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CHAPTER 10

# Solid lipid nanoparticles and microemulsions for drug delivery: the CNS

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**Abstract:** The chapter examined solid lipid nanoparticles (SLN) and microemulsions, chosen as carriers 19 of drugs, administered in vivo to be transported to the central nervous system. Drugs of different 20 structures and for different therapies have been studied such as doxorubicin SLN stealth and nonstealth 21 administered in rats by intravenous route, apomorphine SLN administered in rats by duodenal route, 22 melatonin SLN administered by transdermal and oral routes in humans, and apomorphine microemulsion 23 administered by transdermal route in Parkinson's patients. The pharmacokinetics of the drug, followed in 24 most studies, put in evidence that the many important pharmacokinetic parameters were notably 25 improved versus the drug alone or in a commercial formulation. 26

**Keywords:** solid lipid nanoparticles; microemulsions; drug delivery system; central nervous system 28 29

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Introduction 32

The brain homeostasis is of primary importance 34 for survival so that specific interfaces, also 35 referred to as barriers, tightly regulate the 36 exchange between the peripheral blood circula-37 tion and the cerebrospinal fluid (CSF) circulatory 38 system. These barriers are represented by the 39 choroid plexus epithelium, the arachnoid epithe-40 lium, and the blood-brain barrier (BBB). The 41 concentration and clearance of endogenous and 42 exogenous molecules, essential for the normal 43 brain functions or dangerous because of their toxi-44 city, are strictly regulated by the anatomic and 45 physiologic features of each barrier (Abbott, 46 2002; Segal, 2000). 47

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The presence of the BBB is certainly the most critical issue encountered in brain drug delivery. Among the possible strategies to deliver therapeutic molecules into the brain, namely, intracerebral, intraventricular, and intravascular delivery, the latest represents the most reliable one because of its potential efficacy, safety, and compliance (Silva, 2007).

Brain capillaries, differently from the peripheral capillaries, present no fenestrae, a low amount of pinocytosis vesicles and particular tight junctions also known zonula occludens. Tight junctions are structures that form a narrow and continuous seal surrounding each endothelial and epithelial cell at the apical border and are at strictly regulating the movements the molecules through the paracellular

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<sup>01</sup> pathway. These structures, together with the brain
<sup>02</sup> endothelial cells, make an almost impermeable
<sup>03</sup> barrier for drugs administered through the
<sup>04</sup> peripheral circulation (Kniesel & Wolburg, 2000;
<sup>05</sup> Lapierre, 2000).

A further contribution to the peculiar BBB
functions is given by the periendothelial structures
represented by astrocytes, pericytes, and the basal
membrane (Balahanov & Dore-Duffy, 1998; Lay
& Kuo, 2005).

The presence of BBB transport systems further complicates the scenario. In fact, these transporters may assist or hinder the drug delivery to the brain. The carrier-mediated transport may be able to shuttle drugs or prodrugs into the brain in therapeutic concentrations, mimicking nutrients or endogenous compounds (Conford & Hyman, 18 1999; Pardrige, 1998).

<sup>19</sup> Unfortunately, the presence of active efflux
<sup>20</sup> transporters to the BBB also limits the therapeutic
<sup>21</sup> efficacy of drugs virtually able to access the brain.
<sup>22</sup> The P-glycoprotein (P-gp) is an ATP-dependent
<sup>23</sup> drug transport protein present at the apical mem<sup>24</sup> branes of different epithelial cell types including
<sup>25</sup> those forming the BBB.

Recently, it has been demonstrated, either *in vitro* or *in vivo*, that BBB P-gp can prevent the accumulation of many molecules including a variety of drugs in the brain (Stouch & Gudmundsson, 2002), and P-gp inhibition has been proposed as a possible strategy to enhance the drug penetration (Skinkel, 1999).

Different strategies have been studied for the 33 <sup>34</sup> delivery of drugs to the brain. Indeed most part of 35 the small drug molecules and of large molecules <sup>36</sup> such as recombinant proteins or gene-based molecules are not able to penetrate the BBB and many 37 38 efforts have been spent in the previous years 39 toward delivery and targeting of drugs to the 40 brain (de Boer & Gaillard, 2007). Many investiga-41 tions have been carried out in the previous years <sup>42</sup> to improve brain tumors therapy with nanoparti-<sup>43</sup> culates; there are less number of studies regarding 44 colloidal carriers of drugs for neurological diseases 45 or of diagnostics. Liposomes, polymeric nanopar-46 ticles, and solid lipid nanoparticles (SLN) have 47 been studied, with different approaches, and the

<sup>48</sup> problems of overcoming the BBB.

In this chapter, we consider SLN and microemulsions as carriers for the delivery only of drugs active on the central nervous system (CNS). In particular, examining drugs used for therapy in neurological diseases, as many times their administration gives problems, such as high amount of drug administered by parenteral route, short halflife, high hydrophilicity, and poor transport through the BBB. The aim of all the researchers is to study if some improvements in pharmacokinetic parameters in laboratory animals and/or in humans could be achieved using colloidal formulations; the review considers studies on SLN and microemulsions carrying only drugs active on CNS.

### Solid lipid nanoparticles

Different approaches are followed for the SLN preparation.

They can be prepared by high-pressure homogenization at elevated or low temperatures, via warm microemulsions, by solvent emulsification– evaporation–diffusion, by high-speed stirring, and/ or sonication (Muller, Kader, & Gohla, 2000).

Here we refer only about SLN carrying drugs active on CNS (at brain level).

SLN carrying the lipophilic antipsychotic drug clozapine were prepared by hot homogenization followed by ultrasonication method. Clozapine has a very poor bioavailability (Manjunath & Venkateswarlu, 2005). The SLN were administered by intravenous (IV) and duodenal routes to Swiss albino mice. For the intravenous administration, stearylamine was entrapped with clozapine in SLN; the area under curve (AUC) in the brain increased up to 2.91-fold the one of clozapine suspension.

The same authors (Manjunath & Venkateswarlu, 2006) developed SLN as carriers of the highly lipophilic drug nitrendipine, using different triglycerides for the lipid matrix, soy lecithin, and Poloxamer 188. Positive and negative charged nitrendipine SLN were also produced and then examined to explore the influence of the charge on oral bioavailability. The different kinds of SLN were administered by IV and intraduodenal

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nitrendipine SLN were examined, tissue distribu-02 tion studies were carried out in Swiss albino mice, 03 against that of a nitrendipine suspension. Follow-04 ing IV administration nitrendipine-loaded SLN 05 were found to be taken up to a greater extent in tested organs than nitrendipine suspension. The 07 AUC and MRT of nitrendipine SLN were higher AU3 08 than those of nitrendipine suspension especially in 09 brain and heart. Positively charged SLN were bet-10 11 ter taken up by the brain and moderately taken up by the heart. Reticuloendothelial system (RES) 12 organs such as liver and spleen were compared 13 with the ones after nitrendipine suspension admin-14 istration. The higher levels of the drug were main-15 tained for over 6h in confront to only 3h with 16 nitrendipine suspension. 17 SLN were investigated for their ability to deli-18

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ver quinine dihydrochloride for the management 19 of cerebral malaria (Gupta, Jain, & Jain, 2007). 20 Quinine was incorporated in SLN and successively 21 coupling of SLN with transferrin (Tf) was 22 achieved by a cross-linker. IV administration of 23 Tf-conjugated SLN enhanced the brain uptake of 24 quinine in confront to the SLN loaded of quinine 25 alone. 26

In order to enhance the delivery of atazanavir, a 27 HIV protease inhibitor, spherical SLN carrying 28 the drug were tested at first using a well-charac-29 terized human brain microvessel endothelial cell 30 line (hCMEC/D3). Cell viability experiments 31 demonstrated that SLN exhibit no toxicity on 32 hCMEC/D3 cells up to a concentration corre-33 sponding to 200 nM of the drug. Delivery of <sup>3</sup>H-34 atazanavir by SLN led to a significantly higher 35 accumulation by the endothelial cell monolayer 36 as compared to the drug aqueous solution (Chat-37 topadhyay, Zastre, Wong, Wu, & Bendayan, 38 2008). 39

The transport in situ of lipid nanoparticles to the 40 brain was evaluated by Koziara, Lockman, Allen, 41 and Mumper (2003); the lipidic nanoparticles were 42 prepared by warm microemulsion precursors fol-43 lowed by hot homogenization technique. Their 44 components were emulsified wax (E wax) or Brij 45 72 as matrix, and water and Brij 78 as surfactant. 46 The warm microemulsion was cooled upon stir-47 ring and the lipid SLN were obtained and 48

homogenized. The SLN were labelled with <sup>3</sup>H cetyl alcohol. The transport of the nanoparticles was measured by an "in situ" rat brain perfusion method; significant uptake of SLN was obtained . suggesting CNS uptake. The same group studied also the effect that the addition of a thiamine ligand to NPs, obtained by microemulsion as precursors, causes association with the BBB thiamine transporter (Lockman et al., 2003).

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Muller and coworkers studied the preferential adsorption of blood protein onto intravenously injected particulate carriers from different origins (Luck, Paulke, Schroder, Blunk, & Muller, 1998); in particular, Apolipoprotein E (Apo E) on the surface of P80-coated SLN after their incubation in human plasma citrate. Delivery to the brain using nanoparticulate drug carriers in combination with the targeting principles of "differential protein adsorption" has been proposed (Dehouck et al., 1997). The Pathfinder technology (Muller & Schmidt, 2002) exploits proteins present in the blood which absorb onto the surface of intravenously injected carriers for targeting nanoparticles to the brain. Apo E is one of such targeting molecules for the delivery of nanoparticles to the endothelial cells of the BBB. Apo E can play an important role in the transport of lipoprotein into brain via the low-density lipoprotein receptor present on the BBB. Atoquavone (Muller & Keck, 2004; Scholler et al., 2001) is a drug poorly adsorbed after oral administration, showing poor therapeutic efficacy against toxoplasma encephalitis (TE). Nanocrystals of the drug were produced, their surface was modified with Tween 80 leading to in vivo preferential absorption of Apo E; the nanosuspension was IV administered to a murine model of TE, obtaining the disappearance of parasites and of cysts at dose 10-fold smaller than the one of atoquavone administered by oral route.

### Solid lipid nanoparticles from warm microemulsions

SLN can be achieved from warm microemulsions.

Warm microemulsions are prepared at temperature ranging from 60°C to 80°C by using AU7

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on melted lipids (such as triglycerides/fatty acids) as oil, surfactants such as lecithin, and cosurfactants 02 03 (such as short-chain carboxylates, biliar salts); the warm microemulsions are subsequently dispersed in cold water. The nanodroplets of warm micro-05

emulsion, using this procedure, become SLN; they

are successively washed by tangential flow filtra-07

tion. SLN are spherical in shape and with a narrow 08

<sup>09</sup> size distribution .The zeta potential is normally <sup>10</sup> high (30/40 mV) being positive or negative 11 depending on the starting formulation.

Hydrophilic and lipophilic molecules (drugs or 12 <sup>13</sup> diagnostics) can be incorporated in SLN using <sup>14</sup> different methods.

SLN are able to carry drugs of different struc-15 <sup>16</sup> ture and lipophilicity, such as cyclosporine A 17 (Ugazio, Cavalli, & Gasco, 2002), paclitaxel 18 (Cavalli, Caputo, & Gasco, 2000), doxorubicin (Fundaro, Cavalli, Bargoni, Vighetto, & Gasco, 19 20 2000), tobramycin (Cavalli et al., 2003), short-<sup>21</sup> chain fatty acids (Dianzani et al., 2006), peptides 22 (Morel, Cavalli, & Gasco, 1996), antisense oligo-<sup>23</sup> nucleotides (Brioschi et al., 2008), and melatonin <sup>24</sup> (MT) (Rezzani et al., 2009). Also diagnostic com-

pounds such as iron oxides (Pereira, 2003) have AU10 25 <sup>26</sup> been incorporated into SLN.

SLN can be internalized within 2-3 min into all 28 the tested cell lines (Miglietta, Cavalli, Bocca, 29 Gabriel, & Gasco, 2000; Serpe et al., 2006); admi-30 nistered by duodenal route and are targeted to 31 lymph (Bargoni et al., 1998). SLN stealth can 32 also be prepared to avoid their recognition by <sup>33</sup> the RES, thus prolonging their residence time AU11 <sup>34</sup> (Podio, 2001). SLN drug, unloaded or loaded, <sup>35</sup> stealth/or nonstealth, are transported through the

BBB (Podio, 2001; Zara et al., 2002). 36 37

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#### 39 Drug-loaded solid lipid nanoparticles

<sup>41</sup> In the late 1990s SLN were proposed for brain <sup>42</sup> drug targeting by several groups (Yang, Zhu, Lu, <sup>43</sup> & Liang, 1999; Zara et al., 1999), which studied <sup>44</sup> the pharmacokinetics of two anticancer agents: 45 camptothecin and doxorubicin. After oral and IV administration, they observed drug accumulation 47 into the brain.

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Both stealth and nonstealth stearic acid unloaded labelled SLN were found in rat CSF 20 min after IV administration even though low amount of radioactivity was found in the CSF samples collected from cysterna magna (Podio, Zara, Carazzone, Cavalli, & Gasco, 2000).

When the same kind of SLN were loaded with doxorubicin, significantly higher drug concentrations were found in the brain of the animals treated with stealth SLN as compared to nonstealth SLN and doxorubicin solution. The overall plasma pharmacokinetics of stealth and nonstealth SLN provided to be significantly different from that of the doxorubicin solution (Fundaro et al., 2000).

*R*-apomorphine (10,11-dihydroxyapomorphine) is a well-known potent short-acting dopamine agonist at D1 and D2 dopamine receptors and it was proposed as an antiparkinsonian drug more than a century ago. It significantly reduces the severity and duration of "off" periods and it is able to reverse bradykinesia when administered alone. Despite these favorable clinical effects, the drug's clinical use is somewhat limited by its pharmacokinetic profile: short half-life ( $\sim 30 \text{ min}$ ), rapid clearance from the plasma, lack of storage and retention in brain regions, poor oral bioavailability (5%), and first-pass hepatic metabolism are significant limitations to chronic oral administration. Our group evaluated a new formulation of apomorphine in SLN (submitted data for publishing); the study was designed to investigate the AU12 pharmacokinetics and biodistribution of apomorphine incorporated in SLN, injected orally or intravenously in rats.

In vitro the release over time of apomorphine from the SLN dispersion was almost linear After IV administration the peak plasma concentration was higher after apomorphine solution administration than after apomorphine SLN. However, the total area under curve (AUCtot) was nonsignificantly different after SLN than apomorphine solution. The terminal half-life was significantly longer following apomorphine SLN.

Following intraduodenal administration we found that the  $C_{\text{max}}$  and AUC<sub>tot</sub> were significantly higher with apomorphine SLN compared to apomorphine solution; on the contrary, the clearance

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was shorter after apomorphine solution than after 01 the SLN formulation. 02

In the brain the apomorphine concentration 03 was significantly higher 30 min after apomorphine 04 SLN IV administration versus solution; it was 05 detected only at 4h after apomorphine SLN 06 injection. 07

After duodenal administration the drug was 08 detectable in brain only at 30 min after apomor-09 phine SLN administration. No drug was found 10 neither at 4 h nor at 24 h after injection of either apomorphine SLN or the solution. 12

Furthermore, the free drug concentration was 13 measured in human plasma and we showed that 14 the release started after the absorption of the 15 apomorphine SLN. We also measured the free 16 apomorphine concentration in human blood over 17 time. The amounts in question are relatively low, 18 but may be sufficient to expect clinical effects 19 when administered to parkinsonian patients. 20 After apomorphine solution administration, the 21 amounts of apomorphine determined in the 22 plasma were by far lower than those from SLN, 23 confirming previous studies on the duodenal 24 administration of drug loaded and unloaded SLN 25 (Fig. 1). 26

In order to furnish a general model for SLN-21 based delivery systems of drugs devoid of favor-28 pharmacokinetics, we have recently able 29 30

incorporated MT in SLN (MT-SLN). MT has been chosen for our in vivo study because of its safeness in humans even at high dosages.

MT is a hormone produced by the pineal gland at night, involved in the regulation of circadian rhythms. For clinical purposes (mainly disorders of the sleep-wake cycle and insomnia in the elderly), exogenous MT administration should mimic the typical nocturnal endogenous MT levels, but its pharmacokinetics is not favorable due to its short half-life of elimination (DeMuro, Nafziger, Blask, Menhinick, & Bertino, 2000; Mallo et al., 1990). The pharmacokinetics of MT-SLN has been examined in humans after administration by oral and transdermal route (Priano et al., 2007). Three kinds of freeze-dried MT-SLN containing different amounts of MT were prepared and characterized: (a) MT-SLN: MT = 1.8% for in vitro experiments (average diameter: 85 nm, polydispersity index = 0.135); (b) MT-SLN: MT = 2% for transdermal application (average diameter = 91 nm, polydispersity index = 0.140); and (c) MT-SLN: MT = 4.13% for oral route (average diameter = 111 nm, polydispersity index = 0.189).

In vitro, MT-SLN produced a flux of MT of 1  $\mu$ g/h/cm<sup>2</sup> through hairless mice skin, following a pseudo-zero-order kinetics (45). At the same time, AU13 in vivo study produced very interesting results,





#### 47 Fig. 1. Plasma levels of free apomorphine and total apomorphine after duodenal administration of apomorphine solution or apomorphine SLN in rats. 48

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of confirming in humans that SLN can act as a reservoir that allows a constant and prolonged release 02 of the included drugs (Peira et al., 2003). MT 03 (3 mg) incorporated in SLN was orally adminis-04 tered at 8.30 a.m. to seven healthy subjects; for 05 control purposes, 1 week later the same subjects 06 07 received orally a standard formulation of MT at the same dose (3 mg) and again at 8.30 a.m. Com-08 09 pared to the MT standard solution,  $T_{\text{max}}$  observed 10 after MT-SLN administration was delayed of 11 about 20 min, while mean AUC and mean half-12 life of elimination were significantly higher 13 (respectively 169,944.7 + 64,954.4 pg/mL × hour vs.  $85,148.4 \pm 50,642.6$  pg/mL × hour, p = 0.018; 14 and 93.1 + 37.1 min vs. 48.2 + 8.9 min, p = 0.009). 15 Even more, standard formulation and MT-SLN 16 17 after oral administration produced similar peak plasma levels of MT, even if delayed of about 18 half an hour in the case of MT-SLN. More inter-19 estingly, detectable and clinically significant MT 20 plasma levels after MT-SLN oral administration 21 were maintained for a longer period of time, sug-22 gesting that SLN orally administered to humans 23 can yield a sustained release of the incorporated 24 drug, a feature that could be particularly useful for 25 molecules, such as MT, characterized by unfavor-26 able kinetics (Priano et al., 2007). Previous studies 27 28

in laboratory animals indicated a probable targeting of SLN - either drug-loaded or unloaded to lymph, after duodenal administration (Bargoni et al., 1998). Similarly, the significantly longer half-life of MT observed in the study of Priano et al. (2007) may suggest a targeting of MT-SLN to human lymph, even though the capsules used to administer SLN were not gastro-resistant. In fact, MT half-life of elimination has been calculated in about 40 min after an intravenous bolus and following oral administration low bioavailability and rapid clearance from plasma have been shown, primarily due to a marked first-pass hepatic metabolism. Moreover, pharmacokinetic analysis following transdermal administration of MT-SLN demonstrated that plasma levels of MT similar to those produced by oral administration may be achieved for more than 24 h (50). In 10 healthy subjects, SLN incorporating MT were administered transdermally by applying a patch at 8.30 a.m. and leaving it in place for 24 h. In this delivery system, MT absorption and elimination were slow (mean half-life of absorption =  $5.3 \pm 1.3$  h; mean half-life of elimination =  $24.6 \pm 12.0$  h) so that MT plasma levels above 50 pg/mL were maintained for at least 24 h (Figs. 2 and 3). Tolerability of MT-SLN administered transdermally or by oral



Fig. 2. MT plasma profile in humans after MT (♦) and MT-SLN (■) oral administration.

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<sup>17</sup> Fig. 3. MT plasma levels in humans at baseline and after MT <sup>18</sup> SLN transdermal administration (MT-SLN-TD).

route was good and no adverse effect occurred,
apart from a predictable mild somnolence and
transient erythema after gel application. This
means that, at least at the doses used in that
study (45), SLN administration via the oral or
transdermal routes is safe.

In this context, we also tested transdermal MT-26 SLN for three consecutive nights in five patients 27 suffering from delayed sleep phase syndrome 28 (unpublished data), confirming the safeness of 29 this formulation. Due to the small sample, how-30 ever, the tendency of clinical benefits was present 31 but statistical significance could not be reached, so 32 that further investigations in larger samples are 33 needed in order to evaluate the impact of this 34 new formulation in clinical practice. 35

However, these very favorable results, obtained 36 in humans administering MT-loaded SLN, clearly 37 suggest that SLN can be considered effective in 38 vivo delivery systems that could be suitably 39 applied to different drugs, and in particular to 40 those requiring prolonged high plasma levels but 41 that have unfavorable pharmacokinetics. Finally, 42 it must be stressed that, since doses and concen-43 trations of drugs included in SLN can be varied, 44 different plasma level profiles could be obtained, 45 thus disclosing new chances for sustained delivery 46 systems adaptable to a variety of clinical condi-47

<sup>48</sup> tions (Priano et al., 2007).

Suitability of SLN to convey drugs into CNS is also confirmed by studies regarding baclofen included in SLN. Intrathecal baclofen administration represents the reference treatment for spasticity of spinal or cerebral origin. Nevertheless, surgical involvement together with risk of infection or catheter dysfunction may limit the number of potentially treatable patients (Dario & Tomei, 2004; Perot & Almeida-Silveira, 1994). In order to explore alternative and efficacious routes of administration, we studied a new pharmaceutical preparation characterized by SLN incorporating baclofen (baclofen-SLN) (submitted data for publishing). Baclofen concentration, after reconstitu- AU14 tion with water of freeze-dried SLN, was 1.7 mg/ mL. Groups of Wistar rats were injected intraperitoneally with physiological solution and unloaded SLN at 10 mL/kg (control groups), with baclofen-SLN (baclofen-SLN group), and baclofen solution (baclofen-sol group) at increasing dosages of 2.5, 5, 7.5, 8.5, and 10 mg/kg. At different times up to the fourth hour, efficacy testing was performed by means of H-reflex, while behavioral characterization was obtained using two scales validated for motor symptoms due to spinal lesions and sedation in rat models (Nemethy, Paroli, Williams-Russo, & Blanck, 2002; Tsunoda, Kuang, Tolley, Whitton, & Fujinami, 1998). Rats were sacrificed for detecting baclofen concentration in blood and tissue. Compared to baclofen-sol and control group, H/M amplitude curve after baclofen-SLN injection was characterized by a dose-dependent reduction at the first and second hours, so confirming efficacy, and a rebound increase at the fourth hour, indicating an unexpected belated spinal hyperexcitability (Fig. 4). Similarly, baclofen-SLN effect on behavioral scales was stronger compared to baclofen-sol group, with the maximum effects obtained at the first hour. Moreover, clinical effects were detectable after low dosages of baclofen-SLN (2.5 mg/kg) but only after higher dosages of baclofen-sol (7.5 mg/kg). After 4 h from the injection, only the rats treated with the higher dosages of baclofen-SLN still presented clinical signs consisting in sedation (8.5 mg/kg) or complete paralysis and piloerection (10 mg/kg). On the whole, these data suggest a dose-dependent modulation of spinal reflex excitability, which is

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<sup>14</sup> Fig. 4. H/M amplitude ratios after baclofen-solution (a) and after baclofen-SLN (b), at increasing doses, compared to control animal <sup>15</sup> group. \*p < 0.05; \*p < 0.01; \*\*p < 0.001.

18 not so evident after administration of standard 19 formulation of baclofen. Nevertheless, important 20 cortical effects were also present. Clinical data 21 were related with plasma and tissue concentra-22 tions. In fact, after 2 and 4h only baclofen-SLN 23 administration produced measurable baclofen 24 plasma concentrations, with an almost linear 25 decrease of baclofen appreciable for 4 h. On the 26 contrary, undetectable amount of baclofen in 27 plasma were noticed 2h after administration of 28 baclofen-sol. In brain, both the two formulations 29 (baclofen in solution and in SLN) gave a maxi-30 mum after 2 h but concentrations after SLN were 31 almost twice the ones after solution. This last data 32 might be due partly to the free drug already 33 released and to baclofen-SLN overcoming the 34 BBB. We realize that for clinical purposes this 35 effect of baclofen-SLN is unwished, as it is responsible for sedation. However, baclofen-sol injec-37 tions also produced sedation, even if weaker and 38 corresponding to lower plasma concentrations, 39 compared to baclofen-SLN. In conclusion, higher 40 spinal and cortical effects of baclofen-SLN, com-41 pared to equivalent dosages of baclofen-sol, seem 42 attributable to higher and more prolonged con-43 centrations of drugs in plasma and brain. 44

<sup>44</sup> As previously noted, unloaded SLN adminis-<sup>45</sup> tered by duodenal route are targeted to lymph <sup>46</sup> and the incorporated drug can be partly distribu-<sup>47</sup> ted in the brain; moreover, SLN can also be <sup>48</sup> prepared stealth for increasing their residence time (Bargoni et al., 1998; Fundaro et al., 2000; Podio, 2001; Zara et al., 2002). Other new studies will be directed toward a duodenal administration of baclofen-SLN stealth, not only for prolonging their residence time but also to target them to lymph, enhancing their bioavailability. Further research should also be directed toward the optimization of dosages and concentrations of baclofen included in SLN, in order to preserve the prolonged antispastic effect, peculiar of this new formulation, but devoid of clinically significant cortical effects.

### Solid lipid nanoparticles as potential diagnostics

Superparamagnetic iron oxides are classified as contrast agents for magnetic resonance imaging (MRI). They are able to affect the water relaxation times  $T_1$  and  $T_2$ ; their ability in altering such properties is quantified by the parameter relaxivity. Iron oxides are able to affect preferentially the  $T_2$  relaxation times of tissues (and are called  $T_2$ -relaxing agents) while paramagnetic contrast agents such as Gd complexes affect mainly  $T_1$  and are called  $T_1$ -relaxing agents.

Iron oxides are insoluble in water; therefore, to be clinically used they must be transformed in modified colloids while their magnetic properties <sup>01</sup> should remain unchanged. The surface of the iron
<sup>02</sup> oxide nanoparticles can be modified, covering

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them by hydrophilic macromolecules; such as dex tran in the case of Endorem.

Chapter 10

A research was performed in order to know 05 whether SLN can load iron oxides and whether 06 they are able to reach the brain. Two different 07 SLN, SLN-Fe<sup>A</sup> and SLN<sup>B</sup> containing iron oxides AU16 08 were prepared from warm microemulsions and 09 studied at first in vitro (1). The comparison of 10 Fe-SLN was performed with Endorem. Both the Fe-SLN preparations showed relaxometric prop-12 erties similar to the ones of Endorem. The good 13  $T_2$ -relaxation-enhancing properties allow an *in* 14 vivo study of their distribution by MRI. Fe-15 SLN<sup>B</sup>, at the higher Fe concentration, were 16 administered IV to rats; the comparison was 17 performed with Endorem. Images obtained 18 after Endorem IV administration show early 19 modification, but soon return to baseline; these 20 findings are consistent with short Endorem 21 retention time in blood. Results from SLN-Fe<sup>B</sup> 2.2 show a different behavior. For each part of the 23 AU17 brain, maximal SS is reached in the last images 24 (135 min after administration). SS increase from 25 the first to the last acquisition. This study shows 26 that after inclusion in SLN, Endorem becomes a 27 new type of contrast agent: Endorem is taken by 28 the liver and does not cross the BBB, while 29 Endorem containing SLN-Fe<sup>B</sup> shows CNS 30 uptake. This means that SLN-Fe kinesis is 31 related to SLN and not to their iron oxide con-32 tent as already seen. 33

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### 36 Microemulsions

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Microemulsions are transparent, thermodynami-38 cally stable dispersions of water and oil, usually 39 stabilized by a surfactant and a cosurfactant. They 40 contain particles smaller than 0.1 µm. Microemul-41 sions are often defined as thermodynamically 42 stable liquid solutions; the stability of microemul-43 sions is a consequence of the ultralow interfacial 44 tension between the oil and water phases. A clear 45 distinction exists between microemulsion and 46 coarse emulsions. The latter are thermodynami-47 48 cally unstable, droplets of their dispersed phase

are generally larger than  $0.1 \,\mu\text{m}$  and consequently their appearance is normally milky rather than transparent.

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The limits in the use of microemulsions in the pharmaceutical field are chiefly from the need of all the components to be acceptable, particularly surfactants and cosurfactants — the amounts of surfactants and cosurfactants required to form microemulsions are usually higher than those required for emulsions.

Recently, apomorphine was incorporated into microemulsions to study whether they are a feasible vehicle for transdermal transport of this drug. In the preparatory in vitro study (Peira, Scolari, & Gasco, 2001), two different microemulsions whose components were all biocompatible were studied: the concentration of apomorphine was 3.9% in each. Since apomorphine is highly hydrophilic, to increase its lipophilicity, apomorphine-octanoic acid ion pairs were formed. At pH 6.0, log  $P_{app}$ of apomorphine increased from 0.3 in the absence of octanoic acid to  $\log P_{app} = 2.77$  for a molar ratio 1:2.5 (apomorphine: octanoic acid). The flux of drug from the two thickened microemulsions through hairless mouse skin was, respectively, 100 and  $88 \mu g/h/cm^2$ . The first formulation, having the higher flux, was chosen for in vivo administration to Parkinson's patients.

For the in vivo study, 21 patients with idiopathic Parkinson's disease who presented long-term L-DOPA syndrome, motor fluctuations and prolonged "off" periods were selected (Priano et al., 2004). Here, 10 g of apomorphine hydrochloride (3.9%), included in microemulsion for transdermal delivery (Apo-MTD), was applied to a 100 cm<sup>2</sup> skin area on the chest; the area was delimited by 1-mm-thick biocompatible foam tape and covered with a polyester-based membrane and an occlusive membrane to prevent evaporation. In these conditions, a single layer of microemulsion (1 mm thick) was directly in contact with the skin surface and acted as a reservoir of apomorphine. Apo-MTD was applied at 8.00 a.m. and left for 12 h. In all patients, except two, apomorphine was detected in blood samples after a variable lag time. Pharmacokinetic analysis revealed that epicutaneous-transdermal apomorphine absorption was rapid (mean half-life of absorption = 1.03 h)

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of with a variability among patients (half-life of absorption, SD = 1.39 h). Mean  $C_{\text{max}}$  was above the therapeutic range (mean  $C_{\text{max}} = 42.81 \pm 11.67$ 03 ng/mL), with a mean  $T_{\text{max}}$  of 5.1  $\pm$  2.24 h. Ther-04 os apeutic concentrations of apomorphine were reached after a mean latency of 45 min (range 06 <sup>07</sup> 18–125), and stable concentrations, above the <sup>08</sup> therapeutic range, continued for as long as Apo-<sup>09</sup> MTD was maintained in place. At the 12th hour, <sup>10</sup> Apo-MTD was removed, and the apomorphine II plasma concentration then decreased at a rate <sup>12</sup> comparable to that described for subcutaneous administration (mean half-life of elimination 13 <sup>14</sup> equal to  $10.8 \pm 1.93$  h).  $C_{\text{max}}$  and AUC showed good correlations with the reduction of "off" per-15 iods duration and with the improvement of clinical 16 17 scores evaluating motor performances (r values <sup>18</sup> ranging form 0.49 to 0.56, with p values ranging from 0.02 to 0.04). Apo-MTD overall tolerability 19 was good: systemic side effects were similar to 20 those caused by subcutaneous apomorphine injec-21 tion (sleepiness, mild orthostatic hypotension, and 22 transient nausea), and in the case of nausea, they 23 were strictly related to the highest plasma level of 24 apomorphine. Moreover, regarding local side 25 <sup>26</sup> effects, the large majority of patients (71.4%) pre-27 sented a transient mild erythema at the site of Apo-MTD application, with a complete regression 28 within 48 h, whereas only in two cases the 29 <sup>30</sup> erythema lasted more than 3 days and required 31 local therapy. This study clearly demonstrated 32 that in most Parkinson's patients Apo-MTD is absorbed by the epicutaneous-transdermal route. 33 This result is in contrast with other reports, where 34 the transdermal route did not produce detectable 35 plasma levels of apomorphine, or in which no apo-36 morphine was transported passively through the skin 37 (Gancher, Nutt, & Woodward, 1991; van der Geest, 38 van Laar, Gubbens-Stibbe, Boddé, & Danhof, 39 <sup>40</sup> 1997). Probably, this difference was mainly due to the peculiar pharmaceutical preparation used. Even 41 42 if pharmacokinetic parameters are variable, Apo-<sup>43</sup> MTD demonstrated the feasibility of providing ther-<sup>44</sup> apeutic apomorphine plasma levels for much longer periods of time than previously tested apomorphine 45 <sup>46</sup> preparations (several hours), allowing a more con-47 stant dopaminergic stimulation. These results are 48 encouraging and Apo-MTD might become of

clinical value in some parkinsonian patients suffering from uncontrolled "wearing-off" and prolonged "off" phenomena. On the contrary, because of the lag time of about 1 h before therapeutic concentrations are reached, Apo-MTD may not be the "ideal" preparation for rapid relief of "off" periods.

Since Apo-MTD was found to provide constant drug release over several hours, other studies have been addressed to its use for the nocturnal sleep disorders of Parkinson's patients. Twelve parkinsonian patients underwent standard polysomnography on basal condition and during one night treatment with Apo-MTD (applied to  $100 \text{ cm}^2$ from 10 p.m. until 8 a.m.; Priano et al., 2003). Sleep analysis during APO-MTD treatment in comparison to basal condition showed very favorable findings: 16% increment of total sleep time, 12% increment of sleep efficiency, 16% increment of stage 3 and 4 nonrapid eye movement (NREM), 15% reduction of periodic limb movements index, 22% reduction of arousal index, and 23% reduction of the "cycling alternating pattern" rate, an objective measure of disruption and fragmentation of NREM sleep. Pharmacokinetic analysis confirmed the absorption of apomorphine and the maintenance of therapeutic plasma levels for several hours (mean  $C_{\text{max}} = 31.8 \pm 9.7$  ng/mL; mean  $T_{\text{max}} = 3.1 \pm 1.6$  h; mean half-life of absorption =  $1.2 \pm 1.4$  h; mean half-life of elimination =  $8.8 \pm 1.9$  h). On the whole, this study confirmed that APO-MTD in Parkinson's disease might be able to reduce nocturnal anomalous movements, akinesia, and rigidity, and might be efficacious for reducing the instability of sleep maintenance typical of parkinsonian sleep.

### References

- Abbott, N. J. (2002). Astrocyte-endothelial interactions and bloodbrain-barrier permeability. *Journal of Anatomy*, 200, 629–638.
- Balahanov, R., & Dore-Duffy, P. (1998). Role of the CNS microvascular pericyte in the blood-brain barrier. *Journal* of Neuroscience Research, 53, 637–644.
- Bargoni, A., Cavalli, R., Caputo, O., Fundaro, A., Gasco, M. R., & Zara, G. P. (1998). Solid lipid nanoparticles in lymph and plasma after duodenal administration to rats. *Pharmaceutical Research*, 15, 745–750.
- Brioschi, A., Calderoni, S., Pradotto, L. G., Guido, M., Strada, A., Zenga, F., et al. (2008). Solid lipid nanoparticles

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183

- carrying oligonucleotides inhibit vascular endothelial grow
   factor expression in rat glioma models. *Journal of Nanoneuroscience*, 1, 1–10.
- <sup>03</sup> Cavalli, R., Caputo, O., & Gasco, M. R. (2000). Preparation
- <sup>04</sup> and characterization of solid lipid nanospheres containing
- paclitaxel. European Journal of Pharmaceutical Science, 10,
   305–308.
- Cavalli, R., Bargoni, A., Podio, V., Muntoni, E., Zara, G. P., &
- Gasco, M. R. (2003). Duodenal administration of solid lipid nanoparticles loaded with different percentages of tobramy-
- <sup>o</sup> cin. Journal of Pharmaceutical Sciences, 92, 1085–1094.
- <sup>10</sup> Chattopadhyay, N., Zastre, J., Wong, H. L., Wu, X. Y., & <sup>11</sup> Bendayan, R. (2008). Solid lipid nanoparticles enhance the
- $_{12}$  delivery of the HIV protease inhibitor, atazanavir, by a
- human brain endothelial cell line. *Pharmaceutical Research*, 25, 2262–2271.
- <sup>14</sup> Conford, F. M., & Hyman, S. (1999). Blood-brain barrier per-<sup>15</sup> meability to small and large molecules. *Advanced Drug*
- 16 Delivery Reviews, 36, 145–163.
- <sup>17</sup> Dario, A., & Tomei, G. (2004). A benefit-risk assessment of baclofen in severe spinal spasticity. *Drug Safety*, *27*, 799–818.
- de Boer, A. G., & Gaillard, P. J. (2007). Drug targeting to the
   <sup>19</sup> brain. Annual Review of Pharmacology and Toxicology, 47,
- <sup>20</sup> 323–355. <sup>21</sup> Dehouck, R., Ferrari, L., Dehouck, M. P., Pierce, A.,
- <sup>22</sup> Torpier, G., & Cecchelli, R. (1997). A new function for the LDL receptor. Transcytosis of LDL across the blood brain
- <sup>23</sup> barrier. Journal of Cell Biology, 38, 877–889.
- <sup>24</sup> DeMuro, R. L., Nafziger, A. N., Blask, D. E., Menhinick, A. M.,
- <sup>25</sup> & Bertino, J. S. (2000). The absolute bioavailability of oral melatonin. *Journal of Clinical Pharmacology*, 40, 781–784.
- Diamani C. Cauelli D. Zara C. D. Calliashia M. Lambardi
- <sup>27</sup> Dianzani, C., Cavalli, R., Zara, G. P., Gallicchio, M., Lombardi, G.,
   <sup>28</sup> Gasco, M. R., et al. (2006). Cholesteryl butyrate solid lipid
   <sup>28</sup> nanoparticles inhibit adhesion of human neutrophils to endothe-
- <sup>29</sup> lial calls. *Pritich Journal of Pharmacology* 148, 648, 656
- <sup>29</sup> lial cells. *British Journal of Pharmacology*, *148*, 648–656.
   <sup>30</sup> Fundaro A. Cavalli B. Bargoni A. Vighetto G. P. & G.
- <sup>30</sup> Fundaro, A., Cavalli, R., Bargoni, A., Vighetto, G. P., & Gasco,
- M. R. (2000). Non-stealth and stealth solid lipid nanoparti-
- <sup>32</sup> cles (SLN) carrying doxorubicin: Pharmacokientics and tissue distribution after i.v. administration to rats.
- <sup>33</sup> *Pharmacological Research*, *42*, 337–343.
- <sup>34</sup> Gancher, S. T., Nutt, J. G., & Woodward, W. R. (1991).
- Absorption of apomorphine by various routes in parkinsonism. *Movement Disorders*, 6, 212–216.
- Gupta, Y., Jain, A., & Jain, S. K. (2007). Transferrin conjugate
- solid lipid nanoparticles for enhanced delivery of quinine
- <sup>38</sup> dihydrochloride to the brain. *Journal of Pharmacy and Phar-*<sup>39</sup> *macology*, *59*, 935–940.
- Kniesel, K., & Wolburg, H. (2000). Tight junctions of the blood
   brain barrier. *Cellular and Molecular Neurobiology*, 20, 57–76.
- <sup>42</sup> Koziara, J. M., Lockman, P. R., Allen, D. D., & Mumper, R. J. (2003). In situ blood-brain barrier transport of nanoparticles.
- <sup>43</sup> Pharmaceutical Research, 20, 1772–1777.
   <sup>44</sup> Logistro L. A. (2000). The molecular structure of the tight
- Lapierre, L. A. (2000). The molecular structure of the tight
   junctions. Advanced Drug Delivery Reviews, 41, 255–264.
- 46 Lay, C. H., & Kuo, K. H. (2005). The critical component to
- establish in vitro BBB model: Pericyte. Brain Research
- *Reviews*, 50, 258–265.

- Lockman, P. R., Oyewumi, M. O., Koziara, J. M., Roder, K. E., Mumper, R. J., & Allen D. D. (2003). Brain uptake of thiaminecoated nanoparticles. *Journal of Controlled Release*, 93, 271–282.
- Luck, M., Paulke, B. R., Schroder, W., Blunk, T., & Muller, R. H. (1998). Analysis of plasma protein adsorption on polymeric nanoparticles with different surface characteristics. *Journal of Biomedical Materials Research*, 39, 478–485.
- Mallo, C., Zaidan, R., Galy, G., Vermeulen, E., Brun, J., Chazot, G., et al. (1990). Pharmacokinetics of melatonin in man after intravenous infusion and bolus injection. *European Journal of Clinical Pharmacology*, 38, 297–301.
- Manjunath, K., & Venkateswarlu, V. (2005). Pharmacokinetics, tissue distribution and bioavailability of clozapine solid lipid nanoparticles after intravenous and intraduodenal administration. *Journal of Controlled Release*, 107, 215–228.
- Manjunath, K., & Venkateswarlu, V. (2006). Pharmacokinetics, tissue distribution and bioavailability of nitrendipine solid nanoparticles after intravenous and intraduodenal administration. *Journal of Drug Targeting*, 14, 632–645.
- Miglietta, A., Cavalli, R., Bocca, C., Gabriel, L., & Gasco, M. R. (2000). Cellular uptake and cytotoxicity of solid lipid nanospheres (SLN) incorporating doxorubicin or paclitaxel. *International Journal of Phamaceutics*, 210, 61–67.
- Morel, S., Cavalli, R., & Gasco, M. R. (1996). Thymopentin in solid lipid nanoparticles. *International Journal of Phamaceutics*, 132, 259–262.
- Muller, R. H., Kader, K., & Gohla, S. (2000). Solid lipid nanoparticles (SLN) for controlled drug delivery – a review of the state of the art. *European Journal of Pharmaceutics and Biopharmaceutics*, 50, 161–177.
- Muller, R. H., & Schmidt, S. (2002). Path. Finder technology for the delivery of drugs to the brain. *New Drugs*, *2*, 38–42.
- Muller, R. H., & Keck, C. M. (2004). Challenges and solutions for the delivery of biotech. Drugs — a review of drug nanocrystal technology and lipid nanoparticle. *Journal of Biotechnology*, *113*, 151–170.
- Nemethy, M., Paroli, L., Williams-Russo, P. G., & Blanck, T. J. (2002). Assessing sedation with regional anesthesia: Interrater agreement on a modified Wilson sedation scale. *Anesthesia and Analgesia*, 94, 723–728.
- Pardrige, W. M. (1998). CNS drug design based on principles of blood-brain barrier transport. *Journal of Neurochemistry*, 70, 1781–1792.
- Peira, E., Scolari, P., & Gasco, M. R. (2001). Transdermal permeation of apomorphine through hairless mouse skin from microemulsions. *International Journal of Phamaceutics*, 226, 47–51.
- Peira, E., Marzola, P., Podio, V., Aime, S., Sbarbati, A., & Gasco, M. R. (2003). In vitro and in vivo study of solid lipid nanoparticles loaded with superparamagnetic iron oxide. *Journal of Drug Targeting*, 11, 19–24.
- Perot, C., & Almeida-Silveira, M. I. (1994). The human H and T reflex methodologies applied to the rat. *Journal of Neuroscience Methods*, 51, 71–76.
- Podio, V., Zara, G. P., Carazzone, R., Cavalli, R., & Gasco, M. R. (2000). Biodistribution of stealth and non-stealth solid

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- lipid nanoparticles after intravenous administration in rats.
   *Journal of Pharmacy and Pharmacology*, 52, 1057–1063.
- <sup>60</sup> Priano, L., Albani, G., Brioschi, A., Guastamacchia, G.,
   <sup>63</sup> Calderoni, S., Lopiano, L., et al. (2003). Nocturnal anom-
- <sup>04</sup> alous movement reduction and sleep microstructure ana-
- <sup>05</sup> lysis in parkinsonian patients during 1-night transdermal
- apomorphine treatment. Neurological Science, 24,
   207–208.
- Priano, L., Albani, G., Brioschi, A., Calderoni, S., Lopiano, L.,
- <sup>08</sup> Rizzone, M., et al. (2004). Transdermal apomorphine per <sup>09</sup> meation from microemulsions: A new treatment in Parkin-
- <sup>10</sup> son's disease. *Movement Disorders*, *19*, 937–942. <sup>11</sup> Priano, L., Esposti, D., Esposti, R., Castagna, G., De Medi
- Priano, L., Esposti, D., Esposti, R., Castagna, G., De Medici, C., Fraschini, F., et al. (2007). Solid lipid nanoparticles incor-
- <sup>12</sup> porating melatonin as new model for sustained oral and
- transdermal delivery systems. Journal of Nanoscience and
   Nanotechnology, 7, 3596–3601.
- 15 Rezzani, R., Fabrizio Rodella, L., Fraschini, F., Gasco, M. R.,
- <sup>16</sup> Demartini, G., Musicanti, C., et al. (2009). Melatonin delivery
- <sup>17</sup> in solid lipid nanoparticles: Prevention of cyclosporin A induced cardiac damage. *Journal of Pineal Research*, *46*, 255–261.
- <sup>18</sup> Segal, M. B. (2000). The choroid plexuses and the barriers
   <sup>19</sup> between the blood and the cerebrospinal fluid. *Cellular and*
- 20 Molecular Neurobiology, 20, 183–196.
- Scholler, N., Krause, K., Kayser, O., Muller, R. H., Borner, K.,
   Hahn, H., et al. (2001). Atovaquone nanosuspensions show
- excellent therapeutic effect in a new murine model of reactivated toxoplasmosis. *Antimicrobial Agents and Chemother*-
- <sup>24</sup> apv, 45, 1771–1779.
- <sup>25</sup> Skinkel, A. H. (1999). P-glycoprotein, a gatekeeper in the
- blood-brain barrier. Advanced Drug Delivery Reviews, 36,
   179–194.
- Serpe, L., Cavalli, R., Gasco, M. R., Muntoni, E., Cavalli, R.,
- <sup>28</sup> Panzanelli, P., et al. (2006). Intracellular accumulation and
- <sup>29</sup> cytotoxicity of doxorubicin with different pharmaceutical <sup>30</sup>

formulations in human cancer cells. *Journal of Nanoscience and Nanotechnology*, 6, 3062–3069.

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- Silva, A. S. (2007). Nanotechnology approaches for drug and small molecule delivery across the blood brain barrier. *Surgical Neurology*, 67, 113–116.
- Stouch, T. R., & Gudmundsson, O. (2002). Progress in understanding the structure-activity relationship pf P-glycoprotein. *Advanced Drug Delivery Reviews*, 54, 315–328.
- Tsunoda, I., Kuang, L. Q., Tolley, N. D., Whitton, J. L., & Fujinami, R. S. (1998). Enhancement of experimental allergic encephalomyelitis (EAE) by DNA immunization with myelin proteolipid protein (PLP) plasmid DNA. *Journal of Neuropathology & Experimental Neurology*, 57, 758–767.
- Ugazio, E., Cavalli, R., & Gasco, M. R. (2002). Incorporation of cyclosporin A in solid lipid nanoparticles in solid lipid nanoparticles. *International Journal of Phamaceutics*, 241, 341–344.
- van der Geest, R., van Laar, T., Gubbens-Stibbe, J. M., Boddé, H. E., & Danhof, M. (1997). Iontophoretic delivery of R- apomorphine — II: An in vivo study in patients with Parkinson's disease. *Pharmaceutical Research*, 14, 1804–1810.
- Yang, S. C., Zhu, J. B., Lu, Y., & Liang, C. Z. (1999). Body distribution of camptothecin solid lipid nanoparticles after oral administration. *Pharmaceutical Research*, 751–757.
- Zara, G. P., Cavalli, R., Fundaro, A., Bargoni, A., Caputo, O., & Gasco, M. R. (1999). Pharmacokinetics of doxorubicin incorporated in solid lipid nanospheres (SLN). *Pharmaceutical Research*, 40, 281–286.
- Zara, G. P., Cavalli, R., Bargoni, A., Fundaro, A., Vighitto, D., & Gasco, M. R. (2002). Intravenous administration to rabbits of non-stealth and stealth doxorubicin loaded solid lipid nanoparticles at increasing concentration of stealth agent: Pharmacokinetics and distribution of doxorubicin in brain and in other tissues. *Journal of Drug and Targeting*, 10, 327–335.

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