

1 Preparation and evaluation of solid dispersions of Ibuprofen using 2 Glucosamine HCl as a carrier

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12 **Key words:** Solid dispersions, Ibuprofen, Glucosamine HCl, Solvent Evaporation, Solubility,
13 Dissolution rate.

14

15 Abstract

16 The present work reports the study of preparation and evaluation of solid dispersions of
17 Ibuprofen. As Ibuprofen is sparingly water-soluble drug and has low bioavailability, so to
18 enhance its solubility and improve bioavailability solid dispersions with different drug to carrier
19 ratios (1:1, 1:2 and 1:3) were prepared, as solid dispersion is the most effective method for
20 enhancing the solubility and improving the bioavailability of poorly or sparingly water- soluble
21 drugs. In this study Glucosamine HCl was used as a potential hydrophilic carrier to improve the
22 solubility, dissolution rate and bioavailability of poorly water-soluble drug, Ibuprofen from
23 physical mixtures and solid dispersions. Solid dispersions with different drug to carrier ratios
24 were prepared, using solvent evaporation method. Physical mixtures of Ibuprofen and
25 Glucosamine HCl were also prepared for comparison. All solid dispersions of Ibuprofen and
26 Glucosamine showed considerably higher dissolution rate than corresponding physical mixtures
27 and pure Ibuprofen. Different techniques such DSC, FT-IR, XRD and SEM were used to study
28 the properties of pure Ibuprofen, solid dispersions and physical mixtures. These results
29 confirmed that Glucosamine HCl can increase the solubility and dissolution rate of poorly water-
30 soluble drug, Ibuprofen.

31

32 Introduction

33 Nearly one third of drugs in development are water insoluble which are mostly failed during trail
34 phase of development because of underprivileged pharmacokinetics [1]. Poorly water-soluble
35 drugs belong to class II and Class IV of Biopharmaceutical Classification System (BCS). Poor
36 water solubility of a drug leads to low dissolution, slow absorption, variable bioavailability and
37 gastrointestinal toxicity [1]. Formulation of poorly soluble drugs for orally drug delivery now
38 represent one of the most interesting challenges to formulation scientist in the pharmaceutical
39 industries and for formulation containing poorly soluble drugs, dissolutions is the rate limiting
40 step in the process of drug absorption [2].

41 Ibuprofen [(±)-2-(4'-isobutylphenyl) propionic acid] a phenyl propionic acid derivative, is
42 widely accepted as one of the best tolerated non-steroidal anti-inflammatory and common
43 analgesic drugs available for the treatment of rheumatoid arthritis, osteoarthritis, and mild to
44 moderate pain [3]. The drug has been classified as class II drug as per the Biopharmaceutical
45 Classification System (BCS) having low solubility and high permeability through stomach as it
46 remains 99.9% unionized in stomach, so because of its solubility limitation and fast emptying
47 time from stomach to intestine (30min to 2 hrs) can not enter into systemic circulation. After this
48 time it goes to small intestine where it is solubilized but can not permeate through its membrane
49 because of its pH dependent solubility and permeability [2]. Thus solubility and dissolution
50 become the rate limiting steps for absorption. Drugs with low dissolution rates generally show
51 erratic and incomplete absorption leading to low bioavailability when administered orally. To
52 enhance solubility and improve dissolution rate of Ibuprofen is challenging and rational because
53 its serum concentration and therapeutic effects are correlated, rapid Ibuprofen absorption is
54 prerequisite for the quick onset of action.

55 Several techniques have been reported to improve the solubility and dissolution rates of poorly
56 water soluble drugs which include solid dispersions, micronization, lipid based formulations, melt
57 granulation, direct compaction, solvent evaporation, adsorption, coprecipitation, ordered mixing,
58 inclusion complexation, liquisolid compacts, steam aided granulation, solubilization in
59 surfactant systems, formation of water soluble complex and use of prodrugs [1,4]. Among all
60 these methods and techniques micronization and liquidsolid are most commonly used for class II
61 drugs but these techniques are having some disadvantages as the micronized particles are stick
62 together and make larger agglomerates, consequently leads to a reduction in effective surface
63 area for dissolution [4]. The most effective method for improving dissolution rate is the use of
64 solid dispersion technique. This technique has been widely used for poorly soluble drugs such as
65 nimsulid, ketoprofen, tenoxicam, nifedipine and nimodipine [5-9].

66 Solid dispersion is defined the dispersion of one or more active ingredients in an inert carrier or
67 matrix in a solid state prepared by melting, dissolution in solvent or melting solvent method [10,
68 11].

69 In solid dispersion carrier plays an important role in improving solubility and dissolution rate.
70 Different polymers, superdisintegrants, surfactants are extensively studied in recent years for
71 improving dissolution rate and enhancing solubility but in this study Glucosamine HCl was used
72 as a hydrophilic carrier to increase the solubility and dissolution of poorly soluble drug Ibuprofen
73 because glucosamine HCl is more stable as compared to other salts of glucosamine. The same
74 study was conducted by Al-Hamidi [4], using glucosamine HCl as a carrier for improving
75 dissolution rate and enhancing solubility of poorly water-soluble drug Carbamazepine.

76 **Experimental**

77 *Materials*

78 Model drug Ibuprofen (Gratis sample by drug testing laboratory, Peshawar, Pakistan),
79 Glucosamine HCl (Sigma, UK), Distilled water, ethanol (Fisher Scientific, UK), KH_2PO_4
80 (Sigma, UK), NaOH (Sigma, UK). All chemicals and solvent used in this study were of
81 analytical grade and used as obtained.

82 *Methods*

83 *Preparation of solid dispersions*

84 Solid dispersions of Ibuprofen were prepared with drug and carrier (Glucosamine HCL) ratio
85 1:1, 1:2 and 1:3 by weight, using solvent evaporation technique[12,13]. The drug was dissolved
86 in ethanol followed by the addition of carrier dispersion in ethanol. The solvent was then
87 removed by evaporation keeping at 40° C under stirring condition (100rpm) for 24 hours. The
88 solid dispersions prepared were then collected and kept at room temperature for 48 hours. Then
89 the mass was pulverized in porcelain mortar and pestle and passed through sieve no 100, and
90 stored at room temperature in a desiccator until further use.

92 *Preparation of physical mixtures*

93 For comparative studies of solid dispersions, physical mixtures were also prepared. The physical
94 mixtures prepared were having the same composition of the solid dispersions; however, they
95 were prepared by simple trituration of drugs and carrier in porcelain mortar followed by
96 thorough blending in poly bags. The mixtures were then sieved and stored in desiccator at room
97 temperature until further evaluation.

98 The composition of physical mixtures and solid dispersions of the model drugs is shown in tables
99 1.

101

102 **Table 1 Composition of solid dispersions and physical mixtures of Ibuprofen**

Formulation Code	Carrier	Drug : Carrier	Method
F1IBF	Glucosamine HCL	1:1	Physical mixture (trituration)
F2 IBF	Glucosamine HCL	1:2	Physical mixture (trituration)
F3 IBF	Glucosamine HCL	1:3	Physical mixture (trituration)
F4 IBF	Glucosamine HCL	1:1	Solid dispersion (solvent evaporation)
F5 IBF	Glucosamine HCL	1:2	Solid dispersion (solvent evaporation)
F6 IBF	Glucosamine HCL	1:3	Solid dispersion (solvent evaporation)

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104 ***Evaluation of solid dispersions and physical mixtures***

105 The evaluation of solid dispersion and physical mixture was performed using the following
106 different techniques:

107

108 ***Determination of drug content***

109 The drug content in each formulation was determined by taking the solid dispersions or physical
110 mixtures equivalent to 50mg of the respective model drug (Ibuprofen) and transferring it to
111 volumetric flask of 100ml and then small volume of phosphate buffer (pH 7.4) was added to
112 hydrate the samples. Finally the volume was made upto the mark. The samples were shaken for
113 some time to dissolve the drugs completely and were filtered carefully. The absorbance values of
114 standard (Ibuprofen, supplied by Abbott Laboratory, Karachi, Pakistan) and the samples were
115 determined at λ_{max} 223 nm, using double beam spectrophotometer (UV-1601, Shimadzu,
116 Japan). Three reading were taken and then mean and standard deviation were calculated.

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118 ***Differential scanning calorimetry (DSC) studies***

119 The differential scanning calorimetry (DSC) study of carrier Glucosamine, pure Ibuprofen, the
120 solid dispersions and physical mixtures of the model drug was performed using DSC instrument
121 (Mettler Toledo DSC 822e, Greifensee, Switzerland) equipped with Star^e computer program.
122 Approximately 3-6mg of sample was weighed in aluminum pan and then sealed with punched

123 lid. The temperature ranged between 20-300°C with heating rate of 10°C/min under nitrogen gas
124 flow.

125

126 ***Fourier transform Infrared (FT-IR) studies***

127 The FT-IR spectra of carrier Glucosamine, pure Ibuprofen, the solid dispersions and physical
128 mixtures were taken to observe the drugs-carrier interaction, using FT-IR SpectrumOne
129 spectrophotometer (Perkin Elimer, UK) in the range of 650 to 4000 cm^{-1} . The sample of several
130 milligrams was placed on the stage of machine and then handle of the machine was placed on the
131 sample for generation of enough pressure. Then sharp peaks with reasonable intensities were
132 obtained. The spectra obtained were the result of 4 scans at 1 cm^{-1} resolution.

133 ***X-ray powder diffractometry studies***

134 X-ray patterns of pure Ibuprofen, pure, physical mixtures and solid dispersions were taken using
135 a Philips PW 1830 powder diffractometer (Philips, Eindhoven, Netherlands). The prepared
136 samples were exposed to Cu K α radiation ($\lambda= 1.5418 \text{ \AA}$) in the range of $0^{\circ} \leq 2\theta \leq 50^{\circ}$. The step
137 size was 0.05° and the time for each step was kept two seconds.

138 ***Scanning electron microscope (SEM) analysis***

139 Electron micrographs of carrier Glucosamine, pure Ibuprofen, pure, physical mixtures and solid
140 dispersions were obtained using scanning electron microscope (SEM; Joel JSM-5910, Japan)
141 operating at 10 kV. The samples were mounted on a metal stub using adhesive tape with double
142 sided and coated with gold for conductivity in an organ atmosphere before observation. To study
143 the morphology of active drugs, physical mixture and solid dispersions, micrographs with
144 different magnification were obtained.

145 ***Solubility measurement***

146 The solubility measurements of pure Ibuprofen, physical mixtures and solid dispersions in
147 distilled water were performed according to the well published method by Higuchi and Connors
148 (1965), accordingly, surplus amount (100mg) of Ibuprofen, physical mixtures and solid
149 dispersions were placed in 100ml volumetric flasks and then made the final volume with the
150 distilled water up to 100ml. The flasks were sealed with aluminum foils using rubber bands to
151 avoid solvent loss. Then these flasks were kept on shaking using thermostatically controlled
152 shaking water bath (Shel Lab, 1217-2E, USA) for 24 hours at room temperature (25°C). The
153 oscillation speed was kept at 100 oscillations per minute. After 24 hours all flasks were kept

154 undisturbed on flat surface for three hours. A few ml supernatant from each flask was taken and
155 filtered through membrane filter (0.45 μ m). One ml each filtrate was diluted with the same
156 distilled water up to 25ml to achieve suitable dilutions. The diluted samples were analyzed to
157 determine the Ibuprofen solubility, using a UV/Visible double beam spectrophotometer
158 (Shimadzu, 1601, Japan) at λ max 223nm. The calibration curve was used for the determination
159 of the quantity of soluble drug per ml.

160 ***In –vitro dissolution studies***

161 The *in-vitro* dissolution studies were performed by USP method II (Paddle method) using eight
162 stations dissolution apparatus Pharma Test (PTWS-11/P, TPT, Hunburg, Germany) and the
163 rotation speed of paddles was set at 100 r. p.m. Each station or flask of the dissolution apparatus
164 was filled with 900ml of distilled water used as dissolution medium to study percentage
165 dissolution of model drugs (Ibuprofen), physical mixtures and solid dispersions. The temperature
166 of dissolution medium was kept 37°C \pm 0.5°C. Samples of five ml were withdrawn at selected
167 time intervals (5, 10, 15, 20, 25, 30, 40, 50, 60, 70, 80, 90, 100, 110, 120 min) with the help of
168 syringes consisting of 0.45 μ m filters. After each sampling equal volume of fresh dissolution
169 medium was replaced to maintain the dissolution medium constant. Then after appropriate
170 dilution the samples were analyzed for Ibuprofen, using double beam spectrophotometer (UV-
171 1601, Shimadzu, Japan) at λ max 223nm. Percent drug dissolution of Ibuprofen was calculated by
172 using calibration standard curves of the drug. The study was conducted in triplicate.

173

174 **Results and discussions**

175 ***Preparation of solid dispersions***

176 Different methods such as salt formation, solubilization, particle size reduction, complex
177 formation, solvent evaporation, etc. are used to prepare solid dispersions to enhance the
178 dissolution rate and thereby improve the bioavailability of poorly water soluble drugs[14],
179 however, in this study solvent evaporation method was used due to its inherent ease of handling
180 and no more steps were required. The solid dispersions of model drug (Ibuprofen) with different
181 drug and carrier ratios were prepared. The respective physical mixtures with the same drug and
182 carrier ratios were prepared by simple trituration method for comparative evaluation.

183 For conformation of uniform dispersion of drug in solid dispersions and physical mixtures drug
184 content analysis was performed and it was found between 99.57 ± 0.7 % and 101.3 ± 0.32 %. All
185 the solid dispersions and physical mixtures indicated the high content and uniformly dispersion
186 of drugs. These findings conformed that the solvent evaporation method appears to be
187 reproducible for development and preparation of solid dispersions. Similar studies were
188 conducted by Prasad and Rosario [12,15], who prepared solid dispersions of Tebinanfine
189 hydrochloride and NSAIDs by the same method obtaining good results in terms of content
190 analysis and uniform distribution of the drugs used.

191

192 *Solubility study*

193 As shown in table 2, the aqueous solubility study of pure Ibuprofen, their physical mixtures and
194 solid dispersions was performed in distilled water. The study showed that solubility of Ibuprofen
195 was enhanced in presence of carrier (Glucosamine HCL). This effect of solubility enhancement
196 was more prominent in case of solid dispersions as compared to that of their respective physical
197 mixtures. The enhancement of drugs solubility in presence of solid dispersions may be due to
198 conversion of drugs to amorphous form as amorphous forms of drug are more soluble than their
199 crystalline form [12,16]. The increase in solubility of drugs in solid dispersions might also be
200 due to good wettability and dispersability [16].

201

202 **Table 2 Solubility data of different Ibuprofen formulations.**

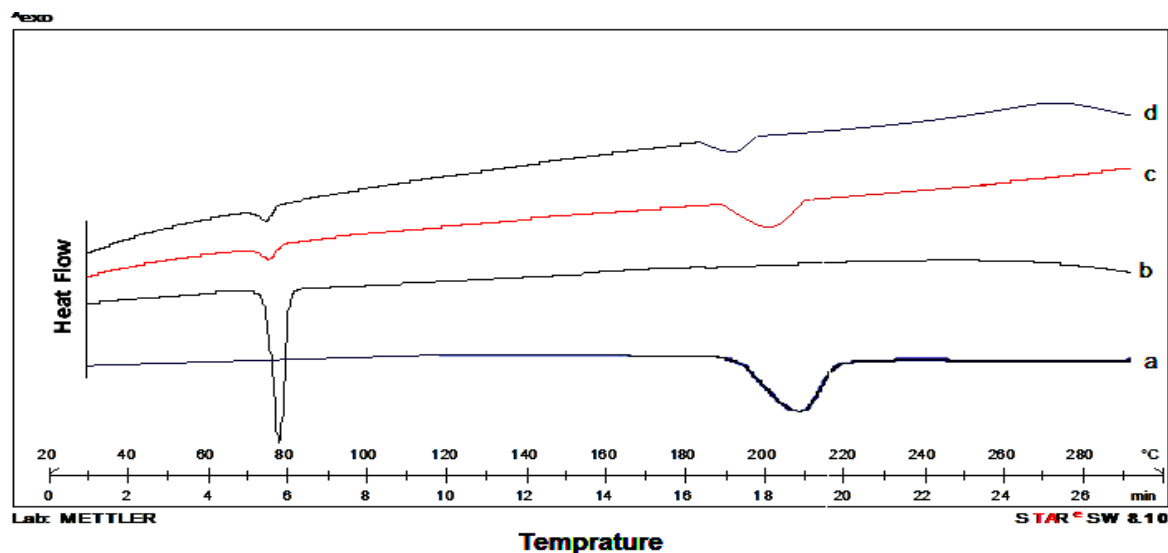
Formulations	Solubility (mg/ml)
Ibuprofen	
IBF Pure	0.285
F1 IBF	0.297
F2 IBF	0.313
F3 IBF	0.333
F4 IBF	0.320
F5 IBF	0.357
F6 IBF	0.398

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204 *Differential scanning calorimetry (DSC) studies*

205 Differential scanning calorimetric (DSC) studies of pure ibuprofen, their physical mixtures and
206 solid dispersions were conducted to investigate the crystallinity and drugs carrier interaction. The

207 DSC run of the pure Ibuprofen and carrier (glucosamine HCL) show sharp endothermic peaks
208 around 76.94°C and 210°C, corresponding to the melting point of Ibuprofen and Glucosamine
209 HCL, respectively (Fig. 1a-d).



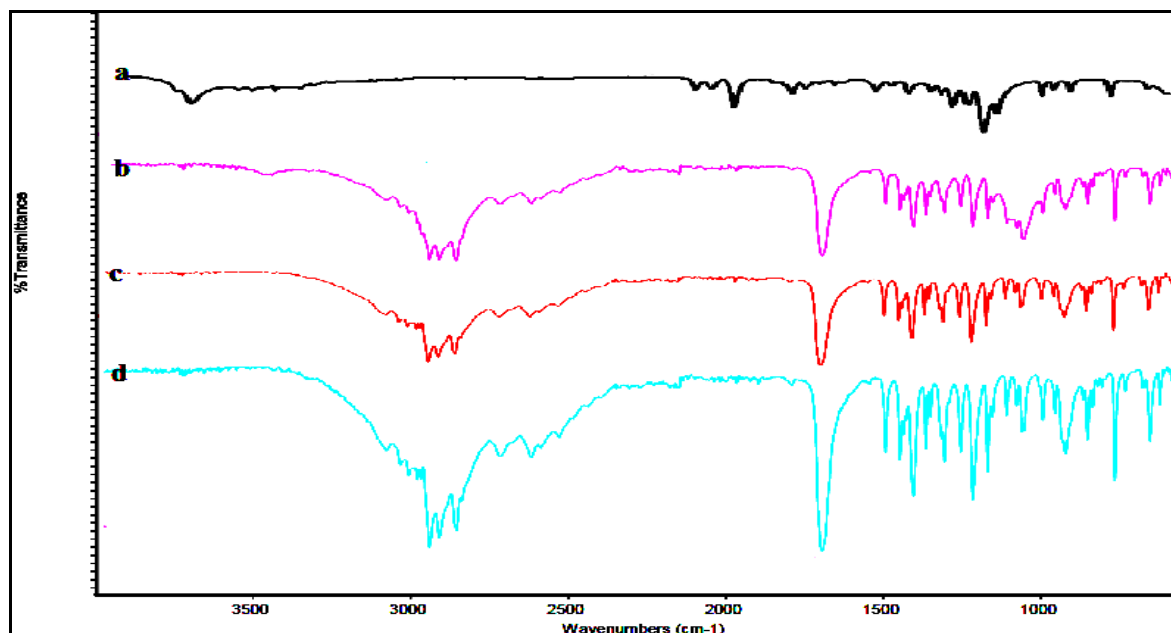
210
211 **Figure 1 DSC Thermograms of (a) Glucosamine; (b) Pure ibuprofen; (c) Physical mixture;**
212 **and (d) Solid dispersions of ibuprofen with glucosamine.**
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214 The endothermic peak of Ibuprofen is of very high intensity, showing the crystalline form of
215 ibuprofen. The DSC thermograms of Ibuprofen-carrier (Glucosamine HCL) physical mixture and
216 solid dispersions showed both the endothermic peaks (Fig. 1c-d) with some changes in the
217 characteristics of the peaks shown by individual components; for example the endothermic peaks
218 of physical mixture and solid dispersions lost its sharpness and distinctive appearance. It showed
219 that no possible interaction was found between drug and carrier but the loss of peaks sharpness
220 may be due to conversion from crystalline form to amorphous form of the drug.

221 ***Fourier transform Infrared (FT-IR) studies***

222 For the conformation of interaction between drugs and carrier in presence of physical mixtures
223 and solid dispersions FT-IR studies were performed. The FT-IR spectrums of pure Ibuprofen,
224 and Ibuprofen-Glucosamine physical mixtures and solid dispersions were obtained as shown in
225 the Fig. 2a-d. Pure Ibuprofen showed sharp characteristic peaks at 1706 cm⁻¹ which corresponds
226 to the carboxyl acid (COOH) present in ibuprofen. Other smaller peaks in the region 1200-1000
227 cm⁻¹ are the indication of benzene ring [17]. These peaks can also be seen in the ibuprofen-
228 carrier physical mixture and solid dispersions, but in this case IR spectrum for Ibuprofen-carrier

229 mixture and solid dispersion shows the overlapping of carboxyl acid group (Fig. 2c-d).
230 Therefore, it can be concluded that no chemical interaction occurred between Ibuprofen and
231 glucosamine HCL.



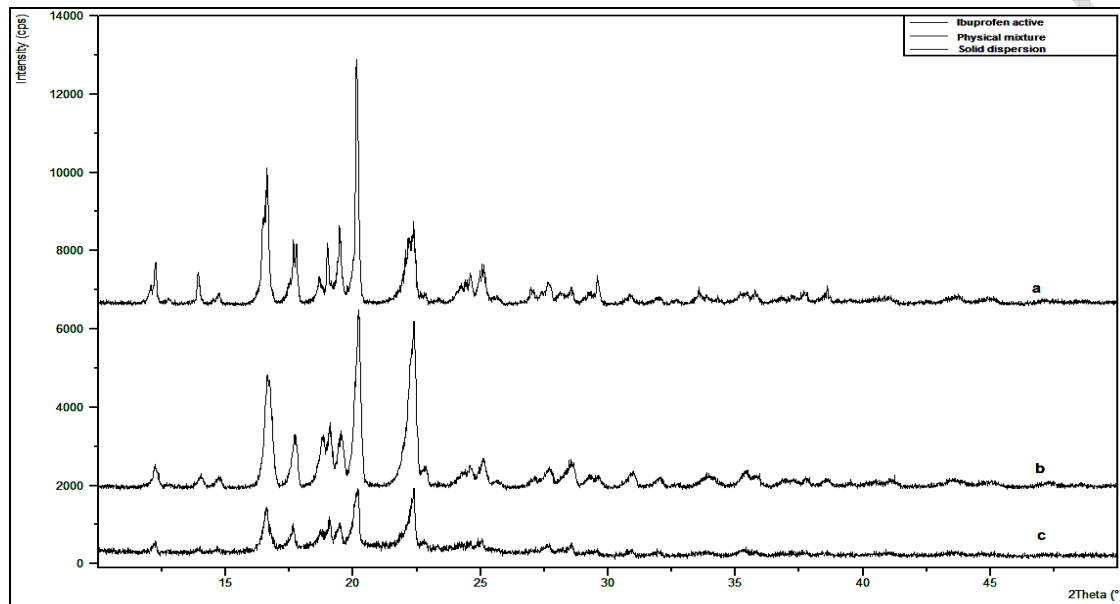
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233 **Figure 2 FT-IR spectra of (a) Glucosamine; (b) Pure ibuprofen; (c) Physical mixture; and**
234 **(d) Solid dispersions of ibuprofen with glucosamine.**
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X-ray diffractometry studies

Figure (3a-c) shows the diffractograms of pure Ibuprofen, Ibuprofen-carrier physical mixture and solid dispersion.



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Figure 3 X-ray diffractograms of (a) Pure Ibuprofen; (b) Physical mixture; and (c) Solid dispersions of Ibuprofen and glucosamine.

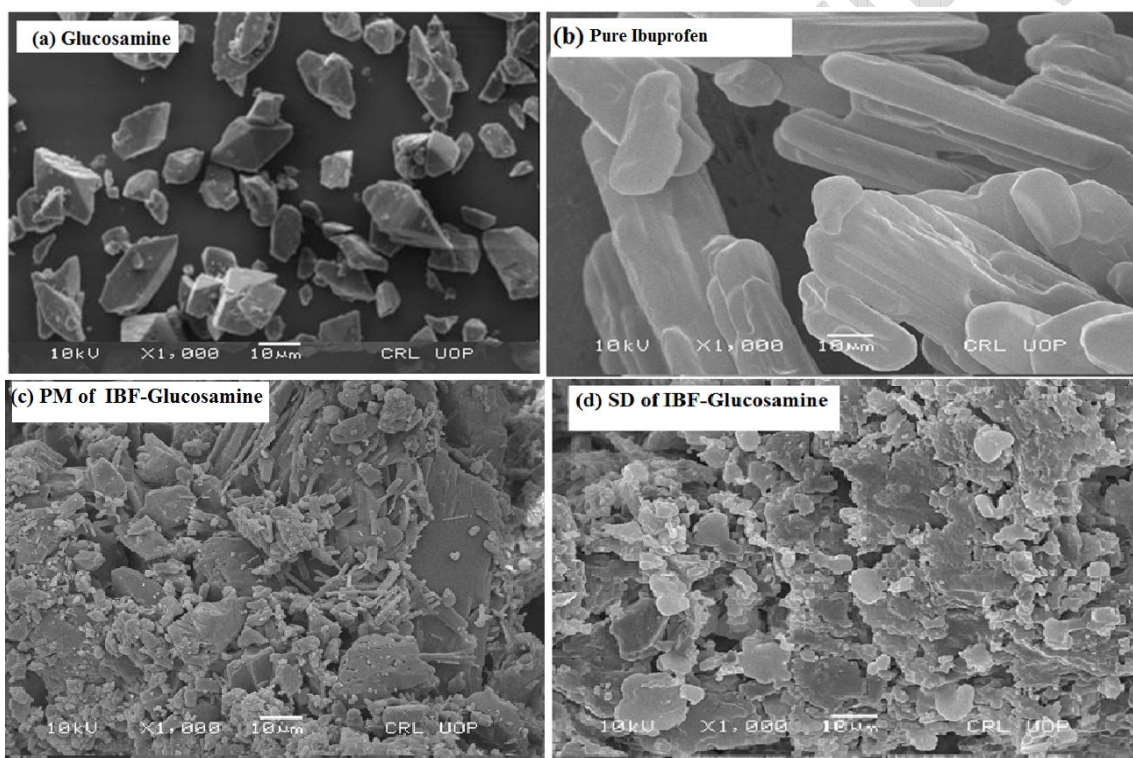
The diffractograms of pure Ibuprofen with numerous distinctive peaks showed that the drug is highly crystalline in nature, conforms the DSC studies as shown in figure (3b). Four peaks with high intensity were present in the diffractogram of Ibuprofen around 17°, 20°, 23° and 25° along with some other peaks of lower intensity. The same peaks were present in the diffractogram of Ibuprofen-carrier physical mixture and solid dispersions, but with lower intensity. This indicates that Ibuprofen crystallinity has been diminished. As compared to pure Ibuprofen and physical mixture of Ibuprofen-carrier, the peaks in the diffractogram of solid dispersions were of much reduced intensities, indicating the amorphous nature of the Ibuprofen in presence of solid dispersions.

Scanning electron microscope analysis

Figure 4a-d shows the scanning electron micrographs of Glucosamine HCL, pure Ibuprofen, Ibuprofen-carrier physical mixture and solid dispersions of Ibuprofen with glucosamine HCL.

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272 After analysis, the scanning electron microscopy (SEM) revealed that Glucosamine has prismatic
273 shape (polygonal) and pure Ibuprofen has irregular crystalline shape. Both of these crystals can
274 easily be identified in the physical mixture, as shown in the Figure 4c. In physical mixture, there
275 are numerous small crystals of Ibuprofen which are responsible for more solubility and enhanced
276 dissolution rate as compared to pure compound, while in case of solid dispersions the crystals of
277 Ibuprofen are in smallest size and they have irregular, circular and plate like shapes. The
278 dissolution rate of Ibuprofen in solid dispersions was rapid and more as compared to pure
279 Ibuprofen and physical mixture because the particle shape irregularity and small particle size
280 increased the specific surface area and enhanced the dissolution rate[18].

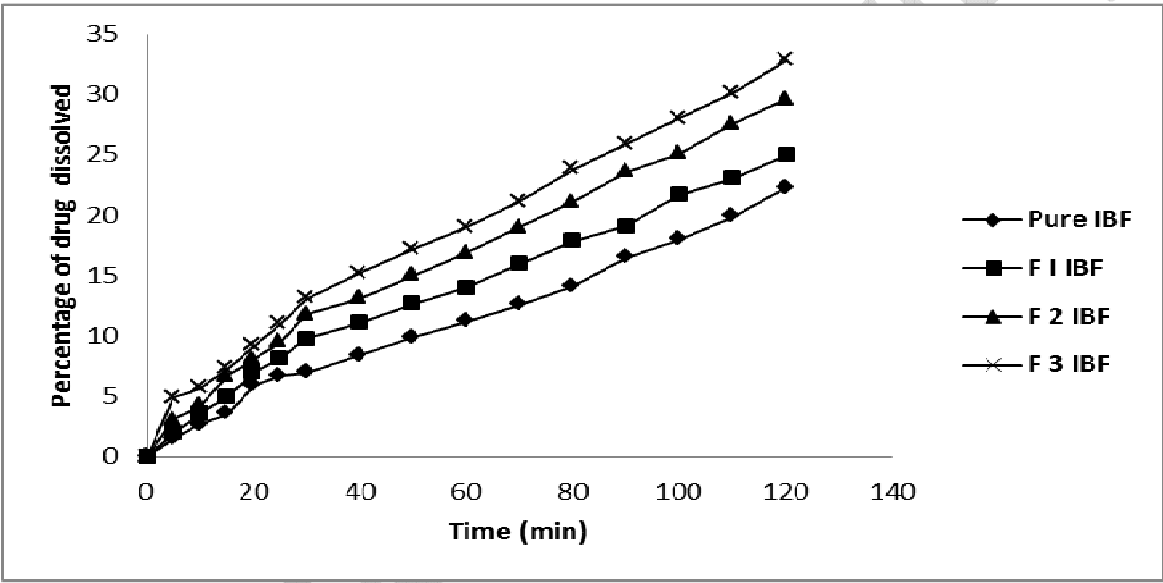


281
282 **Figure 4 Scanning electron photomicrographs of (a) Carrier (Glucosamine HCL); (b) Pure**
283 **Ibuprofen; (c) Physical mixture of Ibuprofen-Glucosamine HCL; (d) Solid dispersion of Ibuprofen-**
284 **Glucosamine HCL.**
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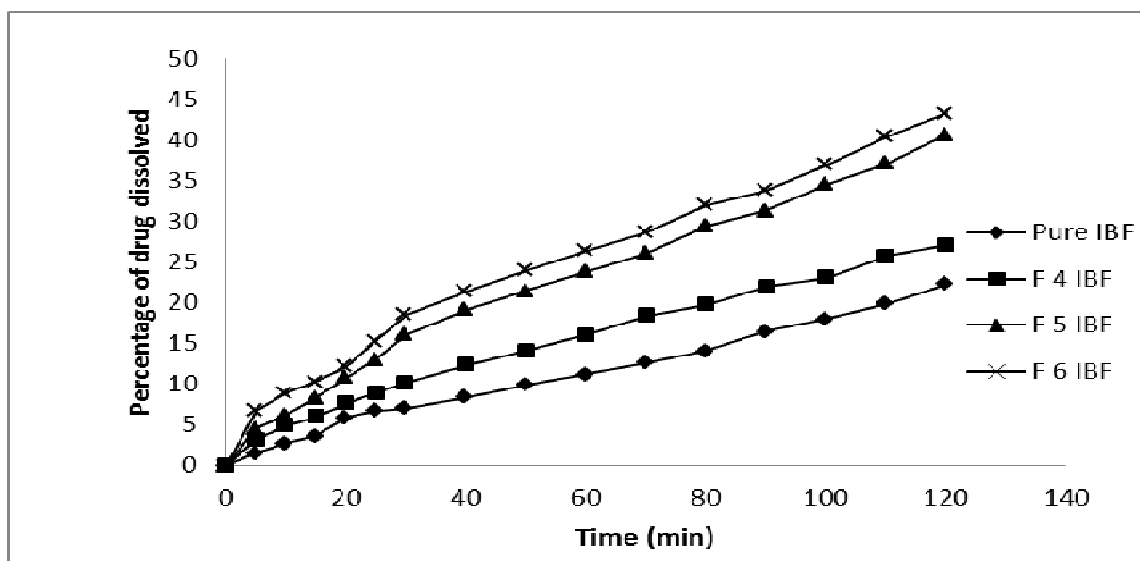
286 *In –vitro dissolution studies*

287 The dissolution profiles of pure Ibuprofen, Ibuprofen physical mixtures and solid dispersions
288 prepared with Glucosamine HCL are shown in Figs. 5 and 6. It is shown that pure Ibuprofen has
289 the slowest dissolution rate and 22.3% of drug was dissolved after 120 minutes, while in case of
290 physical mixtures and solid dispersions with different Drug: Carrier ratios (1:1, 1:2 and 1:3) the

291 dissolution rate was linearly increased and 25%, 29.65, 32.8% and 27.1%, 40.75, 43.3% of drug
292 was dissolved after 120 minutes from formulations F1 IBF, F2 IBF, F3 IBF and F4 IBF, F5 IBF,
293 F6 IBF, respectively. The fastest dissolution rate was obtained for the formulation (F6 IBF) with
294 the D: C ratio of 1:3 in carrier concentration dependent manners. The fast and rapid dissolution
295 rate of Ibuprofen in solid dispersion may be due to the presence of Ibuprofen in amorphous form
296 which is revealed by the results of different techniques as mentioned above. On the other hand it
297 may be that if the percentage of carrier is too high, this may lead to increase in solubility and
298 dissolution rate due to absence of crystallinity of drug [19].
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302 **Figure 5** *In-vitro* dissolution profiles of pure Ibuprofen and physical mixture with different
303 drug-carrier (Glucosamine HCL) ratio.
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 307 **Figure 6** *In-vitro* dissolution profiles of pure Ibuprofen and solid dispersions with different
 308 drug-carrier (Glucosamine HCL) ratio
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310 Conclusion

311 The study shows that the dissolution rate and solubility of sparingly soluble drug Ibuprofen can
 312 be improved and enhanced to great extent by solid dispersion technique, using Glucosamine HCl
 313 as a carrier. The current study also showed that amino sugar could be used as new carrier for
 314 solid dispersion formulations of non-steroidal anti-inflammatory drugs (Ibuprofen).

315 Acknowledgement

316 Abdul Wahab thanks the Higher Education Commission (HEC) of Pakistan for providing PhD
 317 scholarship.

318 References

- 319 [1] Saharan VA, Kukkar V, Kataria M, Gera M, Choudhury PK. Dissolution Enhancement of
 320 Drugs. Part I: Technologies and Effect of Carriers. *Int J Health Res.* 2009; 2: 107-124.
 321 [2] Patel Rajanikant P, Nirav P, Patel NM, Patel MM. A novel approach for dissolution
 322 enhancement of Ibuprofen by preparing floating granules. *Int J Res Pharm Sci.* 2010; 1: 57-
 323 64.
 324 [3] De Brabander C, Vervaet C, Bortel LV, Remona JP. Bioavailability of ibuprofen from hot-
 325 melt extruded mini-matrices. *Int J Pharm.* 2004; 271: 77-84.
 326 [4] Al-Hamidi H, Edwards AA, Mohammad AM, Nokhodch A. To enhance dissolution rate of
 327 poorly water-soluble drugs: Glucosamine hydrochloride as a potential carrier in solid
 328 dispersion formulations, *Colloids Surf. B: Biointerface.* 2009;
 329 doi:10.1016/j.colsurfb.2009.10.030.

- 330 [5] Babu GV, Kumar NR, Himasankar K, Seshasayana A, Murthy KV. Nimesulide-modified
331 gum karaya solid mixtures: preparation, characterization and formulation development. *Drug*
332 *Dev Ind Pharm.* 2003; 29: 855-864.
- 333 [6] Rogers JA, Anderson AJ. Physical characteristics and dissolution profiles of ketoprofen-urea
334 solid dispersions. *Pharm Acta Helv.* 1982; 57: 276-281.
- 335 [7] El-Gazayerly ON. Characterization and evaluation of tenoxicam coprecipitates. *Drug Dev*
336 *Ind Pharm.* 2000; 26: 925-930.
- 337 [8] Vippagunta SR, Maul KA, Tallavajhala S, Grant DJW. Solidstate characterization of
338 nifedipine solid dispersions. *Int J Pharm.* 2002; 236: 111-123.
- 339 [9] Murali Mohan Babu GV, Prasad CHDS, Ramana Murthy KV. Evaluation of modified gum
340 karaya as carrier for the dissolution enhancement of poorly water soluble drug nimodipine.
341 *Int J Pharm.* 2002; 234: 1-17.
- 342 [10] Chiou WL, Riegelman S. Pharmaceutical applications of solid dispersion systems. *J Pharm*
343 *Sci.* 1971; 60: 1281-1302.
- 344 [11] Chaulang G, Patil K, Ghodke D, Khan S and Yeole P. Preparation and Characterization of
345 Solid Dispersion Tablet of Furosemide with Crospovidone. *Research J. Pharm. and Tech.*
346 2008; 1: 386-389.
- 347 [12] Parsad KA, Narayanan N and Rajalakshmi G. Preparation and Evaluation of Solid
348 Dispersion of Terbinafine Hydrochloride. *Int. J. Pharm. Sci. Rev. and Res.* 2010; 3: 130-134.
- 349 [13] Jain R, Jain Kaushal, Setty CM and Patel D. Preparation and Evaluation of Solid
350 Dispersions of Aceclofenac. *Int. J. Pharm. Sci. Drug. Res.* 2009; 1: 32-35.
- 351 [14] Galia E, Nicolaidis E, Horters D, Lobenberg R, Reppas C and Dressman B. Evaluation of
352 Various Dissolution Media for Predicting In Vivo Performance of Class I and Class II Drugs.
353 *Pharm. Res.* 2009; 15: 698-705.
- 354 [15] Rosario P, Maranilla F and Giovanni P. Preparation of Solid Dispersions of NSAIDs with
355 Acrylic Polymers and Studies on Mechanism of Drugs Polymer Interactions. *AAPS.*
356 *PharmSciTech.* 2002; 3: Article 10
- 357 [16] Khan GM and Zhu JB. Ibuprofen Release Kinetics from Controlled-release Tablets
358 Granulated with Aqueous Polymeric Dispersion of Ethylcellulose II: Influence of Several
359 Parameters and Coexcipients. *J Control Rel.* 1998; 59: 127-134.
- 360 [17] Socrates G. Infrared characteristic group frequencies, table and chart, 2nd ed. Wiley, New
361 York. 1994.
- 362 [18] Javadzadeh Y and Nokhodchi A. *Acta Pharm.* 2009; 59: 187.
- 363 [19] Ford JL. The Current states of Solid dispersions. *Pharm. Acta. Helv.* 1986; 61: 69-88.
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