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**Tropospheric NO₂
columns**

A. Lauer et al.

Tropospheric NO₂ columns: a comparison between model and retrieved data from GOME measurements

A. Lauer¹, M. Dameris¹, A. Richter², and J. P. Burrows²

¹DLR Institut für Physik der Atmosphäre, Oberpfaffenhofen, D-82234 Wessling, Germany

²Institut für Umweltphysik, Universität Bremen, D-28359 Bremen, Germany

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Correspondence to: A. Lauer (Axel.Lauer@dlr.de)

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Print Version

Interactive Discussion

© EGS 2001

Abstract

Tropospheric NO₂ plays a variety of significant roles in atmospheric chemistry. In the troposphere it is one of the most significant precursors of photochemical ozone (O₃) production and nitric acid (HNO₃). In this study tropospheric NO₂ columns were calculated by the fully coupled chemistry-climate model ECHAM4.L39(DLR)/CHEM. These have been compared with tropospheric NO₂ columns, retrieved using the tropospheric excess method from measurements by the Global Ozone Monitoring Experiment (GOME) of up-welling earthshine irradiance and the extraterrestrial radiance. GOME is part of the core payload of the second European Research Satellite (ERS-2). For this study the first five years of GOME measurements have been used. The period of five years of observational data is sufficient to enable a comparison based on climatological averages and with global coverage, focussing on the geographical distribution of the tropospheric NO₂, for the first time.

A new approach of analysing regional differences (i.e. on continental scales) by calculating individual averages for different environments provides more detailed information about specific NO_x sources and of their seasonal variations. The results obtained enable the validity of the model NO₂ source distribution and the assumptions used to separate tropospheric and stratospheric parts of the NO₂ column amount from the satellite measurements to be investigated.

1. Introduction

Tropospheric NO₂ plays a key role in both stratospheric and tropospheric chemistry. In the troposphere the photolysis of NO₂ results in the formation of O₃ (e.g. Bradshaw et al., 2000). NO₂ can then be regenerated by catalytic cycles involving both organic peroxy radicals (RO₂), the hydroperoxyradical (HO₂), the hydroxyl radical (OH) and both volatile organic compounds (VOC) and carbon monoxide (CO). Thus NO₂ plays a significant role in determining the oxidising capacity of the troposphere.

Tropospheric NO₂ columns

A. Lauer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Print Version

Interactive Discussion

© EGS 2001

**Tropospheric NO₂
columns**A. Lauer et al.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[I◀](#)[▶I](#)[◀](#)[▶](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

© EGS 2001

Although direct absorption of ultraviolet and visible radiation is not thought to provide a large atmospheric forcing, local maxima of up to 0.1 to 0.15 Wm⁻² can be reached (Velders et al., 2001). As tropospheric O₃ is also a significant greenhouse gas, NO₂ also contributes indirectly to the radiative forcing.

5 The emission of NO_x (NO_x = NO + NO₂) into the troposphere is strongly influenced by human activities, NO_x being produced in significant amounts by industrial combustion and biomass burning (Lee et al., 1997), the natural source of NO_x being lightning and emissions from soils in the troposphere. NO₂ is known to impact on human health and the environment both directly and through the production of O₃ (e.g. EPA, 2000).
10 Overall it is necessary to monitor and understand the global impact of this pollutant on the physics and chemistry of the atmosphere.

The launch of GOME aboard the ERS-2 in April 1995 has enabled the global observation of the distribution of NO₂, which has significant amounts in both the stratosphere and the troposphere to be retrieved (Burrows et al., 1999). Further the development
15 of the tropospheric excess method has enabled tropospheric NO₂ columns to be retrieved on scales up to global for the first time (Burrows et al., 1999), (Richter and Burrows, 2001). This retrieved data product provides a set of long-term observational data, which are well suited for evaluating the quality of the results of chemistry-climate models.

20 In contrast to recent studies by Leue et al. (1999) and Velders et al. (2001) focussing on a single year (1997), in this study climatological averages of the tropospheric NO₂ columns retrieved from GOME have been used. These have been compared with those obtained from the interactively coupled chemistry-climate model ECHAM4.L39(DLR)/CHEM on global and regional scales. Monthly average values of the NO₂ tropospheric
25 columns retrieved using the TEM algorithm (“Tropospheric Excess Method”) from five years of GOME observations (January 1996 to August 2000) and 20 years of model output provide the data base.

This comparison of modelled and measured tropospheric NO₂ column amounts is the first step in evaluating the ability of ECHAM4.L39(DLR)/CHEM to simulate the

**Tropospheric NO₂
columns**A. Lauer et al.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[I◀](#)[▶I](#)[◀](#)[▶](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

© EGS 2001

5 tropospheric NO_x chemistry and to unveil still present deficiencies in chemistry and emission datasets. This first step is necessary to prepare future studies on the global impact of traffic (road and aircraft) induced NO_x emissions on climate and air chemistry as well as their contribution to the global NO_x budget in comparison to other man-made

10 (industry and biomass burning) and natural (soils and lightning) NO_x emissions. Once the chemistry-climate model ECHAM4.L39(DLR)/CHEM has been adjusted and evaluated to reproduce present and past global NO_x measurements, it becomes possible to do prognostic simulations of future scenarios, which cannot be provided by Chemical Transport Models.

15 To achieve optimum comparability of the two different data sources, the satellite data have been fitted to the lower resolution of the model grid. The tropospheric NO₂ columns from the model data have been calculated in two ways:

1. Integration from the surface up to the (thermal) model tropopause (“Thermal Tropopause”-method).
2. Separation of tropospheric and stratospheric NO₂ amount using the method applied to the satellite data (“Tropospheric Excess or Reference Sector Method”).

2. Data

2.1. Tropospheric NO₂ columns retrieved from GOME observations

20 GOME is a spectrometer on board ERS-2, which was launched on the 20th of April 1995 and flies in a sun-synchronous, polar orbit at an average height of 785 km above the earth’s surface (Burrows et al. (1999) and references therein). The GOME instrument observes in nadir viewing geometry the light (UV/visible) scattered back from the atmosphere and reflected at the ground. It also observes the extra terrestrial solar irradiance. The instrument is designed to observe simultaneously the spectral range

25 between 232 and 793 nm. The atmosphere is scanned by a moving mirror resulting in

**Tropospheric NO₂
columns**A. Lauer et al.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[I◀](#)[▶I](#)[◀](#)[▶](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

© EGS 2001

spatial resolution of 320 km x 40 km (across track x along track). As a result of the sun-synchronous orbit, the measurements in low and middle latitudes are always taken at the same local time (LT) (the northern mid-latitudes are crossed at about 10:45 am LT). Nearly global coverage is achieved within three days.

5 The trace gas retrieval of NO₂ is achieved using the DOAS technique (Differential Optical Absorption Spectroscopy). This technique utilises the atmospheric absorption, defined as the natural logarithm of the ratio of the extraterrestrial irradiance and the earthshine radiance, for a selected spectral window. This is compared with reference absorption spectra of gases absorbing in the spectral window and a polynomial of low
10 order. The polynomial describes the scattering and broad absorption in the window. The slant column of a gas is derived from the differential absorption of the gas in question and is the integrated concentration along the light path through the atmosphere. For this study, the spectral window from 425 to 450 nm has been used, the spectra of NO₂, O₃, O₄ and H₂O and a reference Ring spectrum being fitted (Richter and Burrows,
15 2001).

The resultant slant columns of NO₂ can be converted to vertical columns by the application of an air mass factor, AMF. The AMF describes the effective length of the light path through the atmosphere and is derived from radiative transfer calculations. The value of the AMF depends on the viewing geometry and the solar zenith angle, but also
20 on surface albedo, vertical gas profile, clouds and atmospheric aerosol. In this study, a constant vertical profile with all NO₂ in a 1.5 km boundary layer has been assumed. Stratospheric NO₂ is not included in the airmass factor calculation as the tropospheric slant columns have already been corrected for the stratospheric contribution, and the influence of stratospheric NO₂ on the radiative transfer can be neglected. The uncertainties introduced by this and other assumptions are discussed in detail in (Richter
25 and Burrows, 2001) and (Velders et al., 2001).

The Tropospheric Excess or Reference Sector Method for determining the tropospheric columns of NO₂ makes two assumptions:

a) the longitudinal distribution of stratospheric NO₂ is relatively homogeneous. This

**Tropospheric NO₂
columns**A. Lauer et al.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[I◀](#)[▶I](#)[◀](#)[▶](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

© EGS 2001

is reasonable at latitudes below 60° N through the year, because the bulk of the NO₂ in the stratosphere is at a relatively high altitude and as a result determined by photolysis and therefore mainly by day length which is a function of latitude only.

5 b) at remote locations, the tropospheric column of a gas is constant and negligibly small in the case of NO₂.

Thus the TEM tropospheric columns of NO₂ are determined by subtracting the NO₂ vertical column at a selected remote and clean location from that at other locations at the same latitude. In this study the reference clean sector is chosen to be around the international date line at longitude 180° W.

10 For this study, climatological monthly means of the tropospheric NO₂ column amounts (January 1996 to August 2000) have been used. The data were selected to be cloud free i.e. only pixels having a cloud coverage below the threshold value 10% were used to derive the tropospheric NO₂ column amounts from the GOME measurements (see also Sect. 4.2). (Version 1.0 of the IUP/IFE-UB TEM NO₂ Dataset.)

2.2. ECHAM4.L39(DLR)/CHEM

ECHAM4.L39(DLR)/CHEM (hereafter referred to as E39/C) is a spectral interactively coupled atmospheric chemistry - general circulation model. It has a horizontal resolution of T30 (3.75° x 3.75°) and 39 layers in the vertical direction extending from the surface up to 10 hPa (30 km). The chemistry module CHEM (Steil et al., 1998) includes 107 reactions and 37 trace gases in the troposphere and stratosphere. It is connected with the ECHAM4 radiation scheme via H₂O, O₃, CH₄, N₂O and CFCs. The system thereby allows feedbacks between chemistry and the radiation scheme, which in turn, affects dynamics.

25 The current chemical scheme within CHEM does not include the NO_x reservoir species PAN. In addition, CHEM neither includes VOC chemistry nor the heterogeneous reaction of N₂O₅ on the surface of wet aerosols in the troposphere forming

**Tropospheric NO₂
columns**

A. Lauer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I◀

▶I

◀

▶

Back

Close

Print Version

Interactive Discussion

© EGS 2001

HNO₃. This version of E39/C is specialized on stratospheric ozone chemistry. Nevertheless, this first look not only provides a detailed view into the current ability and deficiencies of E39/C to simulate tropospheric NO_x, but also enables the validation of the seasonal variation of currently used NO_x emission data sets as e.g. biomass burning. This step is essential to enable upcoming studies on the present, past and future global impact of man-made NO_x emissions (especially road-traffic and aircraft) in comparison to the natural (soils and lightning) NO_x emissions not only on the climate (e.g. tropospheric O₃ production) but also on air chemistry (e.g. OH budget) and air quality.

Possible effects of the limitations on tropospheric NO_x chemistry are discussed in Sect. 4.1.

For this study an existing dataset from the “1990” control experiment (Hein et al., 2001) has been used. The E39/C data used for this comparison represent the beginning of the 1990s. Therefore the model was run in quasi-equilibrium mode. Gas emissions, Sea Surface Temperature, SST and boundary conditions were assumed similar to those measured or determined for the year 1990. A detailed model description and model applications can be found in (Hein et al., 2001) and (Schnadt et al., 2001).

Table 1 summarizes the Nitrogen Oxide emissions as considered for this model simulation. The total sum equals 39.1 Tg(N)/yr. The emissions from industry and ground based traffic, which are predominantly emitted by the eastern United States, Central Europe and Japan, have the major contribution of about 58% of the global budget. (This dataset is based on version 1A of the GEIA global inventories of the annual emissions of NO_x from anthropogenic sources around the year 1985 (Benkovitz et al., 1996).) Especially in the tropics, biomass burning and lightning are the most important NO_x sources. In addition emissions from soils and aircraft are explicitly taken into account.

3. Model Data analyses and comparisons

In this study two methods have been used to calculate the tropospheric NO₂ columns from the model data. The first approach is to integrate the NO₂ concentration from the surface to the tropopause which is determined by employing the thermal WMO-criterion. This dataset is defined as the “Thermal Tropopause” dataset.

The second approach applies the TEM to the model data in a manner similar to that applied to the GOME observations. The averaged total column over a Pacific sector (170° W to 180° W) as estimate for the stratospheric amount (Richter and Burrows, 2001). The tropospheric column is calculated by subtracting this approximation of the stratospheric amount from the total columns.

The model results have been compared to each other. Although the “TEM” dataset yields smaller absolute values, the qualitative seasonal variation is not affected in any of the cases studied. Figure 1 shows the results of the two methods of calculation applied to the model output of E39/C. To calculate the climatological annual means, all 20 model years are used. As it can be seen easily, all major features of the global pattern are conserved. In both cases, the areas with high NO_x emissions (namely United States, Central Europe and Southeast Asia/Japan) are clearly visible. Even the distribution of the patterns of regions with lower values of the tropospheric NO₂ column amounts (e.g. Africa, South America, Australia) are (in a qualitative sense) similar.

Comparison of the two different methods of calculation indicates that:

1. For the “Thermal Tropopause” dataset, the regions with high values of the tropospheric NO₂ column amounts have a somewhat larger extent and higher maximum values than the results of “TEM”.
2. For the “TEM” dataset, negative values become possible in regions with low NO₂ column amounts, e.g. over the oceans.
3. In regions with very low tropospheric NO₂ column amounts (e.g. the oceans)

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Print Version

Interactive Discussion

**Tropospheric NO₂
columns**A. Lauer et al.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[I◀](#)[▶I](#)[◀](#)[▶](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

© EGS 2001

“TEM” has a large inherent error resulting from subtraction of two similar quantities.

4. The results over the continents are quite reasonable: the annual mean relative difference of both methods being below 30% for most of the examined regions.

5 The reason for the lower values obtained by TEM with model data is that significant concentrations of NO₂ are generated by the model at 170° W to 180° W. This appears to be outflow from the continents which is predominantly the case in the northern mid-latitudes in winter and results in a background of (1 to 7) × 10¹⁴ molec/cm² north of 60° S and < 1 × 10¹⁴ molec/cm² south of 60° S. In the remote maritime boundary layer
10 assuming the height of the PBL to be 2 km this would correspond to 20 to 150 pptv. This may indicate that in the model the NO₂ is not being removed rapidly enough. However, as mentioned before, the seasonal variation of the studied regions is not affected by “TEM”, because (especially in the northern mid-latitudes in winter) the tropospheric NO₂ column amounts are several times higher than the overestimation of stratospheric
15 NO₂ by the NO₂ above the reference sector. Figure 2 shows the averaged tropospheric NO₂ column amounts as modelled by E39/C above the reference sector over the pacific ocean.

3.1. Global comparison

For optimal comparison between the model results and the TEM dataset derived from GOME, in the following only model results obtained using TEM are compared to the GOME data (Fig. 3).
20

In January, both the satellite and the model data clearly show the large northern hemispheric NO_x emission areas. These are caused by anthropogenic emissions from domestic heating, industry and road traffic: USA (particularly the eastern part), Europe and Southeast Asia/Japan. These areas can be easily identified by the high values of
25 the tropospheric NO₂ column amounts.

**Tropospheric NO₂
columns**A. Lauer et al.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[I◀](#)[▶I](#)[◀](#)[▶](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

© EGS 2001

A significant difference can be seen between the model- and satellite data in these regions: E39/C produces larger maxima and the regions of enhanced tropospheric NO₂ column amounts have a larger spatial extent than those observed in the satellite data.

5 On the other hand, the high NO₂ areas in central and southern Africa (caused by biomass burning and lightning produced NO_x) of both data sources are in good agreement: the location and the absolute column amounts being similar.

In July, the tropospheric NO₂ column amounts reflect the reduction in the NO_x emission in North America, Europe and Asia: both the magnitude of the NO₂ clouds and their areas being reduced in size in comparison to those of January. Again, these areas have a larger extent and higher maxima in the model data than shown by the GOME data.

10 These observations are in general consistent with the observation that relatively high NO₂ values are found at 180° E/W and above the oceans in the E39/C dataset. This seems to indicate that the model is not destroying NO_x in the troposphere rapidly enough.

3.2. Analysis of the regional averages of NO₂ tropospheric columns

The tropospheric NO₂ columns exhibit a strong land-sea-contrast. To analyse regional differences and seasonal variations between the model and the TEM NO₂ GOME dataset, several regions of interest are chosen for further analysis (USA, Europe, Africa, Australia, South America, Southeast Asia/Japan) by selecting a suitable boundary. Each data point within this boundary that represents a NO₂ column above land is used to calculate a mean value for the domain. To differentiate between points over land and sea, the land-sea mask which is used by E39/C running at T30 resolution is utilised. This concept has been proven to give more reliable results when studying the seasonal variations than the standard method of calculating zonal means. This is explained primarily by the high spatial variability of the tropospheric NO₂ column amounts.

**Tropospheric NO₂
columns**A. Lauer et al.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[I◀](#)[▶I](#)[◀](#)[▶](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

© EGS 2001

To investigate the sensitivity of the calculated mean tropospheric NO₂ column values to the selection of spatial boundaries, the average values for each domain were calculated for boundaries, which were diminished or enlarged by one or two pixels in each direction. This showed that the relative differences are typically negligibly small and in all cases, the qualitative characteristics of the seasonal variations were not affected (Lauer, 2001).

Figure 4 shows the results of the regional comparison. These can be summarized as follows:

- For the region 'Europe', E39/C and GOME show similar small seasonal variations, but the annual mean of the results of E39/C ("TEM") is 2.65 times greater than that of GOME (Fig. 4a).
- In contrast to the GOME data, the model produces a relatively large and distinctive seasonal variation in the regions 'USA' (not shown) and '(eastern) USA' (Fig. 4b): maximum values occurring during the winter and minimum values during the summer. The results of E39/C show higher column amounts than the GOME data during the whole year, the annual mean is 1.7 times (USA) resp. 1.9 times ((eastern) USA) greater than that of GOME.
- The tropospheric NO₂ column amounts of E39/C and GOME for the region 'Africa' are both qualitatively and quantitatively similar. The annual mean of the GOME data is 1.2 times higher than that of E39/C ("TEM") (Fig. 4c).
- The tropospheric TEM NO₂ column amounts from GOME exhibit a distinctive seasonal variation for the region 'South America', having a minimum in May and a maximum in September. This is not reproduced by E39/C which shows only small seasonal variations. The annual mean of E39/C is 1.4 times greater than that of GOME (Fig. 4d).
- In 'Australia', the TEM NO₂ column data from GOME exhibit only a small seasonal variation, whereas E39/C shows a distinctive seasonal variation with minimum

Tropospheric NO₂ columns

A. Lauer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Print Version

Interactive Discussion

© EGS 2001

values occurring in the winter and maximum values in the summer. The annual mean of GOME is 1.7 times greater than that of E39/C (Fig. 4e).

- For the region 'Southeast Asia/Japan', the tropospheric NO₂ column amounts of E39/C and GOME are in good agreement. The only striking discrepancy is the month February. Whereas the values of GOME go down rapidly from January to February, the decrease between January and February shown by E39/C to a much lesser extent. The annual mean of E39/C is slightly higher than that of GOME, the relative difference is about 35% (Fig. 4f).

Table 2 summarizes the minimum, maximum and average relative difference of the TEM modelled and retrieved tropospheric NO₂ column amounts by E39/C and GOME, respectively. The relative difference *r* is calculated by (1):

$$r = \frac{'E39/C (TEM)' - 'GOME'}{'GOME'} \cdot 100\% \tag{1}$$

4. Discussion of results

There are several possible reasons for the observed differences between model and measurements. These can be divided into three basic error classes:

- model errors and deficiencies,
- errors from the GOME measurements and from the derivation of the tropospheric column amounts of the measured spectra,
- differences arising in the generation of the two data sources.

4.1. Model errors and deficiencies

How accurate and representative the model output is, depends on:

- the description of the atmospheric dynamics;
- the chemical scheme;
- the accuracy of the input data;
- the initialization data.

5 As NO₂ has a relatively short chemical short lifetime, the description of its chemical production and loss and the distribution and magnitude of sources are of significance.

The input sources are based on monthly means, which are assumed to remain constant during the whole period of simulation (20 years), except for lightning NO_x which is related to the model's cloud parameterisation scheme (Hein et al., 2001).

10 The uncertainties of the NO_x emissions in the total amount and the seasonal variation are somehow a measure for the reliability of the calculated NO_x volume mixing ratios, the basis for the calculation of the tropospheric NO₂ column amounts. Here, a simple phase shift of the used biomass burning dataset in the southern hemisphere (particularly South America and South Africa) by about one month could possibly improve the agreement of the modelled and observed seasonal variation of E39/C and GOME significantly (Lauer, 2001). In contrast, the seasonal variation of the biomass burning data set in (Northern) and Central Africa, which is dominating the seasonal variation of the total NO_x emissions in this region seems to be quite good as the seasonal variation of the modelled and the observed tropospheric NO₂ column amounts are in good agreement.

15 In Australia, very low values of the NO₂ column amounts are modelled. As a direct result of the uncertainties of the emissions, the modelled tropospheric NO₂ has large uncertainties. Here, even slight changes of the emissions could give a different seasonal variation. In addition to the general uncertainties of the NO_x emissions, the datasets employed in the model have no daily variation as e.g. caused by the rush hour in the morning and evening in Europe or the United States.

Tropospheric NO₂ columns

A. Lauer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Print Version

Interactive Discussion

© EGS 2001

**Tropospheric NO₂
columns**A. Lauer et al.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[I◀](#)[▶I](#)[◀](#)[▶](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

© EGS 2001

Even more important is the missing VOC chemistry and the missing NO_x sink process 'heterogeneous reaction of N₂O₅ with H₂O forming HNO₃' on the surface of tropospheric aerosols and at the earth's surface.

The missing VOC chemistry prevents the formation of the NO_x reservoir species PAN.

5 A model study by [Kuhn \(1996\)](#) examined the impact of PAN on the NO_x mixing ratios using the global 3-D Chemical Tracer Model, CTM, which has a horizontal resolution of 4° x 5° and 9 vertical layers extending from the surface up to 10 hPa. Two model simulations, with and without taking into account PAN, were compared. The results showed, that PAN decreases (increases) the NO_x mixing ratio in regions with high (low)
10 NO_x mixing ratio, e.g. above the continents (oceans). However, the results suggest the effect on the NO_x mixing ratio is below 10% above continents ([Kuhn, 1996](#)), and therefore probably even less on the tropospheric NO₂ column amounts. In contrast, the missing hydrolysis of N₂O₅ on tropospheric aerosols seems to be quite important, especially for the mid-latitudes.

15 [Dentener and Crutzen \(1993\)](#) studied the impact of the reaction of N₂O₅ on tropospheric aerosols on the global distributions of NO_x, O₃ and OH using the global 3-D chemical tracer model Moguntia, which has a horizontal resolution of 10° x 10° and 10 vertical layers at 100 hPa distance. They took into account the role of night time chemical reactions of NO₃ and N₂O₅ on aerosol surfaces and calculated the resulting loss in NO_x. Their results showed that this additional NO_x sink reduces the NOX
20 (NOX = NO + NO₂ + NO₃ + N₂O₅ + HNO₄) present in the boundary layer above the U.S. and Europe by about 50% in winter and about 20% in summer. The increased heterogeneous NOX removal in winter is due to long darkness and low temperatures during the winter months. For tropical regions e.g. Africa, this process seems to be much less important and reduces the boundary layer NOX only about 10 to 30%. This
25 can be explained by the increased importance of the daytime reaction of NO₂ with OH in tropical regions ([Dentener and Crutzen, 1993](#)).

In Europe, North America and Asia, large quantities of NO_x are released in winter into the planetary boundary layer. In the tropics convective activity pumps some of the

NO_x out of the planetary boundary layer and the lightning source releases NO_x above the planetary boundary layer, where the lifetime of NO_x is increased compared to the surface.

The current limitations within the model, describing the NO_x emissions and the chemistry in the lower troposphere, appear strong candidates to explain the differences between model and retrieved datasets.

4.2. GOME errors

Errors are introduced to the TEM NO₂ column dataset from GOME observations for two types of reason:

- Inherent uncertainties in the measurement itself and the retrieval of the trace gas concentration from the measured spectra.
- Remote sensing specific issues.

The errors from the measurement and the retrieval using DOAS are small in general and can usually be neglected compared to the other error sources.

In contrast, the errors due to remote-sensing specific problems are potentially more significant. To detect NO₂, GOME measures visible light (425 to 450 nm). The presence of clouds prevents the detection of NO₂ below the cloud, and enhances GOME's sensitivity for the detection of NO₂ above the cloud top. To minimize the impact of clouds, the retrieval of the TEM tropospheric NO₂ column amounts is restricted to atmospheric scenes having a cloud coverage below the threshold value 10%.

Errors arising from undetected clouds (i.e. sub pixel scene) and cloud fractions below the threshold value may lead to an underestimation of tropospheric NO₂ by up to 40%, although in most cases, this error is well below this peak value (Richter and Burrows, 2001). The presence of significant aerosols have a similar effect to that of clouds on the detection of NO₂.

A study by Martin et al. (2001) showed the potential of further improvements on the retrieval of tropospheric NO₂ from the GOME measurements. A new approach of using

Tropospheric NO₂ columns

A. Lauer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Print Version

Interactive Discussion

**Tropospheric NO₂
columns**A. Lauer et al.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

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5 a global 3-D model of tropospheric chemistry for the calculation of the vertical NO₂ profiles required to calculate the AMF instead of assuming a globally uniform vertical profile seems to improve the NO₂ retrieval. In addition, especially the new treatment of partly cloudy scenes should improve the retrieval of tropospheric NO₂. This new AMF formulation enables the quantitative retrieval for partly cloudy scenes (which is the case quite often because of the large dimensions of the individual GOME pixels) (Martin et al., 2001).

10 In addition, also the separation of the total column amounts into stratospheric and tropospheric part are a potential error source as additional assumptions have to be made. In order to see the effect on the tropospheric NO₂ column amounts of this method of calculation, the model data have been calculated using “TEM” as well.

4.3. Different assumptions in the creation of the two datasets

There are three major sources, which may result in differences when comparing the model- and satellite data and arise from differences in their:

- 15 – Temporal offset,
- exclusive use of clear sky conditions,
- sun synchronous orbit of ERS-2.

20 As mentioned above, all the input conditions used for this computer simulation are representative of the beginning of the 1990s, whereas the GOME measurements were made between 1996 and 2000. Thus a temporal offset of several years exists between the periods represented by the model and measured by GOME. Changes of the anthropogenic NO_x emissions resulting in different NO₂ concentrations might in part also explain the differences. However the uncertainties on the NO_x emission datasets used as input for the model are in any case large (see Table 1).

25 For the GOME data, only clear sky (threshold value for the cloud coverage of 10%) pixels are accounted for. In contrast, the tropospheric NO₂ column amounts from the

**Tropospheric NO₂
columns**A. Lauer et al.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[I◀](#)[▶I](#)[◀](#)[▶](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

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model output have been calculated without taking cloud effects or the cloud fraction within the model box explicitly into account.

The loss of NO_x by the reaction of N₂O₅ on aerosol and on cloud, which is significant at night, has been mentioned above. In addition as NO₂ is photolysed by the incoming solar ultraviolet radiation, the lack of cloud in the model and presence of some cloud in the measurement is likely to impact on the comparison. This effect is difficult to quantify, too, because of the large dimensions of the model boxes, only very few data points will remain after sorting out all values with a cloud fraction above 10%. Especially for the mid-latitudes, it might become impossible to calculate representative monthly means.

The probably most important fact making the comparison of the model- and satellite data difficult is the sun synchronous orbit of GOME's space platform, the ERS-2 satellite. Because of this, every measurement of GOME is performed at the same local time between 10:05 and 10:55 am. This period coincides with the minimum of the daily lightning activity over the continents as shown by long term observations of the Optical Transient Detector, OTD (Kurz, 2001). The simulation E39/C only provides averaged NO₂ values. Thus the modelled 24 hours average of the lightning produced NO_x will overestimate the NO_x present in the late morning. A 3-D chemical transport model study by Velders et al. (2001) indicates that the NO₂ tropospheric columns at 10:30 am are about 80% of the values averaged over 24 hours for the region Europe and the U.S., about 50 to 70% for the regions South America and Africa.

5. Conclusions

Overall the differences in the NO₂ tropospheric columns as retrieved from GOME observations and calculated by the general circulation model E39/C are within a factor of 2 to 3. The likely overestimation of the NO_x in the GCM is probably best explained by the lack of the heterogeneous loss of NO_x through the reaction of N₂O₅ with H₂O on aerosols and clouds in the lower troposphere. Thus, extending the chemistry module CHEM to properly handle the heterogeneous N₂O₅ chemistry should be the next step

in improving E39/C to perform simulations of tropospheric NO_x.

In spite of the deficiencies and various error sources of both model and satellite data, this investigation shows clearly the potential for testing the current capability of general circulation models to simulate the behaviour of the troposphere using satellite observations.

The major features of this first look can be summarized as follows:

In the northern mid-latitudes (USA, Europe), emissions from industry and ground based traffic are the most important source for NO_x.

In the tropics, especially biomass burning and emissions from lightning are the dominant NO_x sources. Here, the seasonal variation, as well as the quantitative column amounts are in better agreement than for the USA and Europe. This is consistent with the conclusion of Dentener and Crutzen (1993) stating much lesser influence of the still missing additional NO_x sink in the tropics.

Acknowledgements. This work is a contribution to TROPOSAT (EUROTRAC-2). It has been funded in part by the University and State of Bremen, the Ludwig-Maximilians-University München, the German Aerospace Center (DLR) and the European Union. The provision of level 1 GOME data by ESA for this scientific study is acknowledged.

This work was also supported by the project TRADEOFF of the European Union.

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Tropospheric NO₂ columns

A. Lauer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Print Version

Interactive Discussion

© EGS 2001

**Tropospheric NO₂
columns**A. Lauer et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Print Version

Interactive Discussion

© EGS 2001

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Tropospheric NO₂ columnsA. Lauer et al.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[I◀](#)[▶I](#)[◀](#)[▶](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

Tropospheric NO₂ columns

A. Lauer et al.

Table 1. Nitrogen Oxide emissions as used in the E39/C model simulation (Hein et al., 2001) and the range of uncertainty (Bradshaw et al., 2000)

Source	Emissions [Tg(N)/yr]	Range [Tg(N)/yr]	Contribution [%]	Reference
Industry	22.6	16-30	57.8	Benkovitz et al. (1996)
Soils	5.5	3-8	14.1	Yienger and Levy (1995)
Lightning	5.4±0.1 (clim. annual mean)	3.2-26 ¹	13.8	Price and Rind (1992)
Biomassburning	5.0	4-16	12.8	Hao et al. (1990)
Aircraft	0.6	0.5-0.6	1.5	Schmitt and Brunner (1997)
Total	39.1	26.7-80.6	100.0	

¹ free troposphere + near surface

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Print Version

Interactive Discussion

© EGS 2001

Tropospheric NO₂ columns

A. Lauer et al.

Table 2. Summary of the minimum, maximum and mean relative differences r (given by (1)) of the average values of the tropospheric NO₂ column amounts for the selected source regions. Positive (negative) values are corresponding to larger (smaller) column amounts by E39/C

region	mean (%)	min. (%)	month	max. (%)	month
USA	70.1	12.3	(7)	173.6	(2)
USA (eastern part)	90.5	37.6	(7)	162.7	(2)
Europe	164.7	66.6	(1)	287.4	(5)
Africa	-16.6	-9.1	(2)	-24.3	(10)
Central Africa	-3.7	0.1	(8)	-18.6	(11)
South Africa	-18.7	-2.0	(7)	-31.9	(4)
South America	41.7	-5.7	(9)	148.3	(6)
Australia	-40.4	-2.4	(3)	-81.8	(9)
Southeast Asia/Japan	34.8	6.1	(4)	95.4	(2)

[Title Page](#)
[Abstract](#)
[Introduction](#)
[Conclusions](#)
[References](#)
[Tables](#)
[Figures](#)
[Back](#)
[Close](#)
[Print Version](#)
[Interactive Discussion](#)

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**Tropospheric NO₂
columns**

A. Lauer et al.

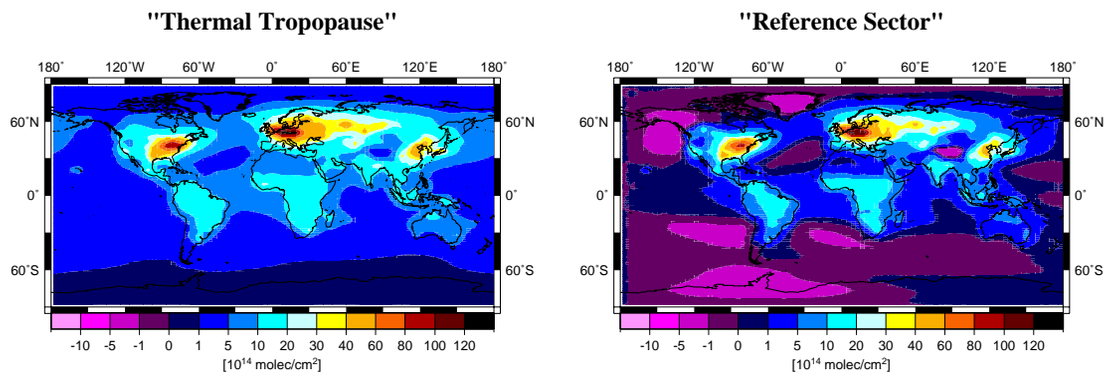


Fig. 1. E39/C, climatological annual means based on 20 years of the modelled tropospheric NO₂ column amounts, showing the results of the two different methods of calculation.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[I◀](#)[▶I](#)[◀](#)[▶](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

© EGS 2001

**Tropospheric NO₂
columns**

A. Lauer et al.

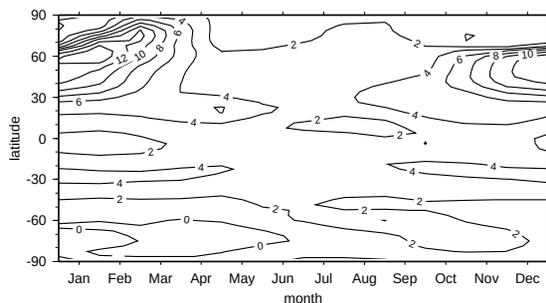


Fig. 2. Seasonal variation of the averaged climatological tropospheric NO₂ column amounts (10^{14} molec/cm²) for the reference sector over the Pacific ocean (170° W to 180° W) as modelled by E39/C (“Thermal Tropopause”-method). In contrast, “TEM” assumes no tropospheric NO₂ being present in the reference sector.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[◀](#)[▶](#)[◀](#)[▶](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

© EGS 2001

Tropospheric NO₂
columns

A. Lauer et al.

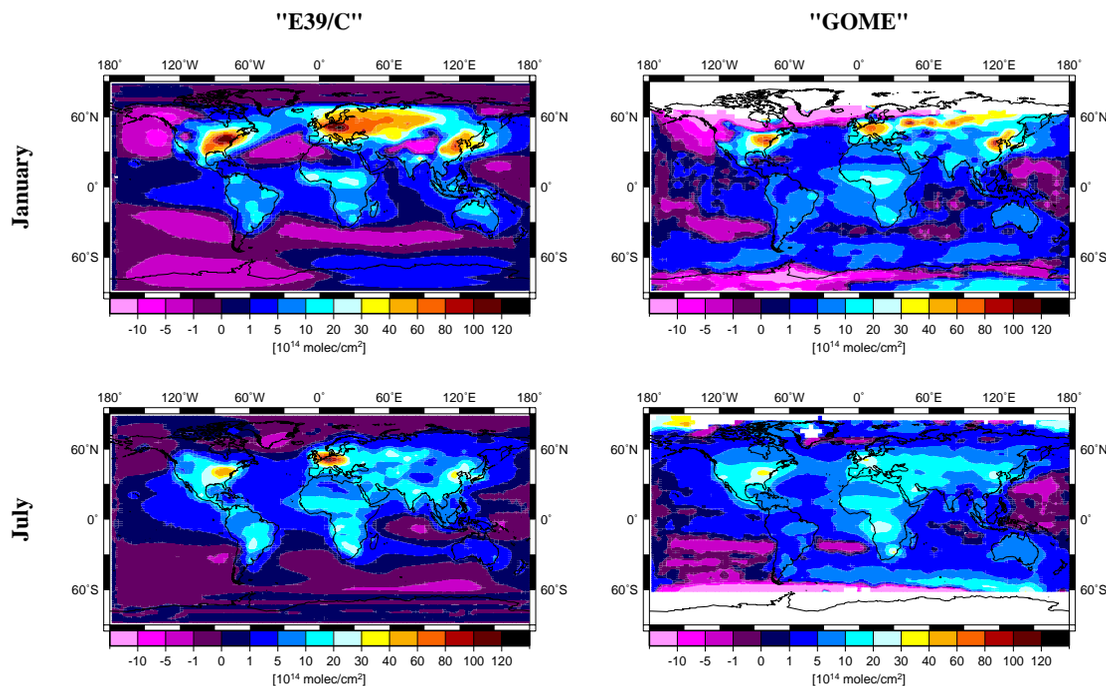


Fig. 3. Climatological monthly means of the tropospheric column amounts calculated by E39/C ('TEM') and derived from GOME measurements for January and July, respectively. Blank (white coloured) areas are data gaps.

[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[I◀](#)[▶I](#)[◀](#)[▶](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

© EGS 2001

Tropospheric NO₂
columns

A. Lauer et al.

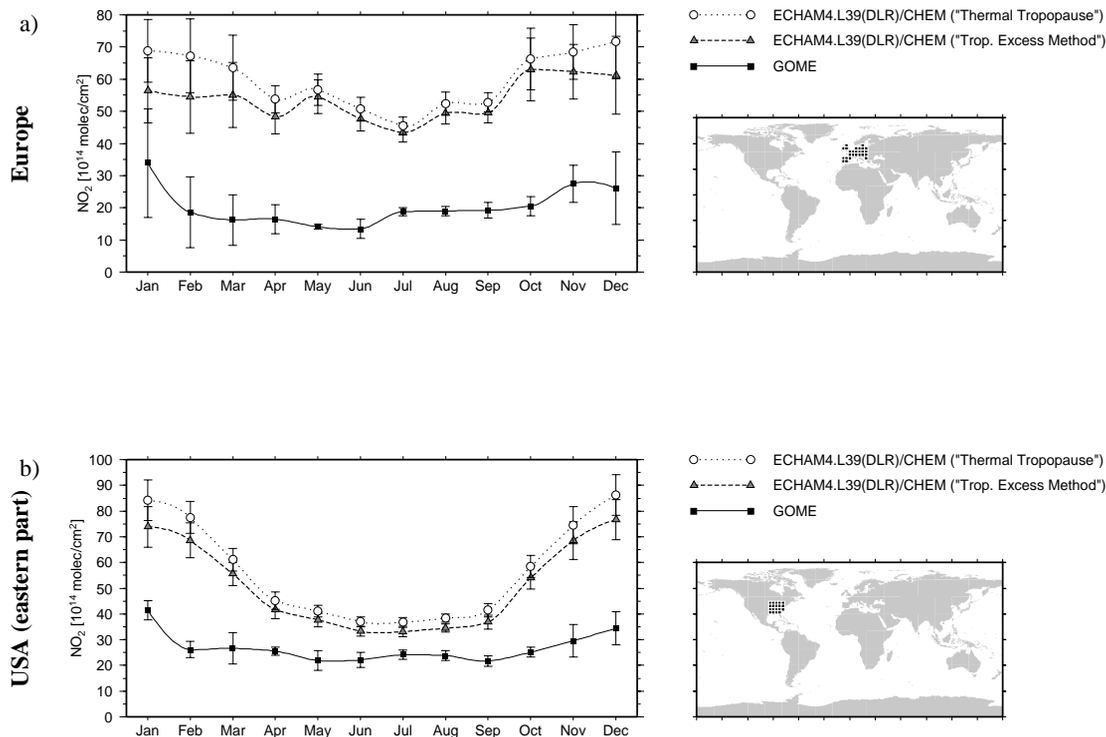


Fig. 4. (a – f): Seasonal variation of the tropospheric NO₂ column amounts for the climatological average values for selected spatial domains. The two sigma standard deviation of the individual monthly means to the climatological monthly means are drawn as errorbars for each data point. The small map to the right depicts the grid cells, that have been used to calculate the average values of the specified domain.

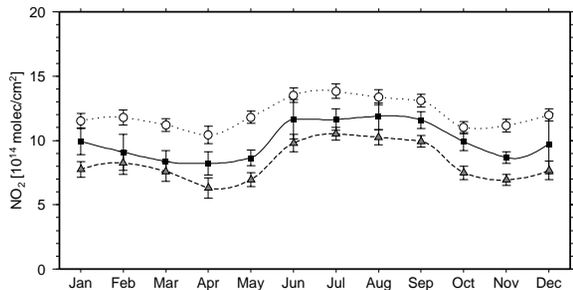
[Title Page](#)[Abstract](#)[Introduction](#)[Conclusions](#)[References](#)[Tables](#)[Figures](#)[◀](#)[▶](#)[◀](#)[▶](#)[Back](#)[Close](#)[Print Version](#)[Interactive Discussion](#)

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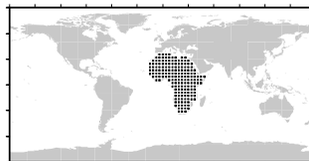
Tropospheric NO₂ columns

A. Lauer et al.

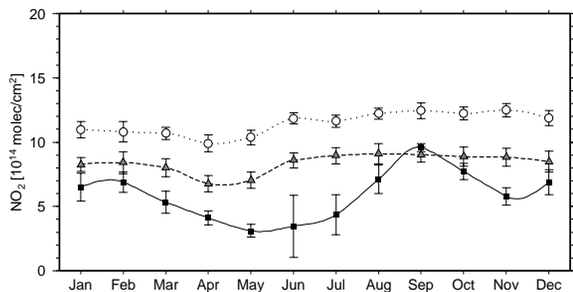
c) Africa



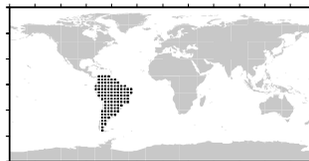
○·····○ ECHAM4.L39(DLR)/CHEM ("Thermal Tropopause")
 ▲- - -▲ ECHAM4.L39(DLR)/CHEM ("Trop. Excess Method")
 ■——■ GOME



d) South America



○·····○ ECHAM4.L39(DLR)/CHEM ("Thermal Tropopause")
 ▲- - -▲ ECHAM4.L39(DLR)/CHEM ("Trop. Excess Method")
 ■——■ GOME



Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

◀

▶

◀

▶

Back

Close

Print Version

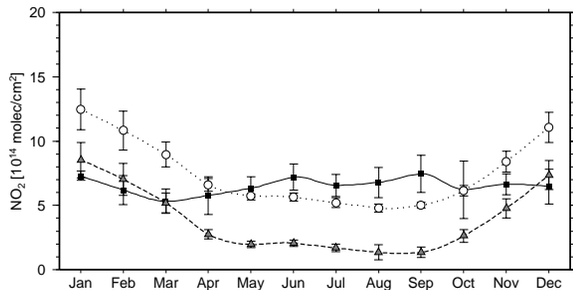
Interactive Discussion

Tropospheric NO₂ columns

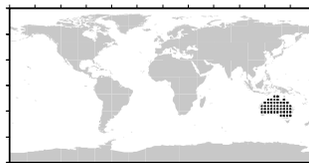
A. Lauer et al.

e)

Australia

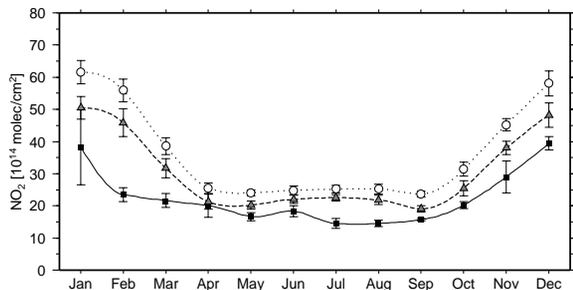


○·····○ ECHAM4.L39(DLR)/CHEM ("Thermal Tropopause")
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 ■——■ GOME

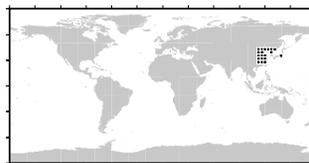


f)

Southeast Asia



○·····○ ECHAM4.L39(DLR)/CHEM ("Thermal Tropopause")
 ▲- - -▲ ECHAM4.L39(DLR)/CHEM ("Trop. Excess Method")
 ■——■ GOME



Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

⏪

⏩

◀

▶

Back

Close

Print Version

Interactive Discussion

© EGS 2001