# Investigation of IR and Raman spectra of species present in formaldehyde-water-methanol systems

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## ABSTRACT

Formaldehyde forms a variety of hydrated and methoxylated species when reacted with water and methanol. Vibrational spectroscopy has been deployed for both remote and in situ sensing of formaldehyde species and it can be a useful tool for process development, monitoring and control at both laboratory and industrial scale, as well as for environmental, atmospheric and space monitoring. While IR and Raman spectroscopic studies of formaldehyde species in solid, liquid or gas phases have been reported, assignments of vibrational frequencies of relevant species in previous literature have been contradictory and incomplete. In this work we report IR and Raman spectra for formaldehyde-water-methanol solutions across a wide range of formaldehyde concentrations and solvent compositions. We present an analysis of vibrational spectra of formaldehyde-water-methanol systems using a combination of experimental measurements and gas phase quantum mechanical density functional theory simulations. For the first time, we explicitly consider spectra of oligomeric mixtures of formaldehyde species in relation to spectra of specific representative hydrated and methoxylated species and we resolve some previously reported contradictions in assignments of vibrational frequencies for formaldehyde systems.

**KEYWORDS**: Formaldehyde, Methylene glycol, methanediol, methoxylated methylene glycol, methoxymethanol, dimethyleneglycol, di(oxymethylene)glycol, FTIR, Raman, vibrational frequencies, vibrational spectra, spectroscopy

#### Introduction

Formaldehyde solutions are commonly used as reactants and precursors in synthesis of many chemical intermediates [1] and polymeric materials, including resins and nanoporous organic gels [2–5]. Formaldehyde-water-methanol solutions are frequently used in both laboratory and industrial environments where IR and Raman spectroscopies can both be deployed in situ and can be invaluable tools for process development, monitoring and control. Both qualitative and quantitative vibrational spectroscopic analysis of hydrated and methoxylated formaldehyde species is thus required. However, assignments of vibrational frequencies of relevant species have been incomplete and to some extent contradictory in previous literature, while the speciation in formaldehyde-water-methanol systems has been only recently quantitatively investigated [4]. Hydrated and methoxylated formaldehyde-based species are also of interest in many other fields, including environmental, atmospheric and space sciences [2,6–8], where monitoring of these species can only be achieved by vibrational spectroscopy-based sensing and accurate measurement of their concentrations is crucial for assessment and validation of climate models as well as understanding of interplanetary and interstellar chemistry.

Formaldehyde hydration to methylene glycol (MG, HO-CH<sub>2</sub>-OH), also called methanediol, and their subsequent polymerisation to diglycol, or dimethylene glycol (DG, HO-CH<sub>2</sub>-O-CH<sub>2</sub>-OH), triglycol (TG, HO-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-OH) and so on is favourable in aqueous solution [4]:

(a) 
$$CH_2O + H_2O \rightleftharpoons HO(CH_2O)H$$
  
(b)  $HO(CH_2O)_{i-1}H + HO(CH_2)OH \rightleftharpoons HO(CH_2O)_iH + H_2O$   $i \ge 2$ 

However, when polymerisation is undesirable (such as in highly concentrated formaldehyde solutions available commercially), methanol is added as a stabiliser. In the presence of methanol, formaldehyde forms methoxylated methylene glycol (MMG, CH<sub>3</sub>O-CH<sub>2</sub>-OH), also called methoxymethanol, and related oligomers, such as methoxy diglycol (MDG, CH<sub>3</sub>O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O-CH<sub>2</sub>-O

(c) 
$$HO(CH_2O)H + CH_3OH \rightleftharpoons HO(CH_2O)CH_3 + H_2O$$
  
(d)  $HO(CH_2O)_{i-1}H + CH_3OH \rightleftharpoons HO(CH_2O)_{i-1}CH_3 + H_2O$   $i \ge 2$ 

Since methoxylation of methylene glycol and its oligomers is relatively favourable compared to polymerisation, the degree of polymerisation is much lower in the presence of methanol. All these reactions are acid and base catalysed but they proceed relatively rapidly in aqueous methanolic solutions even without adding any catalyst [4]. Further methoxylation of MMG to  $CH_3O(CH_2O)CH_3$  is favoured thermodynamically but does not proceed without a sufficient amount of catalyst [9].

Several studies were carried out to investigate the speciation in aqueous formaldehyde solutions in the absence of methanol obtained by depolymerisation of pure formaldehyde polymer (paraformaldehyde or poly(oxymethylene)) using NMR [10,11], while formaldehyde solutions in pure methanol were investigated as well [12]. Despite the widespread use of commercial formaldehyde stock solutions, the quantitative speciation in aqueous formaldehyde solutions containing methanol has only recently been investigated in detail using NMR spectroscopy [4].

Raman spectra of formaldehyde solutions were first reported by Matsuura et al. [13]. Although commercial formaldehyde stock solutions stabilised by methanol were used in this study, Matsuura et al. assumed that these solutions contain only methylene glycol, water and methanol when making assignments of vibrational frequencies, as it was not known at that time that these solutions contain a large fraction of oligomers and methoxylated oligomers in addition to methylene glycol. Further studies on aqueous formaldehyde solutions in the absence of methanol [14,15] suggested assignments for observed Raman spectra over a wide range of concentrations where numerous oligoglycol species were present in solutions. Raman spectroscopic studies of frozen aqueous formaldehyde solutions were also reported [8] as well as studies of phenol-formaldehyde solutions [16]. However, formaldehyde solutions containing methoxylated species have not been investigated previously using Raman spectroscopy.

IR spectra of MG and MMG in gas phase [6,7,17] as well as MMG in methanol [18] were reported as well as liquid phase IR spectra of formaldehyde aqueous solutions over a relatively narrow spectral range [19]. Studies of phenol-formaldehyde resins using IR spectroscopy were also reported [20]. More recently, vibrational frequencies of MG, MMG and DG were computed using quantum mechanical methods [7,21–23]. Similarly to Raman spectroscopy, formaldehyde-water-methanol solutions containing methoxylated species have not been investigated previously using IR spectroscopy.

In this work, we report IR and Raman spectra for commercial formaldehyde stock solutions diluted with either water or methanol across a wide range of formaldehyde concentrations. We present a detailed characterisation of vibrational frequencies observed in formaldehyde-watermethanol systems using a combination of experimental measurements and quantum mechanical density functional theory simulations, while for the first time considering vibrational spectra of complex oligomeric mixtures of formaldehyde species together with those of specific representative species, including MG, MMG and DG.

## Methods

## Sample preparation

Table 1 shows the compositions of mixtures investigated in this research. Required amount of formaldehyde stock solution (37%wt.; Sigma-Aldrich UK) stabilised by 13%wt. of methanol [24] was measured into a glass beaker equipped with a magnetic bar (14x5 mm, PTFE surface) and placed on a magnetic stirrer. After required amount of diluent was added, either de-ionised water from in-house supply (Millipore Elix 5) or methanol ( $\geq$ 99.8%wt., ACS grade, VWR International), the beaker was covered with parafilm to prevent evaporation and the contents were stirred for 5 min. The measurements were performed directly afterwards in the same vessel.

Attenuated Total Reflection (ATR) Fourier Transform Infra-Red (FTIR) Spectroscopy

MB3000 mid-IR Fourier Transform spectrometer manufactured by ABB Inc., equipped with class 3B laser (760 nm, 2 mW output power) and an ATR probe manufactured by Fibre Photonics linked with the FTIR by an optical polycrystalline silver halide fibre cable, were used. Due to the fact that an ATR diamond probe was used in all IR experiments, the range of wavenumbers accessible was limited by a diamond resonance signal at ca. 2000 cm<sup>-1</sup>, and limited

due to poor throughput of light by the optical fibre cable above 2000 cm<sup>-1</sup>, therefore the spectra between 800 cm<sup>-1</sup> and 1700 cm<sup>-1</sup> were collected and analysed. The ATR probe was immersed in the solution, while gently agitating the contents to minimize the effect of local concentration differences. All experiments were carried out at room temperature with the same set of parameters: 16 scans, resolution 8 cm<sup>-1</sup>, using de-ionised water as a background. Background subtraction, baseline correction and peak deconvolution for measured IR spectra was done using Origin software.

#### Raman Spectroscopy

All spectra were collected using a Leica DM/LM microscope equipped with an Olympus 20x/0.4 long working distance objective, which was used to collect  $180^{\circ}$  backscattered light from macrosampler. The spectrometer system was a Renishaw inVia equipped with a helium-neon (He-Ne;  $\lambda$ =632.8 nm) laser coupled with Renishaw Ramascope System 2000. Background subtraction, baseline correction and peak deconvolution for measured Raman spectra was done using Origin software.

#### Simulation of Vibrational Frequencies

Simulated vibrational frequencies were calculated using the Quantum ESPRESSO package, version 5.1 [25], which implements density functional theory (DFT) using a plane wave basis set. Molecules were generated with the aid of the chemical drawing tool Avogadro (http://avogadro.cc/). Projector augmented wave pseudopotentials, C.pbe-n-kjpaw\_psl.0.1.UPF, O.pbe-n-kjpaw\_psl.0.1.UPF and H.pbe-kjpaw\_psl.0.1.UPF, were taken from the Quantum ESPRESSO pseudopotential data base: http://www.quantum-espresso.org/pseudopotentials. The PBE version [26] of the generalised gradient approximation was used for the exchange and correlation functional. Cut-offs of 50 Ry and 200 Ry were used for the wavefunctions and charge densities, respectively. The molecules were ionically relaxed in a cubic cell of side 20 Bohr, using a single k-point, an electronic convergence threshold of  $1x10^{-10}$  Ry, and a force convergence threshold of  $1x10^{-18}$  Ry was used. The vibrational mode characters were assigned by viewing individual vibrational modes in Molden [27].

## **Results and Discussion**

## IR and Raman spectra: qualitative analysis

IR and Raman spectroscopy provide information about the vibrational spectra of species present in given solution matrices. These two techniques are complementary, as certain functional groups in molecules might be invisible in IR due to the fact they have no dipole moment but may be observed in Raman spectra due to their polarizability. Figure 1 shows comparison of IR and Raman spectra for the stock formaldehyde solution (37%wt. formaldehyde and 13%wt. methanol in water), where one can see a number of vibrational frequencies in the region of wavenumbers between 800 and 1700 cm<sup>-1</sup>. A very similar Raman spectrum was reported previously by Matsuura et al., see their Figure 1 [13], although the authors assumed that all signals come from either MG or methanol. However, this is not the case as numerous methylene glycol oligomers and methoxylated oligomers are present in these highly concentrated solutions [4]. A comparable IR spectrum of a stock formaldehyde solution was reported previously by Poljansek and Krajnc, see their Figure 1 [20].



Figure 1. Normalised IR (solid line) and Raman (dashed line) spectra of the stock formaldehyde solution. Peaks A-R, assigned in Table 5 based on experimental spectra and vibrational frequency calculations, are indicated, although some are not apparent in these particular spectra.

In Figures 2 and 3 we show IR and Raman spectra measured for progressively diluted formaldehyde stock solutions, using either water or methanol as diluents.

As stock formaldehyde solutions are progressively diluted with either water or methanol, the distribution of formaldehyde related species changes. Since oligomerisation of methylene glycol is a second order reversible reaction, dilution with solvent results in reverse oligomerisation. In dilution with water, methanol present in the original stock solution is diluted as well, and since methoxylation is also second order reversible reaction, dilution with water results in reverse methoxylation.



Figure 2. IR spectra of formaldehyde stock solutions diluted with water (a) or with methanol (b). See Table 1 for corresponding solution compositions.





а

Figure 3. Raman spectra of formaldehyde stock solutions diluted with water (a) or with methanol (b). See Table 1 for corresponding solution compositions.

Therefore in the limit of large dilutions, monomeric methylene glycol is present as the major formaldehyde containing species, accompanied by minor amounts of MMG and methanol [4], in formaldehyde stock solutions diluted with water. In the case of dilution of stock formaldehyde solutions with methanol, methoxylation is favoured and in the limit of large dilutions methoxylated methylene glycol is present as the major formaldehyde containing species, accompanied by minor amounts of MDG and even less of MG [4], while spectra become dominated by methanol itself (see vibrational spectra of methanol and methanol-water mixtures in Figures S7 and S8 in Supporting Information). Consequently, formaldehyde-related contribution to spectra changes due to decreasing absolute concentration of formaldehyde containing species as well as changing speciation, while methanol-related contribution to spectra increases upon methanol addition. Significantly, peaks at 910 cm<sup>-1</sup>, 1110 cm<sup>-1</sup>, 1195 cm<sup>-1</sup> and 1300 cm<sup>-1</sup> initially increase in the IR spectra upon dilution with methanol (see Figure 2b) and then gradually decrease. However, all of these peaks gradually decrease and then vanish in the IR spectra upon dilution with water (see Figure 2a). This indicates that the corresponding vibrational frequencies are present in methoxylated species (such as MMG and MDG) which are thermodynamically favoured in the presence of methanol. On the other hand, peak at 1025 cm<sup>-1</sup> decreases in both IR and Raman spectra upon dilution with water but persists until the highest dilution, indicating that that the corresponding vibrational frequency is present in monomeric methylene glycol. At the same time, it can be seen that the same peak increases in both IR and Raman spectra upon dilution with methanol, so it is clearly also related to a vibrational frequency present in methanol.

As numerous peaks observed in IR and Raman spectra above may not be uniquely linked to single specific formaldehyde-related species, it is crucial to investigate assignments of vibrational frequencies for most relevant species present in these systems in order to interpret these spectra properly. We will aid this investigation by careful examination of simulated vibrational frequencies and their corresponding vibrational modes using density functional calculations as described in the following section.

## Simulated vibrational frequencies

To gain further insight into the experimental vibrational spectra, we simulated vibrational frequencies for methylene glycol (MG), methoxylated methylene glycol (MMG) and diglycol (DG) using DFT. Formaldehyde and methanol were also calculated for validation purposes and results for these are shown in the Supporting Information. We note that our calculated vibrational spectra correspond to gas phase molecules.

# Methylene glycol (MG)

Methylene glycol has 15 vibrational frequencies. We calculated vibrational frequencies for the lowest energy conformer, corresponding to methanediol-C2 in Barrientos et al. [21]. The vibrational frequencies, intensities and mode assignments are presented in Table 2, with all frequencies listed in ascending order. Vibrational frequencies calculated using second order

Møller-Plesset perturbation theory (MP2) [21], quadratic configuration interaction including single and double substitutions (QCISD) [21] and DFT polarisable continuum model (PCM) [22] are also shown in ascending order in Table 2. Our data is in excellent agreement with the QCISD and DFT PCM data and the correlation between our results and the vibrational frequencies in the literature over the range 800-1700 cm<sup>-1</sup> is shown in Supporting Information. We note that Barrientos et al. [21] and Delcroix et al. [22] used scaling factors of 0.9624 and 0.989 for the QCISD data and DFT PCM implicit solvent model data, respectively, whereas our results have not been scaled. The relative magnitudes of the QCISD and present DFT IR intensities agree well and both agree that the strongest mode is the OCO stretch mode at our calculated frequency of 1003 cm<sup>-1</sup>. For the two lowest frequency torsion modes, QCISD predicts that the lowest mode is more intense whereas DFT predicts that the second lowest is more intense. Our assignments mostly agree with the QCISD assignments and the six assignments given by DFT PCM results. The assignments where the agreement is not clear are the 1345 and 1387 cm<sup>-1</sup>, where our calculations show both CH<sub>2</sub> and OH modes and Barrientos et al. found only CH<sub>2</sub> modes. The MP2 data mostly agree with QCISD except that the MP2 assignments for the 1374 cm<sup>-1</sup> and 1441 cm<sup>-1</sup> modes are CH<sub>2</sub> scissor and CH<sub>2</sub> wag, which are reversed with respect to QCISD.

We have also shown a comparison with the IR spectroscopic study by Lugez et al. [6]. Their data are also in good agreement with the calculated frequencies, although only 8 frequencies were reported by Lugez et al. Their tentative assignments of the 1354-1359, and 1424-1426 cm<sup>-1</sup> modes are COH in-plane bends only, in contrast with the MP2 and QCISD calculations that predicted CH<sub>2</sub> wag or scissor modes. We note again that our data shows both COH bending and CH<sub>2</sub> character for our modes closest to their frequencies.

## Methoxylated methylene glycol (MMG)

Three conformers for MMG were relaxed and their structures are shown in Figure 4. Conformer A is the lowest energy conformer, and conformers B and C are 6.7 and 9.8 kJ/mol higher in energy, respectively. The relative ground state energies of the three conformers are in good agreement with the DFT B3LYP study by Wrobel et al. [7] who predicted that conformers B and C are 8.8 and 9.2 kJ/mol (2.1 and 2.2 kcal/mol) higher in energy.



Figure 4. Three conformers of methoxylated methylene glycol (MMG).

MMG has 24 vibrational modes and Table 3 shows the vibrational frequencies in ascending order for the lowest energy conformer alongside previous DFT and experimental data. Vibrational frequencies for conformer B are similar to those for conformer A, and are given in the Supporting Information. There is reasonable agreement between our DFT PBE frequencies and the B3LYP

frequencies of Wrobel et al, although the deviation increases at higher frequencies, as shown in the correlation plot in Supporting Information. Wrobel et al also measured the IR spectra for MMG in argon matrix isolation experiments at 10 K and these results also agree well with the DFT calculations. Vibrational frequencies for the second lowest energy conformer are shown in Table S3 in Supporting Information and the frequencies are similar to those of conformer A. We also compared the frequencies with a gas phase FT-IR study by Johnson and Stanley [17]. These results were mostly in agreement, except for the 1052 cm<sup>-1</sup> mode. All other studies found a single peak/frequency near 1050 cm<sup>-1</sup> but Johnson and Stanley suggested there were two peaks in the measured spectra, at 1046 and 1052 cm<sup>-1</sup>. However, we believe that there is likely to be only a single peak and that the appearance of the double peak is due to noise in their measured spectra. We also compared our results with an IR spectroscopy study of MMG formed by irradiation of liquid methanol [18]. They only reported 5 frequencies and did not report assignments so we have presented the frequencies in Table 3 beside the nearest frequencies. Our assignments for conformer A are also presented in Table 3. The agreement with the assignments of Wrobel et al. is mixed. In general, there is good agreement regarding the involved groups but often the mode types, e.g., bend vs scissor do not agree. However, we note that some of the modes for MMG are not easy to define and appear to be a mix of different modes. The files containing the mode frequencies and modes are available as electronic Supporting Information.

#### Diglycol (DG)

For DG, we considered four conformations of the molecule, labelled A, B, C and D, as shown in Figure 5. Conformer A has the lowest energy, followed by B, C and D, which are 3.4 kJ/mol, 10.9 kJ/mol and 28.5 kJ/mol higher in energy, respectively. Structures similar to conformers A and B were calculated by Delcroix et al. [23] and they found that conformer A is 1.3 kJ/mol lower in energy. However, they discarded this conformer as they expected that the H-bond between the OH groups would not be predominant in solution.



Figure 5. Four conformers of diglycol (DG).

DG has 27 vibrational frequencies and these are shown in Table 4 in ascending order for conformers A and B. The frequencies for conformers A and B differ by less than 50 cm<sup>-1</sup> over our experimental range of 800-1700 cm<sup>-1</sup>. The study by Delcroix et al. calculated vibrational frequencies for the isolated DG molecule and also for the solvated molecule in implicit and explicit water [23]. Here we have reported their data for the isolated molecule only as they reported more than 27 vibrational frequencies for DG with explicit water. For the anharmonic isolated case they reported only 26 frequencies and for the harmonic isolated case they reported all 27 frequencies and used a scaling factor of 0.989 [23] (also see Supporting Information). Their results are shown alongside the present results in Table 4 and the correlation plot is shown in the Supporting Information. In our experimental range (800-1700 cm<sup>-1</sup>), our predicted frequencies

are systematically lower in frequency than both sets of data by Delcroix et al. The assignments of conformers A and B are similar. The 3<sup>rd</sup> and 4<sup>th</sup> lowest mode assignments are in a different order and a few other modes with mixed assignments differ slightly. Conformer B agrees quite well with Delcroix's assignments for conformer B, with a few differences for the modes with mixed assignments. Delcroix also studied DG in water and found that a few frequencies were significantly different in gas and solvated phases. The only frequency affected that lies within our experimental range is their 1478 cm<sup>-1</sup> mode (corresponding to our frequency at 1442 cm<sup>-1</sup>) that shifts to 1520 cm<sup>-1</sup>.

#### Assignments of vibrational frequencies

In order to ascertain assignments of the peaks observed in IR and Raman spectra measured in formaldehyde-water-methanol solutions (see Figures 1-3), we analysed vibrational spectra of representative species (MG, MMG, DG) based on their simulated and previously measured vibrational frequencies (see Tables 2-4). In Table 5 we show assignments of peaks based on our analysis using DFT PBE as well as trends observed in spectra from dilution experiments (see Figures 2 and 3). We also show comparison with previous experimental data for Raman [14,15] and IR [19] spectra of formaldehyde-water solutions, the formaldehyde stock solution containing methanol [13] as well as MG [6] and MMG [7,17,18] species. We note that there are no experimental measurements of DG or any other related species on their own in a solvent or gas.



Figure 6. Correlation between calculated DFT frequencies and experimental peak wavenumbers based on assignments in Table 5. Discrepancies for methanol are discussed in the text.

Figure 6 shows a correlation between calculated DFT frequencies and experimental peak wavenumbers based on assignments in Table 5. The correlation shows that most significant differences between calculated vibrational frequencies (which were obtained in the absence of solvent) and experimental vibrational frequencies for spectra measured in solutions pertain to those cases where there are known to be large solvent effects. Most notably for methanol, while some frequencies remain mostly unchanged between gas and liquid phase spectra, it has been reported [28] that gas phase modes at 1060 cm<sup>-1</sup> and 1345 cm<sup>-1</sup> shift to 1115 cm<sup>-1</sup> and 1418 cm<sup>-1</sup> respectively, in the liquid state (see Table S2 in Supporting Information). Similarly, the only frequency within our experimental range where DG vibrational frequencies calculated by Wrobel et al. significantly shifted when going from gas to aqueous solutions is our frequency at 1442 cm<sup>-1</sup> that shifts from 1478 cm<sup>-1</sup> to 1520 cm<sup>-1</sup> [7].

The early work of Matsuura et al. reported Raman spectra of commercial formaldehyde solutions (37 %wt. formaldehyde, 10-15 %wt. methanol). When we compare Raman spectra shown in Figure 1 here with the spectra in Figure 1 in Matsuura et al., we find that both spectra are very similar. While analysing their spectra Matsuura et al. assumed that only MG and MeOH were present and results of their computational analysis were interpreted based on this assumption, ignoring oligometric and methoxylated species present in these solutions, as there was only limited knowledge of formaldehyde speciation at that time. This work was later followed up by Mohlmann and Lebrun et al., where Raman spectra of aqueous formaldehyde solutions were collected and speciation information was taken into account from then available literature [10]. Since only relatively concentrated solutions were considered, resolution of respective species was not possible, although some assignments specific for oligomeric species were suggested (see Table 5). We can see from Table 5 that some frequencies that we assigned to MG or DG were reported by neither Mohlmann nor Lebrun et al., (e.g., peaks D, J, P) and some of their assignments are in disagreement with each other (e.g., peaks K, L/M, N). However, based on our analysis, these peaks correspond to combinations of CH<sub>2</sub> twists (peaks K, L) or wags (peak N) and COH bends in DG. Importantly, many peaks have contributions from multiple species, even in the absence of methanol: while peaks A, C, G, J, K, O, P correspond to DG (and likely to other oligomers as well), peaks D, E, F, L, M, N, Q correspond to both MG and DG. In the presence of methanol, this becomes even more complex, as peaks B, E, F, G, H, I, K, M, N, O, P, Q, R have contributions from MMG and peaks E, G, H, N, O, P, Q have contributions from methanol.

Ryabova et al. and to some extent also Lebrun et al. based their assignments on quantitative correlations between areas of deconvoluted peaks and concentrations of specific species in aqueous formaldehyde solutions. While Lebrun et al. have only identified peaks likely to correspond to oligomers (in agreement with our assignments), Ryabova et al. also assigned specific peaks to MG, DG and TG (1050, 1072 and 1100 cm<sup>-1</sup>, respectively, in their IR spectra), in addition to assigning frequencies 992 and 1026 cm<sup>-1</sup> C-O-C stretch in oligomers. While we agree with the last two assignments (our peaks D and E), we note that both these peaks have a contribution from MG as well, and their MG assignment (our peak F) has a contribution from DG as well. Their TG assignment (our peak G) has a contribution plot shown in their Figure 1 and may be an artefact. On the other hand, it was observed by Lebrun et al. that the peak in Raman spectra around 1059 cm<sup>-1</sup> (corresponding to our peak F) shifts to about 1070 cm<sup>-1</sup> as concentration of formaldehyde increases, indicating that there may be two peaks near each other,

each of them possibly corresponding to contributions with varying proportions from similar species.

Overall, we can see that similar vibrational frequencies can be expected for multiple formaldehyde species and there are only very few peaks that may be uniquely assigned to specific groups of species. In aqueous formaldehyde systems (in the absence of methanol) these are: peak H for the hydrated monomer (MG) and peaks A, C, G, J, K, O, P for oligomers (DG and higher). In the presence of methanol, there are only few peaks, B, I and R, specific to just MMG within the spectral range investigated, with others overlapping with MG, DG or methanol. However, since there are numerous vibrational modes that are close to each other, the corresponding peaks in measured spectra would overlap to various degrees and therefore there may be additional peaks that have yet to be identified, e.g., see shoulder around 1380 cm<sup>-1</sup>. Furthermore, there are further minor species present in moderately diluted formaldehyde-water-methanol solutions, such as MDG [4], which have not been specifically considered here; for example, there could be contribution from MDG vibrational modes to peaks A (assigned to DG) or B (assigned to MMG).

We have also attempted to correlate concentrations of formaldehyde-based species obtained previously using NMR [4] with intensities of deconvoluted peaks from IR and Raman spectra obtained from formaldehyde-water-methanol solutions [24]. However, the main challenge has been that the solution matrix composition influences both IR and Raman spectra in terms of the areas under the peaks, as well as vibrational frequencies [29], some of which are sensitive to changes in the water-methanol ratio in the solvent matrix. This can be seen clearly in IR and Raman spectra of pure methanol when compared with those of methanol-water mixtures (see Figures S7 and S8 in Supporting Information).



Figure 7. Ratio of areas of deconvoluted peaks E and G from IR spectra of formaldehyde stock solutions diluted with water (F-W), formaldehyde stock solutions diluted with methanol (F-M) and methanol-water mixtures (M-W).

Figure 7 shows the ratio of areas of deconvoluted peaks E and G observed in IR spectra at 1025 and 1110 cm<sup>-1</sup> collected for formaldehyde-water-methanol solutions, as well as those for methanol-water mixtures. For formaldehyde stock solutions, the ratio of the peak areas increases with increasing dilution with water (as the methanol mole fraction in the solution decreases) and it also increases with increasing dilution with methanol (as the methanol mole fraction in the solution increases). In the case of dilution with water this is expected, as peak E has a contribution from MG, DG, MMG and methanol, while peak G has contribution from DG, MMG, and methanol but not from MG, and upon dilution speciation distribution shifts toward MG. On the other hand, the trend observed for methanol-water mixtures shows that the peaks corresponding to methanol change their relative areas as the solvent matrix changes, so in the case of dilution with methanol the spectrum is gradually approaching that of methanol. Therefore it is not possible to quantify concentrations of individual species bases on simple univariate assumptions about relationships between peak areas and individual species concentrations, especially when the solvent matrix is changing. Furthermore, concentrations of various species which are in mutual equilibria would be strongly correlated, so for example concentrations of MG and MMG would be approximately proportional to each other at a given methanol to water ratio. Therefore more sophisticated multivariate models taking into account the solvent matrix would be required for quantitative interpretation of vibrational spectra of formaldehyde-water-methanol systems.

# Conclusions

We have reported IR and Raman spectra for formaldehyde-water-methanol solutions across a wide range of formaldehyde concentrations (mole fractions from 0.004 to 0.279) and solvent compositions (methanol mole fractions form 0.001 to 0.974). Solutions were prepared by diluting commercial formaldehyde stock solutions (37%wt formaldehyde, 13%wt methanol) with either water or methanol. Resulting solutions contained a wide range of hydrated and methoxylated formaldehyde species, from just methylene glycol (MG) and methoxylated methylene glycol (MMG) at the highest dilution with water and methanol, respectively, to a complex mixture of their oligomers at the highest formaldehyde concentration. We have present a detailed characterisation of vibrational frequencies of formaldehyde-water-methanol systems using a combination of experimental measurements and quantum mechanical density functional theory simulations, while for the first time considering vibrational spectra of oligomeric mixtures of formaldehyde species together with those of specific representative species, including MG, MMG and diglycol (DG).

# Tables

Comula	Formaldehyde	Added of	liluent [ml]		Mole fract	ions
Sample	stock solution [ml]	Water	Methanol	Water	Methanol	Formaldehyde
F-W 1-0	3.00	0	0	0.629	0.092	0.279
F-W 1-1	2.00	2.00	0	0.828	0.043	0.130
F-W 1-3	1.00	3.00	0	0.917	0.021	0.063
F-W 1-6	1.00	6.00	0	0.953	0.012	0.035
F-W 1-10	1.00	10.0	0	0.970	0.007	0.022
F-W 1-33	1.00	33.0	0	0.991	0.002	0.007
F-W 1-67	1.00	67.0	0	0.995	0.001	0.004
F-M 1-0	3.00	0	0	0.629	0.092	0.279
F-M 1-1	2.00	0	2.00	0.416	0.399	0.185
F-M 1-3	1.00	0	3.00	0.248	0.642	0.110
F-M 1-6	1.00	0	6.00	0.155	0.777	0.069
F-M 1-10	1.00	0	10.0	0.103	0.852	0.046
F-M 1-33	1.00	0	33.0	0.035	0.949	0.016
F-M 1-67	0.50	0	33.5	0.018	0.974	0.008

Table 1. Compositions of formaldehyde solutions diluted with water (F-W) or with methanol (F-M). F-W 1-0 and F-M 1-0 correspond to the original formaldehyde stock solution.

Table 2. Vibrational frequencies  $[cm^{-1}]$ , infrared intensities and assignments for methylene glycol (MG). Vibrational modes: torsion (t), stretch (v), symmetric stretch (v<sub>a</sub>), antisymmetric stretch (v<sub>a</sub>), bend ( $\delta$ ), twist ( $\tau$ ), scissor ( $\sigma$ ), wag ( $\omega$ ), rock( $\rho$ ).

Vibration	Vibrational Frequencies				IR intensity		Assignment				
DFT PBE	MP2	QCISD	DFT PCM	Expt	DFT PBE	QCISD	DFT PBE	MP2	QCISD	DFT PCM	Expt
Present	Barrie ntos et al. [21]	[21]	Delcro ix et al. [23]	Lugez et al. [6]	Present	Barrient os et al. [21]	Present	Barriento s et al. [21]	Barriento s et al. [21]	Delcroi x et al. [23]	Lugez et al. [6]
366	333	349	332	-	1.65	161.9	t(OH)	t(COH)	t(COH)	-	-
371	368	366	361	-	3.44	57.5	t(OH)	t(COH)	t(COH)	-	-
539	538	545	538	545-548	1.36	62.7	δ(OCO)	δ(OCO)	δ(OCO)	-	t(CO)
973	1010	985	986	-	0.38	20.5	ρ(CH <sub>2</sub> ) + δ(OH)	ρ(CH <sub>2</sub> )	ρ(CH <sub>2</sub> )	-	-
995	1019	1024	1025	-	1.98	95.2	v <sub>s</sub> (OCO)	v(OCO)	v(OCO)	v <sub>s</sub> (CO)	-
1003	1048	1070	1026	1055-1057	6.55	243.7	v <sub>a</sub> (OCO)	v(OCO)	v(OCO)	$v_a(CO)$	v(CO)
1165	1188	1178	1177	-	0.04	1.6	$\tau(CH_2) + \delta(OH)$	δ(COH)	δ(COH)	-	-
1323	1329	1341	1339	1335	0.55	28.7	$\omega(CH_2) + \delta(OH)$	δ(COH)	δ(COH)	-	ω(CH <sub>2</sub> )
1345	1350	1360	1355	1354-1359	0.07	3.4	$ \begin{aligned} \tau(\mathrm{CH}_2) \\ + \delta(\mathrm{OH}) \end{aligned} $	τ(CH <sub>2</sub> )	τ(CH <sub>2</sub> )	-	δ(COH) in- plane
1387	1374	1413	1407	1424-1426	1.08	60.7	$\omega(CH_2) + \delta(OH)$	σ(CH <sub>2</sub> )	ω(CH <sub>2</sub> )	-	δ(COH) in- plane
1463	1441	1498	1472	-	0.00	0.3	σ(CH <sub>2</sub> )	ω(CH <sub>2</sub> )	σ(CH <sub>2</sub> )	-	-
2951	2807	2956	2959	-	1.34	49.9	v <sub>s</sub> (CH <sub>2</sub> )	v(CH <sub>2</sub> )	v(CH <sub>2</sub> )	$v_s(CH_2)$	-

2999	3020	3007	3016	2978	0.87	33.5	$v_a(CH_2)$	ν(CH <sub>2</sub> )	v(CH <sub>2</sub> )	v <sub>a</sub> (CH <sub>2</sub> )	v <sub>a</sub> (CH <sub>2</sub> )
3683	3648	3710	3753	3564	0.63	47.5	v(OH)	v(OH)	v(OH)	v <sub>a</sub> (OH)	v(OH)
3684	3648	3711	3755	3638-3639	0.58	27.5	v(OH)	v(OH)	v(OH)	v <sub>s</sub> (OH)	v(OH)

Table 3. Vibrational frequencies [cm<sup>-1</sup>], infrared intensities and assignments § of methoxylated methylene glycol (MMG) conformer A.

Vibrationa	Vibrational frequencies				IR Intensity			Assignment §			
DFT PBE	DFT B3LYP	Expt in argon matrix	Expt gas	Expt in liquid methanol	DFT PBE	DFT B3LYP	Expt in argon matrix	DFT PBE	DFT B3LYP	Expt argon matrix	
Present	[7]	[7]	[17]	[18]	Present	[7]	[7]	Present	[7]	[7]	
137	144	-	-	-	0.06	2	-	t(CH <sub>3</sub> )	t(CH)	-	
180	191	-	-	-	0.17	4	-	t(CH <sub>3</sub> )	t(CH <sub>3</sub> )	-	
351	358	-	-	-	1.48	44	-	$t(OH) + \delta(COC)$	t(OH)	-	
383	394	-	-	-	1.48	36	-	t(OH)	-	-	
564	583	576	-	-	0.58	18	14	$\sigma(OCO) + \sigma(COC) + t(OH)$	$\nu(COC) + \delta(OH)$	$\nu(COC) + \delta(OH)$	
904	925	936	930	925	0.93	23	20	v <sub>s</sub> (COC)	v <sub>s</sub> (COC)	v <sub>s</sub> (COC)	
981	1024	1020	1020	-	4.57	100	94	$v(CO) + \delta(OH)$	v(COH)	v(COH)	
1004	1037	1044	1046, 1052 ♭	1067	1.86	56	-	$\rho(CH_2) + \delta(OH) + \nu_a(OCO) + \nu_a(COC)$	$v(COC) + \sigma(CH_2)$	$\nu(COC) + \sigma(CH_2)$	
1097	1134	1125	1129	-	2.79	81	100	$     v_a(COC) + \rho(CH_2) + \delta(OH) $	$\nu(COC) + (CH_2)^{\#}$	$\nu(COC) + (CH_2)^{\#}$	
1130	1171	1150	-	1123	0.12	2	2	ρ(CH <sub>3</sub> )	(CH <sub>2</sub> ) #	-	
1153	1200	1187	1189	1200	0.36	16	34	$\rho(CH_3) + \tau(CH_2) + \delta(OH)$	$(CH_3)^{\#} + (CH_2)^{\#}$	$(CH_3)^{\#} + (CH_2)^{\#}$	

1252	1299	1286	-	1303	0.35	9	9	$\tau(CH_2) + \delta(OH)$	$(CH_3)^{\#} + (CH_2)^{\#}$	$(CH_3)^{\#} + (CH_2)^{\#}$
1334	1379	1355	1350	-	0.22	5	10	$\tau(CH_2) + \delta(OH)$	$(CH_3)^{\#} + \delta(CH_2)$	δ(CH <sub>2</sub> )
1374	1433	1406	1410	-	0.42	13	3	$\omega(CH_2) + \delta(OH)$	ω(CH <sub>3</sub> )	ω(CH <sub>2</sub> )
1414	1473	1444	-	-	0.17	6	4	CH <sub>3</sub> umbrella	δ(CH <sub>3</sub> )	δ(CH <sub>3</sub> )
1428	1486	1452	1450	-	0.14	2	7	d-o(CH <sub>3</sub> )	δ(CH <sub>3</sub> )	-
1446	1506	1470	-	-	0.02	2	6	$\sigma(CH_2) + d - \sigma(CH_3)$	δ(CH <sub>3</sub> )	-
1454	1511	-	-	-	0.23	3	-	$d-\sigma(CH_3) + \sigma(CH_2)$	δ(CH <sub>3</sub> )	-
2921	2994	2878	-	-	1.33	31	13	v <sub>s</sub> (CH <sub>3</sub> )	$v_a(CH_2)$	$v_a(CH_2)$
2932	3017	2929	-	-	1.57	39	15	v <sub>s</sub> (CH <sub>2</sub> )	v <sub>a</sub> (CH <sub>2</sub> )	v <sub>a</sub> (CH <sub>2</sub> )
2991	3064	2964	-	-	0.79	20	13	v <sub>a</sub> (CH <sub>2</sub> )	$v(CH) + v_s(CH_2)$	$v(CH) + v_s(CH_2)$
2995	3073	-	-	-	0.91	25	-	v <sub>a</sub> (CH <sub>3</sub> )	$v(CH) + v_s(CH_2)$	$v(CH) + v_s(CH_2)$
3057	3125	-	-	-	0.41	12	-	v <sub>a</sub> (CH <sub>3</sub> )	$v(CH) + v_s(CH_2)$	-
3682	3815	3631	3695	-	0.67	20	18	ν(OH)	v(OH)	v(OH)

<sup>§</sup> Vibrational mode labels: torsion (t), stretch (v), symmetric stretch (v<sub>s</sub>), antisymmetric stretch (v<sub>a</sub>), bend ( $\delta$ ), twist ( $\tau$ ), scissor ( $\sigma$ ), wag ( $\omega$ ), rock( $\rho$ ). The CH<sub>3</sub> 'umbrella' mode refers to a symmetric in-phase deformation, the CH<sub>3</sub> d- $\sigma$  mode is a degenerate (or nearly) scissor mode, which is an out-of-phase half-methyl deformation [30].

<sup>#</sup> Wrobel labelled these modes as  $\omega$  in the paper, but this did not correspond to a wag mode as wag modes were explicitly labelled 'wag'. Wrobel did not define t and  $\delta$  so we assume these correspond to torsion and bending modes.

<sup>b</sup> The two values reported by Johnson and Stanley are likely to be just a single peak.

Table 4. Calculated vibrational frequencies, IR intensities and mode assignments § for the lowest energy conformer, A, and second lowest energy conformer B, of diglycol (DG).

Vibrational frequencies					IR intensity	Assignment §				
DFT PBE	DFT PBE	DFT PBE	DFT PBE	DFT PBE DFT PBE		DFT PBE	DFT PBE	AIMD		

		anharmonic	harmonic scaled					
Present A	Present B	[23]	[23]	Present A	Present B	Present A	Present B	[23]
121	25	72	75	0.14	0.12	t	t	t
145	161	166	165	0.05	0.36	t	t	t
308	285	302	305	3.49	0.37	t(OH)	t	σ(COC)
393	299	356	351	0.17	2.18	σ(COC)	t(OH)	ω(OH)
464	370	-	372	1.35	2.42	$t(OH) + \sigma(OCO)$	t(OH)	-
508	550	562	583	1.87	0.18	t(OH)	$\sigma(COC) + \sigma(OCO) + t(OH)$	sym σ(OCO)
617	566	582	594	0.40	0.70	$\sigma(COC) + \sigma(OCO) + t(OH)$	$\sigma(COC) + \sigma(OCO) + t(OH)$	asym σ(OCO)
886	896	916	936	1.45	0.79	$v_{s}(COC) + v_{a}(OCO)$	v <sub>s</sub> (COC)	v <sub>s</sub> (CO)
949	935	1011	1002	4.19	8.86	$v_a(COC) + v_a(OCO)$	$v_a(OCO) + v_a(COC) + \delta(OH)$	v <sub>a</sub> (CO)
985	981	1012	1030	2.85	3.88	$\rho(CH_2) + \delta(OH)$	$\frac{\rho(CH_2) + v(CO_H) +}{\delta(OH)}$	v <sub>a</sub> (CO)
1033	1024	1040	1057	4.40	1.14	$v(CO_H) + \delta(OH)$	v(CO <sub>H</sub> )	$v_s(CO) + \delta(CH_2)$
1050	1030	1127	1103	1.06	1.53	$\rho(CH_2) + \delta(OH)$	$\rho(CH_2) + \delta(OH)$	v <sub>a</sub> (CO)
1078	1076	1140	1128	2.35	1.96	$\begin{array}{l} v_a(COC) + \rho(CH_2) + \\ \delta(OH) \end{array}$		$\delta(CH_2) + v_s(CO)$
1211	1211	1245	1262	0.76	0.50	$\tau(CH_2) + \delta(OH)$	$\tau(CH_2) + \delta(OH)$	$\delta(CH_2) + \delta(OH)$
1271	1262	1287	1315	0.41	0.37	$\tau(CH_2) + \delta(OH)$	$\tau(CH_2) + \delta(OH)$	$\delta(CH_2) + \delta(OH)$
1331	1332	1378	1381	0.20	0.20	$\tau(CH_2) + \delta(OH)$	$\tau(CH_2) + \delta(OH)$	$\delta(OH) + \delta(CH_2)$
1343	1336	1374	1384	0.26	0.16	$\omega(CH_2) + \delta(OH)$	$\tau(CH_2) + \delta(OH)$	$\delta(OH) + \delta(CH_2)$
1376	1363	1401	1418	0.53	0.56	ω(CH <sub>2</sub> )	$\omega(CH_2) + \delta(OH)$	sym δ(CH <sub>2</sub> )
1397	1382	1428	1446	0.98	0.27	$\omega(CH_2) + \delta(OH)$	$\omega$ (CH <sub>2</sub> ) + $\delta$ (OH)	δ(CH <sub>2</sub> )

1442	1449	1478	1502	0.05	0.12	σ(CH <sub>2</sub> )	σ(CH <sub>2</sub> )	σ(CH <sub>2</sub> )
1456	1460	1481	1506	0.07	0.02	σ(CH <sub>2</sub> )	σ(CH <sub>2</sub> )	σ(CH <sub>2</sub> )
2883	2950	2901	3037	1.57	2.10	v <sub>s</sub> (CH <sub>2</sub> )	v <sub>s</sub> (CH <sub>2</sub> )	v <sub>s</sub> (CH)
2944	2956	2901	3043	1.36	0.17	v <sub>s</sub> (CH <sub>2</sub> )	v <sub>s</sub> (CH <sub>2</sub> )	v <sub>s</sub> (CH)
3008	3004	2948	3100	0.90	0.77	v <sub>a</sub> (CH <sub>2</sub> )	v <sub>a</sub> (CH <sub>2</sub> )	v <sub>a</sub> (CH)
3051	3053	2950	3102	0.59	0.45	v <sub>a</sub> (CH <sub>2</sub> )	v <sub>a</sub> (CH <sub>2</sub> )	v <sub>a</sub> (CH)
3646	3686	3715	3844	1.68	0.61	v(OH)	v(OH)	v <sub>a</sub> (OH)
3687	3722	3716	3846	0.66	0.90	ν(OH)	ν(OH)	v <sub>s</sub> (OH)

<sup>§</sup> Vibrational modes: torsion (t), stretch (v), symmetric stretch (v<sub>s</sub>), antisymmetric stretch (v<sub>a</sub>), bend ( $\delta$ ), twist ( $\tau$ ), scissor ( $\sigma$ ), wag ( $\omega$ ), rock( $\rho$ ).

Table 5. Assignments of vibrational frequencies for species in formaldehyde-water-methanol solutions.

	Wavenumbers and Assignments §										
Peak label	IR/ Raman	DFT PBE	Raman (aqueous formaldhyde)	Raman* (aqueous formaldehyde )	Raman (stock solution w/ methanol)	IR (aqueous formaldehyde )	IR (MG)	IR (MMG in gas)	IR (MMG in MeOH)	IR (MMG a/b conformers in argon)	
	Present	Present	Möhlmann et al. [14]	Lebrun et al. [15]	Matsuura et al. [13]	Ryabova et al. [19]	Lugez et al. [6]	Johnson et al. [17]	Fanetti et al. [18]	Wrobel et al. [7]	
Α	910/905	<b>DG 886</b> $v_{s}(COC) + v_{a}(OCO)$	927 v <sub>s</sub> (OCO/COC ) in oligomers	910 v <sub>s</sub> (OCO) in oligomer chains	907 v <sub>s</sub> (OCO) in MG						
В	920/915	<b>MMG 904</b> v <sub>s</sub> (COC)						930	925	936/940	

										v <sub>s</sub> (COC)
С	960/950	<b>DG 949</b> v <sub>a</sub> (OCO) + v <sub>a</sub> (COC)		933 v <sub>s</sub> (OCO) in oligomer endgoups						
D	990/995	<b>MG 973</b> ρ(CH <sub>2</sub> ) + δ(OH) <b>DG 985</b> ρ(CH <sub>2</sub> ) + δ(OH)				992 v(COC) in oligomers				
E	1025/ 1025	$\label{eq:constraint} \begin{array}{l} \textbf{MeOH 1008 } v(CO) \\ + \ \delta(OH) \\ \textbf{MG 995 } v_{s}(OCO) \\ \textbf{MMG 981 } v(CO) + \\ \delta(OH) \\ \textbf{DG 1033 } v(CO_{H}) + \\ \delta(OH) \end{array}$		1040 v <sub>a</sub> (OCO) in oligomer chains	<b>1034</b> ν(CO) in liquid MeOH	1026 v(COC) in oligomers		1020		<b>1020/1019</b> v(COH)
F	1060/ 1055	$\begin{array}{c} \textbf{MG 1003 } v_a(OCO) \\ \textbf{MMG 1004 } \rho(CH_2) \\ + \delta(OH) + \\ v_a(OCO) + \\ v_a(COC) \\ \textbf{DG 1050 } \rho(CH_2) + \\ \delta(OH) \end{array}$	<b>1063</b> v <sub>a</sub> (OCO)	<b>1059</b> v <sub>a</sub> (OCO) in oligomer endgoups	<b>1055</b> ν <sub>a</sub> (OCO) in MG	<b>1050</b> v(OCO) in MG ( <b>1072</b> v(OCO) in DG)	<b>1055-</b> <b>1057</b> v(CO)	1046 & 1052 ♭	1067	<b>1044/1064</b> ν(COC) + σ(CH <sub>2</sub> )
G	1110/	<b>MeOH 1052</b> v(CO)+ δ(OH) +		1111	<b>1109</b> ρ(CH <sub>3</sub> ) in liquid	1100		1129		1125/1119

	1110	$\begin{split} \rho(CH_{3}) \\ \textbf{MMG 1097} \\ \nu_{a}(COC) + \rho(CH_{2}) \\ + \delta(OH) \\ \textbf{DG 1078} \nu_{a}(COC) \\ + \rho(CH_{2}) + \delta(OH) \end{split}$		γ(HCH)	MeOH 1111 ρ(CH <sub>2</sub> ) in MG	v(OCO) in TG				v(COC) + (CH <sub>2</sub> )#
H	1155/	MeOH 1128           ρ(CH <sub>3</sub> )           MG 1165 τ(CH <sub>2</sub> ) +           δ(OH)           MMG 1130 ρ(CH <sub>3</sub> )	<b>1138</b> ρ(HCH)*		<b>1157</b> ρ(CH <sub>3</sub> ) in liquid MeOH				1123	1150/1143
I	1195/ 1195	<b>MMG 1153</b> ρ(CH <sub>3</sub> ) + τ(CH <sub>2</sub> ) + δ(OH)						1189	1200	<b>1187/1186</b> (CH <sub>3</sub> ) <sup>#</sup> + (CH <sub>2</sub> ) <sup>#</sup>
J	1210/ 1210	<b>DG 1211</b> τ(CH <sub>2</sub> ) + δ(OH)								
K	1275/ 1265	MMG 1252 τ(CH <sub>2</sub> ) + δ(OH) DG 1271 τ(CH <sub>2</sub> ) + δ(OH)	<b>1268</b> δ(COH) in plane	1257 t(HCH)	<b>1262</b> t(CH <sub>2</sub> ) in MG				1303	<b>1286/1285</b> (CH <sub>3</sub> ) <sup>#</sup> + (CH <sub>2</sub> ) <sup>#</sup>
L	1300/ 1315	MG 1323 ω(CH <sub>2</sub> ) + δ(OH) DG 1331 τ(CH <sub>2</sub> ) +		1315 w(HCH)	<b>1316</b> ω(CH <sub>2</sub> ) in		<b>1335</b> ω(CH <sub>2</sub> )			

		δ(OH)			MG			
М	1330/ 1330	$\begin{array}{c} \textbf{MG 1345 } \tau(CH_2) + \\ \delta(OH) \\ \textbf{MMG 1334 } \tau(CH_2) \\ + \delta(OH) \\ \textbf{DG 1343 } \omega(CH_2) + \\ \delta(OH) \end{array}$	<b>1333</b> τ(HCH) in oligomers			 <b>1354-</b> <b>1359</b> δ(COH) in plane	1350	 <b>1355/1356</b> δ(CH <sub>2</sub> )
N	1420/ 1410	$\begin{tabular}{lllllllllllllllllllllllllllllllllll$	<b>1429</b> ω(HCH)	<b>1408</b> δ(COH) in plane	<b>1412</b> δ(COH) in plane in MG	 <b>1424-</b> <b>1426</b> δ(COH) in plane	1410	 <b>1406/1414</b> ω(CH <sub>2</sub> )
0	1450/ 1450	MeOH 1422 CH3 umbrella           MMG 1414 CH3 umbrella           DG 1397 ω(CH2) + δ(OH)		<b>1440</b> δ(COH) in plane	1450 CH3symdeformationin liquidMeOH1453 δ(COH)in plane inMG	 		 <b>1444/1444</b> δ(CH <sub>3</sub> )
P	1470/ 1460	MeOH 1441 d-           σ(CH <sub>3</sub> )           MMG 1428 d-			<b>1465</b> CH <sub>3</sub> deg. deformation in liquid	 	1450	 <b>1452/1452</b> δ(CH <sub>3</sub> )

		σ(CH <sub>3</sub> )			МеОН			
		<b>DG 1442</b> σ(CH <sub>2</sub> )						
Q	1490/	MeOH 1455 d-	1495	1492	1488	 	 	1470/1470
	1480	σ(CH <sub>3</sub> )	$\sigma(\text{HCH})^*$ in	δ(HCH)	$\sigma(CH_2)$ in			δ(CH <sub>3</sub> )
		<b>MG 1463</b> σ(CH <sub>2</sub> )	oligomers		MG			
		<b>MMG 1446</b> σ(CH <sub>2</sub> )						
		$+ d-\sigma(CH_3)$						
		<b>DG 1456</b> σ(CH <sub>2</sub> )						
R	1500/	MMG 1454 d-				 	 	
	1500	$\sigma(CH_3) + \sigma(CH_2)$						

<sup>§</sup> Vibrational modes: torsion (t), stretch (v), symmetric stretch (v<sub>s</sub>), antisymmetric stretch (v<sub>a</sub>), bend ( $\delta$ ), twist ( $\tau$ ), scissor ( $\sigma$ ), wag ( $\omega$ ), rock( $\rho$ ). The CH<sub>3</sub> 'umbrella' mode refers to a symmetric in-phase deformation, the CH<sub>3</sub> d- $\sigma$  mode is a degenerate (or nearly) scissor mode, which is an out-of-phase half-methyl deformation [30]. The final column shows experimental IR values by Wrobel *et al* and the two wavenumbers correspond to two different conformers (a and b) of MMG.

\*Note that Lebrun *et al* did not define their symbols for mode assignments so we reported their assignments as shown in their paper. The original paper by Mohlmann uses  $\gamma$  to denote rock and  $\delta$  to denote scissor.

<sup>#</sup> Wrobel labelled these modes as  $\omega$  in the paper, but this did not correspond to a wag mode as this was labelled as 'wag'. Wrobel did not define t and  $\delta$  and we assume these correspond to torsion and bending modes.

<sup>b</sup> The two values reported by Johnson and Stanley are likely to be just a single peak, which we have attributed to our peak F.

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