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Molecular modelling as a predictive tool for the development of solid dispersions

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ABSTRACT

In this study molecular modelling is introduced as a novel approach for the development of pharmaceutical solid dispersions. A computational model based on quantum mechanical (QM) calculations was used to predict the miscibility of various drugs in various polymers by predicting the binding strength between the drug and dimeric form of the polymer. The drug/polymer miscibility was also estimated by using traditional approaches such as Van Krevelen/Hoftyzer and Bagley solubility parameters or Flory Huggins interaction parameter in comparison to the molecular modelling approach.

The molecular modelling studies predicted successfully the drug-polymer binding energies and the preferable site of interaction between the functional groups. The drug-polymer miscibility and the physical state of bulk materials, physical mixtures and solid dispersions were determined by thermal analysis (DSC/MTDSC) and X-ray diffraction. The produced solid dispersions were analysed by X-ray photoelectron spectroscopy (XPS), which confirmed not only the exact type of the intermolecular interactions between the drug-polymer functional groups but also the binding strength by estimating the N-coefficient values. The findings demonstrate that QM-based molecular modelling is a powerful tool to predict the strength and type of intermolecular interactions in a range of drug/polymeric systems for the development of solid dispersions.

Keywords: solid dispersions, quantum mechanics, molecular modelling, miscibility, drug-polymer interactions.

INTRODUCTION

Drug-polymer miscibility is a significant factor to consider when designing solid dispersions for pharmaceutical dosage forms¹. The interactions between small molecule drugs and polymers in a polymeric matrix system determines the nature of the drug loading inside the polymer matrix, such as oversaturated or under saturated, thus play a vital role in determining the long-term stability of the formulated product. Therefore, the characterisation and prediction of drug-polymer miscibility and more importantly drug solubility at various conditions has become an emerging topic in pharmaceutical research for the development of solid dispersions¹⁻³. The dispersion of a small drug molecule being miscible into a polymer matrix is mechanistically a complex process. Both the repulsive and attractive intra- and intermolecular forces (e.g. dispersion force, dipole-dipole interaction, and hydrogen bonding forces) present in the system may further complicate the process. As a mechanistic rule, the molecular level mixing requires a step of balancing and/or breaking the crystalline lattice of drug molecules (with high activation energy) in the first instance. Subsequently, the polymer swelling (a process of polymer chain segment absorbing drug molecule) and rebuilding of the possible interactions inside the mixture occur⁴. The drug molecule (either crystalline or amorphous) may have significantly different physical properties within the polymer matrices after processing due to the different energy input and possible inter-molecular interactions that are required to form a solid dispersion. In terms of the random distribution of small drug molecules into the polymeric matrices, the miscible drug content demonstrates amorphous characteristics.

The lattice-based Flory-Huggins (F-H) theory (Flory 1952) is well known for the prediction of polymer-solvent or polymer-polymer interactions based on the Gibbs free energy change before and after mixing⁵. The F-H theory can be successfully applied to determine the interaction parameter therefore signifying the strength of the interactions between two compounds during their melting. Recent publications applying the F-H theory in solid dispersions focus on obtaining the F-H interaction parameter, χ , by the melting point depression method^{6, 7}. The employment of F-H theory for the prediction of the intermolecular interactions in pharmaceutical binary systems can provide important insights in the development of solid dispersions processed by hot melt extrusion (HME) for example. However, the theory assumes random distribution of the segments, which may not be suitable in cases where there are strong polar forces or specific interactions, such as hydrogen

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3 bonding, between the components of the blends ⁸. A deeper understanding of the relatively
4 stronger interactions between the drug and the polymer has to be built on insights of the
5 atomic level interactions. Molecular modelling approach is a powerful addition to the F-H
6 theory to characterise the drug-polymer interactions through both visualisation of the
7 interaction and estimation of the strength of the interactions. To date, a number of molecular
8 mechanics (MM)-based molecular dynamics (MD) simulation studies ⁹⁻¹² have been
9 performed to study various molecules (e.g. dissolution of the molecules from the crystal
10 surface)¹³ or to identify drug-polymer miscibility in solid dispersions ¹⁴⁻¹⁸. Recently, we ¹⁹
11 reported the use of quantum mechanical (QM) calculations with commercially available
12 software (Gaussian 09) to characterise possible drug/polymer interactions. Our calculations
13 indicated the existence of hydrogen bonding between the amine group of the drug molecule
14 and the carboxyl groups of the polymer among different drug/polymer formulations, which
15 was subsequently, confirmed by NMR and X-ray photoelectron spectroscopy (XPS) studies.
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26 In recent years, hot-melt extrusion (HME) has undergone a renaissance in the research
27 and development of pharmaceutical formulations ²⁰. HME is a manufacture processing
28 technology that combines the advantages of a solvent-free solid dispersion and has been used
29 for a wide range of solid dispersion with increased drug solubility, taste masking of bitter
30 APIs, and sustained/controlled release properties ^{20, 21}.
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35 In this study, we developed a new approach to determine the drug-polymer miscibility
36 based on quantum mechanical (QM) calculations, whereby the calculated binding energy
37 between the drug and the polymer provides a strong indication for the formation of stable
38 solid dispersions. The QM approach was compared with the Hansen solubility parameters
39 and F-H estimated values in order to demonstrate the superiority of the molecular modelling
40 approach, which not only provides more accurate description of the drug-polymer interaction
41 energy, but also predicts the interacting functional groups between both substances. The latter
42 was confirmed by X-ray photoelectron spectroscopy (XPS) ^{22, 23} studies on the produced
43 HME solid dispersions. XPS has been reported as a valuable tool for the characterisation of
44 solid dispersions by providing information on the nature (e.g. H-bonding) and specific site of
45 chemical bonding on the surface of the solid dispersion.
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MATERIALS AND METHODS

Materials. Propranolol HCl (PRP) and Diphenhydramine HCl (DPD), Paracetamol (PMOL), Ibuprofen (IBU), Dichlofenac Na (Df-Na) and Hydrocortisone (HCS) were purchased from Sigma Aldrich (London, UK) of $\geq 98\%$ purity. Eudragit L100 (L100), Eudragit EPO (EPO) and Eudragit L100-55 were kindly donated by Evonik Pharma Polymers (Darmstadt, Germany). Kollidon VA64 (VA64) was kindly donated by BASF (Germany). All materials were used as received.

Hot-melt extrusion (HME) processing. Drug/polymer blends (PRP/L100, PRP/L100-55, DPD/L100, DPD/L100-55, PMOL/VA64, IBU/VA64 and IBU/EPO) were blended in 100 g batches for 10 min each with a Turbula (TF2, Basel) mixer. The extrusion of all batches was performed using a Randcastle single-screw extruder (RCP0625) equipped with a 5-mm rod die at $100^{\circ}\text{C}/140^{\circ}\text{C}/155^{\circ}\text{C}/155^{\circ}\text{C}/150^{\circ}\text{C}$ (Feeder \rightarrow Die) temperature profiles for PRP and DPD based formulations, $100^{\circ}\text{C}/113^{\circ}\text{C}/115^{\circ}\text{C}/115^{\circ}\text{C}/115^{\circ}\text{C}$ (Feeder \rightarrow Die) for PMOL and $70^{\circ}\text{C}/85^{\circ}\text{C}/85^{\circ}\text{C}/85^{\circ}\text{C}/80^{\circ}\text{C}$ (Feeder \rightarrow Die) for IBU based formulations. The screw speed was set at 15rpm for all drug/polymer binary compositions. The produced extrudates (strands) were grinded by using a Ball Mill system (Retsch, Germany) to obtain granules ($<500\ \mu\text{m}$) at a rotational speed of 400 rpm for 5 min. The drug-polymers composition consisted of PRP/L100, PRP/L100-55, DPD/L100 and DPD/L100-55 were extruded at ratios of 10/90 w/w. These four extruded formulations (comprising oppositely charged drugs and polymers) were used in XPS analysis to determine possible intermolecular interactions.

Hansen solubility parameters: Predictions of drug/polymers miscibility. The theoretical drug-polymers miscibility was determined by applying the Hansen solubility parameters (δ) for both drugs as well as the polymers. The Hoftyzer and van Krevelen method was used to calculate all drug polymers solubility parameters by considering the chemical structural orientations²⁴ according to the following equation:

$$\delta^2 = \delta_d^2 + \delta_p^2 + \delta_h^2 \quad (1)$$

Here,

$$\delta_d = \frac{\sum F_{di}}{V_i}, \delta_p = \frac{\sqrt{\sum F_{pi}^2}}{V_i}, \delta_h = \frac{\sqrt{\sum E_{hi}}}{V_i}$$

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i = structural groups within the molecule, δ = the total solubility parameter. F_{di} = molar attraction constant due to molar dispersion forces, F_{pi}^2 = molar attraction constant due to molar polarization forces, E_{hi} = hydrogen bonding energy, and V_i = group contribution to molar volume

By using the two – dimensional approach Bagley *et al.*²⁵ introduced the volume – dependent solubility parameter which is defined as δ_v , where δ_p and δ_d were used to predict the combined thermodynamic effects on the drug-polymers miscibility over hydrogen bonding energy, δ_h .

$$\delta_v = \sqrt{\delta_d^2 + d_p^2} \quad (2)$$

Flory Huggins (F-H) theory for the prediction of drug/polymers interaction parameter.

The F–H interaction parameter, χ , was correlated with temperature dependence, as in many simplified cases. The interaction parameter of the model system was determined at two different conditions using the Nishi–Wang Eq. 3²⁶ equation based on melting point depression data and Hildebrand and Scott Eq. 4²⁷ correlations with solubility parameter, respectively. The F-H interaction parameter (χ) of the drug/polymers as shown in Table 2 binary mixtures were calculated by using the following equations. The value determined by Eq. 3 represents the interactions between the two substances, specifically at the melting temperature, which may not be extrapolated to other temperatures.

$$\frac{1}{T_m} - \frac{1}{T_m^0} = -\frac{Rv_{drug}}{\Delta H_{drug}v_{poly}} \left[\ln\phi_{drug} + \left(1 - \frac{1}{m_{poly}}\right) \times (1 - \phi_{drug}) + \chi_{drug-poly} (1 - \phi_{drug})^2 \right] \quad (3)$$

Where, v is the molar volume of the repeating unit, m is the degree of polymerization, ϕ is the volume fraction and χ is the crystalline– amorphous polymer interaction parameter, T_m is the crystalline melting peak of the pure drugs and T_m^0 is the melting endotherm of the drug-polymer physical blends.

F-H interaction parameter (χ) can be also estimated by the method developed by Hildebrand and Scott according to the following Eq. 4²⁷.

$$\chi = \frac{v(\delta_{drug} - \delta_{poly})^2}{RT} \quad (4)$$

Where, R is the gas constant, T is the absolute temperature, and v the volume per lattice site and δ_{drug} and δ_{poly} are solubility parameters of drugs and polymers respectively.

Thermal analysis (DSC and MTDSC). A Mettler-Toledo 823e (Greifensee, Switzerland) differential scanning calorimeter (DSC) was used to carry out thermal analysis of the bulk APIs, polymers, drug/polymer binary physical mixtures and extruded formulations ($n=3$). About 3-5 mg of each samples were placed in sealed aluminium pans with pierced lids. The samples were heated, cooled and reheated at $10^{\circ}\text{C}/\text{min}$ from 0°C to 220°C under an atmosphere of dry nitrogen. In addition modulated temperature differential scanning calorimetry (MTDSC) studies were performed from 20°C to 150°C with an underlying heating rate of $1^{\circ}\text{C}/\text{min}$. The pulse height was adjusted to $1\text{-}2^{\circ}\text{C}$ with a temperature pulse width of 15-30 s.

Molecular Modelling. The dimeric structures of L100, L100-55, EPO and VA64 and the monomeric structures of PRP, DPD, PMOL, IBU, Df-Na and HCS were constructed by program Gaussview²⁸. Hydrogen bonding patterns were identified by placing the drug molecule within the proximity of the dimeric structure of the polymer and then energy optimised to local minima at the M06-2x/6-31G** level and verified by frequency calculations using Gaussian09. A counterpoise procedure²⁹ was employed to correct for the effect of basis set superposition error (BSSE) for the binding energy between the drug molecule and the dimer.

X-ray photoelectron spectroscopy (XPS) analysis. X-ray photoelectron spectra (XPS) were measured on a Kratos Axis Ultra-DLD using a monochromatic Al K_{α} X-ray source (120 W) and an analyser pass energy of 160 eV (survey scans) or 20 eV (high resolution scans); the pressure during analysis was 1×10^{-9} Torr. All data were referenced to the C(1s) signal at 285.0 eV attributable to unsaturated C-C/C-H bonds²². Quantification and curve fitting was performed in CasaXPSTM (Version 2.3.15) using elemental sensitivity factors supplied by the manufacturer. All samples were run in triplicate.

RESULTS AND DISCUSSIONS

Predictions of drug/polymers miscibility: Hansen solubility parameters. Up to date there are several approaches employed to estimate the drug/polymer miscibility based in the determination of the solubility parameter. The theoretical approaches derived from the solubility parameter calculations suggest that compounds (e.g. drug or polymer) with similar

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δ values are likely to be miscible. This is because the energy of mixing which is released from intra-molecular interactions is balanced with the energy of mixing from intermolecular interactions³⁰. It has been reported that compounds with $\Delta\delta < 7 \text{ MPa}^{1/2}$ are likely to be miscible and compounds with $\Delta\delta > 10 \text{ MPa}^{1/2}$ are likely to be less miscible or immiscible³⁰.

It can be seen in Table 1 that the difference between the calculated solubility parameters indicate that all of the drugs used are likely to be miscible with all polymers ($\Delta\delta < 7 \text{ MPa}^{1/2}$). By using the Van Krevelen/Hoftyzer method, the calculated $\Delta\delta$ values for DPD/L100, DPD/ L100-55, PRP/L100 and PRP/L100-55 are 5.0, 3.9, 0.81 and 0.29 $\text{MPa}^{1/2}$, respectively indicating complete drug/polymer miscibility. Similarly, $\Delta\delta$ values estimated for HCS/L100 and Df-Na/L100 are 2.59 and 3.32 $\text{MPa}^{1/2}$, respectively suggesting that both HCS and Df-Na are likely to be miscible with L100. On the other hand PMOL is unlikely to be miscible with EPO as the calculated $\Delta\delta$ falls in the range close to $7\text{MPa}^{1/2}$ threshold while IBU seems highly miscible with EPO ($\Delta\delta$ is 3.59 $\text{MPa}^{1/2}$). However, both PMOL and IBU are miscible with VA64.

Furthermore, to determine the solubility parameters more precisely a two-dimensional approach introduced by Bagley *et al.*²⁵ can be also used to predict drug–polymer miscibility (Table 1). Previous studies showed that, even though Van-Krevelen method has widely been used to screen drug/polymer miscibility, the estimated solubility parameters were not always accurate. For example, Maniruzzaman *et al.*²⁰ reported a study where the difference of the calculated partial solubility parameters of the model drug PMOL and the VA64 polymer was less than $7 \text{ MPa}^{1/2}$. However, PMOL/VA64 were found immiscible and the solid-state characterisation (e.g. DSC, XRPD) showed that PMOL existed as separate amorphous-amorphous phase from the polymer after extrusion. In the same study the Bagley approach provided a more accurate estimation of the drug-polymer miscibility complementing the findings from the solid-state analysis.

Therefore, the two – dimensional approach is used to determine the drug – polymer miscibility by calculating the distance ($R_{a(v)}$) using the Pythagorean Theorem²⁰. By this approach two components are considered miscible when $R_{a(v)} \leq 5.6\text{MPa}^{1/2}$. Table 1 shows that the δ_h values of the polymers are substantially different compared to the δ_p and δ_d . However the distances calculated for DPD/L100, DPD/L100-55, PRP/L100 and PRP/L100-55 blends indicate that both DPD and PRP are miscible with L100 and L100-55 polymers. Similarly, $R_{a(v)}$ values calculated for Df-Na/L100 ($6.97 \text{ MPa}^{1/2}$) and HCS/L100 ($2.95 \text{ MPa}^{1/2}$)

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3 indicated that higher values for hydrogen bonding have favoured HCS to be miscible with
4 L100 while Df-Na to be immiscible. The estimation of the Van Krevelen/Hoftyzer partial
5 solubility parameter indicates that Df-Na is miscible with L100 but the Bagley parameter
6 completely opposes this prediction. However, both the Van Krevelen/Hoftyzer and Bagley
7 solubility parameters indicate that PMOL is not miscible with EPO.
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12 The Hansen solubility parameter is considered a reliable approach to predict possible
13 drug polymer miscibility, but the limited available data for different group contributions
14 render this approach questionable. It does not also take in account the effect of chain
15 conformation, including branching and linkages between monomer units and the molecular
16 weight (MW) of compounds. These are considered important factors for the solubility of the
17 drug molecule into polymer matrixes and thus in many cases the calculated solubility
18 parameters provide misleading estimations of the drug/polymer miscibility.
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24 **Flory Huggins (F-H) theory for the prediction of drug/polymers interaction parameter.**

25 The Flory–Huggins (F–H) theory is another approach that describes the polymer–drug or
26 polymer–polymer miscibility on the basis of Gibbs free energy change before and after
27 mixing. It assumes that each molecule occupies one site in the lattice and assumes random
28 distribution of the segments³¹. The presence of strong bonds or intermolecular forces (e.g. H-
29 bonding) in solid dispersions often limits the mobility of the chains, which are forced into
30 non-random configurations^{6, 31}. Therefore, the presence of such strong interactions in the
31 systems makes the suitability of F-H theory unsatisfactory as it underestimates those
32 interactions.
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40 In order to determine the F-H interaction parameter between the investigated drugs–
41 polymers blends, the heat of fusion and melting peaks of bulk drug substances, polymers and
42 physical blends were determined by thermal analysis. In the DSC thermograms the bulk PRP,
43 DPD, HCS, Df-Na, PMOL and IBU showed sharp melting peaks at 166.65°C, 170.83°C,
44 226.12°C, 289.60°C, 170.00°C and 79.50°C, respectively with heat of fusion / enthalpy (ΔH)
45 values of 126.25 J/g, 124.59 J/g, 114.92 J/g, 161.06 J/g, 137.06 J/g and 151.17 J/g,
46 respectively. The molecular volumes of all drugs and polymers were estimated from the
47 functional group contribution and by using Eqs. 3 & 4 the average value of χ is calculated as
48 shown in Table 2.
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55 The negative values of the interaction parameter in Table 2 indicate that there is a net
56 attraction force (such as dispersion, polar or hydrogen bonding) between the components in a
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binary mixture. Therefore higher absolute values of χ , suggest stronger interactions between drug/polymers blends. In Table 2, it can be seen that L100 facilitates stronger interactions with different drugs compared to those of L100-55. Similarly, PMOL showed a stronger interaction with VA64 compared to that of EPO, which is in agreement with the solubility parameter calculations. As expected, IBU also showed relatively strong interactions with both EPO and VA64.

As mentioned in a previous paragraph, the F-H parameter cannot provide successful estimations when multiple stronger interactions (such as H-bonding) appear in the drug/polymer systems ⁶. Another key limitation is that a range of experimental data is required in order to apply F-H equation, which makes the approach questionable if the data are not accurate ⁸. Furthermore, the F-H interaction parameter cannot reveal the mechanism or site of interaction in the drug-polymer dispersions. Therefore, a new novel method that could provide both the atomistic and quantitative descriptions of drug-polymer interactions is of immense need for the development of solid dispersions for various drug-polymer systems.

Thermal analysis. DSC was used to analyze the glass transition temperature (T_g) of all pure drugs, polymers, drug/polymer binary mixtures and extrudates (Table 3, Supp. Fig. 1-7). Thermal analysis has been used as a valuable tool to predict drug-polymer miscibility ^{19, 32-34}. According to the Gordon–Taylor equation, if a drug and polymer are miscible the mixture will show a single T_g that ranges between the T_g of pure components ²⁰. Thus, the theoretical prediction of Hansen solubility parameters by using Van Krevelen method and F-H interaction parameters was complemented by DSC studies to access drug /polymer miscibility.

The DSC results of bulk drug substances, bulk polymers, physical mixtures and melt extrudates are presented in Table. 3. All physical mixtures exhibited melting peaks of the drug at slightly decreasing melting temperatures due to possible drug/polymer interactions and partial miscibility. In contrast, no drug melting transitions were observed PRP/L100, PRP/L100-55, DPD/L100 and DPD/L100 solid dispersions prepared by HME processing. The solid dispersions exhibited only a single T_g at temperatures between the T_g of the drug and the polymer, which is strong evidence that PRP and DPD are miscible with both L100 and L100-55 polymers. Similar results were also obtained for the other drug/polymer blends with an exception for the PMOL/VA64 extrudates where a phase separation was observed (PMOL and VA64 were present in two distinct amorphous phases).

Molecular Modelling. Molecular modelling has been used extensively to assist in understanding experimental data and to predict the atomic and molecular properties that can provide guidance for experiment design. In the current study, the computation of binding strength between the drug and dimeric form of the polymer was carried out with a QM-based molecular modelling approach using commercial software package Gaussian09. In the QM-based calculations, the total energy of the system is calculated with respect to all atomic coordinates and thus the sum of electronic energy and repulsion energy between the nuclei and electrons. Because all electrons within the system are explicitly considered, the QM approach is capable of characterising with accuracy non-bonded interactions, such as hydrogen bonds and charge-charge interactions. Nonetheless, the present approach focuses on the intrinsic strength of H bonds and charge-charge interactions between the drug molecule and a small fragment of the polymer, which represents predominantly the polar forces present within the complex of drug-polymer carrier (Fig. 1-2). It does not take consideration of the effect of the full polymeric matrix to drug loading and the process of polymer chain swelling to absorb the drug molecules, both of which can also play a part in determining drug-polymer miscibility. Thus, the binding energy obtained from our QM calculations should be an underestimate compared to the free energy estimated using the F-H theory.

A range of polymers (in their dimeric form) and model drugs were selected to study the strength of hydrogen bonding interactions and to probe the feasibility of identifying possible drug – polymer interactions prior the manufacturing of solid dispersions. The drug-polymer interaction patterns were identified by placing the drug molecule within the proximity of the dimer and then energy optimised to a local energy minimum. Based on these optimised configurations of drug-dimer interactions, the binding energy was calculated using the following equation:

$$E_{\text{binding}} = -[E_{\text{complex}} - (E_{\text{drug}} + E_{\text{dimer}})] \quad (5)$$

The term E_{binding} is the electronic energy component and also the dominant component of the enthalpy term (ΔH) for the formation of the drug-dimer complexes. In all of the drug/polymers combinations primarily two different H bonding were detected with the donor-acceptor distance at $\sim 2 \text{ \AA}$. All possible H bonding were shown in dash line in Table 4.

Because the dimers and the drugs interact predominantly through hydrogen bonds, we interpret the calculated binding energy as a reflection of the strength of hydrogen bonds. Overall, the strength of the interactions is dependent on both the type of the donor and

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4 acceptor and the number of hydrogen bonds formed between the drug and the polymer
5 (Figure 3 and Table 4). Comparisons between L100-DPD^(a) and L100-DPD^(b) and between
6 L100-55-DPD^(a) and L100-55-DPD^(b) (superscripts (a) and (b) denote two possible drug-
7 dimer binding orientations) show that, with structurally similar polymers L100 and L100-55,
8 the interactions formed between the tertiary amine and the carboxylate group via a proton
9 (21.1 kcal/mol and 20.0 kcal/mol) are significantly higher than those between the alkoxy (-O)
10 group and the protonated carboxylate group (12.8 kcal/mol and 15.0 kcal/mol). In addition,
11 other than the interactions involved in tertiary amines, the binding energy arising from one
12 hydrogen bond ranges from 12.8 to 15.1 kcal/mol, while the binding energy from two
13 hydrogen bonds ranges from 16.1 to 25.3 kcal/mol. On average, the binding strength of two
14 H-bond interactions can be stronger by ~5 kcal/mol compared to interactions held through
15 one H-bond (Figure 3). Higher binding energy represents more stable drug/polymer
16 intermolecular interactions formed after the extrusion process³⁵. Strong H-bonding
17 interactions predicted by molecular dynamics suggest drug-polymer miscibility as observed
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29 We have also calculated the binding energy between two other polymers, VA64 and
30 EPO with the model drugs PMOL and IBU (Table 4). The results are consistent with those
31 from L100 and L100-55. The binding energy is predicted at 7.0 kcal/mol without hydrogen
32 bond interactions between VA64 and IBU, while it increases to 16.7 kcal/mol when one H
33 bond is formed between VA64 and IBU, and to 23.3 kcal/mol when two H bonds are formed
34 between VA64 and PMOL. It is also noteworthy that again the tertiary amine from EPO and
35 the carboxylate of IBU interacting via a proton forms the strongest interaction (27.9
36 kcal/mol). This could be attributed to the fact that the most characteristic chemical property
37 of amines is their acid neutralizing capacity (act as bases), thus, giving rise to strong binding
38 energy between the amine interacting with the carboxylic acid via a proton³⁵. This is also
39 reflected by comparing between EPO-IBU and VA64-IBU, where EPO-IBU^(a) and VA64-
40 IBU^(b), both held together through one H-bond between the carbonyl (C=O) group and the
41 protonated carboxylate group, exhibit similar binding energy (16.3 kcal/mol and 16.7
42 kcal/mol). When the donor and acceptor type is altered to tertiary amine and the carboxylic
43 acid in EPO-IBU^(b), the binding energy increases to 27.9 kcal/mol. In summary, based on the
44 drug-dimer combinations listed in Table 4, our results indicate that the strength of the
45 interactions is dependent on both the type of the donor and acceptor and the number of
46 hydrogen bonds formed between the drug and the polymer. In general, addition of a hydrogen
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bond between the polymer and the drug may be accompanied by a ~5 to 8 kcal/mol increase in binding energy.

X-ray Photoelectron Spectroscopy (XPS) analysis. XPS analysis was employed to confirm the QM predictions by analysing the extruded solid dispersions of two drug-polymer formulations (PRP/L100, PRP/L100-55, DPD/L100 and DPD/L100-55). The surface elemental ratios of PRP, DPD, polymers and the extruded formulations were determined experimentally by XPS³⁶⁻³⁸ based on various surveys as depicted in Fig.4-5. This was implemented as a comparison with the anticipated theoretical values derived from the structural formula²². According to the surveys the PRP/L100, PRP/L100 55, DPD/L100 and DPD/L100-55 showed the amount of N atoms as 0.63%, 0.23%, 0.53% and 0.21%, respectively simply indicating lower amount of N atom present in the final extruded formulations.

The N (1s) binding energy (BE) of ~402.035 eV (Fig. 6) in PRP and of ~402.6 eV (Fig. 7) in DPD suggests the protonation of the NH⁺ group while the slightly higher values of N (1s) energy (~402.8 eV) in PRP/L100 extrudates represents further protonation effect of N atom as NH₄⁺. This observed N 1s peak at BE= ~402.80 eV is in good agreement with the previously observed protonation of amine group by Beamson and Briggs²². The BE peak at ~402.80 eV (higher than typically observed for amines BE= ~399 eV - 400.5 eV and much more for -NH₂⁺ group) for N1s is an indication of C-O-NH₂⁺ structure whereas the O atom peak at ~534.40 eV shows the same^{39, 40}. These results strongly indicate H-bonding interactions between the amine group of the API and ester/carboxyl group of the polymer (L100) (Fig. 6).

Similarly, N 1s peaks from PRP/L100-55 and DPD/L100-55 also complement the observations from PRP/L100 and DPD/L100 formulations. The N (1s) energy of ~402.9 eV in DPD/L100 formulation suggests protonation of the amine group as observed for aforementioned PRP/L100 formulation. The BE peak at ~402.90 eV (Fig. 7) for N1s is an indication of C-O-NH₂⁺ structure with longer peak shift than that of PRP/L100. As before, we concluded that a strong interaction between the amine group of API and ester/carboxyl group of polymer through H-interactions has taken place⁴⁰⁻⁴².

Furthermore, the calculation of the N-coefficient for all extruded formulations indicated the strength of the intermolecular interactions within the solid dispersions. The calculated N coefficients for all extruded formulations are summarized in Table 5. The values

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in Table 5 clearly show that the N coefficient values of the active L100 extrudates are smaller than those of L100-55 extrudates. It is accepted that the lower the N-coefficient, the higher the amount of protonated N atoms in cationic drugs and thus the stronger the interaction between polymer and APIs.

CONCLUSIONS

In this study, we exploited the use of quantum mechanical (QM) calculations to predict drug-polymer interactions by using the drug molecule and the dimeric form of the polymer. In general, addition of a hydrogen bond between the polymer and the drug is accompanied by ~5 to 8 kcal/mol increase in binding energy. The strongest interactions however are between the tertiary amines and the carboxylate groups via a proton, with their binding energy ranging from 20.0 kcal/mol to 27.9 kcal/mol. The findings from XPS analysis confirmed the site of interaction and the strength of H-bonding between the carboxyl group of anionic methacrylate co-polymers and the amine group of the active substances. The prediction of strong interactions for some of the drug-polymer blends, from the QM calculations, were in agreement with the DSC experimental findings, which proved the presence of the molecular solid dispersions of the same blends. The QM-based calculations enable the study of drug-polymer interactions with great detail at the atomistic level and are particularly suited for characterising drug-polymer interactions that contain strong hydrogen bonds or charge-charge interactions. Thus QM-based molecular modelling can broadly be applied for the development of pharmaceutical solid dispersions where traditional methods such as Van Krevelen solubility parameter or F-H parameter present a limited capacity.

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Table 1: Estimated solubility parameters of different drugs and polymers using Van Krevelen /Hoftyzer and Bagley methods.

Comp.	δ_p (MPa ^{1/2})	δ_d (MPa ^{1/2})	δ_h (MPa ^{1/2})	δ_v (MPa ^{1/2})	δ (MPa ^{1/2})	Van Krevelen ($\Delta\delta$)				Bagley ($R_a(v)$)			
						$\Delta\delta$ L100 (MPa ^{1/2})	$\Delta\delta$ L100- 55 (MPa ^{1/2})	$\Delta\delta$ EPO (MPa ^{1/2})	$\Delta\delta$ VA64 (MPa ^{1/2})	$R_a(v)$ L100	$R_a(v)$ L100-55	$R_a(v)$ EPO	$R_a(v)$ VA64
PRP	3.67	19.30	9.90	19.64	21.94	0.81	0.29	-	-	3.26	4.67	-	-
DPD	4.05	16.39	5.44	16.89	17.75	5.0	3.9	-	-	3.59	4.22	-	-
Df-Na	3.32	24.65	7.82	24.87	26.07	3.32	-	-	-	6.97	-	-	-
HCS	8.02	19.04	14.65	20.67	25.34	2.59	-	-	-	2.95	-	-	-
PMOL	7.93	19.67	13.96	21.20	25.39	-	-	6.49	5.79	-	-	13.16	12.17
IBU	5.76	18.47	10.04	19.35	21.80	-	-	2.90	2.20	-	-	4.21	5.87
L100	0.41	19.31	12.03	19.31	22.75	-	-	-	-	-	-	-	-
L100-55	0.25	18.22	11.69	18.22	21.65	-	-	-	-	-	-	-	-
EPO	0.65	17.89	6.08	17.90	18.90	-	-	-	-	-	-	-	-
VA64	18.0	0.64	18.01	7.73	19.60	-	-	-	-	-	-	-	-

$$R_{a(v)} = \sqrt{(\delta_{v2} - \delta_{v1})^2 + (\delta_{h2} - \delta_{h1})^2}$$

Table 2: Calculation of F–H interaction parameter of different Drug–Polymer extruded formulations.

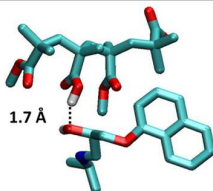
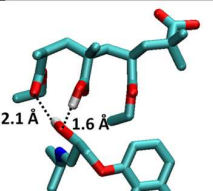
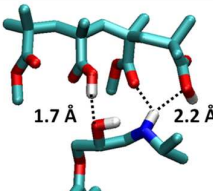
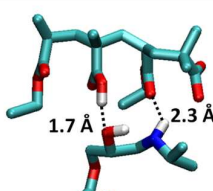
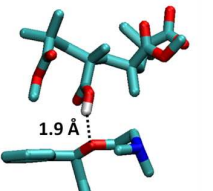
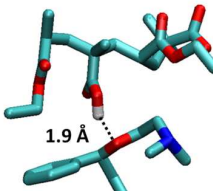
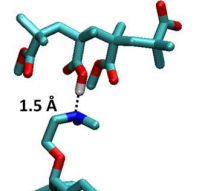
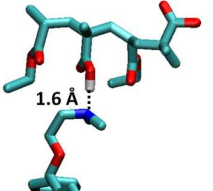
Formulations	Volume Fractions (ϕ)	Nishi-Wang ($-\chi$)	Hildebrand - Scott ($-\chi \times 10^{-4}$)
PRP/L100	0.1:0.9	0.22	2.89
PRP/L100-55	0.1:0.9	0.08	0.37
DPD/L100	0.1:0.9	0.20	1.10
DPD/L100-55	0.1:0.9	0.07	0.67
HCS/L100	0.1:0.9	0.21	1.08
Df-Na/L100	0.3:0.7	0.01	0.15
PMOL/VA64	0.5:0.5	2.12	1.60
PMOL/EPO	0.5:0.5	0.38	0.20
IBU/EPO	0.4:0.6	0.81	0.35
IBU/VA64	0.4:0.6	0.61	0.20

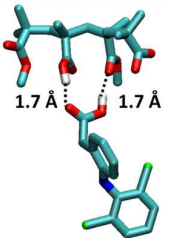
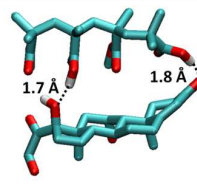
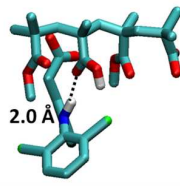
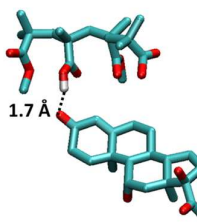
Table 3: Thermal transitions of pure drugs, polymers and binary mixtures.

	Glass transition (°C)		Melting endotherm/ Enthalpy (°C /ΔH, Jg⁻¹)
PRP	36.22		166.65/126.25
DPD	14.13		170.82/124.59
Df-Na	-		289.61/161.06
HCS	82.45		226.12/114.92
PMOL	24.55		168.84/33.40
IBU	-		77.9/121.30
L100	164.38		-
L100-55	83.97		-
EPO	48.38		-
VA64	105.0		-
Physical mixtures (PM) and extruded formulations (EXT)			
	PM (°C)	EXT (°C)	PM (°C)
PRP/L100	98.82	74.84	162.41
PRP/L100-55	71.16	63.36	153.62
DPD/L100	114.46	76.56	139.19
DPD/L100-55	74.15	62.11	153.62
PMOL/VA64*	102.22	27.57, 93.47	131.32
PMOL/EPO	79.38	56.76	159.96
IBU/VA64	54.83	29.49	76.6
IBU/EPO	57.40	15.57	76.35

* Two Tgs were observed for the extruded blend due to phase separation.

Table 4: The calculated drug-dimer binding energy using Gaussian09 (energy in kcal/mol). The top table is for the dimeric forms of polymer L100 and L100-55 and model drugs PRP and DPD, while the bottom table is for the dimeric forms of polymer VA64 and EPO and model drugs PMOL and IBU. Superscripts (a) and (b) denote two possible drug-dimer binding orientations, which were determined by positioning the drug molecule in two different orientations (based on its H-bond donor/acceptor groups) relative to the position of the dimer prior to the QM optimisation.

Drug-Poly Comb.	Conformation	Energy	Drug-Poly Comb.	Conformation	Energy
L100-PRP ^(a)		14.5	L100-55-PRP ^(a)		17.8
L100-PRP ^(b)		16.1	L100-55-PRP ^(b)		16.9
L100-DPD ^(a)		12.8	L100-55-DPD ^(a)		15.0
L100-DPD ^(b)		21.1	L100-55-DPD ^(b)		20.0

L100-Df-Na ^(a)		19.4	L100-HCS ^(a)		25.3
L100-Df-Na ^(b)		15.1	L100-HCS ^(b)		15.1

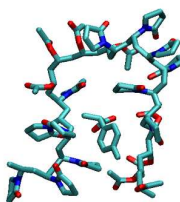
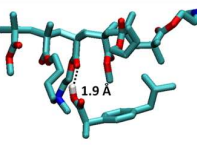
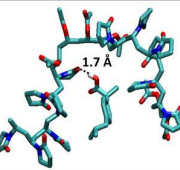
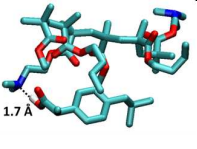
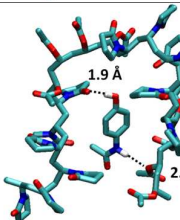
Drug-Poly Comb.	Conformation	Energy	Drug-Poly. Comb.	Conformation	Energy
VA64-IBU ^(a)		7.0	EPO-IBU ^(a)		16.3
VA64-IBU ^(b)		16.7	EPO-IBU ^(b)		27.9
VA64-PMOL		23.3			

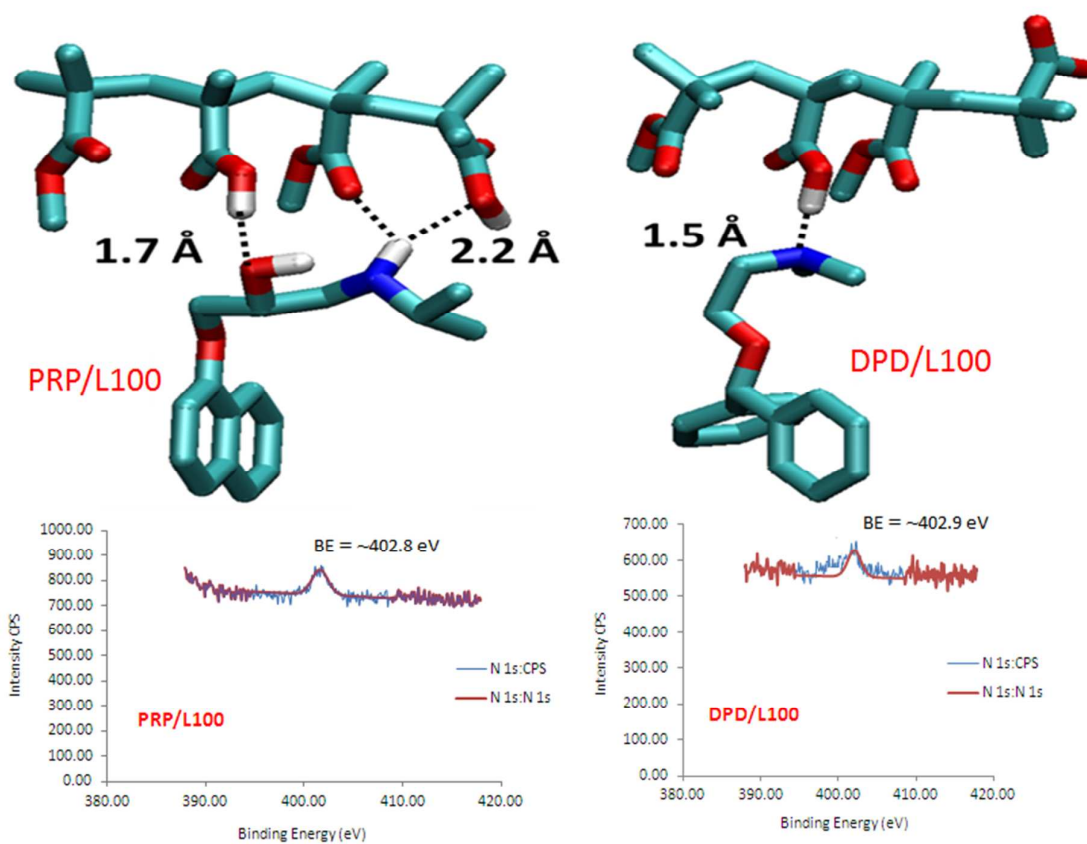
Table 5: N coefficients of the extruded formulations

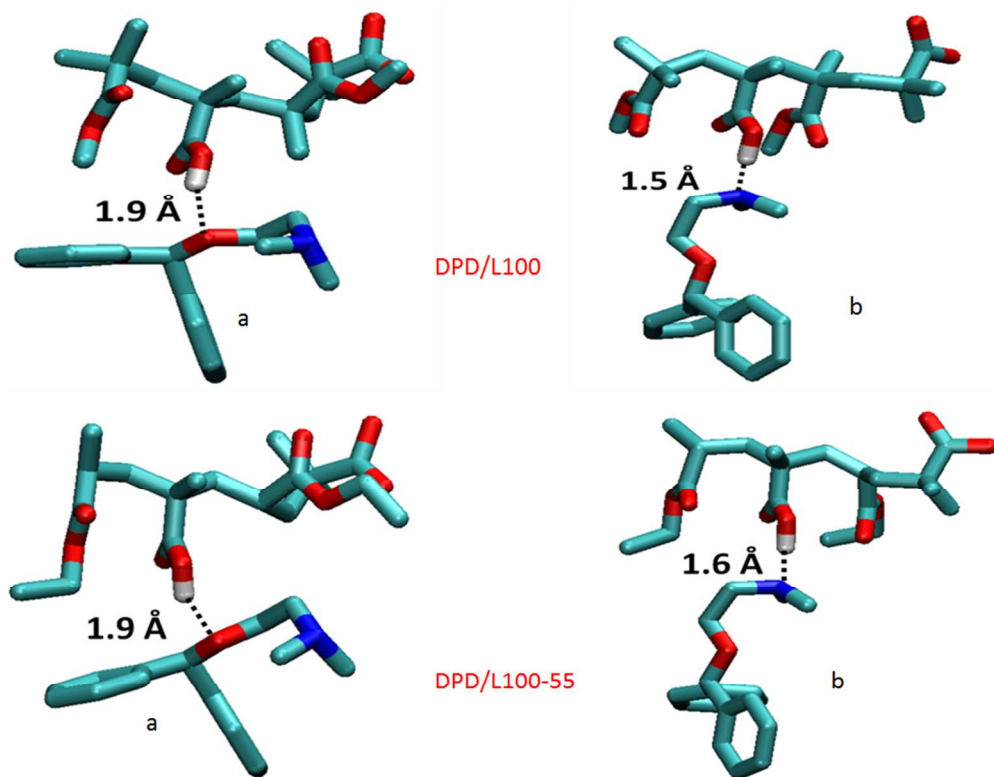
Formulations	N 1s Binding Energy		Calculated N Coefficient
	1 st Fitting (eV)	2 nd Fitting (eV)	
PRP/L100	63	100	0.63
PRP/L100-55	58.51	49.49	1.18
DPD/L100	45.97	54.03	0.85
DPD/L100-55	51.19	48.81	1.05

Figures caption list

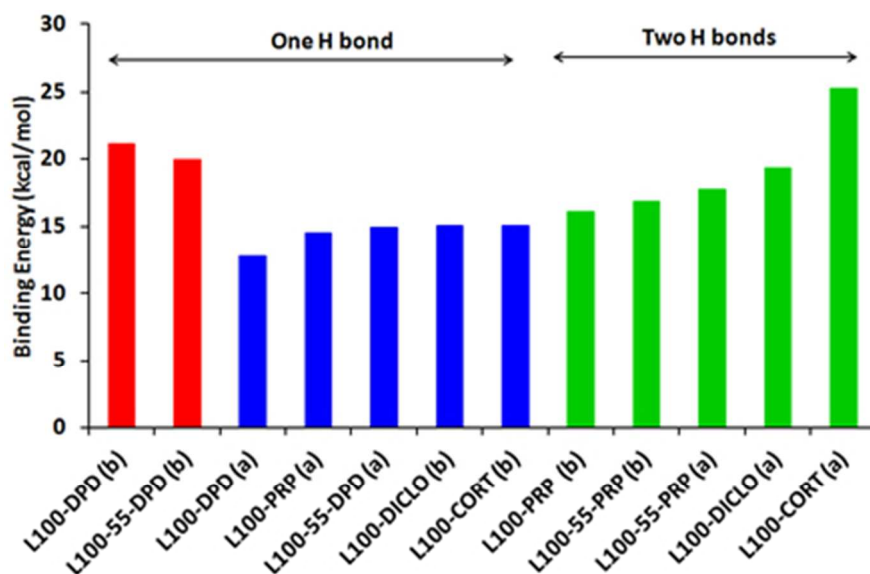
- Fig. 1** Molecular modelling of PRP and L100 or L100-55 pairs (Gaussian09).
- Fig. 2** Molecular modelling of DPD with different polymers (energy optimised to local minima at the M06-2x/6-31G** level and verified by frequency calculations using Gaussian09).
- Fig. 3** Binding energy as arranged based on the number of hydrogen bonds (red and blue columns for one H bond while green for two H bonds) between the dimeric structure of the polymer and the model drugs. Red columns represent the binding energy involved in tertiary amines.
- Fig. 4** XPS surveys of pure PRP, DPD, L100 and L100-55 showing BE peaks of C 1s, O 1s, N 1s and Cl 2p atoms and their area ratios.
- Fig. 5** XPS surveys of DPD and PRP based extruded formulations with L100 and L100-55 polymers (C 1s, O 1s, N 1s and Cl 2p atoms and their area ratios).
- Fig. 6** Molecular structure of PRP and N 1s BE peaks of PRP and extruded formulations (from the peak fitting using CasaXPS™ - Version 2.3.15).
- Fig. 7** DPD molecular structure and its N 1s BE peaks with DPD/L100 and DPD/L100-55 extruded formulations (peak fitting performed using CasaXPS™ - Version 2.3.15).

Graphical Abstract

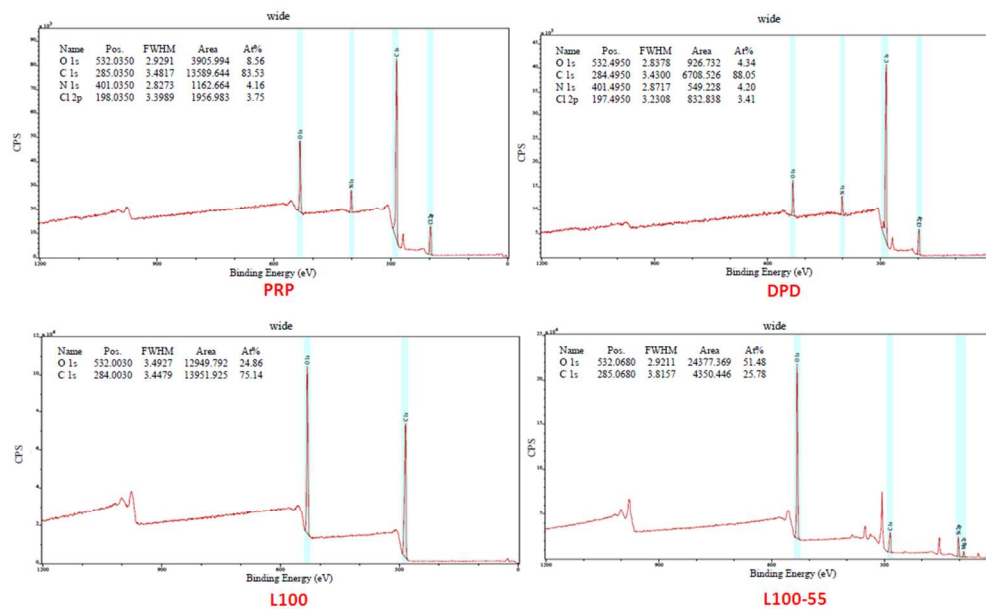




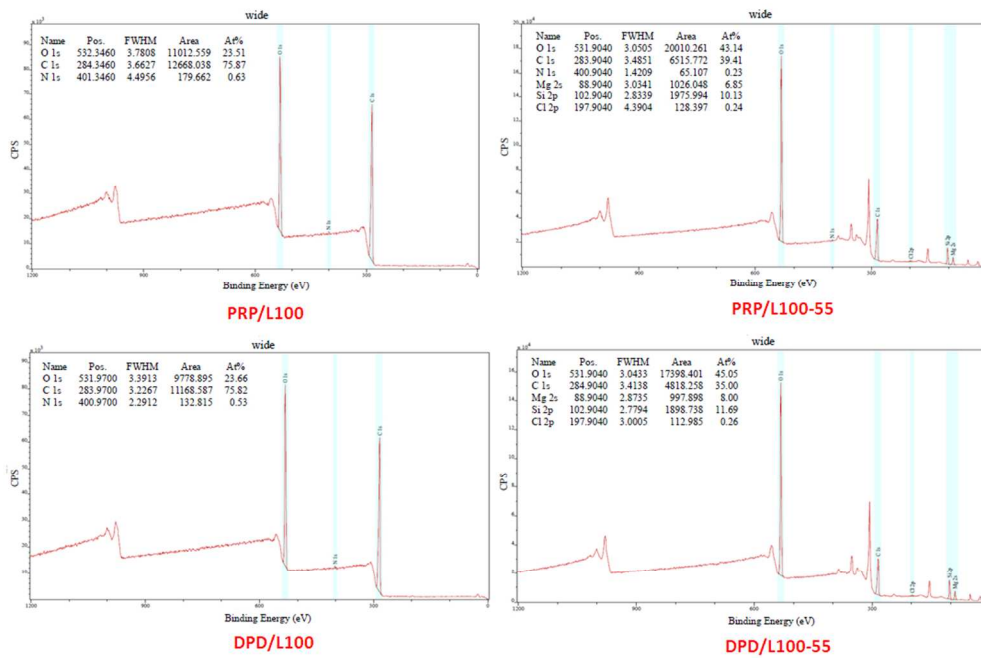
Molecular modelling of DPD with different polymers (energy optimised to local minima at the B3LYP/6-31G* level and verified by frequency calculations using Gaussian09).
260x203mm (96 x 96 DPI)



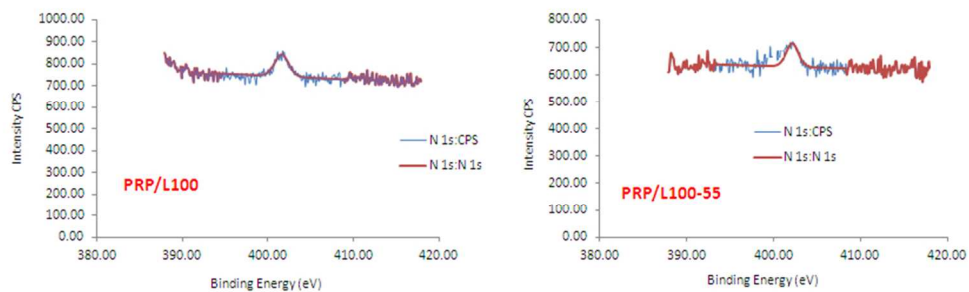
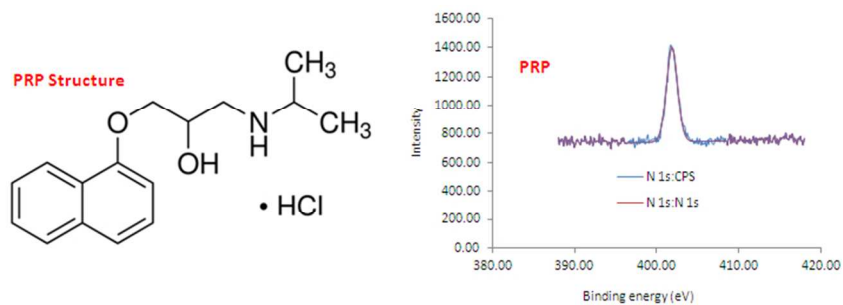
Binding energy as arranged based on the number of hydrogen bonds (red and blue columns for one H bond while green for two H bonds) between the dimeric structure of the polymer and the model drugs. Red columns represent the binding energy involved in tertiary amines.
118x78mm (96 x 96 DPI)



XPS surveys of pure PRP, DPD, L100 and L100-55 showing BE peaks of C 1s, O 1s, N 1s and Cl 2p atoms and their area ratios.
288x174mm (96 x 96 DPI)

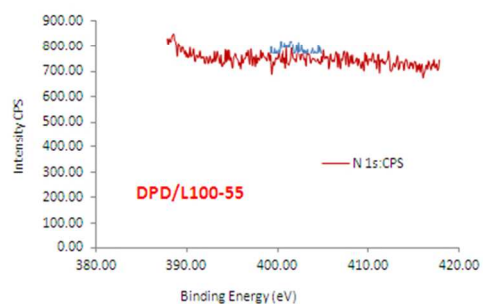
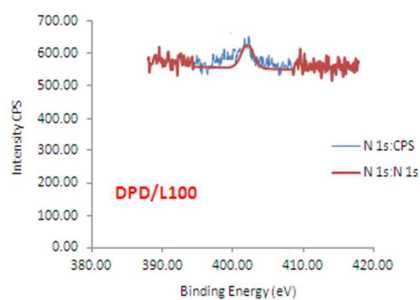
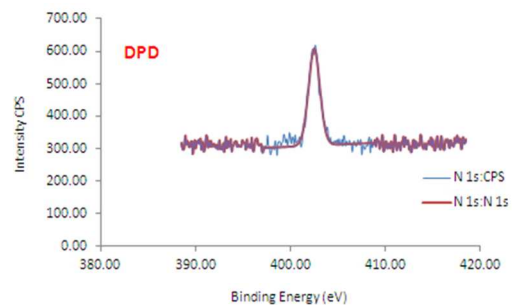
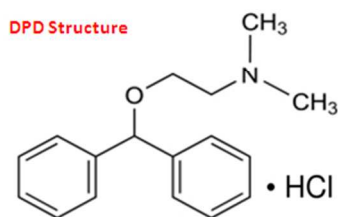


XPS surveys of DPD and PRP based extruded formulations with L100 and L100-55 polymers (C 1s, O 1s, N 1s and Cl 2p atoms and their area ratios).
288x194mm (96 x 96 DPI)



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Molecular structure of PRP and N 1s BE peaks of PRP and extruded formulations (from the peak fitting using CasaXPSTM - Version 2.3.15).
225x139mm (96 x 96 DPI)



DPD molecular structure and its N 1s BE peaks with DPD/L100 and DPD/L100-55 extruded formulations (peak fitting performed using CasaXPSTM - Version 2.3.15).
215x140mm (96 x 96 DPI)