Tropospheric ozone production and chemical regime analysis during the COVID-19 lockdown over Europe

Clara M. Nussbaumer\(^1\), Andrea Pozzer\(^1\), Ivan Tadic\(^1\), Lenard Röder\(^1\), Florian Obersteiner\(^2\), Hartwig Harder\(^1\), Jos Lelieveld\(^1,3\), and Horst Fischer\(^1\)

\(^1\)Max Planck Institute for Chemistry, Department of Atmospheric Chemistry, 55128 Mainz, Germany
\(^2\)Karlsruhe Institute of Technology, 76021 Karlsruhe, Germany
\(^3\)Climate and Atmosphere Research Center, The Cyprus Institute, Nicosia, Cyprus

Correspondence: Clara Nussbaumer (clara.nussbaumer@mpic.de)

Abstract.

The COVID-19 (Coronavirus disease 2019) European lockdowns have lead to a significant reduction in the emissions of primary pollutants such as NO (nitric oxide) and NO\(_2\) (nitrogen dioxide). As most photochemical processes are related to nitrogen oxide (NO\(_x\) \(\equiv\) NO + NO\(_2\)) chemistry, this event has presented an exceptional opportunity to investigate its effects on air quality and secondary pollutants, such as tropospheric ozone (O\(_3\)). In this study, we present the effects of the COVID-19 lockdown on atmospheric trace gas concentrations, net ozone production rates (NOPR) and the dominant chemical regime throughout the troposphere based on three different research aircraft campaigns across Europe. These are the UTOPIHAN campaigns in 2003 and 2004, the HOOVER campaigns in 2006 and 2007 and the BLUESKY campaign in 2020, the latter performed during the COVID-19 lockdown. We present in situ observations and simulation results from the ECHAM5/MESSy Atmospheric Chemistry model which allows for scenario calculations with business as usual emissions during the BLUESKY campaign, referred to as "no-lockdown scenario". We show that the COVID-19 lockdown reduced NO and NO\(_2\) mixing ratios in the upper troposphere by around 55\% compared to the no-lockdown scenario due to reduced air traffic. O\(_3\) production and loss terms reflected this reduction with a deceleration in O\(_3\) cycling due to reduced mixing ratios of NO\(_x\) while NOPRs were largely unaffected. We also study the role of methyl peroxyradicals forming HCHO (\(\alpha_{\text{CH}_3\text{O}_2}\)) to show that the COVID-19 lockdown shifted the chemistry in the upper troposphere/tropopause region to a NO\(_x\) limited regime during BLUESKY. In comparison, we find a VOC limited regime to be dominant during UTOPIHAN.

1 Introduction

COVID-19 (Coronavirus disease 2019) describes the disease accompanying an infection with the SARS-CoV-2 (severe acute respiratory syndrome coronavirus-2) virus. The disease is highly infectious and can have severe health consequences, including premature death, particularly for elderly and people with pre-existing conditions (WHO, 2021). On 11 March 2020, COVID-19 was declared a pandemic by the World Health Organization (WHO, 2020a,b). As a response, in many countries worldwide - including the European continent - governments initiated a shutdown of the daily life for minimizing the spread of the virus, which is referred to as COVID-19 lockdown. Among others, this included a reduction in vehicular and industrial activities as
well as sharp restrictions on air travel accompanied by a reduction in atmospheric pollutants such as nitrogen oxides (NO\textsubscript{x} = NO + NO\textsubscript{2}) (Venter et al., 2020; Kroll et al., 2020; Chossière et al., 2021; Onyeaka et al., 2021; Salma et al., 2020; Matthias et al., 2021; Forster et al., 2020).

NO and NO\textsubscript{2} are important atmospheric trace gases as they are involved in almost all photochemical processes taking place in the earth’s atmosphere. NO\textsubscript{x} directly impacts the production of tropospheric ozone (O\textsubscript{3}) which is a hazard to human, animal and plant health (Nuvolone et al., 2018). Together with volatile organic compound (VOC) oxidation, NO forms NO\textsubscript{2} within the HO\textsubscript{x} cycle, catalyzed by an OH radical. Under the influence of sunlight, NO\textsubscript{2} can subsequently form O\textsubscript{3} through the reaction with molecular oxygen as shown in Reaction (R1) (Crutzen, 1988; Lelieveld and Dentener, 2000; Pusede and Cohen, 2012; Pusede et al., 2015; Nussbaumer and Cohen, 2020).

\[
NO_2 + O_2 \xrightarrow{h\nu} NO + O_3 \quad \text{(R1)}
\]

Various termination reactions such as the formation of HNO\textsubscript{3} from OH and NO\textsubscript{2} or other radical recombinations cause ozone chemistry to be non-linear, which means that a reduction in ambient NO\textsubscript{x} can either increase or decrease O\textsubscript{3} production (Pusede et al., 2015). For low ambient NO\textsubscript{x} levels, a NO\textsubscript{x} reduction usually causes a decrease in O\textsubscript{3} production which is referred to as a NO\textsubscript{x} limited chemical regime. In contrast, a NO\textsubscript{x} reduction increases O\textsubscript{3} production when a VOC limited chemical regime is dominant - usually at high ambient NO\textsubscript{x} levels (Sillman et al., 1990; National Research Council, 1991; Pusede and Cohen, 2012). In the transition region between the two regimes, changes in NO\textsubscript{x} do not (or only slightly) impact O\textsubscript{3} production rates (Wang et al., 2018).

Many different methods enable the determination of the dominant chemical regime, such as the use of the weekend ozone effect which considers the response of O\textsubscript{3} to NO\textsubscript{x} reductions on weekends or the ratio of HCHO to NO\textsubscript{2} with various approaches from in situ observations, remote sensing and model simulations (e.g. Jin et al. (2020); Pusede and Cohen (2012); Nussbaumer and Cohen (2020); Duncan et al. (2010)). We have recently shown that the share $\alpha$ of methyl peroxyradicals (CH\textsubscript{3}O\textsubscript{2}) forming formaldehyde (HCHO) in correlation with ambient NO concentrations is capable of indicating the dominant chemical regime based on three different field campaigns across Europe in Finland (HUMPPA 2012), Germany (HOPE 2012) and Cyprus (CYPHEX 2014) (Nussbaumer et al., 2021). CH\textsubscript{3}O\textsubscript{2} formed from e.g. the oxidation of methanol (CH\textsubscript{3}OH) or methane (CH\textsubscript{4}) can either react with NO or OH radicals to form HCHO or undergo the competing reaction with HO\textsubscript{2} to form methyl hydroperoxide (CH\textsubscript{3}OOH). For more details, please see Figure 1 in Nussbaumer et al. (2021). $\alpha_{CH_3O_2}$ consequently depends on the ambient concentrations of NO, OH and HO\textsubscript{2} and the respective rate constants for the reaction with CH\textsubscript{3}O\textsubscript{2}, the latter of which were taken from the IUPAC Task Group on Atmospheric Chemical Kinetic Data Evaluation (2021). The calculation of $\alpha_{CH_3O_2}$ is presented in Equation (1).

\[
\alpha_{CH_3O_2} = \frac{k_{CH_3O_2+NO} \times [NO] + k_{CH_3O_2+OH} \times [OH]}{k_{CH_3O_2+NO} \times [NO] + k_{CH_3O_2+OH} \times [OH] + k_{CH_3O_2+HO_2} \times [HO_2]} \quad (1)
\]
Low values for $\alpha_{CH_3O_2}$ with a high response to NO are an indicator for a NO$_x$ limited regime whereas high values for $\alpha_{CH_3O_2}$ with little response to changing NO represent a VOC limitation (Figure 11 in Nussbaumer et al. (2021)). Investigating the dominant chemical regime is an important method for analyzing photochemical processes and air quality.

Previous studies have explored changes in air quality, trace gas emissions and the dominant chemical regime during the COVID-19 lockdown in Europe. Menut et al. (2020) reported NO$_2$ reductions between 30 and 50% for various Western European countries in the course of March 2020 with both decreasing and increasing O$_3$ concentrations in response, depending on the location, based on surface in situ observations and model simulations. Ordóñez et al. (2020) observed decreased NO$_2$ and increased O$_3$ concentrations in Central Europe in March and April 2020 based on in situ observations compared to 2015 - 2019. While they found NO$_2$ reductions to be mainly attributed to the COVID-19 lockdown, O$_3$ enhancements were predominantly affected by meteorological changes. Chossière et al. (2021) presented evidence on NO$_2$ reductions during the COVID-19 lockdown in Europe and O$_3$ changes dependent on the dominant chemical regime through investigation of HCHO/NO$_2$ ratios based on in situ and satellite observations. Similar studies were performed by Matthias et al. (2021); Mertens et al. (2021); Balamurugan et al. (2021); Grange et al. (2021) and many more. Besides the changes within the dominant chemical regime through NO$_x$ reductions, i.e. increasing ozone within a VOC limited regime and decreasing ozone within a NO$_x$ limited regime, the COVID-19 lockdown could have potentially changed the dominant chemical regime from VOC to NO$_x$ limited as pointed out by Kroll et al. (2020) and Gaubert et al. (2021). Cazorla et al. (2021) found a lockdown induced change from a VOC to a NO$_x$ limited regime in Quito (Ecuador) based on the share of precursor loss to HNO$_3$ and H$_2$O$_2$. The latter is dominant for NO$_x$ limited chemistry (Kleinman et al., 2001). A change from a VOC to a NO$_x$ limited regime was also reported by Zhu et al. (2021) in China based on HCHO to NO$_2$ ratios (NO$_x$ limitation for ratios above 2 according to Duncan et al. (2010)). Most of the literature on emission reductions during the COVID-19 lockdown focuses on near-surface air quality and only few studies consider the free troposphere. Steinbrecht et al. (2021) and Bouarar et al. (2021) reported decreases in O$_3$ concentrations in the free troposphere based on in situ observations and modeling studies in the northern hemisphere. A similar observation with reduced O$_3$ in the free troposphere around Frankfurt airport was shown by Clark et al. (2021).

In this study, we present atmospheric trace gas concentrations, net ozone production rates and an analysis on the dominant chemical regime based on in situ observations during the research aircraft campaign BLUESKY which took place in May and June 2020 over Europe, and model simulations. During this time period, aircraft activity was still strongly limited due to the COVID-19 lockdown. We compare the results to model simulations assuming business as usual emissions not impacted by government restrictions which we refer to as "no-lockdown scenario". Additionally, we present results on two previous aircraft campaigns which are UTOPIHAN (Upper Tropospheric Ozone: Processes Involving HO$_x$ and NO$_x$) in 2003/2004 and HOOVER (HO$_x$ over Europe) in 2006/2007. While many studies have been published on emissions reductions and the effect on secondary pollutants during the COVID-19 lockdown, only a few studies have investigated changes in the dominant chemical regime and to our knowledge we are the first to report a shift to NO$_x$ limited chemistry in the upper troposphere. This can demonstrate the consequences of emission changes of VOCs (including methane) and NO$_x$ on tropospheric ozone.
2 Observations and methods

2.1 Calculations of net ozone production rates (NOPR)

Besides the chemical regime, production and loss processes of \( O_3 \) are effective tools in exploring relevant photochemistry. As already demonstrated in Reaction (R1), \( O_3 \) is formed via \( NO_2 \) photolysis. Under the assumption of photostationary state, this term can be equated with the reactions of \( NO \) with \( O_3 \), \( HO_2 \) and \( RO_2 \) (Hosaynali Beygi et al., 2011). The resulting term for \( O_3 \) production \( P(O_3) \) is shown in Equation (2) (Tadic et al., 2020; Leighton, 1961). \( j(NO_2) \) is the photolysis frequency of \( NO_2 \) and \( k \) describes the respective rate constant (for this work taken from the IUPAC Task Group on Atmospheric Chemical Kinetic Data Evaluation (2021)).

\[
P(O_3) = [NO_2] \times j(NO_2) = [NO] \times (k_{O_3+NO} \times [O_3] + k_{NO+HO_2} \times [HO_2] + \sum k_{NO+R_zO_2} \times [R_zO_2]) \tag{2}
\]

We assume \( R_zO_2 \) (the sum of all peroxy radicals) to be represented by \( CH_3O_2 \) which was found to be a reasonable approximation for the comparable, though independent aircraft campaign CAFE Africa in 2018 as described in Tadic et al. (2021). \( CH_3O_2 \) can be calculated via Equation (3) as derived by Bozem et al. (2017a). While the model can simulate \( CH_3O_2 \) mixing ratios, Equation (3) is required when working with experimental data as \( CH_3O_2 \) was not directly measured.

\[
[CH_3O_2] = \frac{k_{CH_4+OH} \times [CH_4]}{k_{CO+OH} \times [CO]} \times [HO_2] \tag{3}
\]

\( O_3 \) loss occurs via the reaction with \( NO \), \( OH \) and \( HO_2 \) and via photolysis and can be calculated as presented in Equation (4). The photolysis of \( O_3 \) first yields \( O^{1}D \) which reacts back to \( O_3 \) through collision with \( O_2 \) or \( N_2 \), and causes an \( O_3 \) loss through reaction with \( H_2O \). The share of \( O_3 \) that is effectively lost through \( O_3 \) photolysis is described by \( \alpha_{O^{1}D} \) in Equation (5) (Bozem et al., 2017a). Additional loss due to reactions of \( O_3 \) with alkenes and the loss of \( NO_2 \) due to formation of \( HNO_3 \) or peroxy nitrates are negligibly small, particularly in the upper troposphere.

\[
L(O_3) = [O_3] \times (k_{O_3+NO} \times [NO] + k_{O_3+HO_2} \times [HO_2] + k_{O_3+OH} \times [OH] + \alpha_{O^{1}D} \times j(O^{1}D)) \tag{4}
\]

\[
\alpha_{O^{1}D} = \frac{k_{O^{1}D+H_2O} \times [H_2O]}{k_{O^{1}D+N_2} \times [N_2] + k_{O^{1}D+O_2} \times [O_2] + k_{O^{1}D+H_2O} \times [H_2O]} \tag{5}
\]
Net ozone production rates (NOPR) are then calculated from the difference in $P(O_3)$ and $L(O_3)$ whereas $P(O_3)$ can either be expressed via NO$_2$ or NO reaction terms. The term $k_{O_3+NO} \times [O_3] \times [NO]$ can be neglected for the latter as it is equally present in $P(O_3)$ and $L(O_3)$.

$$\text{NOPR} = P(O_3) - L(O_3)$$

$$= [NO_2] \times j(NO_2) - [O_3] \times (k_{O_3+NO} \times [NO] + k_{O_3+HO_2} \times [HO_2] + k_{O_3+OH} \times [OH] + \alpha_{O_1D} \times j(O_1D))$$

$$= [NO] \times (k_{NO+HO_2} \times [HO_2] + k_{NO+CH_3O_2} \times [CH_3O_2])$$

$$- [O_3] \times (k_{O_3+HO_2} \times [HO_2] + k_{O_3+OH} \times [OH] + \alpha_{O_1D} \times j(O_1D))$$  \hspace{1cm} (6)

### 2.2 Field experiments

We have investigated in situ trace gas observations from three different research aircraft campaigns which are the UTOPIHAN campaigns in 2003/2004, the HOOVER campaigns in 2006/2007 and the BLUESKY campaign in 2020. Figure 1 shows an overview of the flight tracks over Europe. We have filtered the data for the tropospheric region by help of the modeled tropopause pressure (see Section 2.3) and south of 60°N as there were no data points for the BLUESKY campaign further north. Dashed lines show the complete flight tracks during each campaign and solid lines show the data which we have considered in this study. The experimental data were obtained with a time resolution of 1 minute and subsequently adjusted to fit the model resolution of 6 minutes. For this, each sixth experimental data point (which fit the model time scale) and the data points from ±2 minutes were averaged. The remaining data points were discarded.

#### 2.2.1 UTOPIHAN 2003/04

The UTOPIHAN (Upper Tropospheric Ozone: Processes Involving HO$_x$ and NO$_x$) campaigns took place in June/July 2003 and March 2004 starting from Oberpfaffenhofen airport in Germany (48.08 °N, 11.28 °E) with the GFD (Gesellschaft für Flugziel darstellung, Hohn, Germany) research aircraft Learjet 35A (Colomb et al., 2006; Klippel et al., 2011; Stickler et al., 2006). NO and O$_3$ were measured via chemiluminescent detection (CLD 790 SR, ECO Physics, Dürnten, Switzerland). NO data have a precision of 6.5 %, an accuracy of ≤ 25 % and a detection limit of < 0.01 ppbv. O$_3$ data have a precision of 1 % and an accuracy of 5 %. j(NO$_2$) was determined via filter radiometers (Meterologie Consult GmbH, Königstein, Germany) with a precision of 1 % and an accuracy of 15 %. CO measurements were obtained from a tunable diode laser absorption spectrometer with a detection limit of 0.26 ppbv (30 s time resolution) and an accuracy of 3.6 % (6 s time resolution) (Kormann et al., 2005).

#### 2.2.2 HOOVER 2006/07

The HOOVER (HO$_x$ over Europe) campaigns took place in October 2006 and July 2007 using the GFD research aircraft Learjet 35A with the campaign base in Hohn, Germany (54.31 °N, 9.53 °E) (Klippel et al., 2011; Bozem et al., 2017b,a; Regelin et al., 2013). NO and O$_3$ measurements were performed via chemiluminescence (CLD 790 SR, ECO Physics, Dürnten, Switzerland) with a precision of 7 and 4 %, an accuracy of 12 and 2 % and a detection limit of 0.2 and 2 ppbv, respectively.
Figure 1. Overview of the flight tracks of the considered aircraft campaigns UTOPIHAN (2003 & 2004) in green, HOOVER (2006 & 2007) in blue and BLUESKY (2020) in red. Solid lines present the data considered in this study (filtered for troposphere and south of 60°N) and dashed lines show the complete flight tracks.

CO and CH\textsubscript{4} were measured via quantum cascade laser absorption spectroscopy with an accuracy of 1.1 and 0.6 % and detection limits of 0.2 and 6 ppbv, respectively (2 s time resolution) (Schiller et al., 2008). OH and HO\textsubscript{2} measurements were performed via laser-induced fluorescence with the HORUS (HydrOxyl Radical measurement Unit based on fluorescence Spectroscopy) instrument with an accuracy of 18 % and detection limits of 0.016 and 0.33 pptv, respectively (1 min time resolution) (Regelin et al., 2013). Photolysis frequencies were measured using filter radiometers (Meterologie Consult GmbH, Königstein, Germany) with a precision of 1 % and an accuracy of 15 % (1 s time resolution). H\textsubscript{2}O was measured via IR-absorption with a typical accuracy of 1 % (modified LI-6262, LI-COR Inc., Lincoln, USA) (Gurk et al., 2008; LI-COR, Inc., 1996).

2.2.3 BLUESKY 2020

The BLUESKY campaign took place in May and June 2020 over Europe. Eight flights were carried out using the HALO (High Altitude Long Range) research aircraft starting from the campaign base in Oberpfaffenhofen, Germany. The goal of the campaign was to examine the effects of the COVID-19 lockdown on the troposphere and lower stratosphere over European cities, rural areas and the transatlantic flight corridor. More details can be found in Reifenberg et al. (2021) and Voigt et al. (2021). NO was measured via chemiluminescence (CLD 790 SR, ECO Physics, Dürnten, Switzerland) with a total uncertainty of 15 % and a detection limit of 5 pptv (1 min time resolution) (Tadic et al., 2020).
measurements were performed with the FAIRO (Fast AIRborne Ozone) instrument, which allows fast detection via chemiluminescence that is calibrated in situ by UV photometry (2.5 % combined uncertainty, 5 Hz time resolution) (Zahn et al., 2012). CO was measured via the quantum cascade laser spectrometer TRISTAR (Tracer In Situ Tdlas for Atmospheric Research) with an uncertainty of 3 % (1 min time resolution) (Schiller et al., 2008).

2.3 Modeling study

The modeled data were obtained from the ECHAM5 (5th generation European Centre Hamburg general circulation model, version 5.3.02)/MESSy2 (2nd generation Modular Earth Submodel System, version 2.54.0) Atmospheric Chemistry (EMAC) model which is described in Jöckel et al. (2016) and Reifenberg et al. (2021).

We use data of NO, NO$_2$, O$_3$, OH, HO$_2$, CO, CH$_4$, CH$_3$O$_2$, H$_2$O, j(NO$_2$), j(O$^1$D) temperature and pressure, modeled along the flight tracks of the described research aircraft campaigns UTOPIHAN, HOOVER and BLUESKY. The data were filtered for the troposphere using the modeled tropopause pressure. Stratospheric data were discarded. In order to evaluate the impact of reduced emissions during the COVID-19 lockdown, the model was used to simulate a scenario with usual emissions for the BLUESKY campaign which we refer to as "no-lockdown scenario". For details of the model set-up please see the paper by Reifenberg et al. (2021).

3 Results and Discussion

This analysis is structured as follows: As a full set of in situ observations necessary for a regime analysis and calculating net ozone production rates, what includes NO, O$_3$, OH, HO$_2$, CO, CH$_4$, H$_2$O, j(NO$_2$) and j(O$^1$D), is only available for the HOOVER campaign, we first show that the model and experimental data are in close agreement for this campaign. We conclude from this finding that the model is generally capable of reproducing the experimental data and therefore use the model data in our following analysis. In the second step, we provide a comparison between the three campaigns as well as the no-lockdown scenario regarding the individual trace gases and net ozone production rates. We finally present our results of the analysis of the dominant chemical regime, based on $\alpha_{CH_3O_2}$.

3.1 Comparison of Model and Experiment

Figure 2 shows a comparison of in situ observations (orange) and model simulations (blue) for the HOOVER campaign as vertical profiles. The shaded areas present the $1\sigma$ standard deviations and the numbers of data points available for each altitude bin are shown in Table S1 and S2 of the Supplement.

Figure 2a presents the vertical profile of NO which shows the typical tropospheric C-shape distribution with the highest values at the surface (e.g. vehicle and industrial emissions) and the upper troposphere (e.g. aircraft and lightning emissions). Ground-level mixing ratios (0 - 1000 m) were around 0.4 ppbv and decreased with altitude to values of $37 \pm 27$ (1σ) pptv and $47 \pm 32$ pptv for the model and the experiment, respectively, between 3 and 9 km altitude. The only relevant deviation of model and experiment was between 10 and 11 km altitude with mixing ratios of $0.20 \pm 0.03$ ppbv and $0.39 \pm 0.32$ ppbv, respectively.
Figure 2. Vertical profiles of in situ observations and model data of the atmospheric trace gases (a) NO, (b), O₃, (c) CO, (d) HO₂, (e) OH, (f) CH₄ and (g) CH₃O₂ and the photolysis rates (h) j(NO₂) and (i) j(O¹D) during the HOOVER campaign for estimating the model performance. Blue colors show model data and orange colors show experimental data. The orange trace in panel (g) shows the calculation of CH₃O₂ from experimental CH₄, CO and HO₂ via Equation (3). The shaded areas represent the 1σ standard deviation from averaging the data points at each altitude bin. The numbers of data points averaged per altitude bin are displayed in Table S1 and S2 of the Supplement.
Figure 2b shows the measured and modeled O\textsubscript{3} mixing ratios which were lowest at ground levels with 43.7 ± 14.5 ppbv and 36.4 ± 12.8 ppbv for model and experiment and increased with altitude up to 128.1 ± 22.7 ppbv and 97.5 ± 15.6 ppbv, respectively. Model values were approximately 20% higher compared to the measured data, but showed the same vertical shape.

CO vertical profiles are shown in Figure 2c which were highest at the surface with 146.4 ± 63.2 ppbv and 128.0 ± 42.3 ppbv for model and experiment, respectively, and decreased with altitude to around 70 ppbv in the upper troposphere. HO\textsubscript{x} (≡ OH + HO\textsubscript{2}) are presented in Figure 2d and e. HO\textsubscript{2} mixing ratios showed a maximum value of around 20 pptv between 2 and 3 km altitude and decreased aloft to values of around 2 pptv in the upper troposphere. Model and experiment showed close agreement. OH mixing ratios were mostly below 1 pptv. Similar to NO, the main deviation between model and experiment was between 10 and 11 km altitude where measured values were higher by around 0.5 pptv. Nevertheless, the error bars representing the 1σ standard deviation of the averages overlapped at all altitudes.

Figure 2f shows the vertical profiles of CH\textsubscript{4} which did not show any particular trend with altitude. Mixing ratios were 1809 ± 19 ppbv for the model simulation and 1815 ± 40 ppbv for the experiment throughout the campaign. CH\textsubscript{4} is needed for calculating CH\textsubscript{3}O\textsubscript{2} via Equation (3), which we show in Figure 2g in orange compared to the model simulation of CH\textsubscript{3}O\textsubscript{2}.

Figure 2h and i present the photolysis frequencies j(NO\textsubscript{2}) and j(O\textsubscript{1}D) which show close agreement for model and experiment. We show the vertical profiles for H\textsubscript{2}O, temperature and pressure in Figure S1 of the Supplement. Again, model simulation can represent the experimental data well.

For the UTOPIHAN and the BLUESKY campaigns only a limited number of observations is available. Similar to the HOOVER campaigns, NO, O\textsubscript{3} and CO can be well approximated by the model simulations which we present in Figure S2 and S3 of the Supplement. Tropospheric ozone is slightly overestimated, which we attribute to the simplified representation of multiphase chemistry (clouds) in the present model version, which underpredicts chemical ozone loss (Rosanka et al., 2021). Based on these results, we conclude that the model is generally capable of well representing the in situ observations and use the model data for all following analyses.

### 3.2 Campaign Comparison

#### 3.2.1 Trace gases

Figure 3 presents the vertical profiles of some selected trace gases during the research aircraft campaigns UTOPIHAN (green), HOOVER (blue) and BLUESKY (red) which were obtained from model simulations. Yellow lines show the no-lockdown (NL) scenario for the BLUESKY campaign in 2020.

The vertical profiles of NO are presented in Figure 3a. For all campaigns, we observe the typical C-shape as described for the HOOVER campaigns in Section 3.1. Surface (0 - 1000 m) mixing ratios were similar for UTOPIHAN and HOOVER with 0.45 ± 0.37 (1σ) ppbv and 0.43 ± 0.74 ppbv, respectively. In comparison, the ground-level concentration of NO during BLUESKY was 0.12 ± 0.11 ppbv. A possible explanation can be the general emission reduction due to legislative limitation of nitrogen oxides and other hazardous pollutants over the past decades, as the campaigns took place 15 - 20 years apart.
Figure 3. Vertical profiles of the atmospheric trace gases (a) NO, (b), O$_3$, (c) CO, (d) NO$_2$, (e) HO$_2$ and (f) OH for the campaigns UTOPI-HAN (green), HOOVER (blue), BLUESKY (red) and the no-lockdown (NL) scenario (yellow). The shaded areas represent the 1σ standard deviation from averaging the data points at each altitude bin. The numbers of data points averaged per altitude bin are displayed in Table S2 of the Supplement.

Assuming the no-lockdown scenario during BLUESKY, NO ground level mixing ratios were 0.15 ± 0.14 ppbv and therefore 25 % higher compared to actual mixing ratios (20 % emission reduction). This difference between lockdown and no-lockdown mixing ratios is slightly lower compared to the findings by other studies, for example by Donzelli et al. (2021) who found a NO decrease by 35 - 65 % in Valencia, Spain or by Higham et al. (2021) who reported a NO decrease by 55 % in the UK compared to 2019. A possible reason can be that the BLUESKY aircraft campaign took place in May and June 2020 whereas the main lockdown period across Europe occurred rather in March and April. Emission were still reduced in the following months, but likely to a smaller extent. NO was low and similar for all campaigns between 3 and 8 km altitude, a region without any particular NO sources, with most values below 50 pptv. Above 10 km, NO mixing ratios were 0.29 ± 0.19 ppbv for UTOPIHAN, 0.21 ± 0.03 ppbv for HOOVER and 0.08 ± 0.04 ppbv for BLUESKY. In comparison, NO mixing ratios for the no-lockdown scenario were 0.17 ± 0.08 ppbv above 10 km altitude. This corresponds to an emission reduction of 55 % and results in both absolute and relative NO reductions in the upper troposphere being much higher compared to ground-level
reductions. The observed NO reduction in the upper troposphere can be attributed to reduced air traffic which we show in Figure S4 of the Supplement.

Figure 3b presents the \( \text{O}_3 \) vertical profiles. For all campaigns, \( \text{O}_3 \) mixing ratios were lowest at ground levels with values of around 50 ppbv and increased with increasing altitude up to around 140 ppbv above 10 km altitude. No significant differences between the campaigns, including the no-lockdown scenario, can be observed.

CO vertical profiles can be seen in 3c. Ground level mixing ratios were 181.4 ± 39.4 ppbv for UTOPIHAN, 146.4 ± 63.2 ppbv for HOOVER and slightly lower with 103.2 ± 9.2 ppbv for BLUESKY. Mixing ratios slightly decreased with altitude. Above 3 km altitude, CO for HOOVER was lower compared to the other campaigns (mostly between 70 and 80 ppbv). Mixing ratios for UTOPIHAN were slightly higher up to 11 km altitude (between 90 and 110 ppbv) compared to BLUESKY (between 80 and 100 ppbv), but generally, significant differences are not evident.

Figure 3d shows the vertical profiles of \( \text{NO}_2 \) mixing ratios. Similar to NO, ground level \( \text{NO}_2 \) mixing ratios were highest for UTOPIHAN and HOOVER with 1.57 ± 0.77 ppbv and 2.58 ± 2.72 ppbv, respectively. In contrast, mixing ratios for BLUESKY were 0.39 ± 0.30 ppbv and 0.49 ± 0.38 ppbv considering the no-lockdown scenario which yields a 20 % \( \text{NO}_2 \) lockdown reduction, as observed for NO. We show the \( \text{NO}_2 \) range 0 - 1 ppbv for enabling the campaign distinction at low mixing ratios and present the full range in Figure S5 of the Supplement. As expected for \( \text{NO}_2 \), mixing ratios decreased with increasing altitude.

No differences between the campaigns can be observed for mid-range altitudes. In the upper troposphere, \( \text{NO}_2 \) mixing ratios for the individual campaigns showed the same behavior as for NO. Above 10 km altitude, \( \text{NO}_2 \) was on average 100.6 ± 93.2 pptv for UTOPIHAN, 70.5 ± 13.5 pptv for HOOVER and 43.1 ± 23.1 pptv for the no-lockdown scenario for BLUESKY. In comparison, BLUESKY \( \text{NO}_2 \) mixing ratios were 19.9 ± 9.8 pptv which corresponds to a 55 % reduction. In contrast to NO, \( \text{NO}_2 \) reductions were relatively higher in the upper troposphere, but absolutely higher at the surface.

Figure 3e and f show the vertical profiles of \( \text{HO}_2 \). \( \text{HO}_2 \) mixing ratios were highest at mid-range altitudes (2 - 6 km) with values up to 20 pptv and decreased aloft. OH mixing ratios were lowest at the surface (0.1 - 0.2 pptv) and increased with altitude. Above 10 km altitude, OH mixing ratios were 0.62 ± 0.38 pptv for UTOPIHAN, 0.40 ± 0.24 pptv for HOOVER, 0.30 ± 0.06 pptv for BLUESKY and 0.39 ± 0.08 pptv for the no-lockdown scenario.

### 3.2.2 Net ozone production rates

Figure 4 shows the vertical profiles of \( \text{O}_3 \) production and loss terms. All calculations were performed using model data (justified by the findings from Section 3.1) as a full set of in situ observations is only available for HOOVER, but not for UTOPIHAN and BLUESKY. Figure 4a presents net ozone production rates, which were highest at the surface with values between 1 and 2 ppbv h\(^{-1}\), but had large atmospheric variabilities, represented by the 1 σ variability shades from the vertical bin averaging. NOPRs then decreased with increasing altitude. For the HOOVER campaigns, \( \text{O}_3 \) loss dominated between 3 and 6 km altitude with NOPRs of -58.9 ± 73.4 pptv h\(^{-1}\). Negative NOPRs were also found for BLUESKY between 4 and 7 km with -18.7 ± 12.9 pptv h\(^{-1}\) and for UTOPIHAN as well as the no-lockdown BLUESKY scenario between 5 and 6 km. NOPRs were mostly positive and constant aloft. Above 10 km altitude, NOPRs were 91.7 ± 260.9 pptv h\(^{-1}\) for UTOPIHAN (51 data points), 71.2 ± 151.5 pptv h\(^{-1}\) for HOOVER (25 data points), 60.7 ± 39.7 pptv h\(^{-1}\) for BLUESKY (130 data points)
Figure 4. Vertical profiles of (a) net ozone production rates, (b) O$_3$ production via NO$_2$ photolysis, (c) O$_3$ loss via reaction with NO, (d) O$_3$ loss via photolysis, (e) O$_3$ loss via reaction with HO$_2$ and (f) O$_3$ loss via reaction with OH for the campaigns UTOPIHAN (green), HOOVER (blue), BLUESKY (red) and the no-lockdown (NL) scenario (yellow). The numbers of data points averaged per altitude bin are displayed in Table S2 of the Supplement.

and 61.4 ± 99.8 pptv h$^{-1}$ for the no-lockdown scenario. The error ranges are large and overlapping and therefore, significant differences between the campaigns cannot be observed.

Figure 4b shows O$_3$ production. We calculated the P(O$_3$) via the photolysis of NO$_2$. In contrast, NO$_2$ is not available experimentally for the HOOVER campaign in which case the approximation via the extended Leighton ratio as shown in Equation (2) is necessary. Modeled P(O$_3$) via NO$_2$ photolysis and measured P(O$_3$) via reaction of NO with O$_3$, OH and HO$_2$ are in good agreement which we show in Figure S6 of the Supplement. The only relevant deviation is observed at ground levels, where the experimental value is significantly higher compared to the modeled value. However, only three data points were available for the calculation with a 1 σ standard deviation of the averaging of > 100 %. Similar to NOPRs in Figure 4a, ground-level P(O$_3$) shows large variability with absolute values of around 10 ppbv h$^{-1}$ for BLUESKY and values of around 20 ppbv h$^{-1}$ for UTOPIHAN and HOOVER. The production term then decreased with altitude for each campaign. Significant differences
between the campaigns can only be observed at high altitudes. Above 10 km, \( \text{P(O}_3) \text{) was 4.55 ± 3.82 ppbv}\ h^{-1} \) for UTOPIHAN (51 data points) and 2.68 ± 0.90 ppbv h\(^{-1} \) for HOOVER (25 data points). For BLUESKY with the no-lockdown scenario, \( \text{P(O}_3) \text{) was 2.17 ± 0.95 ppbv}\ h^{-1} \) (130 data points) and in comparison, lockdown values were on average 0.97 ± 0.41 ppbv h\(^{-1} \) which corresponds to a 55% reduction in ozone production. We observed the same relative reduction as for NO and NO\(_2\) mixing ratios.

Consequently, net production of ozone was dominated by NO\(_x\) chemistry for all campaigns and variations in production and loss terms corresponded to the mixing ratios of NO and NO\(_2\) as presented in Figure 3. In the campaign comparison, higher NO\(_x\) concentrations (as for example for UTOPIHAN) lead to higher production and loss terms of O\(_3\) and vice versa. For the BLUESKY campaign, this analysis shows that the lockdown did not affect net ozone production rates, but instead impacted the cycling of O\(_3\) such that both production and loss rates were decreased through the reduced availability of NO and NO\(_2\) in the upper troposphere.

### 3.3 Chemical regime

As described above, the share of methyl peroxyradicals forming formaldehyde \( \alpha_{\text{CH}_2\text{O}_2} \) can be a measure for the dominant chemical regime when correlated with NO mixing ratios. Figure 5a shows the vertical profiles of \( \alpha_{\text{CH}_2\text{O}_2} \) for all available
data point for all campaigns based on the model simulation. \( \alpha_{CH_3O_2} \) values were close to 1 at the surface and decreased with altitude up to around 5 km where values of around 0.6 were observed, with no significant differences between the campaigns. \( \alpha_{CH_3O_2} \) increased again aloft whereas it was lowest for the BLUESKY campaign. Above 10 km, \( \alpha_{CH_3O_2} \) was 0.97 ± 0.03 for UTOPIHAN, 0.98 ± 0.01 for HOOVER and 0.96 ± 0.04 for the no-lockdown scenario for BLUESKY. In comparison, \( \alpha_{CH_3O_2} \) was lower for BLUESKY with 0.90 ± 0.06.

Figures 5b and c present \( \alpha_{CH_3O_2} \) in correlation with NO mixing ratios below 2 km altitude and above 10 km altitude, respectively, based on model results. Below 2 km altitude, \( \alpha_{CH_3O_2} \) ranged between 0.5 and 1.0 over the NO range of 0 - 1 ppbv. No significant trends or differences can be observed. We show \( \alpha_{CH_3O_2} \) between 2 and 10 km altitude in Figure S7 of the Supplement which does not present any differences between the campaigns either. In contrast above 10 km altitude, tropospheric \( \alpha_{CH_3O_2} \) showed a different behavior for each campaign. For an easier distinction, we show each campaign in an individual panel in Figure S8 of the Supplement. For UTOPIHAN, \( \alpha_{CH_3O_2} \) was high and almost non-responsive to changing NO mixing ratios with a slope of \( \Delta \alpha / \Delta NO \) [ppbv] = 0.09 ± 0.02 ppbv\(^{-1}\). In contrast, \( \alpha_{CH_3O_2} \) for BLUESKY was between 0.75 and 1. Small changes in NO mixing ratios caused large changes in \( \alpha_{CH_3O_2} \) with a slope of 1.12 ± 0.08 ppbv\(^{-1}\). For the no-lockdown scenario the response of \( \alpha_{CH_3O_2} \) to NO was intermediate between UTOPIHAN and BLUESKY with a slope of 0.37 ± 0.03 ppbv\(^{-1}\). These observations suggest that a VOC limited chemical regime was present during the UTOPIHAN campaign in the upper troposphere and a transition regime during the BLUESKY no-lockdown scenario, likely due to emission control over time. For BLUESKY, we observe a distinct NO\(_x\) limitation in the upper troposphere which is related to the lockdown conditions. Only few data points were available for HOOVER which were observed at similar NO levels and the response of \( \alpha_{CH_3O_2} \) to NO can therefore not be investigated. While the NOPR did not change under lockdown conditions due to compensating effects in the NO\(_x\) chemistry, we can expect impacts on tropospheric ozone from changes in VOCs (including CH\(_4\)) relevant for future emission scenarios.

### 4 Conclusions

In this study, we have presented in situ observations of atmospheric trace gases and model simulations from the EMAC model for three different aircraft campaigns across Europe, the UTOPIHAN campaigns in 2003/04, the HOOVER campaigns in 2006/07 and the BLUESKY campaign in 2020, including a modeled “no-lockdown scenario” with business as usual emissions for the latter. We found that model results can reproduce in situ observations well and thus could be used for further analysis which benefits from a more complete set of parameters and a higher data coverage. While observations for O\(_3\), CO and HO\(_x\) were very similar for all campaigns, NO\(_x\) showed significant differences, particularly in the upper troposphere, where mixing ratios were highest for UTOPIHAN and HOOVER, followed by the no-lockdown scenario for BLUESKY. Observed NO and NO\(_2\) emissions during the BLUESKY campaign were approximately 55% lower compared to the modeled no-lockdown scenario which are attributed to reduced aircraft activity at these altitudes due to the COVID-19 travel restrictions. We found a similar trend in production and loss terms of O\(_3\) which were dominated by NO\(_x\) chemistry. The COVID-19 lockdown caused a significant deceleration in O\(_3\) cycling whereas net ozone production rates were not affected by the emission reductions. Finally,
we showed that chemistry in the upper troposphere was VOC limited during the UTOPIHAN campaign, NO$_x$ limited during the BLUESKY campaign and in a transition regime for the BLUESKY no-lockdown scenario. While ground-level chemistry regimes were not found to be affected, the COVID-19 lockdown caused the predominant chemistry to shift from a transition regime to a clear NO$_x$ limited regime at high altitudes.

We found that the three aircraft campaigns, performed over a period of 17 years, represent the range from VOC to NO$_x$ limited tropospheric ozone chemistry, which can help analyze the impacts of anthropogenic emission scenarios. We encourage future studies to investigate governing chemistry in the upper troposphere, a topic which has not received much attention in literature so far, in order to get a deeper understanding of photochemical processes and the dominant ozone chemistry in a range of the atmosphere which receives its main NO$_x$ emissions from air traffic and lightning. The COVID-19 lockdown has been a unique opportunity to examine the effect of sharp reductions in primary pollutants on our atmosphere and could be a guidepost for future air policy in an effort to decrease anthropogenic emissions and to decelerate global warming.

Data availability. Data measured during the flight campaigns BLUESKY, UTOPIHAN and HOOVER are available upon request at https://keeper.mpdl.mpg.de/ to all scientists agreeing to the respective data protocols. The model results used in this study are available upon request to the author.

Author contributions. CMN and HF had the idea and designed the study. CMN analyzed the data and wrote the manuscript. AP provided the modeling data. IT provided NO data for BLUESKY. CO data for BLUESKY were received from LR. O$_3$ data for BLUESKY were obtained from FO. HH provided HO$_x$ data for HOOVER. JL and HF were significantly involved in planning and operating the research campaign.

Competing interests. Andrea Pozzer is a member of the editorial board of Atmospheric Chemistry and Physics.

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