Preparation and evaluation of novel self-dispersing lipid formulation of nevirapine for oral delivery

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ABSTRACT
Nevirapine, a non-nucleoside reverse transcriptase inhibitor, exhibits pH-dependent solubility resulting in poor dissolution limiting its absorption. In the present study, an attempt is made to prepare a self-dispersing lipid formulation (SDLF) involving the novel combination of two drug solubility enhancing strategies, i.e., self-emulsifying system (SES) and solid dispersion (SD). Nevirapine-SES (NV-SES) consisting of an ideal ratio of oleic acid (oil), Cremophor RH 40 (surfactant), and Transcutol (co-surfactant) exhibited good emulsifying properties with maximum drug loading. The NV-SES developed was mixed with an equal quantity of nevirapine: gelucire 50/13 solid dispersion [Nevirapine Solid Dispersion (NV-SD)] to get a novel nevirapine SDLF (NV-SDLF). The NV-SDLF was characterized for its zeta potential, polydispersity index, and particle size, and the results obtained were found to be satisfactory. NV-SDLF was adsorbed on carrier neuciline U2 to get solid NV-SDLF, and solid-state characterization using Differential Scanning Calorimetry (DSC) and Powder X-Ray Diffraction (PXRD) indicated the amorphous nature of the drug in NV-SDLF. The in-vitro drug release of the NV-SDLF was found to be 87.2% ± 0.96% in comparison with nevirapine 28.8% ± 1.02%. In-vivo pharmacokinetic performance of solid NV-SDLF and pure drug suspension was carried out in rats, and a significant two-fold increase (p < 0.05) in area under the curve (AUC) of solid NV-SDLF was observed with respect to nevirapine AUC. Thus, the in-vitro and in-vivo results confirmed that the combined strategies of SES and SD in the formulation of solid NV-SDLF were found to enhance the bioavailability of nevirapine, which could be a promising novel drug delivery system in the management of Acquired Immune Deficiency Syndrome (AIDS).

INTRODUCTION
Antiviral drug nevirapine is one of the preferred choices for the treatment of AIDS and is effectively used in Highly active antiretroviral therapy (HAART) for the management of the disease. It stops the growth of the virus at a submicromolar concentration by inhibiting the ability of the viral enzyme to transcribe the ribonucleic acid (Jindal and Mehta 2015; Wang et al., 2019). The major concern with nevirapine is its side effects which include Central nervous system (CNS) toxicity, hepatotoxicity, insomnia, depression, nausea, rashes, and hyperlipidemia. It gets metabolized by CYP3A and stimulates mutagenicity and hepatocarcinogenicity (Kuo and Chung, 2011). Also, a high dose has to be given to penetrate viral reservoir sites, which results in intolerance (Shegokar and Singh, 2011).

In addition to the mentioned toxicity, nevirapine belongs to the Biopharmaceutics Classification System (BCS) class II drugs for the reasons of poor water solubility (~0.1 mg/ml) and pH-dependent solubility with a pKa of 2.8 (Chadbha et al., 2010; Moreton et al., 2014). In an acid medium, the solubility is 4.99 g/l and the solubility is reduced by a factor of 10–40 when pH is increased. This observable fact is due to change in the structure of molecules at different pH (de Oliveira et al., 2013). Solubility plays a significant role in in-vitro and in-vivo performances of a drug, and dissolution of a drug depends on ionic strength, pH, and buffer capacity of the Gastro intestine (GI) fluids (Hamed et al., 2016). Due to the above-mentioned reasons, the dissolution of nevirapine has been considered to be the rate-limiting step for its absorption.

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Always there exists a correlation between solubility and bioavailability. A number of formulation strategies could be used to improve the solubility of class II drugs; such as a change in crystal habit, particle size reduction, chemical approaches, and formulation approaches using carriers (Leuner and Dressman, 2000; Vasconcelos et al., 2016). Solid dispersion (SD) is one of the approaches, which enhances solubility by reducing particle size and increasing water solubility (Huang and Dai, 2014). Among the four generations of SD, the third generation SD is prepared using surfactants as carriers; this has the advantage of increased drug release, long stability, and enhanced bioavailability (Vo et al., 2013).

An advance in formulation technology has introduced a range of carriers for the preparation of dispersion. Gelucire belongs to the group of saturated polyglycolized glycerides that has gained more attention as SD carriers due to its surface active properties. It reduces the surface tension between the drug and dissolution medium and consequently enhances the oral bioavailability of the poorly soluble drug (Bertoni et al., 2020; da Fonseca Antunes et al., 2013). It has been successfully used as a carrier in SD of antiviral drugs (Damian et al., 2000; Sinha et al., 2010), ursolic acid (de Oliveira Eloy et al., 2012), carvedilol (Poturi et al., 2011), and Lycopene (Faisal et al., 2013).

In recent years, another approach has gained importance in the delivery of poorly soluble drug called lipid-based drug delivery system, which can enhance solubility and oral bioavailability (Gershanik and Benita, 2000; Sandeep et al., 2013; Singh et al., 2020). Self-emulsifying drug delivery system (SEDDS) is one among them, consisting of a blend of oils, surfactants, and co-surfactants, on mild agitation with GI fluid and consequently enhances the oral bioavailability of the poorly soluble drug (Bertoni et al., 2020; da Fonseca Antunes et al., 2013). It has been successfully used as a carrier in SD of antiviral drugs (Damian et al., 2000; Sinha et al., 2010), ursolic acid (de Oliveira Eloy et al., 2012), carvedilol (Poturi et al., 2011), and Lycopene (Faisal et al., 2013).

In the present study, we have investigated a novel self-dispersing lipid formulation (SDLF) of nevirapine by combining the benefits of both SD and self-emulsifying system (SES). As SES has the advantage of overcoming toxicity associated with the drug, initially SES was designed using various combinations of oil, surfactant, and co-solvent and was characterized for drug loading, microscopic, and self-emulsifying properties. Later, a SD of drug was prepared using gelucire 50/13 as a carrier to improve its solubility and characterized for drug using DSC, XRD, and in-vitro drug release. Subsequently, novel SDLF was developed by combining SD and SES and evaluated for its in-vitro and in-vivo behavior.

**MATERIALS AND METHODS**

Nevirapine was a gift sample from Micro Labs Limited, Bengaluru, India. Gelucire 50/13 and Transcutol HP were gift samples from Gattefose India Pvt. Ltd, Mumbai, India. Neucil US2 and Cremophor RH 40 were gift samples from Strides Arco Labs, Bengaluru, India. Oleic acid was purchased from SD Fine Chemicals, Mumbai, India. All the chemicals used in this study were analytical grade. Buffers were prepared using double distilled water.

**Preparation and characterization of drug-loaded SES**

Oil (oleic acid), surfactant (Cremophor RH 40), and co-surfactant (Transcutol HP) were selected based on drug solubility and emulsifying ability. Nevirapine-SES (NV-SES) formulations were prepared by dissolving nevirapine in different amounts of oil (10%-90%) placed in glass vials at ambient temperature by vortexing on a vortex mixer. To the mixture, previously prepared surfactant co-surfactant mixture (S/CoS mix) in the ratio of 3:1 was added in the range of 10%-90% (Table 1) and vortexed with simultaneous heating up to 40°C until a homogenous transparent preparation was obtained. NV-SES formulations were evaluated for drug loading, phase separation, precipitation, and the emulsifying ability for selecting optimized NV-SES (Cometa et al., 2020; Komal et al., 2015; Shweta et al., 2011). Optimized NV-SES was subjected to aqueous dilution to get microemulsion and evaluated for droplet size, polydispersity index (PDI), and zeta potential (ZP) using Malvern Zetasizer instrument (Nano-ZS, UK). The solubility of the drug in excipients and SES was determined using

**Table 1. Formulation and characterization of NV-SES.**

| NV-SES code | Formulation of NV-SES | Characterization of NV-SES |
|-------------|------------------------|---------------------------|
|             | Oil (%)                | Surfactant (%)             | Co-surfactant (%) | Drug loading (mg/ml) | %Transmittance at 638.2 nm | Emulsification grade<sup>a</sup> |
| F1          | 10                     | 67.5                       | 22.5             | 12.24              | 99.5                       | A                           |
| F2          | 20                     | 60                         | 20               | 15.68              | 94.2                       | A                           |
| F3          | 30                     | 52.5                       | 17.5             | 16.34              | 92.2                       | A                           |
| F4          | 40                     | 45                         | 15               | 16.09              | 63.2                       | A                           |
| F5          | 50                     | 37.5                       | 12.5             | 18.91              | 56.8                       | B                           |
| F6          | 60                     | 30                         | 10               | 16.71              | 44.5                       | B                           |
| F7          | 70                     | 22.5                       | 7.5              | 18.41              | 43.8                       | B                           |
| F8          | 80                     | 15                         | 5                | 19.71              | 42.9                       | C                           |
| F9          | 90                     | 7.5                        | 2.5              | 23.12              | 40                         | C                           |

<sup>a</sup> Emulsification grade: A- microemulsion formed within 5 minutes; B- droplets spread in water to form a turbid emulsion; and C- coalescence of oil droplets.
either DMSO/Methanol as diluent by UV spectrophotometer (Shimadzu, Pharmaspec UV-1700) at wavelength 282 nm.

**Preparation and characterization of drug-loaded SDs**

A SD of nevirapine with Gelucire 50/13 was prepared in the ratio of 1:1, 1:2, 1:3, and 1:4 by the conventional melt method. Nevirapine was dissolved in a small quantity of ethanol and this solution was added to melt gelucire 50/13 (at 50°C) in a china dish. The solution mixture was dried in a rotary evaporator to remove ethanol and cooled rapidly in an ice bath (Faisal et al., 2013; Sinha et al., 2010). Prepared SDs were subjected to *in-vitro* drug release in intestinal pH 6.8. The selected ratio was subjected to DSC and XRD to identify drug excipient interaction.

**Preparation of SDLF**

A novel SDLF containing a combination of SD and self-emulsifying formulation was prepared by mixing (Table 2) an equal quantity of drug loaded self-emulsifying system to get an nevirapine SDLF (NV-SDLF) (Faisal et al., 2013). Adsorbent NeucilUS2 was added to the above formulation to overcome the stickiness and tackiness and to improve flow properties. The prepared solid NV-SDLF was scrapped, pulverized, sieved, and stored in the desiccator.

**Characterization of NV-SES, NV-SD, and NV-SDLF**

**Differential scanning electron microscope**

Thermal analysis was carried out on a DSC Q1000 (TA Instruments, UK). Samples were accurately weighed into aluminum pans and sealed. The sample and reference pans were loaded into the sample chamber at ambient temperature, equilibrated to 25°C, and held at this temperature for 5 minutes. The temperature range for the thermogram was 25°C–250°C, and the samples were heated at a rate of 5°C/minute with an applied modulation of ±1°C every 60 seconds. Samples were purged in a stream of dry nitrogen flowing at 50 ml/minute.

**X-ray diffraction**

Powder XRD was recorded on an X-ray diffractometer (RigakuMiniflex 600) consisting of an X-ray tube operated at 600 watts with a fixed tube current of 15 mA and a voltage of 40 kV. Samples were exposed to radiation, over an angular range of 5°–130° (2θ) at a scanning speed of 0.01°–100°/minute. The diffraction patterns were analyzed by Integrated X-Ray Powder Diffraction Software (PDXL) software (version 1.8.0.3).

**Surface morphology of NV-SDLF**

Surface characteristics of nevirapine, NV-SDLF, and other excipients were assessed using a scanning electron microscope [ZEISS EVO MA18 with Oxford EDS(X-act)]. Magnified images were taken at an excitation voltage of 15–20 kV.

**In-vitro dissolution studies of NV-SD and NV-SDLF**

*In-vitro* dissolutions of NV-SD and NV-SDLF were carried out in 900 ml of phosphate buffer pH 6.8 using USP I dissolution apparatus (Electrolab TDT-08L, India). Samples equivalent to 50 mg nevirapine filled in capsules were used for dissolution study at 50 rpm and 37°C ± 0.5°C. Aliquots of samples were withdrawn at a regular time interval of 10 minutes for 180 minutes. The volume pipetted out was replaced with fresh media to maintain sink condition. Samples taken were filtered through a 0.45 µ membrane and then analyzed using a UV spectrophotometer at 282 nm (Chadha et al., 2010; US Food and Drug Administration).

**Ex-vivo permeability study of solid NV-SDLF**

The permeability study was carried out using chicken intestinal sac (Nipun and Ashraful Islam, 2014) which was collected from a slaughter house. A cleaned intestinal sac of length 3 cm was tied at one end and filled with nevirapine, and the sample was suspended in 1 ml of water. Tying the other end of the sac, we determined the release of the drug using USP I dissolution apparatus containing 900 ml of basic medium 6.8 phosphate buffer at 50 rpm maintained at 37°C ± 0.5°C. Samples were withdrawn at a predetermined interval and each sampling was replaced with a fresh medium. Samples filtered through 0.45 µ were analyzed for drug release by using UV spectrophotometer at 282 nm.

**In-vivo pharmacokinetic studies in the animal model**

The bioavailability of the prepared solid NV-SDLF was assessed in male Wistar rats (220–250 g) after taking institution ethical clearance. Experiments were conducted as per the Committee for the Purpose of the Control and Supervision on Experiments on Animals guidelines. Before the experiments, animals (n = 4 for each group) were fasted for 12 hours. Each animal was cannulated in the right jugular vein in anesthetized condition. Suspensions of solid NV-SDLF and the pure drug were prepared in 0.5% carboxymethyl cellulose with gentle agitation. A test group of animals was administered with solid NV-SDLF suspension and a control group of animals was administered with the pure drug at a dose of 60 mg/kg body weight with oral gavage (Inugala et al., 2015; Moretton et al., 2014; Usach and Peris, 2011).

**Estimation of nevirapine in the blood sample**

Nevirapine was extracted from 200 µl plasma by liquid–liquid extraction using ethyl acetate as the extraction solvent. The extracted dried sample was reconstituted with mobile phase, acetonitrile: 0.1% formic acid (80:20 v/v) containing internal standard (METAXALONE) for introduction into Liquid Chromatography with tandem mass spectrometry (LCMS/MS). LC separation was carried out by introducing 10 µl of the reconstituted sample into High-performance liquid chromatography (HPLC) (Prominence LC-20AT, Shimadzhu, Japan) consisting of C-18 column (Agilent, ZORBAX XB-C18, 5 µm, 4.6 × 50 mm) maintained at 40°C. Throughout isolation, the mobile phase flow rate of 0.500 ml/minute was maintained. The eluted components from the column were delivered to API 4000 Triple Quadruple mass spectrometer (SCIEX, USA), to detect ions.

**Pharmacokinetics analysis**

Non-compartmental pharmacokinetic analysis of plasma data was carried out using WinNonline software. Parameters determined using WinNonline are as follows: half-life (t1/2), area under the curve (AUC) of plasma drug concentration time curve from zero to infinity (AUC∞), AUC from time of administration
to the last plasma concentration (AUC_{last}), maximum plasma concentration (C_{max}), time of maximum concentration (T_{max}), elimination rate constant (K), and relative bioavailability (%F).

Statistical analysis

_in-vivo_ pharmacokinetic data expressed as mean ± SE were analyzed for statistical significant differences between the means of the two groups by unpaired Student’s "t"-test using GraphPad Prism Software. The results were considered significant at p < 0.05.

RESULTS AND DISCUSSION

Preparation of drug-loaded SES

SES has gained attention in the delivery of BCS class II drugs in recent years due to its ability to combat solubility issues of poorly soluble drug and other advantages with respect to increased GI permeability, enhanced lymphatic absorption, and inhibiting first-pass metabolism and Pgp protein efflux (Gurram et al., 2015, Sandeep et al., 2013). In the present study, the development of NV-SES was initiated by screening the various oils, surfactants, and co-surfactants based on the solubility of the drug in the individual vehicle and miscibility among each other. Based on the solubility studies, oleic acid a long-chain triglyceride was selected as oil, Cremophor RH 40 as surfactants, and transcutol HP as co-surfactants, with the solubility of 21.05, 15, and 12.23 mg/ml, respectively. In the preparation of SES, the combination of surfactant and co-surfactant ratio plays a prominent role in the formation of smaller particles. As per the requirement stated by Pouton, the ideal combination of surfactant and co-surfactant is 3:1 (Colin, 1997). Hence, 3:1 ratio of Cremophor RH40:Transcutol HP (surfactant/co-surfactant mix) were mixed and added to the different percentages of oils to get different ratios of SES. Drug loading was conducted for all the ratios and the emulsifying ability of these ratios was determined by dilution method and observed for %transmittance at 638.2 nm. Among the SES ratios, the ratio containing 30% of oil and 70% of S/Co mixture emulsified within a minute with the %transmittance of 92.2 with drug loading 20 mg/ml was selected as optimized SES. The optimized SES was analyzed for particle size and was found to be 108.2 nm, which indicates microemulsion formation after dilution. Measurements of PDI and ZP confirm uniform size distribution of particles and the stability of the colloidal system. The particles in the dispersion with large potential will tend to repel each other and there will be no tendency for particles to aggregate (Gurram et al., 2015). The ZP of the optimized SES was observed to be ±36.5 mV, which confirmed good stability. In support of the ZP result, the polydispersibility index of optimized SES was found to be 0.291 which is less than 0.3, which indicates uniform size distribution of dispersed particles (Zupančič et al., 2016).

Preparation of drug-loaded SDs

SDs are molecular mixtures of hydrophobic drugs in hydrophilic carriers, where the drug release profile is driven by polymer properties. In SD, the drug is dispersed but not fully solubilized in the excipient matrix which facilitates a higher dose loading (Vo et al., 2013). For the present study, gelucire 50/13, belonging to the gelucire family, was selected as a SD carrier because of its ability to enhance the oral bioavailability of the hydrophobic drugs and the ability to self-emulsify on contact with aqueous media forming a fine dispersion, i.e., microemulsion (Damian et al., 2000).

A SD of nevirapine with different ratios of gelucire 50/13 was prepared by the melt method. From the in-vitro release study in 6.8 phosphate buffer (Fig. 1), it was observed that as the gelucire ratio increased, nevirapine solubility also enhanced. With regard to the bulk of the final formulation and drug loading, 1:4 drug to gelucire ratio was used for the preparation of a novel lipid delivery system.

Preparation of SDLF

Although SES has number of justifiable advantages in delivering a poorly soluble drug, in our present study, it was not possible to successfully load the target dose of 50 mg nevirapine within the SES pre-concentrate. With the intention of increasing the drug loading, a new formulation has been designed called SDLF which has the combined benefits of SES and SD. The SDLF was prepared by mixing an equal quantity of optimized SES and the SD. The product SDLF was of semisolid consistency which would have the disadvantage of the storage and the stability issues. The use of adsorbent can minimize the stickiness and tackiness problems of lipid formulations. Neuciline US2, a synthetic, amorphous form of magnesium alumino-metasilicate, exhibits multiple excipient roles due to its porosity, greater specific area of 300 m²/g, and adsorbing capacity of 3.2 ml/g (Gupta et al., 2001). The literature supports its applications in the lipid formulations as an adsorbent in the preparation of solid SEDDS and as the carrier in the formulation of lipid SD (Gumaste et al., 2013; Gupta et al., 2001; Kazi et al., 2017). Hence, neuciline US2 was added to the NV-SDLF to get solid NV-SDLF with desired solid consistency accompanying good stability and flow property.

Characterization of formulations

_Differential scanning electron microscope_

A thermogram of the drugs, gelucire 50/13, neucilin U2, NV-SD, and solid NV-SDLF, are shown in Figure 2f. A sharp endothermic peak at 251.35°C in Figure 2f and 50.36°C in Figure 2e indicates the melting point of nevirapine and gelucire 50/13. The DSC curve of SD (Fig. 2d) revealed a shallow peak without any sharp endothermic peak of the drug at 251.35°C; this confirmed the change in drug property from crystalline to amorphous form. Also, the presence of the endothermic peak of the carrier in SD substantiated that there is no

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**Figure 1.** _in vitro_ drug release of NV-SD in 6.8 phosphate buffer (n = 3).
change in the melting point of the carrier and suggested the formation of a monotectic system (Sinha et al., 2010). DSC of neuciline U2 in Figure 2c showed characteristic endotherms at 91.17°C and 231.72°C. A solid SDLF without drug and solid NV-SDLF exhibited one endothermic peak at 150.55°C and 146.51°C in comparison with neuciline U2 (Fig. 2c). This may be due to the melting of the mixture exhibiting more than one phase, indicating the minor change in the thermal behavior of Neucilin U2. But there is no obvious appearance of drug endothermic peak at 251.35°C which confirms the amorphous nature of the drug in the SDLF.

X-ray diffraction

The physical states of nevirapine in NV-SD and solid NV-SDLF were investigated by the X-ray diffraction method and diffractograms are shown in Figure 3. The XRD patterns of pure nevirapine (Fig. 3f) showed distinct peaks between 5° and 30° (2θ = 9.3°, 13.2°, 13.6°, and 25.6°) indicating crystalline nature of nevirapine (Jindal and Mehta, 2015) and diffractogram in Figure 3d shows typical triglycerides peaks at 19.1° and 23.3° of gelucire 50/13 (Chauhan et al., 2005). Triglyceride peaks were retained in NV-SD (Fig. 3d), with suppressed nevirapine crystalline peaks, confirming the transformation of crystalline nevirapine into amorphous nevirapine. As expected, the XRD pattern of neucilin US2 gave no peaks, indicating the amorphous nature of the solidifier. SDLF and N-SDLF retained sharp characteristic peaks of gelucire 50/13 similar to pure gelucire50/13; this result ruled out the possible interaction of drug and gelucire 50/13. The absence of crystalline peaks of nevirapine in solid NV-SDLF reveals that the physical state of the crystalline drug has been transferred into an amorphous form, which is attributed to greater dissolution rate of solid Nevirapine self-dispersing lipid formulation (NV-SDLF) due to higher internal energy (Venkata and Basanth 2013; Tran et al., 2013).

Surface morphology

The surface morphology of the nevirapine, neusilin US2, and solid NV-SDLF was studied by Scanning Electron Microscopy (SEM), as shown in Figure 4. As expected, it was observed that adsorbent neusilin US2 existed as a porous granular spherical fine powder (Fig. 4b), and Figure 4c clearly shows the adsorption of semisolid NV-SDLF on the surface of neusilin US2.

Dissolution studies

Stomach pH can vary due to the presence of food and in fasting conditions (Hamed et al., 2016), and drugs may get absorbed over an extended period of time and reach the small intestine with a physiological pH between 6.5 and 7.6. Therefore, in-vitro dissolution of the formulation NV-SDLF
was carried out in the pH, the non-enzymatic simulated gastric pH 0.1 N HCl, and simulated intestinal 6.8 pH phosphate buffer. The release profile of a solid NV-SDLF was compared with the pure drug release profile and it was observed that the drug release in acidic pH was found to be 93% ± 0.91%, which is very close to 99.5% ± 0.84% pure drug release (Fig. 5). In basic pH, solid NV-SDLF exhibited 87.2% ± 0.96% drug release, which was 58.4% more than that of pure drug release (28.8% ± 1.02%). The enhanced drug release in basic pH was attributed to gelucire 50/13, which can enhance drug dissolution (de Oliveira Eloy et al., 2012) and good dispersibility properties of the prepared solid NV-SDLF.
Table 2. Composition of NV-SES, NV-SD, and NV SDLF.

| SES components/g          | Solid dispersion components/g | NV-SDLF components/g |
|---------------------------|-------------------------------|----------------------|
| 30% OA                    | GU50/13 = 800 mg              | 15% OA               |
| 70% CRH40:TC (S:Comix 3:1 ratio) |                  | 35% CRH40:TC (S:Comix 3:1 ratio) |
| Drug loaded 20 mg         | Drug loaded 200 mg            | Drug loaded 120 mg   |

Table 3. Pharmacokinetic parameters of solid NV-SDLF and suspension of NV.

| Kinetic parameter | Unit       | Suspension of NV | NV-SDLF |
|-------------------|------------|------------------|---------|
| $C_{max}$         | ng/ml      | 5,412.45 ± 3,965.61 | 7,217.52 ± 3,345.58 |
| $t_{max}$         | hour       | 2 ± 1.63         | 2.5 ± 1.15 |
| $AUC_{0-\infty}$  | µg.hour/ml | 21.70 ± 13.05    | 33.03 ± 10.44 |
| $AUC_{\infty}$    | µg.hour/ml | 26.84 ± 4.087    | 46.54 ± 21.57 |
| $t_{1/2}$         | hour       | 4.01 ± 4.21      | 3.29 ± 1.20 |
| $%F$              | %          | 100              | 173.39  |

$C_{max}$: Maximum plasma concentration; $t_{max}$: Time to maximum plasma concentration; $AUC_{0-\infty}$: Area under the curve between 0 and last time interval; $AUC_{\infty}$: Area under the curve between 0 and infinite time interval; $t_{1/2}$: Plasma elimination half-life; $%F$: $AUC_{\infty}$ of formulation divided by $AUC_{\infty}$ of pure drug.

Results are expressed as mean ± SE ($n = 4$).

$p < 0.05$ versus suspension of NV.

Release kinetics

It was found that the formulated solid NV-SDLF in 0.1N HCl and 6.8 pH phosphate buffer indicated the best agreement with Korsmeyer and Peppas’ release with “n” values of 0.25 ($R^2 = 0.8072$) and 0.080 ($R^2 = 0.9675$), which shows that this release is not governed by the Fickian diffusion. The undefined mechanism of the formulation may be due to the combined effect of SES with SDs.

Particle size and ZP

The in-vivo behavior of the dispersed particle is influenced by particle size. The smaller the size of the particle, the larger will be the surface area, which results in enhanced dissolution and bioavailability (Shweta et al., 2011). As depicted in Figure 6b in the present study the particle size of the dispersed solid NV-SDLF was found to be 100.5 nm, which could justify the enhanced dissolution of the drug in basic pH. Quality of dispersion is assessed by determining PDI. The dispersed NV-SDLF exhibited a polydispersibility index of 0.277, which is less than 0.3 and confirms the good dispersion property of NV-SDLF (Zupančič et al., 2016). The stability of dispersed particles depends on the surface charge and its potential; the greater the ZP value of surface charge, the greater will be the repellant forces. The ZP of the NV-SDLF was measured using a zeta sizer, and it exhibited a ZP of ±32.4 mV. The result signifies the stability of the dispersed system (Fig. 6a).

Ex-vivo permeation studies

Nevirapine belongs to BCS class II drug which does not have permeability issues. But for confirming the absence of influence of formulation excipients on permeability, an ex-vivo permeation study was carried out as described by Nipun and Ashraful Islam (2014), using chicken intestinal sac. The permeation profile of the pure drug was compared with solid NV-SDLF and observed that there is no significant difference in drug diffusion profile of the pure drug and optimized formulation (Fig. 7).

Pharmacokinetic studies

The in-vivo oral performance of solid NV-SDLF was compared with nevirapine suspension obtained by dispersing nevirapine in 0.5% w/v carboxymethylcellulose aqueous solution. In this assay, we evaluated the following pharmacokinetic parameters: maximum plasma concentration ($C_{max}$), time to maximum plasma concentration ($T_{max}$), area under the curve between 0 and last time interval ($AUC_{0-\infty}$), area under the curve between zero to ∞ ($AUC_{\infty}$), and plasma elimination half-life ($t_{1/2}$) using Wistar rats at a dose of 15 mg nevirapine per 1 ml. These parameters of nevirapine were obtained by non-compartmental analysis of plasma concentrations at selected time points. The pharmacokinetic curves of the nevirapine plasma concentration versus time after oral administration are shown in Figure 8. Details of the pharmacokinetic parameters are shown in Table 3. The $AUC_{0-\infty}$ values observed were 33.03 ± 5.22 and 46.54 ± 21.57 µg/ml for solid NV-SDLF and 21.70 ± 13.05 and 26.84 ± 4.087 µg/ml for nevirapine suspension, respectively. The increase in AUC values was found to be statistically significant at $p < 0.05$. Also, it was observed that solid NV-SDLF exhibited an increase in $C_{max}$ value to 72.17 ± 3,345.58 from 54.12 ± 3,965.61 µg/ml for pure drug suspension, which is nearly 34% increase. The increase in AUC and $C_{max}$ was substantiated with increased in-
vitro dissolution of Nevirapine. The result is most likely due to the presence of components in NV-SDLF; oleic acid, Cremophor RH 40, and Gelucire, which enhance the bioavailability of the poorly soluble drug by various mechanisms. The expected mechanisms are lymphatic transport of lipophilic drug (long-chain oleic acid) (Zhang et al., 2021), absorption of the lipophilic drug by inhibiting the efflux pumps, cytochrome P450 3A (Cremophor RH 40) (Taylorouz et al., 2003), and solubilizing and emulsifying ability of gelucire (de Oliveira Eloy et al., 2012).

\[ T_{\text{max}} \] and \( t_{\text{max}} \) of nevirapine remained unchanged for both solid NV-SDLF and nevirapine drug suspensions, respectively. This confirmed that the component used in the formulation of SDLF has not interfered in the elimination of the drug and release of the drug from the dosage form.

**CONCLUSION**

A novel self-dispersing nevirapine formulation was successfully developed by dual formulation approach. Formulated NV-SDLF exhibited enhanced in-vitro dissolution in simulated gastric and intestinal pH and showed good dispersibility properties with desired particle size, PDI, and ZP. NV-SDLF with semisolid consistency was adsorbed onto carrier neuciline US2 to overcome handling issues. Solid NV-SDLF was evaluated for its in-vivo performance in male Wistar rats and characterized by DSC and XRD for drug excipient interaction. A significant increase in AUC was observed for solid NV-SDLF in comparison with pure drug nevirapine, and also solid NV-SDLF showed increased \( C_{\text{max}} \) value. The result proved that added excipient in the formulation of NV-SDLF has the ability to enhance solubility, thereby enhancing bioavailability. At the same time, the added excipients may contribute to the lymphatic transport of the drug and inhibits the efflux pumps and cytochrome P450 3A. Further studies need to be conducted to verify the possibility of converting novel solid NV-SDLF into a suitable dosage form for the effective management of AIDS.

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**CONFLICT OF INTEREST**

The authors declare that there are no conflicts of interest to disclose.

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None.

**AUTHORS’ CONTRIBUTIONS**

All authors have made an active contribution in every step of research work and reviewed the final draft before submission to the journal. All the authors are accountable for the information provided in the manuscript.

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