Global gridded NO_x emissions using TROPOMI observations

Anthony Rey-Pommier^{1,2,a}, Alexandre Héraud¹, Frédéric Chevallier¹, Philippe Ciais^{1,2}, Theodoros Christoudias², Jonilda Kushta² and Jean Sciare².

¹ Laboratoire des Sciences du Climat et de l'Environnement, LSCE/IPSL, CEA-CNRS-UVSQ, Université

Paris-Saclay, 91190 Gif-sur-Yvette, France

² The Cyprus Institute, Climate and Atmosphere Research Center, 2121 Nicosia, Cyprus
 ^a Now at: European Commission, Joint Research Centre, 21027 Ispra, Italy

Correspondence: Anthony Rey-Pommier (anthony.rey-pommier@lsce.ipsl.fr)

Abstract. We present top-down global gridded emissions of NO_x for the year 2022. This dataset is constructed Q from retrievals of tropospheric vertical column densities of NO_2 by the TROPOMI spaceborne instrument associated 10 with winds and atmospheric composition data from ECMWF reanalyses, using an improved version of a mass-balance 11 atmospheric inversion. The dataset has a spatial resolution of $0.0625^{\circ} \times 0.0625^{\circ}$, and delivers a detailed overview 12 of the distribution of emissions. It allows the identification of intense area sources, such as cities, and isolated 13 emitters, such as power plants or cement kilns, but does not correctly represent biomass burning. At global level, 14 the emissions obtained are consistent with the EDGARv6.1 bottom-up inventory, although there are differences at 15 regional level, particularly in emerging countries and countries with low observation densities. The emissions of the 16 three largest emitting countries, China, the United States and India, are 6, 14 and 4% lower than EDGAR estimates. 17 Uncertainties remain high, and a quantitative analysis of emissions over several averaging periods indicates that 18 averaging emissions uniformly across the year may be sufficient to obtain estimates consistent with annual averages, 19 in regions of the world with high retrieval densities. This dataset is designed to be updated with a low latency to 20 help policymakers monitor emissions and implement energy savings and clean air quality policies. The data can be 21 accessed at https://doi.org/10.5281/zenodo.13758447 as monthly files (Rey-Pommier et al., 2025). 22

²⁴ 1 Introduction

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Air pollution is one of the leading causes of premature death in the world. Public health policies, implemented 25 at the scale of countries, regions or cities, often aim to reduce the exposure to several pollutants, such as nitrogen 26 oxides $(NO_x = NO + NO_2)$. Such mitigation plans therefore require a precise knowledge of the emitters, as well as 27 a monitoring of their emission levels over time. Data on NO_x emissions is therefore fundamental for monitoring the 28 implementation of air quality policies. Besides, because NO_x is mainly produced during the combustion of carbon fuels 29 at high temperatures, such data can also be, in conjunction with NO_x/CO_2 ratios derived at the scale of industrial 30 sectors and countries, a tool to measure progress towards carbon neutrality. Gridded emissions with high spatial and 31 temporal resolution are therefore of great scientific and political value. Many of such datasets are emission inventories, 32 i.e. bottom-up models in which emissions are calculated on the basis of known sectoral activities and allocated in 33 time and space, combined with specific emission factors by sector and, possibly, by country. These inventories provide 34 valuable information on long-term trends and large-scale emission budgets, but they suffer from several weaknesses. 35 They hardly represent daily or weekly variations, their activity data may be outdated, and some sources may be 36 misallocated or unknown, which is common in many developing countries. Besides, uncertainties surrounding rapidly 37 changing emissions factors and the generally low temporal resolution of activity data limits, in certain circumstances, 38 the realism of such bottom-up inventories. Finally, they have a data lag of at least three years, which limits their 39 potential as monitoring tools. 40

In this context, increasing efforts have been made to overcome the weaknesses of the inventories in order to obtain independent emission datasets that are homogeneous from one country to another. Such datasets are of the top-down type: they use direct observations of pollution and result from the inversion of an atmospheric chemistrytransport model (CTM) in which these atmospheric observations are assimilated. The observation data may be in-situ measurements or satellite retrievals.

In previous studies, we used a method for detecting and quantifying NO_x emissions from daily observations of NO_2 46 columns by the TROPOMI instrument, onboard the Sentinel 5P satellite. This method, developed for the countries 47 of the Eastern Mediterranean and Middle East region, is based on a two-dimensional simplification of atmospheric 48 chemistry and transport, and does not require the direct use of a full 3D chemistry-transport model. Here, we extend 49 the emissions domain to the whole world for the year 2022, and provide a dataset of averaged NO_x emissions at 50 a resolution of $0.0625^{\circ} \times 0.0625^{\circ}$. We analyse the results by pinpointing emitters and distinguishing between point 51 sources, generally corresponding to isolated industrial facilities, and diffuse/area sources, generally corresponding to 52 megacities. We also compare the results with the bottom-up inventory EDGARv6.1 and assess their reliability using 53 different average horizons. 54

This article is structured as follows: Section 2 details the method used throughout this study, its improvements and simplifications since its previous uses, and the input data in its implementation. Section 3 presents the global NO_x emissions dataset and analyses the different types of emitters. It also compares the results obtained with the EDGARv6.1 bottom-up inventory, and analyses different time horizons for averaging daily emissions in order to obtain representative results. Section 4 analyses the applicability limits of the method and highlights sources of uncertainty.

$_{60}$ 2 Methods

61 2.1 Input data

62 2.1.1 TROPOMI NO₂ column densities

 NO_2 can be observed from space with satellite instruments based on its strong absorption features in the 400–465 nm 63 wavelength region (Vandaele et al., 1998). By comparing observed spectra with a reference spectrum, the amount of 64 NO_2 in a portion of the atmosphere between the instrument and the surface can be derived. The TROPOspheric Mon-65 itoring Instrument (TROPOMI), onboard the European Space Agency's (ESA) Sentinel-5 Precursor (S-5P) satellite, 66 is one of those instruments (Veefkind et al., 2012). This instrument has a large swath width (~ 2600 km), combined 67 with the 15-day orbit cycle of the satellite, leading to a revisit time of one day for every point of the Earth in absence 68 of clouds. Moreover, these daily measurements are always collected during the middle of the day, the satellite crossing 69 the sunlit equator at around 13:30 local time (LT). The high spatial resolution of the instrument (up to 3.5×5.5 km² 70 since 6 August 2019) allows observing fine-scale structures of NO_2 pollution, such as hotspots within medium-size 71 cities or plumes from power plants and industrial facilities. Tropospheric vertical column densities (VCDs, or sim-72 ply "columns") are provided after retrieval of total slant column densities using the Differential Optical Absorption 73 Spectroscopy method (Platt et al., 2008). VCDs represent the integrated number of NO_2 molecules per surface unit 74 between the surface and the tropopause at the corresponding vertical. An algorithm also supplies an air mass factor, 75 which is the ratio between slant and vertical column densities. This factor is derived from the knowledge of many 76 physical quantities such as the vertical distribution of the absorber but also the viewing angle and the albedo of the ob-77 served surface. It comprises a significant part of the uncertainty in NO_2 measurements (Boersma et al., 2004; Lorente 78 et al., 2019), which becomes non-negligible in a polluted atmosphere. Each TROPOMI retrieval is also associated 79 with a quality assurance value q_a , which ranges from 0 (no data) to 1 (high-quality data). We selected NO₂ retrievals 80 with q_a values greater than $q_{a,\lim} = 0.75$, which correspond to clear-sky conditions (Eskes et al., 2022). Here, we use 81 TROPOMI NO₂ retrievals in 2022 (OFFL product using processor version 2.5.0, product version 2.3.1 and 2.4.0 before 82 and after November 2022 respectively). To limit effects due to product of processor version changes, other years are 83 not studied. 84

85 2.1.2 Meteorological and air composition fields

Horizontal wind is taken from the ERA5 data archive, provided by the European Centre for Medium-Range Weather 86 Forecasts (ECMWF). Both components have a horizontal resolution of $0.25^{\circ} \times 0.25^{\circ}$ gridded on 37 vertical pressure 87 levels (Hersbach et al., 2020). We vertically average wind fields using the first two vertical levels, at 975 and 1000 88 hPa, except for representing ground winds, for which the last level at 1000 hPa is used. ECMWF also produces a 89 reanalysis for air composition, under the Copernicus Atmospheric Monitoring Service (CAMS). It provides analyses 90 and forecasts for reactive gases, greenhouse gases and aerosols. These parameters are gridded on 25 vertical pressure 91 levels with a horizontal resolution of $0.4^{\circ} \times 0.4^{\circ}$ and a temporal resolution of 3 hours (Huijnen et al., 2016). Here, 92 concentrations of NO₂, NO, OH, as well as temperature, are taken from CAMS to represent chemical processes in our 93 model. Fields are vertically averaged using the first two vertical levels, at 950 and 1000 hPa. 94

95 2.1.3 Elevation data

For computing altitude gradients, we use the Global Multi-resolution Terrain Elevation Data (GMTED2010, Danielson and Gesch (2011)) with a 0.0625°×0.0625° resolution. Elevation data is re-gridded on the TROPOMI grid, before calculation of the corresponding gradient to derive a corrective "topography-wind" value that is detailed in Section 2.2.2.

¹⁰⁰ 2.2 The mass-balance inversion

¹⁰¹ 2.2.1 Main principle

The flux-divergence method is a mass-balance inversion model calculating the emissions of a given trace gas from 102 observations of the corresponding vertical tropospheric columns, which is particularly well suited to data with high 103 spatial resolution. In the case of NO_2 , this approach was pioneered by Beirle et al. (2019). It has subsequently been 104 implemented differently by other researchers, in different circumstances under simplified forms or, on the contrary, 105 more complex ones (Lama et al., 2020; Rey-Pommier et al., 2022; de Foy and Schauer, 2022; Sun, 2022). The flux-106 divergence method is based on the conservation of mass principle, which makes it possible to calculate emission 107 densities at the pixel scale as a function of a transport term and a sink term. By noting C the local concentration of 108 NO₂ and $\mathbf{w} = (u, v, w)$ the mean wind at the time of measurement, the corresponding emissions E_C are expressed as: 109

$$E_C = \frac{\partial C}{\partial t} + \operatorname{div}(C\mathbf{w}) + S_C \tag{1}$$

Here S_C is the sink term expressing the loss of NO₂ due to chemical reactions. Assuming that the vertical variations in concentration are small compared with the horizontal variations, and considering that most NO₂ remains confined close to the ground, the previous equation can be rewritten in terms of tropospheric columns Ω , which enables, in steady state, the computation of emissions per surface area E, as:

$$E = \frac{\partial(\Omega u)}{\partial x} + \frac{\partial(\Omega v)}{\partial y} + S_{\Omega}$$
⁽²⁾

 S_{Ω} is the sink term expressed by surface unit. $D = \frac{\partial(\Omega u)}{\partial x} + \frac{\partial(\Omega v)}{\partial y}$ is the horizontal advection (transport) term. The assumption of a stationary state and a pollution close to the ground means that the temporal and vertical dimensions of the problem can be ignored, resulting in a purely 2D calculation of emissions. The corresponding reduction in complexity means that inversions can be performed very quickly compared with the conventional use of full-fledged 3D CTMs and without *a priori* knowledge on emissions. While useful, these simplifications come with inherent uncertainties, the main sources of which being on the input tropospheric columns, wind direction and atmospheric composition. It must also be noted that far from strong and localized sources, the underlying assumptions of stationarity and pollution containment are no longer valid.

Finally, the NO₂ production can be converted into NO_x emissions. Performing this conversion is accounting for the portion of NO_x, mainly emitted as NO, which is not converted into NO₂ by reaction with ozone. The reformation of NO by the photolysis of NO₂ during the day leads to an equilibrium between the two compounds. The ratio $\mathcal{L} = [NO_x]/[NO_2]$ usually varies between 1.2 and 1.4, depending on local conditions. NO_x emissions are therefore calculated as:

$$E_{\rm NO_x} = \mathcal{L}E \tag{3}$$

¹²⁷ In most urbanized areas, daytime NO concentrations frequently exceed 20 ppb. Under such conditions, this ratio is ¹²⁸ stabilized in a few minutes (Graedel et al., 1976; Seinfeld and Pandis, 2006). As this time is shorter than the inter-mesh ¹²⁹ transport timescale, the impact of stabilization time on the overall emission composition can be justifiably ignored. ¹³⁰ Here, it is near emission sources that the stationary hypothesis may not be applicable, in which case the value of \mathcal{L} ¹³¹ could be significantly higher than 1.4. The implications of this neglect are discussed in Section 4.1.

132 2.2.2 Refined version

¹³³ In order to consider only anthropogenic pollution located close to the ground, it is necessary to remove any signal of ¹³⁴ natural emissions from the tropospheric columns provided by TROPOMI. In the absence of anthropogenic sources, ¹³⁵ the NO₂ columns that are observed constitute a tropospheric background Ω_b . At the global scale, this background is ¹³⁶ mostly due to soil emissions in the lower troposphere (Yienger and Levy, 1995; Hoelzemann et al., 2004). In the upper ¹³⁷ troposphere, NO₂ sources include lightning, convective injection and downwelling from the stratosphere (Ehhalt et al.,

1992). We remove that background by calculating the 1st tercile in a 200 pixel \times 430 pixel zone around each pixel 138 (along \times across track, i.e. approx. 700 km \times 2360 km). We assume that this zone is sufficiently large whatever the 139 considered pixel so that this tercile corresponds to the typical local value for this background. The large across-track 140 distance is chosen for two reasons. Firstly, it limits the influence of low-quality observations and intermediate pollution 141 levels on the background estimate, which can inflate the estimation when performed on a smaller domain and prevent 142 the detection of smaller emitters. Secondly, it minimizes the mixing of pixels with varying NO_2 backgrounds due to 143 distinct climatic and geographical conditions, such as between arid and temperate regions. Such variations are less 144 pronounced across-track than along-track. Besides, the overpass of Sentinel-5P around 13:00 LT, when diurnal NO_x 145 variations are minimal, limits the mixing of pixels with backgrounds corresponding to different local times. After 146 calculation of the background, it is subtracted from the vertical tropospheric column density, and the resulting lower 147 tropospheric vertical density $\Omega' = \Omega - \Omega_b$ is used in the flux-divergence method. Pixels with columns lower than the 148 calculated background have a corrected column reduced to $\Omega' = 0$. Such assumption can be challenged above macro-149 regions for which soil emissions and wildfires result in high NO₂ values observed by TROPOMI. High tropospheric 150 backgrounds can also arise from localized paths of long-range transport of reactive nitrogen (Zhai et al., 2024) or 151 around shipping lanes, where exhaust emissions directly increase NO₂ levels and may also enhance lightning activity 152 that produces additional NO_x (Thornton et al., 2017). The neglect of such effects is highlighted in Section 4.1. 153

We represent the sink term S_{Ω} by considering only the chemical loss of NO₂ due to its reaction with the hydroxyl 154 radical (OH). This reaction follows a first-order kinetics, and the sink term can be expressed as $S_{\Omega} = k_{\text{OH}+\text{NO}_2}[\text{OH}]\Omega'$ 155 with k_{OH+NO_2} the reaction rate whose value is given by Burkholder et al. (2020). This is equivalent to compute a 156 mixed lifetime $\tau = 1/(k_{\text{OH}+\text{NO}_2}[\text{OH}])$. This lifetime generally ranges between 1 and 14 h in mid-latitude regions and 157 reaches higher values in polar and subtropical zones. Two global maps of lifetimes for winter and summer, computed 158 with this method, are added in the Supplementary Materials. In many studies, this quantity is kept uniform and 159 constant in the use of the flux-divergence method (Beirle et al., 2019; de Foy and Schauer, 2022), because justified by 160 a relatively small domain of interest. Here, a singularity of our version of the flux-divergence method is to account for 161 the temporal variability of OH, which is primarily driven by the amount of UV radiation from the stratosphere, but 162 also for its spatial variability, since OH can also be influenced by NO_x through a non-linear relationship (Valin et al., 163 2011). In this respect, our sink term is heavily reliant on the NO_x sources accounted for in CAMS data. Neglecting 164 a source, or mis-estimating the order of magnitude of its NO_x emissions, therefore results in a wrong OH field whose 165 bias depends on the amplitude of the neglect. Similarly, the coarse resolution in CAMS data $(0.4^{\circ} \times 0.4^{\circ})$ can fail to 166 represent the particular conditions within or downwind power plant plumes, leading to a wrong estimation of the real 167 OH budget. We regard these effects as less pronounced compared to those that would result in representing a constant 168 lifetime for NO_2 which oversimplifies and misrepresents temporal and spatial dynamics by representing all situations 169 the same way, whether they represent emitters or not. 170

Additionally, systematic artifacts concerning advection processes were reported over regions with complex topographies, particularly when high tropospheric vertical column densities are observed over mountainous regions. These high values can hinder the identification and quantification of point sources, possibly due to inaccurate mean wind fields over mountains. A study by Sun (2022) shows that these patterns can also be caused by 3D transport effects which have been ignored in the simplified 2D approach which has been described so far. A "topography-wind" V term can be introduced in Equation 3 in order to correct for this effect using ground wind \mathbf{w}_g , the topography gradient ∇z_0 , and an inverse scale height X_e as follows:

$$V = X_e \Omega' \mathbf{w}_g \cdot \nabla z_0 \tag{4}$$

Here, we choose a uniform and constant value of $X_e = 0.3 \text{ km}^{-1}$. Although this value is about one order of magnitude lower than that used by Beirle et al. (2023), it corresponds to the approximate mean inverse scale height calculated by Sun (2022) in which a variability for X_e was allowed by fitting its value through linear regressions on the basis of selected observations. While we acknowledge that choosing a single value for X_e is a simplification, we note that performing the fit of its value would require an arbitrary selection of the cells used for that fit. We therefore compute the following equation to estimate NO_x emissions:

$$E_{\rm NO_x} = \mathcal{L}(\frac{\partial(\Omega' u)}{\partial x} + \frac{\partial(\Omega' v)}{\partial y} + k_{\rm OH+NO_2}[\rm OH]\Omega' + X_e \Omega' \mathbf{w}_g \cdot \nabla z_0)$$
(5)

Following de Foy and Schauer (2022), we perform the calculation of derivatives directly on the original TROPOMI grid (along-track and across-track) to better handle pixels with low-quality or no data, resulting in lower discontinuities in the calculated transport term. To do so, we re-grid the wind field on the TROPOMI grid and linearly interpolate the estimates at the satellite timestamp. We do the same for all other parameters that are concerned for the calculation of the sink term (concentrations of OH, NO and NO₂, and temperature). Emissions are thus calculated on the TROPOMI grid and are then re-gridded on a regular north-south/east-west grid with a $0.0625^{\circ} \times 0.0625^{\circ}$ resolution.

Finally, the accuracy of TROPOMI retrievals can be compromised by challenges in estimating the air mass factor 190 or local effects, particularly in specific vertical distribution scenarios (Griffin et al., 2019; Lorente et al., 2019; Judd 191 et al., 2020). The latest versions of TROPOMI (v2.x) showed VCD values higher than those of earlier versions (v1.x), 192 with biases up to 40%, depending on pollution levels and seasonal variations (Van Geffen et al., 2022). Additionally, the 193 chemistry-transport model TM5, which is integrated into the operational TROPOMI product, tends to underestimate 194 pollution near the ground, while overestimating NO_2 concentrations at higher altitudes over the sea (Latsch et al., 195 2023; Rieß et al., 2023). To compensate for such effects, studies like Goldberg et al. (2022) or Beirle et al. (2023) 196 corrected the used VCDs by changing the corresponding vertical sensitivity over emitters. In this study, we do not 197 perform such adjustment, while recognizing it could constitute a further step in the improvement of our dataset. On 198 Figure 1, we sum up the functioning of our method. 199

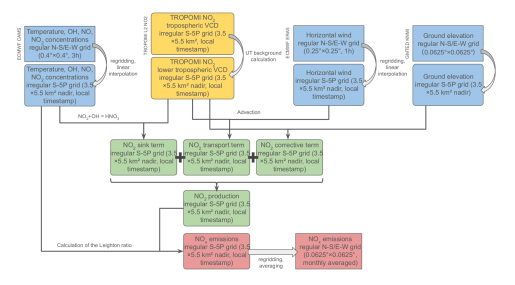


Figure 1: General overview of the mass-balance inversion.

200 2.3 EDGAR bottom-up inventory

Many high-resolution datasets for air quality exist at global (Benkovitz et al., 1996; Granier et al., 2019) or regional 201 scale (Kuenen et al., 2022; He, 2012). Here we compare our averaged emissions for the year 2022 to NO_x emissions 202 provided by The Emissions Database for Global Atmospheric Research (EDGARv6.1) for 2018. It is a global inventory 203 providing $0.1^{\circ} \times 0.1^{\circ}$ gridded emissions of greenhouse gases and air pollutants at the monthly scale, covering different 204 sectors (Crippa et al., 2020). It is based on activity data of different nature (population, industrial processes, energy 205 production, fossil fuel extraction, agricultural outputs, etc.) derived from the International Energy Agency (IEA) and 206 the Food and Agriculture Organization (FAO), and the emission factors corresponding to each of the covered sectors. 207 National and regional information on technology mix data provide a better characterization of these emission factors. 208 End-of-pipe measurements are also used for correcting purposes. The version 6.1 of the inventory covers the years 209 1970-2018. 210

211 **3** Technical validation

$_{212}$ 3.1 Spatial distribution of the global NO_x emissions

The global map of the averaged NO_x emissions for 2022 is shown on Figure 2, while Figure 3 zooms over seven macroregions that cover most of the emitters over land and sea. Emissions are represented as density, i.e. by surface unit. All the analyses carried out in this study are based on the displayed domain, i.e. between latitudes 65°S and 65°N. This discards frequent outliers above these latitudes, resulting from monthly and annual estimates based on too few observations. Significant regional differences appear on these maps. The highest values are concentrated in developing areas such as eastern China, India and the Middle East. High values are also found in Europe, Russia and the United

States, where they correspond to megacities and industrial areas. Transport emissions can also be highlighted where 219 they provide the highest share of emissions, i.e. on highways and shipping lanes which appear in various regions. 220 South America, Oceania and sub-Saharan Africa display low or zero emissions except in a small number of cities and 221 industrial sites. Wildfires, which are frequent in rainforests and savannas (Mebust and Cohen, 2013; Castellanos et al., 222 2014; Ossohou et al., 2019; Opacka et al., 2022), display quasi-zero emissions in Amazonia and low emissions in the 223 Congo basin. Wildfire emissions might be under-estimated due to a wrong estimation of the lifetime, in particular in 224 tropical regions where sinks other than the reaction with OH are important. Such other sinks are developed in Section 225 4.1. It must be noted that at a lower temporal scale, wildfire emissions display an annual variability. The example of 226 the fires in the Congo basin is studied in the Supplementary Materials, with high emissions during summer (JJA). It 227 is thus possible that a large number of smaller wildfires, occurring during other seasons, are too small to be correctly 228 observed from space, as shown by other studies (Ramo et al., 2021; Khairoun et al., 2024). 229

Generally speaking, the maps highlights the industrialized areas, revealing the world's main megacities where 230 several sources of emissions (traffic, power, residential) are mixed. Some industrial facilities and large power plants 231 also appear. Emissions are correctly resolved in most regions of the world. The observed spread of emissions over two 232 to three pixels (i.e. about 12 to 20 km) further away from the exact location of the corresponding emitters is due to 233 the turbulent spread of emissions, which is not considered in our method. Finally, we note that emissions in mid- and 234 high-latitude regions (beyond about 40° from the Equator) are noisy, due to an averaging over a smaller number of 235 clear-sky days throughout the year. On average, countries such as Egypt, Niger and Saudi Arabia are observed about 236 70% of the time with a quality flag higher than $q_{a,\text{lim}} = 0.75$, while Ireland, Canada and Finland are observed less 237 than 20% of the time. This uneven sampling is also present in tropical regions where rainfall is frequent, as there is no 238 measurement during cloudy scenes. Countries like Gabon, Indonesia or Peru are observed less than 30% of the year 239 with quality flags higher than the threshold. In some cases, this low density of observations prevents emissions from 240 intense sources from being quantified correctly at the monthly scale, as it is discussed in Section 3.4. 241

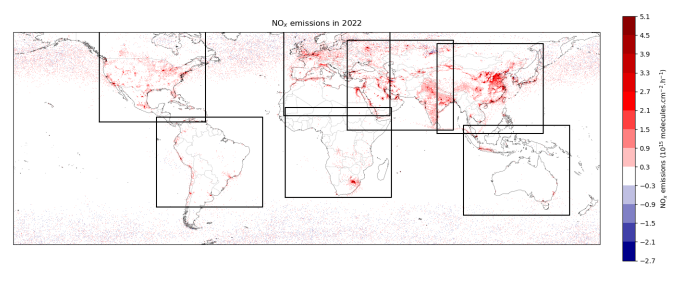


Figure 2: TROPOMI-derived mean daytime NO_x emission rates in 2022 estimated with the flux-divergence method. The seven frames correspond to macro-regions whose emissions are specifically shown in Figure 3.

The statistical distribution of emissions is shown in Figure 4. Four different regimes of emissions can be distinguished in the red curve (note the log-log scale):

Very low values of emission densities (less than ~0.02 Pmolecules.cm⁻².h⁻¹), in practice at places where there are almost no emissions in reality. Note that, as the calculated fluxes represent averaged emissions, such pixels can also represent places where high emissions occurred, but only during a small portion of the year, as it is the case in regions where wildfires frequently happen.

- Residual emission densities (between ~0.02 Pmolecules.cm⁻².h⁻¹ and ~0.2 Pmolecules.cm⁻².h⁻¹), for which it is difficult to determine the corresponding source.
- Low emission densities (between ~ 0.2 and ~ 2 Pmolecules.cm⁻².h⁻¹), generally high enough to be associated with an emitter, but too low for a reliable quantification to be possible unless heavy averaging. The upper limit corresponds approximately to the emission densities observed on smaller power plants.

• High emission densities (higher than 2 Pmolecules.cm⁻².h⁻¹), where the signal-to-noise ratio is high enough to quantify emissions when enough observations are averaged.

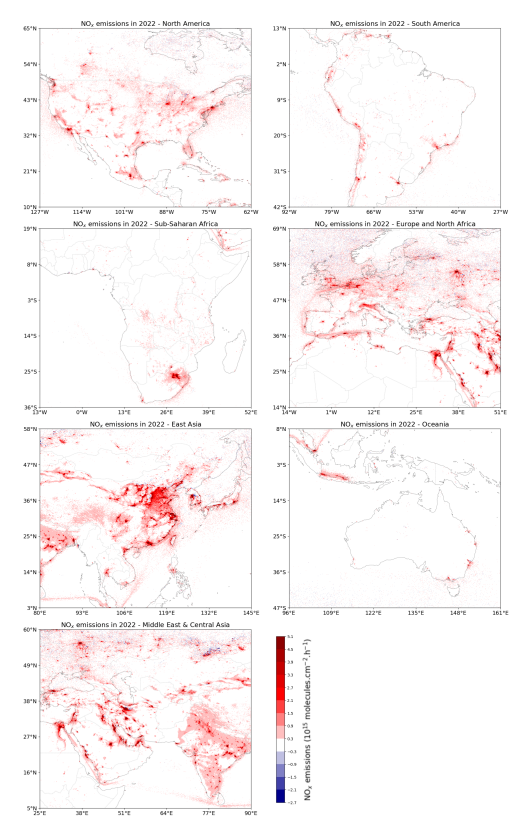


Figure 3: TROPOMI-derived mean daytime NO_x emission rates in 2022 estimated with the flux-divergence method for North America, South America, sub-Saharan Africa, Europe and North Africa, East Asia, Oceania, Middle East and Central Asia.

Figure 4 also shows negative values (blue curve), even though negative emissions are physically impossible. 255 They appear in practice because the transport term, which includes a derivative, can be negative. In calculated 256 emission densities, negative pixels of low absolute value are as numerous as positive pixels of the same amplitude; 257 they correspond to numerical noise and are found in pollution-free zones where the sink term is virtually zero. Higher 258 values for negative pixels are less frequent: we count about 5 times less pixels with emission densities lower than -0.2 259 Pmolecules.cm⁻².h⁻¹ than pixels with emission densities higher than 0.2 Pmolecules.cm⁻².h⁻¹ (yellow and red parts 260 of the graph in Figure 4). The locations where such high values are observed for negative pixels correspond to areas 261 close to anthropogenic sources of NO_x , but in situations for which the absolute transport term has been overestimated 262 or the sink term has been underestimated. Such negative emissions are limited to rare cases, such as Tehran, which 263 is discussed in Section 4.2. 264

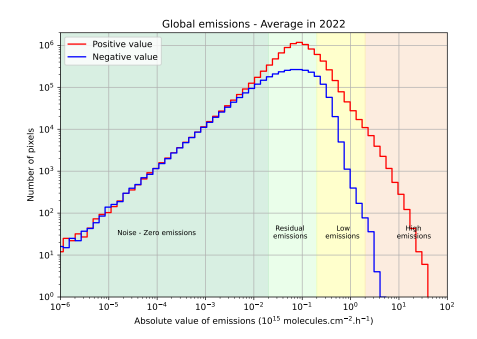


Figure 4: Distribution of positive and negative TROPOMI-inferred NO_x emissions for year 2022. Four regimes can be distinguished (the values defining the thresholds between these regimes are given as order of magnitudes).

²⁶⁵ 3.2 Diffuse sources and point sources

The assimilation of high-resolution observations with the flux-divergence method holds a significant potential for 266 pinpointing emissions at small scale. As a consequence, it reveals the difference between sources that emit pollutants 267 from a localized area, called point sources, from diffuse sources emitting pollutants over a wider area, such as sprawling 268 urban regions like megacities. While the extent of the observed NO_2 pollution created by a point source is primarily 269 determined by advection and turbulent mixing, the spread of the pollution for a diffuse source is above all determined 270 by the spatial extent of the source itself. Point sources are therefore characterized by a dominance of the transport 271 term, while diffuse sources (the term "area sources" is also used) exhibit a balance or dominance of the sink term 272 (Beirle et al., 2019). Within the flux-divergence method, these two types of sources can be identified differently, since 273 274 the main sources of uncertainty come from wind angle in the case of a point source while they come from the OH concentration explaining the sink term for a diffuse source. Because this distinction remains qualitative, to classify a 275 detected source as one or the other type, arbitrary thresholds must be defined, concerning the number of pixels above 276 a certain value of emissions, or the share of the transport term within the emissions in Equation 5. Here, we catalog 277 all sources in the averaged emissions map for 2022. Firstly, we define a source as a cluster of at least 3 contiguous 278 pixels above the value of 2 Pmolecules. cm^{-2} . h^{-1} . We then classify these sources as "point" or "diffuse" according 279 to the number of pixels in the detected cluster, point sources being the clusters comprising 3 to 9 pixels, and diffuse 280 sources those with more than 10 pixels. We detect 436 point sources and 323 diffuse sources, whose locations are 281 displayed on Figure 5. The statistical distribution of the emitters, as well as their detailed location, are provided in 282 the Supplementary Materials and in Rey-Pommier et al. (2025). 283

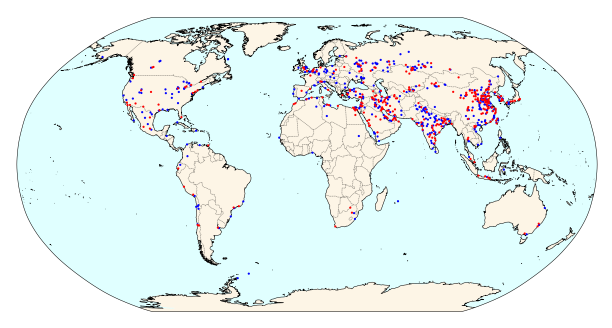


Figure 5: Location of different point sources in blue (between 3 and 9 contiguous pixels above 2 Pmolecules.cm⁻².h⁻¹) and diffuse sources in red (more than 10 contiguous pixels) for 2022.

284 3.2.1 Diffuse sources

Most point sources correspond to facilities such as power stations, cement kilns or mining sites. They can also 285 correspond to concentrated urban areas. Conversely, most diffuse sources correspond to urban areas of megacities, 286 whether they comprise industrial facilities within their extent or not. Exceptions concern mega-emitters like the 287 Medupi and Matimba power plants in South Africa, mentioned in various articles (Reuter et al., 2019; Hakkarainen 288 et al., 2021; Cusworth et al., 2023) or the Ain Sokhna industrial area in Egypt, already mentioned in Rey-Pommier 289 et al. (2022). In both cases, such groups of industrial facilities exhibit particularly high emissions over more than 290 10 pixels and are detected as diffuse sources. Figure 6 displays the emissions of diffuse sources corresponding to 291 megacities: Baghdad (32.3 t.h⁻¹, 198 pixels), Istanbul (15.4 t.h⁻¹, 127 pixels), Mexico City (17.6 t.h⁻¹, 114 pixels), 292 Moscow (19.0 t.h⁻¹, 177 pixels), Riyadh (33.0 t.h⁻¹, 171 pixels) and Shanghai (100.2 t.h⁻¹, 836 pixels). City cores are 293 denoted with dashed lines, and generally correspond to areas where emissions are largely above the cluster-detection 294 threshold. Table 1 shows the 20 diffuse sources with the highest emissions. 295

Number of	Latitude	Longitude	Mean emission density	Output	Emitter	
pixels in cluster	(°N)	(°E)	$(Pmolecules.cm^{-2}.h^{-1})$	$(t.h^{-1})$		
2623	37.617	116.030	2.818	217.89	Beijing urban area, China	
836	31.283	120.354	3.771	100.21	Shanghai urban area, China	
443	35.557	51.321	6.939	93.05	Tehran urban area, Iran	
554	-26.407	28.738	4.217	77.79	Gauteng coal region, South Africa	
417	22.796	113.626	3.691	52.78	Shenzhen & Hong-Kong urban area, China	
364	29.648	31.129	4.064	47.84	Cairo & Beni Suef urban area, Egypt	
302	29.585	47.872	4.496	43.95	Kuwait City urban area, Kuwait	
171	24.650	46.791	5.708	33.01	Riyadh urban area, Saudi Arabia	
198	32.775	44.301	5.223	32.36	Baghdad urban area, Iraq	
255	41.124	123.005	4.281	30.60	Anshan urban area, China	
347	39.339	110.659	2.933	29.31	Ordos mining region, China	
169	25.251	55.348	4.790	27.23	Dubai urban area, United Arab Emirates	
193	37.162	126.822	4.425	25.30	Seoul urban area, South Korea	
157	32.583	51.602	4.796	23.62	Ispahan urban and industrial area, Iran	
124	21.112	39.313	4.886	21.03	Djeddah urban area, Saudi Arabia	
220	37.320	112.088	3.131	20.39	Shanxi urban area, China	
177	55.706	37.508	5.121	19.02	Moscow urban area, Russia	
101	24.120	82.744	5.461	18.73	Jogi Chaura industrial zone, India	
158	39.327	106.809	4.116	18.72	Wuhai/Hainan industrial zone, China	
83	-12.183	-76.853	6.101	18.41	Lima urban area & Pachamac mines, Peru	

Table 1: List and location of the 20 diffuse sources with highest TROPOMI-inferred NO_x emissions (expressed as NO_2), and corresponding size of the cluster and main sector responsible for the emissions.

These six diffuse sources differ greatly from one another: Baghdad, Mexico City and Riyadh are very dense and isolated, allowing their emissions to stand out from the rest of the hotspots, while Moscow and Istanbul are less dense, resulting in lower emission densities. The Shanghai urban area has a large spatial extent, and the associated cluster extends over an area much wider than the city limits. Finally, it should be noted that Moscow and Shanghai experience many cloudy days, resulting in a fairly low level of averaging, leading to numerical noise that is visible on the maps. Many industrial facilities near city centers do not have high emissions, possibly due to an irregular production throughout the year, with high-activity periods covered by clouds.

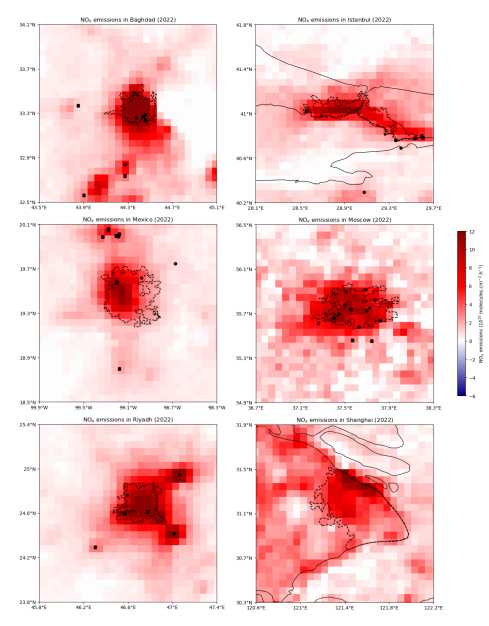


Figure 6: Map of mean daytime TROPOMI-inferred NO_x emissions for 2022 (expressed as NO_2) for six megacities (diffuse sources), clockwise: Baghdad, Istanbul, Moscow, Shanghai, Riyadh, and Mexico City. The approximate boundaries of the city cores are denoted with dashed lines and the approximate location of power plants and other industrial facilities are denoted with circles and squares respectively, except for Shanghai (unavailable data).

303 3.2.2 Point sources

With a manual verification of the 436 detected point sources, we identify 48 outliers, 26 of which being points in places totally empty from any anthropogenic activity, and 22 points in areas with anthropogenic activity but without significant source (no facility of significant size). Most of these outliers are located in high-latitude regions, with 29 of them being located north to the 50°N parallel.

Because a threshold has been introduced to detect emitters, classified sources are isolated from each other. 308 For many of them, emissions peak within the associated cluster. With a threshold set at 2 Pmolecules. $cm^{-2}.h^{-1}$, 309 the corresponding signal-to-noise ratio is generally high enough to perform peak-fitting around the source, enabling 310 accurate emission derivation. While this method works well for most point sources, it is not directly applicable to 311 many diffuse sources. Since the observed spread of emissions around a source is caused by turbulent diffusion, a 312 2D-Gaussian function is applied to fit the detected sources within a 15×15 pixel zone around the maximum emission 313 density within the cluster. Three examples are shown for the city of Medina, Saudi Arabia, the Sohar Industrial zone, 314 Oman and the Western Mountain power plant, Libya on Figure 7. Note that these locations correspond to sources 315 well-isolated from other industrial activities, in countries with frequent cloud-free conditions that allowed an averaging 316 over high number of days in 2022. 317

We acknowledge the fact that the value of 2 Pmolecules.cm⁻².h⁻¹ (corresponding to 37 kg.h⁻¹ for a pixel at 60°N 318 or 60°S, to 74 kg, h^{-1} for a pixel at the equator) to mark the limit between high and low emissions is arbitrary, as other 319 values for this threshold could be used. For instance, the Beijing cluster, identified on Table 1, with a size of 2623 320 pixels, is broken down into 35 smaller clusters (13 diffuse sources and 22 point sources) when changing the threshold 321 from 2 Pmolecules.cm $^{-2}$.h $^{-1}$ to 3 Pmolecules.cm $^{-2}$.h $^{-1}$. These new clusters represent better urban sprawling around 322 the various megacities and industrial facilities in Eastern China. However, in the same region, three point sources 323 disappear when performing this threshold change. To determine the sensitivity of the point source and diffuse source 324 detection and classification method, we carry out the detection by changing this threshold from 2 Pmolecules.cm $^{-2}$.h $^{-1}$ 325 to 3 and 4 Pmolecules.cm⁻².h⁻¹. A comparative map is displayed in the Supplementary Materials. The point sources 326 and diffuse sources are identified, and a fit with a 2D-Gaussian is carried out on point sources to estimate better 327 emissions by accounting for the Gaussian nature of turbulent diffusion around the source. We then count the number 328 of point sources with a fit of correct quality (with a correlation coefficient R^2 higher than 0.4). The results are shown 329 in Table 2 for the different thresholds, and we compare the countries with the most point sources. Note that among 330 the 48 outliers identified in the detected point sources with the threshold of 2 Pmolecules. $cm^{-2}.h^{-1}$, only 11 reached 331 a value of R^2 higher than 0.4. 332

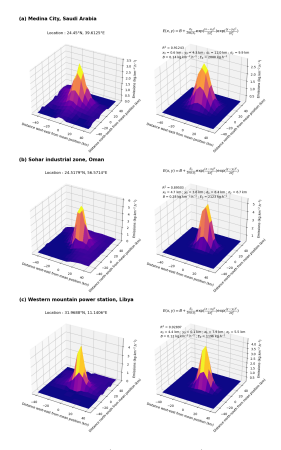


Figure 7: Calculated mean daytime NO_x emissions in 2022 (expressed as NO_2) for different sources (left) and fitted emissions using a 2D-Gaussian function (right) for the city of Medina, Saudi Arabia (a), the Sohar Industrial Zone, Oman (b) and the Western Mountain power plant, Libya (c).

Threshold value	2 Pmolecules.cm $^{-2}$.h $^{-1}$	3 Pmolecules.cm $^{-2}$.h $^{-1}$	4 Pmolecules.cm $^{-2}$.h $^{-1}$
Number of point sources	436	287	163
Point sources with $R^2 > 0.4$	237	179	111
China	23	23	17
India	38	33	23
Russia	29	20	9
United States	17	4	3
Türkiye	8	5	2
Iran	7	8	9
Saudi Arabia	4	6	5
Japan	2	5	4
Australia	3	5	1
Germany	6	1	1
Iraq	6	4	2
Mexico	7	5	3
Kazakhstan	0	6	1

Table 2: Analysis of the number of point sources detected as a function of the threshold applied for cluster detection, and the number of point sources whose fit with a 2D-Gaussian was of acceptable quality ($R^2 > 0.4$). Countries with at least 5 point sources with one of the thresholds are displayed.

As seen with the example of Beijing, moving to a higher threshold can reduce the number of point sources by not 333 including some emitters with lower emissions, but it can also increase the number of detected point sources by 334 reducing the number of pixels corresponding to the cluster and moving certain emitters from the "diffuse source" 335 category to the "point source" category. For example, with a limit of 2 Pmolecules. $cm^{-2}.h^{-1}$, the group of the Ras 336 Laffan power stations in Qatar does not appear as a point source because its emissions are associated to a greater 337 cluster corresponding to a diffuse source which includes the nearby Doha megacity. Conversely, with limits of 3 or 338 4 Pmolecules.cm⁻².h⁻¹, these power plants appear as a point source, and a good quality Gaussian fit provides their 339 total emissions of 1.66 t.h⁻¹, close to the value of 1.86.h⁻¹ reported for the four-year average between 2019 and 2022 340 in Rey-Pommier et al. (2023). Finally, it should be noted that lowering the threshold to 1 Pmolecules. $cm^{-2}.h^{-1}$ also 341 reduces the number of diffuse sources because several nearby urban areas become linked by residual emission zones 342 into a single, larger, diffuse source. Conversely, lowering the threshold detects a very large number of point sources, 343 but many of these additional points are outliers. In the rest of the study, we therefore choose to keep the lowest 344 value of the threshold, i.e. 2 Pmolecules.cm⁻².h⁻¹, to optimise the number of correct emitters we work with. These 345 emitters account for a total output of $2,303 \text{ t.h}^{-1}$ (352 t.h⁻¹ for point sources and 1951 t.h⁻¹ for diffuse sources). 346 This represents about 17% of all emissions with densities higher than 0.2 Pmolecules.cm⁻².h⁻¹ (with a total output 347 of $14,335 \text{ t.h}^{-1}$). As urban areas with more than 1 million inhabitants gather around 16% of the global population 348 (Zimmer et al., 2023), this share of emissions from point and diffuse sources seems consistent with the detection limit of 349 the flux-divergence method using TROPOMI retrievals, as urban areas lower than 1 million inhabitants are generally 350 not detected as diffuse sources here. 351

The full list of the 436 point sources and 323 diffuse sources are given in Supplementary Materials. This list can 352 be compared with the catalog provided by Beirle et al. (2023). Of the 237 point sources for which the Gaussian fit 353 is of correct quality (with $R^2 > 0.4$), 137 also appear in their catalog. For these points, we generally obtain higher 354 emissions (with a median of 441 t.h⁻¹ and an average of 487 t.h⁻¹ in our case, whereas they have a median of 303 t.h⁻¹ 355 and an average of 353 t.h^{-1}). The two datasets have no particular reason to exhibit any clear correlation because they 356 concern different years, and because their approach focused on monthly averages, while ours presents annual averages. 357 For example, a site designated as a point source by Beirle et al. (2023) might not be detected if averaged over a whole 358 year, especially if it stays inactive during certain periods. For instance, their catalog shows 187 occurrences where the 359 signal of NO_x emissions was significant for 6 months out of 12, and 348 occurrences for 5 months. 360

361 3.3 National and regional outputs and comparison with bottom-up emissions

We perform an analysis of emissions at the scale of countries by comparing them to the NO_x emissions provided by EDGARv6.1 for 2018. For our TROPOMI-inferred emissions, we calculate the total mean NO_x output, representing daytime emissions for 2022, for each country using country masks at the $0.0625^{\circ} \times 0.0625^{\circ}$ resolution. To avoid any over-estimation of the total output due to a very high number of pixels with very low emissions, we exclude from the calculation pixels with emission densities below 0.2 Pmolecules.cm⁻².h⁻¹. For emissions in EDGARv6.1, we sum the gridded emissions, representing monthly averages in 2018, for all sectors covered by the inventory and calculate the average flux for the year 2018. The output for each country is calculated using country masks at the $0.1^{\circ} \times 0.1^{\circ}$

resolution. In both cases, we include pixels that directly touch coastlines because marine regions close to the shore 369 receive the spread of anthropogenic emissions due to turbulent diffusion. This can result in over-estimating total 370 emissions for smaller countries, especially those with low emission densities. In order not to account for such outliers, 371 we exclude countries with a population lower than 300,000 inhabitants or with a size lower than $1,000 \text{ km}^2$ from our 372 analysis. This concerns many insular countries in the Caribbean and the Pacific, as well as micro-states like Andorra or 373 Singapore. Overseas territories are considered together with their mainland country. Figure 8 shows the country-wise 374 comparison, covering 165 countries, and Table 3 provides a comparison at the scale of eight different macro-regions: 375 Europe, North America & the Caribbean, South America, the Middle East & North Africa, former USSR countries, 376 Oceania, sub-Saharan Africa and the rest of Asia. For each macro-region, differences are evaluated with the relative 377 bias for the total region, and the mean absolute error (for which each country has the same weight). The use of these 378 different metrics enables to assess the performance of the method on a large scale with respect to an inventory, while 379 simultaneously evaluating its performance on a smaller scale to identify systematic effects that might offset each other 380 at the larger scale. 381

TROPOMI-inferred emissions are generally close to EDGAR estimates for high-income countries, which generally 382 have localized and powerful sources, or countries with a majority of sources located in areas with high observation 383 densities. As a consequence, the macro-regions that perform best with both metrics are Europe, North America & 384 the Caribbean, the Middle East & North Africa, and the rest of Asia. At the scale of countries, TROPOMI-inferred 385 emissions are close to EDGAR estimates for the three largest emitting nations, i.e. China, the United States and India, 386 with TROPOMI-inferred emissions 6, 14 and 4% lower than EDGAR estimates respectively. These three countries 387 account for 45% of global estimated emissions. However, for the fourth highest emitting country, Russia, we estimate 388 emissions 52% higher than in EDGAR. This difference can be due to the low density of observations for major emitters 389 in Russia, leading to the estimation of monthly emissions on the basis of only a few estimates. To illustrate this, the 390 monthly emissions of the two largest Russian cities, Moscow and Saint Petersburg, are studied in the Supplementary 391 Materials. In extreme cases, such key emitters can have no estimates at all for months, making the calculation of 392 the annual average representative of only a part of the year, even when its order of magnitude is correct. Generally 393 speaking, large differences between our top-down estimates and EDGAR emissions are found for many countries that 394 also have low observation densities for this reason. Without prior knowledge of the annual emission profiles in these 395 countries, these biases cannot be corrected, leading to a systematic mis-estimation of total emissions. 396

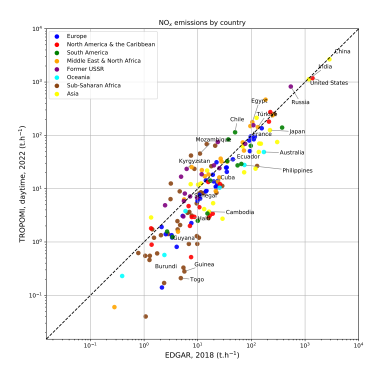


Figure 8: Comparison between TROPOMI-inferred daytime NO_x emissions for 2022 (expressed as NO_2) and mean emissions from EDGARv6.1 in 2018 for all countries, classified by macro-regions.

The other countries for which the difference between our TROPOMI-inferred emissions and EDGAR estimates is significant are low-income countries. For such countries, it is possible that most sources are too small to be detected

with our method, resulting in an under-estimation of emissions. Countries for which TROPOMI-inferred estimates 300 are lower than EDGAR estimates by more than an order of magnitude are Guinea-Bissau, Equatorial Guinea, Togo, 400 Guinea, Gabon, Montenegro, El Salvador, Liberia, Ivory Coast and Myanmar. Of all these countries, only Montene-401 gro and Gabon are not considered low-income countries. Conversely, although no country has TROPOMI-inferred 402 estimates higher than EDGAR estimates by more than an order of magnitude, notable biases exist. In this respect, 403 largest differences are found in central Asia (ratios of 3.8 for Kyrgyzstan, 3.2 for Uzbekistan, 3.4 for Tajikistan), in 404 central and southern Africa (ratios of 5.6 for Zambia, 4.7 for Democratic Republic of Congo, 4.0 for Eswatini, 4.1 for 405 Mozambique, 3.0 for Angola), and Yemen (ratio of 3.2). For these countries, it is also possible that the corresponding 406 EDGAR estimates are imprecise, due to the incomplete or outdated nature of the reported sources in these coun-407 tries. The presence of many sub-Saharan African countries with extreme differences between TROPOMI-inferred and 408 EDGAR estimates explains why the macro-region has the highest mean absolute error despite having the lowest total 409 relative bias. 410

At the global scale, our TROPOMI-inferred daytime emissions for all considered countries (i.e., excluding emissions 411 which take place at sea and smaller countries) reach a total value of $11,168 \text{ t.h}^{-1}$. This value is consistent with that 412 of EDGAR at 12,254 t.h⁻¹, i.e around 107 Mt per year, and lower than the value of 123 Mt calculated by Stocker 413 (2014) for global anthropogenic emissions in 2000 (which include shipping and aircraft emissions). If the lower value 414 can be interpreted as a reduction of NO_x emissions between the two dates, it is also possible that our emissions are 415 under-estimated due to biased-low columns in the TROPOMI NO₂ operational product (Verhoelst et al., 2021). We 416 detail this uncertainty in Section 4.2. We should also note that our TROPOMI-inferred emissions only represent 417 daytime emissions taken around 13:30 LT for each pixel, which are generally lower during mid-day than other times 418 of the day, when pollution peaks in the early morning and late afternoon are reported for traffic in most cities (Menut 419 et al., 2012; Goldberg et al., 2019). For the power sector, emissions at 13:30 are generally similar to the daily mean for 420 power plants used for electricity baseload, but for power plants whose purpose is to meet peak demand, the mid-day 421 emissions can largely differ from the daily mean. For other sectors such as cement, it is difficult to assess whether 422 mid-day emissions are higher than the daily average, since cement production can be driven by factors that are more 423 irregular than those driving power generation or traffic. 424

Region	TROPOMI 2022 (t.h ⁻¹)	$\begin{array}{c} \mathbf{EDGAR} \\ 2018 \ (\mathbf{t}.\mathbf{h}^{-1}) \end{array}$	Relative bias VS EDGAR (weighted average)	Mean absolute error VS EDGAR (unweighted average)	
Sub-Saharan Africa	660	712	-7.4 %	95.0%	
Rest of Asia	4584	5482	-16.4 %	54.4%	
Europe	830	1092	-24.0 %	40.3%	
Middle East & North Africa	1531	1125	36.1 %	52.1%	
North America & the Caribbean	1690	1944	-13.1 %	48.8%	
Oceania	92	282	-67.5 %	62.9%	
South America	514	762	-32.5 %	61.8%	
Former USSR	1268	856	48.2 %	77.4%	
Total	11168	12254	-8.8 %	64.7%	

Table 3: Comparison between TROPOMI-inferred daytime NO_x emissions for 2022 (expressed as NO_2) and mean emissions from EDGARv6.1 in 2018 for macro-regions. For each macro-region, the relative bias between total TROPOMI-inferred emissions and total EDGAR emissions is calculated. The mean absolute bias for all countries of these macro-regions is also calculated.

A source of underestimation can also come from the threshold used to filter out emissions. Here, the limit used of 425 0.2 Pmolecules.cm⁻².h⁻¹ makes it possible to eliminate residual emissions that are difficult to attribute to a source. 426 This filtering also eliminates pixels with negative emissions that are physically impossible. Nevertheless, as negative 427 emissions may represent NO_x incorrectly distributed spatially in the transport term due to errors in the wind field, 428 calculating the sum of emissions without the use of thresholds may be important for identifying countries and regions 429 where the flux-divergence method is limited. In this case, total emissions reach 14,835 t.h⁻¹, which corresponds to an 430 increase of 32.8% compared to the total with the application of the threshold. This estimate is therefore higher than 431 the total EDGAR budget. The differences between the two estimates vary greatly by macro-region: it rises to 149.6% 432 for sub-Saharan Africa, 126.7% for Oceania and 95.5% for South America. The increase is moderate in the Middle 433 East & North Africa region and the North America & the Caribbean, (41.7% and 33.3% respectively). The difference 434 between the two estimates is the lowest in former USSR countries, Europe and the rest of Asia (increases of 16.4%, 435 14.1% and 12.1% respectively). The trends observed previously regarding the reasons for the discrepancy between the 436 TROPOMI-inferred estimates and EDGAR remain unchanged. 437

⁴³⁸ 3.4 Temporal distribution and averaging size

The results presented so far concerned daytime emissions averaged on the entire year 2022. They therefore show a 439 certain potential for mapping the sources of pollution, quantifying the corresponding emissions and characterising 440 their type (by size and country or region). Several studies have shown the possibility to characterise a weekly cycle 441 of NO_x emissions (Stavrakou et al., 2020; Rey-Pommier et al., 2022). The use of geostationary satellites, such as 442 the Geostationnary Environment Monitoring Spectrometer (GEMS) in East Asia (Kim et al., 2020), the Tropopheric 443 Emissions Monitoring of Pollution (TEMPO) in North America (Zoogman et al., 2017) and Sentinel-4 (planned in 444 June 2025) in Europe (Gulde et al., 2017), could also be used to characterise the daily cycle of emissions, leading to 445 a significant improvement of forecasting capabilities. In our case, TROPOMI can only monitor pollution on a daily 446 basis provided that retrievals are of high quality, and the analyses presented so far could theoretically be carried 447 out at this temporal resolution. In the Supplementary Materials, we monitor the daily emissions of the Zaporizhia 448 thermal power plant in Ukraine, whose activity was altered following the ongoing conflict in the country that started in 449 February 2022. However, this type of monitoring remains rare and is more indicative of order-of-magnitude variations 450 rather than precise emission estimates. In practice, the high sensitivity of the method to wind direction and the low 451 signal-to-noise ratio around sources at high latitudes leads to daily emission maps that are very noisy in most cases, 452 making it difficult to precisely monitor activity at this temporal resolution. In general, averaging is therefore required 453 to reduce noise effects and limit the uncertainties associated to emission estimates. Here, we try to evaluate what 454 level of averaging is necessary to limit noise effects and allow a monitoring of emissions. To this end, we consider 455 the average daily emissions obtained for 2022 (i.e. over a maximum of 52 weeks) to be the most accurate estimate of 456 daytime emissions. We compare this maximum averaging value with averages based on a smaller number of estimates. 457 We compare the emissions of various emitters, calculated with an averaging period of 12, 24, 36 and 48 weeks. Figure 458 9 shows the results for eight urban areas, but with different latitudes, populations, levels of development and energy 459 mixes: Ankara (Turkey), Cape Town (South Africa), Madrid (Spain), Portland (Oregon, United States), Chaguanas 460 (Trinidad and Tobago), Saint Petersburg (Russia), Manila (Philippines) and Muscat (Oman). Portland and Manila 461 are urban areas classified as a point sources. Figure 10 shows the results for six industrial facilities, which are all point 462 sources located in Egypt, Australia, Mexico, Chile, India and Germany. The sources were chosen for their relative 463 isolation from other emitters. Calculated emissions correspond to the sum of pixels around the source with densities 464 greater than 2 Pmolecules.cm⁻².h⁻¹. There are two pitfalls to be avoided in this comparison: 465

- The first pitfall would be not to account for the seasonal cycle of emissions, which is very pronounced in some cases, and to compare chronological averages. For example, comparing the first 12 weeks of the year with the first 24 weeks of the same year would not make sense in terms of the difference with emissions averaged over the whole year, because in the first case, emissions would essentially be calculated in boreal winter, whereas in the second case, emissions would be included during spring and summer. To avoid this seasonal bias, emissions averaged over 12 weeks correspond to an average over the first week of each of the 12 months of 2022, emissions averaged over 24 weeks correspond to the first two weeks of these same 12 months, and so on.

- The second pitfall would be not to account for the weekly cycle of emissions. NO_x emissions are generally lower 473 at weekends due to a reduction in human activity in most areas (i.e. on Saturday and Sunday, or Friday and 474 Saturday in most Arabian and North-African countries). It is therefore necessary to ensure that the proportion of 475 weekend days and weekdays in each of the averages calculated remains the same, hence the interest in averaging 476 by weeks (these proportions are therefore 2/7 and 5/7 respectively). We also carry out a fifth set of averaging 477 over 24 days, i.e. 2 days per month. Since the seasonal effect (first pitfall) is generally stronger than the weekly 478 bias (second pitfall), we therefore choose to retain the principle of selecting the same number of days in each 479 month, even if it means making comparisons between averages where the weekend and weekday rates differ from 480 2/7 and 5/7. This last averaging set will be indicated as "irregular". 481

482 In the case of urban areas, the different averages uniformly distributed over time show a similarity in the emissions calculated over the time horizons for Ankara, Muscat, Cape Town, and Madrid. For these cities, the low cloud cover 483 allows a high density of observations and optimal averaging. The 84-day averaging, and to some extent the 24-day 484 irregular averaging, seems sufficient for monitoring emissions. This is not the case for the other urban areas studied, 485 for which the observation density is lower, such as Manila, Saint Petersburg, and Chaguanas. For these cities, a 486 monitoring performed with an averaging below 168 days (or even 252 days in the case of Manila) is therefore limited 487 by noise effects. The limit-case is Portland, which has the larger difference between 84- and 336-days averagings. This 488 is due to a limited number of observations over a small urban area which are not compensated by high-emissions like 489 other point sources shown on Figure 10. For those point sources, similar emissions are observed after an 168-day 490 averaging. In some cases, a 24-day averaging is also sufficient, while in others it is not. The representativeness of 491

492 emissions on such a low level of averaging should be considered with caution, as emissions from industrial plants

⁴⁹³ are always more irregular than those from cities, with the exception of power stations used for baseload electricity

⁴⁹⁴ generation. The averages over 84 days presented here represent emissions that include several days of activity and

⁴⁹⁵ several moments of inactivity.

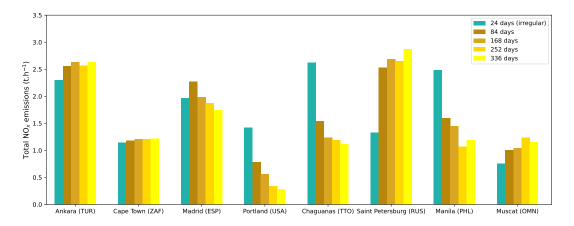


Figure 9: NO_x emissions for 8 different urban areas (diffuse sources), averaged over a period of 24, 84, 168, 252 and 336 days, evenly distributed throughout the year. The proportion of weekend days and weekdays is identical in all the averaging sets except the first one of 24 days. Masses are expressed as NO_2 .

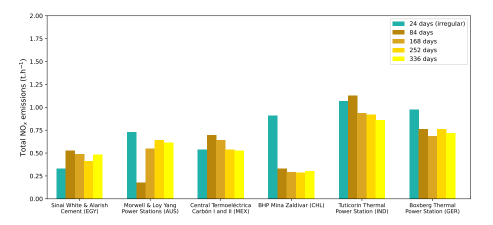


Figure 10: NO_x emissions for 8 different industrial facilities (diffuse sources), averaged over a period of 24, 84, 168, 252 and 336 days, evenly distributed throughout the year. The proportion of weekend days and weekdays is identical in all the averaging sets except for the first set of 24 days. Masses are expressed as NO_2 .

Overall, this analysis seems to indicate that tracking emissions from point sources or diffuse sources using the flux-divergence method requires an averaging effort to limit the noise obtained in the daily emissions. This averaging effort, which is made more difficult for smaller sources, increases with the density of observations, is of about a month in countries with frequent high-quality observations, but of about a year quarter in regions with low observation densities, such as tropical regions and high-latitude regions.

⁵⁰¹ 4 Uncertainties and assessment of results

502 4.1 Model uncertainties

⁵⁰³ Our top-down emissions are calculated here using a flux-divergence model, based on a simplified calculation of a ⁵⁰⁴ transport term, a sink term and a conversion factor from NO_2 to NO_x . This simplicity reduces the computation time ⁵⁰⁵ to calculate emissions and the dependence on external datasets, at the cost of increased model uncertainties. Here, ⁵⁰⁶ although a "topography-wind" term has been introduced in this article to refine the transport term, the sink term ⁵⁰⁷ remains simple and only represents the reaction between NO_2 and OH. While this reaction is the first contributor of

 NO_x loss, other sinks may be significant. For instance, organic peroxy radicals can oxidise NO_x to form peroxy nitrates, 508 making the corresponding sink important in the presence of VOCs (Stavrakou et al., 2013), especially in biomass fires. 509 In different conditions, the formation of peroxyacetyl nitrate from NO_2 (Moxim et al., 1996), can also contribute to a 510 significant share of the NO_x loss. The vertical averaging of is also made simple here, and alhough the sink term varies 511 little with the thickness of the layer within which the temperature and OH concentration are calculated (Rey-Pommier 512 et al., 2022), this assumes the OH field is correctly represented. This assumption may be wrong if large NO_x emitters 513 are not taken into account, as this would distort the corresponding NO_2 field and the subsequent OH field. A possible 514 improvement to our dataset could be to compare the columns calculated from the TROPOMI observations with the 515 NO₂ column represented by CAMS and correct outliers detected from this comparison. Another refined version could 516 infer directly the effective NO_x lifetime from the NO_2 observations themselves, as suggested by Laughner and Cohen 517 (2019).518

Another model uncertainty comes from the calculation of the conversion of NO_2 production to total NO_x . The 519 majority of NO_x is emitted in the form of NO, which is not observed from space. A common assumption is that NO 520 is rapidly transformed into NO_2 through its reaction with ozone, reaching a stationary state within a few minutes. 521 Numerous studies (Beirle et al., 2019; de Foy and Schauer, 2022) assumed a photostationary state in typical urban 522 conditions and used a ratio of 1.32 based on Seinfeld and Pandis (2006). Here, the values of this ratio calculated 523 from CAMS data did not differ much from this value. However, the photostationary state is a hypothesis which is 524 potentially not verified on the scale of a NO_x source like a power plant stack. Li et al. (2023b) calculated values of 525 this conversion ratio correlated with the combustion temperature and energy efficiency for sources in China that are 526 highly intensive in energy such as power plants, and found a median value of 3.3. Biases in the calculation of the 527 $NO_x:NO_2$ ratio can also arise in highly polluted environments, in which the Leighton relationship used to calculate 528 this ratio is no longer valid. In particular, OH can also react with VOCs and form oxygenated VOCs. Further studies 529 estimating this ratio at various spatial and temporal scales would thus provide a better implementation of our model. 530

⁵³¹ 4.2 Data uncertainties

The NO₂ column densities are the main input quantity in our estimation of NO_x emissions, making the its calculation 532 within the TROPOMI product the first element to examine when considering the data uncertainties in our estimates. 533 Columns are calculated from measurements of solar backscattered radiation and comparison with a specific UV-Visible 534 band using the Differential Optical Absorption Spectroscopy method, before being assimilated to derive a tropospheric 535 vertical component. The corresponding uncertainty under polluted conditions is dominated by the sensitivity of 536 satellite observations to air masses near the ground, and is expressed through the calculation of the tropospheric air-537 mass factor (AMF). To assess the significance of such effects, vertical profiles within the TROPOMI product can be 538 replaced by any other profile information, resulting in a new retrieved tropospheric NO_2 column. Douros et al. (2023) 539 replaced the *a priori* TROPOMI OFFL NO₂ profile by high-resolution air quality forecasts for Europe. As compared 540 to the standard TROPOMI NO₂ data, this new product was found to be biased-low by 5% to 12% for most European 541 cities. The air mass factor (AMF) itself can be replaced: for instance, Lama et al. (2022) re-calculated the AMF 542 by replacing the tropospheric AMF of the original TROPOMI OFFL product by an AMF taken from WRF-Chem 543 simulations. Similarly, Beirle et al. (2023) re-calculated the AMF above different emitters from the corresponding 544 averaging kernel based on a peak profile at plume height to better reflect the distribution of NO_2 close to ground, 545 which resulted in an AMF correction of about 1.61. Here, we did not perform any of such corrections, and we consider 546 a relative uncertainty for the column of 30% (Boersma et al., 2004) for pixels corresponding to non-urbanized areas. 547 For pixel corresponding to cities, S-5P validation activities which indicate that TROPOMI tropospheric NO₂ columns 548 are systematically biased low by higher rates (Verhoelst et al., 2021), and a higher relative uncertainty of 50% is used. 549 Such biases seem to run counter to our comparison with the catalog by Beirle et al. (2023), for which this change in 550 sensitivity was performed but leading to emissions generally lower than ours. A more detailed analysis of the concerned 551 emitters seems necessary to better understand the parameters that have the largest impact on the vertical sensitivity 552 of TROPOMI retrievals and our inversion model. 553

Other data uncertainties can arise from other parameters that play a crucial role in the estimation of advection 554 and chemistry effects. An accurate representation of the wind is critical to estimate the transport term correctly. 555 For a given plume, the poor representation of wind speed leads to an under-or over-estimation of transport, but the 556 correct orientation of positive and negative values around the source remains. However, an incorrect representation 557 of the wind direction, such as a non-alignment with the main direction of the plume, fails to represent a correct 558 orientation of positive and negative values. The estimation of the transport term significantly thus relies heavily on 559 the representation of the wind angle. Higher errors are therefore expected to be high in regions having winds that 560 vary rapidly in time, or regions with complex horizontal wind variations, such as mountainous regions. In particular, 561

situations where sub-grid scale-phenomena occur, not accounted for in ERA5 wind fields, might display even higher 562 errors in the estimation of transported NO_x. For instance, Tehran, Iran, has an extremely complex topography, 563 and in the calculated emissions, the transport term is particularly high compared with the sink term, with high and 564 unrealistic negative values on large scales around the Tochal mountain immediately to the north of the city. Other 565 megacities such as Seoul, South Korea, Jeddah, Saudi Arabia, Chittagong, Bangladesh, also exhibit unrealistically 566 high values for the transport term. Besides, these values are not compensated by the topography-wind term, for which 567 an inverse scale height of $X_e = 0.3 \text{ km}^{-1}$ is used based on Sun (2022). For this term to be sufficient to compensate 568 for the negative values observed, a higher inverse scale height would be required. Such observation is consistent with 569 Beirle et al. (2023), who used empirical values of X_e up to 2 km⁻¹ and reduced the amplitude of the negative patterns 570 observed for Los Angeles, United States, Tehran, Iran, or Seoul, South Korea. Finally, errors in the estimation of 571 emissions can also come from a wrong estimation of the air composition when calculating the sink term. The NO_2 572 lifetime relies heavily on the representation of the OH concentration field, which varies with NO_x itself through a 573 non-linear mechanism. An incorrect representation of the sink term can occur at the scale of a plume by not capturing 574 this relationship due to an incorrect knowledge of emitters on the ground. This can also be due to the $0.4^{\circ} \times 0.4^{\circ}$ 575 resolution of CAMS that do not always capture the NO_2 gradients adequately in plumes near a known emitter (Valin 576 et al., 2011; Li et al., 2023a). For the OH concentration, a relative uncertainty of 30% has been used (Huijnen et al., 577 2019), representing the largest component of absolute uncertainty apart from the vertical columns. Large errors in 578 the annual cycle of OH, and therefore in the sink term, can thus be expected. As a consequence, a wrong estimation 579 of wind angle and OH concentration can lead to unrealistically high emissions, or even negative emissions. 580

581 5 Conclusion

In this study, we present a global quantification of NO_x emissions by performing a mass-balance inversion based on 582 the flux-divergence method, based. This approach offers a rapid alternative to traditional 3D inversion methods using 583 Chemical Transport Models. The foundation of this method lies in the observation of tropospheric vertical column 584 densities of NO₂ provided by TROPOMI. Our methodology incorporates several components in the calculation of 585 emissions: a transport term driven by horizontal wind, a sink term largely driven by OH concentrations, and a 586 topography-wind correction term. The emissions calculated represent mean daytime fluxes for the year 2022, allowing 587 us to map emissions on a global scale. The results highlight that the primary sources of NO_x emissions are industrialized 588 and developing countries. Our emission estimates are consistent with global estimates, as well as the EDGARv6.1 589 inventory, though notable discrepancies are observed at the national level, particularly in former USSR countries 590 and sub-Saharan Africa. Besides, we performed a pinpointing of emitters by distinguishing between diffuse sources, 591 typically large metropolitan areas with extensive spatial distribution, and point sources, generally isolated industrial 592 facilities with emissions that often exhibit a Gaussian spread. 436 diffuse sources and 323 point sources are identified. 593 Significant uncertainties remain, especially in regions where OH is not the only source of NO_x removal, regions where 594 wind representation is inaccurate, and regions where TROPOMI data exhibit substantial biases. Nonetheless, this 595 work demonstrates the feasibility of annual NO_x emission monitoring with reduced latency and fewer mis-allocation 596 issues compared to traditional inventories. Our approach enables the monitoring of emissions at the monthly scale in 597 regions with high observation densities, that usually correspond to dry, mid-latitude countries. Conversely, the effect 598 of numerical noise, combined with low-observation densities, restricts such monitoring to a higher averaging period of 599 up to months, generally in tropical and high-latitude regions. Efforts should be made to further develop this method 600 to provide a near-real time monitoring tool a higher temporal resolution for these regions. The results of this study 601 were obtained from the calculation of daily NO_x emissions in 2022 and their annual average. 602

6 Data availability

The monthly NO_x emission maps can be accessed at https://doi.org/10.5281/zenodo.13758447 (Rey-Pommier et al., 2025). Data is made available as emission grid maps as .nc files with emissions expressed in petamolecules per square centimetre per hour (Pmolecules.cm⁻².h⁻¹). Conversion factors to mass terms (expressed as NO₂, NO or N) are included. The lists of diffuse and point sources are also provided.

Author contributions. AR analysed the data, prepared the main software code and wrote the paper. AH improved some aspects of the code and prepared the code for Gaussian fitting. FC, PC, TC, JK and JS contributed to the improvement of the method and the interpretation of the results. All the authors read and agreed on the published ⁶¹¹ version of the paper.

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619 References

- Beirle, S., Borger, C., Dörner, S., Li, A., Hu, Z., Liu, F., Wang, Y., and Wagner, T. (2019). Pinpointing nitrogen oxide emissions from space, sci. adv., 5, eaax9800.
- Beirle, S., Borger, C., Jost, A., and Wagner, T. (2023). Improved catalog of NOx point source emissions (version 2).
 Earth System Science Data, 15(7):3051–3073.
- Benkovitz, C. M., Scholtz, M. T., Pacyna, J., Tarrasón, L., Dignon, J., Voldner, E. C., Spiro, P. A., Logan, J. A.,
 and Graedel, T. (1996). Global gridded inventories of anthropogenic emissions of sulfur and nitrogen. *Journal of Geophysical Research: Atmospheres*, 101(D22):29239–29253.
- Boersma, K., Eskes, H., and Brinksma, E. (2004). Error analysis for tropospheric NO2 retrieval from space. Journal
 of Geophysical Research: Atmospheres, 109(D4).
- Burkholder, J., Sander, S., Abbatt, J., Barker, J., Cappa, C., Crounse, J., Dibble, T., Huie, R., Kolb, C., Kurylo,
 M., et al. (2020). Chemical kinetics and photochemical data for use in atmospheric studies; evaluation number 19.
 Technical report, Pasadena, CA: Jet Propulsion Laboratory, National Aeronautics and Space
- ⁶³¹ Technical report, Pasadena, CA: Jet Propulsion Laboratory, National Aeronautics and Space . .
- Castellanos, P., Boersma, K., and Van Der Werf, G. (2014). Satellite observations indicate substantial spatiotemporal
 variability in biomass burning NOx emission factors for South America. Atmospheric Chemistry and Physics,
 14(8):3929–3943.
- Crippa, M., Guizzardi, D., Muntean, M., Schaaf, E., Solazzo, E., Monforti-Ferrario, F., Olivier, J., and Vignati, E.
 (2020). Fossil CO2 emissions of all world countries. *Luxembourg: European Commission*, pages 1–244.
- Cusworth, D. H., Thorpe, A. K., Miller, C. E., Ayasse, A. K., Jiorle, R., Duren, R. M., Nassar, R., Mastrogiacomo,
 J.-P., and Nelson, R. R. (2023). Two years of satellite-based carbon dioxide emission quantification at the world's
 largest coal-fired power plants. Atmospheric Chemistry and Physics, 23(22):14577–14591.
- Danielson, J. J. and Gesch, D. B. (2011). Global multi-resolution terrain elevation data 2010 (GMTED2010). Technical
 report, US Geological Survey.
- de Foy, B. and Schauer, J. J. (2022). An improved understanding of NOx emissions in south asian megacities using
 TROPOMI NO2 retrievals. *Environmental Research Letters*, 17(2):024006.
- Douros, J., Eskes, H., van Geffen, J., Boersma, K. F., Compernolle, S., Pinardi, G., Blechschmidt, A.-M., Peuch,
 V.-H., Colette, A., and Veefkind, P. (2023). Comparing Sentinel-5P TROPOMI NO2 column observations with the
 CAMS regional air quality ensemble. *Geoscientific Model Development*, 16(2):509–534.
- ⁶⁴⁷ Ehhalt, D. H., Rohrer, F., and Wahner, A. (1992). Sources and distribution of NOx in the upper troposphere at
- northern mid-latitudes. Journal of Geophysical Research: Atmospheres, 97(D4):3725–3738.

- Eskes, H., van Geffen, J., Boersma, F., Eichmann, K., Apituley, A., Pedergnana, M., Sneep, M., Veefkind, J., and
 Loyola, D. (2022). Sentinel-5 Precursor/TROPOMI Level 2 Product User Manual Nitrogen Dioxide.
- Goldberg, D. L., Harkey, M., de Foy, B., Judd, L., Johnson, J., Yarwood, G., and Holloway, T. (2022). Evaluating NOx
 emissions and their effect on O3 production in Texas using TROPOMI NO2 and HCHO. Atmospheric Chemistry
 and Physics, 22(16):10875–10900.
- Goldberg, D. L., Saide, P. E., Lamsal, L. N., de Foy, B., Lu, Z., Woo, J.-H., Kim, Y., Kim, J., Gao, M., Carmichael,
 G., et al. (2019). A top-down assessment using OMI NO2 suggests an underestimate in the nox emissions inventory
 in Seoul, South Korea, during KORUS-AQ. Atmospheric Chemistry and Physics, 19(3):1801–1818.
- Graedel, T., Farrow, L., and Weber, T. (1976). Kinetic studies of the photochemistry of the urban troposphere.
 Atmospheric Environment (1967), 10(12):1095–1116.
- Granier, C., Darras, S., van der Gon, H. D., Jana, D., Elguindi, N., Bo, G., Michael, G., Marc, G., Jalkanen, J.-P.,
 Kuenen, J., et al. (2019). *The Copernicus atmosphere monitoring service global and regional emissions (April 2019 version)*. PhD thesis, Copernicus Atmosphere Monitoring Service.
- Griffin, D., Zhao, X., McLinden, C. A., Boersma, F., Bourassa, A., Dammers, E., Degenstein, D., Eskes, H., Fehr, L.,
 Fioletov, V., et al. (2019). High-resolution mapping of nitrogen dioxide with TROPOMI: First results and validation
 over the canadian oil sands. *Geophysical Research Letters*, 46(2):1049–1060.
- Gulde, S., Kolm, M., Smith, D., Maurer, R., Courrèges-Lacoste, G. B., Sallusti, M., and Bagnasco, G. (2017). Sentinel
 4: A geostationary imaging UVN spectrometer for air quality monitoring: Status of design, performance and
 development. In *International Conference on Space Optics—ICSO 2014*, volume 10563, pages 1158–1166. SPIE.
- Hakkarainen, J., Szeląg, M. E., Ialongo, I., Retscher, C., Oda, T., and Crisp, D. (2021). Analyzing nitrogen oxides to
 carbon dioxide emission ratios from space: A case study of Matimba Power Station in South Africa. Atmospheric
 Environment: X, 10:100110.
- ⁶⁷¹ He, K. (2012). Multi-resolution Emission Inventory for China (MEIC): model framework and 1990-2010 anthropogenic ⁶⁷² emissions. In *AGU fall meeting abstracts*, volume 2012, pages A32B–05.
- Hersbach, H., Bell, B., Berrisford, P., Hirahara, S., Horányi, A., Muñoz-Sabater, J., Nicolas, J., Peubey, C., Radu,
 R., Schepers, D., et al. (2020). The ERA5 global reanalysis. *Quarterly Journal of the Royal Meteorological Society*,
 146(730):1999–2049.
- Hoelzemann, J. J., Schultz, M. G., Brasseur, G. P., Granier, C., and Simon, M. (2004). Global wildland fire emission model (GWEM): Evaluating the use of global area burnt satellite data. *Journal of Geophysical Research:* Atmospheres, 109(D14).
- Huijnen, V., Eskes, H., Wagner, A., Schulz, M., Christophe, Y., Ramonet, M., Basart, S., Benedictow, A.,
 Blechschmidt, A.-M., Chabrillat, S., et al. (2016). Validation report of the CAMS near-real-time global atmospheric composition service: System evolution and performance statistics. Status up to 1 June 2016; https://pure.mpg.de/rest/items/item_2441827/component/file_2441834/content.
- Huijnen, V., Pozzer, A., Arteta, J., Brasseur, G., Bouarar, I., Chabrillat, S., Christophe, Y., Doumbia, T., Flem ming, J., Guth, J., et al. (2019). Quantifying uncertainties due to chemistry modelling-evaluation of tropospheric
 composition simulations in the CAMS model (cycle 43r1). Geoscientific Model Development, 12(4):1725–1752.
- Judd, L. M., Al-Saadi, J. A., Szykman, J. J., Valin, L. C., Janz, S. J., Kowalewski, M. G., Eskes, H. J., Veefkind,
 J. P., Cede, A., Mueller, M., et al. (2020). Evaluating Sentinel-5P TROPOMI tropospheric NO2 column densities
 with airborne and Pandora spectrometers near New York City and Long Island Soun. Atmospheric measurement
 techniques, 13(11):6113–6140.
- Khairoun, A., Mouillot, F., Chen, W., Ciais, P., and Chuvieco, E. (2024). Coarse-resolution burned area datasets
 severely underestimate fire-related forest loss. *Science of the Total Environment*, 920:170599.
- Kim, J., Jeong, U., Ahn, M.-H., Kim, J. H., Park, R. J., Lee, H., Song, C. H., Choi, Y.-S., Lee, K.-H., Yoo, J.-M.,
 et al. (2020). New era of air quality monitoring from space: Geostationary Environment Monitoring Spectrometer
- ⁶⁹⁴ (GEMS). Bulletin of the American Meteorological Society, 101(1):E1–E22.

- Kuenen, J., Dellaert, S., Visschedijk, A., Jalkanen, J.-P., Super, I., and Denier van der Gon, H. (2022). CAMS-REG-
- v4: a state-of-the-art high-resolution european emission inventory for air quality modelling. *Earth System Science Data*, 14(2):491–515.
- Lama, S., Houweling, S., Boersma, K. F., Aben, I., Denier van der Gon, H. A., and Krol, M. C. (2022). Estimation of OH in urban plumes using TROPOMI-inferred NO2/CO. *Atmospheric Chemistry and Physics*, 22(24):16053–16071.

Lama, S., Houweling, S., Boersma, K. F., Eskes, H., Aben, I., Denier van der Gon, H. A., Krol, M. C., Dolman, H.,
 Borsdorff, T., and Lorente, A. (2020). Quantifying burning efficiency in megacities using the NO2/CO ratio from
 the Tropospheric Monitoring Instrument (TROPOMI). Atmospheric Chemistry and Physics, 20(17):10295–10310.

- Latsch, M., Richter, A., and Burrows, J. P. (2023). Improving the detection of global NOx emissions from shipping
 in S5P/TROPOMI data. Technical report, Copernicus Meetings.
- Laughner, J. L. and Cohen, R. C. (2019). Direct observation of changing NOx lifetime in North American cities.
 Science, 366(6466):723-727.
- Li, C., Martin, R. V., Cohen, R. C., Bindle, L., Zhang, D., Chatterjee, D., Weng, H., and Lin, J. (2023a). Variable effects of spatial resolution on modeling of nitrogen oxides. *Atmospheric Chemistry and Physics*, 23(5):3031–3049.
- Li, X., Cohen, J. B., Qin, K., Geng, H., Wu, X., Wu, L., Yang, C., Zhang, R., and Zhang, L. (2023b). Remotely
 sensed and surface measurement-derived mass-conserving inversion of daily NOx emissions and inferred combustion
 technologies in energy-rich northern china. Atmospheric Chemistry and Physics, 23(14):8001–8019.
- Lorente, A., Boersma, K., Eskes, H., Veefkind, J., Van Geffen, J., De Zeeuw, M., Denier Van Der Gon, H., Beirle,
 S., and Krol, M. (2019). Quantification of nitrogen oxides emissions from build-up of pollution over Paris with
 TROPOMI. Scientific reports, 9(1):1–10.
- Mebust, A. K. and Cohen, R. C. (2013). Observations of a seasonal cycle in NOx emissions from fires in african woody
 savannas. *Geophysical research letters*, 40(7):1451–1455.
- Menut, L., Goussebaile, A., Bessagnet, B., Khvorostiyanov, D., and Ung, A. (2012). Impact of realistic hourly emissions
 profiles on air pollutants concentrations modelled with chimere. *Atmospheric environment*, 49:233–244.
- Moxim, W., Levy, H., and Kasibhatla, P. (1996). Simulated global tropospheric PAN: Its transport and impact on
 NOx. Journal of Geophysical Research: Atmospheres, 101(D7):12621–12638.
- Opacka, B., Müller, J.-F., and Stavrakou, T. (2022). Evaluation of soil NO emissions in the tropics using field data
 and TROPOMI NO2 columns. In EGU General Assembly Conference Abstracts, pages EGU22–1287.
- Ossohou, M., Galy-Lacaux, C., Yoboué, V., Hickman, J., Gardrat, E., Adon, M., Darras, S., Laouali, D., Akpo, A.,
 Ouafo, M., et al. (2019). Trends and seasonal variability of atmospheric NO2 and HNO3 concentrations across
 three major african biomes inferred from long-term series of ground-based and satellite measurements. *Atmospheric Environment*, 207:148–166.
- Platt, U., Stutz, J., Platt, U., and Stutz, J. (2008). Differential absorption spectroscopy. Springer.
- Ramo, R., Roteta, E., Bistinas, I., Van Wees, D., Bastarrika, A., Chuvieco, E., and Van der Werf, G. R. (2021).
 African burned area and fire carbon emissions are strongly impacted by small fires undetected by coarse resolution
 satellite data. *Proceedings of the National Academy of Sciences*, 118(9):e2011160118.
- Reuter, M., Buchwitz, M., Schneising, O., Krautwurst, S., O'Dell, C. W., Richter, A., Bovensmann, H., and Burrows, J. P. (2019). Towards monitoring localized CO2 emissions from space: co-located regional CO2 and NO2 enhancements observed by the OCO-2 and S5P satellites. *Atmospheric Chemistry and Physics*, 19(14):9371–9383.
- Rey-Pommier, A., Chevallier, F., Ciais, P., Broquet, G., Christoudias, T., Kushta, J., Hauglustaine, D., and Sciare, J.
 (2022). Quantifying NOx emissions in Egypt using TROPOMI observations. *Atmospheric Chemistry and Physics*, 22(17):11505–11527.
- Rey-Pommier, A., Chevallier, F., Ciais, P., Kushta, J., Christoudias, T., Bayram, I. S., and Sciare, J. (2023). Detecting nitrogen oxide emissions in Qatar and quantifying emission factors of gas-fired power plants-a 4-year study.
- ⁷³⁹ Atmospheric Chemistry and Physics, 23(21):13565–13583.

- Rey-Pommier, A., Héraud, A., Chevallier, F., Ciais, P., Christoudias, T., Kushta, J., and Sciare, J. (2025). Global
 gridded NOx emissions using TROPOMI observations. [Data set].
- Rieß, T. C. V., van Vliet, J., Van Roy, W., de Laat, J., Dammers, E., and Boersma, F. (2023). Aircraft validation
 reveals a 20% low bias in TROPOMI NO2 over sea caused by TM5 a priori profiles. Technical report, Copernicus
 Meetings.
- ⁷⁴⁵ Seinfeld, J. H. and Pandis, S. N. (2006). Atmospheric chemistry and physics from air pollution to climate change.
- Stavrakou, T., Müller, J.-F., Bauwens, M., Boersma, K., and van Geffen, J. (2020). Satellite evidence for changes in
 the NO2 weekly cycle over large cities. *Scientific reports*, 10(1):1–9.
- Stavrakou, T., Müller, J.-F., Boersma, K., Van Der A, R., Kurokawa, J., Ohara, T., and Zhang, Q. (2013). Key chem ical NOx sink uncertainties and how they influence top-down emissions of nitrogen oxides. *Atmospheric Chemistry and Physics*, 13(17):9057–9082.
- ⁷⁵¹ Stocker, T. (2014). Climate change 2013: the physical science basis: Working Group I contribution to the Fifth
 ⁷⁵² assessment report of the Intergovernmental Panel on Climate Change. Cambridge University Press.
- ⁷⁵³ Sun, K. (2022). Derivation of emissions from satellite-observed column amounts and its application to TROPOMI
 ⁷⁵⁴ NO2 and CO observations. *Geophysical Research Letters*, 49(23):e2022GL101102.
- Thornton, J. A., Virts, K. S., Holzworth, R. H., and Mitchell, T. P. (2017). Lightning enhancement over major oceanic
 shipping lanes. *Geophysical Research Letters*, 44(17):9102–9111.
- Valin, L., Russell, A., Hudman, R., and Cohen, R. (2011). Effects of model resolution on the interpretation of satellite
 NO2 observations. Atmospheric Chemistry and Physics, 11(22):11647-11655.
- Van Geffen, J., Eskes, H., Compernolle, S., Pinardi, G., Verhoelst, T., Lambert, J.-C., Sneep, M., Ter Linden, M.,
 Ludewig, A., Boersma, K. F., et al. (2022). Sentinel-5P TROPOMI NO2 retrieval: impact of version v2.2 improvements and comparisons with OMI and ground-based data. *Atmospheric Measurement Techniques*, 15(7):2037–2060.
- Vandaele, A. C., Hermans, C., Simon, P. C., Carleer, M., Colin, R., Fally, S., Merienne, M.-F., Jenouvrier, A., and
 Coquart, B. (1998). Measurements of the NO2 absorption cross-section from 42 000 cm-1 to 10 000 cm-1 (238–1000 nm) at 220 K and 294 K. Journal of Quantitative Spectroscopy and Radiative Transfer, 59(3-5):171–184.
- Veefkind, J. P., Aben, I., McMullan, K., Förster, H., De Vries, J., Otter, G., Claas, J., Eskes, H., De Haan, J.,
 Kleipool, Q., et al. (2012). TROPOMI on the ESA Sentinel-5 precursor: A GMES mission for global observations of
 the atmospheric composition for climate, air quality and ozone layer applications. *Remote sensing of environment*,
 120:70–83.
- Verhoelst, T., Compernolle, S., Pinardi, G., Lambert, J.-C., Eskes, H. J., Eichmann, K.-U., Fjæraa, A. M., Granville,
 J., Niemeijer, S., Cede, A., et al. (2021). Ground-based validation of the Copernicus Sentinel-5p TROPOMI NO2
 measurements with the NDACC ZSL-DOAS, MAX-DOAS and Pandonia global networks. Atmospheric Measurement
 Techniques, 14(1):481–510.
- Yienger, J. and Levy, H. (1995). Empirical model of global soil-biogenic NOx emissions. Journal of Geophysical
 Research: Atmospheres, 100(D6):11447–11464.
- Zhai, S., Jacob, D. J., Franco, B., Clarisse, L., Coheur, P., Shah, V., Bates, K. H., Lin, H., Dang, R., Sulprizio, M. P.,
 et al. (2024). Transpacific transport of asian peroxyacetyl nitrate (PAN) observed from satellite: Implications for
 ozone. *Environmental Science & Technology*.
- Zimmer, A., C. T. N. B. A. G., J. Giezendanner. Palisades, N. N. S. D., and (SEDAC)., A. C. (2023). Global Urban
 Demography Dataset (GUDD) (Preliminary Release). https://doi.org/10.7927/4rjm-vf23.
- Zoogman, P., Liu, X., Suleiman, R., Pennington, W., Flittner, D., Al-Saadi, J., Hilton, B., Nicks, D., Newchurch,
 M., Carr, J., et al. (2017). Tropospheric emissions: Monitoring of pollution (tempo). Journal of Quantitative
 Spectroscopy and Radiative Transfer, 186:17–39.

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