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# Insight into inclusion complexation of indomethacin nicotinamide cocrystals

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7 Abstract The objective of this research was to investi-8 gate the feasibility of the interaction between 9 indomethacin-nicotinamide cocrystals with β-cyclodextrin 10 and hydroxypropyl-\beta-cyclodextrin in the solid-state. The 11 study has emphasized on the possibility of inclusion 12 complex formation and its effect on the dissolution per-13 formance of the cocrystals. The solid systems in the molar 14 ratio of 1:1 of the host and guest molecules were prepared 15 by co-grinding and co-evaporation methods and compared with their physical mixtures. Furthermore, the molecular 16 17 behaviors of the cocrystals in all prepared samples were 18 thoroughly characterized by powder X-ray diffraction, 19 differential scanning calorimetry, Fourier-transform infra-20 red spectroscopy, scanning electron microscopy and 21 in vitro dissolution performance. The results of these 22 studies indicated that complexes prepared by the co-evap-23 oration method with hydroxypropyl-B-cyclodextrin have 24 shown complete inclusion of the cocrystals into the 25 cyclodextrin cavity and a partial inclusion with β-cyclodextrin. Moreover, a significant (p < 0.05; ANOVA/ 26 27 Tukey) higher in vitro dissolution was achieved in co-28 evaporate complex prepared with hydroxypropyl-\beta-cy-29 clodextrin compared to that prepared with β-cyclodextrin, 30 indomethacin-nicotinamide cocrystals and indomethacin 32 itself.

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## Introduction

In the development and manufacture of pharmaceutical 37 products, improving the physicochemical properties of 38 drugs is often essential but can be challenging. Because 39 these improvements can often be achieved by making new 40 solid forms of the drug without altering its chemical 41 structure. Recent research has focused on identifying 42 polymorphs, hydrates, solvates, salts and, more recently, 43 cocrystals of drugs [1]. Cocrystals are composed of mul-44 tiple molecular components, including the drug and a 45 benign, non-toxic 'coformer' molecule. The design, for-46 mation and understanding of the physicochemical 47 properties of cocrystals have received considerable atten-48 49 tion [2-5]. Indeed, pharmaceutical cocrystals are an attractive alternative solid form with the potential to fine 50 tune the physicochemical properties of drugs [6]. 51

Inclusion or host–guest complexes are supramolecular 52 systems where one chemical compound (the *host*) has a 53 cavity, in which molecules of a second compound (the *guest*) are located [7, 8]. The study of non-covalent forces 55 involved in the formation of host–guest complexes is of 56 paramount importance for the design of synthetic inclusion 57 compounds of active pharmaceutical ingredients (APIs). 58

β-Cyclodextrin (β-CD) is a cyclic oligosaccharide consisting of seven glucose units linked by α (1  $\rightarrow$  4) bonds, 60 resulting in a hollow truncated cone shape [9]. In water 61 they have a hydrophilic outer surface and a hydrophobic 62 central cavity able to include a wide variety of lipophilic 63 guest molecules, with suitable polarity and dimensions, 64

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	Article No. : 594	LE	TYPESET
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65 without the formation of any covalent bond [10]. Complexation with  $\beta$ -CD has found extensive applications in 66 67 pharmaceutical technology to enhance the aqueous solu-68 bility, dissolution rate, bioavailability, and stability of 69 poorly water soluble drugs [11, 12]. Unsubstituted  $\beta$ -CD 70 has poor water solubility (16 mg mL<sup>-1</sup> at 25 °C), whereas random substitution of the hydroxyl groups with alkyl or 71 72 hydroxyalkyl groups is able to increase solubility. There-73 fore, several synthetically modified  $\beta$ -CDs were used as 74 multifunctional drug carriers in parenteral formulations, 75 such as hydroxypropyl β-cyclodextrin (HP-β-CD) [13].

Indomethacin (IND) (1-(p-chlorobenzovl)-5-methoxy-2methylindole-3-acetic acid) (Scheme 1a) is a non-steroidal anti-inflammatory drug [14]. It is known to form cocrystals a non-toxic coformer. nicotinamide with (NIC) (Scheme 1b) [15, 16].

81 To the best of our knowledge, indomethacin-nicoti-82 namide cocrystals (INDNIC) complexation with different 83 cylcodextrins has not been studied before. There is only an 84 available report in the literature about formulation devel-85 carbamazepine-nicotinamide opment of cocrystals 86 complexed with y-cyclodextrin using supercritical fluid 87 process [17]. In spite of  $\gamma$ -cyclodextrin is a highly water 88 soluble derivative of cyclodextrins but it is not economic 89 for the pharmaceutical industry. Additionally, the super-90 critical fluid technology is a complex, expensive and time 91 consuming process. So, the objective of the present work 92 was to study the feasibility of interaction of INDNIC with 93 other types of cyclodextrins e.g., β-CD and HP- β-CD in 94 the solid-state. More economic, simple and fast approaches 95 have been applied to achieve inclusion complex formation 96 which are co-grinding and co-evaporation methods. The 97 study was further aimed to characterize the prepared 98 complexes by powder X-ray diffraction (PXRD), differ-99 ential scanning calorimetry (DSC), Fourier-transform 100 infrared (FTIR) spectroscopy and scanning electron 101 microscopy (SEM). An in vitro dissolution study was also 102 conducted to show the effect of cocrystals inclusion com-103 plexation on the dissolution profile of IND after its

#### Scheme 1 The chemical structures of a IND and b NIC

(a)

COOH CH.

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eriod of 6 h.	The solutions	were evaporate	ed at 40 °C i

(y form), meotimannue, emyr acetate,	100
were purchased from Sigma-Aldrich (Cairo,	109
B-cyclodextrin and HP-β-cyclodextrin were pur-	110
com Fluka Chemie (Switzerland). All chemicals	111
ents were of analytical grade and used as received.	112
tion of indomethacin cocrystals	113
acin cocrystals were prepared by slurry crystal-	114

Indomethacin cocrystals were prepared by slurry crystal 114 lization [18]. A total of approximately 3.578 g of IND and 115 1.221 g of NIC in a (1:1 molar ratio) was magnetically 116 stirred in 10 ml of ethyl acetate for 5 days at room tem-117 perature. Solids were filtered, dried and analyzed by 118 PXRD, DSC and FTIR. 119

#### Preparation of the physical mixtures

Physical mixtures of INDNIC with β-CD and HP-β-CD in 121 (1:1 molar ratio) were prepared by simple blending in a 122 glass mortar for 5 min and stored under vacuum in desic-123 cator over calcium chloride for five consecutive days. 124

#### 125 **Preparation of the inclusion complexes**

Co-grinding method

The ground mixtures of INDNIC with each of β-CD and 127 HP-β-CD in (1:1 molar ratio) were prepared using the 128 vibrational uniball mill (VEB leuchtenbau-KM1, Ger-129 many) for 15 min. The ground mixtures were sieved to 130 131 obtain a particle size range of 125-250 µm, and then stored 132 in a desiccator over calcium chloride for five consecutive days. 133

#### Co-evaporation method

Co-evaporates of INDNIC with each of β-CD and HP-β-135 136 CD were prepared by solvent evaporation method in the same molar ratio (1:1) as previously reported [19]. Briefly, 137 IND-NIC was dissolved in a sufficient volume of methanol. 138 Both β-CD and HP-β-CD were dissolved at 40 °C in dis-139 tilled water. INDNIC and respective CD solutions were 140 mixed together with constant stirring and kept at 40 °C for 141 an hour, then gradually cooled to room temperature over a 142 143 under р

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complexation	with	cyclodextrins	compared	with	uncom-	104
plexed (free)	INDN	IC cocrystals	and pure IN	ND.	AQ	105

#### **Experimental**

#### **Materials**

Indomethacin 108 (~ form) nicotinamide othyl acetate methanol Egypt). chased fr and solve

#### Preparat

Journal : Large 10847	Dispatch : 11-1-2016	Pages : 10
Article No. : 594	LE	TYPESET
MS Code : JIPH-D-15-00175	<b>✓</b> <sub>CP</sub>	✓ <sub>DISK</sub>

NH,

(b)

vacuum till a constant weight was achieved. The collected
powders were sieved to obtain a particle range of 125–
250 μm and stored under vacuum in a desiccator over
calcium chloride for five consecutive days.

#### 148 Instrumentation

#### 149 Powder X-ray diffractometry (PXRD)

150 The powder X-ray diffraction patterns of the solid samples 151 were recorded using a Philips 1710 powder diffractometer 152 with Cu Ka radiation (1.54056 Å). A Cu target tube 153 operated at a voltage of 40 kV and a current of 40 mA and 154 a single crystal graphite monochromator were employed. A 155 scanning speed of 0.6°/min and a wide angel diffraction of 156  $4^{\circ} < 2\theta < 60^{\circ}$  were applied. Standard polycrystalline silicon powder was used to calibrate the instrument. 157

### 158 Differential scanning calorimetry (DSC)

159 DSC thermograms were obtained by using a Shimadzu 160 DSC-50 (Japan). Samples of about 5 mg were placed in aluminum pans of 50 µL capacity & 0.1 mm thickness, 161 162 press-sealed with aluminum cover of 0.1 mm thickness. An 163 empty pan sealed in the same way was used as a reference. The thermograms were recorded by heating the samples 164 from 30 to 250 °C at a rate of 10 °C min<sup>-1</sup>, under nitrogen 165 flow of 40 ml min<sup>-1</sup>. Indium was used as a standard for 166 167 calibrating the temperature. Reproducibility was checked 168 by running the samples in triplicate, the standard deviations 169 calculated were found negligible.

### 170 Infrared spectroscopy

171 FT-infrared spectra were collected in triplicate using a 172 Nicolet 6700 FTIR Advanced Gold Spectrometer in the 173 diffuse reflectance mode with potassium bromide as a 174 diluent (1:200), using <10 mg of the solid samples. The 175 spectra were recorded in the range of 400–4000  $\text{cm}^{-1}$  at  $2 \text{ cm}^{-1}$  spectral resolution with the accumulation of 256 176 spectral scans. The instrument was controlled with OMNIC 177 178 8 software. Triplicate spectra were averaged to obtain one 179 spectrum for each sample. All FTIR spectra were exported 180 to the Galactic\* SPC format using GRAMS AI (Version 181 8.0, Thermo Electron Corp. Waltham, MA, USA) with no 182 further processing.

### 183 Scanning electron microscopy

Particles were visualized by scanning electron microscopy(SEM; Jeol 5400 LV, Japan).

Particles were mounted on aluminum stubs (pin stubs,<br/>13 mm), layered with a sticky conductive carbon tab and<br/>coated in gold (10–15 nm) using an EmiTech K 550X Gold186<br/>188Sputter Coater, 25 mA for 3 min.189

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#### **In-vitro dissolution**

Dissolution experiments were carried out in triplicate with 191 USP apparatus II dissolution using paddle at a rotation 192 speed of 100 rpm. Powdered samples of each preparation 193 equivalent to 25 mg of IND were added to the dissolution 194 medium, 500 mL of phosphate buffer, pH 7.4, kept at 195  $37 \pm 0.5$  °C. At appropriate time intervals, 5 mL of the 196 solution were withdrawn using cotton plug from the dis-197 solution medium and replaced with an equal volume of the 198 fresh dissolution medium equilibrated at 37 °C. Then ali-199 quots were injected into the HPLC system. The HPLC 200 system, Knauer, D-14163, Germany, consists of HPLC 201 pump, UV- detector, and integration interface box. Chro-202 matographic separation was carried out using Kromasil 203 C-18 column (250  $\times$  4.60 mm, particle size: 20  $\mu$ m). The 204 detection wavelength, 377 nm, was determined by scan-205 ning the maximum absorbance wavelength of IND in the 206 mobile phase (Methanol: distilled water) using an UV-Vis 207 spectrophotometer (Jenway, Model 6305, UK). 208

#### Statistical analysis

All statistical analysis was performed using One-way 210 analysis of variance (ANOVA) with the Tukey's multiple 211 comparisons was employed for comparing the preparations with each other (Minitab<sup>®</sup>16 Statistical Software). Statistically significant differences were assumed when p < 0.05. 214 All values are expressed as their mean  $\pm$  standard 215 deviation. 216

### Results and discussions 217

The solid-phase identity and purity of INDNIC cocrystals218were verified using PXRD, DSC and FTIR prior to the219inclusion complexation. The PXRD pattern and DSC220melting curve and FTIR spectrum (Figs. 1a, 3a, 5a, b,221respectively) of the INDNIC agreed well with the previously published data [16].223

#### **PXRD** study

The PXRD patterns of INDNIC,  $\beta$ -CD and their physical, 225 co-ground and co-evaporate mixtures were presented in Fig. 1. Those with HP- $\beta$ -CD, in turn, were presented in Fig. 2. The diffractograms of both INDNIC cocrystals and  $\beta$ -CD exhibited series of intense peaks, which are 229

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Fig. 1 The PXRD patterns of a INDNIC, b  $\beta$ -CD, c physical mixture, (D) co-ground mixture and (E) co-evaporate product. C, D and E are with  $\beta$ -CD



Fig. 2 The PXRD patterns of a INDNIC, b HP- $\beta$ -CD, c physical mixture, (D) co-ground mixture and (E) co-evaporate product. C, D and E are with HP- $\beta$ -CD

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indicative of their crystallinity. Further, the diffractograms 230 231 of their physical mixtures, co-ground mixtures still showed the characteristic peaks of both INDNIC and β-CD how-232 ever, some peaks were reduced in their intensities (Fig. 1; 233 traces C-E). Co-evaporate of INDNIC with β-CD showed 234 235 further reduction in other crystalline peaks of INDNIC especially the most intense diffraction peaks at 236  $(2\theta) = 15.35^{\circ}, 23.89^{\circ}$  and  $27.22^{\circ}$  (Fig. 1; trace D). How-237 ever, the diffraction peak of INDNIC at  $(2\theta) = 12.94^{\circ}$  was 238 239 disappeared with the recording of a new diffraction peak at  $(2\theta) = 12.51^{\circ}$  for their co-evaporate product. This may 240 indicate that some sort of interaction has been occurred in 241 case of the co-evaporate system which has led to formation 242 of compound with lower crystallinity as compared with 243 244 INDNIC cocrystals.

In case of INDNIC cocrystals with HP- $\beta$ -CD, in turn, a 245 significant reduction in the diffraction patterns of both the 246 physical and co-ground mixtures of INDNIC/HP-B-CD has 247 been observed (Fig. 2, traces C, D). On the other hand, the 248 co-evaporate mixture of INDNIC and HP-\beta-CD showed 249 new very small peaks at  $(2\theta)$  values of 8.44°, 11.80°, 250 14.44°, 25.36° and 28.36° as compared to their physical 251 and co-ground mixtures (Fig. 2, trace E). The disappear-252 ance of the crystalline peaks of INDNIC and recording of 253 other new peaks could indicate the formation of structure 254 with very low crystallinity [20]. On other words, co-255 evaporate system with HP- $\beta$ -CD could have the potential to 256 257 form an inclusion complex with INDNIC fitted inside the HP- $\beta$ -CD cavity and such complex appeared to be partially 258 amorphous. 259

The inclusion complexes of INDNIC with either  $\beta$ -CD 260 or HP- $\beta$ -CD were polycrystalline and of low crystallinity, 261 respectively. Interestingly, Jambhekar et al. [21] concluded that, the inclusion complexes of IND itself with either  $\beta$ -CD or HP- $\beta$ -CD were polycrystalline and amorphous, 264 respectively. 265

#### DSC study

The DSC thermograms of INDNIC cocrystals,  $\beta$ -CD and 267 their physical mixtures, co-ground and co-evaporate solid 268 systems were presented in Fig. 3. Similarly, those with HP- $\beta$ -CD were presented in Fig. 4. 270

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The peaks corresponding to the evaporation of water 271 vapors from both  $\beta$ -CD and HP- $\beta$ -CD appeared at 85 and 272 60 °C; respectively (Figs. 3, 4; trace B) [22, 23]. 273

The physical and co-ground mixtures of INDNIC with  $\beta$ -CD showed a reduced intensity endothermic peaks at 127.73 and 127.21 °C; respectively (Fig. 3; traces C and D) compared to that at 128.50 °C for INDNIC (Fig. 3; trace A). The presence of such endothermic peak in both the physical and co-ground mixtures still reflect the presence of free INDNIC in the prepared solid systems. Also, it was



**Fig. 3** The DSC thermograms of *A* INDNIC, *B* β-CD, *C* physical mixture, *D* co-ground mixture and *E* co-evaporate product. *C*–*E* are with β-CD



**Fig. 4** The DSC thermograms of *A* INDNIC, *B* HP- $\beta$ -CD, *C* physical mixture, *D* co-ground mixture and *E* co-evaporate product. *C*–*E* are with HP- $\beta$ -CD

281 noted that the water evaporation endothermic peak was 282 shifted from 80 to 60 °C in case of the physical mixture 283 with β-CD (Fig. 3, trace C).

Similarly, it was found that the melting endotherm of INDNIC physical and co-ground mixtures with HP- $\beta$ -CD appeared broader and shifted from 128.50 to 127.50 and 120.26 °C; respectively (Fig. 4, traces C and D). Further, the broadening and reduction of that peak could be attributed to the incomplete complex formation via simple physical and co-ground methods.

291 However, in case of co-evaporate mixtures with  $\beta$ -CD, a 292 small, highly broad endothermic peak was observed at 293 135.74 °C concomitantly with broadening of the water 294 evaporation peak (Fig. 3; trace E). This may indicate that 295 some sort of inclusion into the  $\beta$ -CD cavity was occurred 296 and there was still some INDNIC cocrystals cannot fit 297 completely inside the  $\beta$ -CD cavity which is responsible for 298 this small endothermic broad peak [24].

299 On the other hand, the melting endotherm of co-evap-300 orate with HP- $\beta$ -CD was completely disappeared which 301 indicated that the new solid compound formed has an 302 amorphous structure (Fig. 4; trace E). These findings come 303 in accordance with the above mentioned PXRD results. 304 Moreover, the complete disappearance of the water evaporation peak of HP- $\beta$ -CD in case of co-evaporate 305 mixture with INDNIC could indicate that the cocrystal has penetrated into the HP- $\beta$ -CD cavity and replaced the water 307 molecules. 308

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#### Solid-state FTIR spectroscopic investigation

FTIR spectroscopy was used to assess the interaction310between the cocrystal and the two studied cyclodextrins in311the solid state. The chemical interaction between the host312and the guest molecules often leads to identifiable changes313in the FTIR spectra of complexes. However, some of the314changes are very subtle requiring careful interpretation of315the spectra.316

The FTIR spectra of INDNIC,  $\beta$ -CD, HP- $\beta$ -CD and their 317 respective physical and cog-round mixtures and co-evap-318 orate products exhibited a number of differences in both 319 the fingerprint and high wavenumber regions (Figs. 5, 6). 320 The v (OH) regions in the FTIR spectra appear as very 321 broad bands in the range of 3400-2500 cm<sup>-1</sup> which are 322 superimposed on the v (CH) regions. Both the broad nature 323 and the position of these FTIR bands are characteristic of 324 hydrogen-bonded OH groups [25]. The differences in the 325 shapes of these bands suggest that there may be associated 326 327 variations in hydrogen bonding.

By close inspection of the FTIR spectra of INDNIC,  $\beta$ -328 CD and their physical and ground mixtures and their co-329 evaporate products in the wavenumber range (3400-330  $2500 \text{ cm}^{-1}$ ), it could be noted that the characteristic sharp v 331 (OH) bands at 3413 and 3318 cm<sup>-1</sup> and v (NH<sub>2</sub>) bands at 332 3285 and 3218 cm<sup>-1</sup> of INDNIC were recorded as broader 333 bands in the FTIR spectra of their physical and ground 334 mixtures with reduced intensities but not in the spectrum of 335 their co-evaporate products except the v (OH) band at 336  $3413 \text{ cm}^{-1}$  (Fig. 5a). That may indicate incomplete inclu-337 sion of the cocrystal in the  $\beta$ -CD cavity by physical mixing 338 and co-grinding approaches. The same also may apply to 339 the co-evaporation approach as indicated by the recording 340 of the above weak v (OH) band at 3413 cm<sup>-1</sup> in the FTIR 341 spectrum of the co-evaporate product (Fig. 5a, trace E). 342 Based on the above observations, partial inclusion is more 343 efficient by the co-evaporation approach as compared to 344 the two other methods which supports the previous findings 345 from the PXRD and DSC data. 346

In case of HP- $\beta$ -CD only the sharp  $\upsilon$  (OH) and  $\upsilon$  (NH<sub>2</sub>) 347 bands at 3413 and 3218 cm<sup>-1</sup>, respectively were recorded 348 as broader bands in the FTIR spectra of their physical and 349 ground mixtures with reduced intensities but not in the 350 spectrum of their co-evaporate products (Fig. 6a). That 351 may indicate more efficient partial inclusion of the 352 cocrystal in the HP-\beta-CD cavity by physical mixing and 353 co-grinding approaches. However, in case of the co-354



	Journal : Large 10847	Dispatch : 11-1-2016	Pages : 10	
	Article No. : 594	LE	TYPESET	
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**Fig. 5 a** The FTIR spectra in the wavenumber range (2500– 4000 cm<sup>-1</sup>) of *A* INDNIC, *B* β-CD, *C* physical mixture, *D* coground mixture and *E* coevaporate product. *C*–*E* are with β-CD. **b** The FTIR spectra in the wavenumber range (1550– 1750 cm<sup>-1</sup>) of *A* INDNIC, *B* β-CD, *C* physical mixture, *D* coground mixture and *E* coevaporate product. *C*–*E* are with β-CD



evaporation approach, the inclusion complexation appears to be complete (*cf*. that with  $\beta$ -CD).

By careful inspection of the FTIR spectra of INDNIC,  $\beta$ -357 358 CD, and their physical and ground mixtures and their coevaporate products in the wavenumber range (1550-359 1750 cm<sup>-1</sup>) region, in turn, rather interesting findings were 360 apparent (Fig. 5b). The characteristic v(C=O) bands of 361 INDNIC at 1661 and 1680 cm<sup>-1</sup> were recorded in the FTIR 362 spectra of their physical and ground mixtures. However in 363 the FTIR spectrum of their co-evaporate product they were 364 shifted to 1680 and 1691 cm<sup>-1</sup>, respectively with the 365 recording of a new v(C=O) band at 1735 cm<sup>-1</sup>. These 366 367 significant shifts together with the recording of a new v(C=O) band suggest the formation of new hydrogen 368 bonding between the cocrystal and the OH groups in the 369 370 interior cavity of β-CD during their more efficient partial

inclusion. The same was also noted with HP- $\beta$ -CD 371 (Fig. 6b). These shifts together with the recording of a new  $\nu$ (C=O) band at 1735 cm<sup>-1</sup> further confirm the complete inclusion of the cocrystal with the HP- $\beta$ -CD using the coevaporation approach. These findings further confirm the 375 above PXRD and DSC observations. 376

### SEM analysis

The SEM images of INDNIC, HP- $\beta$ -CD and their coevaporate and co-ground products,  $\beta$ -CD and its co-evaporate product with INDNIC were presented in Fig. 7 (traces A, B, C D, E and F, respectively). 381

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Striking differences in the morphology of the particles382were observed upon careful inspection of the SEM images.383INDNIC particles were appeared as rectangular crystals384

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	Article No. : 594	LE	TYPESET
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**Fig. 6 a** The FTIR spectra in the wavenumber range (2500– 4000 cm<sup>-1</sup>) of *A* INDNIC, *B* HP- $\beta$ -CD, *C* physical mixture, *D* ground mixture and *E* coevaporate product. *C*–*E* are with HP- $\beta$ -CD. **b** The FTIR spectra in the wavenumber range (1550–1750 cm<sup>-1</sup>) of *A* INDNIC, *B* HP- $\beta$ -CD, *C* physical mixture, *D* ground mixture and *E* co-evaporate product. *C*–*E* are with HP- $\beta$ -CD



385 while HP-\beta-CD was presented as aggregated spherical 386 particles (Fig. 7, traces A and B). By comparing the morphology of the co-evaporate and co-ground products of 387 388 INDNIC with HP- $\beta$ -CD, it could be noted that, the co-389 evaporate product appears as large irregular aggregates of 390 thick lumps which differ in morphology than both INDNIC and HP- $\beta$ -CD (Fig. 7, trace C), which could indicate the 391 loss of the crystalline shape of INDNIC and formation of 392 393 another complex structure with a new morphology. The co-394 ground product, in turn, shows the loss of the spherical 395 appearance of HP-\beta-CD particles with the appearance of 396 the cocrystals (Fig. 7, trace D). The loss of the spherical 397 characters could be possibly attributed to the grinding 398 process. These observations are in agreement with our above findings from PXRD, DSC and FTIR studies which 399further confirm the complete inclusion of the cocrystal with the HP- $\beta$ -CD using the co-evaporation approach. 401

To further confirm our findings, the morphology of  $\beta$ -CD particles and their co-evaporate product with INDNIC were also compared. Interestingly, the cocrystals were appeared as a huge network of aggregated filaments on the surface of  $\beta$ -CD particles (Fig. 7, trace E). Such morphology may indicate the incomplete inclusion of the cocrystals using the co-evaporation approach. 402 403 404 405 406 407 408

The variations in the particles morphology among the<br/>above products despite similar processing conditions could<br/>be explained by considerable differences in the crystal-<br/>lization kinetics or crystal lattices.409<br/>410<br/>411

Fig. 7 The SEM images of a INDNIC, b HP-β-CD, c coevaporate product of INDNIC with HP- $\beta$ -CD, **d** co-ground product of INDNIC with HP-β-CD,  $\mathbf{e} \beta$ -CD and  $\mathbf{f}$  co-evaporate product of INDNIC with β-CD





Fig. 8 The dissolution profiles of IND, INDNIC and the co-evaporate mixtures of INDNIC with β-CD and HP-β-CD

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Figure 8 shows the dissolution profiles of IND, INDNIC 414 and co-evaporate mixtures of INDNIC with B-CD and HP-415  $\beta$ -CD in pH 6.8. It was found that the percentages of IND 416 powder dissolved after 15 and 120 min were 19.32  $\%\pm6.5$ 417 and 62.55 %  $\pm$  4.9; respectively. INDNIC cocrystals and 418 co-evaporate mixtures of INDNIC with β-CD and HP-β-419 CD showed a significantly (p < 0.05; ANOVA/Tukey) 420 higher increase in the dissolution rate of IND 421 61.20 %  $\pm$  4.5, 62.96 %  $\pm$  1.8 and 75.16 %  $\pm$  2.7 after 422 15 min; respectively compared with untreated IND powder, 423 19.32 %  $\pm$  6.5, after the same time interval. Further, the 424 co-evaporate mixture of INDNIC with HP-\beta-CD gives a 425 significantly (p < 0.05; ANOVA/Tukey) higher increase in 426

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	Article No. : 594	LE	TYPESET
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427 the dissolution rate of IND 75.16  $\% \pm 2.7$  and 428 86.76 %  $\pm$  0.7 after 15 and 120 min; respectively com-429 pared to INDNIC cocrystals which showed  $61.20 \% \pm 4.5$ 430 and 65.77  $\% \pm 2.8$  IND percentage released after the same 431 time. INDNIC cocrystals significantly increased the dis-432 solution rate of IND compared with the untreated IND 433 powder possibly as previously reported with indomethacin-434 saccharin cocrystals [18]. Furthermore, the effect of HP-B-435 CD on the dissolution rate of certain drugs was higher than 436 that of  $\beta$ -CD due to its higher water solubility and the 437 bigger capacity of its cavity [19, 26, 27]. It is worth to note 438 that, the co-evaporate mixture with HP-B-CD produces a 439 significant (p < 0.05; ANOVA/Tukey) rapid dissolution 440 after 15 and 120 min as compared to that with β-CD co-441 evaporate system. That may be attributed to local solubil-442 isation action operating in the micro-environment or the 443 hydrodynamic layer surrounding the cocrystals particles in 444 the early stages of the dissolution process. Additionally, 445 HP-B-CD dissolves in a short time compared with B-CD 446 thus improves the wettability of the cocrystals and hence 447 their dissolution [23]. Also, one could not neglect that, the 448 formation of the complete inclusion complex with a new 449 partially amorphous structure which could enhance the 450 dissolution process compared to the co-evaporate system 451 with  $\beta$ -CD and this is in agreement with PXRD, DSC and 452 FTIR studies.

#### 453 Conclusions

454 Partial and complete inclusion complexes of INDNIC 455 cocrystals with β-CD and HP-β-CD were prepared suc-456 cessfully by the co-evaporation method in a molar ratio of 457 1:1 of the guest to host molecules. This was confirmed by 458 various analytical techniques. The co-evaporate mixture of 459 INDNIC with HP-β-CD showed new morphological 460 structure as observed from SEM Study. The prepared 461 inclusion complexes enhanced the dissolution rate of IND 462 significantly as compared with the untreated IND and 463 INDNIC cocrystals. Furthermore, the highest improvement 464 in IND in vitro dissolution was observed in the inclusion complex prepared with HP-β-CD. 465

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