Mid-infrared fiberoptic evanescent field spectroscopy for in-situ monitoring of tetrahydrofuran hydrate formation and dissociation

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<td>Manuscript ID</td>
<td>AN-COM-10-2016-002237.R1</td>
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<tr>
<td>Article Type</td>
<td>Communication</td>
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<td>Date Submitted by the Author</td>
<td>n/a</td>
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<td>Complete List of Authors</td>
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Mid-infrared fiberoptic evanescent field spectroscopy for in-situ monitoring of tetrahydrofuran hydrate formation and dissociation

M. Schwenk, a A. Katzir b and B. Mizaikoff a

Tetrahydrofuran is a relevant auxiliary molecule when storing carbon dioxide or hydrocarbons as gas hydrates. The present study demonstrates the applicaton of in-situ mid-infrared fiberoptic evanescent field absorption spectroscopy for studying the formation and dissociation of THF hydrates. Thereby, the utility of this analytical technique providing unique molecular-level insight even at harsh environmental conditions is evidenced.

Gas hydrates - or more general, clathrate hydrates - are crystalline host-guest compounds, which are able to trap small molecules inside a network of hydrogen-bonded water structures. The guest size depends on the cavity diameter and thereby determines the final structure of the clathrate hydrate. In the case of natural gas hydrates, the three main structures are the cubic structures sI and sII, and the hexagonal structure sH. Each structure is composed of a different number of water molecules combined into the respective water cages with an increasing capability to trap larger molecules from sI to sII. 1-2 The formation conditions are generally in the range of several MPa pressure, and at temperatures close to the freezing point of water with the respective values being interrelated. Detailed tabulated phase equilibria data for many systems can be found in literature. 3-4 Next to gaseous guests such as short chained hydrocarbons (e.g., methane, ethane, propane), carbon dioxide, and nitrogen, also small others including tetrahydrofuran (THF) may form hydrates at appropriate conditions. In the case of gaseous clathrate hydrates, the diffusion of gas into the liquid phase, adequate pressure, and sufficient supply of the guest are among the key requirements for hydrate growth. Tetrahydrofuran hydrates on the contrary form directly from liquid mixtures of water and THF at an ideal molar ratio of 1:17 THF:water 4 at atmospheric pressure and at temperatures below 277.4 K. 5 Due to these modest formation conditions, the experimental effort when studying THF hydrates is low compared to e.g., CO2 or hydrocarbon hydrates, as THF hydrates can be synthesized in virtually any kind of reaction vessel connected to a cooling system. In contrast, gas containing clathrates require a high-pressure autoclave along with appropriate gas supply/mixing systems. The resulting structure of pure THF hydrates is known as sI, which comprises 16 dodecahedral (S15) and 8 hexakai decahedral (S166) cages. The larger hexakai decahedral cavities are occupied by THF, while the smaller dodecahedral cavities remain empty. 2,4,6 Next to the capability forming hydrates of its own, THF also has proven to be an excellent thermodynamic promoter during the formation of carbon dioxide and methane gas hydrates. The addition of small amounts of THF actively shifts the formation conditions towards more modest regimes, thereby resulting in lower pressure and higher temperature requirements. 7-12 The utilisation of THF as an assistive molecule for storing hydrogen as hydrate is another interesting concept that is currently being researched. 13-16 These properties render THF hydrates an appealing model system to investigate fundamental processes involved during hydrate formation and dissociation in molecular-level detail for further application as auxiliary molecule in hydrate storage scenarios involving various gases.

In the field of hydrate research, Raman spectroscopy is usually considered the method of choice for monitoring hydrate formation. Accordingly, Prasad et al. investigated THF hydrate across a sizeable temperature range (i.e., 90 – 300 K) via Raman spectroscopy, and observed a splitting of the ring breathing mode at 90 K, which may be attributed to different guest sites within structure sI hydrate. 17 Infrared (IR) spectroscopic techniques are rarely used, which is mainly attributed to the rather strong absorption of water in the mid-infrared (MIR; 3-15 μm) regime usually rendering the detection of additional species difficult. Bertie and Othen investigated for the first time ethylene oxide clathrate hydrate in the range of 20–4000 cm⁻¹. 18,19 Fleyfel and Devlin later applied IR spectroscopy when studying carbon dioxide and THF hydrates.
determining pronounced bands for CO\textsubscript{2} in the small and large cages depending on the present structure, as well as a band characteristic for THF in the large cages of structure sl. 20

More recently, several publications have demonstrated the fundamental potential of IR spectroscopy in the field of hydrate science. 21–25 The application of fiber optic evanescent field absorption spectroscopy (MIR-FFES) was pioneered by the research team of Mizaioff introducing significant advancements for MIR-based monitoring systems in gas hydrate research. The developed gas hydrate monitoring systems enable performing in-situ dynamic IR studies during hydrate formation and dissociation, since fiber optic waveguides may be subjected to elevated pressure conditions. Consequently, methane, ethane and propane containing hydrates have been investigated via MIR-FFES, as reported by Dobbs et al. and Luzinova et al. 26,27 Recently, this technique was adapted towards the investigation of carbon dioxide containing gas hydrates, and for monitoring relevant additives during hydrate formation and dissociation. 28

The present study demonstrates the first application of MIR-FFES for in-situ monitoring of the formation and dissociation processes of THF hydrates. Spectroscopic information at the molecular-level of the relevant reactants (i.e., water and THF) was correlated with continuously recorded temperature traces, thereby introducing MIR-FFES as an innovative analytical strategy for studying THF hydrates. The developed reaction vessel enables the IR spectroscopic investigation of the initial reaction mixture (i.e., water/THF molar ratio 17:1), and its in-situ conversion into crystalline hydrate. MIR-FFES is based on the principles of attenuated total reflection (ATR) spectroscopy, i.e., more general, internal reflection spectroscopy (IRS) introduced by Harrick. 29 While in conventional IR-ATR usually hemispherical single-reflection or trapezoidal multi-reflection crystalline waveguides (e.g., made from ZnSe, ZnS, Ge, Si, diamond, etc.) are applied, MIR-FFES utilizes cylindrically shaped MIR-transparent optical fibers, here, polycrystalline fiberoptics made from AgCl\textsubscript{0.3}Br\textsubscript{0.7} as IRS waveguide. Light is propagated via total internal reflection along the fiber giving rise to an evanescent field at each reflection point, which exponentially decays in intensity into the adjacent medium. Thereby, the detection of IR-active species in the vicinity of the fiber surface is enabled via evanescent field absorption spectroscopy. 30

The experimental setup (Fig. 1) is assembled inside a darkened fume hood to minimize light-induced degradation of the fiber. A custom designed glass reaction vessel (Fig. 1 #3) featured fiber optic feed-through ports sealing the silver halide fiber (diameter 900 \textmu m, AgCl\textsubscript{0.3}Br\textsubscript{0.7}) into the vessel, thus enabling spectroscopic access. Silver halide fibers are mechanically robust while retaining some flexibility due to their polycrystallinity, and feature a wide spectral window in the MIR region. 30 The reaction vessel has been constructed double-walled facilitating effective cooling of the reaction solution via a Huber Ministat 125 cooling system. A Pt100 thermistor (Fig. 1 #4) was immersed into the solution just above the fiber surface for recording the process temperature throughout the entire experiment close to the analytical volume probed by the evanescent field. A Bruker Alpha FTIR spectrometer (Bruker Optik GmbH, Ettlingen, Germany) (Fig.1 #1) in an OEM version was used. Three 2" gold-coated off-axis parabolic mirrors (OAPMs; Edmund Optics Inc., Barrington, NJ, USA) (Fig. 1 #2) guided the IR radiation from the source via the fiber optic towards the detector element of an external Stirling-cooled mercury-cadmium-telluride (MCT) detector (Model K508, Infrared Associates, Stuart, FL, USA) (Fig.1 #6). The detector was connected to the FTIR spectrometer via an impedance-matched MCT-1000 pre-amplifier. Underneath the reaction vessel, a magnetic stirrer (Thermo Fisher Scientific Inc., Waltham, MA, USA) (Fig. 1 #5) has been mounted. The experiments were conducted in a solution of demineralized water and THF (Merck KGaA, Darmstadt, Germany with a purity of \textgreek{29.9\%} at a molar ratio of 17:1. The solution was stirred for approx. 1 h prior to analysis enabling adequate mixing of both liquids. Prior to recording a background spectrum of air, the fume hood comprising the setup was flushed with dry compressed air for 1.5 h. Afterwards, the solution was added to the reaction vessel. The experiment was not started before another 1.5 h ensuring equilibration of the entire environment. The cooling system was programmed such that the solution was cooled to approx. 1 °C during 1 h after reaching an initial temperature of approx. 22 °C during 10 min. Initial fluctuations were resulting from equilibrating the cooling system. Upon reaching the target temperature (1 °C), it was maintained constant for 72 h following heating to 22 °C during 1 h. IR spectra were recorded at intervals of 10 min with a spectral resolution of 2 cm\textsuperscript{-1} between 4000 – 600 cm\textsuperscript{-1}. Each spectrum averaged 100 spectral scans. The reaction solution was stirred throughout the entire measurement. Eventually, the stirrer stopped due to the formation of crystalline hydrate.

Fig. 1 MIR-FFES setup: (1) Bruker Alpha FTIR spectrometer, (2) gold coated off-axis parabolic mirrors (OAPMs), (3) customized glass reaction vessel, (4) Pt100 thermistor, (5) magnetic stirrer, (6) Stirling-cooled mercury-cadmium-telluride (MCT) detector.

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Fig. 2 shows exemplary MIR-FEFS spectra (i.e., solution vs. solid phase) obtained during a hydrate formation experiment. The black spectrum shows the spectral features of the reaction solution at the start of the experiment, whereas the red spectrum reveals the characteristics of THF hydrate obtained later during the experiment. Water has several pronounced absorption bands in the MIR region, i.e., the OH stretch vibration (ν\text{OH}) located at 3750-2750 cm\(^{-1}\), the 3\textsuperscript{rd} overtone of the vibration mode (3ν\text{v}) at 2300-1900 cm\(^{-1}\), the HOH bending vibration (ν\text{b}) at 1850 and 1520 cm\(^{-1}\), and the libration mode (ν\text{l}) located below 1000 cm\(^{-1}\). Each of these features is specifically altered upon the crystallisation of hydrates, as already shown for methane, ethane, propane, and carbon dioxide containing hydrates via MIR-FEFS.\(^{20-28}\) The respective changes are similar to those appearing during cooling or crystallisation of water.\(^{31,32}\) The resulting structure of THF hydrate is cubic sii; hence, the spectroscopic changes during this study are expected to be qualitatively similar to previously conducted experiments with sii mixed CO\(_2\)/THF hydrate.\(^{33}\) As can be derived from Fig. 2, the ν\text{OH} shifts to the red, while the 3ν\text{v} and the ν\text{l} feature shift to the blue. Furthermore, the ν\text{b} decreases in intensity upon hydrate formation.

Next to water, THF also reveals distinctive absorption features, which are evident in the region of approx. 3015-2830 cm\(^{-1}\), and as a feature centered at 1041 cm\(^{-1}\). These spectral features are attributed to the CH stretching vibrations (ν\text{C-H}), and the asymmetric C-O-C stretch vibration (ν\text{as(C-O-C)}), respectively (see Fig. 2).\(^{33}\) The ν\text{as(C-O-C)} of THF appears with a shoulder at approx. 1051 cm\(^{-1}\), as magnified in the inset of Fig. 2. After comparison of this feature with literature it is apparently resulting from a mixture of two different species. Based on the findings of Mizuno \textit{et al.} and regarding the molar fraction of water in the reaction solution (i.e., in this study 0.94), the detected THF feature may be interpreted as a mixture of THF connected to two (1053 cm\(^{-1}\)) or at least three water molecules (1038 cm\(^{-1}\)).\(^{33}\) The inset of Fig. 2 provides a detailed view of the region of interest regarding the ν\text{as(C-O-C)} of THF. In the hydrate spectrum (red), a new feature centered at 1072 cm\(^{-1}\) was observed next to the broad feature of THF in aqueous solution (@ approx. 1051 / 1040 cm\(^{-1}\)). This feature has been assigned by others to the asymmetric C-O-C stretch vibration of THF inside the large 5\textsuperscript{12}6\textsuperscript{4} cages of the resulting sii hydrate structure.\(^{20,23,24,34}\)

The fact that for THF in solution there is still a signal evident may be explained by remaining non-converted

![MIR-FEFS spectra](image_url)

**Fig. 2** Exemplary MIR-FEFS spectra recorded during a THF hydrate experiment; black: spectrum of the sample solution at the start of the experiment; red: spectrum with THF hydrate present. Inset: ν\text{as(C-O-C)} of THF appearing with a shoulder at approx. 1051 cm\(^{-1}\).

![Temperature trace and spectral data](image_url)

**Fig. 3** Temperature trace (a) and spectral data traces of water absorption features with time during a hydrate formation experiment (b - e): red dashed lines indicate the time of nucleation: approx. 22.7 h.
water/THF in the vicinity of the fiber after initiation of crystal formation. Changes of these features with time were monitored via peak integration and peak picking methods of individual characteristic features. 

Fig. 3 displays the evolution of the absorption features of water throughout a hydrate formation experiment. Next to the spectroscopic data traces, the temperature trace is displayed clearly revealing the initiation of nucleation, as the temperature increases to approx. 4.32 °C after approx. 22.7 h into the experiment. In order to compensate interferences of water vapour mainly in the region of the HOH bending absorption (Fig. 2), a compensation algorithm provided by the spectrometer software (Bruker OPUS software package) was applied to the spectra prior to integration. It is clearly evident that simultaneous with hydrate nucleation the spectral data shift sharply (indicated by red dashed line in Fig. 3 b-e) reaching an equilibrated state with no more changes detectable. Only upon heating of the sample solution to approx. 22 °C and thereby dissociating the hydrate these shifts recover to the state prior to nucleation (starting after approx. 73.3 h). These sharp shifts occurring simultaneously to the temperature peak indicate that nucleation occurs in immediate vicinity of the silver halide fiber surface. This direct response renders MIR-FEFS a novel and highly promising analytical technique for studying hydrate formation, and specifically THF hydrates at molecular level detail and in real time. In the following, the spectral changes associated with water are discussed in detail. The υOH feature shifts to the red (Fig. 2), which is interpreted as an increase of highly networked water species in the hydrate phase. 31 Quantitative monitoring of this change was obtained via peak area integration (3310-3099 cm⁻¹; Fig. 3 b). 35 For analysis of the υ2 feature, the area of 1750-1577 cm⁻¹ was integrated. The resulting decreases in intensity (Fig. 2), and as a consequence, the decrease in peak area (Fig. 3 c) upon hydrate formation is likely due to a loss of induced transition dipole moment with increasing solidification of the sample. 31 Upon hydrate formation, the υ1 shifts to the blue (Fig. 2), thereby emerging within the detectable spectral region of the present setup limited by the spectral bandwidth of the Stirling-cooled MCT detector system. Analogous to the υOH absorption, partial peak area integration has been applied to the υ1 at 925-703 cm⁻¹ (Fig. 3 d), which enables monitoring of the shift towards higher wavenumbers, even though the feature appears not fully evident due to the limited detectable spectral range. While these features have been analysed via peak area integration methods, a peak picking method has been utilized to monitor the blue shift of the 3υ1 feature. Prior to peak picking, the wavelength segment of 2250-2000 cm⁻¹ has been smoothed via a moving average method (software package Essential FTIR 3.50.047; Operant LLC, Madison, WI, USA). The resulting peak positions vs. time are displayed in Fig. 3 e. Likewise, a sharp shift is observable at the point of nucleation, which is in excellent agreement with the other monitored features. 

To monitor changes associated with THF with time, the respective spectral features of THF have likewise been integrated. Integration borders were set at 1061-1020 cm⁻¹ for the υνC=O of THF in aqueous solution (Fig. 4, black dataset), and at 1088-1061 cm⁻¹ for the υνC=O of THF in the hydrate structure (Fig. 4; red dataset).

Similar to the water features displayed in Fig. 3, the changes related to the υνC=O feature of THF occur simultaneously with hydrate nucleation at 22.7 hours (Fig. 3 a). A sharp decrease in peak area is evident for the υνC=O feature of THF in solution (Fig. 4, black dataset), while an increase appears for the feature assigned to the hydrated species following this point of time (Fig. 4, red dataset). Hence, this study provides unique continuously monitored spectral evidence that indeed dissolved THF is continuously enclathrated into hydrate ice cages, as the associated spectral features behave complementarily until equilibrium is reached. At equilibrium, hydrate formation is assumed to be completed. Upon dissociation of the THF hydrate by heating it was proven that both features recover their initial state. The incorporation of THF molecules into the hydrate structure results in a depletion of THF in the remaining solution, which is clearly evidenced in the present experiments. The observed sharp changes occurring simultaneously in both datasets display the incorporation of THF molecules into the clathrate structure in real time onwards after 22.7 h into the experiment. In order to substantiate these findings, this experiment has been repeated individually three times with all experiments revealing nucleation simultaneous with sharp changes of the monitored spectral features of water and THF. It should be noted that in comparison to previous reports (see refs. 20,23,24,34) identifying the feature at 1072 cm⁻¹ as the signal of the υνC=O THF feature inside the large hexakaidecahedral sll cage, this feature was evident already at significantly higher temperatures during the present experiments.

Fig. 4 Peak area integration evaluating the evolution of υνC=O of THF vs. time. The feature originating from THF in aqueous solution (black, right ordinate) decreases sharply at the point of nucleation (i.e., @ 22.7 hours, red dashed line), while the feature assigned to THF inside the clathrate structure behaves complementarily and increases sharply (red, left ordinate).
Conclusions
This study demonstrates for the first time the utility of infrared fiber optic evanescent field absorption spectroscopy for in-situ monitoring the formation and dissociation of THF hydrates in real time and at molecular level detail. During THF hydrate formation the spectral changes associated with water and THF could be unambiguously identified, and correlated to the recorded temperature traces. Unique spectroscopic access is facilitated via a silver halide fiber submerged into the reaction solution serving as attenuated total reflection waveguide. Thereby, direct studies of hydrates providing molecular-level insight on the changes occurring during clathrate formation and dissociation for THF-containing hydrates is available for the first time. The present study reveals the potential of MIR-FEFS-based monitoring systems not only for in-situ studies of hydrates, but corroborates the utility of fiber optic IR-ATR spectroscopy in demanding analytical environments by pairing direct spectroscopic access with mechanical robustness.

Acknowledgements
The authors greatly acknowledge the collaboration with A. Katzir and his research team at Tel-Aviv University (Israel) for providing customized silver halide fibers. Furthermore, the machine shop at Ulm University is acknowledged for assistance in establishing the MIR-FEFS gas hydrate setup. Partial financial support of this work by the European Union FP7 Project SCHeMA – Integrated In Situ Chemical Mapping Probes (Grant Agreement Number 614002) is gratefully acknowledged.

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