Modified Lipid Nanocapsules for Targeted Tanshinone IIA Delivery in Liver Fibrosis

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Introduction: Liver fibrosis represents a serious global disease with no approved treatment. Tanshinone IIA (TSIIA) is a phytomedicine with documented activity in treating many hepatic disorders. TSIIA has been reported to have potent anti-inflammatory and antioxidant properties. It can also induce apoptosis for activated hepatic stellate cells, and is thereby considered as a promising herbal remedy for treating fibrotic liver. However, its poor aqueous solubility, short half-life, exposure to the first-pass effect, and low concentration reaching targeted cells constitute the major barriers hindering its effective therapeutic potential. Therefore, this work aimed at enhancing TSIIA systemic bioavailability together with achieving active targeting potential to fibrotic liver via its incorporation into novel modified lipid nanocapsules (LNCs).

Methods: Blank and TSIIA-loaded LNCs modified with either hyalurionate sodium or phosphatidyl serine were successfully prepared, optimized, and characterized both in vitro and in vivo.

Results: The developed LNCs showed good colloidal properties (size ≤100 nm and PDI ≤0.2), high drug-entrapment efficiency (>97%) with sustained-release profile for 24 hours, high storage stability up to 6 months, and good in vitro serum stability. After a single intraperitoneal injection, the administered LNCs exhibited a 2.4-fold significant increase in AUC0–∞ compared with the TSIIA suspension (p≤0.01). Biodistribution-study results proved the liver-targeting ability of the prepared modified LNCs, with a significant ~1.5-fold increase in hepatic accumulation compared with the unmodified formulation (p≤0.05). Moreover, the modified formulations had an improved antifibrotic effect compared with both unmodified LNCs and TSIIA suspension, as evidenced by the results of biochemical and histopathological evaluation.

Conclusion: The modified TSIIA-LNCs could be regarded as promising novel targeted nanomedicines for effective management of liver fibrosis.

Keywords: phytomedicine, hyaluronic acid, phosphatidyl serine, antifibrotic activity

Introduction
Liver fibrosis represents a serious health problem affecting millions of people worldwide. It is considered an ineffective reversible wound-healing response to variable acute and/or chronic stimuli that might progress to complicated cirrhosis.1 Following chronic liver injury, various immune-system cells, responsible for induction of inflammation, infiltrate hepatocytes. Some injured hepatocytes undergo apoptosis as well. The latter release cytokines and reactive oxygen species, which in turn activate resident macrophages (Kupffer cells [KCs]) to further release many proinflammatory cytokines. These changes result in a dramatic phenotypic transformation to hepatic stellate cells (HSCs), called activation, into myofibroblast-like cells. Activated HSCs undergo transformation into proliferative, fibrogenic, and...
contractile cells with overproduction of extracellular matrix (ECM) proteins (mainly collagens type I and III). Moreover, the contractile feature of activated HSCs plays a pivotal role in promoting portal hypertension, which facilitates further progression to cirrhosis.\(^2\)

Currently, no standard treatment exists for liver fibrosis. Elimination of the etiological factor is considered the most effective therapy. No antifibrotic drugs have been approved for clinical use, due to absence of tissue selectivity and low concentrations reaching target cells. It is worth noting that many antifibrotic drugs are currently undergoing clinical trials.\(^3\) Numerous herbal medicines have been reported to have great potential in reversal of liver fibrosis, and among them tanshinone IIA (TSIIA) has superior documented efficacy.\(^4\)–\(^7\) TSIIA has many reported mechanisms of action to resolve liver fibrosis, such as being a potent anti-inflammatory agent.\(^7\)–\(^8\) It has been reported to decrease the expression of TGF\(\beta.\)\(^9\) Moreover, it is a strong antioxidant that attenuates hepatic oxidative stress, preventing damage of hepatic cells resulting from lipid peroxidation,\(^7\) and has been found to induce apoptosis of activated rat HSCs without damaging normal hepatocytes.\(^10\) Despite its huge therapeutic benefits, TSIIA has low systemic bioavailability, due to its poor aqueous solubility, short half-life, and exposure to first-pass metabolism.\(^11\) It is highly lipophilic and extremely insoluble in water, with intrinsic aqueous solubility of 2 ng/mL and log P of 5.8.\(^12\) Among diverse strategies, nanotechnology presents a tailored design that can face the enormous TSIIA-delivery challenges.

Lipid nanocapsules (LNCs) have been widely addressed as an interesting type of lipid-delivery systems that can efficiently entrap many problematic drugs.\(^13\)–\(^17\) Surface modification of these nanocarriers is feasible, allowing protection, controlled release, longer circulation, and improved targeting. LNCs can be produced using a low-energy, solvent-free method and are of good biocompatibility. They can change both pharmacokinetics and biodistribution of loaded drugs after parenteral administration.\(^18\)–\(^20\) Recently, TSIIA-loaded LNCs were
successfully prepared by our research team and reported to greatly enhance its oral bioavailability.\textsuperscript{21}

Targeting various receptors expressed on the surface of activated HSCs and KCs is the mainstay of effective management of liver fibrosis. Accordingly, hyaluronic acid (HA)\textsuperscript{22,23} and vitamin A\textsuperscript{24,25} have been exploited to achieve active targeting to HSCs. Phosphatidyl serine (PS) was previously regarded as a targeting ligand to KCs.\textsuperscript{26,27} HA is a linear polysaccharide abundantly found in animal ECM, connective tissue, and organs. It is a biocompatible, biodegradable, nonimmunogenic, noninflammatory, non-toxic polymer. CD44, a receptor for HA-decorated delivery systems, is overexpressed in hepatic fibrosis. It has an important role in activated HSCs migration during liver injury. Activated HSCs also express elevated levels of CD44.\textsuperscript{22,23} PS is an anionic phospholipid component of cellular membranes. It is present in the inner phospholipid bilayer of the cell membrane. When cellular apoptosis occurs, PS is transferred to the cell surface, permitting recognition by macrophages through scavenger-receptor pathways and subsequent phagocytosis.\textsuperscript{27} Accordingly, it is considered a targeting moiety for hepatic macrophages (KCs).

The current study focused on surface modification of TSIIA-LNCs with either HA or PS, hoping to develop an actively targeted nanomedicine for liver-fibrosis therapy and maximize drug efficacy after parenteral administration. Full physicochemical characterization, hemocompatibility, and serum stability studies were performed. In vivo pharmacokinetic behavior of tailored targeted TSIIA-LNCs was studied. Also, their hepatic targeting potential was investigated via biodistribution and antifibrotic activity studies in a rat model with carbon tetrachloride (CCl\textsubscript{4})–induced liver fibrosis. As far as we know, this study is the first to implement decoration of LNC surfaces with HA and PS to improve their liver targetability.

\textbf{Methods}

\textbf{Preparation of LNCs}

Blank LNCs were prepared by phase inversion method with temperature cycling process\textsuperscript{28} using the same ratios and ingredients reported previously by our research team.\textsuperscript{21} Briefly, 38% w/w labrafac lipophile, 22.5% w/w Kolliphor HS15, 15% w/w Span 80, 22.5% w/w water, and 2% w/w NaCl were mixed under magnetic stirring (IKA Labortechnik, Germany). The mixture was exposed to three cycles of progressive heating and cooling between 55°C and 30°C at a rate of 4°C/min. An irreversible shock was induced with 2-fold dilution with deionized cold water (0°C–2°C) added to the formed O/W emulsion at the phase-inversion temperature. Finally, the LNC dispersion formed was subjected to slow magnetic stirring for 5 minutes, followed by filtration through a 0.45 μm Millipore syringe filter.\textsuperscript{21}

PS-LNCs was prepared by addition of PS (0.25%–0.5% w/v of the final dispersion) to other LNC ingredients at the beginning of the process. The same method of preparation of blank LNCs was adopted, except for raising the temperature in the first cycle to 70°C to ensure complete melting of PS. For preparation of HA-LNCs, a post-insertion technique with some modifications was implemented.\textsuperscript{29} First, positively charged LNCs were prepared using hexadecyltrimethylammonium bromide (CTAB; 0.1% w/v) as reported previously.\textsuperscript{15,16} Then, CTAB-LNCs (1:1 diluted with deionized water) were dropped slowly to different volumes of hyaluronate sodium aqueous solution (6 mg/mL, pH 6.5) under vigorous stirring (800 rpm) for 1 hour at 37°C. The final concentrations of HA in the prepared formulations were 0.55, 1, and 1.4 mg/mL. Effective hyaluronate coating was verified by size and ζ-potential measurements.

Drug-loaded formulations were prepared as the blank formulations using TSIIA (1 mg/mL of the final dispersion volume). TSIIA was dissolved in the mixture of oil and surfactants, followed by addition of other LNC ingredients (water and NaCl).\textsuperscript{21} Finally, all TSIIA-loaded LNC formulations (TSIIA-LNCs, TSIIA-PS-LNCs, and TSIIA-HA-LNCs) were stored away from light at 4°C.

\textbf{Physicochemical Characterization}

The LNCs were subjected to full in vitro characterization of their colloidal properties\textsuperscript{17,21} (Zetasizer, Malvern Instruments, UK), entrapment efficiency,\textsuperscript{21} morphological examination using transmission electron microscopy (TEM; JEM-100 CX, Jeol, Japan)\textsuperscript{16,17} and in vitro drug-release study using the dialysis-bag method.\textsuperscript{21} These methods are mentioned in detail in the \textit{Supplementary Materials}.

\textbf{Hemocompatibility (Hemolysis Test)}

The hemocompatibility of blank and TSIIA-loaded LNCs was assessed according to a method described previously with slight modifications.\textsuperscript{16} In brief, red blood cells (RBCs) were isolated from fresh whole rabbit blood via centrifugation at 2,000 rpm for 15 minutes. The different formulations were incubated with equal volumes of RBCs.
suspension in saline (2% v/v) at 37°C for 1 hour with shaking (200 rpm). Samples were then centrifuged at 3,000 rpm for 5 minutes and hemoglobin content was spectrophotometrically quantified at $\lambda_{\text{max}} = 540$ nm in the supernatant versus negative control (equal volume of RBCs and saline). Another aliquot of 2 mL RBC suspension was treated with a few drops of 1% w/v Triton X-100 to induce 100% hemolysis and used as positive control. Hemolysis was calculated:

$$\% \text{ Hemolysis} = \frac{\text{Absorbance of the sample}}{\text{Absorbance of the negative control}} \times 100$$

$$\frac{\text{Absorbance of the positive control}}{\text{Absorbance of the negative control}} \times 100$$

(Equation 1)

It is worth mentioning that the initial concentrations of both TSIIA and LNCs used in the hemolysis testing were 1 mg/mL and 387.5 mg/mL, respectively.

Serum Stability
To assess the physical stability of LNCs in biologically relevant media, serum stability of different TSIIA-loaded LNC formulations was determined as reported previously with some modifications. Briefly, equal volumes of both tested formulations and FBS (10% w/v in saline) were mixed and then incubated at 37°C under continuous stirring (200 rpm). Samples were withdrawn at different time points (0, 1, 2, 4, and 6 hours), diluted with filtered deionized water (1:60 v/v), and assessed for both particle size and PDI.

Storage Stability
Stability of lyophilized TSIIA-LNC formulations (TSIIA-PS-LNCs and TSIIA-HA-LNCs) containing 2% w/v trehalose as a cryoprotectant kept at 4°C and protected from light was assessed. Lyophilization was performed using a benchtop lyophilizer (Cryodos-50, Telstar, Spain). Samples were evaluated for particle size, $\zeta$-potential, PDI, and EE of TSIIA for 6 months after being reconstituted with deionized water.

In Vivo Studies
In vivo studies were performed using adult male Wistar rats (180–220 g, 8 weeks old). All in vivo studies were conducted in accordance with the guidelines approved by the Institutional Animal Care and Ethics Committee of the University of Alexandria (approval ALEXU-IACUC-42).

Pharmacokinetics
A total of 28 rats were randomly assigned to four groups (seven rats each). The study groups received a single 5 mg/kg dose of TSIIA as TSIIA suspension (in PEG 400:saline 1:9) or LNC formulations (TSIIA-LNCs, TSIIA-PS-LNCs, and TSIIA-HA-LNCs) via intraperitoneal (IP) injection. Prior to administration, all formulations were subjected to filtration sterilization using 0.22 µm Millipore syringe filters. After IP injection, blood samples (1 mL) were collected via the orbital plexus under ether anesthesia at different time points (0, 0.25, 0.5, 1, 2, 5, 7, 24, 48, and 72 hours) in tubes containing EDTA. Plasma was then obtained and TSIIA quantified in isolated samples using liquid LC-MS/MS as reported previously by our group. Plasma concentration–time data were analyzed using a noncompartmental pharmacokinetic model with the help of a Microsoft Excel 2016 pharmacokinetic solver add-in. Key pharmacokinetic parameters for all treated groups were determined: peak plasma concentration ($C_{\text{max}}$), time to reach $C_{\text{max}}$ ($T_{\text{max}}$), total area under the curve (AUC$_{0-\infty}$; calculated using the linear trapezoidal method), elimination half-life ($t_{1/2}$), and mean residence time (MRT).

In Vivo Hepatic Antifibrotic Activity
Liver fibrosis was induced in rats by IP injection of 1 mL/kg CCl$_4$ (1:1 diluted with mineral oil) twice a week for 4 weeks. The study groups were negative-control group (healthy rats injected with saline) and mineral oil–treated group receiving mineral oil (0.5 mL/kg) twice weekly for 4 weeks via IP injection to assess its hepatotoxicity. Rats with induced liver fibrosis were randomly divided into the following groups: positive control (untreated), TSIIA-LNCs, TSIIA-PS-LNCs, TSIIA-HA-LNCs, and TSIIA suspension. The last four treated groups received an equivalent TSIIA dose of 2.5 mg/kg via IP injection three times weekly for 3 weeks. It is important to highlight that all formulations were sterilized through filtration using 0.22 µm Millipore syringe filters before administration.

Animals were observed throughout the study period for any clinical signs of toxicity, their weights recorded weekly, and changes in body weight calculated. At the end of the treatment period, rats were anesthetized using diethyl ether and blood samples were collected from the retro-orbital venous plexus. The recovered serum samples were used for determination of the biochemical markers for hepatic
function and certain proinflammatory cytokines. Animals were then sacrificed and dissected, then livers were isolated, washed with ice-cold saline, and weighed to calculate liver weight relative to body weight. A small part of each liver was fixed in 10% v/v formalin for histopathological examination and another part was employed for the determination of profibrotic biochemical markers.

**Quantitative Assessment of Hepatic Function**
Quantitative colorimetric determination of serum ALT, AST, ALP and albumin was done using suitable assay kits following the instructions. 33–35

**Quantitative Assessment of Hepatic Profibrotic Markers**
For evaluation of both MMP2 and TGFβ, small portions of the isolated livers were cut, weighed, and homogenized in ice-cold PBS (pH 7.4) to prepare 10% w/v homogenates. Liver homogenates were then divided into aliquots for determination of MMP2 and TGFβ by rat MMP2 and TGFβ ELISA kits. 36,37 For hydroxyproline (HP) assay, equal volumes of tissue homogenate (10% w/v in distilled water) and concentrated HCl (~12 N) were mixed, hydrolyzed at 120°C for 3 hours, and finally filtered. HP content was measured using an assay kit according to the manufacturer’s instructions. 38

**Quantitative Assessment of Proinflammatory Cytokines**
Quantification of both TNFα and IL6 levels was performed in serum following the instructions of rat immunoassay kits. 39,40

**Histopathological Evaluation of Liver Fibrosis**
Liver specimens of rats were fixed in 10% v/v formalin solution, dehydrated, and embedded in paraffin wax. Sections of 5 µm thickness were then cut and subjected to both H&E and Masson’s trichrome staining. Each stained section was then examined under a light microscope equipped with a digital camera (Olympus America Inc., USA). The amount of collagen fibers (green stain) in Masson’s trichrome-stained sections was semiquantitatively appraised using ImageJ software (version 1.45s). H&E-stained sections were scored according to Ishak score for the hepatic activity index (HAI) for scoring of necroinflammatory activity in chronic hepatitis. 41,42

**In Vivo Tissue Biodistribution in Fibrotic Rat Model**
In vivo biodistribution and liver-targeting potential of TSIIA-PS-LNCs and TSIIA-HA-LNCs were evaluated by laser-scanning confocal microscopy (Leica TSC SPEII, DMI8 automated, Germany). To allow the in vivo tracing of the prepared LNCs, 1,1-dioctadecyl-3,3,3,3-tetramethylindocarbocyanine perchlorate (DiI; 0.1 mg/mL of the final dispersion volume) was incorporated into the LNC formulations using the method described by Morille et al with slight modifications (for more details see Supplementary Materials). The colloidal properties of the fluorescently labeled LNC dispersions were determined.

For biodistribution study, the groups received a single dose equivalent to 1 mg/kg DiI in the form of free DiI solution (in acetone:water: 1:9), DiI-LNCs, DiI-PS-LNCs, and DiI-HA-LNCs via IP injection. Rats were then sacrificed at 1 hour after administration and organs (liver, kidney, and spleen) harvested and fixed in 10% v/v formalin. Specimens of the tested organs were dehydrated and embedded in paraffin wax before being sectioned into 5 µm–thick sections, mounted on a glass slide, and fluorescently observed using confocal microscopy at excitation and emission wavelengths of 545 and 550–600 nm, respectively. Five sections of each organ per animal were examined, and a minimum of three fields were imaged per section. The amount of DiI distributed in each tissue sample was determined based on the fluorescence-intensity measurement. 44 In brief, mean corrected total fluorescence was determined by calculating the fluorescence intensity across identical areas of the examined tissue sections and subtracting the background autofluorescence, using ImageJ. 44

**Statistical Analysis**
Experiments were performed in triplicate and data are expressed as means ± SD. Statistical analysis was carried out using one-way ANOVA), followed by either Tukey or Duncan post hoc analysis (SPSS 21) for pairwise comparisons. The level of significance was set at p≤0.05.

**Results and Discussion**
**Preparation of Modified LNCs**
In order to increase their therapeutic efficacy in the management of liver fibrosis, surfaces of LNCs were modified by different targeting ligands. This was achieved via inclusion...
of anionic PS in their shell structure or coating their surface with HA. It is worth mentioning that this work is the first attempt to document modification of LNC surfaces with both PS and HA. Negatively charged PS-LNCs were prepared to allow active targeting to liver macrophages using increasing PS concentrations of 0.25%–0.5% w/v. It was observed that addition of PS did not affect the formation of LNCs or their phase-inversion temperature. Regarding HA-LNCs, CTAB (0.1% w/v of the final dispersion) was used for imparting a positive charge to LNCs, a prerequisite for HA coating. This concentration of CTAB was selected based on its reportedly low toxicity and biocompatibility. The coating process of LNC surfaces was mainly based on electrostatic interactions between the cationic CTAB-LNCs and the anionic polysaccharide (HA). Similarly, Zheng et al utilized CTAB-HA nanostructured lipid carriers containing both doxorubicin and gold nanoparticles for targeting breast cancer.

The preparation of HA-LNCs was achieved by a simple titration procedure to avoid tedious and time-consuming washing steps required for the removal of excess HA. Optimization of the added concentration of HA is necessary for uniform coating of LNC surfaces, as excess HA may affect both size and homogeneity of the produced LNCs. Consequently, three concentrations of HA were tried (0.55, 1, and 1.4 mg/mL) to coat diluted CTAB-LNCs (1:1 with water). It was found that both dilution of LNCs and mild heating at 37°C for 1 hour were required for efficient HA coating. Both temperature and LNC concentration were considered important factors governing the post-insertion technique for coating of LNCs, as elevated temperature affects the “brush” conformation of the PEG chains of Kolliphor HS15 present on the surface of LNCs, allowing better penetration of the molecules inserted between the Kolliphor PEG chains. Incubation for 1 hour was found to be sufficient for achieving reversal of ζ-potential values for the coated LNCs. For TSIIA-LNC formulations, TSIIA concentration of 1 mg/mL from the total dispersion volume was selected based on our previous work. This concentration exhibited good stability for the liquid dispersions against drug precipitation during storage at 4°C for 2 weeks.

Physicochemical Characterization of LNCs

Preliminary Colloidal Properties for Optimization of LNC Formulations Modified with PS and HA

Concerning PS-LNCs, the effect of adding different PS concentrations on particle size, PDI, and ζ-potential of the formed LNCs is shown in Figure 1. Particle size and PDI were evaluated using dynamic light scattering. The results indicated that increasing PS concentration led to a decrease in particle size and an increase in PDI. Similarly, the ζ-potential values were found to decrease with increasing PS concentration. For HA-LNCs, the effect of adding different HA concentrations on particle size, PDI, and ζ-potential was also investigated. The results showed that increasing HA concentration led to an increase in particle size and a decrease in PDI. The ζ-potential values were found to increase with increasing HA concentration, indicating an increase in the net negative charge of the LNC surface.
remained unchanged when the PS concentration increased from 0.25% to 0.4% w/v (Figure 1A), suggesting incorporation of PS in the shell of the LNCs. Similar findings have been reported previously, where increasing PS concentration from 4 to 12 mol% did not cause any significant difference in the size of prepared nanostructured lipid carriers, confirming insertion into the phospholipid monolayer.26 However, upon increasing the PS concentration to 0.5% w/v, particle size and PDI markedly increased from 72.4±4.5 nm and 0.09±0.01 to 86.9±4.9 nm and 0.27±0.03, respectively. This corroborated what has previously been reported, where addition of PS to PVA-coated dextran nanoparticles resulted in significant increases in both size and PDI, suggesting its presence on the surface of nanoparticles as an efficient coat.47

Regarding ζ-potential data, Figure 1B shows that increasing PS concentration from 0 to 0.5% w/v resulted in an increase in corresponding negative ζ-potential from −15.8±3.1 to −50.2±1.6, respectively. This in turn further confirmed the efficient integration of PS into the shell of the LNCs. According to colloidal properties obtained for PS-LNCs, the optimum concentration (0.4% w/v) for PS was chosen as it was the highest concentration achieving a high negative ζ-potential value (−42.8±0.6 mV) without affecting particle size or PDI.

In respect of HA-LNCs, inclusion of CTAB (0.1% w/v) in LNCs had an insignificant effect (p>0.05) on particle size and PDI which changed from 69.35±1.95 nm and 0.098±0.003 for LNCs to 73.5±1.89 nm and 0.11±0.007 for CTAB-LNCs. This was in agreement with what was previously found using agents like CTAB or stearyl amine for CTAB-LNCs. This was in accordance with Tran et al15 who attributed the decrease in particle size of solid-lipid nanoparticles after increasing the PS concentration to 0.5% w/v, particle size and PDI, ζ-potential of HA-LNCs. It was found that addition of 0.55 mg/mL HA solution led to aggregate formation and consequently an increase in both particle size and PDI (Figure 1C). On the other hand, upon addition of both 1 and 1.4 mg/mL HA solution, particle size and PDI started to decrease, probably due to electrostatic interaction with the positively charged surface of the CTAB-LNCs. Similarly, Freag et al30 reported a decrease in particle size of a prepared nanosystem with increased concentration of HA solution. They attributed this finding to strong electrostatic interactions between chitosan-coated liquid crystalline nanoparticles and HA at high concentrations, resulting in stiffer and smaller particles.30 Figure 1D shows that increasing the concentration of HA solution from 0 to 1.4 mg/mL resulted in an increase in negative ζ-potential value from 9.3±0.5 mV to −39±0.7 mV, respectively. This suggests coverage of CTAB-LNC surfaces by increasing amounts of HA. Finally, a concentration of 1 mg/mL HA solution was selected in the current study for coating CTAB-LNCs, since it permitted formulation of HA-LNCs of acceptable colloidal properties. Using lower volumes of HA solution (compared to HA concentration of 1.4 mg/mL) resulted in avoidance of excessive dilution of drug in TSIIA-loaded HA-LNCs.

Colloidal Properties of Optimized LNC Formulations

Table 1 shows a summary of the colloidal properties of the optimized LNC formulations under study. All blank and drug loaded formulations exhibited favorable colloidal properties including being relatively small (69.35–100.8 nm) and having a narrow size distribution (PDI 0.096–0.252) and ζ-potential (−12.35 to −42.83 mV), confirming good colloidal stability. Concerning particle size, inclusion of PS in PS-LNCs caused no statistically significant difference in size compared with LNCs (p>0.05). In contrast, the size of HA-LNCs increased significantly compared with LNCs (p≤0.05). This finding was in accordance with Tran et al48 who attributed the increase in particle size of solid-lipid nanoparticles after hyaluronate addition to the successful coating of the

| Table 1 Summary of colloidal properties of optimized blank and TSIIA-LNC formulations |
|-----------------------------------------------|
| **Blank LNCs** | **TSIIA-LNCs** |
| Mean size (nm) | PDI | ζ-Potential (mV) | Mean size (nm) | PDI | ζ-Potential (mV) |
| LNCs | 69.35±1.95 | 0.098±0.003 | −12.35±0.95 | 70.99±1.2 | 0.096±0.005 | −13.5±1.2 |
| PS-LNCs (0.4% w/v) | 69.87±2.86 | 0.14±0.025 | −42.83±0.57 | 74.52±1.4 | 0.157±0.003 | −41.4±1.9 |
| HA-LNCs (1 mg/mL) | 100.8±3.61 | 0.23±0.042 | −30.7±0.47 | 98.23 | 0.252±0.02 | −30.4±0.1 |

Notes: Data presented as means ± SD of three measurements.
negatively charged high-molecular-weight HA onto the nanoparticle surfaces.

Regarding ζ-potential, the slight negative ζ-potential of blank LNCs (−12.3±0.95 mV) could be attributed to the presence of traces of oleic acid in Span 80 together with the formation of intramolecular hydrogen bonding in Kolliphor HS15 creating a negative surface charge. Modification of blank LNCs with PS (0.4% w/v) and HA-LNCs (1 mg/mL) resulted in a significant reduction in ζ-potential to −42.83 mV±0.57 and −30.7±0.47 mV, respectively. It is worth mentioning that, TSIIA loading generally had an insignificant (p>0.05) effect on the colloidal properties of the LNC formulations. This was consistent to what had been previously reported in literature for both miltefosine- and praziquantel-loaded LNCs and was attributed to the presence of the drug in the oily core of the LNCs and not just attached to the surface.16,17

**TSIIA Entrapment Efficiency**

Both modified LNC formulations (TSIIA-PS-LNCs and TSIIA-HA-LNCs) exhibited high TSIIA EE of 98.7% ±0.1% and 98.1%±0.6%, respectively. This could be mainly due to TSIIA lipophilicity and its solubility in LNC oily cores.17,49 In other words, modification of LNCs did not significantly (p>0.05) affect TSIIA EE, an observation previously stated for both miltefosine- and praziquantel-loaded LNCs.16,17

**Morphological Examination by TEM**

TEM was used to visualize the morphology of TSIIA-PS-LNCs and TSIIA-HA-LNCs, as presented in Figure 2A and B. All LNCs were spherical, with size ≤100 nm and narrow size distribution confirming results obtained by the Zetasizer. Their morphological characteristics, particularly in terms of size and shape, were generally similar to those reported in the literature.50,51

**In Vitro Drug Release**

Figure 2C shows that complete release of TSIIA (80.57% ±0.52%) from its suspension occurred after 8 hours, while, the tested LNC formulations showed sustained-release profiles with significantly less release after 24 hours than the TSIIA suspension (p≤0.05), indicating drug entrapment inside the LNCs. Similar profiles have been reported for the release of diverse drugs, such as praziquantel, itraconazole, and quercetin, from LNCs.17,51,52 The sustained-release profiles of lipophilic drugs from LNCs has been attributed to the barrier function of the surrounding aqueous release medium that impedes drug transport owing to the poor aqueous solubility of the drug.21 Additionally, TSIIA-HA-LNCs showed significant less TSIIA release after 24 hours than TSIIA-PS-LNCs (p≤0.01). This finding was in agreement with that previously mentioned for hyaluronate-coated rapamycin–liquid crystalline nanoparticles.30 The lower drug release could be attributed to the thickness of the adsorbed bulky hyaluronate layer acting as an additional diffusion barrier controlling drug release from the hyaluronate-coated nanocarrier.30

**Hemocompatibility (Hemolysis Test)**

Hemolysis is an important problem associated with the bio-compatibility of injectable nanosystems.53 Accordingly, the effect of different LNC formulations on the destruction of erythrocytes was analyzed using a previously reported colorimetric assay.16 It was revealed that the tested formulations showed permissible hemolysis of less than the reported maximum limit (<5%)54 compared to the positive control (100%), as depicted in Figure 3A, indicating their hemocompatibility. This may be attributed to the absence of electrostatic interaction between negatively charged tested LNCs, TSIIA-LNCs, TSIIA-PS-LNCs, and TSIIA-HA-LNCs and erythrocytes of similar negative charge.53 It was also noted (Figure 3A) that TSIIA-PS-LNCs showed the least hemolytic activity, with hemolysis of 0.01%±0.002%. This could be attributed to the fact that TSIIA-PS-LNCs had the highest negative ζ-potential (−42.83 mV±0.57), resulting in the lowest interaction with RBCs.

**Serum Stability**

Assessing serum stability is an important requirement for any injectable delivery system. Following mixing with serum, unstable nanoparticles are expected to adsorb many serum proteins on their surfaces, leading to aggregation.55 This can result in large particles and broad size distribution. Changes in both particle size and PDI of TSIIA-loaded LNCs up to 6 hours after incubation with 10% FBS were monitored and are presented in Figure 3B and C. The results demonstrated no significant change (p>0.05) in particle size or PDI of any of the formulations with time, suggesting their stability in serum. This observation confirmed the impact of negatively charged LNCs to induce repulsive forces against anionic serum–protein interactions, as previously reported in literature.56

**Storage Stability of TSIIA-Loaded LNC Formulations**

The storage-stability results of lyophilized TSIIA-PS-LNCs and TSIIA-HA-LNCs containing 2% w/v trehalose after 6 months at 4°C are presented in Table 2. It was
observed that neither colloidal properties nor EE had changed significantly \( (p>0.05) \) after 6 months storage in the lyophilized form. This in turn reflected the high storage stability of these formulations, as mentioned previously in literature.\(^{16,28}\)

**In Vivo Studies**

**Pharmacokinetics**

Figure 4 shows the mean plasma drug concentration–time curve and Table 3 the main calculated pharmacokinetic parameters \( (t_{1/2}, \ C_{\text{max}}, \ T_{\text{max}}, \ \text{AUC}_{0-t}, \ \text{AUC}_{0-\infty}, \ \text{and MRT}) \). After IP administration, there were highly significant increases in both measured \( C_{\text{max}} \) values (about 2.7-fold) and \( \text{AUC}_{0-\infty} \) (about 2.4-fold) for all tested LNC formulations compared to the TSIIA suspension \( (p\leq0.01) \), indicating a marked increase in the amount of TSIIA reaching the systemic circulation and enhanced systemic bioavailability. Furthermore, both \( t_{1/2} \) and MRT were significantly \( (p\leq0.01) \) prolonged by TSIIA encapsulation in different prepared LNCs, suggesting a long circulation time in vivo, as previously reported in literature.\(^{15,17}\) Statistical analysis of the pharmacokinetic parameters of the three tested TSIIA-loaded LNC formulations (Table 3) indicated that differences among them were not statistically significant \( (p>0.05) \). This implied that modification of TSIIA-LNCs with either PS or HA did not significantly increase TSIIA bioavailability \( (p>0.05) \). However, these additives
were expected to increase the hepatic antifibrotic activity of unmodified TSIIA-LNCs, as is discussed later, via an active targeting mechanism.

In conclusion, the increased AUC$_{0-\infty}$ of all tested formulations after IP administration compared with TSIIA suspension and hence the improved bioavailability might be attributed to the presence of the drug in a molecular state solubilized in the oily cores of LNCs and avoidance of possible hepatic first-pass metabolism, as after IP administration, the drug drains to the portal vein through the mesenteric vessels, then to the liver before reaching the systemic circulation.$^{57,58}$ Therefore, LNCs were postulated to protect the incorporated drug from hepatic metabolism, due to: 1) their ability to reach the liver in structurally intact form, as proven previously,$^{16,17}$ 2) their ability to deliver higher drug concentrations to the liver than the TSIIA suspension, and 3) the inhibitory effect of Kolliphor HS 15 on hepatic cytochrome P450 microsomal enzymes.$^{59}$

**Hepatic Antifibrotic Activity**

**Induction of Liver Fibrosis**
The current study was performed using a CCl$_4$-treated liver fibrosis model, which is one of the most frequently utilized toxin-based models for liver fibrosis induction.$^{24}$ CCl$_4$ is metabolized in hepatocytes to a toxic trichloromethyl radical that mediates many cytotoxic effects via interaction with nucleic acids, proteins, and lipids.

**Table 2** Storage stability study of lyophilized TSIIA-loaded LNC Formulations at 4°C using trehalose (2% w/v) as a cryoprotectant

|                  | TSIIA-PS-LNCs     | TSIIA-HA-LNCs     |
|------------------|-------------------|-------------------|
|                  | Time 0            | 3 Months          | 6 Months         | Time 0            | 3 Months          | 6 Months         |
| Particle size (nm) | 73.5±0.5          | 74±0.6            | 74.6±0.3          | 102.5±1.1         | 101.4±1.3         | 102.8±1          |
| PDI              | 0.15±0.004        | 0.16±0.003        | 0.17±0.02         | 0.26±0.006        | 0.25±0.004        | 0.27±0.008       |
| ζ-potential (mV)  | −43.7±0.65        | −44.1±0.53        | −44.7±0.86        | −30.1±0.6         | −31.6±0.8         | −32.4±1.3        |
| EE (%)           | 98.1±0.55         | 98.2±0.53         | 98.03±0.6         | 98.4±0.83         | 98.6±0.6          | 98.9±1.54        |

Notes: Data presented as means ± SD of three measurements and statistically analyzed using one-way ANOVA followed by Tukey’s test for pairwise comparisons. Significance was set at p≤0.05.
Consequently, fatty degeneration and steatosis are observed. In addition, lipid peroxidation is encountered and results in generalized hepatic damage characterized by inflammation and fibrosis, which can progress to cirrhosis. A preliminary study showed that IP administration of 1 mL/kg CCl$_4$ (1:1 diluted with mineral oil) twice weekly for 4 weeks resulted in significant hepatic impairment based on the Ishak scoring system for determination of HAI.

Upon physical observation, no signs of toxicity were detected throughout the treatment period (3 weeks). Animals' body weight was recorded weekly during treatment and changes in body weight calculated (Table S1). After sacrificing animals, livers were weighed, and % liver weight relative to body weight was determined (Table S1). It was revealed that the positive-control and TSIIA-suspension groups had significantly less change in body weight than the other groups ($p$≤0.05). This could be due to marked hepatic damage, which can affect animals’ food intake. On the other hand, rats that received different TSIIA-loaded LNC formulations showed a progressive increase in body weight throughout the treatment period, statistically similar to the negative-control and oil-treated group ($p$>0.05). This could reflect the improved nutrition of animals upon treatment with these TSIIA nanosystems.

Concerning the measured % liver weight relative to body weight (Table S1), it was found that it significantly increased ($p$≤0.05) for the positive-control group compared to the healthy negative-control rats due to marked CCl$_4$ toxicity in liver tissue, as previously reported in the literature. Similar results were observed for the TSIIA-suspension group, which showed no improvement in hepatic state. For the mineral oil–treated group, no significant difference ($p$>0.05) was found in % relative liver weight.

### Table 3 Calculated pharmacokinetic parameters of TSIIA following IP administration of tested LNC Formulations and TSIIA suspension (5 mg/kg) to healthy rats

|          | TSIIA suspension | TSIIA-LNCs | TSIIA-PS-LNCs | TSIIA-HA-LNCs |
|----------|------------------|------------|---------------|---------------|
| $t_{1/2}$ (h) | 8.6±0.9$^b$ | 11.4 $^{a/b}$±0.4 | 12 $^{a/b}$±1.9 | 13.4 $^{a/b}$±0.5 |
| $T_{max}$ (h) | 1±0 | 1±0 | 1±0 | 1±0 |
| $C_{max}$ (ng/mL) | 50.8 $^{a/b}$±5.7 | 142.8 $^{a/b}$±19.9 | 137.2 $^{a/b}$±18.4 | 133.8 $^{a/b}$±10.3 |
| AUC$_{0-72}$ (ng/mL*h) | 428.4 $^{a/b}$±21.1 | 1,027.2 $^{a/b}$±83.1 | 1,076.9 $^{a/b}$±73.7 | 958 $^{a/b}$±4.5 |
| AUC$_{0-\infty}$ (ng/mL*h) | 436.3 $^{a/b}$±23.9 | 1,045.3 $^{a/b}$±86.6 | 1,115.5 $^{a/b}$±56.4 | 976 $^{a/b}$±6.9 |
| MRT (h) | 10.3 $^{a/b}$±0.2 | 14.1 $^{a/b}$±0.7 | 16.9 $^{a/b}$±0.6 | 14.8 $^{a/b}$±0.6 |
| Relative bioavailability$^d$ | — | 2.4 | 2.5 | 2.2 |

**Notes:** The study was done on Wistar rats with seven animals in each group. Values expressed as means ± SEM. Data analyzed using one-way ANOVA followed by post hoc Tukey’s test for pairwise comparisons. Means of similar symbols are statistically insignificant: $a > b > c$ (p≤0.01). $^d$Relative bioavailability was calculated as a ratio of AUC$_{0-72}$ for all LNC formulations to TSIIA suspension.
compared to the negative-control rats, indicating the safety of the oily vehicle used for dilution of CCl₄. On the other hand, all tested TSIIA-LNC formulations brought about a reduction in % relative liver weight, with the smallest reduction for TSIIA-LNCs, which did not differ significantly when compared to the positive-control and free-TSIIA groups (p>0.05). The actively targeted TSIIA-PS-LNCs and TSIIA-HA-LNCs brought about significantly lower (p≤0.05) % relative liver weight compared to the positive-control and TSIIA-suspension groups, but liver weights did not reach a normal state, as indicated by a significant difference (p≤0.05) from negative-control rats.

In conclusion, TSIIA-LNCs, TSIIA-PS-LNCs, and TSIIA-HA-LNCs brought about an improvement in food consumption, as indicated by statistically significant (p≤0.05) difference in body-weight change when compared to the positive-control and TSIIA-suspension groups. However, concerning % relative liver weight, only the targeted formulations (TSIIA-PS-LNCs and TSIIA-HA-LNCs) brought about a significant (p≤0.05) reduction when compared with the positive-control and TSIIA-suspension groups, reflecting their superiority in treatment, as will be highlighted later.

Quantitative Assessment of Hepatic Function

Serum markers for treated animals were calculated at the end of the study to evaluate hepatic function, and results are shown in Table 4. It has been reported that ALT, AST, and ALP are normally distributed in healthy hepatocytes and that elevated circulating levels of these markers reflect cellular damage and impairment of hepatocellular function. Additionally, albumin levels were measured to assess hepatic biosynthetic function. Positive-control rats showed significant elevation (p≤0.05) in ALT, AST, and ALP as well as hypoalbuminemia when compared with negative-control rats, reflecting considerable hepatocellular damage and impairment in biosynthetic functions of the liver. Administration of the free-TSIIA suspension (2.5 mg/kg) resulted in a statistically insignificant (p>0.05) correction in ALT and albumin, while it caused a significant (p≤0.05) reduction in AST and ALP when compared with the positive-control group, reflecting mild improvement in hepatic state. Treatment with TSIIA-LNCs (2.5 mg/kg) resulted in a significant reduction (p≤0.05) in all tested parameters compared to positive control, but the difference was no significant (p≤0.05) when compared to the TSIIA-suspension group, indicating their moderate therapeutic effect. This could be due to the low dose of TSIIA-LNCs administered (2.5 mg/kg) together with low drug amounts reaching specific cells involved in fibrosis progression (HSCs and KCs). Similar findings have been observed previously, where a low dose of IV-administered TSIIA-loaded globin nanoparticles (2 mg/kg) three times weekly for 3 weeks achieved a similar reduction in tested markers for liver function compared to TSIIA solution in a thioacetamide-induced liver-fibrosis mouse model. However, the difference between the two aforementioned groups was significant at a higher dose (5 mg/kg).

On the other hand, both targeted TSIIA-PS-LNCs and TSIIA-HA-LNCs (2.5 mg/kg) gave rise to significant (p≤0.05) improvement in all measured values on the liver-function test when compared to positive controls, free TSIIA suspension, and untargeted TSIIA-LNCs. Additionally, values returned to normal, as no significant (p>0.05) difference was found between these groups and negative-control rats. Therefore, targeted TSIIA therapy

### Table 4 Effect of tested LNC formulations and TSIIA suspension on markers of liver function after IP administration (2.5 mg/kg) to CCl₄ fibrotic rat model three times weekly for 3 weeks

|                   | ALT (U/mL) | AST (U/mL) | ALP (U/L) | Albumin (g/L) |
|-------------------|------------|------------|-----------|---------------|
| Negative control  | 40 ±2.9    | 53.4 ±4.3  | 2.5 ±0.1  | 42.8 ±1.4     |
| Positive control  | 80.2 ±2    | 97.8 ±9    | 3.4 ±0.03 | 26.4 ±1.4     |
| Mineral oil       | 41.6 ±1.3  | 49.6 ±2.7  | 2.5 ±0.1  | 43.8 ±1.2     |
| TSIIA suspension  | 72.2 ±5.3  | 77.5 ±5.4  | 2.9 ±0.04 | 29.6 ±1.9     |
| TSIIA-LNCs        | 63.8 ±2.5  | 73.8 ±6.7  | 2.9 ±0.08 | 31.2 ±1.4     |
| TSIIA-PS-LNCs     | 45.4 ±1.8  | 61.2 ±4    | 2.5 ±0.04 | 41.2 ±1.2     |
| TSIIA-HA-LNCs     | 48.4 ±2    | 54.6 ±1.6  | 2.5 ±0.03 | 40.6 ±1.4     |

**Notes:** The study was conducted on male Wistar rats with seven animals in each group. Values expressed as means ± SEM. Data analyzed using one-way ANOVA followed by post hoc test (Duncan) for group comparisons. Means of similar symbols are statistically insignificant: a > b > c (p≤0.05).
might be a good choice to improve the antifibrotic therapeutic potential of the drug at the low dose given (2.5 mg/kg).

**Quantitative Assessment of Hepatic Profibrotic Markers**

Elevated hepatic HP is considered a crucial diagnostic marker for liver fibrosis. It is an amino acid present in the collagen constituent of ECM produced by activated HSCs. Additionally, MMP2, a gelatinase enzyme, has been reported to have a dual role in pathogenesis and recovery from liver fibrosis. During liver injury, MMP2 promotes fibrosis by increasing HSCs proliferation. During resolution after hepatic injury subsides, particularly in the initial stages of disease, MMP2 promotes HSCs apoptosis and is required for degradation of the accumulated collagen. Therefore, elevated MMP2 is encountered in liver fibrosis originating from chronic injuries. Elevated TGFβ is also considered a key mediator in liver fibrosis, allowing ECM production. TGFβ is liberated from hepatocytes after injury. After that, it stimulates HSCs activation and proliferation, which in turn produces more TGFβ. The latter stimulates the expression of many ECM proteins, including collagen, downregulates their degradation, and induces hepatocyte apoptosis.

HP, MMP2, and TGFβ were measured in all tested animals at the end of the study, and results are shown in Table 5. Healthy negative-control rats showed relatively small levels of HP, MMP2, and TGFβ. All these markers are normally present in the liver for maintenance of tissue homeostasis, as they affect many cellular activities, such as proliferation, gene expression, fibrogenesis, and inflammation. The mineral oil–treated group showed no significant difference in levels of tested markers compared to negative-control rats ($p>0.05$), further ensuring the safety of the oily vehicle used for CCl4 dilution. Positive-control rats showed significantly elevated levels of all markers (HP, MMP2, and TGFβ) when compared to negative-control group ($p<0.05$). This might indicate abnormal ECM deposition as a result of the imbalance between matrix production and degradation mediated by MMPs. Conversely, TSIIA suspension, unmodified TSIIA-LNCs, and modified TSIIA-LNCs significantly decreased HP, MMP2, and TGFβ to variable degrees compared to positive-control group ($TSIIA$ suspension $< TSIIA$-LNCs $< modified$ TSIIA-LNCs, $p<0.05$). The significant reduction in fibrotic markers exhibited by the TSIIA suspension compared with positive-control rats demonstrated the antifibrotic efficacy of the drug, as previously reported. Untargeted TSIIA-LNCs exhibited a significant difference in levels of tested markers compared with TSIIA suspension ($p<0.05$), except for TGFβ, reflecting the role of LNCs in enhancing the antifibrotic activity of TSIIA. Interestingly, both targeted LNC formulations (TSIIA-PS-LNCs and TSIIA-HA-LNCs) achieved significantly ($p<0.05$) higher reductions in the tested markers than the untargeted TSIIA-LNCs. These results further highlighted the importance of the modified LNC formulations in resolution of liver fibrosis via an active targeting mechanism. The current results were in line with El-Mezayen et al., where targeted imatinib-loaded liposomes had a higher antifibrotic effect than a free-drug solution or untargeted therapy, as indicated by a significant reduction ($p<0.05$) in all fibrosis markers (HP, MMP2, and TGFβ). They attributed their findings to the ability of the targeted therapy to target activated HSCs, which are the main site of synthesis of most fibrogenic mediators.

It is important to highlight that neither TSIIA suspension nor TSIIA-LNC formulations returned markers to normal, as indicated by significant ($p<0.05$) differences from negative-control group (Table 5). Perhaps this might be due the short treatment period (3 weeks only). The latter

| Table 5 Effect of tested LNC formulations and TSIIA suspension on liver profibrotic markers and proinflammatory cytokines after IP administration (2.5 mg/kg) three times weekly for 3 weeks to CCl4 fibrotic rat model |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
|                | HP (ng/g tissue) | MMP2 (ng/g tissue) | TGFβ (ng/g tissue) | TNFα (pg/mL) | IL6 (pg/mL) |
| **Negative control** | 180.2 ±1.8 | 112.4 ±3.1 | 98.2 ±3.3 | 30.2 ±1.4 | 98.2 ±1.4 |
| **Positive control** | 436.4 ±4.7 | 399.8 ±1.7 | 397.4 ±15.5 | 55.4 ±1.3 | 318.4 ±5.7 |
| **Mineral oil** | 178 ±2.7 | 115.8 ±2.1 | 99.8 ±5.7 | 30.0 ±0.7 | 100.8 ±3.7 |
| **TSIIA suspension** | 397.2 ±5 | 362.2 ±8.2 | 271.8 ±5.5 | 50.2 ±1.5 | 281.2 ±2.8 |
| **TSIIA-LNCs** | 350 ±8.8 | 311.6 ±5.4 | 275.2 ±1.5 | 46.0 ±0.7 | 247.0 ±6.0 |
| **TSIIA-PS-LNCs** | 254.4 ±6.5 | 201.2 ±8.9 | 176.2 ±4 | 33.4 ±1.6 | 143.2 ±2.2 |
| **TSIIA-HA-LNCs** | 238.4 ±8.4 | 207 ±8.0 | 177.4 ±2.8 | 33.0 ±1.8 | 139.6 ±1.1 |

**Notes:** The study was conducted on male Wistar rats with seven animals in each group. Values expressed as means ± SEM. Data analyzed using one-way ANOVA followed by post hoc test (Duncan) for group comparisons. Means of similar symbols are statistically insignificant: a > b > c > d > e ($p>0.05$).
was selected based on a pilot study that revealed increased mortality, due to the induced progressive liver damage, in positive control group if the study exceeded 3 weeks.

**Quantitative Assessment of Proinflammatory Cytokines**

Many cytokines have a significant role in inflammatory liver diseases promoting fibrosis, such as IL6 and TNFα.\(^7^0\) TNFα is a proinflammatory cytokine produced by various immune-system cells, such as macrophages. It can trigger numerous signaling pathways associated with inflammation, proliferation, and apoptosis. Additionally, it is implicated in the pathogenesis of chronic liver inflammation resulting in liver fibrosis.\(^7^1\) Similarly, IL6 is a cytokine with various biological activities related to immunoregulation, inflammation, and hematopoiesis. Following chronic inflammation, it promotes the synthesis of many acute-phase proteins by the liver, resulting in induction of fibrosis.\(^7^2\)

Therefore, the level of these proinflammatory cytokines in the sera isolated from all animals of the tested groups was determined, and results are shown in Table 5.

Positive-control rats showed significantly (\(p \leq 0.05\)) elevated levels of both tested inflammatory mediators (TNFα and IL6) when compared with negative-control group, consistent with prior research, where CCl\(_4\)-induced acute hepatic injury activated hepatic nonparenchymal cells and increased the production of both TNFα and IL6.\(^7^3\) Hepatic macrophages (KCs) produce TNFα in rapid response to tissue injury, which then upregulates the expression of IL6. The overexpression of IL6 inhibits hepatocyte growth and causes further liver injury.\(^7^3\) The mineral oil–treated group had similar levels of inflammatory cytokines to the negative-control group (\(p > 0.05\)), indicating no hepatotoxicity for the oily vehicle used for CCl\(_4\) dilution. In contrast, all groups that received TSIIA treatment — either free suspension or encapsulated into LNCs — exhibited a significant decrease in levels of TNFα and IL6 when compared to the positive-control group. This might point to an underlying antifibrotic mechanism of TSIIA, which is mediated via suppression of inflammation. A similar observation was stated previously in the literature, where IV-administered TSIIA solution caused a marked decrease in proinflammatory cytokines in a fibrotic rat model.\(^7^\)

As shown in Table 5, reductions in inflammatory markers for the treated groups were in the order: TSIIA suspension < TSIIA-LNCs < TSIIA-PS-LNCs = TSIIA-HA-LNCs. Additionally, the difference between TSIIA suspension and TSIIA-loaded LNC formulations was statistically significant (\(p \leq 0.05\)), indicating the crucial role of LNCs in enhancing the in vivo performance of the encapsulated drug. The difference between unmodified TSIIA-LNCs and the two targeted LNC formulations was also statistically significant (\(p < 0.05\)), confirming again the importance of the active targeting approach in suppression of inflammation induced in liver fibrosis. It is worth mentioning that after 3 weeks’ treatment, the targeted formulations had succeeded in returning TNFα back to normal levels, as indicated by the insignificant difference from the negative-control group (\(p > 0.05\)).

**Histopathological Evaluation of Liver Fibrosis**

The findings of the animal-observation study and biochemical tests were additionally verified via histopathological examination of liver tissue. Liver sections taken from animals were stained with H&E for evaluation of any histopathological changes, especially hepatocellular damage (Figure 5A–H). Stained liver sections were scored using Ishak HAI score, and results were presented in (Figure 5I). Regarding H & E stain, the parenchyma of all negative-control livers revealed classical histological structure (Figure 5A). Liver sections of positive-control rats showed loss of classical hepatic architecture with pseudolobulation. Hepatocytes showed variable degenerative changes, including centrilobular confluent necrosis, apoptotic hepatocytes, and mononuclear cell infiltration. Dilated sinusoids with increased KCs and macroversicular fatty degeneration were also observed (Figure 5B and C). Liver sections of mineral oil–treated rats revealed more or less normal structure in most samples (Figure 5D). Necrotic cells and periportal cellular infiltration were occasionally seen. These findings and the previous results of biochemical tests confirmed the safety of the oily vehicle used for dilution of CCl\(_4\). The TSIIA-suspension group (Figure 5E) showed moderate hepatic degenerative changes, indicating the slight amelioration of the diseased hepatic state caused by the drug, being a hepatoprotective agent.\(^7^4–7^6\) On the other hand, all TSIIA-loaded LNC groups presented improvement in the hepatic diseased state, but to variable degrees. In this respect, TSIIA-LNCs accomplished the lowest efficacy, as indicated by the moderate correction in the impaired hepatic state...
ANOVA followed by post hoc test (Duncan) for group comparisons. Means of similar symbols were statistically insignificant: a > b > c > d > e (for hepatic activity index for scoring of necroinflammatory activity in chronic hepatitis for the study groups. Data expressed as means ± SEM (n=3). Data were analyzed using one-way ANOVA followed by post hoc test (Duncan) for group comparisons. Means of similar symbols were statistically insignificant: a > b > c > d > e (p≤0.05).

(Figure 5F). In contrast, the administration of both targeted formulations resulted in preservation of normal hepatic architecture in most samples, with classical hepatocytes containing vesicular nuclei. Different signs of degeneration were rarely encountered (Figure 5G and H). The latter finding ensured the superior effect of the targeted TSIIA therapy in improving the impaired hepatic state compared to both the untargeted one and TSIIA suspension.

For more illustration, the degree of hepatic impairment was quantified using Ishak HAI score for necroinflammatory activity in chronic hepatitis, and results are recorded in Figure 5I. The negative-control group exhibited the lowest total HAI score (0.75±0.5), which was statistically insignificant (p>0.05) compared to the mineral-oil group (2 ±0). This again confirmed the safety of the vehicle used for CCl4 dilution. The positive-control group had the highest total HAI score (13.7±0.7), which was statistically significant (p≤0.05) when compared with all other study groups. Both TSIIA suspension and unmodified TSIIA-LNCs were statistically similar (p>0.05) in achieving significant reductions in HAI score compared with positive-control group (p≤0.05). Similarly, El-Mezayen et al24 reported similar antifibrotic efficacy for both untargeted imatinib-loaded liposomes and free-imatinib solution, indicated by the observed deposition of collagen fibers in examined liver sections. This was attributed to the inability of the untargeted therapy to achieve a high
drug concentration reaching cellular targets (HSCs). Interestingly, both targeted TSIIA-LNCs succeeded in causing a significant decrease in HAI score compared to the untargeted formulation and free-TSIIA suspension (p≤0.05). However, TSIIA-PS-LNCs had the best efficacy, as evidenced by the significantly lower HAI score than TSIIA-HA-LNCs (p≤0.05). This might be attributed to the relatively higher abundance of KCs targeted by PS (~30% of nonparenchymal liver cells) than HSCs targeted by HA (<10% of nonparenchymal liver cells). It is worth mentioning that neither targeted LNC formulation returned HAI score to normal state, as indicated by the statistically significant difference (p≤0.05) when compared to negative-control group. This could be due to the short treatment period (3 weeks), as previously mentioned.

To further assess the degree of collagen-fiber deposition, a marker for liver fibrosis, liver sections isolated from all study groups were stained with Masson’s trichrome, coloring the collagen fibers green (Figure 6A–H). The amount of collagen fibers in each stained section were semiquantitatively evaluated using ImageJ to facilitate comparison among study groups, and the results are shown in Figure 6I. The stained liver specimens showed collagen fibers distributed mostly around blood vessels and portal tracts in the negative-control group (Figure 6A). On the other hand, liver sections of the positive-control rats showed massive increases in collagen fibers extending between portal tracts forming pseudolobulation around all necrotic areas and on areas of macrovesicular fatty infiltration (Figure 6B and C). The mineral oil–treated group (Figure 6D) exhibited a slight increase in collagen-fiber deposition when compared with negative-control rats, further assuring the safety of this oily vehicle. Examination of the TSIIA-suspension group (Figure 6E) revealed marked increase in collagen deposition, extending mainly from one portal tract to another, starting pseudolobulation of hepatic tissue. Administration of TSIIA-LNCs (Figure 6F) slightly ameliorated the degenerative changes, with marked collagen deposition in portal tracts. Fortunately, administration of both TSIIA-PS-LNCs and TSIIA-HA-LNCs (Figure 6G and H) caused limited increase in collagen-fiber distribution in both groups, indicating superior antifibrotic efficacy.

**Figure 6** (A–H) Photomicrographs of liver sections of groups and (I) semiquantitative analysis of fibrous tissue (green color) using ImageJ software. (A) Negative-control group showed classical distribution of collagen fibers mainly around blood vessels, especially in portal tracts (Pt). (B and C) Positive-control group demonstrated a massive increase in collagen fibers extending between Pt forming pseudolobulation around all central veins (CVs). Arrows in C point to existing necrotic areas. (D) Mineral-oil group revealed a slight increase in collagen distribution around Pt. (E) TSIIA suspension–treated rats showed a marked increase in collagen deposition extending mainly from one Pt to the other, starting pseudolobulation (arrows). (F) TSIIA-LNC group presented a marked increase in collagen fibers extending around hepatic tissue. (G) TSIIA-PS-LNC group exhibited a mild increase in collagen fibers around Pt. (H) TSIIA-HA-LNC group demonstrated mild increase in collagen fibers around Pt and CVs and extending between the hepatocytes. Masson’s trichrome stain Mic magnification 100×. (I) Semiquantitative analysis of fibrous tissue (green) in study groups using ImageJ software. Data expressed as means ± SEM (n=7). Data were analyzed using one-way ANOVA followed by post hoc test (Duncan) for group comparisons. Means of similar symbols were statistically insignificant: a > b > c > d (p<0.05).
The semiquantification of collagen fibers in each image (Figure 6I) further confirmed the results obtained from histopathological scoring. In this context, the positive-control group had the highest mean color intensity, which was statistically significant ($p \leq 0.05$) compared to other study groups, reflecting the highest deposition of collagen fibers. The TSIIA-suspension group showed a significant reduction ($p \leq 0.05$) in the mean color intensity compared with positive-control rats, verifying its effect in decreasing fibrous tissue formation. Although treatment with TSIIA-LNCs achieved lower mean color intensity than the TSIIA suspension, the difference between the two groups was insignificant ($p > 0.05$). This could be attributed to the low dose of TSIIA used, as previously mentioned. On the other hand, both targeted LNC formulations achieved significantly ($p \leq 0.05$) less collagen-fibers deposition, as indicated by the lowest mean color intensity, when compared to both the TSIIA suspension and untargeted TSIIA-LNCs. Nevertheless, only TSIIA-PS-LNCs succeeded in returning hepatic tissue to a normal healthy state, as indicated by their statistical similarity ($p > 0.05$) to the negative-control group, most probably due to the higher abundance of KCs, the main target for TSIIA-PS-LNCs.

In conclusion, combined results of biochemical and histopathological assessment illustrated that hepatocellular damage, inflammation, and necrosis were slightly corrected in rats administered TSIIA suspension or untargeted TSIIA-LNCs, and sometimes TSIIA-LNCs had higher efficacy. However, targeted LNC formulations achieved the highest antifibrotic efficacy. The difference in antifibrotic activity between unmodified and modified TSIIA-LNC formulations might be explained by the difference in TSIIA doses delivered to target nonparenchymal hepatic cells involved in fibrosis progression (HSCs and KCs). The higher the drug concentration reaching the target, the higher the efficacy.

In Vivo Tissue Biodistribution in Fibrotic Rat Model

We then explored organ biodistribution following IP administration of fluorescently labeled untargeted blank LNCs (Dil-LNCs) versus targeted blank LNCs (Dil-PS-LNCs and Dil-HA-LNCs), as well as Dil solution as control. As a lipophilic indocarbocyanine dye, Dil was selected for labeling of the prepared LNCs because it exhibited good labeling stability, as previously confirmed. It is worth mentioning that addition of Dil to LNCs did not change their colloidal properties, indicating efficient incorporation in the oily core.

Figure 7A presents the photomicrographs of organ sections (liver, kidney, and spleen) at 1 hour following IP

![Figure 7A](https://doi.org/10.2147/IJN.S331690)

**Figure 7** In vivo tissue biodistribution in rats with induced liver fibrosis. (A) Confocal laser-scanning microscopy of tissue sections at 1 hour after IP administration of different fluorescently labeled formulations (1 mg/kg) to CCl$_4$ fibrotic rat model at excitation and emission wavelengths of 545 and 550–600 nm, respectively. Magnification 63×. (B) Calculated mean corrected total fluorescence intensity of tested formulations detected in various tissue organs after 1 hour from IP administration (1 mg/kg) to CCl$_4$ fibrotic rat model. Data expressed as mean ± SEM (n=7), analyzed using one-way ANOVA followed by post hoc test (Duncan) for group comparisons. Means of similar symbols were statistically insignificant: a > b > c ($p \leq 0.05$).
injection of the tested fluorescently labeled formulations. Mean fluorescence intensity was analyzed using ImageJ (Figure 7B). Results demonstrated that the free-dye solution exhibited a nonspecific organ distribution to both liver and spleen, as there was no significant difference in measured fluorescence intensity for these organs \( (p>0.05) \). Its distribution to the kidney was significantly lower than other organs \( (p<0.05) \) owing to its lipophilicity. For Dil-LNCs, Dil-PS-LNCs, and Dil-HA-LNCs, Figure 7B shows that fluorescence intensity was significantly higher \( (p<0.01) \) for the liver than other organs (spleen and kidney), indicating specific distribution pattern of LNCs. Additionally, free-Dil solution demonstrated the least hepatic accumulation compared to other formulations \( (p<0.05) \). This illustrated the role of prepared LNCs in increasing hepatic deposition of the encapsulated dye. A similar finding was reported by Hirsjarvi et al.\(^\text{26}\) where LNCs exhibited higher hepatic distribution than other organs 24 hours after IV administration to mice. Moreover, Lacoeuille et al.\(^\text{79}\) found that the organ biodistribution of paclitaxel-loaded LNCs was higher in the liver compared to other organs. Their results indicated that LNC clearance from the blood was mainly due to the liver. This could be due to the action of hepatic macrophages that captured the nanoparticles.\(^\text{79}\) In addition, the smallness of the prepared LNCs \(<100\) nm helped them to diffuse from blood to hepatocytes through fenestration of the hepatic sinusoidal endothelium.\(^\text{79}\)

Another finding (Figure 7B) was that there were significant 1.4- and 1.6-fold increases \( (p<0.05) \) in hepatic accumulation after IP administration of Dil-PS-LNCs and Dil-HA-LNCs, respectively, compared to untargeted Dil-LNCs. This strongly reflected the role of both PS and HA as suitable ligands for LNCs achieving active targeting to hepatic fibrosis. These results are in agreement with Wang et al.\(^\text{27}\) where curcumin-loaded PS-modified nanostructured lipid carriers displayed an enhanced liver-targeting ability compared to an unmodified formulation. This could be due to anionic PS actively targeting hepatic macrophages.\(^\text{27}\) The role of HA in actively targeting overexpressed CD44 receptors in liver fibrosis was demonstrated by Thomas et al.\(^\text{22}\) They prepared losartan-loaded HA-modified micelles and studied their biodistribution in both a normal and hepatic fibrotic mouse model. They found that accumulation of their prepared system in the liver was significantly higher for fibrotic mice than normal ones, due to the overexpression of CD44 receptors on activated HSCs, which mediated endocytosis of the HA-decorated micelles.\(^\text{22}\) To sum up, the results of tissue biodistribution study proved the active targeting achieved by both PS and HA-LNCs to fibrotic liver and further confirmed the enhanced antifibrotic effect of these formulations compared to the untargeted one.

### Conclusion

No antifibrotic drugs have been approved for clinical use, due to absence of tissue selectivity and low concentrations reaching the target cells. In the current work, two novel modified LNCs (Ps-LNCs and HA-LNCs) were developed for targeted delivery of TSIIA in liver-fibrosis therapy. Modified TSIIA-loaded LNCs were successfully prepared and subjected to full in vitro and in vivo characterization. The prepared nanosystems possessed good in vitro characteristics in terms of their colloidal properties, TSIIA EE, TSIIA release, hemocompatibility, serum stability, and storage stability. In vivo efficacy and biodistribution results obtained implied the liver-targeting potential of the modified TSIIA-loaded LNCs. Consequently, the targeted TSIIA-LNCs look to be a promising nanocarrier that could circumvent the problems encountered with injectable TSIIA delivery and pave the way for their clinical use in liver-fibrosis treatment.

### Data Sharing

The authors confirm that the findings obtained in this study are available within the article.

### Disclosure

The authors confirm that they have no conflicts of interest associated with this work.

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