Solid dispersions: a technology for improving bioavailability

Abstract

An unending challenge in pharmaceutical industry is related to poor solubility of maximum drugs. To overcome this problem various technologies have been developed but none appears to be a promising one. Solid dispersion is a solubilization technology emphasizing basically on drug-polymer two component systems in which drug dispersion and its stabilization is the key for formulation development. Therefore this technology has been realized as extremely useful tool in improving the dissolution properties of poorly water-soluble drugs and in recent years, a great deal of knowledge has been accumulated about solid dispersion, but their commercial application is limited. This review summarizes our current understanding of various methods used for the preparation of solid dispersions emphasizing the elementary aspects of this significant technology.

Keywords: solid dispersions, poorly soluble drug, solid solution, amorphous state, bioavailability

Introduction

Modern drug discovery techniques, with advances in combinatorial chemistry and high throughput screening, continue to fill drug development pipelines with a high number of poorly soluble New Chemical Entities (NCEs). It is estimated that over the years about 40%–70% of NCEs are poorly water soluble and large number of scientists are engage in invention of NCEs and the success rate is poor. A drug with poor aqueous solubility will typically exhibit dissolution rate limited absorption, and a drug with poor membrane permeability will typically exhibit permeation rate limited absorption. Hence, two areas of pharmaceutical research that focus on improving the oral bioavailability of active agents include: enhancing solubility and dissolution rate of poorly water soluble drug and enhancing permeability of poorly permeable drugs.1

The main possibilities for improving dissolution is to increase the surface area available for dissolution by decreasing the particle size of the solid compound and/or by optimizing the wetting characteristics of the compound surface, to decrease the boundary layer thickness, to ensure sink conditions for dissolution and, last but definitely not least, to improve the apparent solubility of the drug under physiologically relevant conditions. Of these possibilities, changes in the hydrodynamics are difficult to invoke in vivo and the maintenance of sink conditions will depend on how permeable the gastrointestinal mucosa is to the compound as well as on the composition and volume of the luminal fluids. Although some research effort has been directed towards permeability enhancement using appropriate excipient, results to date have not been particularly encouraging.2

Administration of the drug in the fed state may be an option to improve the dissolution rate and also to increase the time available for dissolution; the likely magnitude of the food effect can be forecasted from dissolution tests in biorelevant media.

In the Biopharmaceutical Classification System (BCS) drugs with low aqueous solubility and high membrane permeability are categorized as Class II drugs. Therefore, solid dispersion technologies are particularly promising for improving the oral absorption and bioavailability of BCS Class II drugs.3 The basic principle involved in enhancing the poor solubility of drug with solid dispersion includes complete removal of drug crystalline structure and its molecular dispersion in a hydrophilic polymeric carrier.

When the solid dispersion is exposed to aqueous media, the carrier dissolves and the drug releases as fine colloidal particles. This increases surface area of dissolution rate and hence bioavailability of poorly water soluble drugs. Drug in soluble hydrophilic carrier improves the dissolution rate by reducing particle size and increasing the particle porosity. Therefore by improving the drug release profile of these drugs, it is possible to enhance their bioavailability and reduce side effects.4

Advantages of solid dispersion

Improving drug bioavailability by changing their water solubility has been possible by chemical or formulation approaches. Chemical approaches to improving bioavailability without changing the active target can be achieved by salt formation or by incorporating polar or ionizable groups in the main drug structure, resulting in the formation of a pro-drug. Solid dispersions appear to be a better approach to improve drug solubility than these techniques, because they are easier to produce and more applicable.5

In molecular dispersions, solid dispersions represent the last state on particle size reduction, and after carrier dissolution the drug is molecularly dispersed in the dissolution medium. Solid dispersions apply this principle to drug release by creating a mixture of a poorly water soluble drug and highly soluble carriers. A high surface area is formed, resulting in an increased dissolution rate and, consequently, improved bioavailability.6 Solid dispersions also provides particles with improved wettability as it was observed that even carriers without any surface activity, such as urea improved drug wettability. Carriers with surface activity, such as cholic acid and bile salts, when used, can significantly increase the wettability property of drug.

Particles in solid dispersions have been found to have a higher degree of porosity. The increase in porosity also depends on the carrier properties; for instance, solid dispersions containing linear polymers
produce larger and more porous particles than those containing reticular polymers and, therefore, result in a higher dissolution rate. The increased porosity of solid dispersion particles also hastens the drug release profile.9

Poorly water soluble crystalline drugs, when in the amorphous state tend to have higher solubility. The enhancement of drug release can usually be achieved using the drug in its amorphous state, because no energy is required to break up the crystal lattice during the dissolution process. In solid dispersions, drugs are presented as supersaturated solutions after system dissolution, and it is speculated that, if drugs precipitate, it is as a metastable polymorphic form with higher solubility than the most stable crystal form. For drugs with low crystal energy (low melting temperature or heat of fusion), the amorphous composition is primarily dictated by the difference in melting temperature between drug and carrier.9 For drugs with high crystal energy, higher amorphous compositions can be obtained by choosing carriers, which exhibit specific interactions with them. Drug is formulated with hydrophilic carrier as a solid dispersion to increase its aqueous solubility and dissolution. Then supersolventant (e.g. croscarmellose sodium) is used in tablet formulation to achieve rapid disintegration of tablets prepared by wet granulation method. These rapidly disintegrating tablets can be used as an alternative to parenteral therapy enabling patient for self-medication even without the aid of water.9

Disadvantages of solid dispersions

Despite extensive expertise with solid dispersions, they are not broadly used in commercial products, mainly because of the possibility of the change of amorphous state to crystalline state during processing (mechanical stress) or storage (temperature and humidity stress).

The effect of moisture on the storage stability of amorphous pharmaceuticals is also a significant concern, because it may increase drug mobility and promote drug crystallization. Moreover, most of the polymers used in solid dispersions can absorb moisture, which may result in phase separation, crystal growth or conversion from the amorphous to the crystalline state or from a metastable crystalline form to a more stable structure during storage. This may result in decreased solubility and dissolution rate. Therefore, exploitation of the full potential of amorphous solids requires their stabilization in solid state, as well as during in vivo performance.

Another drawback of solid dispersions is their poor scale-up for the purposes of manufacturing. Strategies to overcome the manufacturing process drawbacks will be discussed later.10

Classification of solid dispersions

Depending on the molecular arrangement, solid dispersions can be of the following types:

**Eutectic mixtures**: solid eutectic mixtures are usually prepared by rapidly cooling the co-melt of the two components in order to obtain a physical mixture of very fine crystals of the two components.

**Solid solutions**: Depending on the miscibility, the two types of solid solutions are:

Continuous solid solutions: In continuous solid solutions, the components are miscible in all proportions i.e. the bonding strength between the components is stronger than the bonding between the individual component.

Discontinuous solid solutions: In discontinuous solid solutions, the solubility of each of the component in the other component is limited in nature.11

**Solid solutions**

Depending on the distribution of the solvates in the solvent, solid solutions can be of two types:

**Substitution crystalline solution**: These are solid solutions which have a crystalline structure, the solute molecules substitute for the solvent molecules in the crystal lattice.

**Intersticial crystalline solid solution**: These are those solid solutions in which the dissolved molecules occupy the interstitial spaces between the solvent molecules in the crystal lattice.

**Amorphous solid solutions**: In amorphous solid solutions, the solute molecules are dispersed molecularly but irregularly within the amorphous solvent.

**Glass solutions and glass suspension**: A glass solution is a homogenous system in which the solute dissolves in the glassy solvent. The glassy state is characterized by transparency and brittleness below the glass transition temperature. The term glass refers to a pure chemical or a mixture of pure chemicals in the glassy state.12

**Classification of solid dispersion on the basis of recent advancement**

**First generation solid dispersion**: These solid dispersions are prepared by using crystalline carriers. Urea and sugars were the first crystalline carriers that were used in the preparation of solid dispersions. These have a disadvantage of being thermodynamically unstable and they do not release drug at a faster rate.

**Second generation solid dispersion**: These solid dispersions are prepared using amorphous carriers instead of crystalline carriers. The drug is molecularly dispersed in the polymeric carrier. The polymeric carriers are divided into two groups:

a. **Synthetic polymer**—povidone, polyethylene glycols and polyethylene glycol (PEG)

b. **Natural polymers**—hydroxypropylmethylcellulose, ethyl cellulose, starch derivatives like cycloextrin.

**Third generation solid dispersion**: These solid dispersions contain a surfactant carrier, or a mixture of amorphous polymers and surfactants as carriers. These achieve the highest degree of bioavailability for the drugs that are having poor solubility. The surfactants being used in the third generation solid dispersion are such as inulin, poloxamer 407 etc.15

**Polymers**

A wide variety of polymers are available which have tremendous potential in the area of solid dispersions.

**Polyethylene glycol (PEG)**

These are compounds that are obtained from a reaction of ethylene glycol with ethylene oxide. PEGs whose molecular weight is above 300000 are commonly termed as polyethylene oxides. For the manufacture of solid dispersions and solutions, PEGs with molecular weights of 1500-20,000 are usually employed. As the MW rises, so...
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Cyclodextrins are primarily used to enhance the release rate of griseofulvin. Organic acids and their derivatives: Organic acids such as succinic acid and citric acid have also been used as carriers in solid dispersions, originally to enhance the release rate of griseofulvin. Sugar, polyols and their polymers: Although sugars and related compounds are highly water soluble and have few, if any, toxicity issues, they are less suitable than other carriers for the manufacture of solid dispersions. The melting point of most sugars is high, making preparation by the hot melt method problematic, and their solubility in most organic solvents is poor, making it difficult to prepare coevaporates. Even with these drawbacks, several attempts have been reported to prepare solid dispersions using sugars and their derivatives. Mannitol, which has a melting point of 165-168°C and decomposes only above 250°C, can be employed in some cases to prepare dispersions by the hot melt method.

Organic acids and their derivatives: Organic acids such as succinic acid and citric acid have also been used as carriers in solid dispersions, originally to enhance the release rate of griseofulvin method. Cycloextrinsics: Cycloextrinsics are primarily used to enhance solubility, chemical protection, taste masking and improved handling by the conversion of liquids into solids by entrapment. Advantages of Cycloextrinsics:

a. Increasing the stability of the drug
b. Release profile during gastrointestinal transit

d. Release site and time profile
e. Decreasing local tissue irritation.
f. Masking unpleasant taste.

Methods for preparing solid solutions

There are various manufacturing methods for solid dispersions that have been reported in literature. These are given below in Figure 1.

Kneading technique

In this method, carrier is permeated with water and transformed to paste. Drug is then added and kneaded for particular time. The kneaded mixture is then dried and passed through sieve if necessary.

Solvent method

This method is also known as solvent evaporation method in which physical mixture of the drug and the carrier is dissolved in common solvent and is evaporated until a clear solvent free film is obtained. The main advantage is that the thermal decomposition of the drug or the carrier can be prevented because the organic solvents require a low temp for evaporation. The disadvantage in this method is difficulty in removing the solvent and higher cost of preparation.
**Co-precipitation method**

Required amount of drug is added to the solution of carrier. The system is kept under magnetic agitation and protected from the light. The formed precipitate is separated by vacuum filtration and dried at room temperature in order to avoid the loss of the structure water from the inclusion complex.\(^1\)

**Melting method**

The melting method is suitable for heat stable materials with low melting points. The basic principle of the method consists of melting together the drug and carrier at a temperature slightly above their eutectic point, mixing the liquefied components. It is then cooled to acquire a congealed mass. It is crushed and sieved.\(^2\)

Ex. albendazole and urea solid dispersion was prepared by this method.

**Co-grinding method**

Physical mixture of drug and carrier is mixed for some time employing a blender at a particular speed. The mixture is then charged into the chamber of a vibration ball mill steel balls are added. The powder mixture is pulverized. Then the sample is collected and kept at room temperature in a screw capped glass vial until use. Ex. chlordiazepoxide and mannitol solid dispersion was prepared by this method.\(^3\)

**Spray-drying method**

Drug is dissolved in suitable solvent and the required amount of carrier is dissolved in water. Solutions are then mixed by sonication or other suitable method to produce a clear solution, which is then spray dried using spray dryer.\(^22\)

**Lyophilization technique**

Freeze-drying involves transfer of heat and mass to and from the product under preparation. This technique was proposed as an alternative method to solvent evaporation. Lyophilization has been thought of a molecular mixing technique where the drug and carrier are co dissolved in a common solvent, frozen and sublimed to obtain a lyophilized molecular dispersion.\(^24\)

**Electrospinning method**

In this technique electric force is used to withdraw a nano size fibre thread from the polymer sol/polymer melt. This a combination of solid dispersion with nanotechnology use in polymer industry. Stream of Polymer solution /melt is subjected to electric force (5 to 30kv) which cause body of the liquid becomes charged, and electrostatic repulsion counteracts the surface tension. This made a strong cohesive force between the particle or droplets of polymer and a stream of fibre is formed. Then thinning and stretching of fibre to nano diameter is done by using whipping process called electrostatic repulsion lead to formation of uniform fibre in nano diameter. This process all depend on rate of feeding surface tension and electric force used.\(^25\)

**Dropping method solution**

The dropping method, developed to facilitate the crystallization of different chemicals, is a new procedure for producing round particles from melted solid dispersions. This technique may overcome some of the difficulties inherent in the other methods. For laboratory-scale

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**Figure 1** Different strategy for planning of strong scatterings.
preparation, a solid dispersion of a melted drug-carrier mixture is pipette and then dropped onto a plate, where it solidifies into round particles. The use of carriers that solidify at room temperature may aid the dropping process. The dropping method not only simplifies the manufacturing process, but also gives a higher dissolution rate. It does not use organic solvents and, therefore, has none of the problems associated with solvent evaporation.

**Melt extrusion method**

Solid dispersion by this method is composed of active ingredient and carrier, and prepare by hot-stage extrusion using a co-rotating twin-screw extruder. The concentration of drug in the dispersions is always 40% (w/w). Melt extrusion technique is used in the preparation of diverse dosage forms in the pharmaceutical industry e.g. sustained-release pellets.

**Melt agglomeration process**

This technique has been used to prepare Solid Dispersion where the binder acts as a carrier. SD(s) are prepared either by heating the binder, drug and excipient to a temperature above the melting point of the binder or by spraying a dispersion of drug in molten binder on the heated excipient by using a high shear mixer. A rotary processor has been shown to be alternative equipment for melt agglomeration because of easier control of the temperature and because higher binder content can be incorporated in the agglomerates.

**Super critical fluid (SCF) technology**

SCF is a substance above its critical temperature and pressure. Critical point represents the highest temperature and pressure at which the substance exists as vapour and liquid in equilibrium. In this technique SCF is used to form solid dispersion of insoluble material/polymer with drug cause increase in dissolution property. It is superior over conventional technique(spray drying, hot melt etc.), in this technique SCF carbon dioxide is mainly used which cause very rapid precipitation of solid mixture giving no time for separation of drug and polymer in preparation of solid dispersion. It form very stable small particle with higher surface area for good flow and low organic solvent residual. In recent Solid dispersion of carbamazepine with PEG-4000 are made using SCF carbon dioxide in precipitation vessel. Resulting in formation of carbamazepine with increase rate and extent of dissolution with low solvent residual.

**Characterization of solid dispersion**

A combination of two or more techniques is required to study its complete picture.

**Drug-carrier miscibility**

- Hot stage microscopy
- Differential scanning calorimetry
- Powder X-ray diffraction
- NMR 1H Spin lattice relaxation time

**Drug carrier interactions**

- FT-IR spectroscopy
- Raman spectroscopy
- Solid state NMR

**Physical Structure**

- Scanning electron microscopy
- Surface area analysis
- Surface properties
- Dynamic vapor sorption
- Inverse gas chromatography
- Atomic force microscopy
- Raman microscopy

**Amorphous content**

- Polarised light optical microscopy
- Hot stage microscopy
- Humidity stage microscopy
- DSC (MTDSC)
- ITC
- Powder X-ray diffraction

**Stability**

- Humidity studies
- Isothermal Calorimetry
- DSC (Tg, Temperature recrystallization)
- Dynamic vapor sorption
- Saturated solubility studies

**Dissolution enhancement**

- Dissolution
- Intrinsic dissolution
- Dynamic solubility
- Dissolution in bio-relevant media

Powder X-ray diffraction can be used to qualitatively to detect the presence of crystalline forms in solid dispersion and to determine factors influencing recrystallization during storage stability studies. Sharper diffraction peaks indicate more crystalline material. In Infrared spectroscopy (IR) you can detect the variation in the energy distribution of interactions between drug and matrix. Sharp vibrational bands indicate crystallinity. Fourier Transformed Infrared Spectroscopy (FTIR) was used to accurately detect crystallinity ranging from 1 to 99% in pure material.

Water vapor sorption can be used to discriminate between amorphous and crystalline material when the hygroscopicity is different. This method requires accurate data on the hygroscopicity of both completely crystalline and completely amorphous samples. Dissolution calorimetry measures the energy of dissolution, which is dependent on the crystallinity of the sample. Usually, dissolution of crystalline material is endothermic, whereas dissolution of amorphous material is exothermic.

Macroscopic techniques that measure mechanical properties that
are different amorphous and crystalline material can be indicative for the degree of crystallinity. Density measurements and Dynamic Mechanical Analysis (DMA) determine the modulus of elasticity for and viscosity and thus affected by the degree of crystallinity. However, also these techniques require knowledge about the additivity of these properties in intimately mixed binary solids.29 In Differential Scanning Calorimetry (DSC) samples are heated with a constant heating rate and the amount of energy necessary for that is detected. With DSC the temperatures at which thermal events occur can be detected. Thermal events can be a glass to rubber transition, (re)cristallization, melting or degradation. Furthermore, the melting- and (re)cristallization energy can be quantified. The melting energy can be used to detect the amount of crystalline material.

In vitro dissolution studies

In vitro dissolution studies are done for the find out dissolution behavior. The in-vitro dissolution study can be used to demonstrate the bioavailability or bioequivalence of the drug product through in vitro-in vivo correlation (IVIVC). On the other hand if absorption of the drug is dissolution rate limited that means the drug in the gastrointestinal fluid passes freely through the bio-membranes at a rate higher than it dissolves or is released from the dosage form. The specifically designed in-vivo dissolution study will be required in solid dispersion system to access the absorption rate, and hence its bioavailability and to demonstrate the bioequivalence ultimately.

There are some apparatus used in United States pharmacopoeia for dissolution testing these are following.31

Conclusion

As with increasing number of poorly soluble drug candidates the need for improvements in the drug manufacturing technology also increases. Therefore solid dispersion technology seems to be a promising solution for improving the dissolution characteristics of such drugs. Aspects that still need to be addressed in the next years include further improvements in manufacturing on a large scale, and better predictions of whether a particular drug/carrier combination will lead to a true solid solution or to a partly crystalline dispersion as well as whether the dispersion will remain physically stable during further processing and storage. Last but not least, although this article has been devoted to the use of solid dispersions for the improvement of the release rate and oral bioavailability, by judicious choice of the carrier it is also possible to delay or slow down the release pattern of a drug by formulating it as a solid dispersion. The availability of a wide variety of polymers that are themselves poorly soluble or which swell under aqueous conditions suggests that solid dispersions have tremendous potential in the area of controlled release dosage forms.

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Conflicts of interest

The author declares that there is no conflict of interest.

References


