Pharmacokinetic aspects and *in vitro–in vivo* correlation potential for lipid-based formulations

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**Abstract** Lipid-based formulations have been an attractive choice among novel drug delivery systems for enhancing the solubility and bioavailability of poorly soluble drugs due to their ability to keep the drug in solubilized state in the gastrointestinal tract. These formulations offer multiple advantages such as reduction in food effect and inter-individual variability, ease of preparation, and the possibility of manufacturing using common excipients available in the market. Despite these advantages, very few products are available in the present market, perhaps due to limited knowledge in the *in vitro* tests (for prediction of *in vivo* fate) and lack of understanding of the mechanisms behind pharmacokinetic and biopharmaceutical aspects of lipid formulations after oral administration. The current review aims to provide a detailed understanding of the *in vivo* processing steps involved after oral administration of lipid formulations, their pharmacokinetic aspects and *in vitro in vivo* correlation (IVIVC) perspectives. Various pharmacokinetic and biopharmaceutical aspects such as formulation dispersion and lipid digestion, bioavailability enhancement mechanisms, impact of excipients on efflux transporters, and lymphatic transport are discussed with examples. In addition, various IVIVC approaches

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**Abbreviations:** ADME, absorption/distribution/metabolism/elimination; AUC, area under the curve; BCS, biopharmaceutics classification system; BDDCS, biopharmaceutics drug disposition classification system; CACO, human epithelial colorectal adenocarcinoma cells; C<sub>max</sub>, maximum plasma concentration; CMC, critical micellar concentration; CYP, cytochrome; DDS, drug delivery systems; FaSSGF, fasted-state simulated gastric fluid; FaSSIF, fasted-state simulated intestinal fluid; FeSSIF, fed-state simulated intestinal fluid; GIT, gastrointestinal tract; IVIVC, *in vitro in vivo* correlation; LCT, long chain triglyceride; LFCS, lipid formulation classification system; logP, n-octanol/water partition coefficient; MCT, medium chain triglyceride; MDCK, Madin–Darby canine kidney cells; NCE, new chemical entity; P-app, apparent permeability; P-gp, permeability glycoprotein; SCT, short chain triglyceride; SEDDS, self-emulsifying drug delivery system; SIF, simulated intestinal fluid; SMEDDS, self-microemulsifying drug delivery system; SNEDDS, self-nanoemulsifying drug delivery system; Vit E, vitamin E

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1. Introduction

Approximately 40% of the currently marketed formulations and more than 70% of pipeline molecules from top pharmaceutical companies today contain drugs that are poorly soluble. However, the superior therapeutic efficacy of these poorly soluble molecules (BCS-II and IV) may be the reason that they cannot be always avoided in drug development, and optimal formulation strategies are required to handle them so as to enhance their availability in systemic circulation. Even though there are conventional approaches available for handling poor aqueous solubility, very often advanced drug delivery systems (DDS) are required for developing a stable and acceptable dosage form. The most important category in advanced DDS is lipid-based formulations such as lipid solutions, lipid suspensions and self-emulsifying lipid formulations. The lipid-based formulations in general are well recognized as a frontline formulation technology to handle the poorly water-soluble compounds. These systems can be designed to present and keep the drug substance in a solubilized state thereby preventing the solubilization and subsequent dissolution step of a poorly water-soluble compound. The extensive research work done by Pouton and Porter in the area of lipid formulation development has resulted in increased awareness and understanding about lipid formulations in both industry and academia.

The preparation of lipid-based formulations is considered an easy process when compared with other solid oral dosage forms such as tablets and capsules. The excipients used in lipid formulations include lipids (natural/synthetic origin), surfactants (hydrophilic/hydrophobic), hydrophilic solvents and co-solvents. Once prepared, the lipid-based systems can be administered as solutions after dilution with suitable juices or dietary fluids or in the form of liquid-encapsulated soft gelatin capsules or liquid-filled hard gelatin capsules. The general process for development of lipid formulations along with the pharmacokinetic importance of each step is presented in Fig. 1. Due to the wide variety of excipients available for preparing lipid-based formulations, Pouton et al. introduced a lipid formulation classification system (LFCS) in order to harmonize the understanding about these formulations. As per LFCS, the lipid-based formulations can be classified into four different categories: Types-I, II, III (A and B) and IV. The compositions of these formulation types along with their characteristics, advantages, disadvantages and pharmacokinetic aspects are presented in Table 1. Out of these four systems, Type-II formulations are named as self-emulsifying drug delivery systems (SEDDS, coarse emulsions) and Type-III formulations are named as self-microemulsifying drug delivery systems (SMEDDS, microemulsions) due to their ability to form instantaneous emulsions with minimal energy input.

Out of the lipid-based formulations available in the present market, Neoral® and Sandimmun Neoral® are considered to be the first commercial successes. The complete list of all commercially available lipid formulations is outlined by Strickley. The data clearly indicate that despite the multiple advantages and extensive research work in academia and industries, there are very few commercially successful products available in the market today. From one side of the coin, this problem can be attributed to scale-up and stability challenges, marketability concerns, lack of in-house soft gelatin manufacturing capabilities, and non-acceptability of soft gelatin capsules in a few countries. From the other side of the coin, critical problems arise due to the lack of availability of in vitro tests that can describe the in vivo behavior of the lipid formulations. There are numerous in vitro ADME processes involved after intake of lipid-based formulations which make the concepts more complex for designing the in vitro tests. Additionally no clear IVIVC relationship has been established for lipid-based formulations, indicating the difficulties in correlating in vitro results with in vivo behavior. This clearly indicates the need for understanding the pharmacokinetic aspects and other related systemic processes for lipid-based formulations. An extensive literature review revealed that a few articles have described the pharmacokinetic aspects from different perspectives but no single article described all pharmacokinetic and IVIVC aspects of lipid-based formulations in detail. In this context, the objective of the present article is to outline the pharmacokinetic aspects and in vivo processing steps occurring after administration of lipid formulations so as to provide a better understanding of the lipid-based formulations from a pharmacokinetic point of view. In addition, multiple IVIVC concepts and methodologies are covered which can further aid in development of successful in vitro prediction tests.

2. Pharmacokinetic aspects of lipid-based formulations

2.1. In vivo drug solubilization and processing

Even though in the present context lipids are described as core excipients in the lipid-based formulations, they are an essential group of constituents in the food we take everyday. The processing of lipids containing drug is essentially similar to that of the dietary lipids present in food or in any other related source. Ingestion of lipid formulations results in increase in the total amount of lipids available in GI tract. These larger quantities of lipid (> 2 g, equivalent to two soft gelatin capsules) are capable of stimulating secretion of additional bile through gallbladder contraction thereby increasing the luminal concentration of bile salts. The increased levels of endogenous bile salts, phospholipids and cholesterol in the presence of lipid and surfactants provide a lipidic microenvironment to form emulsion droplets which will further transform into various components such as vesicular and micellar phases. The poorly water-soluble drug initially dissolved in the formulation will partition into these vesicular and micellar phases. The drug then will partition into the micelles formed due to the combination of bile salts, phospholipids and cholesterol which in turn are called “mixed micelles”. The formation of mixed micelles is an important step for solubilization
and absorption of a poorly water-soluble drug. The partitioning of drug into the oil core of micellar systems creates a concentration gradient across the luminal wall required to drive the absorption process. The inherent bile salts can also act as surfactants along with formulation-derived surfactants and help the drug in solubilization by improving wetting. This process provides intestinal environment with a high solubilization capacity for poorly water-soluble drug thereby enhancing the bioavailability of the compound. Various studies in the literature attempted to study the drug solubilization behavior in different phases generated during lipid digestion using a combination of phase studies coupled with solubility studies\(^{13,14}\). In addition, the distribution of drug across the colloidal species produced during in vivo processing was monitored with help of electron paramagnetic resonance spectroscopy\(^{15}\). In in vitro experiments the phases occurring during luminal processing can be studied by addition of bile salts, phospholipids and cholesterol to the intestinal fluid. It was also observed that at very high lipid levels, various liquid crystalline structures such as unilamellar, multilamellar and cubic crystalline structures were present during in vivo processing.

It has also been observed that many formulation-related factors can also contribute to lipid processing. Selection of lipid excipients in the formulation is considered an important factor since the structures formed during lipid processing can vary with the types of lipids ingested (such as long chain triglycerides and medium chain triglycerides). Long chain triglycerides are found to be digested slowly compared with medium chain triglycerides, indicating that lipase activity is a function of lipid chain length\(^{16}\). Hence, depending on the lipid chosen, the rate of digestion will vary and thereby contributing to the enhancement of bioavailability. Another related important factor from excipient point of view is the selection of surfactants due to their ability to inhibit the activity of lipases\(^{17}\). This clearly indicates that improving the dispersibility of a formulation by addition of surfactants does not necessarily improve the in vivo performance of the formulation. From other side, the efficiency of lipases on the exposed surface area was studied, which was further dictated by the number of droplets and their size in the emulsion indicating the importance of the dispersibility of the formulation\(^{18}\).

2.2. In vivo dispersibility and impact on pharmacokinetics

The in vivo dispersibility and dilution behavior are two important characteristics of the lipid-based formulation which can have significant impact on the in vivo bioavailability\(^{19}\). The in vivo
dispersibility is a measure of the size and polydispersity of the droplets formed during the self-emulsification process aided by gastric agitation. This can be measured in vitro after diluting the formulation in various biorelevant fluids such as FaSSGF (pH 1.6), FaSSIF (pH 6.5) and FeSSIF (pH 5.0) and measuring the droplet size using techniques such as Dynamic Light Scattering (such as Malvern Zetasizer)\(^2\). This would provide a clear understanding about the dilution behavior of the formulation under the in vivo conditions in stomach and intestine. In general, the smaller the droplet size, the greater the surface area and the more the lipid digestion. It has been proved that formulations producing lesser droplet size will produce more bioavailability with reduced inter-individual pharmacokinetic variability. The influence of droplet size on bioavailability has been observed for various formulations in the literature. The self-emulsifying formulation of vitamin E has resulted in approximately a 3-fold increase in absorption when compared with vitamin E solubilized in soybean oil, and it was attributed to the finer dispersion size in self-emulsifying formulation\(^6\). Similarly for cyclosporine, enhanced bioavailability (about 6.5 times) was observed for a microemulsion formulation when compared with a conventional formulation. This was mainly attributed to the smaller droplet size in the self-microemulsifying formulation of cyclosporine\(^2\). However, there were some differences with these findings in the literature with halofantrine. The size of the formulation did not have significant impact on the bioavailability when both the formulations (SEDDS and SMEDDS) were prepared using medium chain triglycerides\(^2\). Similarly for atorvastatine the droplet size of the formulation was described as a poor indicator of in vivo performance since various lipid formulations containing different surfactants resulted in different droplet sizes but there was no significant difference in the in vivo results\(^2\). Similar findings regarding the droplet size were observed for danazol and onntazolast\(^2\). It can be concluded that droplet size is not an important indicator of the in vivo performance and fate of the poorly water-soluble drug depends mainly on the dilution and digestion as they determine solubilization or precipitation of the

### Table 1 Lipid formulation classification system overview: pharmacokinetic aspects.

| Characteristics | Type-I | Type-II | Type-III | Type-IV |
|-----------------|--------|---------|----------|---------|
| **Materials**   | Oil: 100% | Oil: 40%–80% | Oil: 40%–80% | Oil: <20% | Water soluble |
|                 | Surfactants (water insoluble, HLB <12): 20%–60% | Surfactants (water soluble or insoluble): 20%–40% | Co-solvents: 0%–40% | Surfactants: 80%–100% | Surfactants: 80%–100% |
| **Dispersion behavior** | No or limited dispersion due to bile salts in GIT porine | SEDDS | SMEDDS/SEDDS | SMEDDS | Micellar solution |
| **Particle size after dispersion** | Very coarse | 100–250 nm | 100–250 nm | 50–100 nm | <100 nm |
| **Significance of aqueous dilution** | Not important due to lack of surfactants | Retains solvent capacity due to the absence of water soluble components | Some loss of solvent capacity leading to drug precipitation | Significant loss of solvent capacity due to the presence of higher quantities of water soluble components | Significant loss of solvent capacity due to surfactant dilution |
| **Significance of lipid digestion** | Highly important since it is the only mechanism to release drug | Likely to occur but not very important | Not important, but may be inhibited due to own digestion products | Good and excellent solvent capacity for many drugs | Not required and not likely to occur |
| **Advantages** | Simple system, good compatibility with capsules | Good solvent capacity, prevents drug precipitation after dilution | Clear dispersion with lesser droplet size, no requirement for digestion | Extensive precipitation of drug after dispersion | Extensive precipitation of drug after dispersion |
| **Disadvantages** | Poor solvent capacity unless drug is highly lipophilic | Turbid emulsions and in vivo fate depends on digestion | Precipitation of drug likely to occur after dispersion and digestion | Bioavailability may be enhanced (depending on the extent of precipitation after digestion and dissolution), less interindividual variability due to less particle size formed after dispersion | May not yield higher bioavailability due to extensive drug precipitation under in vivo conditions |
| **Pharmacokinetic behavior** | May enhance bioavailability but can result in high interindividual variability due to lack of dispersion | May greatly enhance bioavailability but can result in high interindividual variability due to formation of coarse emulsion | Bioavailability may be enhanced (depending on the extent of precipitation after dispersion and digestion), less interindividual variability due to less particle size formed after dispersion | Bioavailability may be enhanced (depending on the extent of precipitation after dispersion and digestion), less interindividual variability due to less particle size formed after dispersion | May not yield higher bioavailability due to extensive drug precipitation under in vivo conditions |
| **Marketed products** | Calcitrol (Rocaltrol\(^6\), Roche) | Cyclosporin A (Sandimmune\(^6\), Novartis) | Cyclosporin A (Neoral\(^6\), Novartis) | Tipranavir (Aptivus\(^6\), Boehringer Ingelheim) | Ritonavir (Norvir\(^6\), Abbott, Amprenavir (Agenerase\(^6\), Glaxosmithkline) |

\(^{21}\)Indicates the dilution behavior of the formulation in various biorelevant fluids such as FaSSGF (pH 1.6), FaSSIF (pH 6.5) and FeSSIF (pH 5.0) and measuring the droplet size using techniques such as Dynamic Light Scattering (such as Malvern Zetasizer). This would provide a clear understanding about the dilution behavior of the formulation under the in vivo conditions in stomach and intestine. In general, the smaller the droplet size, the greater the surface area and the more the lipid digestion. It has been proved that formulations producing lesser droplet size will produce more bioavailability with reduced inter-individual pharmacokinetic variability. The influence of droplet size on bioavailability has been observed for various formulations in the literature. The self-emulsifying formulation of vitamin E has resulted in approximately a 3-fold increase in absorption when compared with vitamin E solubilized in soybean oil, and it was attributed to the finer dispersion size in self-emulsifying formulation. Similarly for cyclosporine, enhanced bioavailability (about 6.5 times) was observed for a microemulsion formulation when compared with a conventional formulation. This was mainly attributed to the smaller droplet size in the self-microemulsifying formulation of cyclosporine. However, there were some differences with these findings in the literature with halofantrine. The size of the formulation did not have significant impact on the bioavailability when both the formulations (SEDDS and SMEDDS) were prepared using medium chain triglycerides. Similarly for atorvastatine the droplet size of the formulation was described as a poor indicator of in vivo performance since various lipid formulations containing different surfactants resulted in different droplet sizes but there was no significant difference in the in vivo results. Similar findings regarding the droplet size were observed for danazol and onntazolast. It can be concluded that droplet size is not an important indicator of the in vivo performance and fate of the poorly water-soluble drug depends mainly on the dilution and digestion as they determine solubilization or precipitation of the
drug. This clearly indicates the importance of maintaining drug in the solubilized state for enhanced absorption rather than precipitating after dilution in the larger quantities of biological fluids in the stomach and intestine.

2.3. In vivo dilution and impact on pharmacokinetics

In vitro dilution in biological fluids followed by optical microscopic examination during the formulation development can provide an understanding about the precipitation of a drug after dilution in GI tract. In addition, the saturation solubility in water added to formulations can be studied which would provide an understanding about drug precipitation due to water intake. Increasing the solubilization capacity of the formulation significantly over the desired drug loading can also help to avoid precipitation under in vivo conditions. The amount of drug precipitated depends on the formulation factors as well as physiological factors of the individual under medication. While formulation factors can be controlled, physiological factors cannot be controlled indicating the importance of ensuring the solubilized state using various in vitro tools. The physiologically related drug precipitation would lead to inconsistent pharmacokinetic profiles leading to higher inter-individual variability. Dai et al. provided an excellent approach for studying the precipitation behavior of poorly water-soluble drug in lipid-based formulations at development stage using only milligrams of NCEs. The authors have suggested an in vitro precipitation determination using a 96-well-plate method with varying dilution factors and durations in biological fluids. Based on this approach, the authors have categorized formulations as no-precipitation, fast-precipitation and slow-precipitation. The in vivo administration of these formulations to dogs resulted in plasma profiles which are in agreement with in vitro precipitation profiles. In another similar study, the ability of Type-II, III-A, III-B and IV systems to maintain the model drug fenofibrate in the solubilized state after dispersion in aqueous solution was examined. The authors concluded that even though Type-II systems produced turbid emulsions after dilution, they are able to maintain the drug in the solubilized state with very slow precipitation. The Type-III-A systems maintained fenofibrate in a metastable state after dilution in water for several hours to days. The Type-III-B systems led to rapid precipitation due to the presence of higher quantities of water-soluble surfactants. Type-IV systems totally failed to maintain the drug in the solubilized state resulting in extensive rapid precipitation. The results indicate the importance of selection of suitable lipid-based systems for maintaining a drug in solubilized state and also indicate that the formulation scientists should not only consider the physical appearance as ultimate aspect but should also consider precipitation of poorly water-soluble drug in the system. While formulating the drug in a Type-II system seems to be an attractive choice for preventing drug precipitation, various other aspects are also studied for maintaining the drug in a supersaturated state without in vivo precipitation. The most interesting approach for formation of supersaturated drug solution in vivo is by incorporation of hydrophilic polymeric excipients in the formulation that can act as precipitation inhibitors. This approach is very useful for drugs exhibiting lower solubility in lipid excipients and thereby requiring higher clinical doses. The precipitation inhibitors can significantly delay the precipitation times and if the precipitation times are greater than the mean absorption time, the extent of absorption of a poorly water-soluble drug can be increased thereby enhancing the bioavailability.

Most of the commonly used polymer excipients in this category are hydroxypropyl methyl cellulose (HPMC), methyl cellulose (MC), sodium carboxymethyl cellulose (NaCMC) and polyvinyl pyrrolidone (PVP). The use of HPMC as a precipitation inhibitor for paclitaxel in a supersaturable SEDDS resulted in a 9-fold increase in bioavailability in rats. The use of PVP as precipitation inhibitor for carbamazepine in a supersaturable SMEDDS that enhanced the bioavailability by 5-fold in dogs when compared with a commercially available tablet has also been reported. The use of polymers as precipitation inhibitors is not only limited to lipid-based formulations but can also be extended to other formulations such as solid dispersions and controlled release tablet formulations. Yamashita et al. reported the use of HPMC as precipitation inhibitor for delivery of tacrolimus in the form of solid dispersion wherein it showed 10-fold increase in area under the curve (AUC) and C\text{max} when compared with crystalline powder.

2.4. Absorption and bioavailability enhancement

The basic property by which optimized lipid formulations such as SEDDS and SMEDDS enhance the absorption and bioavailability of poorly water-soluble drugs is by presenting the drug in the solubilized form throughout the transit through gastrointestinal tract with minimal precipitation. Various mechanisms have been stated in the literature for bioavailability enhancement such as stimulation of pancreatic and biliary secretions, prolongation of GI residence time to slow down the delivery system to the absorption site and increase the time available for dispersion or dissolution, stimulation of lymphatic transport by overcoming the first pass metabolism, increasing the intestinal wall permeability by membrane fluidization and opening the tight junctions, reduced intestinal metabolism by encapsulating the drug in the form of micelles, and reduced efflux transporter activity by incorporating excipients that can act as efflux inhibitors. These processes are provided in pictorial representation in Fig. 2. These numerous mechanisms clearly indicate the advantages of using lipid-based formulations for enhancing bioavailability. It is also evident that even if two to three mechanisms from the above stated work for a drug formulated in a lipid system, it would definitely lead to enhanced bioavailability thereby enhancing pharmacological activity. A literature review revealed numerous examples in which the absorption and bioavailability of poorly water-soluble drugs were enhanced by administration in the form of lipid formulations. Selected examples are discussed here and Table 2 summarizes the other cited studies.

The most important and classical example in the bioavailability enhancement by lipid formulations is Sandimmun® and Sandimmune Neoral® formulations of cyclosporine A by Novartis®. Initially Sandimmune® was introduced into European market in 1981 as a SEEDS formulation containing Labrafil M1944CS, olive oil and ethanol. When dispersed in water, this formulation forms an oil-in-water coarse macroemulsion with high polydispersity values. In 1994 again cyclosporine was introduced as a SMEDDS formulation containing Cremophor RH40, corn oil glycerides, propylene glycol and ethanol under the brand name of Sandimmune Neoral®. This formulation forms a spontaneous microemulsion with droplet size less than 100 nm. Compared with Sandimmune®, Sandimmun Neoral® provided more extent in oral absorption and bioavailability. The improved dispersion
The optimized formulation consisted of triethylcitrate, Tween 20, Span 20, triethanolamine and benzyl alcohol. The SMEDDS formulation demonstrated higher saturation solubility values (>150 times) and dissolution rates for pranlukast. In rats, the absorption of pranlukast improved by about 3-fold when dosed in SMEDDS, compared with a suspension formulation and it was attributed to increased dissolution rates in the form of SMEDDS.

2.5. Role of excipient selection on bioavailability

In general, there is a wide choice of excipients available for formulating poorly water-soluble drugs in the lipid formulations. The typical excipients present in lipid-based formulations are: (a) lipids: synthetic or natural lipids; (b) surfactants: non-ionic lipophilic or hydrophilic surfactants; (c) hydrophilic solvents: for better dispersion and to increase solvent capacity; (d) cosolvents: for reducing the viscosity of a formulation and to facilitate dispersion
t. From a formulation perspective, the excipient selection will have an impact on the drug load, dispersion characteristics, solubilization of drug and, importantly, on stability. However, it is also important to consider excipients from pharmacokinetic aspects which can ultimately affect the bioavailability.

2.5.1. Lipids

The basic function of lipids is to solubilize the drug and keep it in a dissolved state throughout the transit in the GI tract. If lipid digestion leads to drug precipitation, it needs to be resolubilized in intestinal fluids to get absorbed. This is especially important for BCS class-II and IV54. In vitro, the lipolysis can be simulated using a pH stat titration apparatus and the in vivo conditions are simulated by adding the lipid formulation into the biorelevant buffers (fasted or fed condition simulating) and subsequently adding lipases for digestion55. Many factors can contribute to the results obtained from in vitro lipolysis experiment and these factors along with their importance in predicting in vivo data are represented schematically in Fig. 36,57.

![Figure 2: Mechanisms by which lipid formulations can enhance bioavailability.](Image)
**Table 2**  Bioavailability enhancement by lipid based formulations.

| Drug/BCS class | Formulation (Excipients) | Test system | Results | Mechanism |
|----------------|--------------------------|-------------|---------|-----------|
| Acyclovir (III) | Microemulsion (Labrafac, Labrasol, Plurail oleique, water) | Sprague-Dawley rats | Relative bioavailability enhanced by 12.78 times when compared with tablet | Enhanced solubilization in microemulsion |
| Atorvastatin (II) | SMEDDS (a) Labrafac, Cremophor RH40, propylene glycol (b) Estol, Cremophor RH40, propylene glycol (c) Labrafac, Cremophor RH40, propylene glycol | Beagle dogs | Significant increase in relative bioavailability (about 1.5 times) for all three formulations when compared with tablet | Enhanced intestinal solubility and mucosal permeability |
| Carvedilol (II) | SEDDS (Labrafac M1944CS, Tween 80, Transcutol) | Beagle dogs | Relative bioavailability enhanced by 4.1 times when compared with tablet | Enhanced solubility and dissolution rate |
| Coenzyme Q10 (II) | SEDDS (Mylvacet 9-45, Labrafac CM-10, Lauraglycol) | Coonhound dogs | Relative bioavailability enhanced by 2 times when compared with powder | Enhanced aqueous solubilization |
| Cyclosporin (II) | SEDDS-Sandimmune (corn oil, ethanol) SEDDS-Neoral (corn oil glycerides, Cremophor RH40, ethanol) | Humans | Increased relative bioavailability and $C_{\text{max}}$ in humans. Neoral superior to Sandimmune in terms of reduced food effect, dose linearity and reduced interindividual variability | Due to formation of microemulsion after aqueous dilution for Neoral when compared with Sandimmune |
| Danazol (II) | LCT solution (soyabean oil) LCT-SMEDDS (soyabean oil, Maisine 35-1, Cremophor EL, ethanol) MCT-SMEDDS (Captep 355, Capmul MCM, Cremophor EL) | Beagle dogs | Relative bioavailability in the order of LCT solution > LCT-SMEDDS > MCT-SMEDDS > micronized powder. | Enhanced intestinal solubilization resulted in increased relative bioavailability. Significant drug precipitation in MCT-SMEDDS |
| Griseofluvin (II) | Corn oil emulsion, corn oil suspension | Rats | Relative bioavailability in the order of corn oil emulsion > corn oil suspension > aqueous suspension | Enhanced solubilization in emulsion resulted in more relative bioavailability |
| Halofantrine (II) | MCT-SEDDS (Captep 355, Capmul MCM, Cremophor EL, ethanol) MCT-SMEDDS (Captep 355, Capmul MCM, Cremophor EL, ethanol) LCT-SMEDDS (soyabean oil, Maisine 35-1, Cremophor EL, ethanol) | Beagle dogs | Higher relative bioavailability from LCT SMEDDS compared with other formulations. All formulations enhanced bioavailability by 6–8 times when compared with solid tablet formulation | Enhanced solubilization and prevention of precipitation after aqueous dispersion |
| Ibuprofen (II) | Microemulsion (MCT oil, DGMO-C, HCO-40) | Rats | Bioavailability in microemulsion comparable with organic solution and higher than aqueous suspension | Enhanced solubility of compound in oil |
| Indomethacin (II) | SEDDS (Tween 85, ethyl oleate) | Sprague-Dawley rats | Relative bioavailability of SEDDS 1.57 times higher when compared with aqueous suspension | Improvement in the solubility and dissolution |
| Itraconazole (II) | SEDDS (Transcutol, Pluronic L64, tocopherol acetate) | Sprague-Dawley rats | Relative bioavailability of SEDDS was significantly higher than marketed capsule and reduced food effect when administered in SEDDS | Enhanced dissolution by incorporation in SEDDS |
| Ontazolast (II) | SEDDS (Gelucire 44/14, Peceol) | Charles River CD rats | Absolute bioavailability increased at least by 10 times from SEDDS formulations. SEDDS formulations enhanced lymphatic transport | SEDDS formulation improved dissolution and solubility, and also enhanced bioavailability through lymphatic absorption thereby bypassing extensive hepatic metabolism |
2.5.2. Surfactants

Cremophor EL and Cremophor RH40 are the most common and most studied excipients in the category of nonionic surfactants. Even though Cremophor EL and Cremophor RH40 are from the same family of surfactants, Cremophor EL has lower degree of ethoxylation compared with Cremophor RH40 and it is unsaturated. However, similar to lipid excipients, most important differences from the in vivo digestion point of view need to be considered for their use in the formulation. With regard to both the surfactants, Cremophor RH40 is less readily digested when compared with Cremophor EL. The reasons for the differences in digestion are not clear but have been attributed to the differences in the reactivities of the saturated castor oil glyceride backbone in Cremophor RH40 leading to different reaction products when compared with Cremophor EL. Due to its lesser digestion properties, Cremophor RH40 seems to be an attractive choice for keeping the drug in the solubilized state for longer time when compared with Cremophor EL. As observed with danazol, SMEDDS containing Cremophor RH40 maintained more drug in the aqueous phase throughout digestion process when compared Cremophor EL, resulting in a 3-fold increase in bioavailability with Cremophor RH40. Even though the above theories look convincing, under in vivo conditions, the relative affinity of lipases towards lipid and surfactant will ultimately decide the degrees of digestion and drug solubilization, indicating the importance of formulation optimization using in vitro digestion experiments. From another perspective, the reduction in bioavailability of danazol in dogs when the proportional content of Cremophor EL is increased in the formulation has been reported. This was primarily attributed to the fact that reduction of lipid increased the chances of drug precipitation due to dilution of Cremophor EL under the in vivo conditions, which ultimately failed to maintain the drug in the solubilized state. This clearly indicates the necessity to optimize the ratio of lipid to surfactant for enhanced bioavailability. In addition, excess of surfactant can lead to micelle formation and it can also hinder the absorption process due to the dramatic increase in the molecular weight of the whole complex. This can be an important aspect from an in vitro studies point of view (Caco-2, MDCK cell line studies for estimation of permeability) and may not be relevant in vivo due to dilution of surfactants in the larger gastric fluids.

On the other side, long term use of surfactants is not recommended due to their ability to disrupt the cell membrane and their toxic and adverse effects (e.g., Cremophor EL). Several antiviral protease inhibitors such as ritonavir are available in few other cases (e.g., dexamethasone) where in no significant differences in bioavailability were noted with respect to administration in different chain lengths. In addition to the above mentioned aspects of lipid digestion capabilities, the absorption profiles of drug can be changed depending on the type of lipid selected. For example, MCT promotes portal vein absorption through liver and LCT promotes lymphatic absorption thereby bypassing the first pass metabolism (discussed in detail in later sections).

### Table 2 (continued)

| Drug/BCS class | Formulation (Excipients) | Test system | Results | Mechanism |
|----------------|--------------------------|-------------|---------|-----------|
| Peniclovemide (Not available) | LCT (soybean oil, Triolein), MCT (Triotanol, SCT (Tributyrin), liquid paraffin emulsion | Rats | Bioavailability is in the order of MCT>LCT>paraffin emulsion>SCT>aqueous suspension | Higher bioavailability in MCT is due to reduced drug precipitation during lipid digestion |
| Phenytoin (II) | Corn oil emulsion, corn oil suspension | Beagle dogs | Relative bioavailability in rats is in the order of emulsion>oil suspension>aqueous suspension | Enhanced solubility in lipid emulsion |
| Progesterone (IV) | SEDDS (mono-di-glycerides, Polysorbate 80) | Beagle dogs | Relative bioavailability of SEDDS is 9 times higher than the aqueous suspension | Enhancement of solubility and permeability when administered in SEDDS |
| Seocalcitol (Not available) | LCT-SMEDDS (sesame oil, Pecceol, Cremophor RH40) | Rats | Absolute bioavailability LCT-SMEDDS is equal to MCT-SMEDDS | Similar solubility values of drug in SIF may have resulted in similar bioavailability |
| Silymarin (Not available) | SMEDDS (Tween 80, ethyl alcohol, ethyl linolate) | Rabbits | Relative bioavailability of SMEDDS is 1.88 and 48.82 times higher than PEG solution and suspension respectively | Alternative pathways such as lymphatic transport contribution to enhanced bioavailability |
| Simvastatin (II) | SMEDDS (Caproyl 90, Cremophor EL, Carbitol) | Beagle dogs | Relative bioavailability is 1.5 times higher in SMEDDS compared with conventional tablet | Enhanced solubility in lipid excipients |
| Tocotrienols (II) | Two SEDDS (Tween 80 and Labrasol) | Humans | Relative bioavailability enhanced by 2–3 times when compared with non-self-emulsifying formulation | Enhanced solubility and finer dispersion properties |
| Vitamin E (II) | SEDDS (Tween 80, Span 80 and palm oil) | Humans | Relative bioavailability of SEDDS is 2-folds higher than oil solution | Presence of surfactant in SEDDS led to higher bioavailability when compared with oil solution |
market as oral solution and soft gelatin capsules containing huge amounts of surfactants. These formulations are taken in large numbers on a daily basis due to the dosage requirements. Experience from these formulations suggests that these dosage form regimens are well tolerated but long term effects need to be evaluated. Even though there are no official guidelines for recommended daily permissible intake of these excipients in humans and special populations such as pediatrics, it should be ensured that these excipients should be used as little as possible in the formulation so as to avoid adverse/toxic effects until regulatory limits are available.

2.5.3. Hydrophilic solvents

The hydrophilic solvents such as triethyl citrate, polyethylene glycol 400 (PEG400) and propylene glycol are used in smaller quantities in lipid-based Type-III formulations to aid in self-emulsification through a phenomenon called “dispersion and stranding”66. Keeping excipient related toxicities aside, only few of the hydrophilic solvents have significant pharmacokinetic concerns. One excipient in this category is PEG400, due to its influence on gastric motility and subsequent effect on drug absorption when ingested in larger quantities (≥2.5 g). The decreased intestinal transit time is due to poor absorption of this excipient from intestine that results in PEG400-induced osmotic activity due to water retention, thereby stimulating intestinal motility and transit. Since the small intestine is the major site for absorption of many compounds, reduction in transit time will decrease the time available for absorption for many of the poorly soluble and slowly dissolving drug compounds such as ranitidine67. However, at lower concentrations such as 1 g, PEG400 was shown to significantly enhance the absorption of ranitidine probably due to modulation of intestinal permeability68. Even though the amounts of hydrophilic solvents used in the lipid-based formulations are minimal, they should be selected with caution due to the above mentioned impact on the pharmacokinetics.

2.5.4. Cosolvents

In general, cosolvents such as ethanol are used at very low concentration to reduce the viscosity and to aid in the dispersion process. At these lower concentrations the cosolvents do not possess any pharmacokinetic importance.

2.6. Lymphatic transport

Lymphatic transport is an attractive absorptive pathway for enhancing bioavailability of lipophilic drugs. The main advantage of lymphatic absorption is bypassing hepatic metabolism and thereby increasing blood concentrations and therapeutic efficacy for extensively metabolized drugs73. Among the various animal models available for study of the lymphatic transport of small molecules, a rat model with a cannulated mesenteric lymph duct is widely used. Other models such as sheep and dog are widely used for studying lymphatic transport after parenteral and oral administrations, respectively74.

While lymphatic transport is directly proportional to lipophilicity, a drug molecule with a partition coefficient (log P)>5 and triglyceride solubility >50 mg/mL can be forced to be absorbed via lymphatic transport using lipid-based formulations rather than through portal vein and liver75. It has been shown that drugs solubilized in MCT and SCT (number of carbons C<12) are primarily absorbed into the portal blood and encounter hepatic enzymes, and drugs which are solubilized in LCT (C>12) are transported via intestinal lymphatic system and subsequently are absorbed to systemic circulation. During lipolysis in the GI tract, digestion products such as fatty acids and monoglycerides with C>12 are combined with phospholipids and cholesterol present in enterocyte to form triglycerides. The formed triglycerides along with entrapped drug are packed into chylomicrons and subsequently enter the lymphatic system thereby circumventing the liver and preventing hepatic metabolism. Fatty acids and monoglycerides with C<12 are not combined with chylomicrons and hence are absorbed into the portal vein. Triglyceride solubility of the drug is a driving factor for partitioning into lipid-enriched chylomicrons and hence the more the triglyceride solubility, the more the partitioning.

This is the fundamental underlying mechanism for lymphatic absorption mediated by formulating the drug with LCT rather than with MCT and SCT76. The degree of saturation and amount of lipid are also found to have an effect on the lymphatic transport of drug molecules.

Charman and Stella77 studied the lymphatic transport of DDT (log P, 6.19) and hexachlorobenzene (log P, 6.53) in an in situ rat model system. When administered with oleic acid, the lymphatic transport of DDT and hexachlorobenzene was 33.5% and 2.3%, respectively, in the rat in situ model. Even though the log P values for both compounds are similar, the difference in the extent of lymphatic transport was attributed to 13-fold difference in the triglyceride solubility values. In another study by Dahan and Hoffman78 the in vivo performance of vitamin D3 was evaluated in formulations consisting of LCT, MCT and SCT. Even though an in vitro lipolysis experiment estimated rank order of MCT>LCT>SCT, based on the drug retained in aqueous part after lipolysis, in vivo results have provided a different rank order LCT>MCT>SCT. This was primarily attributed to the fact that...
lymphatic transport of vitamin D3 is enhanced in LCT thereby resulting in more in vivo extent in absorption. This is also clear from the fact that when lymphatic transport is blocked by pre-administration of cycloheximide, the in vivo data correlated with in vitro lipolysis data\(^6\). This clearly indicates the limitation of in vitro lipolysis that it cannot predict the in vivo performance for the compounds which are absorbed via lymphatic route. In another study, a good linear correlation between ex vivo uptake of lipophilic drugs by chylomicrons with reported intestinal lymphatic bioavailability in rats was reported\(^7\). In contrast, the most recommended log \(P\) and solubility in triglycerides provided only moderate correlation with lymphatic bioavailability. Based on these results, they have also developed a simple screening model which considers the degree of association of lipophilic drugs with isolated chylomicrons that can be used to predict intestinal lymphatic transport. Similarly Gershkovich and Hoffman\(^7\) have also observed that log \(P\) and triglyceride solubility values do not necessarily guarantee lymphatic transport. Similar contradicting results were reported where poor lymphatic transport (3% of dose transported into intestinal lymph) of peniclomedine was observed even though it has log \(P\) 5.48 and triglyceride solubility of 175 mg/mL\(^7\). Despite these contradictory results, intestinal lymphatic transport still serves as an excellent opportunity to enhance the oral bioavailability of lipophilic drugs.

### 2.7. Food effect reduction

Food effect can be described as an increase in the rate and extent of drug absorption in presence of food. The food effect is most studied over the past few decades and it depends on a number of factors arising from physiology, dosage form and, importantly, physicochemical properties\(^7\). Various aspects such as enhanced solubility in the presence of fat components of food, stimulation of bile secretion, surfactant effects by food components, delay in gastric emptying to enhance the absorption time, and inhibition of efflux transporters by food components are recognized as potential mechanisms for bioavailability enhancement of poorly soluble drugs in the presence of food\(^7\).

The lipid components present in the food can perform the same function as lipid components present in the lipid-based formulations. The dietary lipids present in the food can act as solubilizers for drug and subsequently undergo lipid digestion by gastric and pancreatic lipases similar to the process described for lipid-based formulations so as to produce various micellar species. It has also been shown that a few lipid components in the food can enhance chylomicron production and can stimulate the lymphatic transport to enhance the bioavailability\(^7\). Hence it is evident that if proper lipids and surfactants are chosen for formulating a poorly water-soluble drug, they can also act as surrogates for food components and thereby nullifying the food effect and increasing patient compliance by avoiding the high fat meal consumption with dosage form.

The reduction in food effect for itraconazole using SMEDDS formulation was reported in healthy volunteers\(^20\). The SMEDDS formulation exhibited enhanced \(C_{\text{max}}\) and AUC values under both fasted and fed conditions when compared with conventional capsule formulation and demonstrated similar plasma concentration time profiles under both fasted and fed conditions, indicating an insignificant food effect. Similarly, the reduced food effect for the highly lipophilic compound torcetrapib after administration in lipid-based formulations was demonstrated. The optimized formulations showed enhanced fasted state exposure and reduced food effect from 5-fold to 3-fold in dogs at dose levels of 90 mg\(^8\). The optimized formulations were also found to show reduced inter-individual pharmacokinetic variability.

### 2.8. Bioavailability enhancement by transporter inhibition

The efflux transporters present in the intestine are known to reduce the bioavailability of many drugs. The Biopharmaceutics Drug Disposition Classification System (BDDCS) demonstrated that drugs with poor solubility (class-II) will not be able to saturate the efflux transporters such as P-glycoprotein present in the intestine, hence limiting their bioavailability\(^2\). It was also reported that the bioavailability and pharmacological activity of many anticancer agents have been reduced drastically due to the action of drug efflux transporters such as P-glycoprotein, breast cancer resistant protein and multidrug resistance protein present in the intestine that flushes out the absorbed drug from enterocyte back into the intestine. This might also lead to the problem of multidrug resistance thereby reducing potency of a drug with repeated administration\(^2\).

It has been reported in the literature that formulating drugs in lipids (e.g., LCT and MCT) and surfactants (e.g., Cremophor EL, Cremophor RH40, Tween 80, \(\alpha\)-tocopherol) polyethylene glycol 1000 succinate (Vit E TPGS) can enhance the bioavailability by inhibiting the efflux transporters such as P-glycoprotein (P-gp)\(^83,84\). It has been reported that inhibition of efflux will lead to increase in concentration and residence time of the drug in the enterocyte leading to increased drug available for lymphatic transport. In addition, many of the synthetic and semi-synthetic lipids can inhibit the drug efflux transporters similar to the effects shown by lipid components present in fat enriched food. Various mechanisms have been proposed for inhibition of transporters by excipients, but the main concept behind inhibition is membrane fluidization-induced conformational changes and cholesterol depletion. In addition, the micelles formed during emulsification can mask the active drug thereby not exposing it to the efflux transporters. A few hydrophilic solvents, such as PEG400, are also known to inhibit certain efflux transporters so as to enhance the bioavailability\(^85\). Hence, when the above mentioned excipients such as lipids, surfactants and hydrophilic solvents are used in lipid-based formulations they may enhance the bioavailability not only due to their solubilization capabilities, but also due to the inhibition of efflux transporters.

Paclitaxel is one of the most important chemotherapeutic agents and it is a substrate for P-gp and CYP3A4. The bioavailability of paclitaxel is very limited and hence it is formulated in a 1:1 ratio of Cremophor EL and dehydrated ethanol for intravenous administration (Taxol\(^8\)). Various efforts have been undertaken to enhance the oral bioavailability of paclitaxel through formulating in nanoparticles and supersaturated SEDDS containing HPMC as a precipitation inhibitor along with P-gp inhibitors such as Vit E TPGS and Cremophor EL\(^8\). This formulation resulted in 3-fold higher bioavailability when compared with the traditional Taxol\(^8\) formulation. Even though bioavailability has increased with this formulation it did not yield sufficient therapeutic levels, indicating that either the SEDDS formulation was not able to circumvent the P-gp effect or due to metabolism by CYP3A4. To evaluate this possibility, the SEDDS formulations were dosed along with cyclosporine that further enhanced the bioavailability of paclitaxel when compared with Taxol\(^8\). This clearly indicates that pure SEDDS formulation enhanced the bioavailability of paclitaxel.
either due to solubilization or due to P-gp inhibition or due to combination of both.

3. IVIVC potential for lipid-based formulations

The use of in vitro–in vivo correlation for various Biopharmaceutics Classification System (BCS) classes is described in the seminal article of Amidon et al.\(^8\) that resulted in the birth of BCS. BCS class I drugs are proved to be excellent candidates and class-IV drugs are considered to be poor candidates for IVIVC due to their solubility and permeability characteristics. While the application of IVIVC to class-III drugs is limited due to their poor permeability, the solubility/dissolution properties of poorly water-soluble class-II compounds can be greatly enhanced by formulation approaches such as solid dispersion and lipid formulations such as SEDDS and SMEDDS, thereby making them behave as class-I compounds in vivo. The rapid dissolution of class-II compounds will enable the time required for complete dissolution to be significantly less than gastric emptying time, thereby enhancing the possibility of achieving strong IVIVC correlation\(^9\). Out of the different lipid formulations available, Type-II and III formulations disperse rapidly in the in vivo fluids thereby reducing the time required for complete solubilization in the in vivo biological fluids, which increases the potential for achieving IVIVC for poorly soluble compounds. The following sections detail the various approaches used for developing the in vitro and in vivo correlations for various lipid-based formulations.

3.1. In vitro dispersion/precipitation test and in vivo data

The in vitro dispersion or precipitation test is an attractive and simple test for determining the amount of the drug in a solubilized state after dispersing the lipid formulation into aqueous fluids. The rationale for using this test to predict the in vivo performance is based on the fact that the completely solubilized drug will be readily absorbed whereas precipitated drug will not be available for absorption. Hence, when using this test to screen different lipid formulations, the percentage of solubilized drug in biological fluids in vitro can be correlated with the amount of the drug absorbed in vivo or with AUC, thereby possibility of achieving an IVIVC or rank ordering formulations. The dispersion and precipitation behavior of a poorly soluble NCE in various lipid-based formulations using 96-well microtiter plate in various biological fluids such as SIF, FaSSIF and FeSSIF was reported\(^10\). The authors have short-listed three formulations based on precipitation kinetics (fast, slow and no precipitation) and studied the pharmacokinetics in dogs. There was a good correlation between the precipitation kinetics observed in biological fluids versus AUC values, where the rank ordering was fast < slow < no precipitation. In addition, the in vitro precipitation in FaSSIF correlated well with in vivo performance indicating that lipid excipients can act as surrogates for food components. The IVIVC for BCS class-II drug cyclosporine from SMEDDS formulation was reported\(^11\). In this work, a new generic SMEDDS formulation for cyclosporine was developed and compared with two reference SMEDDS formulations. All three formulations were evaluated for in vitro dissolution (in pH 1.2, 4.5 and 6.8) and in vivo in dogs. Strong Level A correlations were achieved for fraction dissolved versus fraction absorbed. The author concluded that BCS class-II compounds will behave as class-I compounds when administered in optimized SMEDDS formulation. Similar Level A IVIVC correlations were established for various poorly water-soluble drugs such as ritonavir and lopinavir administered as lipid formulations in soft gelatin capsules\(^12\). Various IVIVC examples from the literature using in vitro dispersion/dissolution/precipitation and in vivo data are presented in Table 3.

Even though achieving correlation with this test appears simple, there are many limitations involved. Firstly, if the precipitated drug redissolves at later points in the intestine due to pH change or solid state change of the precipitated drug, the obtained IVIVC correlation or rank ordering will not be valid. Secondly and most importantly it is not only the dispersion but also the lipid digestion by gastric and pancreatic lipases that plays an important role in maintaining the drug in the solubilized state. Hence, even if the drug remains in a solubilized state after dispersion, it can very well precipitate due to lipid digestion. The role of lipid digestion in achieving IVIVC for lipid formulations is described in the following section.

3.2. In vitro lipolysis and in vivo data

Out of all the methods used for establishing IVIVC for lipid-based formulations, in vitro lipolysis plays a major role due to its capability to simulate the most important steps in the absorption of lipid-based formulations such as dispersion and digestion. The limitation of the in vitro dispersion/precipitation test to predict the in vivo performance can be overcome by incorporating in vitro lipolysis. In in vitro lipolysis, the formulation is added to the buffer solution containing various components such as bile salts, thereby simulating the fasted/fed after ingestion of lipid-based formulation. Immediately thereafter enzymes such as pancreatic lipase are added and the lipolysis process is initiated. Samples are taken at various time points and centrifuged and the aqueous phase is analyzed for drug content. Correlation between the amount of drug present in the aqueous phase at different time points and the in vivo plasma concentration time profiles, and percentage of solubilized drug at the end of digestion versus pharmacokinetic parameters such as AUC or C\(_{\text{max}}\) can be used to develop IVIVC relationships for lipid-based formulations.

Various studies have demonstrated the utility of in vitro lipolysis to predict in vivo data. The relative bioavailability of halofantrine formulated using MCT and LCT-based formulations was reported\(^13\). In this study, the authors attempted to correlate in vitro drug solubilization with digestion data to the in vivo data. The authors found strong correlations between the data only when lower lipid masses were used. Out of LCT and MCT, the superior bioavailability was observed with the LCT-based formulation. The authors concluded that the solubilization capacity of the formulations is highly dependent on the nature of the lipid used, and the predictivity of the in vitro lipolysis is dependent on the amount of lipid used in the experiment. This indicates the importance of optimization of various parameters in lipolysis experiments for better prediction of in vivo bioavailability. Reymond et al.\(^14\) demonstrated correlations between in vitro and in vivo data for cyclosporine formulated in MCT and LCT. The in vitro lipolysis indicated that higher amounts of solubilized cyclosporine present the aqueous phase in case of MCT compared with LCT. However in in vivo studies, LCT resulted in higher bioavailability compared with MCT. The authors concluded that since LCT is digested slowly compared with MCT, after 12 min of lipolysis more cyclosporine was present in the MCT. However in the in vivo situation, cyclosporine has highest affinity toward the oil phase.
containing triglycerides and hence affinity towards LCT is higher compared with MCT. This case study clearly explains that not only the design of *in vitro* lipolysis should be considered, but also the inherent properties of the drug when developing IVIVC.

In another similar study, the poorly water-soluble drug danazol was formulated in three formulations: LCT-solution, LCT-SMEDDS and MCT-SMEDDS. *In vitro* lipolysis indicated significantly higher solubilized danazol for LCT-SMEDDS when compared with MCT-SMEDDS and LCT-solution. The *in vivo* results are in agreement with *in vitro* lipolysis wherein higher absorption was observed for LCT-SMEDDS formulations when compared with MCT-SMEDDS. The utility of *in vitro* dynamic lipolysis and neuro-fuzzy networks for establishing IVIVC for SMEDDS formulations was reported for probucol. In this study, probucol was formulated in an oil solution, two SMEDDS and one nano-emulsifying (SNEDDS) formulation. Higher percentages of solubilized probucol were observed for SMEDDS and SNEDDS formulations when compared with oil solution (rank order SMEDDS > SNEDDS > oil solution). The *in vivo* results demonstrated good correlation with *in vitro* data and similar rank order was observed. The developed neuro-fuzzy model achieved significantly higher prediction abilities for different formulations.

The *in vitro* lipolysis experiment can be used to predict the food effect for lipid-based formulations. The fasted and fed state gastrointestinal lipolysis experiment to establish IVIVC for SNEDDS solidified SNEDDS and conventional tablet formulation was reported for cinnarizine. During the lipolysis experiment the fed state was simulated by addition of 3.5% of whole fat milk to the lipolysis media. In the fasted state lipolysis model the rank order for the percentage solubilized cinnarizine in aqueous phase was SNEDDS > solidified SNEDDS > tablet, which was in good agreement with the *in vivo* data in dogs. In the fed state model similar amounts of cinnarizine were solubilized for all the formulations. The fed state *in vivo* dog study indicated similar performance for SNEDDS and solidified SNEDDS whereas the performance of the conventional tablet was improved. This clearly indicates that conventional tablet formulation showed a food effect whereas SNEDDS and solidified SNEDDS showed an absence of a food effect. This example demonstrates the utility of *in vitro* lipolysis experiments to predict food effects for lipid-based formulations. Various IVIVC examples from the literature using *in vitro* lipolysis and *in vivo* data are presented in Table 4.

### 3.3. *Ex vivo* intestinal permeability and *in vivo* data

*Ex vivo* intestinal permeability study includes the use of an animal intestine to study the transport of drug from the mucosal to the serosal layer. In this study, the permeability characteristics of pure drug (or formulation) can be studied using an experimental set up named as “Ussing chamber system”. This system has been widely used in academics and in pharmaceutical industries for evaluation.

### Table 3 Selected IVIVC examples based on *in vitro* dispersion/precipitation/dissolution and *in vivo* pharmacokinetic data.

| Drug/Formulation | *In vitro* dispersion/precipitation/dissolution data | *In vivo* data | IVIVC |
|------------------|---------------------------------------------------|----------------|------|
| Cyclosporin (Soft gelatin capsule) | *In vitro* dissolution (in pH 1.2, 4.5 and 6.8) for one generic and two reference formulations | *In vivo* bioavailability in dogs | Level A correlation between *in vivo* fraction absorbed versus *in vitro* fraction dissolved for test and reference formulations ($R^2=0.992$) |
| JNJ-25894934 (NCE, Soft gelatin capsule) | Formulations categorized based on precipitation kinetics: fast (solubility: 76.37 mg/mL), slow (105.61 mg/mL) and non-precipitating (96.04 mg/mL) formulations in FaSSIF and FeSSIF | Pharmacokinetic results in Mongrel dogs showed lowest bioavailability for fast precipitating and similar bioavailability for slow and non-precipitating formulations in fasted state and fed state | Good agreement between *in vitro* precipitation and *in vivo* pharmacokinetic data, good correlation between fed state precipitation kinetics and *in vivo* data compared with fasted state precipitation kinetics |
| Ritonavir (Soft gelatin capsule) | *In vitro* dissolution data in various media containing water and sodium lauryl sulfate (SLS-0.3%, 0.5%, 0.7%, 1%) | *In vivo* pharmacokinetic data in humans | Strong Level A correlations were obtained in between percent dissolved versus percent absorbed ($R^2$ values: 0.952, 0.935, 0.993, 0.963 for 0.3%, 0.5%, 0.7%, 1% SLS respectively) |
| Lopinavir (Soft gelatin capsule) | *In vitro* dissolution data using media 2.3% SLS at pH 6.0 and USP Type-I apparatus at 25 rpm | *In vivo* pharmacokinetic data in humans | Strong Level A correlation was obtained ($R=0.997$). The equation obtained was Fraction absorbed = $-0.0019+1.0075 \times$ Fraction dissolved |
| Arundic acid (Soft gelatin capsule) | *In vitro* dissolution data in pH 8.0 dissolution medium or the pH 6.8 dissolution medium containing 2% sodium dodecyl sulfate | *In vivo* pharmacokinetic data in humans | IVIVC was established by plotting *in vitro* dissolution time versus *in vivo* absorption time for both dissolution media, pH 6.8 dissolution media produced stronger correlations |
| Fenofibrate | *In vitro* dissolution of three SMEDDS formulations was performed in FaSSGF pH 2 and FaSSIF-V2(PO4) | *In vivo* data in human volunteers | *In vitro* dissolution and *in silico* simulations were used to predict *in vivo* human plasma profiles. The point estimates for $C_{max}$ and AUC fell within range of 0.8–1.25 indicating the accurate simulation of *in vivo* profiles from *in vitro* data |
The apparent permeability characteristics ($P_{app}$) of a NCE. Recently this technique was utilized for lipid-based formulations to predict or rank order for in vivo performance. The ex vivo permeability technique was used to study the transport of adefovir dipivoxil formulated in solid SNEDDS formulation whereas the solid SNEDDS formulation showed superior permeability characteristics compared to drug suspension, thereby enhancing the chances of improving bioavailability. In another study, the ex vivo permeability characteristics of the poorly water-soluble drug talinolol formulated in SNEDDS was evaluated using porcine small intestine and the developed SNEDDS formulation showed enhanced permeability characteristics compared to suspension. These results were very well correlated with the in vivo results in Wistar rats. The correlation between ex vivo rat intestinal studies and in vivo bioavailability for fexofenadine formulated in lipid surfactants such as Gelucire 44/14 and Vit E TPGS was reported. The authors observed increased ex vivo permeability characteristics (A-to-B) for lipid formulations when compared with a pure drug formulation. The authors also reported decreased B-to-A permeability for lipid formulations indicating the inhibition of efflux transporters. The results from ex vivo permeability studies correlated very well with the results obtained from in vivo bioavailability studies in male Wistar rats. The ex vivo permeability characteristics of lipolytic products for lipid formulations of dexamethasone and griseofulvin in LCT, MCT and SCT were reported. Superior permeability characteristics were found for lipolytic products produced from SCT formulations when compared with LCT and MCT formulations. However, the in vivo bioavailability results correlated well with the in vitro lipolysis results where the rank order of MCT>LCT>SCT was observed for Griseofulvin and LCT=MCT=SCT was observed for dexamethasone respectively. The authors concluded that formulating in SCT might enhance the permeability characteristics but overall intelligent optimization of lipid formulations can only be achieved with in vitro lipolysis, and most importantly, permeability does not correlate with in vivo bioavailability for class-II compounds. Various IVIVC examples from the literature using ex vivo permeability and in vivo data are presented in Table 5.

Overall, the summary of different ways to achieve IVIVC for lipid-based formulations is represented in Fig. 4.

### 4. Conclusions and future perspective

Even though lipid-based formulations demonstrate multiple advantages over available formulation technologies today, they only constitute a 2%-4% share in the commercially available formulations in US, UK and Japan. The reasons for significantly lesser market share can be attributed to multiple reasons but most important, contributing factors are lack of availability of standardized in vitro tests and poor understanding of the pharmacokinetic behavior of lipid formulations after oral ingestion. While the standardization and harmonization of in vitro tools such as lipolysis have already been initiated as a part of LFCS (Lipid Formulations Classification System), the knowledge about the pharmacokinetic aspects and processes involved after ingestion of lipid formulations remains a gray area. Understanding the pharmacokinetic aspects will not only help in designing the optimum lipid-based formulations, but also will help in designing the in vitro tests that can be used to predict in vivo behavior.
correctly, thereby establishing IVIVC relationships. From the in vivo processing point of view, dispersion and digestion are found to have major impact on the availability of a drug for absorption. With the in vitro tools such as dissolution/precipitation in biological fluids and in vitro lipolysis, these effects can be predicted to a certain extent but not completely. Out of all the in vitro tests available today for evaluation of lipid formulations, only in vitro lipolysis is found to be more relevant to describe the in vivo pharmacokinetic data, but it still has limitations such as inability to predict in vivo fate of drugs which are transported through lymphatic route and also when drug efflux transporters come into play. Despite these limitations and hurdles, lipid-based formulations still remain an attractive area for enhancing the bioavailability of poorly water-soluble drugs. In addition, with the advent of novel lipid-based formulations such as dried emulsions, spray-dried emulsions, solid SMEDDS, supersaturated SMEDDS and self-emulsifying solid dispersions, the poorly water-soluble drug can be delivered as a solid dosage form by formulating with minimal amounts of lipid and surfactants, which bypasses the limitations of conventional lipid formulations but yet without compromising the bioavailability.

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