Viewpoint

Far reaching potentials of far infrared spectroscopy in catalysis research

Huixiang Li, Z. Conrad Zhang *

1. Introduction

Fourier transform infrared (FTIR) spectroscopy has played an important role in identifying molecular finger prints of reactants, products and intermediates, in elucidating the mechanisms of a broad range of reactions, and in probing the fundamental nature of active sites of many catalysts by both in-situ and ex-situ measurements [1,2]. Specifically, FTIR spectroscopy measures the absorption of chemical bonds or groups in the 10–12500 cm⁻¹ spectral region. The study of FTIR is commonly performed in three spectral segments, near infrared (NIR, 4000–12500 cm⁻¹), middle infrared (MIR, 400–4000 cm⁻¹), and far infrared (FIR, 10–400 cm⁻¹) (Fig. 1). MIR spectroscopy has been well established and most widely applied to study various bonds among C, H, O, N, S, B, X (halides). MIR application covers a wide range of fields [3–6], including chemicals, food safety, environmental monitoring, materials, etc. The application of MIR in catalysis is relatively mature in following the transformation of chemical bonds and groups involved in reactions and over catalyst surfaces.

However, for structural investigation of chemical bonds involving heavier elements, particularly those in inorganic compounds such as metal oxides or salts, spectroscopy in the MIR and NIR regions is not applicable, because the vibration energies of these bonds fall in the FIR spectral region. Therefore, FIR spectroscopy can be used for the characterization of some inorganic compounds [7,8] and some metal coordination complexes [9,10] that may be catalytically active. FIR in the low spectral energy region has been shown to be well suited to study lattice vibrations [11], hydrogen bonds [12], as well as the skeleton vibration of aromatic molecules [13]. It should be emphasized that FIR remains much under developed and the least applied infrared tool, especially in the study of catalytic mechanisms. This mini review is therefore intended to draw attentions to the potential applications of FIR in catalysis research by discussing some important features of early studies and of some recent literatures. Potential future directions are suggested in the development of FIR for catalysis studies.

2. FIR characterization of catalyst structures

Catalysts are commonly categorized according to their physical state in corresponding reaction systems. Homogeneous catalysts refer to those that are fully miscible and disperse molecular level, while heterogeneous catalysts refer to those in which only surfaces contribute to catalytic activity. When organometallic complexes are used as homogeneous catalysts in solvent media in which reactions take place, such complexes and the reactants are both dissolved in the solvents. Although enzymes as catalysts are also homogeneous in aqueous reaction media together with reaction substrates, absorption bands by water in the IR spectral region (both MIR and FIR) is so intense that IR is not a suitable tool for such systems. Inorganic solid catalysts are mostly present as a heterogeneous phase in either liquid or gaseous reaction phase. Bonds among heavier elements, mostly metals in metallic or higher oxidation states often play major roles during catalysis. Techniques such as extended X-ray absorption fine structure (EXAFS), X-ray crystallography, mass spectrometry, nuclear magnetic resonance (NMR), UV-Vis spectroscopy, X-ray powder diffraction (XRD), X-ray photoelectron spectroscopy (XPS) and other specialized methods are often used for characterization of homogeneous catalysts or heterogeneous catalyst [14,15]. These techniques have been proven versatile and informative pertaining to obtain critical mechanistic insights on the role of the catalysts in catalyzing reactions of interest. However, it should be noted that some of these tools are only available in specialized laboratories that are not readily accessible to many researchers.
Even though FIR has not been widely applied as a catalyst characterization tool, like other IR tools, it is simple to operate, and is suited for the characterization of a wide variety of solid catalysts and organometallic complexes. As an underdeveloped spectroscopic tool, FIR offers the potentials to become a low-cost technology to provide structural information related to catalyst functions.

FIR is suited to study organometallic structural configurations. FIR studies of square planar trialkylphosphine complexes of the types cis and trans MX2L2 (M = Pd, Pt; X = Cl, Br, I; L = Me3P, Et3P) have shown that metal-phosphorus stretching frequencies v(M–P) occur in the narrow range 400–440 cm–1 [16]. Metal-halogen v(M–X) frequencies however, are found in a wider frequency range and are markedly dependent on the cis or trans geometry of the complex [17]. Boorman et al. [18] studied tetrahedral and square planar adducts of nickel (II) halides with tertiary phosphines and di-phosphines with FIR. Correlations of v(Ni–X) and v(Ni–P) with stereo-chemistry indicate that FIR is a useful technique to distinguish tetrahedral from cis or trans square planar complexes.

Metal clusters consisting of several metal atoms show spectra unique for each cluster size and exhibit fingerprints of the cluster’s structure [19–21]. Fielicke et al investigated the FIR spectra of neutral and cationic niobium clusters of Nb50+/+. The experimental spectra were recorded in the 85–600 cm–1 region that covered the structure-specific vibrational fingerprint range for these metal clusters. A comparison of the experimental and calculated Far-IR spectra allows for the identification of the cluster structures. The comparison with theory revealed that the overall geometries for the cations and neutrals are very similar [21].

Important to heterogeneous catalysts, FIR was shown to be able to differentiate structures of solid state catalysts [22–24]. For examples, the alumina phases, boehmite, diaspor, gibbsite and bayerite, can be distinguished by FIR spectroscopy because the low-frequency modes of FIR are generally the most sensitive to the changes of AIO4 units. Boehmite has two characteristic FIR bands at 366 and 323 cm–1, while diaspor has five bonds at 354, 331, 250, 199 and 158 cm–1. Gibbsite shows three characteristic FIR bands at 371, 279 and 246 cm–1, whereas bayerite shows six bonds at 383, 345, 326, 296, 252 and 62 cm–1 [22]. Thus FIR allows for the study and the differentiation of the stretching of AIO4 units to characterize these four alunina phases in bauxites.

Microporous materials, particularly zeolites, are widely used as selective adsorbents for separations and as heterogeneous catalysts for catalytic processes. IR spectroscopy of adsorbed probe molecules is a widely used technique for zeolite characterization [25]. CO is the most frequently used molecule to probe zeolite structural and surface properties by monitoring the variation of its IR absorption wavenumber in reference to that of CO in gas phase. Otero Areán et al. [26] studied the stretching vibration of weak M–CO bond (M = Li, Na, K, Rb, Cs) formed upon CO adsorption in different alkali-metal exchanged zeolites at 77K. As shown in Fig. 2, the wavenumber of the cation-carbon bond vibration v(M–C) increases linearly with increasing the inverse of the square of the cation radius. The adsorption wavenumber of CO on an alkali ion does not vary appreciably in different zeolites, such as ZSM-5, Y, X and FER. The largest Na–CO vibration wavenumber is 141 cm–1. The high sensitivity of the v(M–CO) to the metal ions (M = Li, Na, K, Rb, Cs) in the FIR region suggests that it would be possible to quantitatively determine the relative abundance of these metal ions in zeolites.

For CO adsorbed (at 210 K) on ZnO, Saussey et al. [27] reported the ν(Zn–CO) fundamental stretching at 215 cm–1. The higher wavenumber of the cation-carbon bond v(M–C) observed for the CO/ZnO system than the CO/alkali-metal exchanged zeolites can be explained by the stronger Zn–CO bond strength due to the higher electric charge density of the Zn2+ ion. Thus FIR spectroscopy of appropriate probe molecules may reveal structural details of surface sites by accurately measuring the low energy modes. Furthermore, Engrström and Ryberg [28] compared the vibrational properties of the low-energy modes of a molecular and an atomic adsorbate: CO and O on Pt(111) surface by far infrared spectroscopy. For oxygen atoms an antiasorption dip associated with the frustrated translation was observed, while in contrast, no such dip was observed for the frustrated rotations of CO. Obviously, MIR, FIR spectroscopies become complementary in studying gas adsorption process in catalysis [25,26]. They can be used to study a wide range of energy from 10 to 4000 cm–1 and to track light variations of the molecular vibration of adsorbed interaction with the substrate through the whole adsorption process.

3. In-situ FIR study in catalysis

In-situ IR has been widely used for catalyst characterization to gather information about catalyst structure, concentration of reactants, intermediates and products. In-situ FIR could lift, to some extent, limitations in the region of middle infrared spec-
trum to allow for characterization of some catalytic reactions. For example, MIR only follows the changes in reactants, intermediates and products by monitoring the specific chemical bonds or structures in reactions catalyzed by homogeneous catalysts, while FIR may be suited to study the interactions between the catalysts and the reactants, intermediates and products. The FIR spectroscopy may therefore provide rich information for understanding the catalytic mechanism.

E.g., a considerable number of investigations have been focused on glucose catalytic conversion to 5-hydroxymethylfurfural (5-HMF) [29], a platform chemical that can be converted to various chemicals, biofuels and biomaterials [30].

A considerably large number of publications have appeared that reported results of the reaction mechanism with various analysis techniques [31–33] since the discovery of a new catalytic system (Scheme 1) that enabled the conversion of glucose to 5-HMF [34]. Typical techniques such as NMR, XRD, UV-Vis, MIR are not capable of revealing the catalytic mechanism for this complex system. Most recently, an in-situ FIR spectroscopy tool has been successfully applied to the study of the coordination chemistry of various metal chlorides in an ionic liquid, 1-butyl-3-methylimidazolium chloride, or [BMIM]Cl [10].

3.1. Important factors in FIR analysis

In-situ FIR spectroscopy has limitations for two reasons. (1) FIR spectroscopy has been recorded with a DTGS/Polyethylene detector, the sensitivity of which is much lower than that of the MIR spectral detector, and (2) the energy of light source is lower in FIR region, so that the spectral noise becomes greater. Furthermore, water vapor in air greatly impacts on the quality of the far-infrared spectra. Water vapor molecule has many rotating absorption peaks in the far infrared region. When the FIR absorption spectrum of the sample is weak, strong moisture absorption makes it difficult to clearly distinguish peaks due to absorption by the sample. Therefore, in measuring in-situ infrared spectroscopy, care must be taken to minimize the impact of water vapor from the absorption spectrum. For a non-vacuum FTIR spectrometer, dry air or nitrogen purge of the optical bench is critically important. In our experience, a customized sample compartment lid provides an efficient protective physical barrier by reducing the impact of ambient air on the samples under investigation. Care must be taken during sample transfer to minimize air exposure. It should be noted that spectral resolution and the water vapor spectrum are directly related. The number of rotational absorption peaks, the peak shape and peak position of water vapor molecules change with the resolution of the measurement: the higher the resolution, the larger the number of water vapor absorption bands is. An 8 cm⁻¹ resolution may be sufficient for the routine study in FIR spectrum. For a small amount of sample, the absorbance measured could be very low. In this case, a resolution of 16 cm⁻¹ could be used [35].

3.2. In-situ FIR study of the mechanism of metal chlorides catalyzed glucose conversion to 5-HMF

Four representative metal chlorides, CrCl₃, VCl₃, PtCl₂, and FeCl₃ were chosen based on their conversion, yield (Fig. 3) for glucose conversion to 5-HMF and studied by in-situ FIR in order to understand the distinctively different performance characteristics of the catalysts in correlation to their coordination chemistries.

Then in-situ FIR spectroscopy was employed to follow the trend of metal-Cl absorption band variation in the complexes in the process of the glucose conversion catalyzed by the metal chlorides in [BMIM]Cl (Figs. 4 and 5).

The dissolved MClₓ (M = Cr, V, Pt, Fe; x = 2, 3) were found to form new M complexes in the MClₓ/[BMIM]Cl/glucose reaction system under typical reaction conditions, as shown in Fig. 4. The absorbance of Cr-Cl band at 302 cm⁻¹ decreased gradually at the beginning (Fig. 4(a)) and then showed a limited restoration (Fig. 5(a)) after an extended period of reaction, due to the consumption of glucose. Meanwhile, the peak intensity of Cr-O (from glucose) coordination bond at 497 cm⁻¹ increased gradually (Fig. 4(a)) and then decreased slowly (Fig. 5(b)). Thus the FIR spectra indicate that Cr(III) center coordinated with glucose first and catalyzed the isomerization of glucose to fructose. In addition, in-situ far-infrared spectra of the VClₓ/[BMIM]Cl/glucose system (Fig. 4(b)) indicate that the V-Cl bond absor-

![Scheme 1](image1.png)

**Scheme 1.** The proposed pathway of glucose conversion to 5-HMF in 1-ethyl-3-methylimidazole chloride ([EMIM]Cl). Reproduced with permission from Ref. [29].

![Fig. 3](image2.png)

**Fig. 3.** The catalytic characteristics of metal chlorides for glucose conversion to 5-HMF in [BMIM]Cl at 96 °C. Reproduced with permission from Ref. [10].
ance at 287 cm⁻¹ declined much more than the Cr–Cl bond absorbance in the CrCl₃/[BMIM]Cl/glucose system with time, and with concomitant change in V–O bond FIR absorbance. It is possible that the vanadium ion coordinates with more than one glucose molecule, and can coordinate with the oxygen of a carbonyl group and a glycolaldehyde structure at the same time, resulting in increased side reactions dominated by humins. The spectra in Fig. 4(c) show the FIR features of the PtCl₂/[BMIM]Cl/glucose system. Both the glucose absorption peak at 554 cm⁻¹ and the Pt–Cl stretch vibration band at near 310 cm⁻¹ showed a less pronounced change in 80 min compared to that in CrCl₃/[BMIM]Cl/glucose (Fig. 4(a)). Evidently, replacement of Pt–Cl bond by Pt–O bond is not favored as indicated by the FIR spectra. As a result, PtCl₂ displays rather low catalytic activity for glucose conversion (Fig. 3). In the FeCl₃/[BMIM]Cl/glucose system, the absorbance of the anion...
[FeCl₄]⁻ at 381 cm⁻¹ decreased sharply and the absorbance of [FeCl₄]²⁻ [36] at near 280 cm⁻¹ increased gradually at the same time. Neither the absorbance at 381 cm⁻¹ or 280 cm⁻¹ was restored, even after 2 h (Fig. 4(d)). Evidently the strong Fe–O bonds contribute to the non-selective catalytic performance of Fe(III) catalyst due to the formation of dominantly humins.

3.3. In situ FIR study of the interaction strength between substrate and catalytic center

FIR spectroscopy was also applied to the study the coordination strength of metal chloride CrCl₃ in [BMIM][Cl] in the presence of model compounds with different oxygen sources. The results were correlated with the performance of CrCl₃ for glucose conversion in the presence of the same model compounds [10].

The Cr–Cl bond in [CrCl₆]³⁻ complex absorbs at 302 cm⁻¹. The intensity of this band decreased in response to the added model compounds, n-butanol (Fig. 6(a)), cyclohexanone (Fig. 6(b)), H₂O (Fig. 6(c)). The intensities of these bands were able to gradually restore by evaporation of the model compound at 100 °C. The coordination to Cr(III) by the oxygens from n-butanol, cyclohexanone, or water is so weak that a temperature of 100 °C was sufficient to disrupt the Cr–O model compound coordination bond and to reverse fully to the stable [CrCl₆]³⁻ complex. However, when excess glycolaldehyde was added to the CrCl₃/[BMIM][Cl] system, the Cr–Cl bond absorbance at 302 cm⁻¹ was decreased by nearly 1/2 (Fig. 6(d)), due to the formation of Cr–O (coming from glycolaldehyde) coordination bonds and Cr–Cl–Cr bridged bond [36]. Thus Cr(III) center is strongly coordinated with glycolaldehyde. And the strong interaction with the glycolaldehyde competes with glucose coordination to the Cr(III), resulted in a reduced glucose conversion and 5-HMF yield, as supported by the catalytic results in Fig. 7.

![Fig. 6](image_url)

**Fig. 6.** (a) FIR spectra of the Cr–O stretch vibration in the CrCl₃/[BMIM][Cl]/n-butanol system; (b) FIR spectra of the Cr–Cl stretch vibration in the CrCl₃/[BMIM][Cl]/cyclohexanone system; (c) FIR spectra of the Cr–O stretch vibration in the CrCl₃/[BMIM][Cl]/water system. (d) FIR spectra of the Cr–Cl stretch vibration in the CrCl₃/[BMIM][Cl]/glycolaldehyde system. The FIR spectra were recorded at 80 °C; the arrows in (a), (b) and (c) represent the variation trend of the Cr–O coordination bond during evaporation of the model compound. The background spectra of CrCl₃/[BMIM][Cl] system were taken before the addition of the model compound. Reproduced with permission from Ref. [10].

![Fig. 7](image_url)

**Fig. 7.** The effect of different probing model compounds on glucose conversion. (1) None; (2) Glycolaldehyde; (3) n-Butanol; (4) Cyclohexanone. Reproduced with permission from Ref. [10].
4. Conclusion and prospect

FIR spectroscopy is shown to be a convenient technique broadly applicable to characterize the catalyst structure, the process of gas adsorption on catalyst surface, and the determination of metal ions by using a probe molecule such as CO. The in-situ FIR technique is particularly suited to follow the progress of catalyst evolution and the products in catalytic reactions. The lattice vibration energies of metal clusters and metal oxides catalysts lie in FIR region, as well as the vibration energy of coordination complex catalysts. The force between adsorbent and adsorbate is typically weak so that it falls in the low energy mode within the FIR region. Furthermore, in-situ FIR spectroscopy could also track the changes of homogeneous catalyst in the course of catalytic reaction and explore the coordination strength between reagent and catalytic center. FIR spectroscopy could complement the application of MIR, NIR spectroscopy with the advantage of capable of measuring these low energy vibration modes accurately. In addition, there are various low energy vibration modes, such as hydrogen bonding, framework vibration of aromatics, and the rotation of the gas molecules. Thus FIR has a much far-reaching potentials in the study of catalysts and catalytic mechanisms. We expect much progress can be made by devoting efforts in expanding the potential scope and applications in the characterization of catalysts and in the mechanism study of catalytic processes.

Z. Conrad Zhang

State Key Laboratory of Catalysis, Dalian National Laboratory for Clean Energy, Dalian Institute of Chemical Physics, Chinese Academy of Sciences, Dalian 116023, Liaoning, China
Tel: +86-411-84379462
Fax: +86-411-84379462
E-mail: zc.zhang@yahoo.com
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References


Graphical Abstract

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Huixiang Li, Z. Conrad Zhang * 
Dalian Institute of Chemical Physics, Chinese Academy of Sciences; University of Chinese Academy of Sciences

Far infrared spectroscopy complements the application of mid-infrared, near-infrared spectroscopy with the advantage of capable of measuring these typical lower energy vibration modes accurately.