High-resolution NO$_2$ remote sensing from the Airborne Prism EXperiment (APEX) imaging spectrometer

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Abstract. We present and evaluate the retrieval of high spatial resolution maps of NO$_2$ vertical column densities (VCD) from the Airborne Prism EXperiment (APEX) imaging spectrometer. APEX is a novel instrument providing airborne measurements of unique spectral and spatial resolution and coverage as well as high signal stability. In this study, we use spectrometer data acquired over Zurich, Switzerland, in the morning and late afternoon during a flight campaign on a cloud-free summer day in June 2010. NO$_2$ VCD are derived with a two-step approach usually applied to satellite NO$_2$ retrievals, i.e. a DOAS analysis followed by air mass factor calculations based on radiative transfer computations. Our analysis demonstrates that APEX is clearly sensitive to NO$_2$ VCD above typical European tropospheric background abundances ($>1\times10^{15}$ molec cm$^{-2}$). The two-dimensional maps of NO$_2$ VCD reveal a very convincing spatial distribution with strong gradients around major NO$_x$ sources (e.g. Zurich airport, waste incinerator, motorways) and low NO$_2$ in remote areas. The morning overflights resulted in generally higher NO$_2$ VCD and a more distinct pattern than the afternoon overflights which can be attributed to the meteorological conditions prevailing during that day with stronger winds and hence larger dilution in the afternoon. The remotely sensed NO$_2$ VCD are also in reasonably good agreement with ground-based in-situ measurements from air quality networks considering the limitations of comparing column integrals with point measurements. Airborne NO$_2$ remote sensing using APEX will be valuable to detect NO$_2$ emission sources, to provide input for NO$_2$ emission modelling, and to establish links between in-situ measurements, air quality models, and satellite NO$_2$ products.

1 Introduction

Nitrogen dioxide (NO$_2$) is an important reactive trace gas in the troposphere. NO$_2$ acts as an ozone and aerosol precursor and can directly or indirectly affect human health (e.g. pulmonary or cardiovascular diseases) (Brunekreef and Holgate, 2002) and ecosystem functions and services (e.g. damage of leaves, reduction of crop production, acidification) (Bell and Treshow, 2002). Besides natural sources such as lighting and soil emissions, the major fraction of tropospheric NO$_2$ is related to anthropogenic activities, notably fossil fuel combustion by traffic and industry. Despite significant improvements of air quality in European countries during the past two decades, air quality thresholds are still frequently exceeded and further efforts are needed particularly regarding reductions of particulate matter, ozone, and nitrogen oxides (NO$_x$ = NO + NO$_2$). Measurements of NO$_2$ in the troposphere are performed with various in-situ, airborne, and spaceborne instruments. Tropospheric vertical column densities (TVCD) retrieved from satellites (e.g. from the Scanning Imaging Absorption Spectrometer for Atmospheric Chartography (SCIAMACHY), the Global Ozone Monitoring Experiment (GOME(-2)), or the Ozone Monitoring Instrument (OMI)) have largely contributed to a better understanding of the global distribution of NO$_2$ as well as its sources and trends (e.g. Boersma et al., 2004; Richter et al., 2005; van der A et al., 2008; Zhou et al., 2012). The spatial resolution of satellite products on the order of multiple tens of kilometers is only sufficient to detect aggregate sources like entire cities (Beirle et al., 2011) and individual sources like emissions from power plants (Kim et al., 2006) or ships (Beirle et al., 2004) if they are sufficiently separated in space from other sources. Ground-based in-situ instruments, on the other hand, provide accurate and continuous trace gas emissions.
measurements but lack of homogeneous geographical coverage. Airborne remote sensing observations can in this regard provide a valuable link between ground-based and space-borne NO$_2$ information. For example, airborne multi-axis differential optical absorption spectroscopy (AMAXDOAS) was used to retrieve NO$_2$ TVCD (Wang et al., 2005), NO$_2$ profile information (Bruns et al., 2006), or to validate SCIAMACHY NO$_2$ TVCD (Heue et al., 2005). Heue et al. (2008) demonstrated the capability of an imaging DOAS instrument to retrieve two-dimensional NO$_2$ distributions over the highly polluted Highveld plateau in South Africa.

The Airborne Prism EXperiment (APEX) imaging spectrometer is a state-of-the art instrument with an unprecedented combination of high spectral and spatial resolution, good two-dimensional geographical coverage, and high signal stability. Two NO$_2$ distribution maps were retrieved from imaging spectrometer data acquired over Zurich, Switzerland, in the morning and the late afternoon of 26 June 2010. Our results are considered as one of the first spatio-temporal investigations of the NO$_2$ distribution on a regional to local scale. In particular, we present the first high-resolution maps of NO$_2$ VCD in a city measured by an airborne imaging spectrometer. The specific objectives of this study are (i) the presentation of a retrieval scheme to obtain NO$_2$ quantities from APEX imaging spectrometry data and (ii) the qualitative and quantitative assessment of the NO$_2$ products, considering amongst others in-situ measurements of NO$_2$.

2 Instrument and data acquisition

2.1 The APEX instrument

APEX is a dispersive pushbroom imaging spectrometer for environmental monitoring developed by a Swiss-Belgium consortium in the framework of the ESA-PRODEX programme (Itten et al., 2008). APEX consists of the imaging spectrometer itself, a Calibration Home Base (CHB) for instrument calibration, and a data processing and archiving facility (PAF) for operational product generation (Jehle et al., 2010). Table 1 gives a brief overview of the sensor characteristics.

The APEX imaging spectrometer consists of a CCD detector for the visible and near infrared (VNIR) and a CMOS detector for the shortwave infrared (SWIR) wavelength region. The NO$_2$ retrieval is based on absorption bands in the UV/VIS spectral domain, and, hence, only the VNIR specification is discussed hereinafter. Since atmospheric trace gases exhibit spectrally narrow absorption features, the (spectrally) unbinned configuration was applied to provide highest spectral resolution. The spectral sampling interval (SSI) and the full width at half maximum (FWHM) are non-linear functions of the wavelength and increase with longer wavelengths. According to pre-flight sensor calibration, the SSI increases from 0.66 to 1.42 nm and the FWHM from 1.00 to 1.95 between 420 nm and 520 nm where NO$_2$ slant column densities (SCD) are usually derived. APEX is a pushbroom scanner and measures radiances in 1000 spatial pixels across-track. The extent of the flight line along-track depends on the pre-defined flight pattern. The spatial resolution in across-track direction is determined by the sensor’s instantaneous field of view (IFOV) of 0.028°. The spatial resolution along-track depends on the integration time. This unparalleled combination of high resolution, geographical coverage, and high signal stability makes APEX very attractive for a range of remote sensing applications, e.g. in the fields of vegetation, atmosphere, limnology, geology, or natural hazard studies.

2.2 Test site and data

APEX acceptance flight activities took place in Belgium and Switzerland in June/July 2010 using a Dornier Do-228 aircraft operated by the German Aerospace Center (DLR) (Jehle et al., 2010). Image data were collected in more than 42 flight hours for a variety of studies of the land-surface and the atmosphere. Six of these image data sets, acquired in unbinned mode over Zurich, Switzerland, on Saturday 26 June under cloud free conditions (cf. Fig. 1) were used in this study. Three of them were flown around 10:00 local time (08:00 UTC) and three in the late afternoon around 17:30 local time (15:30 UTC) at a cruise level of 5400 m above sea level (asl) and a flight heading of 45° and 225°, respectively. The data integration time of APEX can be adjusted. The definition of an adequate integration time for the unbinned mode is, however, critical because the incoming radiation in the unbinned bands is comparatively small. An integration time of 57 ms was found to be a reasonable compromise between radiometric performance and the resulting pixel size of 2.5 m across-track and approximately 6 m along-track, respectively. This setting, in particular, ensures a sufficient signal stability in these spectral bands (i.e. the visible part from 370–500 nm) and avoids signal saturation in other parts of the sampled spectrum (e.g. near infrared (NIR) from 700 nm onwards).

Zurich is Switzerland’s largest city with about 400 000 inhabitants surrounded by an agglomeration of more than one million inhabitants. The test site includes a wide range of surface types like buildings, roads, parks, forests, and part of Lake Zurich (Fig. 1, cf. also APEX true colour composite in Fig. 7a). It also includes more rural areas with decreased atmospheric pollution levels like the Uetliberg mountain range. Several major NO$_x$ sources were covered. Three national motorways surround the city to the north, west and south (A1, A3, A4), and major transit roads lead through the city with a usually high traffic volume. In addition, two waste incinerators as well as part of the approach corridor of Zurich airport are located in the test site area. Zurich was also selected for the flight experiments because of the dense ground-based air quality network with eight stations within the range covered by APEX (Fig. 1). The National Air Pollution Monitoring
Table 1. Selected characteristics of the APEX imaging spectrometer.

<table>
<thead>
<tr>
<th></th>
<th>VNIR</th>
<th>SWIR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spectral range</td>
<td>380.5–971.7 nm</td>
<td>941.2–2501.5 nm</td>
</tr>
<tr>
<td>Spectral bands</td>
<td>up to 334</td>
<td>198</td>
</tr>
<tr>
<td>Spectral sampling interval</td>
<td>0.55–8 nm</td>
<td>5–10 nm</td>
</tr>
<tr>
<td>Spectral resolution (FWHM)</td>
<td>0.6–6.3 nm</td>
<td>6.2–11 nm</td>
</tr>
<tr>
<td>Spatial pixels (across track)</td>
<td>1000</td>
<td></td>
</tr>
<tr>
<td>Field of view</td>
<td>28°</td>
<td></td>
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<tr>
<td>Instantaneous field of view</td>
<td>0.028° (~ 0.5 mrad)</td>
<td></td>
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<tr>
<td>Spatial sampling interval (across track)</td>
<td>2.5 m at 5000 m AGL</td>
<td></td>
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Fig. 1. Map of Zurich and surrounding areas (data source: Bundesamt für Landestopographie (Swisstopo)). The three morning flight lines are overlayed in black (the afternoon flight lines are almost identical and therefore omitted for clarity). The three white boxes correspond to the position of the examples shown in detail in Fig. 13. An APEX true color composite corresponding to this flight lines is provided in Fig. 7a.

Network (NABEL) provides half-hourly averaged measurements of classical air pollutants like NO$_2$, O$_3$, SO$_2$, PM$_{10}$ at Zurich Kaserne (urban background, situated in a park near the city center) and Duebendorf (suburban). Additional air quality measurements and meteorological parameters are available from the inter-cantonal network OSTLUFT which maintains eight sites in the area of interest from which six sites provided data for 26 June 2010: Heubeeribuehl (periphery), Wettswil Filderen (rural, close to motorway), Wettswil Weieracher (rural), Stampfenbach (urban, kerbside, moderate traffic), Schimmelstrasse (urban kerbside, high traffic volume), and Opfikon (kerbside, motorway).
2.3 Data preparation

APEX data were acquired in unbinned mode to provide highest spectral resolution. This instrumental setting, however, is at the expense of the signal stability. To increase the signal-to-noise ratio (SNR), the imaging spectrometer data were spatially aggregated. A box size of $20 \times 20$ pixels was applied which is expected to increase the SNR 20 times (or $\sqrt{400}$ assuming uncorrelated noise). An image-based SNR estimation applied to the original unbinned APEX data revealed a SNR of 158.5 at 490 nm for dark surfaces (water). Spatial aggregation, hence, increases the SNR to approximately 3170 for dark surfaces. It is relevant to note that image-based SNR estimates usually overestimate noise or underestimate the SNR respectively, as surface variability is inherent in the image statistics. The spatial averaging resulted in a decreased pixel size of around $50 \times 120 \text{m}^2$.

The APEX spectrometer is spectrally and radiometrically calibrated pre-flight. However, it is worth mentioning that this calibration does not compensate for certain effects occurring in-flight. For example, pushbroom sensors are typically affected by slight spectral misregistrations across-track, e.g. spectral smile effects (D’Oдорico et al., 2010) which may depend on the specific flight conditions. In order to minimize these effects, the NO$_2$ retrieval was performed on geometrically uncorrected data to allow a scan-line by scan-line wise processing of the data. In addition, NO$_2$ is determined from raw data (digital numbers or DN) and a spectral calibration is performed directly as part of the retrieval algorithm to account for spectral effects under flight conditions (cf. Sect. 3.1).

In order to obtain surface reflectance as an important input parameter of the retrieval algorithm, a software binning was applied to transform the unbinned data to the standard binning pattern and the data were subsequently calibrated to at-sensor radiances using the APEX-PAF. The re-binned radiance data were then atmospherically corrected using the ACTOR-4 software tool (Richter and Schläpfer, 2002) to obtain hemispherical-conical-reflectance (HCRF) data. In a last step, the unbinned and binned APEX data were geometrically corrected using the PARGE orthorectification software (Schläpfer and Richter, 2002). This processing step is needed to re-project auxiliary data (e.g. a digital elevation model (DHM25, http://www.swisstopo.admin.ch/internet/swisstopo/en/home/products/height/dhm25.html, last access: 12 March 2012)) to the raw geometry of the APEX data to allow a scan-line by scan-line wise processing as mentioned before. Further, the geocorrected data are essential to relate the APEX NO$_2$ vertical column density (VCD) to the in-situ measurements of NO$_2$ concentrations and to identify objects of interest like large NO$_x$ point sources.

3 NO$_2$ retrieval

The derivation of NO$_2$ maps from APEX follows the two step approach usually applied to satellite NO$_2$ retrievals. In a first step, differential slant column densities (dSCD) are derived by the well-known differential optical absorption spectroscopy (DOAS) technique (Platt and Stutz, 2008). Subsequently, the dSCD are converted to VCD by means of air mass factors (AMFs) calculated with a radiative transfer model. Detailed information about the physical principles, applications, and accuracies of DOAS and AMF computations can be found elsewhere (Palmer et al., 2001; Boersma et al., 2004; Platt and Stutz, 2008).

3.1 DOAS analysis

Differential slant column densities (dSCD) were derived with the QDOAS software (http://uv-vis.aeronomie.be/software/QDOAS/, last access: 12 March 2012, Fayt et al., 2011). DOAS analysis requires reference spectra which we obtained from the imaging spectrometer data themselves. Earthshine reference spectra were selected for each individual column of each overflight separately. This results in 50 different reference spectra per overflight and allows minimizing errors in the DOAS analysis caused by spectral miscalibration (e.g. spectral smile) and optical imaging imperfections. Based on visual inspection, areas in the individual overflights were determined which are assumed to only contain a background abundance of NO$_2$ (pollution free), cf. in the forested and elevated area to the south of the city (Fig. 7a). The highlighted areas span over ten rows which were additionally averaged in the columnar direction to increase the SNR of the reference spectra.

NO$_2$ absorption cross sections (at 293 K, Voigt et al., 2002) were subsequently fitted to the differential optical depth derived from the APEX measurements and the reference spectra. Spectrally slowly varying signatures (e.g. from aerosols or surface reflectance) were accounted for by including a fifth-order polynomial in the fit, and instrumental effects such as dark current and/or straylight are dealt with including a ring cross section computed by QDOAS (cf. Fayt et al., 2011), and the interference with O$_3$-O$_2$ was accounted for by fitting an appropriate absorption cross section (Herms et al., 2002). Smallest fitting errors were found by using the 470–510 nm wavelength region (cf. red rectangle in Fig. 2). The usage of a window at shorter wavelengths (e.g. 420–470 nm) led to increased fitting errors, probably due to the lower signal levels and higher noise. We also tested integrating O$_3$ and H$_2$O columns in the DOAS fit which led to distinctly worse results probably due to a too small fitting window relative to the number of cross sections. Retrieved slant O$_3$ and H$_2$O columns reached unrealistic values and were correlated (O$_3$)
or anti-correlated (H₂O) with NO₂. These interfering gases were therefore omitted hereinafter.

Accurate wavelength calibration is an important prerequisite for the DOAS analysis. As indicated above, raw DN (digital number) data were used to keep the highest sensitivity of the measurements for the NO₂ retrieval. The spectral calibration was therefore performed with the QDOAS algorithm itself. A high-resolution solar spectrum (Chance and Kurucz, 2010) was applied to obtain spectral calibration information which was subsequently used to convolve and shift the high-resolution absorption cross sections to the APEX specifications. Two exemplary (pre-processed) APEX spectra from the VNIR detector are illustrated in Fig. 2. The corresponding slant column fit of the pixel over the residential area is presented in Fig. 3 which also gives an impression of the sensor’s spectral resolution. The quality of the DOAS fit depends on the spectral and radiometric characteristics of the instrument. The spectral calibration step in QDOAS disclosed some differences to the pre-flight reported APEX specifications (Sect. 2.1 and Table 1) in the 470–510 nm wavelength range. The DOAS analysis indicated that the individual channels are positioned roughly 0.6 nm higher than reported pre-flight and the (over wavelength) averaged across-track difference of their corrected position (spectral smile) is ~ 0.25 nm. This is in line with recent findings from an in-flight and scene-based APEX performance assessment (D’Odorico et al., 2011). Furthermore, the QDOAS wavelength calibration and slit function characterisation point to a FWHM about double the spectral resolution measured in the laboratory at the CHB. The reasons for this discrepancy are not fully understood yet, and further analysis is currently being carried out. It is noteworthy that this increase in FWHM on the other hand reduces problems associated with spectral undersampling expected for the smaller spectral resolution defined in the pre-flight specification. Finally, note that negative dSCD can occur when the SCD from the reference spectrum is larger than that from the fitted spectrum or due to noise.

3.2 Air mass factor calculation

The AMF expresses the ratio between slant and vertical column of a trace gas:

$$\text{SCD} = \text{VCD} \times \text{AMF}$$

and is a measure of the average backscatter path through the atmosphere of the photons observed by the sensor. The AMF can be calculated as follows (Palmer et al., 2001; Boersma et al., 2004):

$$\text{AMF} = \frac{\sum m_L(\hat{b}) \chi_a \cdot L}{\sum \chi_a \cdot L}$$

where the subscript $L$ denotes a specific atmospheric layer and $m_L$ is the (box) air mass factor per layer. Besides the layer subcolumns of the a priori NO₂ profile ($\chi_a \cdot L$), the AMF depends on forward model parameters ($\hat{b}$) such as the solar and viewing zenith and azimuth angles, surface reflectance, aerosol extinction profile, and surface pressure. The box air mass factors were calculated using the linearized discrete ordinate radiative transfer model (LIDORT, Spurr, 2008):

$$m_L = -\frac{1}{I} \frac{\partial I}{\partial \tau_L}$$

where $I$ is the intensity of the backscattered radiance and $\tau_L$ the optical depth of layer $L$.

With regard to radiative transfer computations, the surface reflectance for every pixel was derived from re-binned and atmospherically corrected APEX data themselves for the central wavelength of the fitting window (490 nm). Surface height was taken from the digital elevation model...
DHM25 previously projected to the raw geometry of the individual flight lines. Surface pressure for every pixel was subsequently obtained applying the US Standard Atmosphere 1976 (http://www.pdas.com/coesa.html, last access: 12 March 2012). The a priori NO$_2$ profile was taken from the EURAD chemical transport simulations (http://www.eurad.uni-koeln.de/, last access: 12 March 2012) over Switzerland at 5 × 5 km$^2$. The coarse resolution profile was subsequently scaled to the corresponding surface height of the aggregated APEX grid cell according to Zhou et al. (2009). Aerosol optical depth (AOD) at 500 nm was taken from the Aerosol Robotic Network (AERONET, Holben et al., 1998) site Laergeren which is approximately 20 km from the city of Zurich. The AOD was converted to an extinction profile for every pixel assuming an exponential decrease with height and a scale height of two kilometers. The box air mass factors from three exemplary (aggregated) APEX pixels are depicted in Fig. 4. APEX’s sensitivity toward a NO$_2$ signal is highest in the atmospheric layer below the aircraft (red horizontal line) and is decreasing toward the surface and toward higher atmospheric layers. Among all the above-mentioned parameters, the surface albedo has the largest impact on the AMF. For example, the bright surface (albedo of 0.27) of case (c) in Fig. 4 highly increases the APEX sensitivity toward surface NO$_2$. This is supported by Fig. 5 which displays the computed AMF versus surface albedo from the easterly morning flight line and which emphasizes the importance of a good quality surface albedo data set in our NO$_2$ VCD determination.

Since we are using earthshine spectra as reference, the results of the DOAS fit are differential slant column densities ($d$SCD) which can be written as

$$d$SCD = (VCD$_{P}$ × AMF$_{P}$ + SCD$_{STR}$) − (VCD$_{R}$ × AMF$_{R}$ + SCD$_{STR}$) \tag{4}$$

where the subscripts $P$ and $R$ refer to tropospheric quantities under polluted and clean (reference) conditions, respectively. The stratospheric contribution to dSCD can reasonably be assumed to be constant for the polluted and reference spectra in our study region. Hence, the (additive) stratospheric SCD (SCD$_{STR}$) cancels out on the right hand side of Eq. (4) and AMF$_{P}$ and AMF$_{R}$ are calculated using only atmospheric levels up to the tropopause in Eq. (2). Rearranging Eq. (4) finally yields the VCD$_{P}$:

$$VCD_{P} = \frac{d$SCD + VCD_{R} × AMF_{R}}{AMF_{P}} \tag{5}$$

where the VCD$_{R}$ has to be estimated. In our case we assume $1 \times 10^{15}$ molec cm$^{-2}$ which is in the range of previously reported rural/background tropospheric columns for European summer conditions from OMI data and an ensemble of regional air quality models (Huijnen et al., 2010). Note that dSCD varies primarily due to different NO$_2$ below the aircraft, mainly in the boundary layer where NO$_2$ profiles usually peak close to the source. Free tropospheric NO$_2$ and particularly the tropospheric NO$_2$ above the aircraft is expected to contribute only very little as it is probably similar for the reference and sample observations, and in addition the NO$_2$ concentrations and above-aircraft AMF values are low.

### 3.3 Post-processing

Missing pixels due to a failed dSCD fit were replaced with the mean NO$_2$ VCD value of the nearest neighbours (less than 5% of all retrievals were affected). The resulting NO$_2$ maps were subsequently de-striped in order to correct for artefacts introduced by parameters varying (randomly) across-track, e.g. the reference VCD or remaining
sensor artefacts. Assuming that NO\textsubscript{2} VCD averaged in along-track (column) direction varies smoothly across-track (row), a fifth degree polynomial was fitted to the column averages. The residuals per column were finally subtracted from the initially retrieved NO\textsubscript{2} VCD field.

4 Results

The capability of APEX to sense NO\textsubscript{2} and the NO\textsubscript{2} product itself are assessed and discussed in the following subsections. The NO\textsubscript{2} SCD fitting and the two-dimensional VCD distribution are analyzed in detail. Further, the APEX NO\textsubscript{2} maps are compared with ground-based in-situ measurements and yearly averaged modelled NO\textsubscript{2} surface concentration fields.

4.1 SCD analysis

Histograms of derived NO\textsubscript{2} dSCD and the corresponding dSCD error are depicted in Fig. 6. Selected statistical parameters obtained by the DOAS analysis can be found in Table 2. The average dSCD for the morning flight lines is \(9.2 \times 10^{15} \text{ molec cm}^{-2} (\pm 8.31 \times 10^{15} \text{ molec cm}^{-2})\) and is about 2.5 times higher than the average dSCD of the afternoon flights \((3.87 \times 10^{15} \text{ molec cm}^{-2} \pm 5.09 \times 10^{15} \text{ molec cm}^{-2})\). The minimum dSCD is on the same order for both overflight times \((-1.58 \times 10^{16} \text{ molec cm}^{-2} \text{ and } -1.68 \times 10^{16} \text{ molec cm}^{-2}, \text{respectively})\), whereas the maximum dSCD for the morning is almost twice the afternoon value \((4.71 \times 10^{16} \text{ molec cm}^{-2} \text{ versus } 2.86 \times 10^{16} \text{ molec cm}^{-2})\). Several random and systematic error sources affect the APEX-based dSCD fitting, e.g. instrumental noise, wavelength calibration, or temperature dependency of the absorption cross sections (Boersma et al., 2004). A detailed assessment of these error sources is beyond the scope of this study. Rather, we concentrate here on the overall dSCD error. About 14% of all retrievals lead to negative dSCD for the morning overflight and 22% for the afternoon overflight (Table 2). The averaged fitting error for the morning is \(2.37 \times 10^{15} \text{ molec cm}^{-2} (\pm 6.47 \times 10^{14} \text{ molec cm}^{-2})\) and for the afternoon \(2.42 \times 10^{15} \text{ molec cm}^{-2} (\pm 4.53 \times 10^{14} \text{ molec cm}^{-2})\). This corresponds to 24% and 47% of the respective absolute dSCD. The absolute dSCD fitting errors and their standard deviations are very similar for the morning and afternoon results. In general, the dSCD errors do not reveal any geographical pattern (not shown) like e.g. correlation with albedo or surface type. SCD errors on the order of \(0.7 \times 10^{15} \text{ molec cm}^{-2}\) are reported for different satellite NO\textsubscript{2} retrievals in the literature (Boersma et al., 2004, 2007; Valks et al., 2011; Valin et al., 2011). These lower SCD errors can be explained by the better characteristics of these sensors specifically designed for trace gas remote sensing (e.g. SSI, FWHM, SNR, fitting window at shorter wavelength with stronger NO\textsubscript{2} signal possible) as compared to APEX (cf. Sect. 2.1 and Table 1).

4.2 NO\textsubscript{2} spatial distribution

The spatial distributions of NO\textsubscript{2} VCD over Zurich for the morning and afternoon overflights are depicted in Fig. 7b and c, respectively. In addition, a comparison between the morning APEX NO\textsubscript{2} VCD and modelled yearly averages of surface NO\textsubscript{2} concentrations for 2010 is presented in Fig. 8. The simulation is based on a high-resolution (100 \(\times\) 100 m\(^2\)) emission inventory combined with a Gaussian plume dispersion model (SAEFL, 2004). Overall, APEX NO\textsubscript{2} VCD are considerably higher for the morning than for the afternoon overflights. Especially the morning mosaic of the three flight lines reveals very distinct and plausible spatial NO\textsubscript{2} patterns. Higher NO\textsubscript{2} VCD can be found in residential areas (specifically over the city), over motorways, and around the international airport of Zurich to the north of the scene. Interestingly, the enhanced NO\textsubscript{2} VCD values in the northeastern part of the scene correspond to a large shopping area west of the motorway A4 (cf. Fig. 1) which is known to have a large traffic volume during shopping hours, especially on Saturdays. Higher NO\textsubscript{2} in the southwestern part of the image area corresponds to a motorway junction where three motorway tunnels intersect open air. In general, the motorways as a prominent source of NO\textsubscript{2} do not show up as clearly in the APEX NO\textsubscript{2} VCD maps as in the model surface NO\textsubscript{2} (Fig. 8). One has to consider that these data were acquired on a Saturday where traffic reveals different characteristics than during weekdays, e.g. generally less traffic volume with more private transport, less commuter traffic, and fewer trucks. Lower NO\textsubscript{2} mainly occurs in remote and/or forested areas, e.g. at the Uetliberg mountain range to the south-west of the city or the forested area just east of the airport. Further, the three different flight lines per mosaic generally superimpose well. This is underlined by Fig. 9 which shows the N-S transects of the APEX NO\textsubscript{2} VCD in the overlapping region (cf. map in Fig. 1) of the central and eastern morning overpasses. The two curves are in very good agreement with a correlation coefficient of around 0.95. However, the values from the central line are biased against the eastern line. The mean difference between these two curves is around \(1.9 \times 10^{15}\) (or 28\%) which might be due to several reasons. For example, the NO\textsubscript{2} columns in the “pollution-free” areas where the reference spectra are selected can slightly differ due to terrain variations such that using reference spectra from lower altitudes (higher NO\textsubscript{2} columns) leads to relatively lower NO\textsubscript{2} VCD than those from higher altitudes. In addition, the two flight lines do not observe exactly the same air mass at the same location due to the varying observation geometry and the time lag between the two measurements.

Overall, the spatial NO\textsubscript{2} distribution and the above-mentioned NO\textsubscript{2} features are in good agreement with the modelled surface concentration in Fig. 8 underlying the capability of APEX to detect tropospheric NO\textsubscript{2}. It should be kept in mind that the APEX data represent the NO\textsubscript{2} distribution at a given time under a specific weather situation while the
Table 2. Statistics for the NO\textsubscript{2} VCD retrievals from the morning and afternoon overflight (OF = overflight, Av. = average, Stddev = standard deviation). The units are molecules cm\textsuperscript{-2} where not otherwise indicated.

<table>
<thead>
<tr>
<th></th>
<th>Av. dSCD</th>
<th>Stddev dSCD</th>
<th>Min dSCD</th>
<th>Max dSCD</th>
<th>Av. dSCD error</th>
<th>Stddev dSCD error</th>
<th>% negative dSCD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Morning OF</td>
<td>9.20 × 10\textsuperscript{15}</td>
<td>8.31 × 10\textsuperscript{15}</td>
<td>−1.58 × 10\textsuperscript{16}</td>
<td>4.71 × 10\textsuperscript{16}</td>
<td>2.37 × 10\textsuperscript{15} (24 %)</td>
<td>6.47 × 10\textsuperscript{14}</td>
<td>14</td>
</tr>
<tr>
<td>Afternoon OF</td>
<td>3.87 × 10\textsuperscript{15}</td>
<td>5.09 × 10\textsuperscript{15}</td>
<td>−1.68 × 10\textsuperscript{16}</td>
<td>2.86 × 10\textsuperscript{16}</td>
<td>2.42 × 10\textsuperscript{15} (47 %)</td>
<td>4.53 × 10\textsuperscript{14}</td>
<td>22</td>
</tr>
</tbody>
</table>

Fig. 6. Histograms of dSCD (upper panel) and dSCD errors (lower panel) for the morning (left panel) and afternoon (right panel) APEX flights. Note the much shorter scale of the x-axis in the lower panel.

model distribution is an annual mean estimate. APEX data also represent vertical below-aircraft columns, whereas the model data are concentrations at the surface. A closer look at some exemplary and specific areas is given in Sect. 4.3.

Differential SCD, AMF as well as the VCD are illustrated in Fig. 10 for the central morning flight stripe. As already underlined by Fig. 5, the surface reflectance has the largest impact on the AMF. For example, the lowest AMF values can be found over the dark forested areas and the highest ones over man-made structures such as buildings and roads. In our study region, therefore, significant correlation between bright surfaces and enhanced NO\textsubscript{x} due to emissions from motorways and residential or industrialized areas can be found. Darker surfaces, in contrast, corresponded to comparatively clean vegetated areas. Without accounting for the varying surface reflectance, the contrast between polluted and clean areas would therefore be clearly overestimated. Figure 10 further demonstrates that spatially varying AMF also has an impact on small-scale NO\textsubscript{x} features like e.g. in the lower (southern) part of the flight stripe around the in situ site Wettswil Filderen (cf. map in Fig. 1). However, overall the largest part of the VCD variability can clearly be linked to the variability of the dSCD.

In general, spatial gradients of NO\textsubscript{2} VCD are more pronounced for the morning, but enhanced NO\textsubscript{2} VCD around the airport, around the motorway junction, over the city and decreased NO\textsubscript{2} VCD in remote areas and over the lake are also detectable in the afternoon maps.

The above-mentioned differences are due to the different meteorological conditions in the morning and afternoon which affected the transport and dilution of NO\textsubscript{x} downwind from its sources and possibly also its lifetime. The diurnal evolution of surface NO\textsubscript{2} concentrations measured at eight NABEL and OSTLUFT sites is plotted in Fig. 11. The values during the APEX overflights range from low (4.1 µg m\textsuperscript{-3}) to polluted (43.55 µg m\textsuperscript{-3}) depending on the location and time of day. All sites show a strong decrease of NO\textsubscript{2} during the morning hours and more or less stable concentrations in the afternoon (after about 12:00 UTC). This behaviour is typical for sunny summer days and is to a large extent driven by the evolution of the boundary layer leading to decreased surface concentrations in the afternoon due to enhanced vertical mixing. However, variations in vertical mixing may not...
explain the differences in the vertical columns observed by APEX. One reason for lower VCD in the afternoon could be enhanced chemical loss of NO$_2$ due to the reaction with the hydroxyl radical (OH). However, since the solar zenith angle was approximately the same during the morning and afternoon overpasses and since OH-levels depend on NO$_x$ concentrations in a strongly non-linear way (e.g. Jaeglé et al., 1998), it is not clear whether OH-levels were on average higher or lower in the afternoon.

A more likely explanation for the lower VCD in the afternoon is stronger dilution due to stronger winds. Wind speed and direction measured at two different sites maintained by the Swiss Federal Office of Meteorology and Climatology (Fig. 12) show that the meteorological situation changed distinctly shortly after the morning APEX flights, i.e. wind speed sharply increased by nearly four times until noon and the wind direction switched from northwest to northeast typical of a bise situation. The stronger winds lead to a stronger dilution and more rapid transport of NO$_2$ to regions.
Fig. 9. North-south transect of retrieved NO$_2$ VCD of the overlapping region of the central (green line) and eastern (black line) morning overflights (the exact position can be found in the map in Fig. 1).

Fig. 10. Differential SCD (a), AMF (b), and VCD (c) of the central flight stripe from the morning overflights (cf. Figs. 7 and 8).

downwind of its main emission sources. The lower winds in the morning may also explain the larger NO$_2$ variability and therewith better detectability of NO$_2$ sources for the APEX sensor during the morning overflights. These meteorological conditions together with the diurnal cycle of NO$_2$ are also of particular interest with regard to the planning of future flight campaigns.

Different parameters related to the sensor characteristic and retrieval algorithm may also partly account for the reported differences. The selected reference spectra should ideally be determined from a pollution-free area with a high albedo and therefore high SNR. However, such an ideal reference spectra could not be found in the current test area. The used reference spectra over a forested area characterized by a relatively low albedo might cause a decreased dSCD fitting quality (cf. Sect. 4.1). Further, the contrast in NO$_2$ between the area where the reference spectra were chosen and the other areas is rather small for the afternoon case for reasons discussed above. This reduced contrast leads to distinctively more negative dSCD derived for the afternoon (cf. Sect. 4.1).

Forward model parameters vary during the course of the day and therewith also the AMF. The average AMF increased from 1.51 in the morning to 1.71 in the afternoon. The major part of this change can be assigned to an increase in the aerosol optical depth from 0.19 to 0.26 (at 500 nm) based on sun photometer measurements from the nearby AERONET site Laegeren. The solar zenith angles were quite similar for the morning (at 08:00 UTC $\sim$ 48.2$^\circ$) and afternoon (around 15:30 UTC $\sim$ 52.9$^\circ$), but these differences nevertheless slightly affect the AMF. Uncertainties in AMF are known to be a major source of uncertainty in NO$_2$ VCD retrievals. For example, several studies (Boersma et al., 2004, 2007; Zhou et al., 2010) estimated AMF uncertainties in the range of 30% for OMI and SCIAMACHY. A somewhat lower AMF uncertainty can be expected in the presented retrieval because we do not have to deal with cloud contamination which is an important part of the AMF uncertainty (Boersma et al., 2004; Popp et al., 2011). An additional influence on the accuracy of the AMF can be expected from retrieval input parameters with insufficient spatial resolution not matching the high resolution of APEX. Heckel et al. (2011), for example, studied the impact of coarse-resolution retrieval input parameters (a priori NO$_2$ profile, surface reflectance, and aerosol information) on satellite retrievals of tropospheric NO$_2$ VCD with significantly smaller pixel size. They identified the a priori profile and surface albedo to have the largest impact on the retrieval uncertainty. In our approach, we therefore derive the surface reflectance directly from the APEX data at high resolution. Profiles of NO$_2$ and aerosols, on the other hand, are taken from coarse-resolution data sets which represents an important remaining error source. For an improved retrieval and error quantification, future flight campaigns should therefore aim at flying vertical NO$_2$ and aerosol profiles with complementary in situ instrumentation. Aerosol information (e.g. AOD) can potentially be derived from APEX data itself in the future (Seidel...
et al., 2011). The high spatial resolution of the APEX data also has implications on the radiative transfer calculations. For example, photons from neighbouring pixels can be scattered into the instantaneous field-of-view. Considering such three-dimensional effects in the radiative transfer computations would potentially improve future APEX NO₂ retrievals.

4.3 Examples of source identification

A more detailed view of specific NO₂ features detected by APEX is illustrated in the basis of three examples in Fig. 13 where the APEX VCD are shown in the left panel (Figs. 13a, d, g), the yearly averaged surface concentration in the middle panel (Figs. 13b, e, h), and some specific zoom-ins in the right panel (Figs. 13c, f, i). The first example (Fig. 13a and b) focuses on an area around the southeastern part of Zurich international airport. The enhanced surface NO₂ concentrations around the end of the two runways are well captured by the APEX VCD map. These two features are most likely caused by emissions from aircraft landing and taking off. The aircrafts took off towards the south on 26 June 2010 (cf. http://www.flughafen-zuerich.ch, last access: 11 July 2012). The enhanced NO₂ VCD at the end of the north-south runway thus likely reflects the increased NO₂ emissions by aircraft during take-off. Additionally, the APEX RGB image (Fig. 13c) acquired in the morning shows different aeroplanes in motion around the hangar and one at the engine test stand (encircled red) close to the area of maximum NO₂ VCD in this example. In contrast, APEX VCD does not mirror the enhanced surface NO₂ concentrations above the motorway A51 which might be explained by the different traffic characteristics on a Saturday morning. The second example (Fig. 13d and e) displays the area around the waste incinerator Hagenholz situated between the city and the airport (cf. map in Fig. 1). The plume of the waste incinerator pointing in the southwest direction does clearly emerge in the RGB subscene (Fig. 13f). The increased NO₂ around the chimney (red and white areas) is well detectable in the APEX VCD maps while the representing surface concentrations does not reflect the emissions from this elevated source. The third example shows an area around the lower part of Lake Zurich, including the national motorway A3 and the harbour/dockyard at the shore of Lake Zurich. In general, the spatial gradients of the yearly average surface NO₂ distribution are reproduced well by APEX VCD. Also the NO₂ abundance from traffic emissions along the motorway A3 is detectable in this example. In contrast to the first example, this specific leg of the motorway A3 is located in a N-S oriented valley, which likely channelled emitted NO₂.

The smeared-out patterns of NO₂ over and around the lake basin are a result of locally transported NO₂. The NO₂ hot spot in the southern part of example three is situated over the harbour/dockyard area of Zurich. The APEX RGB image (Fig. 13i) shows a vessel on the ramp of the dockyard during the morning overflight. However, if specific engine tests were performed or not is unknown.

4.4 Comparison to ground-based measurements

The APEX-derived NO₂ VCD are subsequently compared to the ground-based in situ measurements. Such a comparison is not straightforward since APEX NO₂ VCD is a columnar product (in molec cm⁻²) while the in-situ measurements represent a trace gas concentration at a single point (in µg m⁻³). Therefore, we convert the in situ measured concentrations to an “in situ VCD” (VCD₉S) assuming that the a priori NO₂ profile correctly described the true NO₂ profile:

\[
VCD₉S = \frac{VCD_{AP} \times NO₂_{28}}{NO₂_{AP}}
\]

where VCD₉S is the total of the sub-columns of the a priori NO₂ profile from the ground to the flight altitude, NO₂₂₈ the concentration measured at the ground and NO₂₉₈ the concentration of the a priori profile in the lowest layer.

The comparison is shown in Fig. 14 where the APEX retrieved NO₂ VCD are plotted as a function of the VCD₉S at the eight NABEL and OSTLUFT sites. The Pearson correlation coefficients for the morning and afternoon results separately are almost identical (R = 0.61) suggesting that the variability between the in situ sites corresponds to some extent to the spatial pattern observed by APEX. When disregarding the match-ups corresponding to the in situ sites located immediately next to a road (cross symbols), a rather good agreement is found and the APEX-derived NO₂ VCD are in a similar range as the VCD₉S, and the morning and afternoon results fit well. However, almost all match-ups are located below the 1:1 line suggesting that the a priori NO₂ profile is not perfectly suitable to convert the in situ measurements to a columnar quantity. This is particularly true for the sites Opfikon, Stampfenbach, and Schimmelstrasse located directly at a road where the true atmospheric NO₂
profile is probably much more strongly peaked at the surface than the a priori profile leading to a strong overestimation of the NO$_2$ VCD constructed from the in situ data. This further underlines the need for accurate vertical profile information, e.g. by flying vertical NO$_2$ and aerosol profiles with complementary in situ instrumentation. Finally, the chosen approach to construct “in situ VCD” from surface measurements cannot account for the influence of upstream sources present in the APEX-derived column which partly explains the remaining discrepancies.

5 Conclusions
We presented the first highly detailed two-dimensional NO$_2$ fields derived from the airborne APEX imaging spectrometer taking advantage of its unique combination of high spectral and spatial resolution and high number of pixels. Radiance data were acquired twice over Zurich, Switzerland, in the morning (10:00 LT) and afternoon (17:30 LT) of a cloud-free summer day in June 2010. The main results and conclusions from this study can be summarized as follows:

1. Although not primarily designed to retrieve atmospheric constituents, APEX is clearly sensitive toward NO$_2$ above typical European background concentrations. The DOAS analysis revealed a mean dSCD of
of longer integration times were found to be less critical for this application. On the other hand, a longer integration time will potentially lead to signal saturation in the NIR spectral region which may cause problems for the detector read-out and the smearing correction. Specific NIR filters might be used to compensate this effect. Further, pollution-free regions should be present in the imaged area in order to select appropriate reference spectra. These spectra should, if possible, be acquired over bright surfaces. The spectral calibration and slit function characterisation with QDOAS pointed to a doubling of the spectral resolution in-flight compared to pre-flight laboratory characterisation. Possible causes for this discrepancy are currently under investigation, and such findings might help to optimize future instrument setups. Finally, future flight campaigns should also aim at measuring NO₂ and aerosol profiles which would valuably support the retrieval as well as the interpretation of the results.

The findings of this study clearly reveal a high spatio-temporal variability of NO₂. Airborne-based NO₂ retrieval, such as presented in this study, allows detecting NO₂ emission sources, provides valuable input for NO₂ emission modelling, and helps to strengthen the link between air quality models and satellite NO₂ products. This all together enables to increase the knowledge on processes and characteristics of NO₂ tropospheric distribution. Additional and complementary APEX flights have been carried out in the meantime and others are currently in planning. The growing database of observations will be used to enhance the APEX NO₂ retrievals but also to gain knowledge on spatio-temporal NO₂ distribution.

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