

Validation of Ozone Monitoring Instrument Nitrogen Dioxide Columns

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Abstract. We review the nitrogen dioxide (NO₂) data product, which is based on measurements made in the spectral region 415–465 nm by the Ozone Monitoring Instrument (OMI) on the EOS-Aura Satellite. A number of ground- and aircraft-based measurements have been used to validate the data product's three principal quantities: stratospheric, tropospheric, and total NO₂ column densities. It is found that OMI overestimates the stratospheric column by about 10%, and underestimates both the total and tropospheric columns by 15–30%. Because some of the techniques have not themselves been validated, and because of problems inherent in comparing point measurements from the ground to large area averaged satellite measurements, and because some of the measurements have been made only over short periods of time, validation results are sometimes only qualitative.

1. Introduction

The Ozone Monitoring Instrument (OMI) is a space borne spectroradiometer that uses a 2-dimensional CCD array detector to simultaneously measure the spectra of the earth shine radiance at large number of viewing angles, approximately transverse to the Aura spacecraft's flight track. OMI measures in three broad spectral regions (UV-1, UV-2, and VIS), with a spectral resolution on the order of 0.5 nm. Applying spectral fitting techniques to the OMI data permits the simultaneous retrieval of a wide

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22 range of atmospheric trace gas concentrations as well as
23 cloud and aerosol properties and loadings. Among the
24 trace gases that can be retrieved, ozone (O_3) and nitro-
25 gen dioxide are identified as essential measurements, both
26 for the ongoing monitoring of the Earth's stratospheric
27 ozone layer and for the monitoring of tropospheric air
28 quality. A more extensive discussion of the OMI instru-
29 ment itself can be found in *Levelt and Bhartia* [2007].

30 The OMI NO_2 data production algorithm is designed to
31 retrieve total vertical column densities of NO_2 and sepa-
32 rate stratospheric and tropospheric column densities; this
33 enables the improvement in the calculation of the total
34 vertical column. The stratosphere-troposphere separa-
35 tion is achieved using a low-pass spatial filtering tech-
36 nique; the small-gradient portion of the initial estimate of
37 the total NO_2 field is identified as the background strato-
38 spheric field. Measurements that exceed the constructed
39 stratospheric field are taken to indicate significant tro-
40 pospheric pollution. This separation is important (and
41 possible) because the chemistry and dynamics of NO_2 are
42 different between the stratosphere and the troposphere.
43 Accurate measurements of the tropospheric NO_2 are sig-
44 nificant for the characterization of air quality, a primary
45 objective of the Aura and OMI missions.

46 A number of efforts toward NO_2 validation have been
47 initiated, in which measurements are made coincident
48 with OMI overpass measurements. The purpose of this
49 paper is to provide an overview of results from these ef-
50 forts. A variety of instruments and techniques have been
51 used, each with its characteristic sensitivity to strato-
52 spheric, tropospheric, or total column NO_2 . This paper
53 will address the advantages and, in some cases, the limi-
54 tations of the various measurements.

55 Several of the techniques described are new, and have
56 not been well-validated. In addition, there is an essential
57 difference between observations of NO_2 taken from the
58 ground, and observations averaged over a satellite field
59 of view (FOV). Spatial inhomogeneity, characteristic of
60 airborne constituents emitted at (possibly moving) point
61 sources, and subject to surface-level winds, implies that a
62 single point measurement will often not be a representa-
63 tive sample within a "collocated" a satellite FOV covering
64 a region of the order of several hundred square kilome-
65 ters. Monthly average comparisons of ground-based and
66 satellite measurements can remove much of the variabil-
67 ity due to FOV-point measurement differences. A pre-
68 liminary measurement of horizontal inhomogeneity in the
69 NO_2 field is presented in the DANDELIONS overview paper
70 [*Brinkma et al.*, 2007], using a set of simultaneous tropo-
71 spheric NO_2 measurements made at different azimuths.
72 *Veeffkind et al.* [2007] show a comparison of regrid-
73 ded OMI NO_2 data with ground-based observations by the Dutch
74 national air quality network. This network distinguishes
75 regional stations, and city and street stations, which are
76 close to source regions. For the period of the satellite
77 data, NO_2 reported by 35 stations around the Nether-
78 lands, averaged between 11:00 and 14:00 h local time,
79 was compared to the collocated OMI measurements. A
80 strong correlation ($R = 0.94$) between the satellite data
81 and the regional station data was found. By contrast,
82 correlations with urban stations are weak, because local
83 conditions may vary strongly over a few hundred meters,
84 far smaller than the spatial resolution of OMI. Because
85 of both the novelty of some of the techniques, and such
86 spatial-scale effects, specific results are sometimes limited
87 to qualitative, or order-of-magnitude, conclusions.

88 Among the validation studies discussed here are
89 ground based observations made within the SAOZ and
90 DOAS networks. These are zenith-sky, twilight measure-

91 ments, which are sensitive to the stratospheric NO₂ col-
92 umn (as explained in section 4.1.1). In addition, the loca-
93 tions are very often pristine areas, or at elevation. Even if
94 the instrumentation were sensitive to tropospheric NO₂,
95 the lack of pollution would lead to measurements domi-
96 nated by the stratospheric NO₂ amount.

97 To focus on polluted areas, where satellite NO₂ re-
98 trievals are most challenging, novel, or as yet unvalidated
99 techniques must be used.

100 Measurements of scattered light by the MultiAxial
101 DOAS (MAX-DOAS) technique, using a range of viewing an-
102 gles, from nearly horizontal through zenith, are sensitive
103 to the tropospheric part of the column, and provide both
104 total and tropospheric NO₂ amounts. We present results
105 from MAX-DOAS measurements taken in a polluted area,
106 but away from immediate local sources, in section 4.2.1.

107 Direct-sun ground based measurements, made with
108 a Brewer spectrophotometer and with newly-developed
109 direct-sun instruments, including a high-resolution
110 Fourier-transform UV-FTS technique, and a number of
111 direct-sun DOAS-type measurements, are sensitive to the
112 total NO₂ column. We will briefly review some prelimi-
113 nary results from these methods in section 4.3.

114 Validation of the OMI NO₂ data should take account of
115 the sensitivity of the numerous geophysical and geomet-
116 ric algorithmic inputs. These include the *a priori* profile
117 shapes, surface albedo, and measured and assumed cloud
118 properties. These, in particular, greatly affect the air
119 mass factors (AMF; the ratio of slant-column density of
120 the absorber along the optical path to the vertical column
121 density) the algorithm calculates. Tropospheric NO₂ pro-
122 files have been measured with lidar in the Netherlands,
123 during a number of days in September 2006, and with
124 airborne instrumentation during various validation cam-
125 paigns.

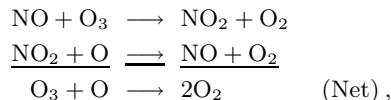
126 During the Polar Aura Validation Experiment (PAVE)
127 (flights from New Hampshire, January and February
128 2005) the TD-LIF instrument [Thornton *et al.*, 2000;
129 Cleary *et al.*, 2002] was used for *in situ* sampling of NO₂,
130 during the aircraft flights. The NASA DC-8 performed
131 two flight legs at 300 m altitude, near the top of the
132 boundary layer. When flights entered the boundary layer,
133 strongly enhanced concentrations of NO₂ were found.

134 During the INTEX-B campaign (flights from Houston,
135 Texas, March 2006 and from Honolulu, Hawaii, and An-
136 chorage, Alaska, April and May 2006) the TD-LIF instru-
137 ment measured NO₂ *in situ*. Spirals were flown by the
138 NASA DC-8 during several flights in spatial and tempo-
139 ral collocation with OMI observations.

140 Besides INTEX-B, a small number of other airborne
141 campaigns have been carried out, measuring NO₂ *in*
142 *situ*, and have been applied to satellite validation [He-
143 land *et al.*, 2002; Martin *et al.*, 2006].

1.1. Nitrogen dioxide in the stratosphere

144 Nitrogen dioxide participates both directly and indi-
145 rectly in the catalytic destruction of ozone in the strato-
146 sphere. Direct ozone destruction occurs *via* the reactions



147 while NO₂ concentrations indirectly control ozone loss
148 through other catalytic cycles by controlling, for exam-
149 ple, the distribution of chlorine between its catalytically

150 active (ClO) and inactive, reservoir (ClONO₂) species:

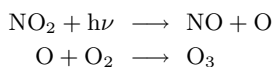


151 In the stratosphere, NO₂ concentration has a distinctive
152 diurnal cycle that is dependent on the reactions
153 shown in Figure 1. At night, all the photolysis reactions
154 stop, shifting the steady state to NO₂. NO₂ is converted,
155 through ozonolysis, to NO₃, which can further combine
156 with NO₂ to form N₂O₅. This results in a slow decrease
157 of NO₂ over the course of the night. When the air is again
158 sunlit, the N₂O₅ rapidly redissociates to NO₂ and NO₃,
159 which photolyzes instantaneously, mostly to NO. Mean-
160 while, NO₂ photolyzes very rapidly, and so decreases
161 very rapidly at sunrise. While in daylight, the domi-
162 nant processes are the interconversion between NO and
163 NO₂. There are a number of specific mechanistic path-
164 ways that can contribute to this interconversion. The
165 most typical atmospheric states (conditions of tempera-
166 ture and ozone concentration, latitude, and season) lead
167 to a slow increase in NO₂ concentration over the course
168 of the daylight hours. At sunset, the photolysis reactions
169 again switch off, and the NO₂ concentration rises rapidly.
170 In addition to the chemical and photochemical processes,
171 transport by the winds, particularly in the vicinity of the
172 polar jets may mean that the air that one is measuring
173 has not had the photochemical history one would expect,
174 based on location and local time, alone. Some caution is
175 therefore needed in matching satellite measurements to
176 the ground based measurements.

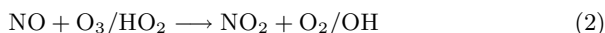
177 The time-dependence of the stratospheric NO₂ con-
178 centration has important implications for the validation
179 of the space based NO₂ measurements. If the ground
180 based measurements are not collocated in time with the
181 OMI measurements, they need to be corrected, using pho-
182 tochemical and transport models, to account for the
183 time difference. In addition, if the ground-based mea-
184 surements entail an optical path that is more horizontal
185 than vertical, view and solar geometries must be taken
186 into consideration when identifying “collocated” mea-
187 surements.

1.2. Nitrogen dioxide in the troposphere

188 In the troposphere, nitrogen oxides are a significant
189 contributor to poor air quality. Both NO and NO₂ are
190 harmful to lung tissue, and, as a powerful oxidizing agent,
191 NO₂ is harmful to biological tissue generally. Besides
192 its direct effects, photolysis of NO₂ contributes to ozone
193 production according to



194 Nitrogen oxides are produced in high-temperature
195 processes in the atmosphere, most notably in combus-
196 tion (fossil fuels and biomass burning) and in lightning.
197 Nearly all the NO_x (NO + NO₂) that is significant for
198 human health is produced by industrial and urban activi-
199 ty, including transportation and power generation. As
200 a rule, the higher the combustion temperature, the more
201 NO_x is produced; the primary reactions necessary for
202 NO_x production are the thermolysis of N₂ and O₂. The
203 most common species directly formed in combustion is
204 NO, however NO readily oxidizes in air to NO₂:



205 Gaseous NO₂ is red in color, and gives rise to the char-
206 acteristic brownish cast of polluted air. NO₂ is removed
207 through conversion to HNO₃, nitric acid, which readily
208 dissolves in any available water droplets. NO₂ plumes are
209 detected only up to about 100 km from their industrial
210 or urban source. In the neighborhood of industrial or
211 urban sources, there is a distinct diurnal pattern in the
212 production and loss of NO_x. The diurnal signal at any
213 location, particularly in the boundary layer, is the result
214 of a complex interplay between the emission source field
215 in space and time, photochemical effects, advection by
216 boundary layer winds, and the concentrations of chem-
217 ical sinks for NO_x species. As mentioned before, these
218 also give rise to spatial inhomogeneities on a sub-100 m
219 scale. At mid- to low-latitudes, where a polar-orbiting
220 satellite passes over a given location is only once or twice
221 a day, the satellite only sees a “snapshot” of the state of
222 the polluted atmosphere at the overpass times. In the
223 mid-to-upper latitudes, inconsistent measurements, from
224 one orbit to the next, over some location may well result
225 from significant changes in the NO₂ concentrations over
226 the intervening 100 minutes, as well as from other rapid
227 geophysical changes, e.g. in cloud cover.

1.3. OMI measurement of NO₂

228 The Aura satellite is a polar-orbiting, sun-synchronous
229 satellite, whose orbital period is 99 minutes. Aura flies
230 over the entire surface of the Earth every 14–15 orbits.
231 Using the 2-dimensional CCD array detector, with pixel
232 binning factors chosen to optimize the signal-to-noise ra-
233 tio, the instrument measures earthshine radiance spectra
234 simultaneously in 60 effective fields of view (FOV), ap-
235 proximately transverse to the flight track, every 2 seconds
236 (the CCD is read out every 0.4 s, and co-added in groups
237 of 5), over a range of angles 57° either side of nadir. This
238 gives a sufficient “push-broom” width to view the entire
239 sunlit surface of the Earth, even in the tropics, with mul-
240 tiple orbital overlaps for much of the mid- to high-latitude
241 regions.

242 During normal operations, OMI measures the solar ir-
243 radiance spectrum once every 24 hours. The ratio of
244 the earthshine radiance to solar irradiance, the *spectral*
245 *albedo*, is calculated for each FOV. The OMI NO₂ algo-
246 rithm starts by fitting the spectral albedo to a set of
247 reference spectra to get slant column densities (SCD). A
248 simple air mas factor (AMF), calculated based on the as-
249 sumption of unpolluted conditions, where most of the
250 NO₂ is in the stratosphere, is used to obtain an initial
251 vertical column density (VCD). The data from up to 15
252 consecutive orbits are assembled and analyzed in order
253 to construct a “background” field (essentially, the unpol-
254 luted, stratospheric field, with a small contribution from
255 the upper troposphere). The individual FOV initial VCDs
256 are compared to the background field, and where they
257 exceed the background field, significant tropospheric pol-
258 lution is inferred. The VCDs are then recalculated, using
259 an AMF derived using an *a priori*, model-derived tropo-
260 spheric NO₂ profile shape. This is used to recalculate the
261 total column, and hence the tropospheric column.

1.4. Data availability

262 The OMI NO₂ data product is available in a number of
263 different geospatial forms:

- 264 • Level 2 orbital swath (L2);
- 265 • Daily global gridded, 0.25° × 0.25° (L2G);
- 266 • Station and regional overpass (OVP).

267 The L2 and L2G datasets and associated documen-
268 tation are freely available through the NASA God-

269 dard Earth Sciences Data and Information Services Center
270 (GES-DISC, URL: [http://daac.gsfc.nasa.gov/Aura/](http://daac.gsfc.nasa.gov/Aura/OMI/index.shtml)
271 [OMI/index.shtml](http://daac.gsfc.nasa.gov/Aura/OMI/index.shtml)).

272 The OVP data, generated on a daily basis for
273 over 100 locations around the world and also in sup-
274 port of validation and regional pollution studies, are
275 available through the Aura Validation Data Center
276 (AVDC, URL: [http://avdc.gsfc.nasa.gov/Data/Aura/](http://avdc.gsfc.nasa.gov/Data/Aura/OMI/OMNO2/index.html)
277 [OMI/OMNO2/index.html](http://avdc.gsfc.nasa.gov/Data/Aura/OMI/OMNO2/index.html)). The subsetted data used for
278 this paper were generated at the AVDC using the recom-
279 mended usage quality flags [Celarier *et al.*, 2006].

280 Both the L2G and OVP data products are derived from
281 the L2 data set, and not all of the fields found in the L2
282 data may be found in the derived data products. The
283 L2 data are available as Hierarchical Data Format–Earth
284 Observing System (HDF-EOS) format files, which consist
285 of three data groups: Geolocation Fields, Data Fields,
286 and File Attributes. Each data file has a corresponding
287 metadata file; copies of all the metadata are also included
288 in the data file. The data fields include all the values of
289 intermediate variables calculated en route to the principal
290 data fields. Complete details concerning the contents of
291 the Level 2 files are available in *Veeffkind and Celarier*
292 [2006].

293 The L2G files are also HDF-EOS files, and contain a
294 $0.25^\circ \times 0.25^\circ$ grid data structure. Each cell of the grid
295 contains a stack of data values for all the FOVs whose
296 centers fell within that cell. For each FOV a subset of
297 the available L2 fields is stored. Because it is organized
298 geographically, the L2G data set should be suitable for
299 users who wish to study specific geographic locations,
300 as, for example, in the case of validation against ground
301 based measurements, or for regional air quality studies.

302 Field campaign data, and other ground-based mea-
303 surement data are also available through the AVDC.

304 Though all the data have been publicly released and
305 are freely available, prospective data users are strongly
306 encouraged to contact the principal investigators respon-
307 sible for the data sets.

2. OMI Measurement

308 Atmospheric NO₂ column densities are retrieved using
309 spectral measurements of the solar irradiance and earth
310 shine radiance in the wavelength region 415–465 nm, us-
311 ing the instrument’s VIS detector. The measurements are
312 made with a spectral resolution of ~ 0.5 nm. Daily mea-
313 surements of the solar irradiance have been made since
314 the instrument became operational, with the exception of
315 the period 2006 February 28 through 2006 March 3, when
316 a problem with the instrument’s folding mirror prevented
317 making daily irradiance measurements. Using measured
318 irradiance spectra has resulted in the appearance of stripe
319 structure in virtually all the data products, in which the
320 retrieved quantities have different means at each of the
321 60 cross-track positions. This has necessitated the imple-
322 mentation of “destriping” algorithms.

323 The OMI instrument design and performance have been
324 described by *Levelt et al.* [2006b, a]. *Dobber et al.* [2006]
325 have discussed the calibration of the instrument, and the
326 origin of the striping, or cross-track bias.

3. OMI Algorithm

327 In this section we present the essential details of the
328 algorithm. A much more detailed description of the OMI
329 NO₂ algorithm, and its theoretical underpinnings may be
330 found in [*Bucsela et al.*, 2006; *Boersma et al.*, 2002].

331 The OMI NO₂ algorithm proceeds in three steps. In

332 the first step, spectral fitting (the DOAS method) is used
333 to fit the logarithm of the ratio of radiance to irradiance
334 to a set of laboratory-measured absorption spectra
335 of the trace gases, plus a function that models the effect
336 of the rotational raman effect, plus a cubic polynomial
337 to model the wavelength dependence of Rayleigh and
338 aerosol scattering. The coefficients thus obtained give
339 estimates of the slant column densities (SCD) of the vari-
340 ous trace gases. To proceed further, an initial estimate of
341 the vertical column density (VCD; V_{init}) is made using an
342 AMF that is computed using a typical profile containing
343 very little tropospheric NO_2 . In the second step, a num-
344 ber of candidate AMFs are computed, based on assumed
345 NO_2 profile shapes for polluted and unpolluted scenes,
346 and for clear and cloudy conditions.

347 In the third step, data are assembled from each orbit
348 and the orbits occurring within ± 12 hours of it. Within
349 1° latitude bands a wave analysis (up to wave-2) is per-
350 formed on the V_{init} after masking data from known persis-
351 tent strong NO_2 sources, as well as any algorithmically-
352 determined outliers. The resulting model is then taken
353 to model the unpolluted, or “background” field. The
354 value of V_{init} for each field-of-view is then compared to
355 the background field. If it significantly exceeds the back-
356 ground field, then the VCD is recomputed using an AMF
357 that is computed assuming an NO_2 profile that has sub-
358 stantial tropospheric concentration.

359 The following subsections provide more detail about
360 these computations.

3.1. Slant column densities

361 The first part of the calculation of NO_2 columns con-
362 sists in calculating the slant column densities. Since the
363 OMI-measured radiance and irradiance spectra, and the
364 laboratory spectra are all measured on different wave-
365 length scales, the measured spectra are interpolated onto
366 a common scale. The spectral albedo, R , is then fit by a
367 nonlinear least-squares technique onto the function

$$R(\lambda) = P_3(\lambda) \cdot \exp(-c_{\text{NO}_2} \cdot \sigma_{\text{NO}_2}(\lambda) - c_{\text{O}_3} \cdot \sigma_{\text{O}_3}(\lambda)) \cdot (1 + c_{\text{ring}} \cdot \sigma_{\text{ring}}(\lambda)), \quad (3)$$

368 where σ is the absorption cross section of the indicated
369 species, and P_3 is a third-order polynomial in the wave-
370 length, which models the component of the spectrum that
371 is smoothly varying, due to Rayleigh and Mie scattering.
372 Literature spectra are used for σ_{NO_2} [Vandaele *et al.*,
373 1998], σ_{O_3} [Burrows *et al.*, 1999], and σ_{ring} [Chance and
374 Spurr, 1997]. These spectra were convolved with a model
375 OMI instrument slit function prior to use in the fitting al-
376 gorithm. In all, each measured spectrum is subjected
377 to a nonlinear least-squares fit with a total of seven free
378 parameters (c_{NO_2} , c_{O_3} , c_{ring} , and the four coefficients in
379 $P_3(\lambda)$). The algorithm also estimates the uncertainties
380 in each of the fit parameters, as well as the χ^2 error and
381 R.M.S. error of the fit.

3.2. Initial vertical column densities

382 Initial estimates of the vertical column density (V_{init})
383 are calculated using AMFs derived from typical climato-
384 logical profile shapes, with a nominal amount of NO_2
385 assumed in the troposphere (AMF_{init}). That is, the ini-
386 tial vertical columns are computed under the assumption
387 that the troposphere is not polluted.

3.3. Stratosphere-troposphere separation

388 At the core of the OMI NO_2 algorithm is a procedure to

389 identify fields-of-view (FOV) where there is significant tropo-
390 spheric NO₂. This is required because the air mass fac-
391 tor depends upon the profile shape (though not the total
392 amount, since the trace gas is optically thin): FOVs where
393 there is significant tropospheric NO₂ require a different
394 AMF to compute the VCD from the SCD. It is observed
395 [Gordley *et al.*, 1996] that the stratospheric NO₂ field
396 has relatively small gradients, particularly in the zonal di-
397 rection. Our procedure for the stratosphere-troposphere
398 separation essentially identifies the slowly-varying com-
399 ponent of the total NO₂ field as the stratospheric field,
400 and the rest as the tropospheric field.

401 Each orbit is treated as follows. The “target” orbit’s
402 data are read in, along with the data from all other avail-
403 able orbits that were measured within ± 12 hours of the
404 target. Each FOV is identified with a grid cell on a $1^\circ \times 1^\circ$
405 grid in latitude and longitude. For all the FOVs that are
406 identified with a particular grid cell, a “cost” is com-
407 puted from the initial AMF and uncertainty estimate for
408 the V_{init} ; the value of V_{init} having the lowest cost is saved
409 in its associated grid cell. A “mask” identifying grid cells
410 where there are known, persistent sources of NO₂ was
411 developed for use in the algorithm; no V_{init} values are
412 stored in masked grid cells. The V_{init} values are averaged
413 in the meridional direction with a boxcar function of half
414 width 5° . For each 1° latitude band, a wave analysis
415 is performed, fitting waves 0, 1, and 2, to give a preliminary
416 background field. Grid cells whose V_{init} value exceeds the
417 preliminary background field by more than one standard
418 deviation are then excluded, and the wave analysis is re-
419 done. The result of this is a background field (V_{bg}) that
420 has been influenced very little by the presence of regions
421 of high NO₂ concentration. Since the V_{init} values were
422 obtained using an AMF that is appropriate to a profile
423 having most of the NO₂ in the stratosphere, no further
424 correction to the background field is required.

3.4. Vertical column densities

425 For each FOV, the value of V_{init} is compared to the
426 evaluated background field at that location. If V_{init} is less
427 than the background field, then the final value of V (the
428 total NO₂ column amount) is taken to be V_{init} . If V_{init}
429 is larger than the background field, then the “polluted”
430 part ($V_{\text{init}} - V_{\text{bg}}$) is scaled by the ratio $\text{AMF}_{\text{init}}/\text{AMF}_{\text{pol}}$,
431 where AMF_{pol} is obtained using the climatological GEOS-
432 CHEM-modeled profile [Bey *et al.*, 2001; Martin *et al.*,
433 2002]. This procedure gives the total column, the back-
434 ground column, and the polluted column. In addition, a
435 tropospheric column, equal, in the polluted case, to the
436 polluted column plus the amount of the unpolluted pro-
437 file that exists below the tropopause (assumed to be at
438 200 hPa). Finally, if, according to the standard cloud
439 product, the cloud fraction is larger than 0.1, then the
440 “below cloud amount” (the amount of NO₂ that is in-
441 ferred to be below the visible surface of the clouds) is
442 also computed.

3.5. Destriping

443 Due to radiometric calibration and dark-current drift
444 in OMI’s CCD detectors, which affects the radiance mea-
445 surements differently from the irradiance measurements,
446 nearly all OMI Level-2 data products show some degree
447 of cross-track bias, which appears as stripes of systemat-
448 ically elevated or diminished values at certain cross-track
449 scan positions, persisting throughout each orbital track
450 [Dobber *et al.*, 2006]. While the origin of much of the
451 cross-track bias is now understood, and an improvement
452 in the Level-0 to Level-1 processing algorithm is being

453 implemented, the data available for the purpose of vali-
454 dation to date have had significant cross-track bias.

455 A “destriping” procedure has been implemented in the
456 OMI NO₂ algorithm. In this procedure, the NO₂ SCDs and
457 AMFs are collected for the 15 orbits (or fewer, depending
458 on data availability) used to construct the background
459 field. These are then used to construct separate SCD cor-
460 rection offsets for the northern and southern hemispheres:

$$\Delta_i = \overline{\text{SCD}_i} - \overline{\text{AMF}_i} \cdot \frac{\langle \text{SCD} \rangle}{\langle \text{AMF} \rangle}, \quad (4)$$

461 where i is the cross-track scan position (1 to 60), the
462 overlines indicate averages for single scan positions, and
463 angle-brackets indicate averages over all scan positions.
464 The Δ_i are subtracted from the SCDs before applying the
465 final air mass factors.

466 One concern about this procedure has been that it
467 could introduce an unknowable bias in the computed NO₂
468 column densities. This will be discussed in light of the
469 ground-based validation data.

4. Validation of OMI NO₂ columns

4.1. Stratospheric column

4.1.1. SAOZ and DOAS instruments in the NDACC network

472 The NDACC (Network for the Detection of Atmo-
473 spheric Composition Change) is an international cooper-
474 ative network that coordinates the operations and data
475 analysis at more than 30 stations at various latitudes
476 on the globe from 76°S to 79°N. The ground-based
477 UV-Visible zenith-sky spectrometers include both SAOZ
478 (*Système d’Analyse par Observations Zénithales*) as well
479 as DOAS instruments, which provide ozone and NO₂ ver-
480 tical columns at sunrise and sunset using the Differen-
481 tial Optical Absorption Spectroscopy (DOAS) technique
482 [Platt, 1994] in the spectral range 410–530 nm. Zenith-
483 sky measurements made at solar zenith angles between
484 86–91° are averaged to give estimates of the column NO₂.
485 Because of the optical geometry of the measurement,
486 the retrieved NO₂ column is much more sensitive to the
487 stratospheric NO₂ column than to the tropospheric col-
488 umn. Most of the instruments are located in remote ge-
489 ographical regions, far from any significant source of tro-
490 pospheric NO₂. Figure 2 shows the geographical distri-
491 bution of the SAOZ stations. Only the instruments at the
492 Observatoire Haute-Provence (OHP), France, and Bauru,
493 Brazil, are in any proximity to presumed anthropogenic
494 sources of NO₂. Measurements from the SAOZ instru-
495 ments have been previously used to compare with NO₂
496 measurements by the space borne GOME and SCIAMACHY
497 instruments [Ionov *et al.*, 2006a, b, 2007; Piders *et al.*,
498 2006; Lambert *et al.*, 2001].

499 Stratospheric NO₂ exhibits a pronounced diurnal cycle
500 due to its daytime photolysis into NO and nighttime con-
501 version into N₂O₅, as described in Section 1.1. The NO₂
502 daily cycle starts with a fast drop shortly after sunrise,
503 followed by a quasi-linear slow increase during the day, a
504 fast increase at sunset, and finally a slow decrease during
505 the night. The diurnal cycle has been simulated with a
506 photochemical box model derived from the SLIMCAT 3D
507 chemical-transport model [Denis, 2005]. It includes 98
508 chemical and 39 photochemical reactions, including het-
509 erogeneous chemistry on liquid and solid particles. Cal-
510 culations are made at 17 altitude levels with a time step
511 of 1 minute. The NO₂ total column is obtained by in-
512 tegrating the profile assuming a constant density in each
513 layer. Figure 3 shows the results of simulations at two

514 SAOZ stations, OHP at mid-latitude and and Scoresby
515 Sund in the Arctic, for summer and winter. Using this
516 photochemical model, a diurnal time series of the ratio
517 $\text{NO}_2(\text{sunrise})/\text{NO}_2(t)$ was calculated for each month, and
518 at each SAOZ location. As SAOZ is an average of measure-
519 ments between 86° and 91° SZA the NO_2 column at 88.5°
520 SZA is taken as the sunrise reference. All OMI measure-
521 ments were normalized to corresponding sunrise values
522 using these ratios.

523 The optical geometry of the twilight SAOZ measure-
524 ments is such that the light paths traverse rather large
525 distances through the stratosphere, so the stratosphere
526 is sampled at some distance from the measurement site.
527 This should be taken into account when seeking “match
528 up” satellite FOVs corresponding to the ground-based
529 measurements, especially in regions with large strato-
530 spheric NO_2 gradients.

531 Finally, the OMI NO_2 algorithm provides total column
532 NO_2 , and the tropospheric column NO_2 . Since SAOZ
533 measurements are roughly 50 times as sensitive to the
534 stratospheric column as to the tropospheric column, it
535 is of interest to compare the SAOZ-derived values to the
536 difference of the total and the tropospheric columns.

537 Figure 4 shows the time series of the difference be-
538 tween the sunrise SAOZ measurements and the match-
539 ing OMI measurements from eight SAOZ sites, adjusted to
540 account for the difference between the satellite overpass
541 time and sunrise. The statistical characteristics of these
542 differences are presented in Table 1. Besides a compari-
543 son to just the stratospheric column, the table presents a
544 comparison between the OMI total column and the SAOZ
545 instrument measurements. It is seen that, at virtually
546 all latitudes, there is very good agreement, on average,
547 between the ground-based and satellite-based measure-
548 ments of the total stratospheric NO_2 column. However,
549 a small annual cycle is apparent in the time-series for the
550 higher latitudes, with lower values in the winter than in
551 the summer. This cycle, which appears in both the north-
552 ern and southern hemisphere high latitudes, may be re-
553 lated to the OMI sampling under those conditions, or may
554 reflect a sensitivity to the choice of matching OMI FOV cor-
555 responding to a given ground-based observation, or may
556 be due to a bias either in the satellite measurement at
557 high solar zenith angle, or in the ground-based measure-
558 ments as the sunrise azimuth tends poleward. While the
559 influence of the seasonal cycle on the overall statistics is
560 fairly small, understanding it may be an avenue of further
561 study.

562 The correlation coefficients between the SAOZ and OMI-
563 measured stratospheric NO_2 columns are better, and the
564 mean absolute differences smaller, for the mid-to-high
565 latitude sites than for the tropical sites. Since the strato-
566 spheric NO_2 concentrations are smaller in the tropics in
567 the first place (annual mean of about $2.5 \times 10^{15} \text{ cm}^{-2}$,
568 compared to an annual mean of $4 - 5 \times 10^{15} \text{ cm}^{-2}$ at
569 the high latitude sites), the *relative* differences are much
570 greater in the tropics, and even the mid-latitude sites
571 (OHP and Kerguelen), than at the high-latitude sites.

4.2. Tropospheric column

4.2.1. MAX-DOAS

573 The MultiAxis DOAS (MAX-DOAS) technique is an ex-
574 tension of the zenith-sky DOAS technique described in
575 Section 4.1.1, but having much greater sensitivity to
576 lower tropospheric layers. In brief, a MAX-DOAS typi-
577 cally consists of two main parts: A grating spectrome-
578 ter mounted inside a thermostatted box that is located
579 inside a building, and one or more scanning telescopes
580 connected to the spectrometer via fiber optics. Consecu-

581 tive measurements at increasing elevation angles are per-
582 formed in an acquisition cycle that always contains ob-
583 servations at a number of low elevations, and a zenith
584 observation.

585 From each of the measurements, a slant column is
586 retrieved using the DOAS method described in Sec-
587 tion 4.1.1 [Platt, 1994]. Besides NO₂, a number of other
588 absorbers, plus the Ring effect, are included in the fit,
589 as are a multiplicative polynomial and an additive poly-
590 nomial for stray light correction. In order to account for
591 the temperature dependence of the NO₂ absorption spec-
592 trum, a second cross-section (295 K and 221 K) may be
593 introduced in the retrieval to improve the fit and correct
594 the derived vertical column. This potentially allows one
595 to derive the effective air temperature at the centroid of
596 the NO₂ profile. It should be noted that not all retrievals
597 used NO₂ cross-sections at two temperatures, so system-
598 atic differences between different data sets may exist.

599 The lowest-elevation measurements have a large sen-
600 sitivity to absorption in the boundary layer, while the
601 zenith measurements are used as background reference
602 spectra which contain Fraunhofer structures and the
603 stratospheric absorption features. Since photon scat-
604 tering largely occurs below the tropopause, the photons
605 collected from different elevation angles have essentially
606 the same stratospheric path, but different light paths
607 in the troposphere. The difference between successive
608 off-axis line-of-sight (LOS) and zenith measurements is
609 therefore only sensitive to the troposphere. For NO₂ re-
610 trieval, radiative transfer simulations show that under
611 polluted conditions, the stratospheric contamination is
612 generally smaller than 1%. A more in-depth description
613 of the MAX-DOAS measurements, as they were done at the
614 DANDELIONS campaign, can be found in *Brinkma et al.*
615 [2007].

616 An application that is under development, is the re-
617 trieval of boundary layer profile information [Wittrock,
618 2006]. This is done by applying an optimal estimation
619 method to the observations from different elevation an-
620 gles, yielding profile information for roughly the first
621 2.5 km, with about 5 independent pieces of information.
622 Experimental NO₂ profiles were retrieved from the 2005
623 Bremen MAX-DOAS data, but are not yet ready for pub-
624 lication (F. Wittrock, private communication).

625 During the DANDELIONS campaigns [Brinkma et al.,
626 2007], various MAX-DOAS instruments operated quasi con-
627 tinuously from the Cabauw Experimental Site for Atmo-
628 spheric Research [Russchenberg et al., 2005] throughout
629 May through mid July 2005 and throughout September
630 2006. These instruments were provided and operated by
631 BIRA-IASB, the University of Bremen, and the Univer-
632 sity of Heidelberg.

633 The Heidelberg MAX-DOAS instrument differs from the
634 others in that it has a set of three movable telescopes,
635 which enable simultaneous measurement cycles in three
636 azimuth viewing directions. The individual quartz fibers
637 from each bundle associated with the three different tele-
638 scopes are arranged in a vertical column at the entrance
639 slit of the spectrograph (with two gaps between the three
640 fiber bundles, see *Wagner et al.* [2004]).

641 For the Bremen instrument [Wittrock et al., 2004], the
642 zenith direction is viewed without a mirror, while the
643 other elevation angles in the measurement cycle are se-
644 lected through a rotating mirror inside the telescope. The
645 range of elevation angles is 0° – 30°. In the 2006 DANDE-
646 LIONS campaign, separate UV and VIS instruments were
647 operated by the Bremen group.

648 4.2.2. Agreement Between MAXDOAS Instru-

650 The level of agreement achieved between the MAX-
651 DOAS instruments is quantitatively summarized in cor-
652 relation plots (Fig. 5) where tropospheric NO₂ columns
653 from the BIRA instrument are compared to those from
654 the other groups, for the 2005 campaign. Very good
655 agreement is found between the BIRA and Bremen data
656 sets (Pearson correlation coefficients of 0.9 and slope of
657 1.1), and also between BIRA and the three Heidelberg
658 telescopes (correlations between 0.82 and 0.91), espe-
659 cially considering that the BIRA instrument was 200 m
660 away from the other two instruments.

661 When the BIRA and Heidelberg data sets were re-
662 trieved using identical NO₂ absorption cross-sections,
663 an even better agreement is found, reaching a correlation
664 coefficient of 0.92 and a slope of 0.99. This level of agree-
665 ment is only achieved when considering the southwest-
666 pointing Heidelberg telescope: this is approximately in
667 the same direction as the two other instruments. This
668 highlights the importance of horizontal inhomogeneities
669 in the NO₂ field that in some cases strongly affect the
670 agreement between ground based and satellite based mea-
671 surements.

672 **4.2.3. Heterogeneity of the NO₂ Field**

673 Assuming that the tropospheric NO₂ layer is horizon-
674 tally homogenous, the observed NO₂ SCDs for the differ-
675 ent azimuth angles observed from the three telescopes of
676 the Heidelberg instrument should have been similar. In
677 turn, from the observed differences for the various view-
678 ing directions, the horizontal heterogeneity of the NO₂
679 concentration field can be estimated. Such an estimation
680 is very important for the validation of satellite instru-
681 ments with ground based observations. In cases of strong
682 horizontal gradients, ground based observations may not
683 be representative for the average value within a satellite
684 ground pixel.

685 Since the horizontal extension of the absorption paths
686 along the line of sight is largest for low telescope ele-
687 vation, we used those at 3° to estimate the heterogene-
688 ity of the tropospheric NO₂ concentration field. We did
689 this by evaluating the SCD in the three azimuthal view-
690 ing directions at 3° elevation, and calculating the ratio
691 of the maximum and the minimum. A horizontally ho-
692 mogenous concentration field yields a ratio of one; the
693 more this ratio deviates from unity, the larger are the
694 horizontal gradients. In addition to the strength of the
695 horizontal gradients, the direction of the NO₂ gradient
696 was estimated, though in a limited way, since the Hei-
697 delberg MAX-DOAS was measuring in only three azimuth
698 directions. Fig. 6 displays the time series of the ratios
699 at daily noon. High ratios indicate strong gradients, and
700 the color of the points indicates the direction of positive
701 gradient.

702 For the interpretation of the retrieved information on
703 the gradient of the tropospheric NO₂ concentration field,
704 it is important to consider two effects that can affect the
705 observed SCDs, especially for low elevation angles: First,
706 the sensitivity to the relative azimuth angle (between the
707 telescope and the sun). This dependency becomes more
708 pronounced for increasing solar zenith angle (SZA) and in-
709 creasing aerosol load [Wagner *et al.*, 2004]. Second, the
710 effect of the atmospheric aerosol load on the atmospheric
711 visibility, and thus on the horizontal extents of the ab-
712 sorption paths along the line of sight. Thus, depending
713 on the aerosol load, the calculated ratio represents in-
714 formation on gradients over areas of different horizontal
715 extent. The dependence on the azimuth angle was found
716 to be below 15%, for SZA between 20° and 80°. Almost
717 all observed ratios of the maximum and minimum NO₂

718 SCDS (see Fig. 6) were much larger than this. Effective
719 path lengths are enhanced by aerosols above about 1 km,
720 and diminished by aerosols below 1 km. For an elevation
721 angle of 3°, the effective path length is about 19 km in
722 a pure Rayleigh-scattering atmosphere, but this can be
723 reduced to as little as 4.5 km by surface-level aerosols, or
724 enhanced to 25 km by higher-altitude aerosols [Brinksma
725 *et al.*, 2007; Deutschmann and Wagner, 2006; Wagner
726 *et al.*, 2007, 2004].

727 4.2.4. Comparisons with OMI tropospheric NO₂

728 The different MAX-DOAS data sets have been compared
729 to the OMI level 2 and level 4 cloud-free data (O₂-O₂ cloud
730 fractions in the OMI products less than 20%) for 2005. In
731 order to produce the correlation plots presented in Fig. 7,
732 the MAX-DOAS data have been linearly interpolated to
733 the satellite overpass time. The corresponding regression
734 analysis parameters are given in Table 2.

735 For the two OMI algorithms, the vertical columns were
736 generally distributed in values ranging from 0 to about
737 2.5×10^{16} molec cm⁻². In one case the ground based MAX-
738 DOAS column (3×10^{16} molec cm⁻²) significantly exceeded
739 the corresponding satellite values, possibly due to a lo-
740 cal enhancement of the NO₂ concentration at Cabauw.
741 A second outlier (reported between parentheses in the
742 Figure) was noted in the OMI Level 4 products. In this
743 outlier case, the reported NO₂ column was twice as large
744 as the corresponding value in the OMI Level 2 product.
745 Because of its obvious anomaly, this point has been ex-
746 cluded from the regression analysis. The strong overesti-
747 mation of the tropospheric NO₂ column obtained in this
748 case with the OMI Level 4 product points to a possible
749 algorithm problem, which might eventually be related to
750 the model profile shape used for the AMF calculation and
751 the calculation of the ghost column for the cloudy part
752 of the pixel. More work is needed to better understand
753 the origin of the problem.

754 The regression analyses show that similar results were
755 achieved with the BIRA and the Bremen data sets, the
756 correlation coefficient between ground based and satel-
757 lite data being about 0.6 for OMI level 2 and about 0.5
758 for OMI level 4. A lower correlation was obtained with the
759 Heidelberg data when considering only the southwest di-
760 rection measurements (closest to the viewing direction
761 of both Bremen and BIRA instruments), possibly due
762 to the smaller number of coincidences with this instru-
763 ment, and also the shorter integration time used, which
764 may increase the sensitivity to local inhomogeneities in
765 the NO₂ field. In order to further explore the impact
766 of possible horizontal smoothing effects on the compar-
767 ison results, the Heidelberg measurements simultaneously
768 recorded from all three directions have been averaged and
769 again compared with satellite data. The resulting corre-
770 lation coefficients, also given in Table 2, have significantly
771 improved and are now the highest of the three MAX-DOAS
772 instruments. This suggests that the scatter in MAX-DOAS
773 versus satellite comparisons is, indeed, largely dominated
774 by spatial (and temporal) averaging effects.

775 As is evident from the regression results, the OMI tro-
776 pospheric NO₂ columns seem to be systematically lower
777 than the MAX-DOAS results, for both OMI products con-
778 sidered. However, it must be noted that the correlation
779 coefficients are rather poor in all cases, which might be
780 due to several reasons including uncertainties in both
781 ground based (geometrical approximation) and satellite
782 retrievals (AMF sensitivity to errors in aerosols, clouds
783 and NO₂ profile shape). As already mentioned, the main
784 reason for the poor correlation is probably related to the
785 spatial mismatch between the ground based MAX-DOAS
786 observation (essentially local) and the satellite measure-

787 ments (averaged over the OMI footprint). One expects
788 that the collection of more comparison points will help
789 in improving the statistical significance of the compar-
790 isons. Hence further studies will be conducted bring-
791 ing in measurement data from the second DANDELIONS
792 campaign. This and detailed validation of the satellite
793 retrievals during the campaigns is the topic of another
794 publication (G. Pinardi et al., in preparation). That pa-
795 per will look into different algorithm results, strength of
796 collocation criteria, role of ghost columns, and, for OMI,
797 also dependence on FOV cross-track angle.

4.3. Total column

4.3.1. Brewer

798 *Cede et al.* [2006] have described a method for retrieval
799 of total-column NO_2 from direct-sun measurements us-
800 ing a Brewer MK-III double monochromator spectropho-
801 tometer. The Brewer MK-III instrument was primar-
802 ily designed to make measurements of ozone from wave-
803 lengths below 320 nm, and can measure spectral irradi-
804 ance and radiance from 285 to 365 nm. Its measurement
805 modes include a spectral scan mode, where the gratings
806 are moved and any wavelength can be selected, and a
807 slit mask mode, in which a slit mask is introduced in the
808 optical path allowing nearly simultaneous measurements
809 at 6 wavelengths, spaced about 3 nm apart. The spacing
810 of the slits in the slit mask was chosen to optimize the
811 ozone retrievals between 303 and 320 nm, but in the 345–
812 365 nm range the measured wavelengths fall very nearly
813 on maxima and minima in the NO_2 absorption spectrum
814 (see Figure 8), which permits the retrieval of total col-
815 umn NO_2 . These measurements have been made at the
816 NASA Goddard Space Flight Center, on a nearly contin-
817 ual basis since August 2004, with measurements made
818 every half hour during the sunlit hours.

819 The retrieved NO_2 columns have a large instrumental
820 noise, so data must be averaged over several hours time
821 in order to make meaningful comparisons to the OMI-
822 measured values. However, the location of the instru-
823 ment, which is situated 3 km from the Washington Cap-
824 ital Beltway and 2 km from the Baltimore-Washington
825 Parkway, on the outskirts of a major metropolitan area,
826 is such that there are often substantial sub-hour time
827 variations in the actual tropospheric NO_2 concentrations.
828 The combination of the intrinsic variability of the mea-
829 surements with the frequent occurrence of significant ac-
830 tual concentration variations within a given time-window
831 used for collocation with OMI overpasses complicates the
832 process of using the Brewer data for validation of OMI
833 NO_2 measurements. Comparisons having useful statisti-
834 cal significance can be made using monthly averages of
835 the Brewer and OMI datasets.

836 Figure 9 shows the comparison between the monthly
837 mean Brewer-measured and OMI-measured NO_2 columns.
838 In the Washington DC area, early afternoon NO_2
839 columns are dominated by the boundary layer columns.
840 The difference that is seen, with OMI-measured columns
841 that are about 35% smaller than the Brewer-measured
842 columns, can thus be largely attributed to the tropo-
843 spheric NO_2 . In Figure 10 the daily and monthly mean
844 values are plotted, along with the line of linear regres-
845 sion to the monthly means. The regression analysis, per-
846 formed on the monthly means, and weighted according
847 to the standard deviations, gives a slope of 0.67, with a
848 correlation coefficient $R = 0.95$.

4.3.2. MultiFunction DOAS (MFDOAS) measure- 851 ments

852 The MF-DOAS instrument observes scattered skylight
853 with a 1° vertical FOV at varying viewing azimuth and el-

854 evation angles, as well as direct sunlight in the UV-visible
855 spectral region. From these measurements are retrieved
856 NO₂, O₃, SO₂, and CH₂O slant columns. The MF-DOAS
857 spectrograph is a single pass commercial Czerny-Turner
858 spectrograph of focal length 300 mm. The instrument
859 covers a wavelength range from 280 nm to 490 nm with a
860 spectral resolution of 0.82 nm (6 pixels FWHM). Scattered
861 sky light is collected by a 12 cm telescope and passes into
862 the spectrograph through two filter wheels that contain
863 depolarizers, spectral flattening filters, and UV cutoff fil-
864 ters. Direct sunlight is fed into a spectralon integrating
865 sphere of diameter 8 cm before passing through the fil-
866 ter wheels and results in a signal level similar to that
867 from the scattered sky. A two-dimensional CCD detector
868 (512 × 2048 pixels) is used in the focal plane. Spectro-
869 graph stray light is reduced by a spectral flattening filter,
870 which reduces the long wavelength throughput of the in-
871 strument relative to the short wavelength signal. A solar
872 tracker moves the entire instrument for positioning and
873 sun tracking. An instrument schematic is presented in
874 Figure 11.

875 This ground-based MF-DOAS instrument was fielded
876 in a prototype form during the INTEX-B campaign for
877 Aura/OMI validation. It was positioned on the roof of
878 a building at Pacific Northwest National Laboratory in
879 Richland, WA (PNNL; 46.3409°N, 119.2787°W), located
880 in an urban area known as Tri-Cities (the merged cities
881 of Kennewick, Pasco and Richland, WA) with total pop-
882 ulation of approximately 150,000 in an area of 250 km².
883 PNNL is situated approximately 15 km north of the center
884 of Richland, and northwest of the area’s population cen-
885 ter. The major source of local NO₂ pollution is vehicular
886 exhaust.

887 NO₂ differential slant columns (DSCD) were derived us-
888 ing the DOAS technique, based on Beers law. A nonlin-
889 ear least squares algorithm was used to fit our measured
890 spectral cross sections of NO₂, O₃, instrument spectral
891 polarization, and Ring effect in the spectral region 400–
892 419 nm. A polynomial was included to remove slowly
893 varying Rayleigh and Mie scattering spectral shapes. The
894 reference solar spectrum used for the data analysis was
895 measured at zenith at local noon on April 30, 2006, a
896 day with very low pollution levels. Raw spectra were cor-
897 rected for detector dark background and flat field. Figure
898 12 shows typical residual optical densities after the least
899 squares fitting procedure for observations taken on May
900 9, 2006 at 5° elevation and 4 azimuth angles. The LI-
901 DORT radiative transfer code [Spurr, 2001; Spurr et al.,
902 2001] was used to calculate the air mass factors (AMF) to
903 convert the DSCD to vertical column density (VCD). As
904 an example of the results, Figure 13 presents the spatial
905 and temporal variation of NO₂ differential slant column
906 for May 9, a polluted day. Higher column densities were
907 observed to the south and east, toward the urban center,
908 as expected. Measurements taken at 5° elevation showed
909 higher NO₂ tropospheric column compared to 15° and
910 45° angles, as expected. These elevated NO₂ slant col-
911 umn densities were particularly pronounced during the
912 morning rush hour.

913 The Aura satellite flies over Tri-Cities area around
914 1330h with spatial resolution approximately 13 km × 24
915 km. Figure 14 shows contour plots of OMI tropospheric
916 NO₂ VCD for May 9, derived from the Level-2 OMI data
917 product. OMI tropospheric NO₂ vertical column densities
918 “integrated” over several pixels in the MF-DOAS observa-
919 tion direction were compared to MF-DOAS tropospheric
920 NO₂ VCD using *a priori* differential AMFs for clear days
921 at PNNL from the LIDORT radiative transfer code. Fig-
922 ure 15 shows results for the time period April 30 through

923 May 13, 2006, with reasonable correlation observed for
924 these clear days.

925 The slope of the data in Figure 15 shows that OMI de-
926 terminations of tropospheric NO₂ VCD are 0.81 ± 0.11 of
927 that determined from MF-DOAS with a correlation coeffi-
928 cient R^2 of 0.92. Thus, OMI measures a somewhat smaller
929 VCD than that determined from MF-DOAS.

930 4.3.3. Pandora-1 Direct Sun DOAS measurements

931 The lightweight, portable Pandora-1 spectrometer sys-
932 tem measures direct-sun irradiances from 270 to 500 nm
933 at ~ 0.5 nm resolution. The outdoor head sensor is
934 mounted on a tracking system and holds a single strand
935 fiber optic cable, which collects the light passed through
936 a collimator (1.6° FWHM field of view) and a filter wheel.
937 The other end of the fiber is connected to a 75 mm fo-
938 cal length symmetric Czerny-Turner grating spectrome-
939 ter using a 1024×1 pixel CMOS detector, stabilized to
940 $20^\circ \pm 1$. The total NO₂ column is retrieved by the DOAS
941 method, in the 400–440 nm window, using a fixed refer-
942 ence spectrum determined from Pandora-1 data obtained
943 over an extended period of at least 2 weeks. To estimate
944 the NO₂ amount in the reference spectrum, a bootstrap
945 method as described in *Cede et al.* [2006] was applied, on
946 the assumption that a few measurements were obtained
947 when there were low tropospheric NO₂ amounts (e.g.,
948 just after sunrise). Figure 16 shows Pandora-1 data dur-
949 ing the SCOUT campaign in July 2006 at Thessaloniki,
950 Greece. Excellent agreement is seen between the OMI
951 and Pandora-1 measurements, though the OMI overpass
952 times seem to occur just before or just after the mid-day
953 maximum in NO₂ concentration; this limits the range of
954 NO₂ concentration values explored in this comparison.
955 A number of further field campaigns are planned, during
956 which Pandora-1/OMI comparisons will be done.

957 4.3.4. Direct Sun DOAS (BIRA)

958 During the second DANDELIONS campaign, a direct-sun
959 DOAS instrument was operated in addition to the MAX-
960 DOAS instrument. The well-defined optical path and air
961 mass factor make this instrument equally sensitive to ab-
962 sorption along the whole optical path and so provides
963 accurate NO₂ total columns.

964 The instrument is similar in concept to the MAX-DOAS:
965 Inside the building, in a thermo regulated box, a grating
966 spectrometer covering the UV-Vis region is coupled to a
967 cooled CCD detector, connected by depolarizing fiber op-
968 tic bundle to the external optical head. Outside, along-
969 side the MAX-DOAS scanning telescope, a collimating op-
970 tic tube is mounted on a BRUSAG commercial sun-tracking
971 system, holding the fiber.

972 The retrieval is also done using the DOAS approach:
973 The ratios of the measured radiance spectra to a reference
974 spectrum are analyzed with respect to a set of reference
975 spectra, in the 425–450 nm window, including laboratory
976 spectra of O₃, H₂O, O₂–O₂, the computed the Ring ef-
977 fect spectrum, and NO₂ cross-sections at two different
978 temperatures. In contrast to the analysis of MAX-DOAS
979 data, a fixed reference spectrum (measured on 7 Septem-
980 ber 2006) has been used for the whole time-series. The
981 NO₂ residual slant column amount included in this refer-
982 ence spectrum has been obtained by analysing it with
983 respect to the Kurucz solar atlas [*Kurucz et al.*, 1984],
984 which was assumed to be free of NO₂ absorption. The
985 Kurucz solar spectrum was convolved with a precisely
986 measured instrument slit function to match the instru-
987 ment's spectral resolution. Based on this analysis, total
988 absolute slant columns could be derived from direct sun
989 measurements; these were transformed into total vertical
990 columns using geometrical AMFs.

991 Figure 17 shows the time series for the BIRA DOAS mea-

992 surements of the total column NO₂ (filled dots), which
993 provides a good idea of the diurnal variation of NO₂ lev-
994 els. The open squares show the collocated OMI measure-
995 ments (one or two per day). The OMI snapshots of ver-
996 tical column NO₂, for the most part, appear to be in
997 quite good agreement with the ground-based measure-
998 ments. Note that the OMI data are filtered for clouds
999 (cloud fraction $\leq 20\%$).

1000 Figure 18 shows the correlation plot of the collocated
1001 data (the point nearest in time to the OMI overpass). A
1002 linear regression, constrained to pass through the origin,
1003 gives a slope of 0.84 ± 0.05 . The scatter in the data
1004 ($R = 0.68$), and the relatively small number of data
1005 points ($N = 26$) do not permit a statistically significant
1006 estimation of an additive bias.

1007 4.3.5. FTUVS measurements at Table Mountain, 1008 California

1009 Another instrument that has been used to validate OMI
1010 NO₂ total column measurements uses the Fourier Trans-
1011 form Ultraviolet Visible Spectrometer (FTUVS), a UV-VIS-
1012 NIR interferometer, at the Table Mountain Facility (TMF),
1013 north of Los Angeles, California, at $34^{\circ}22.9'$ N, $117^{\circ}40.8'$
1014 W, at an altitude of 2290 m (7300') [Cageao *et al.*, 2001].
1015 Spectra are recorded in the direct solar absorption mode
1016 with a spectral resolution of 0.0013 nm, which is suffi-
1017 cient to resolve NO₂ vibronic features. By measuring
1018 the doppler-shifted spectra from the east and west solar
1019 limbs, and taking the ratio of the two, one can remove
1020 the solar Fraunhofer lines; there is no need to measure a
1021 high-sun reference spectrum, as in a number of the other
1022 methods described in this overview.

1023 The instrument is not readily transportable. The
1024 FTUVS observation site overlooks the Antelope Valley,
1025 north of the Los Angeles Basin. This area is character-
1026 ized by relatively clean air under most conditions, but is often
1027 influenced by polluted air from Los Angeles in the after-
1028 noon, advected through the Cajon Pass. While consider-
1029 ably above the tropospheric background under these con-
1030 ditions, the NO₂ column abundance values rarely exceed
1031 1×10^{16} molecules cm⁻², which is considerably smaller
1032 than values measured directly downwind of a polluted
1033 urban area (see Fig. 9). Because the altitude of TMF is
1034 about 2500 feet above the Antelope Valley, FTUVS column
1035 abundance measurements of NO₂ will be biased relative
1036 to the center of the OMI footprint. The bias is small re-
1037 lative to the total column, and will not have a significant
1038 effect on the slope of the OMI-FTUVS correlation. The OMI
1039 data used for validation were sorted by distance from the
1040 TMF site, in order to mitigate somewhat the possible ef-
1041 fects of the distribution of elevations within a FOV. It
1042 was found that a minimum distance of about 10 km is
1043 required for good intercomparison.

1044 The FTUVS instrument time is shared with other Aura
1045 validation activities. On average, measurements were ac-
1046 quired twice a week over the period March–November,
1047 2006.

1048 The slant column NO₂ amounts are retrieved by fitting
1049 the measured absorption spectra to laboratory spectra at
1050 a number of temperatures [Nizkorodov *et al.*, 2004], in
1051 windows containing 10 to 20 NO₂ rotational lines. Ge-
1052 ometric AMFs were used to convert the SCDS to VCDs.
1053 Figure 19 presents the comparison of the OMI-derived
1054 and FTUVS-derived measurements of total column NO₂.
1055 In this figure, the points where the OMI FOV center fell
1056 within 10 km of the Table Mountain Facility site are col-
1057 ored red. The linear regression line shown is fit only to
1058 those points. As shown, this line has a slope of 0.77 ± 0.41 ,
1059 and it does not go through the origin. This data set sug-
1060 gests that the OMI NO₂ totals are underestimated in the

1061 middle of the data range, but that there may also be a
1062 positive additive bias.

4.4. NO₂ Profile measurements

1063 As pointed out in previous sections, and in *Boersma*
1064 *et al.* [2002] and *Bucsela et al.* [2006], the shape of the
1065 vertical profile of NO₂ influences the (physical) air mass
1066 factors. The OMI NO₂ algorithm uses a set of assumed
1067 profiles, which were derived from model studies; these as-
1068 sumed profiles thus affect the retrieved total and tropo-
1069 spheric NO₂ amounts. It is therefore important to evalu-
1070 ate how well the assumed profiles approximate the actual
1071 profiles, *vis-à-vis* the air mass factor calculation. There
1072 have been very few efforts to measure NO₂ profiles [*He-*
1073 *land et al.*, 2002; *Martin et al.*, 2006]. Recent efforts in-
1074 clude measurements during the September 2006 DANDE-
1075 LIONS campaign (lidar, *in situ* at two altitudes, and MAX-
1076 DOAS at two altitudes, see section 4.4.1), and aircraft-
1077 based *in situ* measurements taken during the INTEX-B
1078 campaign in North America.

4.4.1. NO₂ lidar

1079 NO₂ profiles were measured by a lidar system, de-
1080 veloped at RIVM, during the DANDELIONS campaign in
1081 September 2006 [*Brinkma et al.*, 2007]. The lidar con-
1082 sists of an emitter and a receiver unit. The entire system
1083 is housed in a truck, constituting a fully self-supporting
1084 mobile laboratory. The emitter unit consists of a pulsed
1085 pump laser-dye laser combination, running at 30 Hz.
1086 The dye laser is tuned to 449.10 nm and detuned to
1087 448.31 nm every other pulse. The latter wavelength is
1088 absorbed more strongly by NO₂ than the former. The
1089 laser pulses, 40 mJ in energy, 10 ns in duration, are di-
1090 rected into the atmosphere, where they are scattered by
1091 gas molecules and aerosol particles. The receiver unit
1092 collects the backscattered light, through a 280 mm tele-
1093 scope, onto a photomultiplier tube, with an interference
1094 filter to block daylight. A digitizer samples the signals
1095 with a range resolution of 3.75 m.

1096 The NO₂ concentration at a certain altitude is derived
1097 from the log of the ratio of the backscattered signals at
1098 the two wavelengths, using the differential absorption li-
1099 dar (DIAL) method. Since the laser pulses are not emitted
1100 from the center of the telescope, the laser beam is not in
1101 view of the telescope at close range, and thus the lidar is
1102 effectively blind for the first 500 m. When measurements
1103 starting near the surface are required, the emitter section
1104 and receiving telescope are tilted through various eleva-
1105 tion angles; the measurements are combined into a single
1106 profile, where elevations close to the horizontal yield NO₂
1107 concentrations at low altitudes but pertaining to a cer-
1108 tain horizontal extent away from the instrument (for a
1109 near-horizontal measurement, typically about 2500 m),
1110 while a zenith observation is performed exactly above
1111 the truck. Completing one vertical profile typically takes
1112 50 minutes, providing data in a altitude range of a few
1113 meters up to approximately 2500 m, with an accuracy
1114 of 0.2–0.4 $\mu\text{g m}^{-3}$. Range and accuracy depend on at-
1115 mospheric conditions. The vertical resolution of a profile
1116 varies, and typically is about 15 m at the lowest altitude,
1117 increasing to over 500 m at the highest altitude. The
1118 resolution arises from averaging of data over an altitude
1119 range, based on signal-to-noise considerations.

1120 A paper describing the lidar and other time-resolved
1121 three dimensional observations of NO₂ during the 2006
1122 DANDELIONS campaign is in preparation (H. Volten *et al.*,
1123 in preparation).

1124 Figure 20 presents examples of profile measurements
1125 for a relatively clean day, September 9, 2006, and for a
1126 polluted day, September 12, 2006; in both cases, there
1127

1128 was little-to-no cloud cover. The concentration of NO₂
1129 is high at ground level, and drops to zero (within the ac-
1130 curacy of the measurement) above the boundary layer.
1131 The boundary layer heights, provided by the boundary
1132 layer lidar at Cabauw, are indicated in Figure 20 by a
1133 dashed line. The figure shows that the day-to-day vari-
1134 ations in NO₂ at the surface may be considerable, from
1135 around 3 μg NO₂ m⁻³ on a clean day to more than 50
1136 μg NO₂ m⁻³ on a polluted day. Large diurnal variations
1137 may also occur.

1138 4.4.2. In-situ aircraft measurements

1139 *In situ* measurements of NO₂ from the DC-8 aircraft
1140 were obtained during the INTEX-A (summer 2004), PAVE
1141 (winter 2005) and INTEX-B (spring 2006) campaigns.
1142 These have been discussed by *Bucsela et al.* [2007]. The
1143 NO₂ profiles from these experiments are useful for vali-
1144 dating both the shapes of the model profiles used in the
1145 OMI retrieval algorithm, and, in turn, the tropospheric
1146 column amounts from the satellite retrievals. The air-
1147 craft profiles obtained during INTEX-A and PAVE were
1148 combined into composite land and ocean profiles. The *in*
1149 *situ* profiles were seen to be very similar to the annual
1150 mean GEOS-CHEM profiles used to retrieve tropospheric
1151 NO₂ columns from OMI, and the AMFs computed from
1152 the measured profiles were slightly larger than those cal-
1153 culated using the model profiles. A more quantitative
1154 analysis was performed using a set of approximately 70
1155 profiles measured during INTEX-B. Error-weighted lin-
1156 ear regressions comparing the AMFs yielded a slope of
1157 1.10 ± 0.10 (*in situ* profile AMF greater than that used by
1158 the OMI algorithm). This means that the OMI VCD would
1159 overestimate the actual VCD by 10% ($\pm 10\%$).

1160 *In situ* measurements of NO₂ can also be used to vali-
1161 date tropospheric column amounts from OMI. The INTEX-
1162 B data were used for this analysis by *Bucsela et al.* [2007]
1163 (see also *Boersma et al.* [2007]). Two representative pro-
1164 file analyses are shown in Figure 21. The full set of pro-
1165 files from INTEX-B where used. The correlation between
1166 the aircraft and OMI data sets was good ($R = 0.83$). This
1167 comparison is shown in Figure 22. The integrated *in situ*
1168 tropospheric columns were found to be somewhat larger
1169 than the OMI Level-2 columns, as indicated by the slope
1170 of 1.10 ± 0.08 . Although some of the *in situ* columns re-
1171 quired significant extrapolations, sensitivity studies indi-
1172 cated that the overall results were generally robust with
1173 respect to the choices made for the profile binning, in-
1174 tegration and extrapolation, as well as being relatively
1175 insensitive to the errors assumed for the weights. The
1176 insensitivity to extrapolation is consistent with findings
1177 in a similar aircraft study by *Heland et al.* [2002].

1178 5. Conclusions and discussion

1178 This paper has presented a number of results of experi-
1179 ments where ground- and aircraft-based measurements of
1180 NO₂ can be compared with collocated measurements and
1181 retrieval by OMI. Since some measurements estimate the
1182 stratospheric column, others the tropospheric column,
1183 and still others the total column, their results can be
1184 used to validate the OMI NO₂ standard data product's
1185 estimates of these columns.

1186 Table 3 summarizes the results of the numerous vali-
1187 dation studies that have been discussed in this overview.

1188 On the basis of the SAOZ and DOAS measurements,
1189 which are most sensitive to the stratospheric NO₂
1190 columns, the OMI stratospheric NO₂ appears to agree
1191 with the ground-based measurements to within $\sim 10\%$.

1192 The OMI tropospheric column appears to be consis-
1193 tently lower than the various ground-based measure-

1194 ments, though there is some inconsistency amongst those
1195 ground-based measurements. Though many of the vari-
1196 ous instruments and methods for measuring tropospheric
1197 and total NO₂ have not themselves been validated, it is
1198 noteworthy that they all give NO₂ estimates that are on
1199 average greater than those retrieved from OMI. This may
1200 indicate a bias in the OMI retrieval. However, a number
1201 of cases have been studied, where average differences be-
1202 tween OMI and ground based measurements decrease as
1203 the geographic match up criterion is tightened. It is likely
1204 to be due to the inhomogeneity of the tropospheric NO₂
1205 field, and, in particular, the fact that ground-based mea-
1206 surements are often made in or near regions of moderate
1207 to strong sources of NO₂: The OMI FOV that includes the
1208 site will also include a substantial ($\sim 10^2$ km²) regions
1209 where much lower NO₂ concentrations prevail. The was
1210 borne out in the Brewer studies [Cede *et al.*, 2006] and
1211 in the correlation studies of Veeffkind *et al.* [2007] (see
1212 also Section 1.) However, the data taken at TMF (Section
1213 4.3.5) were mostly obtained under conditions of relatively
1214 clean tropospheric air, and these data also suggested a
1215 negative bias for OMI retrieval.

1216 Potential biases can arise at any of the steps in the
1217 algorithm. Instrumental artifacts are known to give
1218 rise to the cross-track bias (striping) and the destrip-
1219 ing process can certainly give rise to a general bias. The
1220 stratosphere-troposphere separation is based on an initial
1221 AMF, and any bias in that AMF will result in a bias
1222 in the background (mostly stratospheric) field. After
1223 the stratosphere-troposphere separation, a new AMF is
1224 constructed, based on model-based-climatology derived
1225 *a priori* profiles. The aircraft *in situ* measurements of
1226 NO₂ profile shape suggest that the *a priori* profile shapes
1227 are essentially correct, in that the two do not give appre-
1228 ciably different AMFs. The AMF is also sensitive to the
1229 surface albedo. The OMI algorithm uses a climatologi-
1230 cal surface albedo, and this may be a worthy subject for
1231 future validation studies.

1232 It should be mentioned that all the validation studies
1233 reviewed here focused on mostly cloud-free conditions.
1234 However, while OMI FOVs are considerably smaller than
1235 those of earlier atmospheric remote sensing instruments,
1236 they are still large enough that very few can be expected
1237 to be completely uncontaminated by clouds.

1238 Acknowledgments

1239 Part of this research was carried out at the Jet Propul-
1240 sion Laboratory, California Institute of Technology, un-
1241 der contract with the National Aeronautics and Space
1242 Administration.

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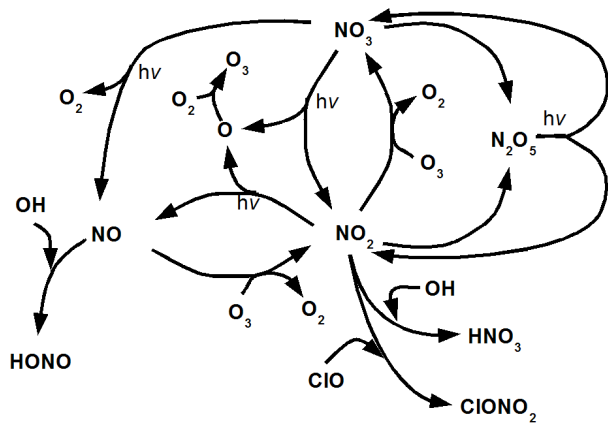


Figure 1. Nitrogen reaction network

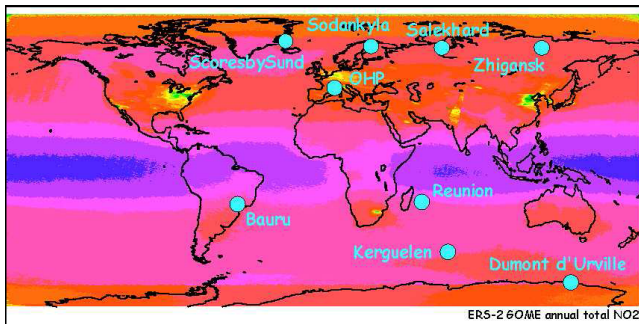


Figure 2. Geographical distribution of SAOZ stations in the NDACC network

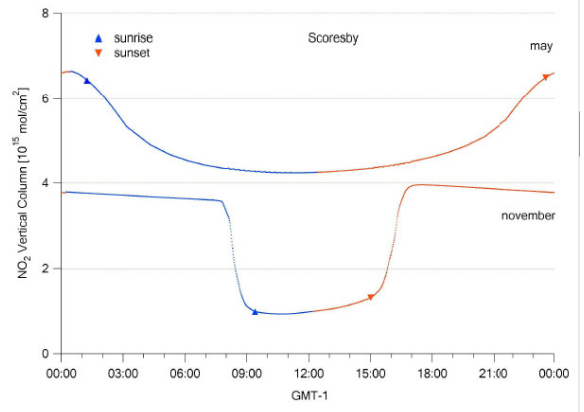
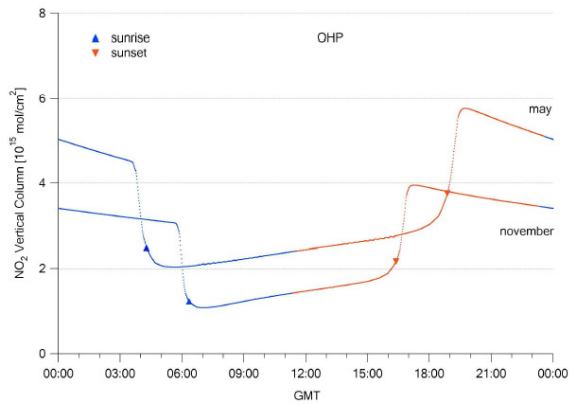


Figure 3. Simulated time-history of stratospheric NO₂ at a mid-latitude station (OHP), and a high-latitude station (Scoresbysund), for summer and winter.

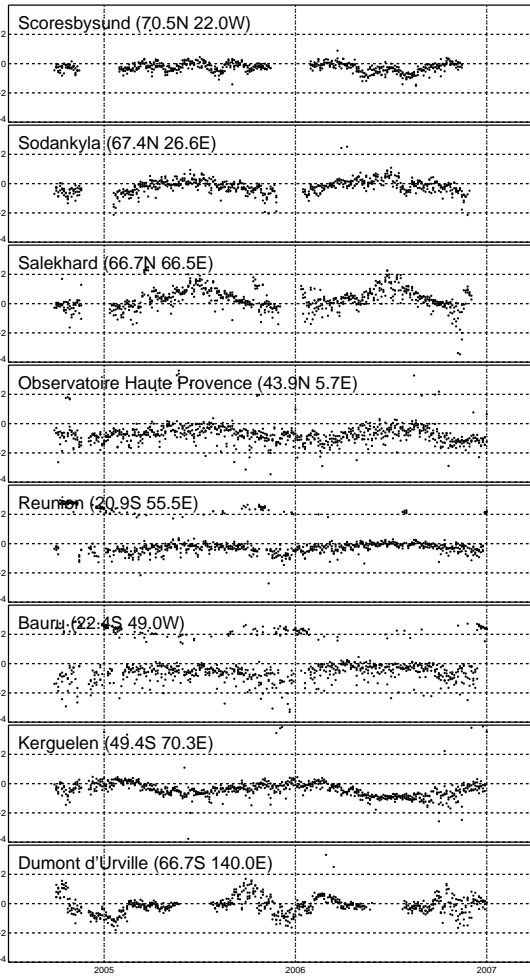


Figure 4. Time series of the difference between OMI and SAOZ-measured stratospheric NO₂ in units of 10^{15} cm^{-2} . The sites are ordered from North to South.

Table 1. Absolute average and r.m.s. difference (Δ , σ ; $\times 10^{16}$ cm $^{-2}$), and correlation (R) between ground-based SAOZ and satellite data, adjusted to sunrise OMI total and stratospheric NO $_2$ (2004-2005)

Station	SAOZ-OMI total column			SAOZ-OMI strat. column		
	Δ	σ	R	Δ	σ	R
Scoresby Sund	+0.09	0.82	0.92	-0.26	0.41	0.99
Sodankyla	+0.64	2.05	0.71	-0.22	0.54	0.97
Salekhard	+1.04	1.57	0.86	+0.25	0.74	0.97
OHP	+1.36	2.35	0.44	-0.83	1.04	0.67
Reunion	+0.29	0.77	0.29	-0.32	0.46	0.65
Bauru	+0.74	2.06	0.14	-0.65	0.86	0.56
Kerguelen	-0.12	0.61	0.87	-0.45	0.66	0.88
Dumont d'Urville	+0.37	1.17	0.87	-0.12	0.61	0.96
OVERALL:	+0.56	1.57	0.70	-0.34	0.70	0.91

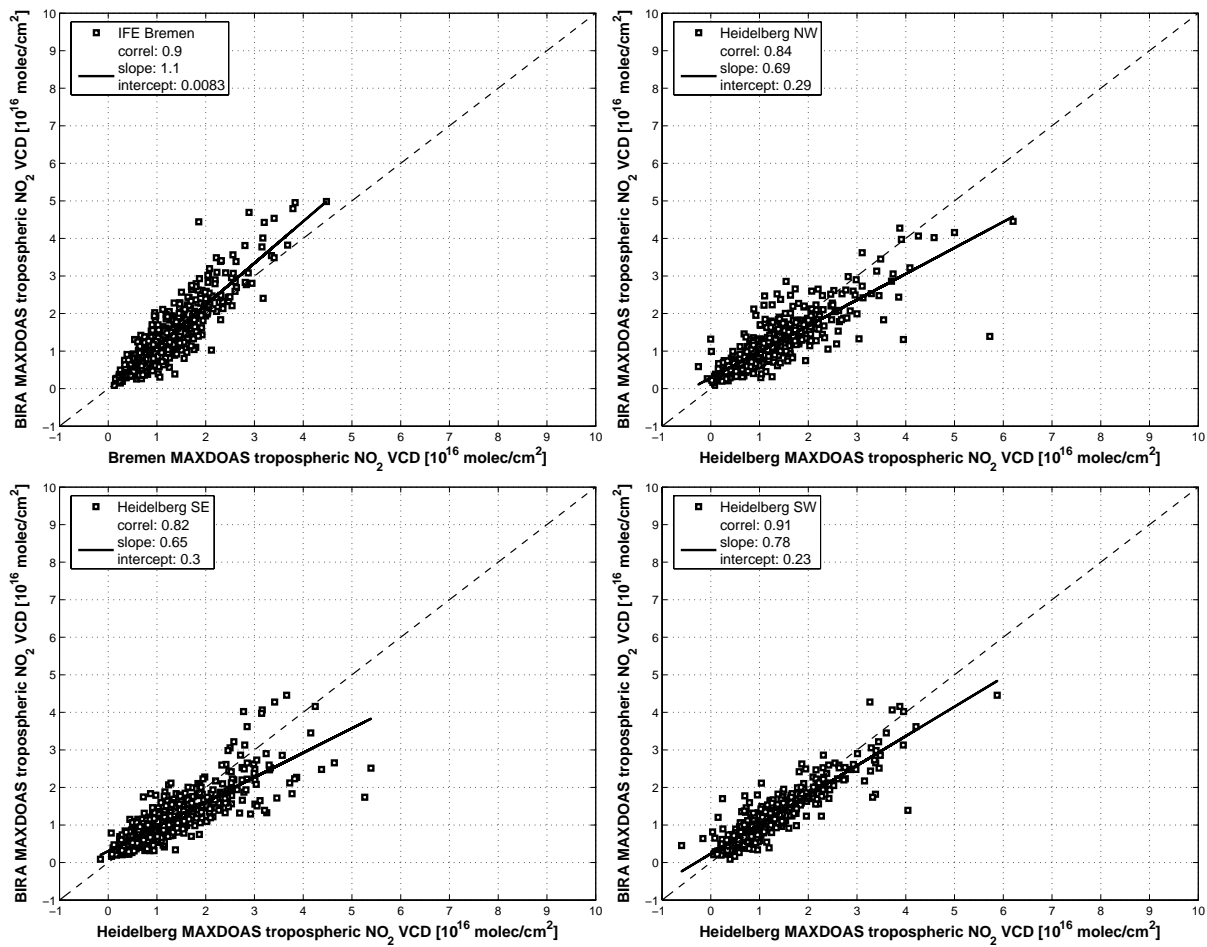


Figure 5. Scatter plots of the tropospheric NO₂ columns retrieved during the 2005 campaign from the BIRA MAXDOAS instrument and, respectively, the Bremen MAXDOAS (top left), the Heidelberg MAXDOAS for the 3 pointing directions North-West (top right), South-East (bottom left) and South-West (bottom right). The regression analysis parameters are given in the legends. It has to be noted that agreement with the Heidelberg observations can be further improved after homogenisation of the retrieval settings (see text).

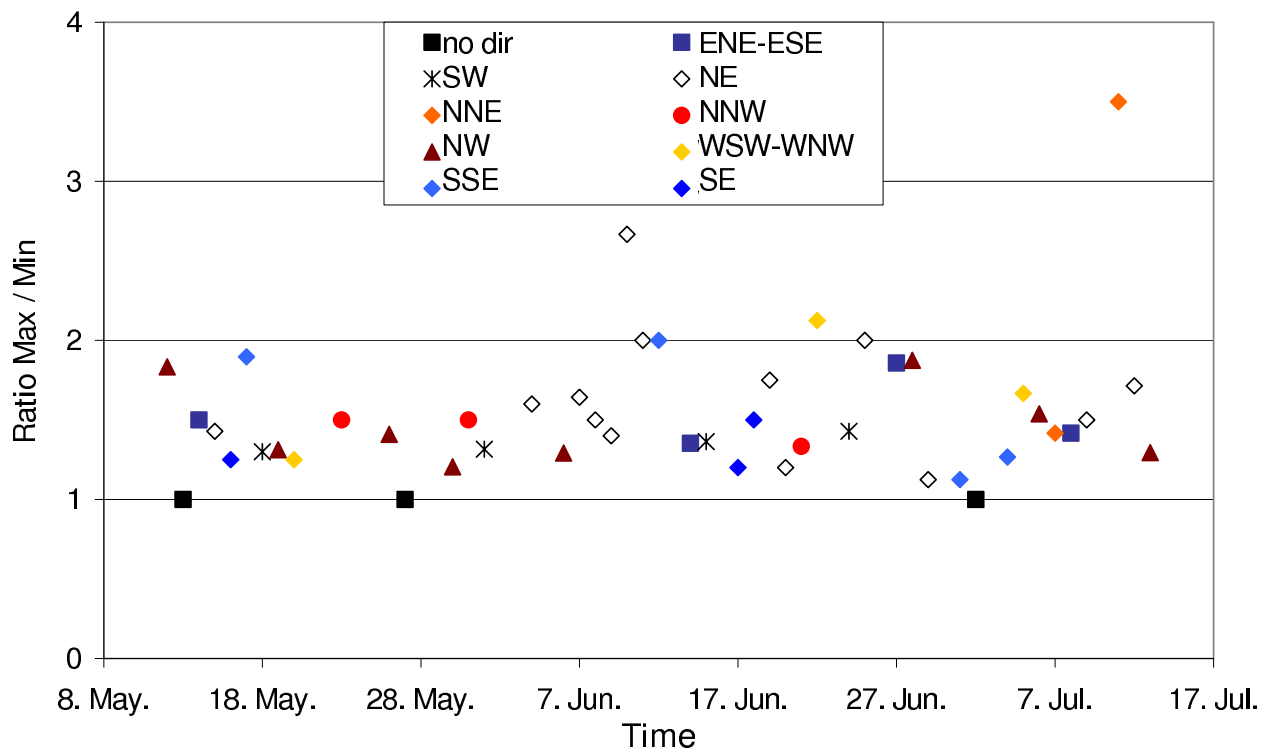


Figure 6. Maximum and minimum NO₂ SCD observed for an elevation angle of 3° of the Heidelberg MAX-DOAS telescopes observing under three different azimuth angles at Cabauw during the DANDELIONS campaign in 2005. High ratios indicate large horizontal gradients of the tropospheric NO₂ concentration field, colors indicate the direction of the gradient (directed toward higher values).

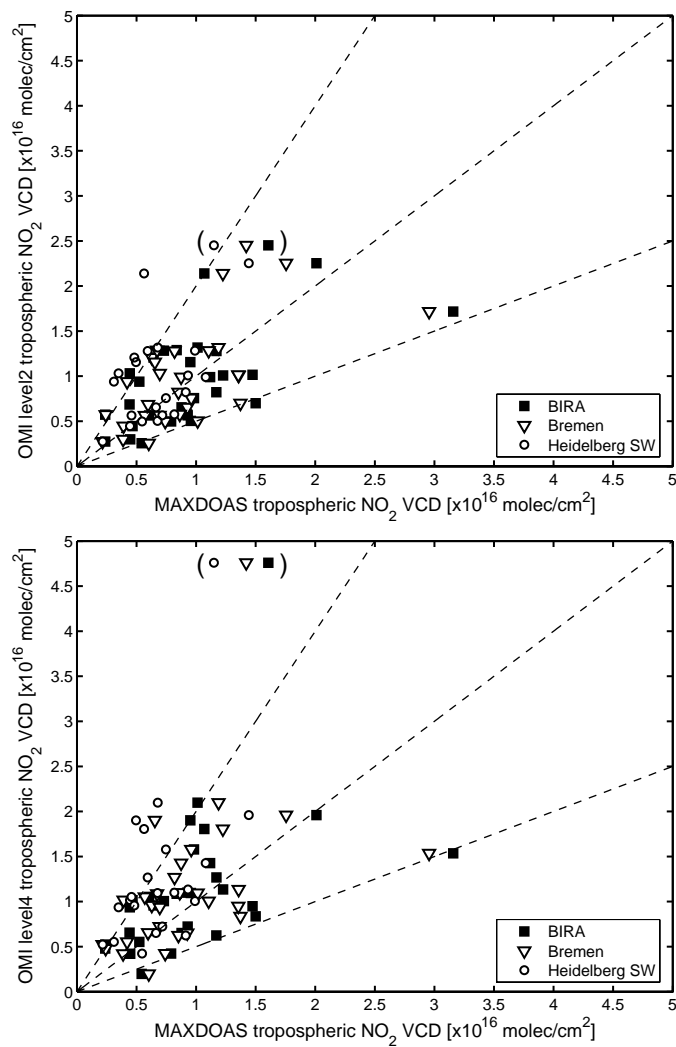


Figure 7. Correlations between tropospheric NO₂ from the three MAX-DOAS instruments at Cabauw (BIRA, Bremen, and Heidelberg SW direction) and OMI-L2 (top panel), or OMI-L4 (bottom panel). OMI data are included if cloud fractions were less than 20%. Correlation and regression coefficients are summarized in Table 2

Table 2. Statistical analysis of comparisons between tropospheric NO₂ from MAX-DOAS data and OMI (L2 and L4)

	BIRA South						Bremen South West					
	N	R	I	slope	rms	rms2	N	R	I	slope	rms	rms2
OMIL2	29	0.6	4.29	0.52	4.82	52%	29	0.63	3.93	0.59	4.44	48%
OMIL4	29	0.51	6.16	0.44	5.35	52%	29	0.52	5.99	0.48	5.18	50%

	Heidelberg South West						Heidelberg spatial average					
	N	R	I	slope	rms	rms2	N	R	I	slope	rms	rms2
OMIL2	21	0.45	4.27	0.8	5.38	56%	21	0.65	1.99	0.85	3.89	40%
OMIL4	21	0.39	6.84	0.67	6.50	57%	21	0.57	4.89	0.71	4.74	42%

^{*} N denotes number of collocations, R is Pearson correlation coefficient, I is intercept in units of $10^{15}m^{-2}$, slope denotes result of linear regression analysis, rms denotes rms difference between groundbased and OMI, in units of $10^{15}m^{-2}$, and rms2 denotes the same in percent relative to the average OMI value.

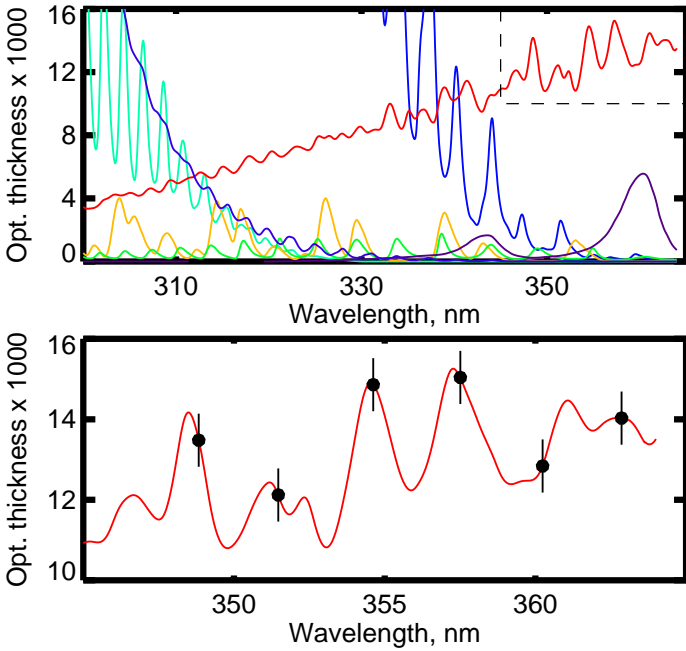


Figure 8. Typical optical depths of the main trace gases in the Brewer MK-III wavelength range. Lower: NO₂ optical depth for 1 DU ($= 2.7 \times 10^{16} \text{ cm}^{-2}$), 6 slit positions with noise estimates.

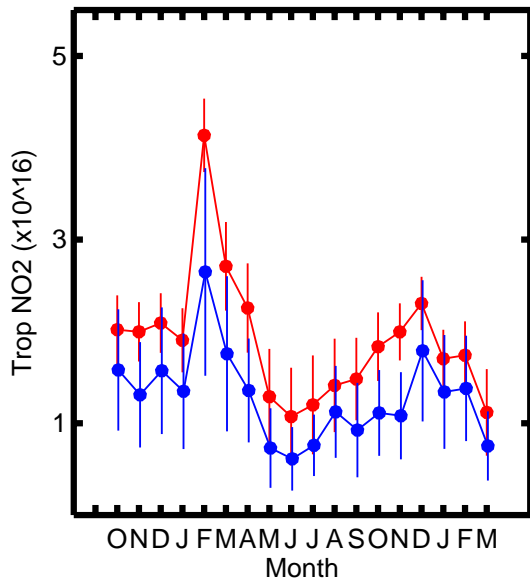


Figure 9. Comparison between the monthly mean Brewer-measured and OMI-measured NO₂ total columns.

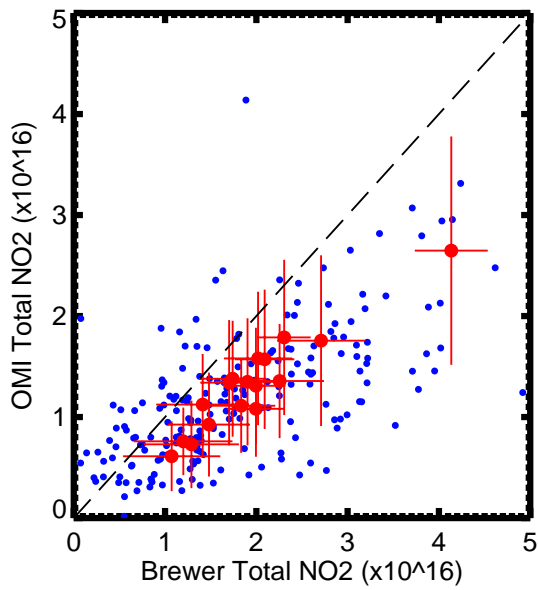


Figure 10. Daily mean and monthly mean values of NO₂ total column measured by the Brewer instrument and OMI. The line of linear regression is also shown.

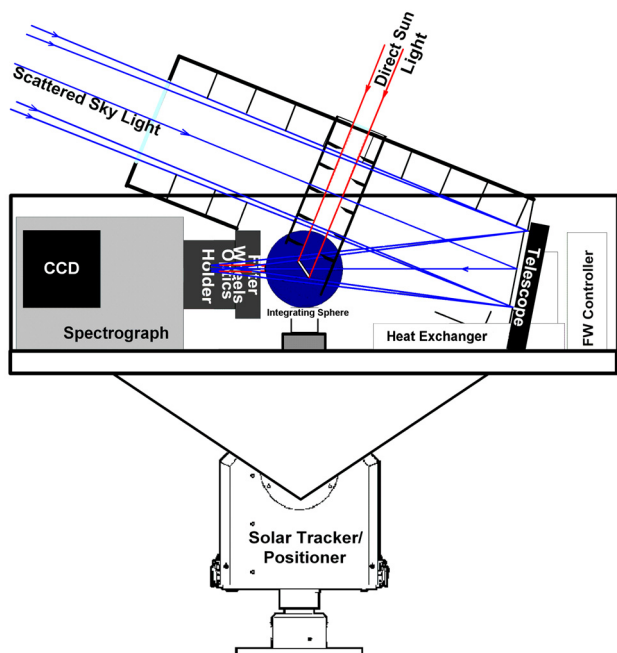


Figure 11. Schematic drawing of the MF-DOAS instrument.

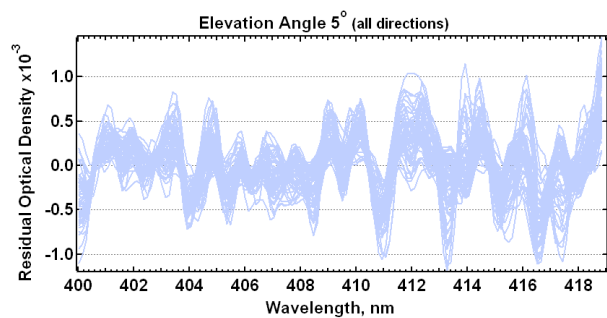


Figure 12. Representative residual optical densities for observations of May 9, 2006 at 5° elevation and 4 azimuth angles.

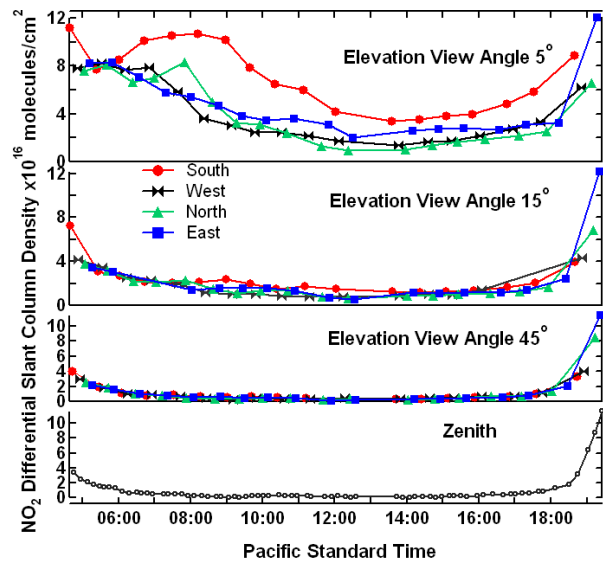


Figure 13. Example of measured spatial and temporal variation of MF-DOAS-measured NO_2 differential SCD for May 9, 2006

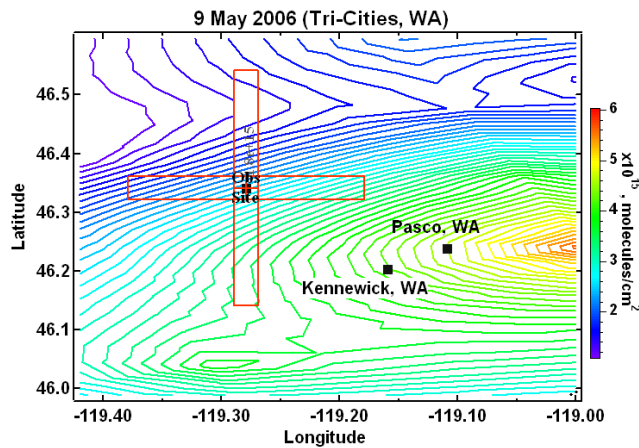


Figure 14. Tropospheric NO_2 VCD over the Tri-Cities area of Washington State on May 9, 2006. The contour map is derived from the individual OMI FOV measurements.

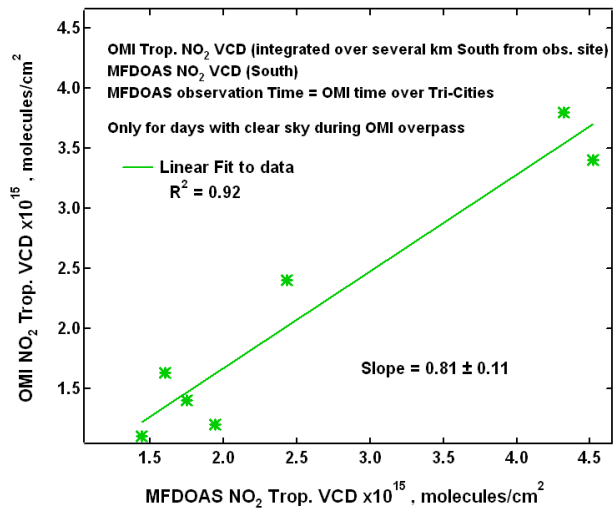


Figure 15. Correlation between OMI tropospheric NO₂ vertical column and MF-DOAS NO₂ tropospheric VCD for measurements made from April 30 through May 13, 2006

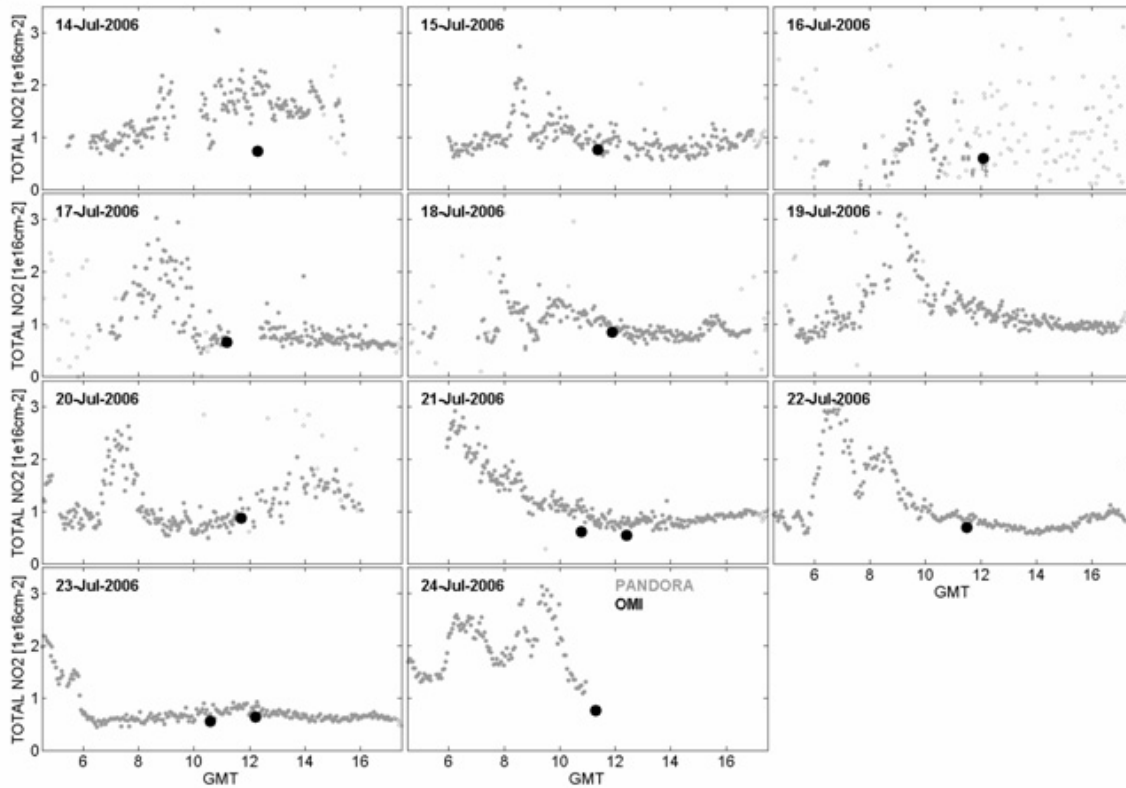


Figure 16. Pandora-1 Data measured during the period 14–24 July 2006, in Thessaloniki (grey dots) and OMI overpass data for the site (large black dots).

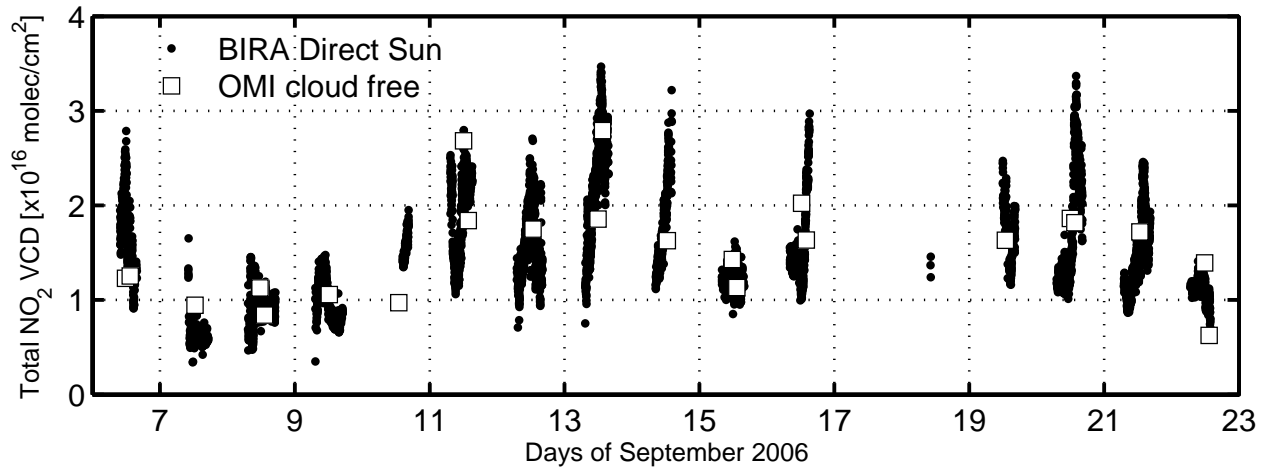


Figure 17. Time series of NO_2 VCD measured by the BIRA direct-sun DOAS instrument. The open squares show the collocated OMI measurements.

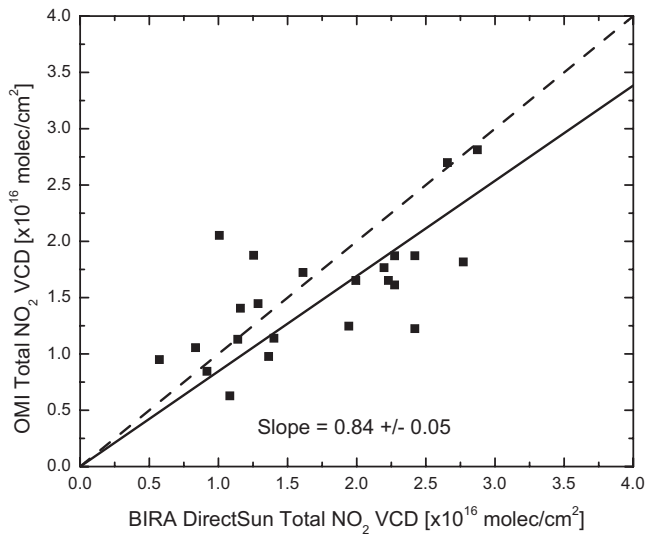


Figure 18. Correlation plot showing the collocated OMI and BIRA direct-sun DOAS instrument measurements (open squares in Figure 17).

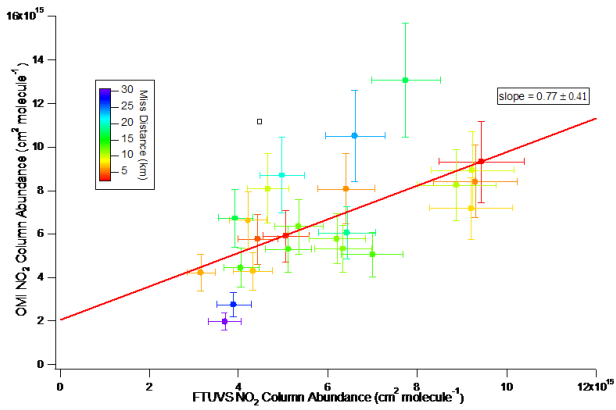


Figure 19. OMI versus FTUVS measurements of total column NO₂, binned by distance between TMF and the centroid of the OMI FOV (distance indicated by color, see inset scale)

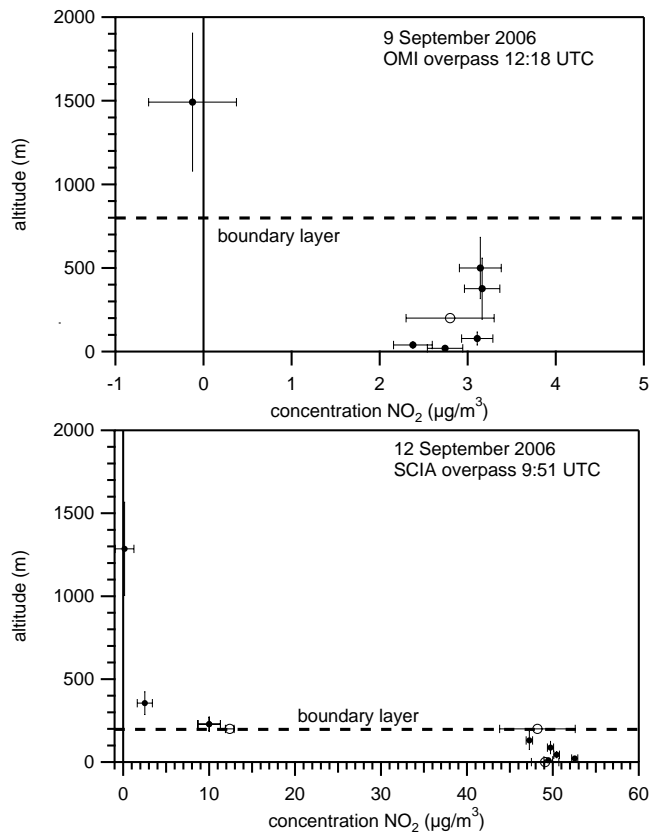


Figure 20. Lidar NO₂ profile (black circles) and NO₂ monitor value (open circle) measured at Cabauw. Horizontal bars indicate two-sigma values for the concentration. For the lidar data, vertical bars indicate the height intervals over which concentrations have been determined. The boundary layer height is indicated by a dashed line. Top panel: Clean day, September 9, 2006 (lidar measurement from 12:04–12:52 UT). On this day, the NO₂ monitor at ground level was not operational. NO₂ monitor data at 200 m were averaged over the lidar integration time. Bottom panel: Polluted day, September 12, 2006 (lidar measurement from 9:37–10:26 UT). For the NO₂ monitor data at ground level an average was made for the time the monitor was operational during this interval, from 10:03–10:17. For the data of the NO₂ monitor at 200 m two averages were determined; the lower average, for 9:37–10:17 UT, is for the situation that the NO₂ monitor is above the boundary layer, the higher value, for 10:16–10:26 UT, is for the situation that the NO₂ monitor is situated below the boundary layer.

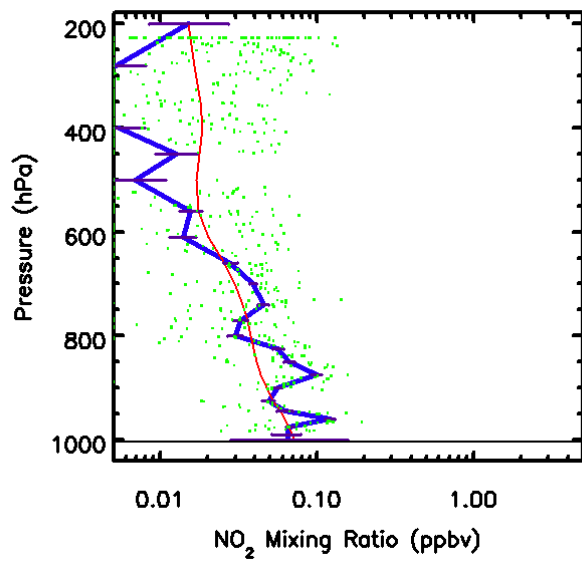
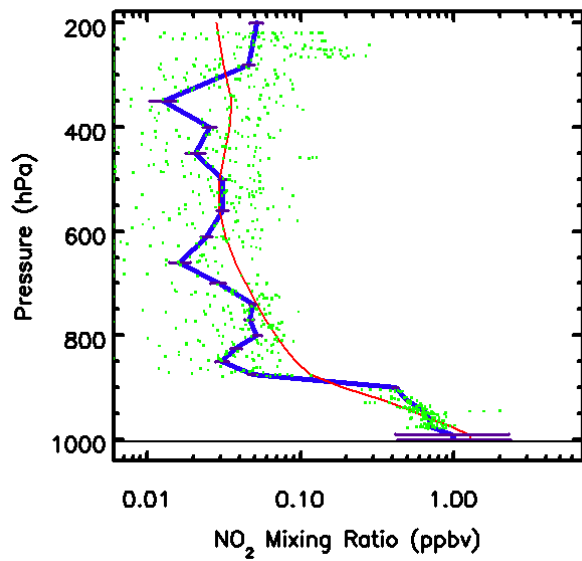


Figure 21. Two INTEX-B NO_2 profiles. The upper panel shows data for a profile measured over the Mississippi-Alabama border (32.0 N, 88.3 W), while the lower panel shows data for a profile measured over the Gulf of Mexico (23.0 N, 91.1 W). Green dots are original measurements, the blue line is binned profile, and the red line is the annual mean GEOS-CHEM model profile for that location.

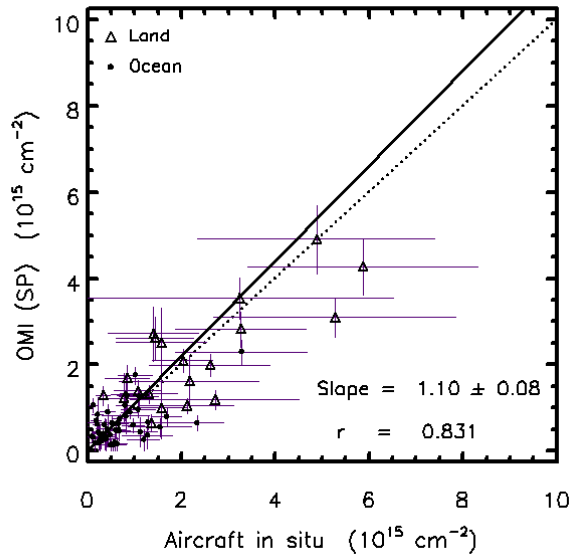


Figure 22. Tropospheric NO₂ columns with *in situ* columns from INTEx-B. The symbols indicate land measurements (triangle) or ocean measurements (dot). The dotted line is a 1:1 ratio and the solid line is the fit to the data.

Table 3. Summary of validation study results for OMI NO₂ data product.

Instrument	Column	Group	OMI, relative to GB	Remarks
SAOZ	Strat.	CNRS	+10%	As large as 50% in tropics.
MAX-DOAS	Trop.	BIRA, etc.	-15%	Large scatter in the data, so this number is quite uncertain.
Brewer	Total	GSFC	-33%	Large <i>N</i> , large scatter.
MF-DOAS	Total	WSU	-19%	Very small <i>N</i> .
Pandora-1	Total	GSFC	-15%	Very small <i>N</i> .
DS-DOAS	Total	BIRA	-16%	Small <i>N</i> , large scatter.
FTUVS	Total	JPL	-23%	Small <i>N</i> , but good correlation.
Aircraft <i>in situ</i>	Trop.	UC Berkeley, GSFC	+10%	Large scatter.