Abstract. Salt formation is a good method of increasing solubility, dissolution rate and consequently the bioavailability of poor soluble acidic or basic drugs [Polymorphism in Pharmaceutical Solids, Drugs and the Pharmaceutical Sciences, Vol. 192, 2nd edn, Informa Healthcare, New York, 2009]. The aim of this study was to obtain and to investigate the structural properties of the compound obtained by solvent drop grinding (SDG) method [Chem. Commun. (Camb.) 20 (2002), 2372–2373] at room temperature starting from the 1:1 molar ratios of ambazone (AMB) and niflumic acid (NIA). The thermal behavior of the obtained compound (AMB·NIA) was investigated by differential scanning calorimetry (DSC) and thermogravimetry (TG), the thermal diffusivity and effusivity parameters were determined by photothermal radiometry (PTR). The structural characterization was performed with X-ray powder diffraction (XRPD) and infrared spectroscopy (FTIR). XRPD data and FTIR spectra demonstrated a new structure for AMB·NIA compound as compared to those of the starting materials.

Keywords: Ambazone, solid forms, X-ray diffraction, FTIR spectroscopy, thermal analysis

1. Introduction

The most active areas of modern solid state chemistry represent the identification and characterization of different crystal forms (polymorphs, solvates, salts and co-crystals) of the same molecule. The investigation of crystal forms impacts on fundamental science because different crystal forms may display a range of different physical–chemical properties, which may affect application and utilization of the solid materials. Traditionally, the solid form selection process was limited to the free drug or pharmaceutically accepted salts. The solid form influences relevant physical–chemical parameters such as solubility, dissolution rate, chemical stability, melting point and hygroscopic parameter which can result in solids with superior properties [5,23]. In recent years, much of the research has been carried out on the preparation of pharmaceutical solid forms. Usually, there are two methods by which solid forms may be prepared: solution-based crystallization and grinding. Mechanical chemical methods [8,9,17,20,22], more commonly and usefully described as grinding, have been employed extensively in the preparation of solid forms. The range of grinding conditions has been extended by the addition of solvents in the “solvent-drop” method [5] and this may represents the introduction of solution conditions on a limited scale to the grinding process. More recently, the use of so-called “solvent-drop grinding” has been developed, in which a small quantity of a solvent is added to the solid substance or mixture prior to grinding.

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Ambazole monohydrate, \([\{4-(2\text{diaminomethylidene)hydrazinyl})\text{phenyl}\} \text{iminothiourea}\), \(\text{C}_8\text{H}_{11}\text{N}_7\text{S} \cdot \text{H}_2\text{O}\) (AMB, see Fig. 1) is one of the oldest antimicrobial chemicals. The studies performed during the 1950–1960 decade have shown the local antibacterial properties when it is administrated at the buccal pharyngeal cavity level, being an efficient antimicrobial drug [11].

The ulterior re-evaluation of the antibacterial AMB properties evidenced an antibacterial activity spectrum similar to that of sulfamides [18]. Recently, the antineoplasm properties of AMB were also demonstrated [1,2,4,10,12,15,16], that accelerated the researches on this substance, without mutagenic effects and unpleasant reactions characteristic to other oncostatic drugs [18].

Niflumic acid, \((2\{-3\text{(trifluoromethyl)phenyl}\text{amino}\text{nicotinic acid}\})\), \(\text{C}_{13}\text{H}_{9}\text{F}_3\text{N}_2\text{O}_2\) (NIA, see Fig. 1), an analgesic and nonsteroidal anti-inflammatory drug [25] that also blocks or modifies the gating of many ion channels, is used in the treatment of rheumatoid arthritis [6].

The aim of this study was to obtain a salt of ambazole with niflumic acid (AMB·NIA) and to characterize its physical–chemical and structural properties using several investigation methods such as: X-ray diffraction, FTIR spectroscopy and several thermal methods (differential scanning calorimetry, thermogravimetry and photothermal radiometry).

2. Experimental procedure

2.1. Materials and preparation

AMB was obtained from Microsin SRL Bucharest, Romania, NIA commercially available was obtained from Alfa Aesar, Germany and these compounds were used without further purification. Solvent-drop grinding (SDG) experiments were performed by placing 255.3 mg AMB with 282.22 mg NIA (1:1 molar ratio) and grinding this mixture in an agate mortar by adding in drops 2 ml ethanol at room temperature, until a dried compound was obtained. The resulting sample was analyzed using X-ray powder diffraction (XRPD), FTIR spectroscopic technique and thermal analysis methods (DSC, TG, PTR).

2.2. X-ray powder diffraction

XRPD patterns were obtained by a Bruker D8 Advance diffractometer, the measurements were performed at 20°C and were controlled by computer. The experimental conditions were: the 2θ range between 2.5 and 45°, CuKα1 radiation (\(\lambda = 1.5406 \text{ Å}\)), (40 kV; 40 mA), Ge 111 monochromator on the diffracted beam. The step scan mode was performed with a step width of 0.01 at a rate of 1 step/s. The samples were mildly pre-ground with a pestle in an agate mortar to make them homogeneous, to control crystals size and to minimize the preferred orientation effects.
2.3. FTIR spectroscopy

The FTIR spectra were recorded with a JASCO 6100 FTIR spectrometer (number of scans, 256; resolution, 4 cm\(^{-1}\); range, 4000–400 cm\(^{-1}\)). The KBr pellets were prepared by mixing 0.8 mg of sample and 150 mg KBr and pressing the mixture into a 13 mm disks at 12 tones pressure. The spectra were analyzed using Spectra Analysis and Origin software.

2.4. Thermal analysis DSC–TG

Differential scanning calorimetry (DSC) was carried out by means of a Shimadzu DSC-60 calorimeter, the sample was heated in the range of 30–350\(^{\circ}\)C with a heating rate of 10\(^{\circ}\)C/min in crimped aluminum sample cell. The purge gas was nitrogen with 60 ml/min flow. For data collection and analysis the Shimadzu TA-WS60 and TA60 2.1 software were employed.

TG curves were obtained with a TGA/SDTA 851e thermobalance. The samples were weighed (Mettler Toledo balance) directly in alumina pans (approximately 5 mg), and scanned between 25 and 400\(^{\circ}\)C with a heating rate of 10\(^{\circ}\)C/min under dynamic nitrogen atmosphere (50 ml/min).

2.5. Photothermal radiometry

PTR technique has been used to measure the dynamic thermal parameters (thermal diffusivity and effusivity) of the pressed powder samples. The PTR technique consists in irradiating the sample with an optical radiation (YAG Laser) and measurement (with a HgCdTe sensor) of the emitted IR radiation.

3. Results and discussion

3.1. X-ray powder diffraction

X-ray powder patterns for AMB, NIA and AMB · NIA are shown in the Fig. 2.

From powder pattern indexing by using Dicvol method [19] it was established that AMB · NIA crystallizes in monoclinic system having following lattice parameters: \(a = 14.61\ \text{Å}, \ b = 5.07\ \text{Å}, \ c = 38.22\ \text{Å}\) and \(\beta = 111.62^\circ\).

The X-ray powder diffraction patterns of the AMB, NIA and AMB with NIA are clearly different, from these one can be see that the powder diffraction pattern of AMB · NIA solid form presents different features comparing with both of starting materials, that means a solid form of AMB with NIA was obtained an ambazone niflumate salt.

3.2. FTIR spectroscopy

The band at \(\sim 3404\ \text{cm}^{-1}\) can be assigned to N–H stretching from primary amine in pure ambazone (see Fig. 3); the FTIR spectrum contains also [13,14] NH stretching band in the 3320–3180 cm\(^{-1}\) spectral range, i.e., 3226 cm\(^{-1}\) [13,24]. The new band which appears at \(\sim 3513\ \text{cm}^{-1}\) in the spectrum of AMB · NIA can be explained by the hydrogen bond breaking between H\(_2\)O molecule and N–H group of AMB after the salt formation. Also it has been shown a change in the NH stretching absorption of amines [13]; it was observed that the sharp strong band at \(\sim 3226\ \text{cm}^{-1}\) due to the NH stretching is shifted to 3254 cm\(^{-1}\) and reduced in intensity. The band at 3147 cm\(^{-1}\) corresponds to the NH vibration
Fig. 2. X-ray powder diffraction patterns for AMB, NIA and AMB·NIA obtained by SDG. (Colors are visible in the online version of the article; http://dx.doi.org/10.3233/SPE-2012-0570.)

Fig. 3. FTIR spectra of AMB, NIA and their salt obtained by SDG method, 4000–2500 cm\(^{-1}\) spectral range. (Colors are visible in the online version of the article; http://dx.doi.org/10.3233/SPE-2012-0570.)

[13] for pure AMB, this band shifted at 3158 cm\(^{-1}\) in the salt spectrum. A new shoulder appeared at \(\sim 2937\) cm\(^{-1}\) probably due to the protonated secondary amine.

The spectrum of the pure NIA contains the NH stretching vibration at 3321 cm\(^{-1}\) and a broad signal at 3090 cm\(^{-1}\) assigned to stretching vibration of CH groups from benzene ring [3]. These signals do not appear in the spectrum of the AMB·NIA.

Primary amine has an absorption band of medium intensity at \(\sim 1613\) cm\(^{-1}\) (see Fig. 4), being shifted to \(\sim 1601\) cm\(^{-1}\) by salt formation [13]. Cleaves and Plyler [7] correlated the spectral bands at 1625–1516 cm\(^{-1}\) with NH deformation vibration. The pure ambazone spectrum contains the secondary amine vibration at 1508 cm\(^{-1}\) which is shifted in AMB·NIA spectrum to 1516 cm\(^{-1}\) and greatly reduced its intensity.
Fig. 4. FTIR spectra of AMB, NIA and their salt obtained by SDG method, 1800–1000 cm\(^{-1}\) spectral range. (Colors are visible in the online version of the article; http://dx.doi.org/10.3233/SPE-2012-0570.)

In the spectrum of the pure NIA a strong absorption band appears at the 1668 cm\(^{-1}\) corresponding to the carboxylic CO stretching vibration \([3]\); this band shifted in the AMB \(\cdot\) NIA spectrum to 1686 cm\(^{-1}\), due to the deprotonation of the carboxyl group for aryl compounds \([19]\) (Fig. 4). This absorption located at \(\sim\)1686 cm\(^{-1}\), can be assigned also to deformation vibration of the protonated secondary amino group \([7]\). The benzene ring stretching vibration at 1614 cm\(^{-1}\) is not shifted and appeared as a shoulder on the spectrum of the AMB \(\cdot\) NIA. At 1428 cm\(^{-1}\) an intense spectral band attributed to carboxylic C=O and C–C stretching and OH rocking vibrations appears, it decreases in intensity, probably due to the carboxyl involving in intermolecular interactions. The shift of the vibrational mode at \(\sim\)1448–1454 cm\(^{-1}\) can be due to the influence of the deprotonated carboxyl group on the pyridine ring.

3.3. Thermal analysis DSC–TG

The DSC thermograms of the pure AMB, NIA and of the AMB \(\cdot\) NIA compound obtained by SDG are presented in Fig. 5. The curve for the pure AMB revealed a broad endothermic signal from 105 to 143\(^\circ\)C, with a maximum at 125\(^\circ\)C and \(\Delta H = 36\) kJ/mol, that corresponds to the water molecules loss of the AMB monohydrate structure, followed by a sharp exothermic signal at 204.52\(^\circ\)C, \(\Delta H = 75\) kJ/mol due to the melting with decomposition of AMB.

The thermogram of the pure NIA revealed a sharp endothermic signal with maximum at 204.75\(^\circ\)C, and \(\Delta H = 30.54\) kJ/mol, that corresponds the melting of the NIA \([21]\).

The DSC curve of AMB \(\cdot\) NIA presents a broad exothermic signal between 185 and 198\(^\circ\)C, with a maximum at 188.5\(^\circ\)C and \(\Delta H = 69.43\) kJ/mol, corresponding to the melt with start of decomposition of the sample.

The thermal behaviour characteristics of AMB, NIA and AMB \(\cdot\) NIA obtained by SDG procedure, have been investigated by thermogravimetry (TG) and are shown in Figs 6–8.

The TG curve for the pure AMB shows the 7.24% mass loss between 97–150\(^\circ\)C corresponding to the evaporation of water. In the 184–208\(^\circ\)C temperature range significant mass loss appeared representing 25.41% from mass of AMB due to the starting of decomposition. The next mass losses, 6.91 and
Fig. 5. DSC thermograms of pure AMB, NIA and AMB·NIA, obtained by SDG. (Colors are visible in the online version of the article; http://dx.doi.org/10.3233/SPE-2012-0570.)

Fig. 6. The TG thermogram of AMB. (Colors are visible in the online version of the article; http://dx.doi.org/10.3233/SPE-2012-0570.)
Fig. 7. The TG thermogram of NIA. (Colors are visible in the online version of the article; http://dx.doi.org/10.3233/SPE-2012-0570.)

Fig. 8. TG thermogram of AMB·NIA, obtained by SDG. (Colors are visible in the online version of the article; http://dx.doi.org/10.3233/SPE-2012-0570.)
3.17% in the 216–288°C temperature range, respectively 294–386°C correspond to the elimination of the volatile components resulted from decomposition. The temperature ranges where mass losses occur are in agreement with DSC results.

The trace of thermogravimetric behaviour of the NIA shows an 87.04% mass loss in the 182–340°C temperature range, corresponding on the DSC curve to two endothermic peaks: a sharp one due to the melting, and a broader one due to decomposition of the NIA.

The thermogravimetric curve of the AMB·NIA shows the first mass loss, 7.82%, in the range 168–192°C, followed by 4.09% mass loss in the range 213–224°C. Last mass loss is the most significant, 31.26% and occurs in the 238–362°C temperature range.

The TG data are in good agreement with the DSC results and support the new obtained compound formation.

3.4. Photothermal radiometry

The PTR signal is a function of samples’ thermal parameters and chopping frequency of radiation, see Fig. 9. The results (room temperature values of thermal diffusivity and effusivity) were obtained through a fitting procedure with the two mentioned thermal parameters as fitting parameters.

The PTR investigations leaded to values of dynamic thermal parameters out of the range of the values of these parameters for the starting materials. Such behavior can be ascribed to two phenomena: formation of a new compound and/or packing effects. Due to the fact that the pressure used to prepare the pellets was the same, we choose as more possible the first alternative. In fact, the formation of a new compound is also supported by the previous investigations.

4. Conclusions

- The FTIR spectra indicate the appearance of vibrations bands characteristic to NH$_2^+$ and NH$_3^+$ groups corresponding to the protonation of the secondary and primary amine. Also new vibration
bands appear. Consequently, a salt of ambazone and niflumic acid formation was confirmed.

- Thermal behaviour of pure substances and of AMB.NIA compounds are different, the distinct melting point and the decomposition mode certifies the new compound formation.
- New solid form that was prepared with SDG method combining the ambazone and niflumic acid. Based on X-ray powder diffraction the lattice parameters and the space group for the new compound were determined.
- The results obtained by photothermal radiometry can be considered also a support for the formation of a new compound.

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References


