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Broadband cavity-enhanced absorption spectroscopy in the ultraviolet spectral region for measurements of nitrogen dioxide and formaldehyde

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Received: 20 August 2015 – Published in Atmos. Meas. Tech. Discuss.: 24 September 2015
Revised: 25 November 2015 – Accepted: 8 December 2015 – Published: 15 January 2016

Abstract. Formaldehyde (CH2O) is the most abundant aldehyde in the atmosphere, and it strongly affects photochemistry through its photolysis. We describe simultaneous measurements of CH2O and nitrogen dioxide (NO2) using broadband cavity-enhanced absorption spectroscopy in the ultraviolet spectral region. The light source consists of a continuous-wave diode laser focused into a Xenon bulb to produce a plasma that emits high-intensity, broadband light. The plasma discharge is optically filtered and coupled into a 1 m optical cavity. The reflectivity of the cavity mirrors is 0.99993 ± 0.00003 (1 − reflectivity = 700 ppm loss) at 338 nm, as determined from the known Rayleigh scattering of He and zero air. This mirror reflectivity corresponds to an effective path length of 1.43 km within the 1 m cell. We measure the cavity output over the 315–350 nm spectral region using a grating monochromator and charge-coupled device array detector. We use published reference spectra with spectral fitting software to simultaneously retrieve CH2O and NO2 concentrations. Independent measurements of NO2 standard additions by broadband cavity-enhanced absorption spectroscopy and cavity ring-down spectroscopy agree within 2 % (slope for linear fit = 1.02 ± 0.03 with \( r^2 = 0.998 \)). Standard additions of \( \text{CH}_2\text{O} \) measured by broadband cavity-enhanced absorption spectroscopy and calculated based on flow dilution are also well correlated, with \( r^2 = 0.9998 \). During constant mixed additions of NO2 and CH2O, the 30 s measurement precisions (1σ) of the current configuration were 140 and 210 pptv, respectively. The current 1 min detection limit for extinction measurements at 315–350 nm provides sufficient sensitivity for measurement of trace gases in laboratory experiments and ground-based field experiments. Additionally, the instrument provides highly accurate, spectroscopically based trace gas detection that may complement higher precision techniques based on non-absolute detection methods. In addition to trace gases, this approach will be appropriate for measurements of aerosol extinction in ambient air, and this spectral region is important for characterizing the strong ultraviolet absorption by brown carbon aerosol.

1 Introduction

Formaldehyde (CH2O) is the most abundant carbonyl compound in the atmosphere, with typical tropospheric concentrations between 0.1 and 10 parts per billion (ppbv; Grosjean et al., 1996; Fried et al., 2002). Its sources include secondary photochemical production from biogenic and anthropogenic volatile organic compounds (VOCs) and primary emissions from combustion sources. A Bayesian inverse model of satellite columns shows that the largest global CH2O source is oxidation of isoprene and other biogenic VOCs, followed by biomass burning and oxidation of anthropogenic VOCs (Shim et al., 2005). Formaldehyde is lost through photolysis...
and reaction with OH, with an atmospheric lifetime of a few hours. These processes contribute to net ozone production and radical recycling.

The existing in situ measurement techniques for formaldehyde can be broadly categorized as wet chemistry, mass spectrometry, and optical spectroscopy. The wet chemical methods include chromatographic detection of formaldehyde products with 2,4-dinitrophenylhydrazine (Fung and Grosjean, 1981; Lee et al., 1998; Spaulding et al., 2002) and fluorimetric detection of formaldehyde products from Hantzsch (Kelly and Fortune, 1994; Junkermann and Burger, 2006) or other reactions (Kok et al., 1990; Fan and Dasgupta, 1994). These wet chemical methods are sensitive, but they lack rapid time response and are susceptible to chemical interferences (Hak et al., 2005). Formaldehyde has been measured by mass spectrometry using proton transfer reaction mass spectrometry (PTR-MS; Karl et al., 2003; Steinbacher et al., 2004; Wisthaler et al., 2008; Warneke et al., 2011). These measurements are rapid and sensitive but are complicated by strong humidity dependence (Warneke et al., 2011). Spectroscopic measurements of formaldehyde exploit its strong absorption features in the infrared or ultraviolet spectral regions, with detection by Fourier transform infrared spectroscopy (Yokelson et al., 1996), tunable diode laser absorption spectroscopy (Fried et al., 1997, 2002), difference frequency generation absorption spectroscopy (Weibring et al., 2007), and quantum cascade laser spectroscopy (Herndon et al., 2007). Formaldehyde has also been measured spectroscopically by observing its fluorescence near 354 nm (Mohlmann, 1985; Hottle et al., 2009). Remote sensing instruments for ground-based (Platt et al., 1979; Heckel et al., 2005) and satellite-based (Bovensmann et al., 1999; Burrows et al., 1999) observations use differential optical absorption spectroscopy (DOAS).

Broadband cavity-enhanced absorption spectroscopy (BBCEAS) is a sensitive technique for direct absorption measurements. Compared to wet chemistry, mass spectrometry, and fluorescence, a key advantage of this approach is the inherent accuracy of direct absorption measurements. BBCEAS uses a broadband light source, typically a light emitting diode (LED) or arc lamp, coupled to a high-finesse optical cavity (Fiedler et al., 2003). The light output from the cavity is dispersed with a monochromator and measured using a multichannel detector. Conceptually, it is similar to a conventional optical absorption instrument with a multipass cell, but the high-finesse cavity allows an effective path length of several kilometers in the ultraviolet spectral region and tens or hundreds of kilometers in the visible spectral region. Standard spectral fitting methods can be used to retrieve multiple absorbers because absorption information is acquired simultaneously for a wide wavelength region (Ball et al., 2004; Langridge et al., 2006; Venables et al., 2006; Washenfelder et al., 2008; Thalman and Volkamer, 2010) and near-ultraviolet spectral range (Gherman et al., 2008; Langridge et al., 2009; Axson et al., 2011; Kahan et al., 2012; Washenfelder et al., 2013). However, only a few measurements have been demonstrated in the ultraviolet spectral region below 340 nm. These include measurements of O$_3$, O$_4$, and SO$_2$ at 335–375 nm (Chen and Venables, 2011), measurements of nitrophenols at 320–450 nm (Chen et al., 2011), and measurements of acetone near 280 nm (Islam et al., 2013).

Despite the distinct advantages of BBCEAS that have made it a powerful technique for trace gas measurements in the visible spectral region, there are multiple challenges in the ultraviolet that have precluded the measurement of atmospheric trace gases or aerosol extinction at ambient levels. First, LED sources are widely used at visible wavelengths because they are spectrally powerful, inexpensive, lightweight, and consume little power. However, LEDs with high optical power are unavailable at wavelengths shorter than 365 nm. Second, the optical bandpass filters required to reject out-of-band light from broadband sources are less efficient in the ultraviolet spectral region. Third, Rayleigh scattering by ambient air increases inversely with the fourth power of wavelength, $\lambda^{-4}$, and can become a significant loss process that limits light intensity and optical path length for an optical cavity in the ultraviolet spectral region. Finally, the thin film coatings used in the fabrication of high reflectivity cavity mirrors are less efficient in the ultraviolet spectral region, with greater absorption and scattering losses in the material. To the extent that these challenges can be overcome, BBCEAS measurements in the ultraviolet spectral region may be a powerful technique for atmospheric spectroscopy due to the large number of trace gases with structured absorption in this region and to the importance of aerosol absorption.

In this work, we report the first laboratory measurements of formaldehyde by BBCEAS in the ultraviolet spectral region at 315–350 nm. The approach uses a novel laser-driven Xe plasma as the high-power ultraviolet light source. This source eliminates some of the issues associated with conventional arc lamps, including large size, heat dissipation, and instability in the arc location. Although challenges remain for atmospheric sampling, this approach brings the CH$_2$O detection limit into the range of ambient concentrations. The subsequent sections describe instrumentation, data analysis, and results. The design requirements for ultraviolet BBCEAS to achieve trace gas detection at background CH$_2$O mixing ratios with 1 min time resolution are discussed in the conclusions.
2 Experimental

2.1 Description of the BBCEAS instrument

As shown in Fig. 1a, the BBCEAS optical system consists of a laser-driven arc lamp, collimating optics, optical filters, cavity, and grating spectrometer with charge-coupled device (CCD) array detector. This is similar to the laboratory instrument described previously (Washenfelder et al., 2008; Axson et al., 2011; Kahan et al., 2012), although the light source, collimating optics, optical filters, and grating spectrometer have been replaced to optimize measurements from 315 to 350 nm.

We use a broadband light source (EQ-99FC LDLS; Energetiq, Woburn, MA, USA), consisting of a continuous-wave diode laser at 974 nm that pumps a Xenon plasma (Islam et al., 2013) with a total electrical power draw of 125 W. The resulting plasma size is less than $100 \times 200 \mu m$, with spectral output from 170 to 2100 nm. Further details about laser-driven arc lamps and a comparison to conventional Xenon arc lamps can be found in Zhu and Blackborow (2011). The light source is air-cooled, but temperature drift caused the output intensity to vary as much as 10% in 1 h. To eliminate this drift, we constructed custom temperature control using water circulation through an attached aluminum plate. Change in lamp intensity as a function of wavelength is an additional consideration, and we measured the relative intensity change over 315–350 nm to be less than 0.3% in 1 h. The light source is purged with a continuous flow of N$_2$ gas to eliminate ozone production in the lamp housing. Inside the housing, the light is collected using an ellipsoidal reflector and a 600 µm diameter fiber, resulting in a manufacturer-specified power output of 130 µW nm$^{-1}$ across the 315–350 nm spectral region. This light is collimated and coupled into the cavity using an off-axis parabola with 0.36 numerical aperture (RC04SMA-F01; Thorlabs, Newton, NJ, USA). Prior to entering the cavity, the light passes through two colored glass filters (Schott Glass WG320 and UG11).

The optical cavity is formed by two 2.5 cm diameter, 1 m radius of curvature mirrors (Layertec GmbH, Mellingen, Germany), with manufacturer reported reflectivity of 0.9995 (1−reflectivity = 500 ppm loss) at their nominal center wavelength of 330 nm. The mirrors were mounted at either end of a 100 cm long Teflon cell (2.5 cm outside diameter), with gas ports for input, output, and mirror purge flows. The optical cavity is mechanically stabilized using 2 cm diameter carbon fiber rods and custom optical mounts. The light exiting the cavity is imaged by a 2.54 cm diameter F/3.1 lens onto a 0.5 cm F/2 lens (74-UV; Ocean Optics, Dunedin, FL, USA) that couples the light into one lead of a custom fiber bundle. There is an additional shortpass filter to eliminate stray light (XUS0350; Asahi Spectra USA, Inc., Torrance, CA, USA).
The optical fiber bundle consists of two leads, with seven 200 µm diameter fibers each. These are arranged linearly along the slit axis of the grating spectrometer, with each lead illuminating a separate region of the CCD. Spectra were acquired using a 203 mm focal length grating spectrometer (IsoPlane-160; Princeton Instruments, Trenton, NJ, USA) with an internal shutter, tunable grating, and manual entrance slit. The astigmatism of the IsoPlane-160 is less than 100 µm over the 26 mm wide focal plane. During these experiments, we used a 1200 line mm\(^{-1}\) grating (300 nm blaze) and a 100 µm entrance slit width. The CCD array is a 16 bit, back-illuminated detector with 2048 × 512 pixels (PIXIS 2kBUV; Princeton Instruments, Trenton, NJ, USA), which is thermoelectrically cooled to −70 °C to reduce thermal noise. The PIXIS 2kBUV had negligible time-dependent thermal noise, and its dark background was dominated by the offset voltage and readout noise.

The two regions of the CCD were used to measure the light output by the laser-driven arc lamp and the intensity through the optical cavity, as shown in Fig. 1. This allowed monitoring for drift in the light intensity. Unlike in previous experiments, the signal from each region was not accumulated in hardware to generate two spectra. Instead, the signal for the entire CCD was read out and the two regions were summed in software, which allowed for greater integration times and protected the shutter assembly from overheating during mirror purges. The required warm up time could be reduced in the future with improved temperature control. Dark spectra were acquired with identical integration time (1.2 or 2.0 s) as the sample spectra and 5–10 dark spectra were averaged together. The total acquisition time for each sample spectrum was 1 min, which included the shutter compensation and CCD readout time, allowing 35 spectra with 1.2 s integration time to be acquired in 1 min (equivalent to a duty cycle of 70 %). This duty cycle does not include the intermittent measurement of dark spectra.

2.2 Cavity ring-down spectrometer for NO\(_2\) validation

Concentrations of NO\(_2\) were independently measured using cavity ring-down spectroscopy at 406.4 nm, as described previously (Washenfelder et al., 2008; Fuchs et al., 2009). Briefly, a continuous-wave diode laser is modulated at 1.6 kHz and coupled into a high finesse cavity. The output light is measured with a photomultiplier tube and recorded with a data acquisition card. Measurements of the laser line shape showed that it was centered at 406.4 nm and was approximately Gaussian with FWHM of 0.48 nm. The laser line shape was convolved with a high-resolution NO\(_2\) reference cross section (Vandaele et al., 1998).

2.3 Preparation and delivery of CH\(_2\)O and NO\(_2\)

Formaldehyde was prepared by purification of paraformaldehyde, following the general method described in Meller and Moortgat (2000). Paraformaldehyde was gently heated in a glass tube connected to a vacuum system. Water from the paraformaldehyde decomposition was trapped in a cold finger, while formaldehyde was collected in a 12 L glass bulb. The bulb was subsequently diluted with N\(_2\), for a final concentration of 1–2 ppmv CH\(_2\)O in N\(_2\). Because formaldehyde readily polymerizes at high concentrations (Meller and Moortgat, 2000), the diluted formaldehyde sample was stored in dark conditions at room temperature. Nitrogen dioxide was prepared by flow dilution from a standard cylinder containing 15.8 ppmv NO\(_2\) in N\(_2\) (Scott-Marrin, Inc., Riverside, CA, USA).

The experimental flow system is shown in Fig. 1b. The BBCEAS and CRDS instruments were connected in series. Prior measurements under dry conditions have shown negligible losses to Teflon and metal surfaces for NO\(_2\) (Fuchs et al., 2009) and glyoxal (a dialdehyde with greater Henry’s Law constant than formaldehyde; Washenfelder et al., 2008; Min et al., 2015). A study of formaldehyde sampling artifacts similarly concluded it is relatively inert to adsorption by Teflon (Wert et al., 2002). Pressure (626 A; MKS Instruments, Andover, MA, USA) and temperature (K-type thermocouple; Omega Engineering, Inc., Stamford, CT, USA) were monitored at the exit of the CRDS cavity and used to calculate number density for the BBCEAS and CRDS. Over the experimental concentration range, CH\(_2\)O and NO\(_2\) did not measurably affect mirror cleanliness or reflectivity, so mirror purges were not used for the BBCEAS instrument. Mirror purges were necessary for the CRDS due to its existing cell design. Typical flows of 2.0 standard L min\(^{-1}\) (slpm) of zero air; 0–20 standard cm\(^3\) min\(^{-1}\) (scm) of 15.8 ppmv NO\(_2\); and 0–40 scm of 1–2 ppmv CH\(_2\)O were introduced to the cells and allowed to exhaust to ambient atmospheric pressure.

2.4 Operation

During normal laboratory operation, the BBCEAS and CRDS were turned on at least 2 h prior to beginning measurements. This allowed sufficient time for the laser-driven arc lamp and its temperature to stabilize. The required warm up time could be reduced in the future with improved temperature control. Dark spectra were acquired with identical integration time (1.2 or 2.0 s) as the sample spectra and 5–10 dark spectra were averaged together. The total acquisition time for each sample spectrum was 1 min, which included the shutter compensation and CCD readout time, allowing 35 spectra with 1.2 s integration time to be acquired in 1 min (equivalent to a duty cycle of 70 %). This duty cycle does not include the intermittent measurement of dark spectra. He and N\(_2\) spectra were measured for 1 min each to calculate the mirror reflectivity of the BBCEAS channel. During standard additions, reflectivity measurements were repeated at 15–20 min intervals, and zero air measurements were repeated between each standard addition.
3 Data analysis

3.1 Determination of mirror reflectivity and extinction

Extinction in a BBCEAS cavity can be described using an infinite sum for the light transmission through the cavity (Fiedler et al., 2003) or a differential equation for light input and output (Washenfelder et al., 2008). These two approaches give an equivalent expression that relates the optical extinction, $\alpha(\lambda)$, to the observed change in transmitted light intensity through the cavity. If we define the reference spectrum as a cavity filled with zero air, the expression is

$$\alpha(\lambda) = \left(1 - \frac{R(\lambda)}{d}\right) + (\lambda)_{\text{Rayleigh,ZA}} \left(\frac{I_{\text{ZA}}(\lambda) - I(\lambda)}{I(\lambda)}\right),$$

where $\lambda$ is the wavelength of light, $d$ is the cavity length, $R(\lambda)$ is the mirror reflectivity, $\alpha(\lambda)_{\text{Rayleigh,ZA}}$ is the extinction due to Rayleigh scattering of zero air, $I_{\text{ZA}}(\lambda)$ is the reference spectrum, and $I(\lambda)$ is the measured spectrum at each wavelength. If mirror purge volumes were included, the right-hand side of Eq. (1) would require a multiplicative factor equal to $d/L$, where $L$ is the length over which the absorbing sample is present. Eliminating the mirror purges in the laboratory instrument described here eliminates a source of uncertainty.

Calculating extinction from Eq. (1) requires knowledge of the mirror reflectivity, $R(\lambda)$. This can be determined by introducing a well-known extinction into the cavity. Rayleigh scattering is convenient because it varies slowly with wavelength across the entire region of interest, and requires only measurements of temperature and pressure. Here, we use measurements of the Rayleigh extinction for helium and zero air to calculate $R(\lambda)$. Based on the recommendation of Thalmann et al. (2014), we have used theoretical Rayleigh scattering cross sections from Sneep and Ubachs (2005; see Eqs. 11, 12, 23, and 24) for N$_2$ and O$_2$. The Rayleigh scattering cross section for He is based on an empirical fit to measurements by Shardanand and Rao (1977), $\sigma_{\text{Rayleigh,He}} = 1.336 \times 10^{-17} \times \lambda^{-4.1287}$. Because the mirror reflectivity is calibrated using well-known Rayleigh scattering cross sections (uncertainty ± 4% in the UV), the derived trace gas absorption or aerosol extinction retains high accuracy, similar to conventional direct absorption or extinction instruments.

3.2 Spectral retrieval method

The measured extinction in Eq. (1) contains contributions from each of the absorbers in the cavity:

$$\alpha(\lambda) = \sum_{i}^{n} \sigma_{i}(\lambda) N_{i} = \sigma_{\text{CH}_2\text{O}}(\lambda)\left[\text{CH}_2\text{O}\right] + \sigma_{\text{NO}_2}(\lambda)\left[\text{NO}_2\right] + \ldots,$$

where $n$ is the number of absorbers, $\sigma_{i}(\lambda)$ is the absorption cross section, and $N_{i}$ is the number density of the $i$th absorber. The number density of each absorber can be determined from the measured extinction, $\alpha(\lambda)$, and reference spectra for each absorber, $\sigma_{i}(\lambda)$.

For the spectral retrieval, reference spectra for CH$_2$O (Meller and Moortgat, 2000) and NO$_2$ (Vandaele et al., 1998) were convolved with a Gaussian line shape with FWHM 0.47 nm. These reference spectra were used with DOAS Intelligent System (DOASIS) spectral fitting software (Kraus, 2006) to retrieve number densities of CH$_2$O and NO$_2$. The spectra were fit from 315 to 350 nm, and a fourth-order polynomial was included in each fit to account for drift in the light intensity and cavity throughput.

4 Results and discussion

4.1 Measured spectra and calculated mirror reflectivity

Figure 2a shows spectra obtained with the absorption cell filled with He and zero air for 24 spectra with 2.0 s integration time. The spectral intensity is determined by the product of the laser-driven arc lamp spectrum (170–2100 nm), the three optical filters, and the transmission of the cavity mirrors. The intensity is lower near 330–340 nm where the mirror reflectivity is higher and the mirror transmission is lower. The accumulated counts at 330 nm were approximately $4.3 \times 10^8$ in 2.0 s. Figure 2b shows the mirror reflectivity calculated from Eq. (1). The peak reflectivity is 0.99930±0.00003 (700 ppm loss) at 338 nm and 0.99922 ± 0.00004 at 330 nm. This is slightly lower than the manufacturer’s specifications of 0.9995 at 330 nm. The repeatability of this result, even after standard procedures for careful mirror cleaning, suggests that it does not arise from surface deposits on the mirrors. This discrepancy has been reported previously for BBCEAS instruments and has been attributed to photons propagating off-axis in the cavity or populating very high-order transverse modes with greater losses than lower-order modes (Varma et al., 2009).

For a 100 cm cavity containing 830 hPa of zero air with $R = 99.930\%$ mirrors at 338 nm, the per pass fractional losses are 700 ppm mirror loss and 73 ppm Rayleigh scattering loss. The measured transmission through a single mirror is 210 ppm at 338 nm, indicating that absorption and scattering are 490 ppm and dominate the losses in the cavity. This...
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Figure 2. (a) Spectral measurements of \( I_{\text{He}} \) and \( I_{\text{ZA}} \), and the ratio of \( I_{\text{ZA}}/I_{\text{He}} \). (b) Cavity loss, \( 1 - R(\lambda) \), and effective pathlength, \( d/(1 - R(\lambda)) \), determined from the Rayleigh extinction of He and zero air spectra, as described in the text. (c) Reference spectra for \( \text{CH}_2\text{O} \) (light blue; Meller and Moortgat, 2000) and \( \text{NO}_2 \) (light red; Vandaele et al., 1998) convolved with a Gaussian lineshape with FWHM 0.47 nm (\( \text{CH}_2\text{O} \) dark blue; \( \text{NO}_2 \) dark red). The \( \text{CH}_2\text{O} \) cross sections have been multiplied by three to put them on the same scale with those of \( \text{NO}_2 \).

the calculated extinction for different \( \text{CH}_2\text{O} \) concentrations, with fits determined by DOASIS, are shown in Fig. 5 with fitting residuals in the upper panel. The average fit uncertainty is 400 pptv \( \text{CH}_2\text{O} \) in 1 min. The maximum absolute absorption cross section of \( \text{CH}_2\text{O} \) at this resolution is \( 7.6 \times 10^{-20} \text{ cm}^2 \text{ molecule}^{-1} \) at 326.0 nm. \( \text{CH}_2\text{O} \) is a relatively weak absorber compared to other species, such as glyoxal, whose peak cross section at 455 nm is 7 times greater. The weak absolute cross section of \( \text{CH}_2\text{O} \) makes it one of the more challenging UV absorbers to retrieve, despite its strong differential absorption in the ultraviolet spectral region, where the differential cross section is 91.0 % of the absolute extinction at 326.0 nm.

The correlation plot for \( \text{CH}_2\text{O} \) is shown in Fig. 6. In this case, \( \text{CH}_2\text{O} \) is compared to a scaled concentration based on flow dilution, rather than to an independent measurement, because the \( \text{CH}_2\text{O} \) concentration in the bulb is not known accurately. For this comparison, \( r^2 = 0.9998 \).

4.4 Simultaneous measurements of \( \text{NO}_2 \) and \( \text{CH}_2\text{O} \)

The dominant ambient lower atmospheric absorbers in this spectra region are \( \text{NO}_2 \) and \( \text{CH}_2\text{O} \), and ambient samples will likely contain a mixture of both. We introduced mixtures of these two species to the BBCEAS and CRDS cells using the flow system shown in Fig. 1b. A time series with the sequential introduction of 9.7 ppbv \( \text{NO}_2 \), 15.6 ppbv \( \text{CH}_2\text{O} \), 5.5 ppbv \( \text{NO}_2 \), and 7.5 ppbv \( \text{CH}_2\text{O} \) is shown in Fig. 7. Each contrasts sharply with higher quality mirrors available in the visible spectral region, where cavity measurements at 455 nm have demonstrated 34 ppm mirror loss and 19 ppm Rayleigh scattering (Washenfelder et al., 2008).

4.2 Measurements of \( \text{NO}_2 \)

Using the flow system shown in Fig. 1b, standard additions of \( \text{NO}_2 \) diluted with zero air were introduced to the BBCEAS and CRDS instruments in series. The calculated extinction for different \( \text{NO}_2 \) concentrations, with fits determined by DOASIS spectral fitting, are shown in Fig. 3, with fitting residuals shown in the upper panel. The average fit uncertainty is 360 pptv \( \text{NO}_2 \) in 1 min.

We note that the differential absorption features for \( \text{NO}_2 \) are weaker in this spectral region compared to the visible region of the spectrum. For example, the maximum differential \( \text{NO}_2 \) cross section in this region is 24.7 % of the absolute extinction at 348.5 nm (spectral resolution 0.47 nm). In contrast, the differential cross section is 54.0 % of the absolute extinction at 448.1 nm, and our previous measurements at 440–470 nm demonstrate a detection sensitivity (1 \( \sigma \)) of 40 pptv in 5 s (Washenfelder, 2008, 2011; Min et al., 2015). Part of this difference is due to the spectrally intense LED light source and lower loss mirrors, but part is also due to the larger differential \( \text{NO}_2 \) cross section in the visible spectral region.

The correlation plot of the independent \( \text{NO}_2 \) measurements is shown in Fig. 4. The slope is 1.024 ± 0.027 and the intercept is −0.20 ± 0.32 ppb, with \( r^2 = 0.998 \). Despite the weaker differential absorption features, the ultraviolet BBCEAS instrument accurately quantifies the \( \text{NO}_2 \) concentrations relative to the more precise CRDS instrument and is within the combined uncertainty of the Rayleigh scattering mirror reflectivity calibration and the \( \text{NO}_2 \) absorption cross section (±5.1 %; see Sect. 4.5).
Figure 3. Measured $\alpha(\lambda)$ for 1 min standard additions and calculated spectral fits for seven NO$_2$ concentrations from 6 to 39 ppbv. Two fitting residuals are shown in the upper panel. The average fit uncertainty for all concentrations shown is 360 pptv NO$_2$.

Point represents the average of 17 spectra with 1.2 s integration time (30 s total acquisition time). The decrease in the measured CH$_2$O concentration during the first addition is due to drift in the output from the pressurized CH$_2$O bulb. During the second set of additions, the 1σ standard deviations for NO$_2$ and CH$_2$O are 140 and 210 pptv, respectively. This represents the detection limit and experimental precision for these measurements. Increasing the averaging time to 10 min would improve the detection limits for NO$_2$ and CH$_2$O to 30 and 50 pptv, respectively, assuming that the averaging follows the square root of the number of points. Experience from prior ground-based field measurements of glyoxal and nitrous acid showed that their measurement precision scaled linearly with the square root of averaging time over 10–60 min periods, when zeros were acquired at more frequent intervals.

Figure 4. Correlation plot comparing NO$_2$ measurements from the BBCEAS and CRDS instruments during 1 min standard additions. The slope is 1.024 ± 0.027 with intercept of $-0.20 \pm 0.32$ ppbv (black line) and $r^2 = 0.998$.

Figure 5. Measured $\alpha(\lambda)$ and calculated spectral fits for CH$_2$O concentrations from 17 to 65 ppbv. Two fit residuals are shown in the upper panel. The average fit uncertainty for all concentrations shown is 400 pptv CH$_2$O.
Figure 6. Correlation plot comparing CH₂O measurements from the BBCEAS with the dilution scaling during 1 min standard additions. The absolute CH₂O concentration is not well known, so the slope is not calculated. The \( r^2 \) value is 0.9998 and intercept is \(-0.6 \pm 0.2 \) ppbv.

Figure 7. (a) Time series of 30 s measurements of sequential additions of 9.7 ppbv NO₂, 15.6 CH₂O, 5.5 ppbv NO₂, and 7.5 CH₂O during a 17 min period. BBCEAS NO₂ and CH₂O concentrations are compared to CRDS NO₂ concentrations and calculated CH₂O dilution factors. (b) Measured \( \alpha(\lambda) \) and calculated spectral fits for 30 s measurements with different CH₂O and NO₂ concentrations.

The absolute accuracy depends on the accuracy of the Rayleigh scattering cross section for zero air (±4 %), Rayleigh scattering cross section for helium (negligible contribution), absorption cross section for NO₂ (±3 %; Vandaele et al., 1998), absorption cross section for CH₂O (±5 %; Meller and Moortgat, 2000), sample pressure (±0.5 %), and temperature (±0.7 %). Summed in quadrature, the to-

Figure 6. Correlation plot comparing CH₂O measurements from the BBCEAS with the dilution scaling during 1 min standard additions. The absolute CH₂O concentration is not well known, so the slope is not calculated. The \( r^2 \) value is 0.9998 and intercept is \(-0.6 \pm 0.2 \) ppbv.

Figure 7. (a) Time series of 30 s measurements of sequential additions of 9.7 ppbv NO₂, 15.6 CH₂O, 5.5 ppbv NO₂, and 7.5 CH₂O during a 17 min period. BBCEAS NO₂ and CH₂O concentrations are compared to CRDS NO₂ concentrations and calculated CH₂O dilution factors. (b) Measured \( \alpha(\lambda) \) and calculated spectral fits for 30 s measurements with different CH₂O and NO₂ concentrations.

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tal calculated uncertainty is 5.1% for NO$_2$ and 6.5% for CH$_2$O. This expected measurement accuracy is consistent with the comparison of the NO$_2$ retrievals from the ultraviolet BBCEAS instrument to the 405 nm CRDS instrument.

5 Summary and conclusions

With a 2$\sigma$ detection limit of 300 pptv CH$_2$O in 1 min, the current demonstration of this measurement technique is appropriate for laboratory studies of CH$_2$O, ground-based field measurements in regions with high CH$_2$O mixing ratios (for example, regions with large emissions and rapid oxidation of anthropogenic or biogenic VOCs), and ground-based field measurements of cleaner environments with increased averaging time (>5 min).

Although the current precision is not competitive with the best spectroscopic in situ CH$_2$O instruments, the ultraviolet BBCEAS is comparable to current ground- and satellite-based DOAS instruments, which use signal averaging to achieve lower detection limits. State-of-the-art DOAS instruments report CH$_2$O detection limits of approximately 150 pptv for data acquisition times of 5–10 min (Warneke et al., 2011), while CH$_2$O satellite data are typically analyzed as monthly or seasonal averages (Chance et al., 2000; Wittrock et al., 2006). With signal averaging, the ultraviolet BBCEAS could be competitive with these direct absorption instruments, while providing a true in situ measurement. Previous work has shown that it is possible to construct inlets for formaldehyde that have minimal sampling artifacts (Wert et al., 2002). The BBCEAS method may also be useful as a lower-time-resolution validation for wet chemical, mass spectrometry, and fluorescence methods, because it is essentially free from the potential interferences or artifacts of those methods.

Ultraviolet BBCEAS measurements would be appropriate for more rapid sampling of ambient, background CH$_2$O concentrations if its 2$\sigma$ detection limit could be improved by approximately a factor of 3 to achieve 100 pptv in 1 min. This limit would require an increase of the effective path length or an increase of the optical power at the detector. The first possibility would require a factor of 9 decrease in absorption and scattering losses in the mirror coating, with unchanged transmission and reflectivity. Figure 9 demonstrates the relationship between $\alpha_{min}$, mirror transmission, and the sum of mirror absorption and scattering, calculated from Eq. (3). The second possibility would require a factor of 9 increase in optical power at the detector, for a system following the shot noise limit. This would need to be achieved through improvements in the light source power, geometric light collection, and optical filtering. More powerful laser-driven light sources or ultraviolet LEDs may be available in the future to partially satisfy this requirement. Super-continuum laser light sources are available in the visible spectral region and
offer much higher optical power output and spatial coher-
eance, allowing them to be more efficiently coupled into an
optical cavity. Such sources are not available in the ultravio-
let spectral region but may be in the future.

The ultraviolet light source and measurements presented
here have strong potential for ambient, high time resolution
measurements of aerosol optical extinction in the ultravi-
oret spectral region, where brown carbon absorption is im-
portant. Previous measurements have reported aerosol extinc-
tion and retrievals of complex refractive indices from 360 to
420 nm (Washenfelder et al., 2013; Flores et al., 2014a, b).
The ultraviolet BBCEAS instrument will allow those mea-
surements to be extended to 315 nm. The 1σ precision of
1.8 × 10⁻⁸ and 6 × 10⁻⁹ cm⁻¹ (1.8 and 0.6 Mm⁻¹) per min
for single pixels (0.05 nm) at 315 and 330 nm determined
from the Allan deviation is appropriate for aerosol extinc-
tion measurements even in clean environments. These val-
ues would be further improved by averaging multiple pixels.
Field measurements of dry aerosol extinction and angstrom
exponent at 360–420 nm measured in the rural southeast-
ern USA (Washenfelder et al., 2015) indicate that the ex-

tinction at 315 nm would be $1 \times 10^{-7} - 2 \times 10^{-6}$ cm⁻¹ (10–
200 Mm⁻¹). These values are easily measurable with the cur-
rent detection limit, with signal-to-noise ratio of 6–110 (17–
300) at 315 nm (330 nm) for 1 min ground measurements and
6–110 (12–250) for 1s aircraft measurements.

Acknowledgements. We thank Max McGillen for preparation of
formaldehyde samples. We thank Andrew Langford for technical
discussions. We thank Ryan Thalman and three anonymous
reviewers for their discussion comments. R. A. Washenfelder,
A. R. Attwood, and S. S. Brown acknowledge financial support
from the NOAA Atmospheric Chemistry and Climate Program.
This research was supported by research grants from the USA-
Israel Binational Science Foundation (BSF) grant no. 2012013.
Y. Rudich acknowledges support from the Grand Center at the
Weizmann Institute and the Dollond Charitable Trust.

Edited by: D. Heard

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