



Research Article

Incorporation of lipid nanosystems containing omega-3 fatty acids and resveratrol in textile substrates for wound healing and anti-inflammatory applications

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Abstract

In the present work, lipid nanosystems containing omega-3 fatty acid (nanostructured lipid carriers, NLCs) or omega-3 fatty acid and resveratrol (liposomes) were developed to improve cotton textile substrates as dressings with anti-inflammatory properties for wound healing applications. Lipid nanosystems were incorporated into woven, non-woven and knitted cotton substrates by exhaustion and impregnation. Based on physical–chemical characterization of the textile substrates, the textile structure and type of lipid nanosystems dictated the adsorption efficiency. In the case of NLCs, the woven substrate functionalized by exhaustion had a higher omega-3 release being the most promising for wound dressing application. Whereas for liposomes, the most adequate textile was the cationized knitted fabric functionalized by impregnation, that showed a more prolonged release profile of resveratrol.

Keywords Textiles lipid nanosystems · Liposomes · Nanostructured lipid carriers (NLCs) · Wound healing · Anti-inflammatory effects

1 Introduction

Wound healing is a complex physiological process involving several phases, including inflammation, proliferation and regeneration. Textiles are still the most used material to assist this complex process. Nowadays, there is a demand for smart wound dressings that, besides basic mechanical protection, must improve the healing process by interacting with the wound through the release of bioactive molecules and/or by maintaining the favorable conditions for the re-establishment of skin integrity and homeostasis [1]. Omega-3 fatty acids are natural bioactive compounds with proven anti-inflammatory activity [2].

Despite the literature reported benefits of omega-3 fatty acids, they still have some issues regarding the hydrophobicity, high propensity for oxidation, unpleasant smell and taste and low bioavailability. Lipid nanosystems have been pointed out as a successful strategy to overcome some of those limitations [3, 4].

The use of bioactive compounds with anti-inflammatory action such as resveratrol (RSV) have also reported benefic effects in wound healing processes [5, 6]. One of RSV's limitations is its rapid metabolization by the human body [7] and also its propensity to suffer oxidation and produce the inactive cis-isomer. To overcome this drawback, the encapsulation of RSV in liposomes, that might

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provide protection against early degradation and metabolism, is a strategy that has been investigated [8, 9].

Based on the galenic (i.e. the smell of fish oil) and stability (i.e. oxidation of both omega-3 and RSV) issues, the main goal of this study was to functionalize textile substrates with lipid nanosystems containing omega-3 (NLCs) or omega-3 and RSV (liposomes) which are intended to be used in wound healing.

Liposomes and nanostructured lipid carriers (NLCs) were produced and used to functionalize cotton textile substrates, by exhaustion and impregnation processes. The materials were characterized based on their Fourier-transform infrared spectroscopy with attenuated total reflectance (FTIR-ATR) and differential scanning calorimeter (DSC) profiles, as well as air and water vapor permeability properties. The bioactive release profiles from lipid-based nanosystems were also studied.

To our knowledge, this is the first study regarding textile substrates functionalized with lipid nanosystems containing RSV and omega-3.

2 Materials and methods

2.1 Materials

Precirol® ATO 5 (glyceryl palmitostearate), was donated by Gattefossé (France). Omega-3, composed by 70% of eicosapentaenoic acid (EPA), 18% of α -linolenic acid (ALA) and 12% of docosahexaenoic acid (DHA), was acquired from MyProtein (Portugal). Tween® 80 was purchased from Acofarma® (Spain) and benzalkonium chloride from Merck Schudart (Germany). Hexane and resveratrol (RSV) with a purity > 99% were obtained from Sigma-Aldrich® (Portugal). 1,2-Distearoyl-*sn*-glycero-3-phosphocholine (DSPC) and 1,2-Dioleoyl-*sn*-glycero-3-phosphoethanolamine (DOPE) with a purity > 99% were acquired from NOF laboratories (Japan). Three textile substrates 100% cotton were used: woven (5 × 5 cm), knitted fabrics (5 × 5 and 25 × 40 cm) and non-woven (disks with a diameter of 5.7 cm).

All lipid colloidal dispersions were prepared using ultrapure water, obtained from a Milli-Q purification system (Millipore Corporation).

2.2 Production of lipid nanosystems

NLC dispersions were prepared by ultrasound technique, as reported by Silva et al. [10]. The lipid phase was composed by Precirol® ATO 5 (7 wt%) and omega-3 (3 wt%) as solid and liquid lipids, respectively. The aqueous phase contained Tween® 80 (2.5 wt%) and benzalkonium

chloride (0.5 wt%), that act as surfactants, and ultrapure water (87 wt%).

Liposomes containing DSPC (0.4 wt%), DOPE (0.2 wt%) and omega-3 (0.1 wt%) were prepared by the lipid film hydration method using ultrapure water (99.3 wt%) followed by extrusion [8, 9]. RSV (200 μ M final concentration, regarded as suitable for obtaining anti-inflammatory effect [10, 11]) was incorporated in the preformed liposomes by an incubation method as previously reported [8, 9]. For sake of clarity the lipid nanosystems will be simply referred as NLCs (designating NLCs containing omega-3 fatty acid) and Liposomes (designating liposomes containing omega-3 fatty acid and RSV).

2.3 Characterization of the lipid nanosystems

2.3.1 Physical–chemical characterization and stability in storage conditions

To avoid the multi-scattering effect of light, lipid nanosystems were diluted with purified water (1:10) and were characterized regarding their size and distribution (measured by the polydispersity index, PDI) by dynamic light scattering in a Zetasizer Nano ZS laser scattering device (Malvern Panalytical, Parabol, Portugal) at a backscattering angle of 173°. Zeta-potential (ZP) of the lipid nanosystems was evaluated by electrophoretic light scattering (ELS) in a Zetasizer Nano ZS (Malvern Panalytical, Parabol, Portugal).

To evaluate physical stability of the lipid nanosystems under storage conditions (at 4 °C) the abovementioned physical–chemical characterization was made during a time span of two months.

2.3.2 In vitro cellular evaluation of anti-inflammatory effect

The excess production of nitric oxide (NO) has been associated with acute and chronic inflammation and consequent tissue damage [12, 13]. Therefore, the anti-inflammatory properties of the lipid nanosystems was evaluated in vitro using the murine macrophage (RAW 264.7) cell line, according to the method previously described by Sobral et al. [14]. The results were expressed in percentages of inhibition of NO production in comparison with the negative control (100%).

2.4 Functionalization of textile substrates

Textile substrates (woven, non-woven and knitted cotton fabrics) were functionalized with NLCs and liposomes by exhaustion and/or impregnation. In the exhaustion functionalization process, textile substrates (5 × 5 cm) were immersed in 5 mL of NLCs aqueous dispersion or

liposomes. They were maintained for 24 h at room temperature (RT), under mechanical agitation (60 rpm). To dry, the textile substrates were kept at 25 °C for 24 h. All samples were weighted before and after functionalization.

In the impregnation functionalization process, knitted fabrics with 25 × 40 cm size were immersed in 100 mL of each formulation and then pass through the foulard (1 m min⁻¹ and 2 bar). Each impregnated fabric was wrapped in a plastic tube, covered with cling film and left at RT for 24 h. To dry, the textile substrates were unwrapped and left at RT for 24 h. All samples were weighted before and after functionalization.

2.5 Characterization of the textile substrates

2.5.1 Chemical characterization using FTIR-ATR

The functionalized textile substrates were characterized by FTIR-ATR, DSC and air and water vapor permeability. For the FTIR-ATR analysis, the spectra were recorded between 400 and 4000 cm⁻¹, with increments of 4 cm⁻¹ and 45 scans/min, in FTIR Shimadzu I Raffinity.

2.5.2 Thermodynamic characterization using DSC

DSC measurements were carried in liquid nitrogen atmosphere using DSC-822e instrument (Mettler Toledo). The calibration was made with indium as standard. Samples were weighed (2.5 ± 0.2 mg) and sealed in aluminum pans. Then, they were heated from 25 to 350 °C, at a scanning rate of 10 °C min⁻¹. Data were treated using LAB Mettler Star SW 8.1 software (Mettler-Toledo International Inc., Swiss).

2.5.3 Air and water vapor permeability

The air/water vapor permeability is a basic physical property of wound dressings that may influence the wound healing process [15, 16], and for that reason should be investigated. Textile substrates functionalized by impregnation were used in air and water vapor permeability tests. The substrates functionalized by this technique were the only ones with the minimum dimensions required to perform these tests. Air permeability tests were carried using FX3300 instrument (TEXTTEST Instruments) in accordance with BS 7209:1990, with test head 20 (20 cm²). Ten measurements were made, all in different areas of the samples, with a pressure of 200 Pa.

Water vapor permeability tests were carried using M261 instrument (SDL) in accordance with EN ISO 9237:1995.

2.5.4 Release of bioactive compounds from textile substrates

To assure that the bioactive compounds encapsulated in the lipid nanocarriers have a therapeutic effect, it is necessary that they are released from the tissues' scaffolds. Therefore, it is necessary to study the release profile of such compounds. The release medium was chosen to provide the necessary chemical affinity for bioactives, so that release is simulated. RSV has moderate lipophilic character [9] and its solubilization is favored in amphiphilic media. Hence, RSV will only be released from liposomes if it has chemical affinity to the surrounding medium. As such, RSV release from liposomes was evaluated in micellar media (pH = 5.5) simulating the hydrophilic content and pH values of skin [17], mimicking the actual environment surrounding the tissue scaffold upon application on skin. Contrastingly, omega-3 is a highly lipophilic bioactive, and its release from NLC would have only been possible to lipid enriched domains of skin by fusion of the lipid nanosystems in contact with skin surface. As this situation is not easy to simulate, we have opted to use a nonpolar release medium (*n*-hexane) which is one of the widely used lipid-like organic phase in partition coefficient studies [18]. Similarly, it has been reported the use of other nonpolar organic solvents to evaluate the in vitro release of fish oils from microcapsules [19]. The use of such solvents as release media does not mimic the in vivo release conditions but allows to provide a chemical environment that is similar to the nonpolar media encountered in high lipid content of skin where the omega-3 fatty acids will solubilize. Furthermore, the release into an organic media allows to demonstrate that the fish oils were successfully encapsulated.

The functionalized textile substrates samples (5 × 5 cm) were immersed in 10 mL of release medium. The systems were disposed in an orbital shaker incubator at 37 °C and 80 rpm. At predetermined timepoints, sample aliquots were withdrawn and replaced with the same volume of fresh medium to maintain the sink conditions. The amount of omega-3 or RSV released was measured by UV-Vis spectrophotometry using a UV-2501 PC spectrometer (SHIMADZU), at λ_{max} of 269 nm and 305 nm for assessment omega-3 and RSV, respectively. Results were presented as cumulative percentage of released compound (omega-3 or RSV) as a function of time.

3 Results and discussion

3.1 Characterization of the lipid nanosystems

3.1.1 Physical–chemical characterization and stability in storage conditions

Immediately after preparation (t_i) lipid nanosystems containing RSV and omega-3 presented adequate sizes (Table 1) for skin penetration through intercellular route of *stratum corneum*, since it has been established that liposomal vesicles of 50–500 nm are able to use this penetration route [20–22]. Polydispersity index (PDI) values provide an indication of the sizes' distribution within a nanosystem suspension. Accordingly, NLC can be considered as a fairly homogenous dispersion ($PDI < 0.3$) [23–25]. Contrastingly, liposomes present higher PDI values typical of heterogeneous size's distribution. Considering that it is aimed a skin topical application of the lipid nanosystems, the higher values of PDI are not so relevant and could result from the insertion of more fluid lipid components (such as omega-3 and DOPE) in a more rigid lipid matrix (composed of DSPC). This strategy originates less uniformed packed vesicles, that by being more deformable originate heterogeneous dispersions. This deformability can be however useful for cutaneous application favoring the penetration through the *stratum corneum* [26]. Concerning the nanosystems surface charge, evaluated in terms of zeta-potential values (ZP), the presence of the cationic surfactant (benzalkonium chloride) explained the positive charge of the developed NLC dispersions (positive ZP values in Table 1). Liposomes are negatively charged (negative ZP values in Table 1) due to the ionization of the carboxylic acid group of omega-3.

During a shelf storage period of 2 months (t_f), NLCs remained stable in terms of mean particle size and PDI that is maintained under 0.30. However, ZP values of NLC became less negative, which could predict loss of stability of the formulation as values of ZP lower than -30 mV or higher than $+30$ mV are usually required to assure colloidal stability by surface charge repulsion [27, 28]. Despite this fact, since the charge neutralization was not

followed by a great increase in the size of nanoparticles, it is expected that Tween® 80 added to NLC composition might have conferred stability to the formulation being able to prevent the aggregation of nanoparticles by steric mechanisms [29]. Liposomes that were more disordered and heterogeneous when first prepared (t_i) apparently suffered lipid packing and/or reordering during the storage. Indeed, after 2 months of storage, liposomes present lower sizes and PDI. Notwithstanding the PDI improvement, ZP values become less negative meaning that as a future prospect the formulation should be improved to include a stabilizing element which prevents liposomal aggregation (for example by coating systems with polyethylene glycol, PEG).

3.1.2 In vitro cellular evaluation of anti-inflammatory effect

The inflammatory response following tissue injury plays important roles both in normal and pathological healing [30]. However, recent studies also suggest that the inflammatory process should be contained as failure to do this, plays a role in the pathogenesis of non-healing wounds [31, 32]. Accordingly, the production of excessive amounts of nitric oxide (NO) contributes to inflammation and tissue injury [33]. Regarding this rational, we hypothesized that RSV and omega-3 fatty acids can ameliorate wound healing due to an anti-inflammatory effect by inhibition of NO production, an effect that has been reported in previous studies [34, 35]. NLCs presented a NO production inhibition of $64.14 \pm 2.08\%$ demonstrating an effect of approximate magnitude of phenylethanoids (NO production inhibition of 6.3–62.3%) that for the extent of NO production inhibition were reported to possess anti-inflammatory properties [36]. The NO production inhibition effect obtained with NLCs is also in accordance with the results obtained by Lima et al. [34]. These authors reported that high concentrations of EPA and DHA lead to NO production inhibition [34].

Liposomes presented an inferior NO production inhibition ($21.97 \pm 1.27\%$), which was however within the NO production inhibition effect achieved by phenylethanoids [36]. The smaller effect in this case must be also related

Table 1 Characterization of liposomes (loaded with RSV and omega-3) and NLCs (loaded with omega-3)

Formulation	Size (nm)		PDI		ZP (mV)	
	t_i	t_f	t_i	t_f	t_i	t_f
Liposomes	152 ± 17	113 ± 11	0.58 ± 0.10	0.39 ± 0.02	−26 ± 1	−19 ± 1
NLCs	129 ± 1	135 ± 9	0.19 ± 0.004	0.24 ± 0.007	+25 ± 1	+8 ± 1

All values represent the mean ± standard deviation ($n=3$). t_i represents characterization immediately after preparation and t_f represents characterization perform after storage at 4 °C for 2 months

with the smaller concentrations of omega-3 fatty acid in this formulation.

All in all, both formulations were effective reducing NO production, which can be regarded as a promising advantage in the wound healing process.

3.2 Functionalization of textile substrates

Figure 1 presents the mass gain for the substrates functionalized with NLCs and liposomes by exhaustion and impregnation techniques.

In the substrates functionalized with NLCs by exhaustion, the mass gain was significantly higher than the non-functionalized ones (control), which confirms the incorporation of NLC into the textiles tested (Fig. 1a). This incorporation is however due to the ability of the textile substrates to absorb and retain the nanosystems dispersions and it does not result from the affinity between dispersions-substrate since all textile substrates are composed exclusively of cotton fibers. Accordingly, the non-woven fabric was the substrate with the highest mass gain, due to its higher absorption capacity (97.35%) (Fig. 1a). No mass gain occurred in the knitted fabric functionalized with NLCs by impregnation, maybe because this method does not use a long incubation period that might be required for the retention of the nanosystems.

In all substrates functionalized with liposomes by exhaustion, there was no significant differences in the mass gain relatively to the control (Fig. 1a). Since liposomes possess a negatively charged surface (Table 1) two knitted textile substrates were tested: non-cationized and cationized (Fig. 1b), however only cationized knitted fabric impregnated with liposomes demonstrated a mass gain about 8% higher than that recorded in the control. The reasons why the cationized knitted fabric functionalized by exhaustion with liposomes did not present any mass gain are not clear at this point. One might hypothesize

that the electrostatic interaction between negatively charged liposomes and the positively charged fabric tissue may induce liposomal fusion and vesicle disaggregation that motivates the loss of vesicles adsorbed to the fabric, especially during the prolonged incubation period used in exhaustion technique. Liposomal fusion onto surfaces has been described on similar incubation conditions [37, 38]. When cationized knitted fabrics are functionalized by impregnation there is not such a prolonged incubation period which might justify the successful electrostatic adsorption of liposomes to the fabric.

Figure 2 presents, as an example, the FTIR-ATR spectra obtained from knitted textile substrates functionalized by exhaustion (a) and impregnation (b). Spectra obtained by exhaustion for other substrates (woven and non-woven) are very similar and are presented as supplementary data.

All the spectra present similar vibration modes with few notorious differences, which can be justified by the common chemical moieties encountered either in the lipidic constituents of the nanosystems [39–45] or in the bioactive omega-3 [42] or even in the textile substrate [43]. The presence of the bioactive RSV, normally detected by the presence of trans olefinic bands at 986 and 964 cm^{-1} [44], was also not possible to assign. This is probably due to two factors: (1) the low concentration of RSV in formulation and (2) the fact of RSV being encapsulated in the liposomes thus justifying the lack of intensity of its ATR-bands which strongly depend on the depth of penetration the evanescent light source. Moreover, in agreement to the current study, other reported of RSV-loaded liposomes, also did not present any differences in FTIR spectra with and without RSV [44].

Despite of the abovementioned resemblance of chemical groups, spectra of the textile substrates functionalized with NLCs (blue spectra), either by exhaustion (Fig. 2a) or impregnation (Fig. 2b), present differences in comparison with the respective controls (grey

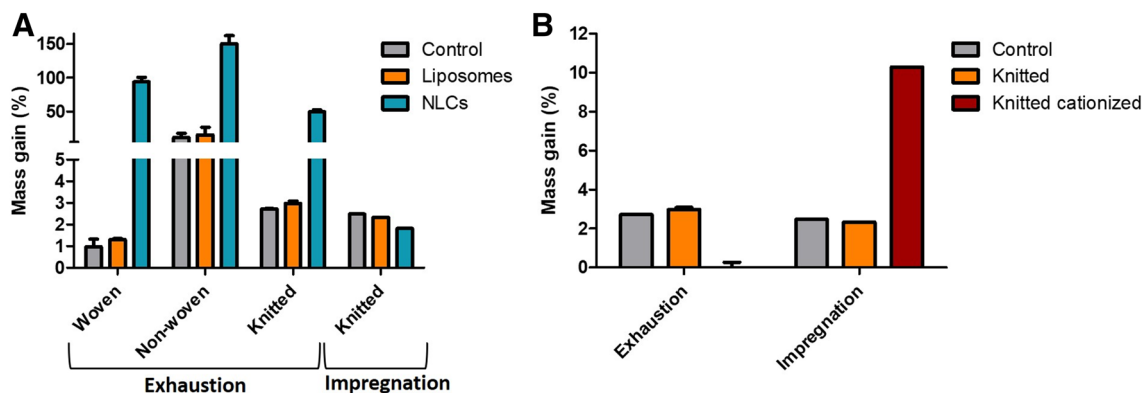


Fig. 1 a Mass gain and respective standard deviation of textile substrates functionalized with lipid nanosystems (NLCs and liposomes) using exhaustion and impregnation technique. b Mass gain of knitted substrates (cationized and non-cationized) functionalized with liposomes

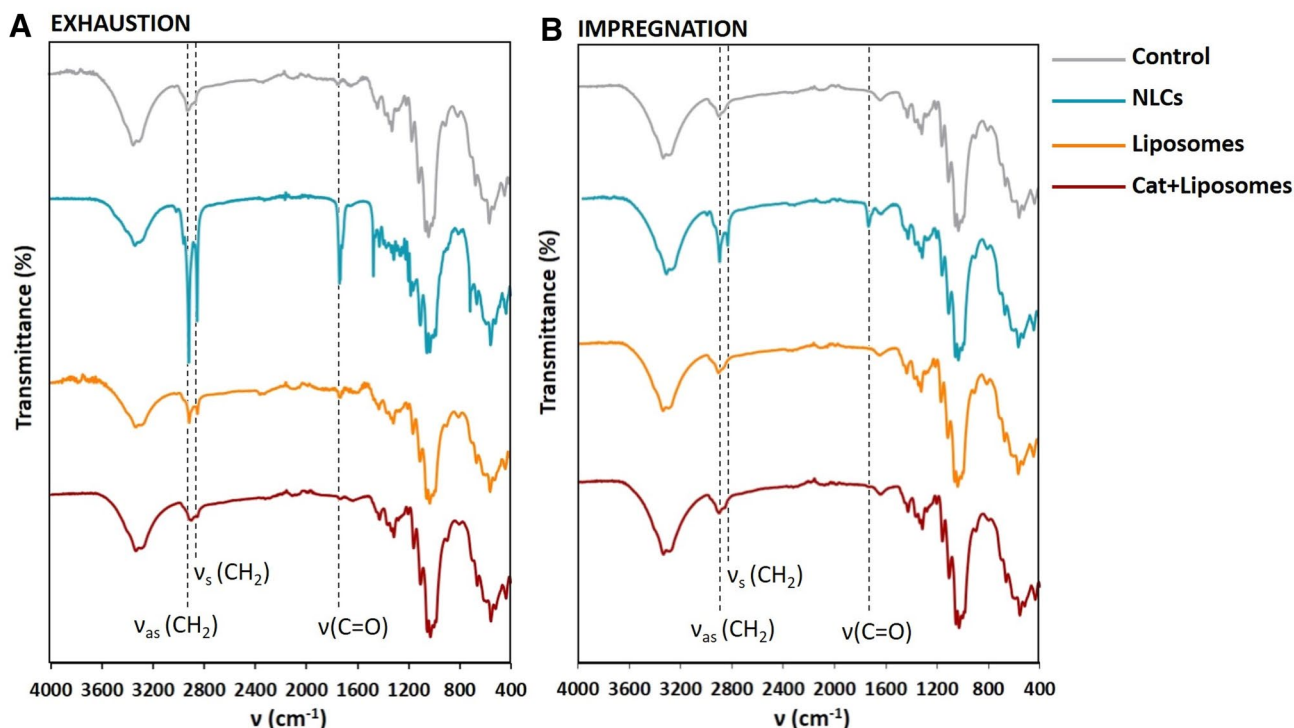


Fig. 2 FTIR-ATR transmittance spectra obtained from knitted fabrics functionalized by **a** exhaustion or **b** impregnation. Dashed lines indicate the vibration modes of CH₂ and C=O. Control are the bare knitted fabric spectra (grey colour); NLCs are the spectra of knitted

fabric functionalized with NLCs (blue colour); Liposomes are the spectra of knitted fabric functionalized with liposomes (orange colour) and Cat + liposomes are the spectra of cationized knitted fabric functionalized with liposomes (wine colour)

spectra). The first marked difference corresponds to the appearance of more intense bands in NLCs spectra (blue spectra) at 2912 and 2850 cm⁻¹ that can be respectively attributed to CH₂ asymmetric [$\nu_{as}(\text{CH}_2)$] and symmetric [$\nu_s(\text{CH}_2)$] stretching vibration modes. Although textile substrate possesses CH₂ groups in its composition, these groups appear in higher number in the long aliphatic tails of fatty acids constituents of lipids (e.g. Precirol[®] ATO 5) [39] or omega-3 [42], therefore originating more intense bands. The second difference noticed is a band at 1732 cm⁻¹ characteristic of C=O bonds that are also part of the ester region of both Precirol[®] ATO 5 and omega-3 [39, 42]. The vibration modes characteristic of NLCs spectra are evident in both methods of functionalization, but are less intense in the impregnation method (Fig. 2a) which maybe related with the smaller efficiency of this method to functionalize textiles with NLCs as corroborated by the lower mass gain of the textile substrate (Fig. 1a).

As explained above, the incorporation of liposomes in the textile substrates did not shown relevant differences in FTIR-ATR spectra either due to the low concentration of liposomes or by coincidence of bands in liposomes' spectra with bands in textiles' spectra (Fig. 2a—orange spectrum).

Figure 3 presents, as an example, the DSC thermograms obtained from knitted textile substrates functionalized by exhaustion (a) and impregnation (b). Thermograms obtained for other substrates (woven and non-woven) functionalized by exhaustion are very similar and are presented as supplementary data.

All thermograms present an endothermic transition between 53 and 60 °C. This transition is believed to be the result of water evaporation and the melting/decomposition of waxes and lipids present on the textile substrates [46]. Near the end of the thermograms, the beginning of a new endothermic transition, around 350 °C, can also be detected. This transition was also reported by Scacchetti [47] and is associated to the thermal decomposition process of cellulose.

In the thermograms of the knitted textile substrates functionalized with NLCs (blue thermogram) either by exhaustion or by impregnation, there are two endothermic transitions between 51 and 64 °C which correspond to the endothermic transition of the cotton that occurred in the control, and to the endothermic transition of the solid lipid used (Precirol[®] ATO5). The endothermic transition of the solid lipid is in agreement with the obtained by Avasthi et al. (63.8 °C) [48]. Wang et al. [49] demonstrated that, when Precirol[®] ATO5 is incorporated into NLC dispersions,

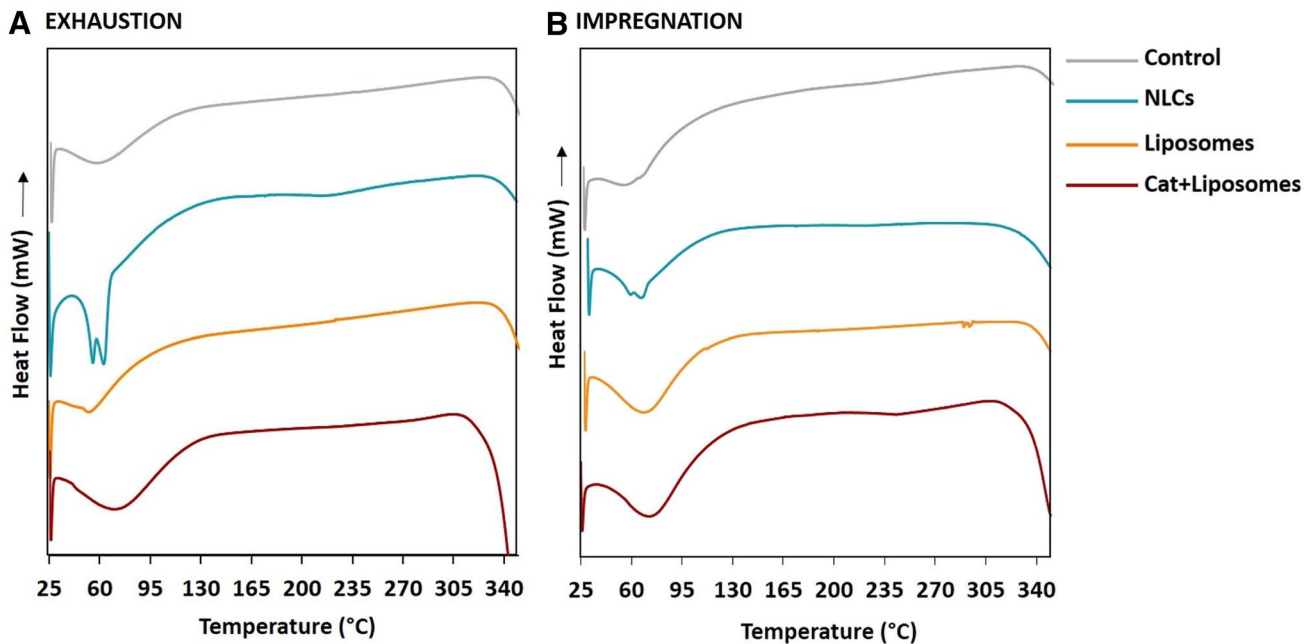


Fig. 3 DSC thermograms obtained from knitted fabrics functionalized by **a** exhaustion or **b** impregnation. Control are the bare knitted fabric spectra (grey colour); NLCs are the spectra of knitted fabric functionalized with NLCs (blue colour); Liposomes are the

spectra of knitted fabric functionalized with liposomes (orange colour) and Cat + liposomes are the spectra of cationized knitted fabric functionalized with liposomes (wine colour)

the transition appeared at slightly lower temperatures (57.8 °C), compared to the thermogram of the pure lipid.

No thermal transition related to the presence of omega-3 was observed, which was already expected since this compound is liquid at room temperature. Together with a reported study that has proved thermal stability of fish oils until 180 °C [50], this result allows us to deduce that omega-3 does not undergo significant changes in the temperature analysis, so it will be stable at the temperature at which NLC production occurs.

For the textile substrates treated with liposomes, no major differences were observed comparing to the controls. From all formulation constituents, those that have melting temperatures within 25–350 °C, are DSPC ($T_m = 55$ °C) and RSV ($T_m = 254$ °C). DSPC endothermic transition, may be covered by the textile transition referred above. On the other hand, the transition related to the melting of RSV is not detectable probably due to the low RSV concentration used.

Figure 4 represents the air and water vapor permeability of knitted fabric before and after impregnation technique.

In wound healing, air permeability is important because it prevents overheating, limiting bacterial proliferation and improving comfort [51]. The non-functionalized knitted fabric (control) possesses adequate air permeability (Fig. 4a grey bar) and the impregnation of the lipid nanosystems (Fig. 4a blue bar—NLCs and orange

bar—liposomes) did not alter the so called “breathability” of the tissue, which is important for wound healing proposes.

The impregnation of cationized knitted fabric with liposomes revealed a 13% increase in air permeability which is probably due to fiber degradation during the cationization process. The high standard-deviation associated to the air permeability measured in cationized knitted fabric (508 ± 49) confirms that the cationization process was not homogeneous and, therefore, the impregnation was not homogeneous too.

Dressings that produce a moist wound environment, thereby promoting cell migration, have long been recognized as desirable for wound healing purposes [52]. Therefore, to attain satisfactory conditions at the wound site, the functionalization of the textile substrates should not affect its water vapor permeability. In comparison with the control (Fig. 4b grey bar), none of the impregnated samples have shown a significant reduction of water vapor permeability. Therefore, in wound healing, this fabric will be able to prevent dehydration of the wound, but it might not be the best choice to wounds with a high amount of exudate.

Figure 5 presents omega-3 (a) and RSV (b) release profile from textile substrates.

In the case of woven and knitted fabrics functionalized by exhaustion technique, a burst of omega-3 release is observed in the first 4 and 2 h, respectively (Fig. 5a). In the

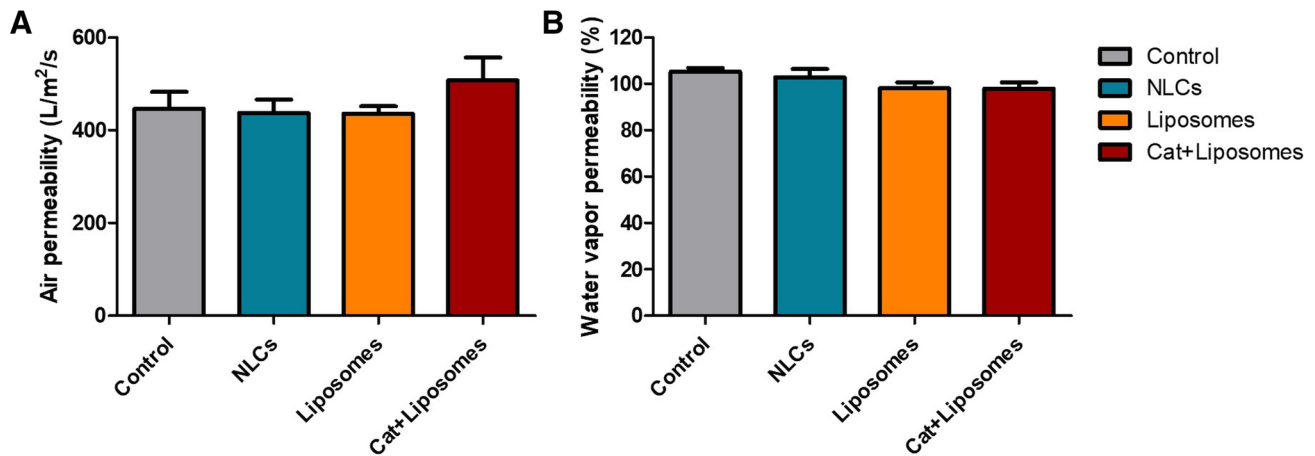
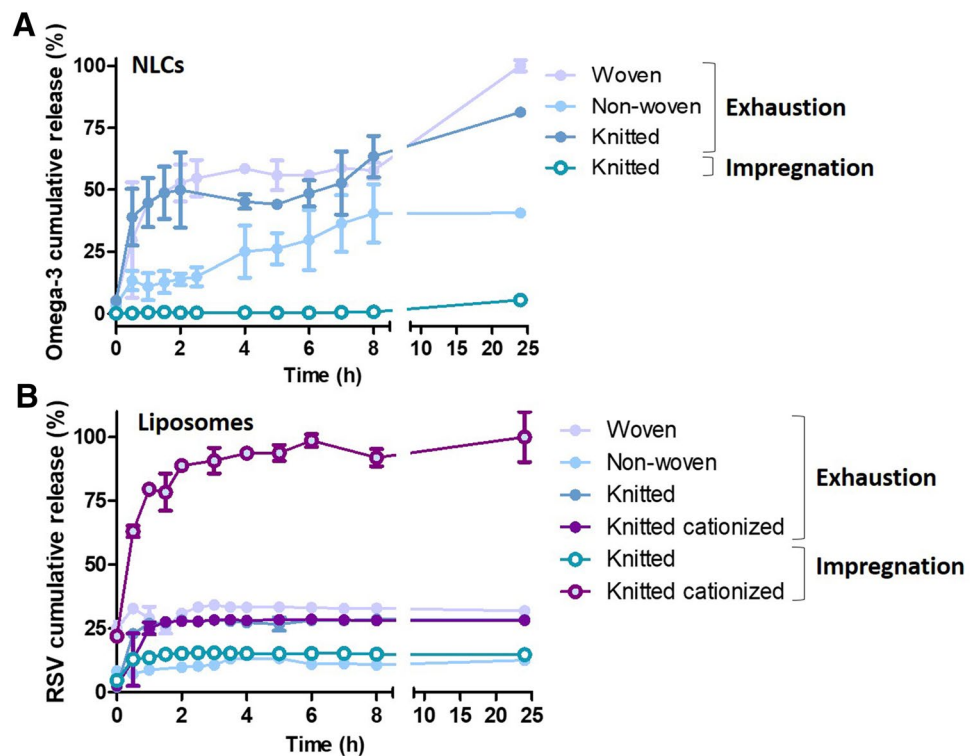


Fig. 4 Air permeability (a) and water vapor permeability (b) of knitted fabric functionalized by impregnation. Control are the bare knitted fabric spectra (grey colour); NLCs are the spectra of knitted fabric functionalized with NLCs (blue colour); Liposomes are the

spectra of knitted fabric functionalized with liposomes (orange colour) and Cat + liposomes are the spectra of cationized knitted fabric functionalized with liposomes (wine colour)

Fig. 5 Omega-3 (a) and RSV (b) cumulative release during a 24 h period



last 16 h of the test, omega-3 release still occurred so it is important to extend this experiment to understand when the maximum release is reached.

With the non-woven fabric, omega-3 suffered a small burst release during the first 8 h. After 24 h, the amount of omega-3 released is similar to the observed after the first 8 h (Fig. 5a). Therefore, it can be concluded that 8 h is

enough time to achieve the maximum release from non-woven fabric.

Despite being the one with higher mass gain (Fig. 1a), non-woven fabric had a lower amount of omega-3 released (Fig. 5a), comparing with woven and knitted fabrics functionalized by exhaustion. This can be explained by the macrostructure of this fabric which makes the contact

with the release medium more difficult when compared with the flat structures (woven and knitted fabrics).

The impregnated knitted fabric presented very low amounts of omega-3 release. This proves the weak functionalization using this technique, confirming the results obtained by mass gain (Fig. 1a) and FTIR-ATR spectra (Fig. 2b).

In summary, all textiles functionalized with NLCs by exhaustion presented a fast omega-3 release suggesting that these textiles are suitable as wound dressing for acute wounds. To conclude about the suitability of these textile in chronic wounds, a more extended release study is mandatory. Woven fabric functionalized with NLCs by exhaustion appears to be the most promising substrate since it presented the higher amount of released omega-3 during the 24 h test.

Almost all substrates functionalized with liposomes had a similar release profile with a maximum of RSV being released in the first hour (Fig. 5b). Cationized knitted fabric functionalized by impregnation of liposomes was the only exception. The higher affinity of negative charged liposomes to the cationized knitted fabric can explain the prolonged release profile. Moreover, in this substrate, not only the amount of RSV released was higher, but the maximum released was only reached after 6 h. The higher amount of RSV released is in accordance with the mass gain results. In comparison with other substrates and except from the non-woven fabric, the cationized knitted fabric showed the highest mass gain obtained (Fig. 1b).

In conclusion, knitted fabric cationized and functionalized by impregnation is the most promising technique to functionalize textiles with liposomes, since the amount of RSV released from this textile is higher and more prolonged.

4 Conclusions

The amount of NLCs and liposomes adsorbed was related with textile substrates adsorption capacity, being the non-woven textile the most efficient. The exhaustion technique was more suitable for NLCs incorporation whereas the impregnation technique was adequate for liposomes. The FTIR-ATR and DSC profiles confirmed the NLCs presence on the textile samples. Otherwise, no significant differences were observed between FTIR-ATR and DSC profiles samples functionalized with liposomes. Lipid nanosystems maintained the textiles permeability to water vapour. Moreover, functionalization of the textiles did not affect the permeability to air. Both characteristics are favorable for the healing process. Lipid nanosystems also present an inhibitory effect of NO production, regarded as important to combat inflammation and promote tissue healing.

The woven fabric functionalized by exhaustion with NLCs was the most promising for wound dressings application, since it had the most suitable omega-3 release profile (i.e., the higher amount released). Regarding the liposome's application, the most promising textile was the cationized knitted fabric functionalized by impregnation, since it was the substrate capable of retaining more RSV and showed a more prolonged release profile of this bioactive.

In conclusion, lipid nanosystems, such as NLCs and liposomes, can be used to improve textiles and enhance their application for anti-inflammatory effects in wound healing.

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Compliance with ethical standards

Conflict of interest On behalf of all authors, the corresponding author states that there is no conflict of interest.

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