Temporal variations in rainwater methanol

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Abstract. This work reports the first comprehensive analysis of methanol concentrations in rainwater. Methanol concentrations measured in 49 rain events collected between 28 August 2007 and 10 July 2008 in Wilmington, NC, USA, ranged from below the detection limit of 6 nM to 9.3 µM with a volume-weighted average concentration of 1 ± 0.2 µM. Methanol concentrations in rainwater were up to ∼200 times greater than concentrations reported previously in marine waters, indicating wet deposition as a potentially significant source of methanol to marine waters. Assuming that these methanol concentrations are an appropriate proxy for global methanol rainwater concentrations, the global methanol wet deposition sink is estimated as 20 Tg yr⁻¹, which implies that previous methanol budgets underestimate removal by precipitation. Methanol concentrations in rainwater did not correlate significantly with H⁺, NO₃⁻, and NSS, which suggests that the dominant source of the alcohol to rainwater is not anthropogenic. However, methanol concentrations were strongly correlated with acetaldehyde, which has a primarily biogenic input. The methanol volume-weighted concentration during the summer (2.7 ± 0.9 µM) was ~3 times that of the winter (0.9 ± 0.2 µM), further promoting biogenic emissions as the primary cause of temporal variations of methanol concentrations. Methanol concentrations peaked in rainwater collected during the time period 12 p.m.–6 p.m. Peaking during this period of optimal sunlight implies a possible relationship with photochemical methanol production, but there are also increases in biogenic activity during this time period. Rain events with terrestrial origin had greater concentrations than those of marine origin, demonstrating the significance of the continental source of methanol in rainwater.

1 Introduction

Methanol is the second most abundant volatile organic compound (VOC) in the troposphere, where it plays a significant role in atmospheric chemistry by producing O₃, CO, and HCHO and decreasing OH concentrations (Millet et al., 2008). Tie et al. (2003), using a global chemical transport model, reported that methanol emissions produce approximately a 1–2 % increase in O₃, a 1–3 % decrease in OH, and a 3–9 % increase in formaldehyde. The GEOS-chem 3-D model of atmospheric chemistry indicates that methanol emissions account for 20 % of CO and HCHO production rates during spring and early summer (Hu et al., 2011). Methanol reacting with OH can also lead to production of formic acid (Monod et al., 2000), which in turn increases acidification of rain. These various atmospheric interactions and subsequent impacts on atmospheric chemistry have led to the investigation and quantification of sources of methanol emissions.

Methanol sources are primarily biogenic (e.g., plant growth and plant decay), but there are a wide range of less significant anthropogenic sources including biofuel burning, gasoline additives, vehicle exhaust, solvent use and many industrial processes (as summarized by Wells et al. 2012). Whereas it is generally agreed that, globally, biogenic sources account for the majority of methanol emissions, the actual percentage attributed to biogenic emissions is still under investigation (Millet et al., 2008). Global budgets reviewed by Jacob et al. (2005) report a biogenic source range of 63 to 91 % but in a source tracer study at a site in the upper Midwest US, up to 70 % of wintertime methanol was attributed to anthropogenic sources (Hu et al., 2011). The anthropogenic contribution may be more substantial in urban areas that have larger methanol concentrations (Heikes et al.,...
2002) and are less prone to biogenic emissions than rural areas. Due to the atmospheric lifetime of methanol (5 to 12 days) (Jacob et al., 2005), emissions from anthropogenic or biogenic sources can possibly travel across large continental regions, making it difficult to apportion various methanol source contributions to ambient methanol concentrations.

Efforts have been made to refine the estimated quantities of sources and sinks (primarily OH oxidation) of atmospheric methanol (Heikes et al., 2002; Millet et al., 2008; Wells et al., 2012). These efforts have been predominately based on methanol concentration data consisting of aircraft and surface air analysis. The limited number of measurements and the lack of concentration data in the condensed phase lead to wide discrepancies in global methanol budget models, which in turn have led to the reported total global methanol source ranging from 75 to 490 Tg yr\(^{-1}\) and sink from 40 to 570 Tg yr\(^{-1}\) (Singh et al., 2000a; Galbally and Kirstine, 2002; Heikes et al., 2002; von Kuhlmann et al., 2003; Tie et al., 2003; Jacob et al., 2005).

There is currently a paucity of condensed phase methanol concentration data and a lack of information of methanol’s role in the global biogeochemical cycling. In fact, there are currently no detailed studies of methanol levels in precipitation. This uncertainty has resulted in a wide range in the predicted wet deposition sink of methanol from 4 to 50 Tg yr\(^{-1}\) (Singh et al., 2000a; Galbally and Kirstine, 2002; Heikes et al., 2002; von Kuhlmann et al., 2003; Tie et al., 2003; Jacob et al., 2005).

The purpose of the current study was to define the ranges and patterns of variation in the abundance of rainwater methanol, including such factors as the influence of air mass back trajectory on concentrations. To gain better insight into the possible anthropogenic and biogenic origins of methanol, concentrations were also intercorrelated with a variety of other rainwater components as well as seasons. Data generated in this study are requisite to the generation of the first total global wet deposition sink of methanol based on measured aqueous-phase concentrations.

2 Materials and methods

2.1 Sample collection

Wilmington rainwater samples were collected on an event basis on the campus of the University of North Carolina at Wilmington (UNCW) from 28 August 2007 to 10 July 2008 (\(n = 49\) events). The collection site at UNCW is a large open area of approximately 1 hectare and is made up of a turkey oak, long leaf pine and wire grass community. This area is typical of the inland coastal area of southeastern North Carolina. The site (34°13.9′N, 77°52.7′W) is approximately 8.5 km from the Atlantic Ocean. Due to the close proximity of the collection site to the laboratory, methanol analysis or filtration and refrigeration of samples could be done within minutes of collection. This reduces the possibility of compositional changes between the times of collection and analysis. If it was not possible to analyze the rain samples within 2 h, they were frozen immediately and stored in a \(-80 \, ^\circ\text{C}\) freezer. Event rain samples were collected using Aerocoem-Metrics (ACM) model 301 automatic sensing wet/dry precipitation collectors containing 4 L Pyrex glass beakers that were precleaned by combusting at 450 °C for 4 h to remove organic impurities. Rainwater concentrations are reported as volume-weighted concentrations with volume-weighted standard deviations. This is the mathematical equivalent to mixing all rain within a specified time period and reporting the analytical result for that composite sample.

2.2 Formaldehyde and methanol

Formaldehyde concentrations in rainwater samples were determined by derivatization with 2,4-dinitrophenylhydrazine followed by separation and detection by HPLC (Kieber et al., 1999). Samples and standards reacted with 2,4-dinitrophenylhydrazine (DNPH) for one hour in the dark, forming a hydrazone, which was separated from interfering substances by HPLC and quantified by UV detection at 370 nm. Derivatized samples (100 \(\mu\)L) were injected onto a reversed phase Luna 100 mm × 4.60 mm 3 \(\mu\)C18 Phenomenex column with a 100 Å pore size at 10°C. The mobile phase was a 1:1 mixture of 0.1% trifluoroacetic acid (TFA) in acetonitrile and 0.1% TFA in deionized water at a flow rate of 1.00 mL min\(^{-1}\).

Methanol was determined from a second aliquot by oxidation of the alcohol to formaldehyde via alcohol oxidase obtained from the yeast Hansenula sp. (Kieber et al., 2013). The enzyme was prepared by dissolution of 100 units of alcohol oxidase in 5 mL of 0.1M potassium phosphate buffer (pH 9.0). The sample (1000 \(\mu\)L) was combined with 10 \(\mu\)L of buffer along with 100 \(\mu\)L of an enzyme working reagent (0.18 units mL\(^{-1}\)) and allowed to react at 40 °C for 40 min before addition of 10 \(\mu\)L of DNPH. The concentration of methanol was determined after HPLC analysis by the difference in formaldehyde concentration in samples with and without added enzyme. Milli-Q water is analyzed for methanol during each analysis and the methanol blank concentration is subtracted from the sample. This method has a detection limit of 6 nM and a precision of 2% relative standard deviation.

2.2.1 Reagents and standards for methanol analysis

Alcohol oxidase (100 units) from the yeast Hansenula sp. was purchased from Sigma (St. Louis, MO). Water was purified using a Millipore Q-water system (Millipore Corp., Bedford, MA) and used to prepare all solutions. Reagent grade 2,4-dinitrophenylhydrazine (DNPH) was purchased from Sigma (St. Louis, MO), triply recrystallized from acetonitrile and kept refrigerated in the dark. Acetonitrile
(HPLC grade, Burdick and Jackson, Muskegon, MI), 12 M hydrochloric acid (Reagent Grade, VWR International, West Chester, PA) and carbon tetrachloride (HPLC grade 99.9 %, Sigma, St. Louis, MO) were used in the preparation and purification of the DNPH reagent solution.

Formaldehyde (37.69 % CH₂O) and paraformaldehyde (94.19 %, containing no methanol) were obtained from the Wright Chemical Company (Wilmington, NC). A 1 M formaldehyde stock solution was prepared before each rain event. Methanol (HPLC grade, Burdick and Jackson, Muskegon, MI) and Milli-Q water were used to prepare a 1 M alcohol stock solution before each rain event. ACS grade (99.0 %) potassium dihydrogen phosphate and reagent grade potassium hydrogen phosphate (Alfa Aesar, Ward Hill, MA) were used in the preparation of all buffer solutions.

2.3 Supporting analyses

These supporting data were used to characterize rain events and to evaluate whether the patterns of variation observed for methanol co-vary with any of these analytes. These supporting data also allow comparison with rain collected elsewhere.

Hydrogen peroxide was analyzed at the time of sample collection by a fluorescence decay technique involving the peroxidase-mediated oxidation of the fluorophore scopoletin by H₂O₂ in rain buffered at a pH of 7 with a phosphate buffer (Mullaugh et al., 2012). The dissolved organic carbon content in the rainwater samples was determined with a Shimadzu TOC 5000 carbon analyzer (Shimadzu, Kyoto, Japan) equipped with an ASI 5000 autosampler (Willey et al., 2000). Inorganic anions (Cl⁻, NO₃⁻, and SO₄²⁻) were analyzed using suppressed ion chromatography. A Ross electrode with low ionic strength buffers was used for pH analysis. Organic acid concentrations were measured with a Dionex 4000i/SP ion chromatograph with a SP4290 integrator. Dionex Ion-Pac AS11 4 mm analytical column, AG11 4 mm Guard column and anion micromembrane suppressor model AMMS-11 (Avery et al., 2001). Acetalddehyde concentrations in rainwater samples were determined by derivitization with 2,4-dinitrophenylhydrazine followed by separation and detection by HPLC (Kieber et al., 1999).

2.4 Rain event origin definitions

Precipitation events were categorized using air-mass back trajectories generated using version 4 of the Hybrid Single Particle Lagrangian Integrated Trajectory Model (HYSPLIT) developed at the National Oceanic Atmospheric Administration – Air Resources Laboratory (NOAA/ARL, 2013). Trajectories were generated using a stand-alone PC version of the model and calculated using pre-processed gridded horizontal and vertical wind fields generated at 6-hour intervals from the National Center for Environmental Prediction’s Global Data Assimilation System using the Medium Range Forecast model to produce the forecast wind fields. Single back trajectories were run for each measured precipitation event collected at UNCW, starting at the recorded onset of precipitation. Trajectories were run starting at the 500m level to represent the air mass near the well-mixed boundary layer likely to contribute more heavily to in-cloud processes contributing to wet deposition (Walker et al., 2000). They were then visually categorized based on origin (compass direction) and path (terrestrial, marine, coastal, or mixed). Terrestrial and marine air masses are those whose pathway for the 120h period preceding the rain event was predominantly over a landmass or the ocean, respectively. Mixed trajectories were those that were determined to have the same potential for oceanic as well as terrestrial influence, based on a visual analysis of their pathway (Kieber et al., 2005). Coastal trajectories followed the Atlantic coastline.

GIS shape files produced by the HYSPLIT program were used to plot trajectories for each rain event type on a US county (county is defined as a geological subdivision of a US state) basemap that portrays size-normalized county-level biogenic methanol emissions. The methanol emissions used to make the map were from the 2008 National Emissions Inventory (NEI, 2008).

2.5 Results and discussion

Methanol concentrations in the samples for the 49 rain events ranged from below the detection limit (< 6 nM) to 9.3 µM with a volume-weighted average concentration of 1 ± 0.2 µM, a simple average concentration of 1 ± 2 µM and a median of 0.9 µM. The range in reported gas-phase methanol concentrations (0.03 to 47 ppbv (Heikes et al., 2002; Jacob et al., 2005)) and a range of Henry’s Law constants (140 to 230 mol kg⁻¹ bar⁻¹ (NIST 2014)) were used to investigate if the rainwater concentrations are in thermodynamic equilibrium with gas-phase methanol. If the rainwater is in equilibrium with the gas phase, the calculated range of rainwater methanol concentrations is 4.2 nM to 10.8 µM. This range is similar to the observed range of < 6 nM to 9.3 µM, and suggests equilibrium, but disequilibrium cannot be ruled out. The average methanol concentration in this study is almost 2 times the average in rainwater (690 nM) collected at an Arizona site in 1982 (Snider and Dawson, 1985). These are the only other known methanol concentrations in rainwater, and may be much lower than the Wilmington average, because Arizona is a much more arid region that probably has lower biogenic emissions. Direct comparison to this earlier study should be viewed with caution however, as the reported blank (625 nM) was very near the reported average concentration, and it lacked sufficient analytical sampling frequency (n = 4) to allow for detailed analysis of temporal variation or air-mass back trajectory influences on methanol concentrations.
2.6 Rain event origin

Rain events were classified based on their back trajectory to determine how methanol concentrations were affected by continental influences (Fig. 1). Terrestrial, mixed, coastal and marine rain events had volume-weighted average methanol concentrations of 1.5 ± 0.5 µM, 1.6 ± 0.2 µM, 0.7 ± 0.4 µM, and 1.1 ± 0.2 µM, respectively (Fig. 1). Figure 2 shows individual rain event trajectories plotted over US county-level biogenic methanol emissions. Rain events with terrestrial back trajectories had greater methanol concentrations than those with marine back trajectories, which is consistent with previous findings that gas-phase atmospheric methanol concentrations reported over land (0.03–47 ppbv) are greater than those reported over the ocean (0.3–1.4 ppbv) (Heikes et al., 2002; Jacob et al., 2005). This also agrees with global budgets that suggest methanol has a primarily terrestrial biogenic source.

Although rain event types with marine influence (coastal, marine) have smaller methanol concentrations than those with terrestrial influence, the difference is not significant (ANOVA: p = 0.15), and methanol is present in substantial amounts in samples from rain events with coastal and marine origin. This may be due to several reasons: (1) the ocean may be a source of methanol emissions. Millet et al. (2008) report the ocean as an overall methanol sink, but consider the ocean biosphere a large enough source to cause detectable concentrations in the atmosphere; (2) methanol has an average atmospheric lifetime of 5 to 12 days (Jacob et al., 2005), allowing for transport of methanol from land masses to the ocean; and (3) methanol at the rain collection site could include some methanol scavenged from local terrestrial sources. It should be noted that the previous gas-phase methanol concentration studies that show much greater methanol concentrations over land than over the ocean sampled air in the remote ocean (Yang et al., 2013). The aqueous-phase methanol concentrations associated with marine rain events were collected on coastal land, not over the open ocean, so it is expected that the large concentration differences seen between gas-phase methanol concentrations collected at ocean and terrestrial sites will not be mimicked in this aqueous-phase study.

2.7 Diurnal variations

The methanol concentrations measured in this study were divided into four time periods in order to examine if the concentrations underwent short-term temporal variations (Fig. 3) similar to what has been observed for other analytes in rainwater at this location (Kieber, 2004; Kieber et al., 2001a, 2001b). Events were excluded from classification if they occurred during more than one time period. Each bar in Fig. 3
represents the average volume-weighted methanol concentration during the given time period. Methanol concentrations peaked during time period III (12 p.m.–6 p.m.), with an average volume-weighted concentration of 2.6 ± 0.7 µM. Peaking during this period of optimal sunlight implies a possible relationship with photochemical methanol production (e.g., the oxidation of methane, and the methylperoxy radical reacting with itself and higher organic peroxy radicals) (Jacob et al., 2005), but there are also increases in plant activity and anthropogenic activity (e.g., vehicle use, industrial processes) during this time period. The concentration peak during this time period corresponds to the peak of numerous previous diurnal measurements of methanol flux over varying vegetation, and is attributed to light causing stomata to open, in turn releasing methanol (Bamberger et al., 2010; Brunner et al., 2007; Custer and Schade, 2007; Harley et al., 2007; Karl et al., 2003; Schade and Goldstein, 2002). Daytime light usually produces a temperature increase, which is also reported to increase biogenic methanol emissions exponentially (Folkers et al., 2008; Hu et al., 2011).

The approximate six-fold decrease in methanol concentration between day and night in this study falls within the reported range (0.3 to 7 times) of decrease (Jacob et al., 2005), which could indicate dry deposition at night and adsorption to vegetation and soils. Other than a likely decrease in methanol emissions from vegetation, a possible mode for the nighttime decrease in the aqueous-phase concentration of methanol is the advection of “cleaner” marine air to the rain collection site. This was suggested as an explanation for smaller nighttime concentrations of other rainwater components (e.g., H$_2$O$_2$, formaldehyde, formic acid, NO$_3^-$) at this collection site (Avery et al., 2001).

2.8 Seasonal variations

The methanol concentration data collected in this study were divided into four seasonal time periods in order to examine seasonal variations in the analyte concentrations (Fig. 4). The seasons were defined as winter (1 December–29 February), spring (1 March–31 May), summer (1 June–31 August), and fall (1 September–31 November). The average volume-weighted concentrations of the methanol observed in this study during the spring and summer months were 1.0 ± 0.3 µM and 2.7 ± 0.9 µM, respectively, and were not significantly different from each other (ANOVA: $p = 0.66$). Summer concentrations were however significantly different than fall and winter (ANOVA: $p = 0.01$). Methanol concentrations were expected to increase in the spring as plant growth increases. A slight increase was seen, but methanol emissions from vegetation may have been hampered by an unusually dry spring. Methanol concentrations in rainwater were greatest in the summer, which was to be expected since the plant growing season occurs during summer, and increasing summer temperatures can lead to exponentially increasing methanol emissions from plants (Folkers et al., 2008; Hu et al., 2011). The early summer peak in the methanol concentrations agrees with simulated methanol seasonal cycles, and more specifically the monthly rainwater methanol concentration peak in June (3.6 ± 0.9 µM) is consistent with the peak seen in several seasonal cycle simulations (Wells et al., 2012).

Since methanol has a large biogenic source, gas-phase methanol concentrations are expected to have smaller concentrations in the winter and fall months, when some plants are dormant. The volume-weighted average concentration of methanol during the winter and fall rain events was 0.9 ± 0.2 µM and 0.7 ± 0.2 µM, respectively. Various previous studies have reported winter gas-phase methanol concentrations to be much smaller than those in summer, and these studies suggest a greater percent contribution from anthropogenic methanol sources during the winter (Millet et al., 2005; Legreid et al., 2007; Jordan et al., 2009; Hu et al., 2011). For instance, using toluene, benzene, and CO as anthropogenic tracers at a rural Midwestern US site, Hu et al. (2011) estimated that up to 70% of wintertime methanol emissions had an anthropogenic origin. Rainwater methanol concentrations reported in this current study are consistent with these previous gas-phase methanol results in that winter rainwater methanol concentrations are 1/3 of the summer concentrations. It might be expected that fall concentrations would increase due to plant decay, but this source of methanol is relatively small when compared to reported contributions from the plant growth source (7 to 27% of plant growth source total) (Galbally and Kirstine, 2002; Heikes et al., 2002; Jacob et al., 2005; Singh et al., 2000b). A possible explanation for smaller than expected concentrations of methanol during the fall rain events may be that the drought conditions of the previous summer were severe to extreme
occur for methanol. The increase in methanol concentrations with rainfall amount (Table 1, Fig. 5), suggesting it is a ubiquitous component of precipitation. Rain events with terrestrial origins had greater concentrations than those of marine origin, indicating the potential for a significant continental source of methanol in rainwater. Correlation analysis with other rainwater analytes and greater growing season concentrations suggest this continental source is primarily biogenic rather than anthropogenic at this location. The lack of washout behavior implies a mode of methanol resupply during rain events possibly involving photochemical processes or transpiration from plants. Increases in methanol during daytime and summer rain events further underscore the importance of biogenic activity and photochemical production in the biogeochemical cycling of methanol in precipitation.

Methanol concentrations represent 1.5 ± 0.3% (range: 0.08 to 14.4%) of the DOC content, indicating that it is not a significant contributor to the overall carbon budget in precipitation. The lack of correlation with the dissolved organic carbon content of samples suggests that the fraction of DOC that is methanol is also variable.

Methanol levels in rainwater measured in this study were up to ~200 times greater than concentrations observed in marine waters (48 to 296 nM) (Dixon et al., 2011), indicating wet deposition of methanol as a potentially significant source to marine waters. The important role by rainwater deposition of atmospherically enriched analytes such as methanol in oceanic concentrations was demonstrated in an earlier study of hydrogen peroxide on surface seawater at the Bermuda Atlantic Time Series Station (Kieber et al., 2001a). The large increase in surface water concentrations after precipitation events indicated that rainwater deposition was a significant, and in some cases the dominant source of hydrogen peroxide to open ocean seawater (Kieber et al., 2001a). It is also likely that the rainwater flux of methanol to freshwater systems is episodically significant, similar to what has been observed for formaldehyde at this location (Kieber et al., 1999). During the high irradiance summer months, formaldehyde contributes as much as 30 times the resident amount present in freshwater lakes of southeastern North Carolina (Kieber et al., 1999).

One of the important issues in the global biogeochemical cycling of methanol is the magnitude and uncertainty of its wet depositional sink. Assuming the annual volume-weighted average reported at the Wilmington, NC, USA sampling site is an appropriate proxy for a global average rainwater concentration of methanol, and the global annual precipitation volume is ~5.36 × 10^11 L (Pidwirny, 2008), the global methanol wet deposition sink is estimated as 20 ± 3.3 Tg yr⁻¹. This estimate falls within the theoretical range (4–50 Tg yr⁻¹) produced by many global methanol budget models (Galbally and Kirstine, 2002; Heikes et al.,

Figure 5. Methanol concentration vs. rainfall amount for all rain events.

(NC Drought 2008), killing vegetation that would otherwise have decayed during the fall.

2.9 Intercorrelation

Formaldehyde, acetaldehyde, rain amount, H⁺, NO₃⁻, H₂O₂, NSS (non-sea-salt sulfate), formate, and acetate concentrations were determined in addition to methanol in this study. All rainwater components were analyzed by intercorrelation (Table 1). Methanol concentrations did not correlate with H⁺, NO₃⁻, and NSS, which suggests that the dominant source of the methanol to rainwater is not anthropogenic. There was no correlation between methanol and the dissolved organic carbon (DOC) content of rainwater samples, indicating that the methanol makes up a variable fraction of the organic carbon pool. Methanol concentrations were strongly correlated with acetaldehyde, which has a primarily biogenic input (Millet et al., 2010), suggesting that the potential biogenic source of methanol is consistent with the larger concentration of methanol observed during the growing season (Fig. 5).

There was no significant correlation of methanol concentrations with rainfall amount (Table 1, Fig. 5), suggesting it is not washed out of the atmosphere at this location, but rather is resupplied during a rain event. A possible mode of resupply is transpiration from plants. Rainwater initiates the transpiration stream in plants; methanol, being highly soluble, is transported in the transpiration stream and is emitted via the stomata (Fall and Benson, 1996; Niinemets et al., 2003). This methanol emission via transpiration may continue throughout rainfall events and lead to a local resupply that in turn is scavenged by continuing wet deposition. Formaldehyde also does not exhibit washout at this location (Kieber et al., 1999), which the authors attribute to in situ photochemical production from chromophoric dissolved organic matter in rainwater (Southwell et al., 2010). The increase in methanol concentrations during periods of peak sunlight intensity (Fig. 3) indicates a similar photochemical production mechanism may occur for methanol.

3 Implications

This work reports the first detailed analysis of methanol concentrations in rainwater. The presence of methanol in all rainwater samples analyzed suggests that it is a ubiquitous component of precipitation. Rain events with terrestrial origins had greater concentrations than those of marine origin, indicating the potential for a significant continental source of methanol in rainwater. Correlation analysis with other rainwater analytes and greater growing season concentrations suggest this continental source is primarily biogenic rather than anthropogenic at this location. The lack of washout behavior implies a mode of methanol resupply during rain events possibly involving photochemical processes or transpiration from plants. Increases in methanol during daytime and summer rain events further underscore the importance of biogenic activity and photochemical production in the biogeochemical cycling of methanol in precipitation.

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Table 1. Intercorrelations among methanol, formaldehyde, acetaldehyde, and various other rainwater components. Bold-faced values indicate significance at $p < 0.001$. Asterisk (*) indicates significance at $p < 0.05$. Number of samples equals 47 for acetaldehyde, nitrate, and sulfate, 27 for formate and acetate, and 49 for others.

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<th>CH$_2$O</th>
<th>CH$_3$CHO</th>
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2002; von Kuhlmann et al., 2003; Jacob et al., 2005) and is double the reported representative best estimates (9–12 Tg yr$^{-1}$) (Galbally and Kirstine, 2002; Heikes et al., 2002; von Kuhlmann et al., 2003; Tie et al., 2003, Jacob et al., 2005). Results of this study are significant because they suggest that previous methanol budgets may be underestimating removal by precipitation. Additional studies of methanol concentrations in rainwater should be carried out in various regions of the Earth to better constrain the global wet depositional sink of this biologically and chemically labile analyte.

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