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Article

Raman and Terahertz Spectroscopic Investigation of Cocrystal Formation Involving Antibiotic Nitrofurantoin Drug and Coformer 4-aminobenzoic Acid

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Abstract: Cocrystallizaiton could improve most physicochemical properties of specific active pharmaceutical ingredients, which has great potential in pharmaceutical development. In this study, the cocrystal of nitrofurantoin and 4-aminobenzoic acid was prepared with solid-state (solvent-free or green-chemistry) grinding approach, and the above cocrystal has been characterized by Raman and terahertz vibrational spectroscopic techniques. Spectral results show that the vibrational modes of the cocrystal within the whole spectral region are different from those of the corresponding parent materials. The dynamic process of such pharmaceutical cocrystal formation has also been monitored directly with Raman spectra. These results offer us unique means for characterizing the cocrystal conformation from molecule-level and also provide us rich information about the reaction dynamic of cocrystal formation within pharmaceutical fields.

Keywords: nitrofurantoin; active pharmaceutical ingredient; 4-aminobenzoic acid; cocrystallization; intermolecular interaction; Raman spectroscopy; terahertz spectroscopy

1. Introduction

Cocrystallizaiton has recently gained great attention as an attractive method to obtain new forms of active pharmaceutical ingredients (APIs) with improved physical and physicochemical properties of drugs [1-6]. Cocrystal design for a specific API is based on evaluating possible synthons with reliable hydrogen bonding motifs (intermolecular interactions) sustaining crystal structures or other non-covalent and nonionic interactions within APIs and cocrystal formers (CCFs). The presence of functional groups acting as hydrogen bonds donors and acceptors in the molecules of APIs and CCFs may lead to the formation of cocrystallized substances. Cocrystallizaiton is particularly important in the pharmaceutical fields as a result of its impact on drug candidate performance, such as solubility, physical and chemical stability, melting point, as well as bioavilability of APIs, whereas the pharmacologic activity of the APIs is generally not influenced [4.5].

Nitrofurantoin (NF) is a well known antibacterial drug extensively used as an oral treatment for genitourinary tract infections, and its molecular structure is shown in Figure 1. NF belongs to a biopharmaceutical classification system (BCS) class II antibiotic drug, and has both low solubility and permeability problems [7-10]. The physicochemical properties, such as the dissolution rate, aqueous solubility, stability and bioavailability of NF in commercial tablets would decrease during its storage under different extraneous conditions [7,8]. These characteristics would significantly impact the therapeutic activity of NF drug. Multi-component crystals or also named cocrystals of NF with various CCFs based on the molecular attributes present have been reported recently [11-15].

The ability of these cocrystals has been shown in enhancing the photo-stability, and also clinically relevant physicochemical properties [11-13]. Two cocrystals NF-4-aminobenzoic acid and NF-urea come under the category of pharmaceutical cocrystals [12-15] because of the generally recognized as safe status of the coformers.

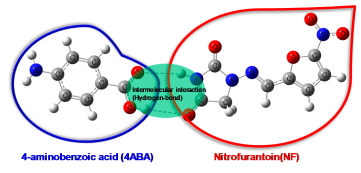


Figure 1. Molecular structures of nitrofurantoin (NF, right) and its conformer 4-aminobenzoic acid (4ABA, left), and the green shadow shows the intermolecular interaction due to hydrogen-bond effect between NF and 4ABA molecules.

Differences in crystal structures of cocrystals could be observed and evaluated by lots of research technologies. Traditional techniques that could be employed include powder X-ray diffractormetry (PXRD) [5,16-18,22], thermal analysis such as differential scanning calorimetry (DSC), and also solid-state nuclear magnetic resonance spectroscopy (ss-NMR) [19-22]. These established techniques are sometimes time consuming or lack in specific structural information about the crystalline and/or molecules, especially the effect from the intra-molecular and weak inter-molecular interactions. Vibrational spectroscopic techniques include mid-infrared, Raman and terahertz (THz) spectroscopy, which have been already used to explain cocrystallization of different solid-state pharmaceuticals successfully [15,22-35]. It is of great importance to use such specific techniques to probe the cocrystal characteristics in order to directly obtain the structural and/or dynamic information at molecular level to better know the structural information of the specific cocrystal and also its formation mechanism [22,24-26,29,30,33-36].

In this work, Raman and THz vibrational spectroscopy of typical cocrystal system formed between NF with cocrystal coformer 4-aminobenzoic acid (4ABA) compounds under solid-state grinding method has been reported. Significant difference among vibrational spectra of the formed cocrystal and involved parent starting molecules is observed. Moreover, the real formation process of NF-4ABA cocrystal could be monitored directly based on the experimental Raman and THz spectral results. The results indicate that vibrational spectroscopy can offer us attractive experimental methods to identify and characterize pharmaceutical cocrystals, and also provide potential tools for further monitoring the real-time reaction dynamic of cocrystals in-line to better know the corresponding reaction mechanism at molecule-level in pharmaceutical industry.

2. Results and Discussion

2.1. Raman spectral characterization of NF, 4ABA and its corresponding NF-4ABA cocrystal

Raman spectroscopy, as a valuable technique for the investigation of solid-state drugs, provides pretty rich intra-molecular structural information and also inter-molecular interaction (e.g. hydrogen bonding effect) interactions between molecules from molecular level. Raman spectra of NF, 4ABA, their physical mixture and cocrystal recorded in the wave-number range of $200 \sim 1700 \, \mathrm{cm}^{-1}$ are shown in Figure 2 (A) and (B).

Experimental results show that the Raman spectrum of physical mixture is almost identical to the linear combination of the individual spectra for the two involved starting components, NF and 4ABA respectively. It means that there is no intermolecular interaction effect between NF and

4ABA without grinding or other extraneous force. As we all know that the intensities and/or vibrational modes shown in Raman spectra are associated with the change in polarizability of specific chemical bonds within interested molecules. Due to the much more rigid and symmetrical structure of 4ABA molecule, it shows relative simple Raman bands which can be seen in Figure 2b, comparing with Raman spectra of NF shown in Figure 2a. Meanwhile Raman spectra of the corresponding cocrystal formed with grinding method can be easily differentiated from that of physical mixture which could be observed in Figure 2. Several band shifts and new characteristic features are observed in the whole spectral region as shown in Figure 2c and 2d, in which the band locations are given in next to characteristic peaks.

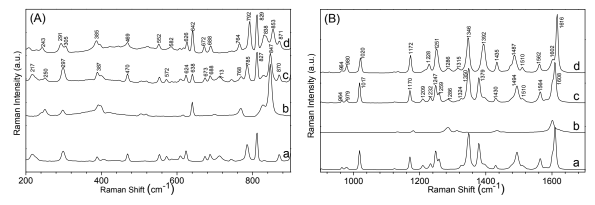


Figure 2. Raman spectra of NF (a), 4ABA (b), physical mixture (c) and cocrystal (d) in different spectral regions (A) 200–900 cm⁻¹; (B) 900–1700 cm⁻¹.

Within spectral region of 200 ~ 900 cm⁻¹ shown in Figure 2(A), both starting raw materials have band at 297 cm⁻¹. Upon the formation of cocrystal between NF and 4-ABA, this band would split into two peaks at 291 and 305 cm⁻¹, respectively. Some bands, such as 217 and 713 cm⁻¹, would disappear, while band at 838 cm⁻¹, the intensity becomes stronger and the peak width shows much broader after cocrystallization process. Pure NF Raman spectrum has bands at 785, 827 cm⁻¹ and the Raman spectrum of pure 4-ABA shows peak at 847 cm⁻¹, however all of these three characteristic features is up-shifted in the spectrum of the cocrystal, at 792, 829 and 853 cm⁻¹, respectively. As for the spectral region from 900 to 1700 cm⁻¹ shown in Figure 2(B), the intensities of 4ABA vibrational modes are pretty weak comparing with those of NF. The bands at 1209, 1269 cm⁻¹ shown in the physical mixture would disappear in cocrystal, meanwhile the characteristic bands at 1379 and 1608 cm⁻¹ would be blue-shifted to be at 1392 and 1616 cm⁻¹ upon the cocrystal formation.

Raman experimental results indicated that the carbonyl, carboxylic and amino groups of NF and 4ABA participate in hydrogen bonding formation during cocrystallization process. It also confirmed that the crystalline phase of the cocrystal is not simply combination of the individual starting components but the totally different crystal phase due to the intermolecular interaction effects, especially the hydrogen bonding interaction, between NF and 4ABA compounds.

2.2. THz spectral characterization of NF, 4ABA and its corresponding NF-4ABA cocrystal

The mechanism leading to THz absorption in molecular systems is dominated by the excitation of intra-molecular as well as inter-molecular vibrations. Therefore, the spectral features in THz spectra are usually considered to depend on intra-molecular and inter-molecular behavior (such as weak non-covalent bonds like hydrogen bonds or other non-valence bonds). It has especially promising applications in the pharmaceutical field for the analysis of pharmaceutical products throughout the drug development process. Using this technology it is possible to directly differentiating and quantifying different solid-state forms of APIs and their cocrystals, and also exploit structural information for such cocrystals whose molecular and/or crystal structures are stabilized by weak inter-molecular interactions.

THz absorption spectra of NF, 4ABA, and their physical mixture recorded in the range $0.20 \sim 1.70$ THz at room temperature are shown in Figure 3, with the THz signal transmitting through the nitrogen as the reference waveform for the corresponding samples. It can be clearly seen from Figure 3 that both individual components present significantly different absorption features in the recorded range. The absorption bands of 4ABA centered at 0.61, 0.82, 0.98, 1.34 and 1.59 THz, and the absorption intensity of NF is very weak in the whole frequency range, with a little broad band at around 1.26 and 1.60 THz position. The contrary intensities of NF and 4ABA shown in above Raman and terahertz spectra indicate that the selection rule of Raman scattering and is totally different from that of terahertz absorption phenomenon. Meanwhile, the physical mixture shows the major absorption peaks at the same region and exhibits some broader than that of 4ABA, which is due to the adding effect from the weak absorption of NF in the same region. The spectrum of physical mixture is similar to the linear addition of the individual absorption spectra for the two involved starting reactants.

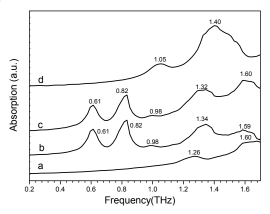


Figure 3. THz spectra of NF (a), 4ABA (b), physical mixture (c) and cocrystal (d) in 0.2 – 1.7 THz spectral region.

The spectra of cocrystals obtained from chemical-green grinding methods (shown in Figure 3d) are different compared with those of physical mixture, showing peaks which are not observed in the physical mixture and individual components. The absorption spectra of grinding cocrystal distinguish at frequencies 1.05 and 1.40 THz, with pretty broader bands comparing with that of physical mixture. It is well-known that the cocrystallization occurs as the result of intermolecular hydrogen bonding, nonionic or other nonconvalent interactions between two or more molecules of the different components [14-16]. So the experimental results indicate that the vibrational modes observed in these spectra are mostly intermolecular character between 4ABA and NF molecules during the cocrystal formation process.

The information obtained from THz spectra is in agreement with that of Raman spectra. It confirms that both Raman and THz spectroscopy could be promising alternatives to mainstream traditional analytical tools such as X-ray diffraction or thermal analysis for further study of solid-state cocrystal reactions in pharmaceutical fields.

2.3. Dynamic information during NF-4ABA cocrystal formation process

To achieve the rich dynamic information during cocrystal formation process between NF and 4ABA, the solid-state reactions were performed by grinding drily together equimolar involved parent components with different grinding times. Typical Raman and THz spectra showing the NF-4ABA cocrystal formation for up to 120 minutes grinding time can be seen in Figure 4 and 5. Overall, upon grinding, significant changes in both Raman and THz spectra can be detected with various grinding times shown in figures.

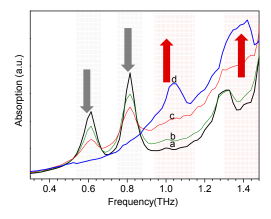


Figure 4. THz spectral comparison of the formed cocrystal between NF and 4ABA using grinding method under typical reaction times (a 0 min, b 15 min, c 30 min, d 80 min, and the shadow area shows the typical THz band change during grinding process).

In THz spectral comparison of the formed NF-4ABA cocrystal shown in Figure 4, the intensity of the characteristic absorption peak of the physical mixture (at 0.61, 0.82, and 1.34 THz, shown with grey arrows) decreases, while that of the cocrystal peaks (at 1.05 and 1.40 THz) increases gradually with typical grinding time during its formation process. There are two clear isobestic points appearing at around 0.67 and 0.87 THz position in Figure 4, which indicates the change of absorption peaks due to physical mixture and cocrystal in the reaction process. It means that the starting parent mixture was consumed while a continuation of the cocrystal formation takes place. After around 120 minutes grinding time, the THz absorption spectrum only shows characteristic peaks of cocrystal and characteristic features from the physical mixture (e.g. 0.61, 0.82 THz) disappear completely, which indicates a thorough cocrystallization between NF and 4ABA occurs finally.

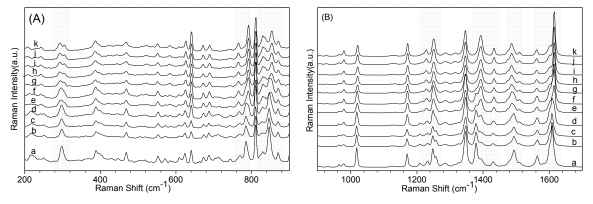


Figure 5. Raman spectral comparison of the formed cocrystal between NF and 4ABA using grinding method under typical reaction times (a 0 min, b 5 min, c 15 min, d 25 min, e 30 min, f 40 min, g 50 min, h 60 min, i 70 min, j 80 min, k 120 min, and the shadow area shows the typical Raman band change during grinding process) in different spectral regions (A) 200–900 cm⁻¹; (B) 900–1700 cm⁻¹.

According to Raman spectral result, the evolution of cocrystal characteristic peaks could also be clearly observed with increasing grinding time shown in Figure 5 (shown with grey arrows), meanwhile the intensity of characteristic peaks of NF and 4ABA is accompanied by a decrease or even disappearance. The more the grinding time performed, the higher the characteristic vibrational peaks of NF-4ABA cocrystal are obtained. This demonstrates that the longer-time grinding process might strongly induce and promote the cocrystallization transformation between NF and 4ABA. The Raman spectrum of cocrystal formation at 80 minutes is almost same as that of even longer 120 minutes and it indicates that the solid-state reaction is already completed around 80 minutes grinding time, and this result is also consistent with that of above THz spectra show in Figure 4.

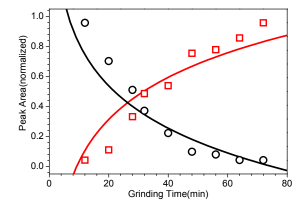


Figure 6. Observed normalized change of the relative peak area for the features at 1379 cm⁻¹ (characterized feature for the physical mixture, shown with black circle and black fitting-curving line) and 1392 cm⁻¹ (characterized feature for the cocrystal, shown with red square and red fitting-curving line) as the grinding time.

Due to the strong baseline influence and also pretty broad band shown in THz spectral results, it is not easy to reveal the accurate dynamic information with quantitative analysis for cocrystal transformation between NF and 4ABA. While Raman spectra are good choice to obtain such information because of its better signal-noise ratio and also pretty weak baseline influence within the whole spectral region which could be seen in Figure 5. Cocrystal formed from NF and 4ABA has characteristic Raman spectral features that could be used for quantification to obtain dynamic information so that the reaction mechanism could be achieved during cocrystallization process. Further analysis of the tentative changes in the Raman spectra shown in Figure 5 may reveal the dynamic of the NF-4ABA cocrystal formation with different grinding times.

Two Raman bands at 1379 and 1392 cm⁻¹(shown in Figure 5B), which are characterized for the physical mixture and cocrystal respectively, are chosen as an indicator. The dynamic information about time-dependent cocrystallization process of NF and 4ABA could be deduced by fitting the data between the normalized relative peak areas and grinding times with single exponential functions. The decay rate from physical mixture characteristic peak ($\tau_{1379} \approx 25.5$ mins) and the growth rate of cocrystal ($\tau_{1392} \approx 26.0$ mins) are consistent with each other, which indicates the successive promotion of conversion of NF-4ABA cocrystal from the starting raw materials. From the tentative changes shown in Figure 6 such solid-state pharmaceutical cocrystal transformation process and the corresponding dynamic information can be directly quantified and investigated.

3. Materials and Methods

NF and 4ABA samples were purchased from Sigma-Aldrich (structures shown in Figure 1) and used without further purification. Physical mixture was obtained by gently mixing two compounds above at a 1:1 molar ratio in a glass vial by using a vortex mixer. Grinding cocrystal was performed by co-grinding NF with 4ABA at 1:1 molar ratio in 25 mL stainless steel milling jars using a planetary ball mill (QM-3SP, gear type, Nanjing University Instrument Plant) with a frequency of 25 Hz at room temperature. A measured amount of product produced in the solid-state reaction at different grinding times was taken out for further spectra analysis.

All the samples were weighted into ~150 mg aliquots and poured into a steel die and subjected to ~4 MPa pressure for several minutes. The sample discs ~13 mm in diameter, ~1.5 mm thick were obtained and sealed in plastic bags before spectroscopic analysis.

Raman spectra of all samples were obtained using Fourier Transform Raman spectrometer (Thermo Nicolet 960, USA) with diode pumped 1064 nm solid-state laser as the near-IR source. Spectra were acquired over 500 scans at 2 cm⁻¹ resolution.

Terahertz time-domain spectroscopy (THz-TDS) measurement was performed with a commercial *Zomega-2* time-domain THz spectrometer (Zomega Terahertz Corp., Troy, USA). The

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THz beam is produced by a Ti:Sapphire oscillator ultrafast laser with a 75 MHz repetition rate, 780 nm center wavelength and ~100 fs pulse duration. The THz region of the spectrometer was purged with nitrogen gas to reduce the absorption effect from ambient water vapor. A total of three THz spectra (representing three complete sets of sample and reference measurements) were averaged for each final spectrum. The time-domain of the THz electric field was recording for the reference (without sample) and each sample. THz absorption spectra were obtained with the fast Fourier transform (FFT) operation.

4. Conclusions

Using Raman and THz-TDS spectroscopy, Raman scattering and THz absorption spectra of NF, 4ABA, the corresponding physical mixture and also cocrystals formed from chemical-green grinding methods have been recorded. Both spectra obtained in the measurement region provide finger-print information of starting raw components and cocrystals, which may be useful for the discrimination of these different solid-state pharmaceutical molecular systems. The dynamic process of NF-4ABA cocrystal formation can also be monitored directly with Raman and THz spectra. From the analysis of tentative changes in Raman spectra, such solid-state pharmaceutical cocrystallization transformation reaction and the corresponding kinetics information can be directly quantified and investigated. The reported results offer us the unique means and benchmark to identify and characterize the cocrystal conformation from molecule-level, and also further real-time monitor the reaction dynamic process of cocrystals in pharmaceutical fields.

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Conflicts of Interest: The authors declare no conflict of interest.

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