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Satellite observations of formaldehyde over North America from GOME

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Abstract. Formaldehyde (HCHO) is an important indicator of tropospheric hydrocarbon emissions and photochemical activity. We present HCHO observations over North America for July 1996 from the GOME instrument on-board the ESA ERS-2 satellite. Slant columns are determined to \( \leq 4 \times 10^{15} \text{ molecules cm}^{-2} \) sensitivity by directly fitting GOME radiance measurements. These show a distinct enhancement over the southeastern United States, consistent with a large regional source from oxidation of non-methane hydrocarbons including in particular isoprene. Conversion of slant to vertical columns is done by combining species vertical distribution information from the GEOS-CHEM 3-D tropospheric chemistry and transport model with scattering weights from the Smithsonian Astrophysical Observatory LIDORT multiple scattering radiative transfer model. The results demonstrate the ability to measure HCHO from space in typical continental atmospheres, and imply that space-based measurements of HCHO may provide valuable information on emission fluxes of reactive hydrocarbons.

1. Introduction

Formaldehyde (HCHO) is a principal intermediate in the oxidation of hydrocarbons in the troposphere. Oxidation of methane (CH\(_4\)) by the hydroxyl radical (OH) is the principal source of HCHO in most of the troposphere; loss of HCHO is mainly by photolysis and reaction with OH. The corresponding atmospheric lifetime of HCHO is of the order of hours [Logan et al., 1981]. Since CH\(_4\) concentrations are relatively constant in the troposphere and photolysis is the dominant HCHO sink, when CH\(_4\) is the dominant source a strong correlation is expected between HCHO concentrations and 24-hour average OH concentrations. Under these conditions, HCHO serves as a proxy for tropospheric OH [Staffelbach et al., 1991].

In continental boundary layers, non-methane hydrocarbons (NMHCs) emitted by biogenic and anthropogenic sources usually dominate over CH\(_4\) as a source of HCHO [Munger et al., 1995; Lee et al., 1998] and make a large contribution to the HCHO atmospheric column. There is considerable uncertainty regarding NMHC emission fluxes. National U.S. emission estimates of biogenic isoprene, an important precursor for ozone (O\(_3\)) smog, have increased by a factor of 5 over the past decade [Guenther et al., 2000]. Emissions of other biogenic NMHCs are even more uncertain. Inventories of NMHC emissions from automobiles are also difficult to constrain [Pierson et al., 1990]. These uncertainties represent a major weakness in our ability to model ozone production [McKeen et al., 1991], OH concentrations [McKeen et al., 1997], formation of organic nitrates [Moxim et al., 1996], and formation of organic aerosol [Penner, 1995]. Because HCHO is a major product of NMHC oxidation and its lifetime is short, it provides a diagnostic of integrated NMHC emissions and thus a test of emission inventories.

The Global Ozone Monitoring Experiment (GOME), launched on the ERS-2 satellite in April, 1995, collects about 30,000 Earth radiance spectra each day. Spectra cover the ultraviolet (237–405 nm at 0.2 nm resolution) and the visible (407–794 nm at 0.4 nm resolution) [Burrows et al., 1993; European Space Agency, 1995]. Current operational data products available for GOME include total columns of O\(_3\) and NO\(_3\), and cloud fractional coverage.

Preliminary HCHO measurement capability has been presented for GOME in Thomas et al. [1998]. We demonstrate here that it is now possible to determine HCHO columns from GOME spectra to sufficient precision (a factor of 4 to 5 improvement) for it to be used in detailed process studies as a tracer of NMHC emissions to the troposphere. The analysis method described here is applied to HCHO over North America in July 1996, where the source from oxidation of isoprene is expected to be particularly large over the southeastern U.S. [Lee et al., 1998]. Under these conditions, most of the HCHO atmospheric column seen by GOME is in the continental boundary layer (this, and the details of the air-mass factor determination, are described in Palmer, P.I., D.J. Jacob, K. Chance, R.V. Martin, R.J.D. Spurr, T.P. Kurosu, I. Bey, R. Yantosca, A. Fiore, and Q. Li, Air-mass factor formulation for spectroscopic measurements from satellites: Application to formaldehyde retrievals from GOME, submitted to J. Geophys. Res., 2000 - hereafter, PIP), so that the GOME measurements offer a proxy for emissions of NMHCs including in particular isoprene.

2. Spectrum Fitting

Vertical columns amounts of HCHO are determined in a two-step procedure. GOME radiance spectra are fitted first using a nonlinear least-squares inversion based on the Levenberg-Marquardt method [Press et al., 1986]. The fitting process determines the slant column amounts that minimize the \( \chi^2 \) error between observed and calculated GOME radiances. Second, a combination of radiative transfer calculations using the Smithsonian Astrophysical Observatory (SAO) LIDORT code [Spurr et al., 2000] and results from the GEOS-CHEM global 3-D model of tropospheric chemistry and transport [Bey et al., 1999] is used to determine the appropriate air-mass factors (AMFs) which convert the...
levels. It is improved by using cross-correlation with the Fraunhofer reference spectrum [Caspar and Chance, 1997].

The GOME instrument suffers from substantial spectral undersampling. In regions where the atmospheric absorption is weak (e.g., that employed for HCHO fitting) this can largely be corrected using an undersampling correction spectrum generated by resampling an independent, high-resolution Fraunhofer reference spectrum [Chance, 1998; Slijkhuis et al., 1999]. The resulting HCHO fits have residual rms values of typically 5x10^{-4} of the full-scale GOME radiance spectrum.

HCHO slant columns are determined to <4.0x 10^{15} molecules cm^{-2} fitting precision (1σ uncertainties are used throughout). Overall accuracies are determined by adding 5% uncertainty for the HCHO cross sections [Canrell al., 1990] to the uncertainty from the fitting procedure, which is determined from the covariance matrix of the fitting errors and the rms of the fitting residuals. The fitting uses 300K cross sections. An additional 5% uncertainty is included to account for the temperature dependence of the cross sections; their integral changes by 5% over our spectral fitting range in going from 300K to 270K. This temperature range is appropriate for the HCHO observations presented here, considering that the bulk of the HCHO column is in the lower troposphere (PIP). Figure 1 shows spectral fitting examples fitted slant columns to vertical column abundances. The GEOS-CHEM model uses assimilated meteorological data for July 1996. Vertical profiles of HCHO are sampled from that model for the time and location of each GOME observation. The AMF calculations are described in a separate paper (PIP).

HCHO slant columns are determined from GOME spectra by direct fitting of GOME radiances, as described in Chance [1998]. Individual radiances are fitted, with no smoothing or filtering applied, over the wavelength region 337.35-356.12 nm. This fitting window was determined as the optimum compromise between large intensity for the HCHO cross sections and small spectral interference; it is slightly different from that used in Thomas et al. [1998]. Reference spectra for HCHO, including linear temperature dependence, are from Cantrell et al. [1990]. The fitting includes the interfering species O_3, NO_2, BrO, and the O_2-O_2 collision complex. O_3 and BrO are fitted in separately optimized spectral windows, where their stronger absorption features are located, in order to determine their contributions in the HCHO fitting window as accurately as possible. The Ring effect is included as described in Chance and Sprütt [1997]. Wavelength calibration of the GOME radiance and irradiance spectra in the level 1b operational data products is not sufficiently accurate to permit fitting to the required accuracy.
for low, moderate, and high concentrations of HCHO, plotted as the fitted absorption of HCHO with and without the fitting residuals.

3. Results

HCHO columns have been retrieved for 0°–75°N, 45°–180°W for the entire month of July 1996. Results for the North American region, 10°–60°N, 65°–130°W, selected for cloud fractional coverage of less than 40% and binned on a 2°x2.5° grid, are shown in Figure 2a, after division by geometric air-mass factors AMF, where

\[
\text{AMF} = \sec \theta_s + \sec \theta_v, \quad \theta_s \text{ is the solar zenith angle, and } \theta_v \text{ the satellite viewing zenith angle.}
\]

The solar zenith angles and solar times corresponding to these measurements range from 15°–35° and 1026–1154, respectively; the range of satellite viewing angles is 0°–31°. The resulting AMF values range from 2.0 to 2.4.

Figure 2b shows the vertical columns of HCHO obtained after determination of AMF that account for Rayleigh scattering, as described above. These AMFs are in the range 0.7–1.2, about half of the geometric AMFs because most of the HCHO is in the lower troposphere where scattering inhibits viewing. Variability in the shape of the vertical profile of HCHO in the GEOS-CHEM model induces ~20% variability in the AMFs. We find a large HCHO enhancement over southeastern United States, as expected from the strong regional source contributed by oxidation of isoprene [Lee et al., 1998]. This enhancement is manifest from the spectral analysis alone (Figure 2a) and is magnified by application of the scattering AMF (Figure 2b). This reflects the a priori information from the GEOS-CHEM model that most of the HCHO column is in the boundary layer (PIP). The HCHO columns over the southeastern U.S. in Figure 2b are in the range 2–3x10\(^{16}\) molecules cm\(^{-2}\). Assuming a typical HCHO scale height of 2 km [Lee et al., 1998], these columns correspond to boundary layer concentrations of 4–6 ppbv, which are consistent with in situ observations in that region [Munger et al., 1995; Lee et al., 1998].

In the future, this work will be expanded to include detailed studies of HCHO emissions over the U.S. in other years and seasons, and in greater temporal detail, to test estimates of isoprene emission fluxes and identify hot spots. The July 1996 data shown here contain much more detail about local production and transport than is provided in the present overview. We will also extend our analysis to the tropics, where biogenic and biomass burning sources of NMHCs are highly uncertain.

Future work should include improvement in the retrieval of HCHO. With improved HCHO sensitivity, or by merging results over selected regions, we will investigate HCHO in the remote marine troposphere. Better quantification of HCHO requires improvements in cloud and aerosol determination. Clouds mask the HCHO from observation. Scattering by aerosols alters the effective path length for HCHO absorption, enhancing or decreasing the observed HCHO absorption, depending on the measurement geometry, the vertical distribution of the aerosol relative to that of HCHO, and the surface albedo. For severe haze events, with scattering aerosol optical depths at 340 nm (\(\tau_{340}\)) of the order of 2 in the boundary layer, the AMF may increase by 40% (PIP). Improved cloud parameterization will come from the Cloud Retrieval Algorithm for GOME (CRAG) [Kurosu et al., 1999], which has been developed to provide cloud fractional coverage, cloud-top height, and optical thickness for the GOME scenes.

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