Mobile DOAS Observations of Tropospheric NO₂ Using an UltraLight Trike and Flux Calculation

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Abstract: In this study, we report on airborne Differential Optical Absorption Spectroscopy (DOAS) observations of tropospheric NO₂ using an Ultralight Trike (ULT) and associated flux calculations. The instrument onboard the ULT was developed for measuring the tropospheric NO₂ Vertical Column Density (VCD) and it was operated for several days between 2011 and 2014, in the South-East of Romania. Collocated measurements were performed using a car-DOAS instrument. Most of the airborne and mobile ground-based measurements were performed close to an industrial platform located nearby Galati city (45.43° N, 28.03° E). We found a correlation of R = 0.71 between tropospheric NO₂ VCDs deduced from airborne DOAS observations and mobile ground-based DOAS observations. We also present a comparison between stratospheric NO₂ Slant Column Density (SCD) derived from the Dutch OMI NO₂ (DOMINO) satellite data product and stratospheric SCDs obtained from ground and airborne measurements. The airborne DOAS observations performed on 13 August 2014 were used to quantify the NO₂ flux originating from an industrial platform located nearby Galati city. Measurements during a flight above the industrial plume showed a maximum tropospheric NO₂ VCD of (1.41 ± 0.27) × 10¹⁶ molecules/cm² and an associated NO₂ flux of (3.45 ± 0.89) × 10⁻³ kg/s.

Keywords: mobile DOAS; airborne observations; nitrogen dioxide; emission flux

1. Introduction

Nitrogen dioxide (NO₂) is a chemical gaseous compound with an important role in the Earth’s atmosphere. NO₂ is a key trace element in the chemistry of ozone, since it is involved in the catalytic destruction of ozone in the stratosphere [1], while in the troposphere its photolysis leads directly to the formation of ozone (O₃) in the presence of VOCs (volatile organic compounds). NO₂ is released in the atmosphere from natural sources (soil, lightning, solar cosmic rays) and anthropogenic emissions (fossil fuels and biomass burning, industrial activities). Long-term exposure to NO₂ may affect the respiratory system and lead to coronary diseases. NO₂ can lead to acidification of the aquatic ecosystem following the oxidation to HNO₃.

The Differential Optical Absorption Spectroscopy (DOAS) technique [2] has been used for NO₂ atmospheric measurements since the early 1970s [3,4]. Nowadays, besides ground-based zenith sky measurements, DOAS techniques have developed into Multi-Axis Differential Optical Absorption Spectroscopy (MAX-DOAS) observations [5]. The mobile DOAS technique was recently used on...
several platforms such as: cars [6,7], airplanes [8–10], Unmanned Aerial Vehicles (UAVs) [11] or satellites [12–16].

Airborne observations have a number of important advantages for atmospheric research such as: the flexibility during the flights and the possibility to access remote areas such as oceans, deserts, rural or areas without roads.

The ULMs (Ultralight Motorized) are airborne platforms with an important scientific potential for atmospheric research. So far, ULMs have been used to study the ultraviolet actinic radiation flux [17], formaldehyde distribution [18], aerosol profiles [19], SO₂, NO₂ and ozone distribution [20–23].

This work highlights the capability of a low-cost system (ULT-DOAS) used for measurements of tropospheric NO₂ VCD and associated flux calculations. This study presents airborne DOAS observations of tropospheric NO₂ using an Ultralight Trike (ULT) and associated flux calculations. The work presented here was motivated by the need to further assess the intrapixel variability of NO₂ detected by UV-VIS DOAS instruments onboard satellites. This work comes in the context of a validation programme of the future Atmospheric Sentinels, starting with the Sentinel-5 Precursor to be launched in summer 2017. A similar system based on DOAS onboard the ULM was used during the Airborne Romanian Measurements of Aerosols and Trace gases (AROMAT) campaign, held in Romania in August 2015 [24]. The AROMAT campaign was conducted under the aegis of the European Space Agency (ESA) in the framework of a series of ESA field campaigns.

2. Methodology

2.1. Experimental and Instrumental Descriptions

The mobile DOAS observations were performed onboard of an Ultralight Trike (ULT), in the South-East of Romania (Figure 1) during several days between 2011 and 2014. All measurements were performed under clear-sky conditions (see Table 1). The Mobile DOAS system used for measurements will be described in the following as the ULT-DOAS system. The measurements took place in an area around Galati (located at 45.43° N, 28.03° E), Braila (45.26° N, 27.95° E), and close to the industrial areas of Slobozia (44.56° N, 27.35° E). Note that an operational steel mill factory is located in the vicinity of Galati, while Slobozia was chosen due to the presence of a fertilizer factory. Due to security concerns, direct flights above the cities or industrial platforms were not performed. Most of the airborne DOAS observations were performed in nadir geometry. Details about the ULT-DOAS measurements are presented in Table 1.

![Figure 1](image.png)

Figure 1. The main locations where airborne and/or ground-based mobile Differential Optical Absorption Spectroscopy (DOAS) observations were performed.
Airborne-DOAS measurements were accompanied by car-DOAS measurements and static twilight observations on 21 July 2014. Static twilight DOAS measurements, used for determination of the NO$_2$ content from the reference spectra, were performed in a rural area close to Galati city. The car-DOAS measurements were performed right before or after the experimental flights.

The aircraft used for all the flight experiments presented in this paper is a double-seated, open-cockpit ultralight aircraft, trike type (model Fanagoria 21, produced by Plovdiv Air Bulgaria). The flexible wing (Atlant-21, produced by Plovdiv Air Bulgaria) has an area of 16 m$^2$. The ULT is powered by a Subaru EA81 engine with 75 HP. The cruise speed is 75 km/h and the maximum speed is 100 km/h relative to the ground. The aircraft has a maximum total weight at take-off of 450 kg.

The ULT-DOAS instrument consists of a compact Czerny-Turner spectrometer (AvaSpec-ULS2048XL-USB2, of 175 × 110 × 44 mm dimensions and 855 g weight) placed in the Ultralight Trike. Figure 2 presents the instrumental DOAS set-up. The spectral range of the spectrometer is 280–550 nm with 0.7 nm resolution (FWHM—Full Width at Half Maximum) with a focal length of 75 mm. The entrance slit is 50 µm and the grating is 1200 L/mm, blazed at 250 nm. The spectrometer is connected to the telescope through a 400 µm chrome-plated brass optical fiber. The telescope achieves a 2.3° field-of-view with fused silica collimating lenses. Each spectrum is recorded by a laptop and georeferenced by a GPS receiver. The spectrometer and the GPS receiver are powered by the laptop USB ports. The entire set-up is powered by 12 V of the ULT through an inverter. Each measurement is a 10-second average of 10 scans accumulations at an integration time between 50 and 150 ms.

This work is mostly based on nadir-DOAS observations but we also present zenith-sky observations onboard ULT for stratospheric NO$_2$ measurements. The same DOAS system was used in the case of the zenith sky car-DOAS observations.

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**Table 1.** Coordinates and temporal coverage of the mobile DOAS measurements.

| Day                  | Time Interval UTC * | Route of ULT-DOAS ** Measurements | NO$_2$ Source Target     |
|----------------------|---------------------|-----------------------------------|--------------------------|
| 1 September 2011     | 8.31–9.45           | Galati–Braila                     | Braila urban area        |
| 25 August 2012       | 7.53–8.89           | Galati–Braila                     | Braila urban area        |
| 21 July 2014         | 9.51–10.96          | Galati–Slobozia                   | Slobozia industrial area |
| 13 August 2014       | 7.32–8.19           | Galati                           | Galati industrial area   |

* Coordinated Universal Time. ** ULT-DOAS = Ultralight Trike-Differential Optical Absorption Spectroscopy.

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![Figure 2. Schematic of the ULT-DOAS measurement principle.](image-url)
2.2. Determination of the NO$_2$ Tropospheric Vertical Column and Flux Calculation

2.2.1. Retrieval of NO$_2$ Slant Column

The analysis of the measured spectra was performed using the QDOAS software [25]. For the NO$_2$ fit, the spectral window of 425–490 nm was used. The NO$_2$ spectral analysis included five absorption cross sections: the NO$_2$ cross sections at 298 K and 220K [26], the O$_3$ cross section at 223 K [27], the O$_4$ cross section [28] and a Ring spectrum [29]. A fifth-degree polynomial to account for scattering processes and broad-band absorption in the atmosphere was used in the DOAS analysis. The direct result of the spectral analysis is a differential slant column density (DSCD), which is the integrated trace gas concentration along the light path through the atmosphere. The DSCD is the difference between the slant column densities in the measured spectra (SCD$_{\text{meas}}$) and the Fraunhofer reference spectrum (SCD$_{\text{ref}}$). The NO$_2$ amount in the Fraunhofer reference spectrum is unknown and its retrieval is important for the determination of the SCD$_{\text{meas}}$ (Equation (1)).

$$\text{SCD}_{\text{meas}} = \text{DSCD} + \text{SCD}_{\text{ref}}$$ (1)

Figure 3 presents a typical DOAS fit using a spectrum recorded during the experiment close to the Galati steel factory, on 13 August 2014.

The Slant Column Density (SCD) is converted to a Vertical Column Density (VCD) by means of an Air Mass Factor (AMF), which is defined as the ratio between SCD and VCD (Equation (2))

$$\text{AMF} = \frac{\text{SCD}}{\text{VCD}} = \frac{\tau_{\text{SCD}}}{\tau_{\text{VCD}}}$$ (2)

where $\tau_{\text{SCD}}$ and $\tau_{\text{VCD}}$ are the optical thickness for the slant column (SCD) and vertical column (VCD), respectively.

Since the measured spectra contain information about both stratospheric and tropospheric NO$_2$ content, the SCD$_{\text{meas}}$ can be written as:

$$\text{SCD}_{\text{meas}} = \text{AMF}_{\text{tropo}} \cdot \text{VCD}_{\text{tropo}} + \text{AMF}_{\text{strato}} \cdot \text{VCD}_{\text{strato}} - \text{SCD}_{\text{ref}}$$ (3)
The above equations can be further simplified assuming that SCD_{ref} is dominated by stratospheric NO\textsubscript{2}. Using this assumption, the stratospheric contributions can be canceled by the NO\textsubscript{2} amount in the reference spectrum (Equation (4)) [8].

\[
VCD_{tropo} = \frac{DSCD}{AMF_{tropo}}
\]  

(4)

The assumption presented above is valid if the reference spectrum (needed for the spectral evaluation) is recorded at noon, in an area with a very low NO\textsubscript{2} content and if SCD_{strato} does not vary in time. The Fraunhofer reference spectrum could also be a zenith-sky spectrum recorded at high altitude over the boundary layer [30]. However, in this work the tropospheric NO\textsubscript{2} VCD is based on calculations using Equation (3).

2.2.2. Deduction of the SCD_{ref} and VCD_{strato}

To avoid systematic errors due to the use of multiple reference spectra, only one spectrum will be used for the spectral analysis of all DOAS observations presented in this paper.

The NO\textsubscript{2} amount in the reference spectrum was calculated using a photo-chemically modified Langley plot [6,31]. The SCD_{ref} corresponds to a zenith spectrum recorded at noon, in a clean rural area close to Galati city. The spectrum was recorded on 13 August 2014 (9.70 UTC and solar zenith angle (SZA) = 31.55\degree).

The photo-chemically modified Langley plot was applied for the twilight sunrise observations performed on 21 July 2014. By applying the Langley plot method, we calculated the SCD_{ref} as \(4.1 \times 10^{15}\) molecules/cm\(^2\) of NO\textsubscript{2}.

The stratospheric contribution used for the retrieval of the VCD\textsubscript{tropo} is derived from the assimilated vertical stratospheric columns simulated by Dutch OMI NO\textsubscript{2} (DOMINO). Table 2 shows the satellite overpass data sets that were used for the retrieval algorithm presented in this work.

| Day          | Orbit Nr. | Overpass Time UTC | Stratospheric VCD \([\times 10^{15}\text{ molecules/cm}^2]\) |
|--------------|-----------|-------------------|--------------------------------------------------|
| 1 September 2011 | 37,923    | 11:04:28          | 3.76                                             |
| 25 August 2012  | 43,151    | 11:11:07          | 3.75                                             |
| 21 July 2014    | 53,272    | 11:17:36          | 4.14                                             |
| 13 August 2014  | 53,607    | 11:23:47          | 3.74                                             |

VCD = Vertical Column Density.

Figure 4A presents the SCD determined at twilight sunrise on 21 July 2014 compared with the SCD_{strato} derived from the DOMINO Level 2 product [32] scaled with a chemically modified AMF calculated by PSCBOX [33,34]. More details about the retrieval of the SCD_{strato} using twilight observations and model simulations are presented in [6]. A good agreement between the two types of SCD_{strato} determinations is obtained, which gives confidence in the stratospheric SCD measured by our static DOAS observations.

Figure 4B shows the SCD_{strato} derived from DOMINO compared with the SCDs determined by car-DOAS zenith-sky observations and ULT-DOAS measurements performed in the zenith geometry, for the same day of 21 July 2014. From this plot, one can see that the car-DOAS measurements are dominated by tropospheric NO\textsubscript{2} while the zenith-sky ULT-DOAS observation presents a low amount of NO\textsubscript{2} close to the stratospheric NO\textsubscript{2} SCD derived from OMI. This is due to the fact that zenith-sky ULT-DOAS observations are performed above the NO\textsubscript{2} plume or above the planetary boundary layer.
2.2.3. Radiative Transfer Calculation

In order to determine the VCD, the SCD retrieved with the DOAS method has to be converted using an appropriate AMF. The geometric approximation for the airborne DOAS observations assumes a simple reflection of the sunlight on the earth’s surface. In this case (neglecting the earth’s sphericity) the nadir AMF can be described as a function of the solar zenith angle (SZA) as:

$$ \text{AMF}_{geo} = 1 + \frac{1}{\cos(SZA)} $$  \hspace{1cm} (5)

Since the ULT-DOAS measurements were performed in the open atmosphere using scattered sunlight radiation, the radiative transfer during the observations needs to be modeled to interpret the retrieved data. In this work, the AMF was calculated using the radiative transfer model (RTM) UVspec/DISORT [35], which is a fully spherical model. This RTM has been validated using six other different codes [34].

The general assumptions made for the radiative transfer calculation using the RTM UVspec/DISORT are introduced in Table 3.

| Trace Gas | Nitrogen Dioxide |
|-----------|------------------|
| wavelength| 440 nm           |
| flight altitude | 1000 m 1500 m |
| albedo    | 0.1              |
| visibility| 1 km 5 km 20 km  |
| line of sight | 0°              |

Results of AMF simulations for the nadir flight performed on 13 August 2014, using the input parameters presented in Table 3, are displayed in Figure 5. The geometric AMF for the nadir view is also shown. The visibility parameter accounts for the effect of aerosols.
3. Results and Discussions

The airborne DOAS observations were designed to determine the distribution of tropospheric NO$_2$ from the South-East of Romania from urban, industrial and rural areas and associated flux.

The first flights were performed on 1 September 2011 and 25 August 2012, and aimed at measuring the NO$_2$ around the industrial area of Galati city and from Braila city. The flight performed on 21 July 2014 aimed to measure the NO$_2$ emitted by a fertilizer factory located nearby Slobozia city; unfortunately, during the DOAS flight the fertilizer factory was not operational. Figure 6 presents the horizontal distribution of the tropospheric NO$_2$ determined in nadir geometry for 1 September 2011, while Figure 7 depicts the results during a similar flight, but on 25 August 2012. During this experiment, the plume from the industrial platform was not fully crossed by the optical instrument onboard the ULT. The wind was northerly resulting in local increases of the NO$_2$ amount detected by the spectrometer. The maximum tropospheric NO$_2$ VCD detected during this experiment was $(1.1 \pm 0.24) \times 10^{16}$ molecules/cm$^2$ while the minimum tropospheric NO$_2$ VCD was $(2.1 \pm 0.81) \times 10^{15}$ molecules/cm$^2$.

Figure 5. AMF simulations obtained from RTM calculations using UVspec/DISORT for various input parameters, for 13 August 2014. AMF = Air Mass Factor.

Figure 6. Map of tropospheric NO$_2$ VCD determined on 1 September 2011 using ULT-DOAS observations.
The trajectory of the flight on 13 August 2014 gave us the opportunity of calculating the NO$_2$ flux emissions around the industrial area of Galati city. This was not possible for the other flights because encircling the NO$_2$ source was not authorized.

On 1 September 2011, the NO$_2$ amount was low relative to the other day. The flight comprised almost two complete circles around Braila; however, the NO$_2$ displayed no clear variation. The horizontal distribution of NO$_2$ was quite homogenous over Braila city on this day.

A double experiment was performed on 13 August 2014, using both a ULT-DOAS and a car-based DOAS system. The mobile ground-based DOAS observations were performed using the same equipment during 9.75–10 UTC, while the ULT-DOAS observations were performed during 7.30–8.15 UTC.

Figure 8 shows the tropospheric NO$_2$ VCD derived along the trajectory of the ULT-DOAS measurements. The right plot shows a photograph of the plume crossed by the ULT flights. During the same day, approximately 1 h after the acquisition of the ULT-DOAS measurements, a zenith-sky car-DOAS system was used to sample the NO$_2$ plume at the same location as the ULT-DOAS observations. We assume that the quantity of the NO$_2$ emitted by the steel factory was almost constant during the airborne and car-DOAS system.

**Figure 7.** Map of tropospheric NO$_2$ VCD determined on 25 August 2012 using ULT-DOAS observations.

**Figure 8.** The tropospheric NO$_2$ VCD along the flight trajectory using the ULT-DOAS system on 13 August 2014 (left); Photography of the NO$_2$ plume determined on the same day (right).
Figure 9 presents the NO$_2$ VCD derived from the nadir airborne DOAS observations performed over the industrial area of Galati city compared with zenith-sky ground-based mobile DOAS measurements performed over the same area in the same day (13 August 2014). In these figures, we show the original SCDs (A) and the retrieved tropospheric NO$_2$ VCD (B). Figure 9A shows that the DSCDs determined from the ULT-DOAS system are ~30% higher than the DSCDs determined using car-DOAS observations. This difference is attributed to the different observation geometries. After appropriate AMF calculation (see Section 2.2.2), both observations show close results.

The ULT flight above the industrial plume led to the detection of a maximum tropospheric NO$_2$ VCD of $\left(1.41 \pm 0.27\right) \times 10^{16}$ molecules/cm$^2$ while the car-DOAS observation shows a maximum tropospheric NO$_2$ VCD of $\left(1.36 \pm 0.21\right) \times 10^{16}$ molecules/cm$^2$. Figure 10 displays the correlation between the tropospheric NO$_2$ VCD retrieved by the ULT-DOAS and the car-DOAS instrument, where closest spatially coincident data were selected. A Pearson correlation coefficient $R = 0.71$ was obtained between ground mobile DOAS observations and airborne DOAS measurements.

![Figure 9](image1)

**Figure 9.** Comparisons between ULT-DOAS and car-DOAS observations performed on 13 August 2014. (A) The results of the preliminary DOAS analysis (DSCDs) and (B) after determination of the vertical columns (VCDs).

![Figure 10](image2)

**Figure 10.** Correlation between tropospheric VCDs measured by the ULT-DOAS and the car-DOAS instrument on 13 August 2014.
NO₂ Flux Calculation

The NO₂ flux above the industrial platform located nearby Galati city was calculated by performing upwind and downwind measurements around the point source using ULT-DOAS observations on 13 August 2014. The calculation of the emission flux is based on the following parameters: the NO₂ VCD determined from the transect over the plume, the wind speed and the wind factor correction, taking into account the angle between the flight direction and wind direction (Equation (6)), [20,36,37]:

\[
\text{Flux}_{\text{NO}_2} = \sum_i \text{VCD}_{\text{NO}_2}(s_i) \cdot v \cdot \sin (\alpha_i) \cdot \Delta s_i
\]  

where VCD_{NO2} is the NO₂ tropospheric vertical column, \(v\) is the wind speed, \(\alpha\) the angle between wind direction and driving route, \(i\) is the observations index, and \(\Delta s_i\) is the distance between two successive spectra.

The wind data used for the NO₂ flux calculation rely on measurements of the automatic weather station (Davis Vantage Pro2) located in the campus university of Galati city (45.44° N, 28.05° E), while vertical wind profiles come from the Hybrid Single Particle Lagrangian Integrated Trajectory Model (HYSPLIT) [38] model using archived dataset GDAS 0.5° × 0.5°. The weather station is located at 30 m height and acquires data every 30 min. Since no atmospheric sounding was possible during the experiments, the wind measured on the ground was scaled to the output of HYSPLIT model simulation at 1000 m altitude. During the period of the ULT-DOAS measurements, the mean NO₂ emission flux was determined to be \((3.45 \pm 0.89) \times 10^{-3}\) kg/s. A local environmental report indicates ~600 tons/year NOₓ emissions emitted by the steel factory [39]. Using a Leighton ratio \((L = [\text{NO}] / [\text{NO}_2])\) of 0.3, we calculated that the steel factory has emitted a mean of \(~10 \times 10^{-3}\) kg/s of NO₂. The difference between the two types of estimation may be attributed to the fact that the NOₓ emissions from the steel factory are dependent on the quantity of the steel produced, which can vary from one day to another. Also, the derived NO₂ fluxes must be dealt with some care because of the probably incomplete NO to NO₂ conversion [40].

4. Conclusions

Ultralight-trike DOAS observations were performed in the South-East of Romania during four days between 2011 and 2014. The first two flights were focused over Braila city, the third aimed at measurements of the NO₂ plume emitted by a fertilizer factory near Slobozia city. Unfortunately, during the DOAS flight the fertilizer factory was not operational. The last flight, performed on 13 August 2014, was focused over the industrial area of Galati city. Nadir observations were performed around the industrial platform of Galati city aiming at measuring the tropospheric NO₂ VCD around the source and at evaluating the associated NO₂ flux. To retrieve the tropospheric NO₂ VCD from ULT-DOAS observations, complementary ground- and space-based measurements were used.

We showed that the tropospheric NO₂ VCD deduced from the ULT-DOAS observations are consistent with measurements performed from the ground using a zenith-sky car-DOAS system. Although two hours separated the two experiments, a correlation coefficient of \(R = 0.71\) was found between the two results, a tropospheric NO₂ VCD of \((1.41 \pm 0.27) \times 10^{16}\) molecules/cm² and an estimated associated flux of \((3.45 \pm 0.89) \times 10^{-3}\) kg/s was measured close to the industrial area of Galati city on 13 August 2014, the only day that it was possible to determine the NO₂ flux.

Also, we showed that the stratospheric SCD derived from ground-based and airborne measurements correlates well with stratospheric NO₂ derived from observations by the OMI satellite sensor.

Based on this study, we conclude that the ULT is an efficient tool which allows determining with a high resolution the NO₂ distribution around urban or industrial sources. Also, the ULT-DOAS system is a very efficient tool for measuring fluxes due to its flexibility during the flights and the
possibility to access remote areas. The ULT-DOAS system might also constitute a promising tool for satellite validation and calibration under clear-sky conditions, especially for upcoming high-resolution sensors such as the TROPOMI/Sentinel-5 Precursor instrument to be launched in summer 2017.

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**Author Contributions:** Daniel-Eduard Constantin and Alexis Merlaud conceived and designed the study; Daniel-Eduard Constantin analyzed the data; and Mirela Voiculescu, Carmelia Dragomir, Lucian Georgescu, Gaia Pinardi, François Hendrick and Michel Van Roozendael improved the paper.

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**References**

1. Solomon, S.; Portmann, R.W.; Sanders, R.W.; Daniel, J.S.; Madsen, W.; Bartram, B.; Dutton, E.G. On the role of nitrogen dioxide in the absorption of solar radiation. *J. Geophys. Res.* **1999**, *104*, 12047–12058. [CrossRef]

2. Platt, U. Differential optical absorption spectroscopy (DOAS). *Chem. Anal. Ser.* **1994**, *127*, 27–83.

3. Brewer, A.W.; McElroy, C.T.; Kerr, J.B. Nitrogen dioxide concentration in the atmosphere. *Nature* **1973**, *246*, 129–133. [CrossRef]

4. Noxon, J.F. Nitrogen dioxide in the stratosphere and troposphere measured by ground-based absorption spectroscopy. *Science* **1975**, *189*, 547–549. [CrossRef] [PubMed]

5. Hendrick, F.; Müller, J.-F.; Clémer, K.; Wang, P.; De Mazière, M.; Fayt, C.; Giezen, C.; Hermans, C.; Ma, J.Z.; Pinardi, G.; et al. Four years of ground-based MAX-DOAS observations of HONO and NO₂ in the Beijing area. *Atmos. Chem. Phys.* **2014**, *14*, 765–781. [CrossRef]

6. Constantin, D.-E.; Merlaud, A.; van Roozendael, M.; Voiculescu, M.; Fayt, C.; Hendrick, F.; Pinardi, G.; Georgescu, L. Measurements of Tropospheric NO₂ in Romania Using a Zenith-Sky Mobile DOAS System and Comparisons with Satellite Observations. *Sensors* **2013**, *13*, 3922–3940. [CrossRef]

7. Dragomir, C.; Constantin, D.-E.; Voiculescu, M.; Georgescu, L.; Merlaud, A.; Roozendael, M.V. Modeling results of atmospheric dispersion of NO₂ in an urban area using METI–LIS and comparison with coincident mobile DOAS measurements. *Atmos. Pollut. Res.* **2015**, *6*, 503–510. [CrossRef]

8. Merlaud, A.; van Roozendael, M.; van Gent, J.; Fayt, C.; Maes, J.; Toledo-Fuentes, X.; Ronveaux, O.; de Mazière, M. DOAS measurements of NO₂ from an ultralight aircraft during the Earth Challenge expedition. *Atmos. Meas. Tech.* **2012**, *5*, 2057–2068. [CrossRef]

9. Meier, A.C.; Schönhardt, A.; Bösch, T.; Richter, A.; Seyler, A.; Ruhtz, T.; Constantin, D.-E.; Shaiganfar, R.; Wagner, T.; Merlaud, A.; et al. High-resolution airborne imaging DOAS-measurements of NO₂ above Bucharest during AROMAT. *Atmos. Meas. Tech. Discuss.* **2016**. [CrossRef]

10. Tack, F.; Merlaud, A.; Iordache, M.-D.; Danckaert, T.; Yu, H.; Fayt, C.; Meuleman, K.; Deutsch, F.; Fierens, F.; van Roozendael, M. High resolution mapping of the NO₂ spatial distribution over Belgian urban areas based on airborne APEX remote sensing. *Atmos. Meas. Tech. Discuss.* **2017**. [CrossRef]

11. Merlaud, A.; Constantin, D.; Fayt, C.; Maes, J.; Mingireanu, F.; Mocanu, I.; Georgescu, L.; Roozendael, M. Small whiskbroom imager for atmospheric composition monitoring (SWING) from an unmanned aerial vehicle (UAV). In Proceedings of the 21st ESA Symposium on European Rocket and Balloon Programmes and related Research, Thun, Switzerland, 9–13 June 2014; pp. 1–7.

12. Bovensmann, H.; Burrows, J.P.; Buchwitz, M.; Frerick, J.; Noël, S.; Rozanov, V.V.; Chance, K.V.; Goede, A.H.P. SCIAMACHY—Mission objectives and measurement modes. *J. Atmos. Sci.* **1999**, *56*, 127–150. [CrossRef]

13. Leveled, P.; van den Oord, G.; Dobber, M.; Malik, A.; Visser, H.; de Vries, J.; Stammes, P.; Lundell, J.; Saari, H. The ozone monitoring instrument. *IEEE T. Geosci. Remote.* **2006**, *44*, 1093–1101. [CrossRef]

14. Munro, R.; Eisinger, M.; Anderson, C.; Callies, J.; Corpaccioli, E.; Lang, R.; Lefebvre, A.; Livschitz, Y.; Albinana, A.P. GOME-2 on MetOp. In Proceedings of the 2006 EUMETSAT Meteorological Satellite Conference, Helsinki, Finland, 12–16 June 2006; p. 48.

15. van der A, R.J.; Eskes, H.J.; Boersma, K.F.; van Noije, T.P.C.; van Roozendael, M.; de Smedt, L.; Peters, D.H.M.U.; Meijer, E.W. Trends, seasonal variability and dominant NOₓ source derived from a ten year record of NO₂ measured from space. *J. Geophys. Res.* **2008**, *113*, D04302. [CrossRef]
16. Varotsos, C.; Christodoulakis, J.; Tzanis, C.; Cracknell, A.P. Signature of tropospheric ozone and nitrogen dioxide from space: A case study for Athens, Greece. Atmos. Environ. 2014, 89, 721–730. [CrossRef]

17. Junkermann, W. An ultralight aircraft as platform for research in the lower troposphere: System performance and first results from radiation transfer studies in stratiform aerosol layers and broken cloud conditions. J. Atmos. Ocean. Technol. 2001, 18, 934. [CrossRef]

18. Junkermann, W. An ultralight aircraft as platform for research in the lower troposphere: System performance and first results from radiation transfer studies in stratiform aerosol layers and broken cloud conditions. J. Atmos. Ocean. Technol. 2001, 18, 934. [CrossRef] [PubMed]

19. Chazette, P.; Sanak, J.; Dulac, F. New Approach for Aerosol Profiling with a Lidar Onboard an Ultralight Aircraft: Application to the African Monsoon Multidisciplinary Analysis. Environ. Sci. Technol. 2007, 41, 8335–8411. [CrossRef] [PubMed]

20. Wang, P.; Richter, A.; Bruns, M.; Burrows, J.P.; Scheele, R.; Junkermann, W.; Heue, K.-P.; Wagner, T.; Platt, U.; Pundt, I. Airborne multi-axis DOAS measurements of tropospheric SO$_2$ plumes in the Po-valley, Italy. Atmos. Chem. Phys. 2006, 6, 329–338. [CrossRef]

21. Grutter, M.; Basaldud, R.; Rivera, C.; Harig, R.; Junkermann, W.; Caetano, E.; Delgado-Granados, H. SO$_2$ emissions from Popocatépetl volcano: Emission rates and plume imaging using optical remote sensing techniques. Atmos. Chem. Phys. 2008, 8, 6655–6663. [CrossRef]

22. General, S.; Pöhler, D.; Sihler, H.; Bobrowski, N.; Frieß, U.; Zielcke, J.; Horbanski, M.; Shipson, P.B.; Stirm, B.H.; Simpson, W.R.; et al. The Heidelberg Airborne Imaging DOAS Instrument (HAIDI)—A novel imaging DOAS device for 2-D and 3-D imaging of trace gases and aerosols. Atmos. Meas. Tech. 2014, 7, 3459–3485. [CrossRef] [PubMed]

23. Liu, L.; Flatøy, F.; Ordóñez, C.; Braathen, G.O.; Hak, C.; Junkermann, W.; Andreani-Aksoyoglu, S.; Mellqvist, J.; Galle, B.; Prévôt, A.S.H.; et al. Photochemical modelling in the Po basin with focus on formaldehyde and ozone. Atmos. Chem. Phys. 2007, 7, 121–137. [CrossRef]

24. Airborne Romanian Measurements of Aerosols and Trace Gases (AROMAT). Available online: http://uv-vis.aeronomie.be/aromat/index.php (accessed on 15 January 2017).

25. Danckaert, T.; Fayt, C.; van Roozendael, M.; de Smedt, I.; Letocart, V.; Pinardi, G. QDOAS Software user manual Version 2.111-April 2016, UV-Visible DOAS Research Group of the Royal Belgian Institute for Space Aeronomy Web Site. Available online: http://uv-vis.aeronomie.be/software/QDOAS/QDOAS_manual.pdf (accessed on 15 December 2016).

26. Vandaele, A.; Hermans, C.; Simon, P.; Carleer, M.; Colin, R.; Fally, S.; Mérienne, F.; Jenouvrier, A.; Coq, B. Measurements of the NO$_2$ absorption cross-section from 42000 cm$^{-1}$ to 10000 cm$^{-1}$ (238–1000 nm) at 220 K and 294 K (220 K). J. Quant. Spectrosc. Radiat. Transf. 1998, 59, 171–184. [CrossRef]

27. Bogumil, K.; Orphal, J.; Homann, T.; Voigt, S.; Spieß, T.; Fleischmann, O.C.; Vogel, A.; Hartmann, M.; Kromminga, H.; Bovensmann, H.; et al. Measurements of molecular absorption spectra with the SCIAMACHY pre-flight model: Instrument characterization and reference data for atmospheric remote-sensing in the 230–2380 nm region. J. Photochem. Photobiol. A Chem. 2003, 157, 167–184. [CrossRef]

28. Thalman, R.; Volkamer, R. Temperature dependent absorption cross-sections of O$_2$–O$_2$ collision pairs between 340 and 630 nm and at atmospherically relevant pressure. Phys. Chem. Chem. Phys. 2013, 15, 15371–15381. [CrossRef] [PubMed]

29. Chance, K.V.; Spurr, R.J.D. Ring effect studies: Rayleigh scattering, including molecular parameters for rotational Raman scattering, and the Fraunhofer spectrum. Appl. Opt. 1997, 36, 5224–5230. [CrossRef] [PubMed]

30. Baidar, S.; Oetjen, H.; Coburn, S.; Dix, B.; Ortega, I.; Sinreich, R.; Volkamer, R. The CU Airborne MAX-DOAS instrument: Vertical profiling of aerosol extinction and trace gases. Atmos. Meas. Tech. 2013, 6, 719–739. [CrossRef]

31. Vaughan, G.; Quinn, P.T.; Green, A.C.; Bean, J.; Roscoe, H.K.; van Roozendael, M.; Goutail, F. SAOZ measurements of stratospheric NO$_2$ at Aberystwyth. J. Environ. Monit. 2006, 8, 353–361. [CrossRef] [PubMed]

32. Dirksen, R.J.; Boersma, K.F.; Eskes, H.J.; Ionov, D.V.; Bucsel, E.J.; Levelt, P.F.; Kelder, H.M. Evaluation of stratospheric NO$_2$ retrieved from the Ozone Monitoring Instrument: Intercomparison, diurnal cycle and trending. J. Geophys. Res. 2011, 116, D08305. [CrossRef]
33. Hendrick, F.; Barret, B.; van Roozendael, M.; Boesch, H.; Butz, A.; De Mazière, M.; Goutail, F.; Hermans, C.; Lambert, J.-C.; Pfeilsticker, K.; et al. Retrieval of nitrogen dioxide stratospheric profiles from ground-based zenith-sky UV-visible observations: Validation of the technique through correlative comparisons. *Atmos. Chem. Phys.* 2004, 4, 2091–2106. [CrossRef]

34. Hendrick, F.; van Roozendael, M.; Kylling, A.; Petritoli, A.; Rozanov, A.; Sanghavi, S.; Schofield, R.; von Friedeburg, C.; Wagner, T.; Wittrock, F.; et al. Intercomparison exercise between different radiative transfer models used for the interpretation of ground-based zenith-sky and multi-axis DOAS observations. *Atmos. Chem. Phys.* 2006, 6, 93–108. [CrossRef]

35. Mayer, B.; Kylling, A. Technical note: The libRadtran software package for radiative transfer calculations—Description and examples of use. *Atmos. Chem. Phys.* 2005, 5, 1855–1877. [CrossRef]

36. Rivera, C.; Sosa, G.; Wöhrnschimmel, H.; de Foy, B.; Johansson, M.; Galle, B. Tula industrial complex (Mexico) emissions of SO2 and NO2 during the MCMA 2006 field campaign using a mobile mini-DOAS system. *Atmos. Chem. Phys.* 2009, 9, 6351–6361. [CrossRef]

37. Johansson, M.; Rivera, C.; de Foy, B.; Lei, W.; Song, J.; Zhang, Y.; Galle, B.; Molina, L. Mobile mini-DOAS measurement of the outflow of NO2 and HCHO from Mexico City. *Atmos. Chem. Phys.* 2009, 9, 5647–5653. [CrossRef]

38. Stein, A.F.; Draxler, R.R.; Rolph, G.D.; Stunder, B.J.B.; Cohen, M.D.; Ngan, F. NOAA’s HYSPLIT atmospheric transport and dispersion modeling system. *Bull. Am. Meteorol. Soc.* 2015, 96, 2059–2077. [CrossRef]

39. *Studiu Privind Calitatea Aerului in Municipiul Galati;* Municipiul Galati: Galati, Romania, 2016.

40. Frins, E.; Bobrowski, N.; Osorio, M.; Casaballe, N.; Belsterli, G.; Wagner, T.; Platt, U. Scanning and mobile multi-axis DOAS measurements of SO2 and NO2 emissions from an electric power plant in Montevideo, Uruguay. *Atmos. Environ.* 2014, 98, 347–356. [CrossRef]

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