (baseline) and the differences between emission in 2020 (lockdown) and 2019. To infer the magnitude of emissions in 2020, the second row should be added to the first one. Both TROPOMI HCHO and NO₂ observations are jointly used to estimate these numbers. Averaging kernels (mean values based on both 2019 and 2020 estimates) describe the level of credibility of the estimate which is heavily dependent on the TROPOMI signal-to-noise ratios.

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985 Figure 4. Scatter maps of relative changes in surface NO₂ concentrations suggested by the 986 European air quality network (first row), and the constrained model (second row). Results are 987 daily-averaged. We only consider grid cells whose averaging kernels (from the NO_x inversion) 988 are above 0.5. Furthermore, grid cells having more than 2 stations are only included to partly 989 account for the spatial representivity factor. Surface concentrations are not accounted for the 990 NO_z family interferences.

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0

[%]

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991

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-30

-20

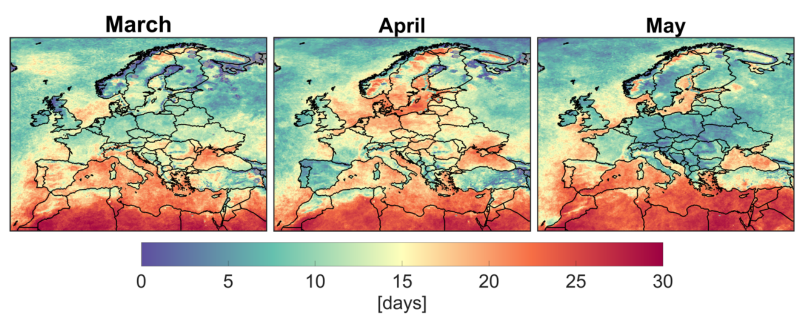
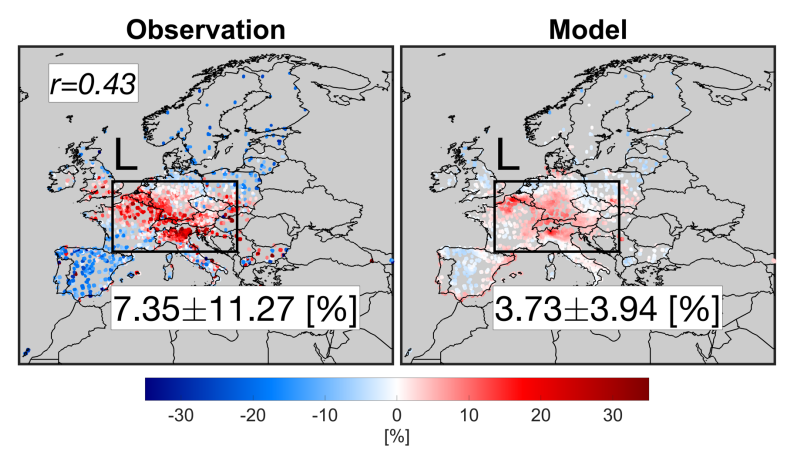


Figure 5. The average number of good quality (qa_flag>0.75) TROPOMI tropospheric NO₂ days

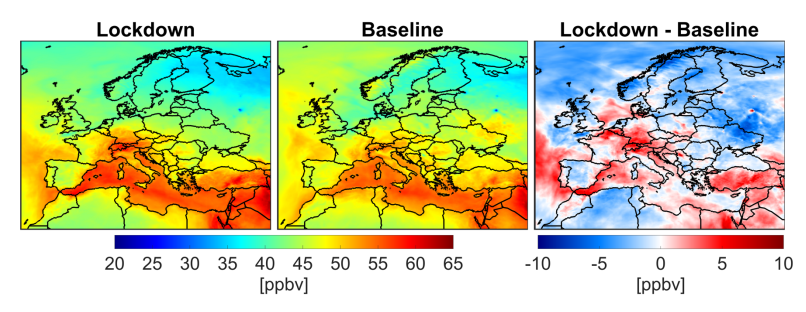
observed at 15×15 km² in 2019 and 2020. These numbers are heavily affected by cloudiness.



1002 Figure 6. Changes in surface MDA8 ozone concentrations suggested by the observation (left),

1003 and the constrained model (right) in April 2020 relative to those in 2019. The numbers are based

- 1004 on the box L region.



1007 Figure 7. Simulated surface MDA8 ozone concentration using the constrained model in the month
1008 of April 2020 (lockdown), April 2019 (baseline), and their difference.
1009



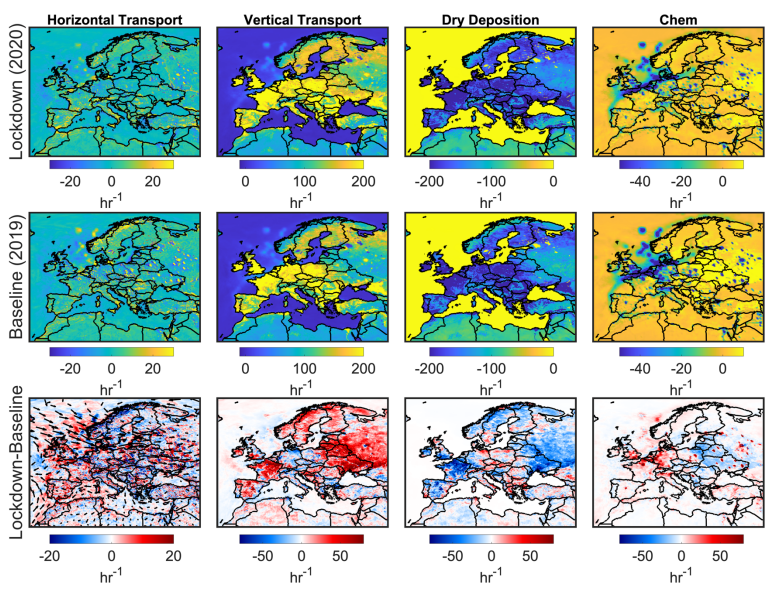
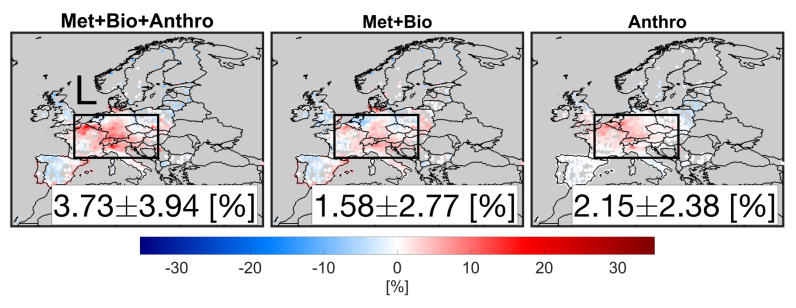


Figure 8. Surface process tendencies (hr⁻¹) including horizontal transport (advection plus

- 1014 diffusion), vertical transport (advection plus diffusion), dry deposition, and chemistry. Positive
- 1015 (negative) values mean source (sink) of ozone. These outputs are based on the constrained
- 1016 model. Wind vectors are the difference.

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1021



1023 Figure 9. Simulated MDA8 surface ozone difference between April 2020 with respect to April

1024 2019 including (left) dynamical meteorology, biogenic and anthropogenic emissions, (middle)

1025 dynamical meteorology and biogenic emissions, and (right) the subtraction of the previous

1026 scenarios isolating dynamical anthropogenic emissions. Emissions used for these experiments

- 1027 are based on the top-down estimates.
- 1028



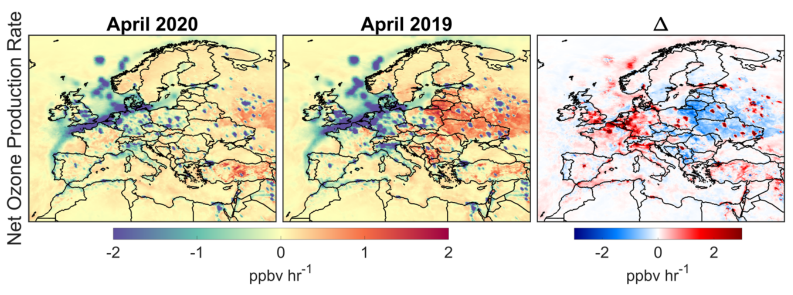
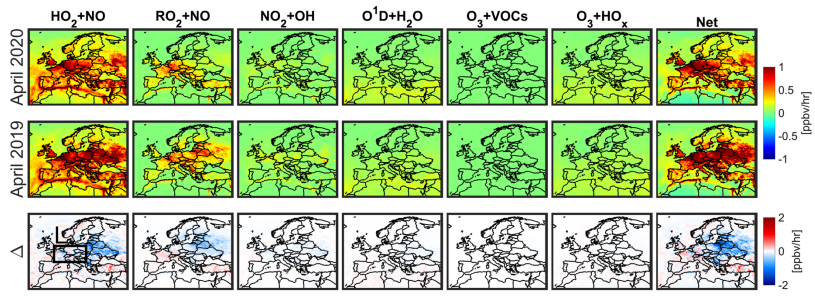
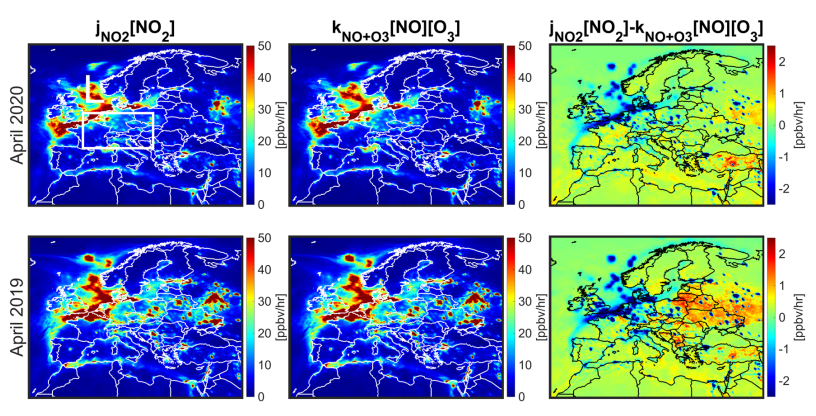


Figure 10. Numerically-solved net ozone production rates based on the WRF-CMAQ simulations
using the constrained emissions by the satellite data in April 2020, 2019, and the difference. These
values are over the surface and are averaged during the MDA8 hours.



1038 **Figure 11.** Surface chemical processes involved in equation 5 (ppbv hr⁻¹) pertaining to the

- 1039 production and loss of ozone in April 2020 (lockdown) and 2019 (baseline) during MDA8 hours.
- 1040 These outputs are based on the constrained model.
- 1041



1043 Figure 12. Surface chemical processes involved in equation 6 (ppbv hr⁻¹) pertaining to the O₃-

1044 NO-NO₂ partitioning in April 2020 and 2019 during MDA8 hours. The constrained model by the

- 1045 satellite observations is used to derive these outputs.
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- 1048