Development of bioactive films based on chitosan and *Cynara cardunculus* leaves extracts for wound dressings

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**Abstract**

The development of natural-based, effective and protective wound dressings associated to local treatment applied on chronic wounds, represents a major challenge nowadays. In this work chitosan-based films were prepared with different concentrations of ethanolic ultrasound-assisted extracts from *Cynara cardunculus* leaves (EPIUAE). The physico-chemical film properties revealed that extract incorporation influences the volumetric swelling capacity and mechanical properties of the films, leading to materials with a lower fluid absorption capacity and more fragile. However, no impact was detected on the thermal properties of the films, as well as on their dense structure characterized by Scanning Electron Microscopy (SEM) analysis. Biological screening of chitosan-based films show that chitosan with 1% (w/w) and 5% (w/w) EPIUAE loading did not induce cytotoxicity on normal human skin fibroblasts (BJ-5ta cell line), mainly attributed to cyanaropicrin (extract key active compound) present in the film below its IC_{50} value. Nevertheless, chitosan-based films with 5% (w/w) EPIUAE presented an interesting anti-inflammatory activity; BJ-5ta cells stimulated with lipopolysaccharides (LPS), presented a reduction of 86% on IL-6 cytokine levels, after exposure to chitosan with 5% EPIUAE film extract. The results obtained in this study open up the possibility of successfully using chitosan films doped with EPIUAE for development of chronic wound dressings, with the advantage of using naturally-sourced materials with anti-inflammatory activity.

1. Introduction

Defined as a disruption resultant from physical or thermal damage, in the continuity of the epithelial lining of the skin or mucosa, a wound can be defined as acute or chronic according to its healing time [1]. Comprising a dynamic and complex process of tissue regeneration and growth, wound healing progresses through four different phases: coagulation and haemostasis, inflammation, proliferation and maturation [2]. Chronic wounds fail to progress through the normal stages of the healing cascade and are not repaired in an orderly and timely manner. Such failure may result from repeated tissue insults or underlying physiological conditions such as diabetes and malignancies, persistent infections, poor primary treatment, and other patient-related factors, presenting a biological, psychological, social, and financial burden on both individual patients and healthcare systems worldwide [3].

The development of advanced biomaterials for transdermal drug delivery is a growing area of biomedicine wherein intensive research is carried out, especially for chronic wound healing and/or efficient delivery of active pharmaceutical ingredients across skin. In fact, wound dressings are an important segment of the medical and pharmaceutical wound care market, with the Wound Dressing Market valued at USD 639.62 billion in 2018 and expected to reach up to USD 11168.93 billion by 2025 with a Compound Annual Growth Rate (CAGR) of 8.50% over the forecast period, according to the Global Advanced Wound Dressing Market Report, by Brandessence Market Research and Consulting Pvt. Ltd [4].

Considering a proper and timely wound healing, dressing selection should be mostly based on its ability to: i) provide or maintain moist environment; ii) enhance epidermal migration and promote angiogenesis, as well as connective tissue synthesis; iii) allow gas exchange between
wounded tissue and the environment, towards maintenance of appropriate tissue temperature, in order to improve blood flow to the wound bed; iv) protect against bacterial infection; v) be non-adherent to the wound and easy to remove after healing; vi) provide debridement action to enhance macrophage migration and support the accumulation of enzymes; and vii) be sterile, non-toxic and non-allergic [5].

Chitosan is one of the most studied biopolymers for wound dressings' application, mainly due to its biocompatibility and biodegradability, mostly arising from the presence of primary amines along the chitosan backbone [6]. Produced by chitin deacetylation, chitosan is a linear polysaccharide composed of glucosamine and N-acetyl glucosamine units linked by β (1-4) glycosidic bonds. Chitosan is an excellent candidate for wound dressing applications due to its non-toxicity, antimicrobial and anti-inflammatory properties, as well as its capacity to stimulate haemostasis and accelerate tissue regeneration [7-9]. Chitosan can be processed into different forms, such as hydrogels, sponges, oil or films, and the latter are characterized by being thin, less obtrusive with the ability to act as a vehicle for transdermal drug delivery [10,11].

Given their biological properties, safety and relatively low cost, bioactive plant extracts have been the subject of particular interest in the context of wound healing and transdermal drug delivery, namely as potential therapeutics for chronic wounds [12-15]. Comprising more than 5000 compounds, sesquiterpene lactones (SL) are one of the major groups of plant secondary metabolites presenting a high biological potential, namely anti-inflammatory [16-20]. Cynaropicrin, a guaianolide SL, has raised high scientific interest due to biological activities such as regulation of major adhesion molecules CD29 and CD59 [21] and anti-inflammatory response [20]. Recently, Brás et al. described cynaropicrin extraction from Cynara cardunculus leaves and their optimization process using ethanol as solvent and pulsed ultrasound as assisted methodology (EtPUEAE), followed by fractionation by membrane separation processes [22-24].

To the best of our knowledge, this is the first report on the development and characterization of chitosan films doped with Cynure cardunculus extracts, and evaluation of their potential as wound dressing. In the present work, EtPUEAE extracts from Cynara cardunculus leaves were incorporated into chitosan films, at different cynaropicrin/chitosan loadings (1, 5 and 10% w/w). The obtained films were characterized for their chemical and mechanical properties, as well as biological potential, with special focus on anti-cytotoxic and inflammatory activities.

2. Materials and methods
2.1. Reagents and standards
Chitosan ≥75% deacetylation degree was provided by Sigma Aldrich (USA). Acetic acid ≥99.8%, sodium chloride (≥99.5%) and potassium chloride (p.a.) were provided by Honeywell (USA), ethanol absolute ≥99.8% and potassium acetate were provided by Bodi Probabo (France), acetonitrile (HPLC grade) by Merck (France), and cynaropicrin was provided by Extrasynthese (France). Sodium bromide (99.0%) was purchased from Alfa Aesar (USA) Dimethyl sulfoxide (DMSO) cell culture grade was obtained from Applichem (Germany). 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) and propidium iodide was purchased from Calbiochem (USA). Acridine orange was purchased from Invitrogen (USA), DMEM, Medium 199, fetal bovine serum (FBS) suitable for cell culture, bovine serum albumin (BSA) (> 96% purity), lipopolysaccharides from Escherichia coli O111 B4 (LPS) and Piroxicam (≥98%) were purchased from Sigma Chemicals Co (Spain). Human IL-6 EUSA development kit (HRP) was purchased from Mabtech (Sweden).

2.2. Raw material
Cynara cardunculus L. (DC) leaves were collected in June 2016 at the Experimental Center of Agriculture School of Instituto Politécnico de Beja, Portugal and preserved at −80 °C. Before extraction, samples were freeze-dried in a freeze-drier (Scicac coolsafe, Labogene, Denmark). Freeze-dried leaves, with moisture content below the detection limit of 1%, were ground using a domestic mixed grinder (Moulinex, France), in order to obtain a powder constituted by particles with a size lower than 500 μm. After gridding, the freeze-dried powder was stored under vacuum at room temperature.

2.3. Cynaropicrin extraction
Ultrasound assisted extraction (UAE) experiments were performed using a 20 kHz UAE device (Bandelin HD3200, Germany), equipped with a 13 mm diameter titanium alloy (Ti-Al6-V4, Bandelin, Germany). Extraction was performed as described by Bis et al. [23]. Briefly, a pulsed UAE with a duty cycle of 25%, a solid/liquid (S/L) ratio of 1:27 (g/mL), extraction temperature of 44 °C and amplitude of 67% (1.43 μm) was performed during 30 min. After extraction, the ethanolic extract (EtPTUAE extract) was filtered using a glass filter funnel, with porosity G4 and a 10–16 μm maximum nominal pore size. The solvent was evaporated at low pressure on a rotary evaporator (Hei-VAP Advantage, HeiDolf, Germany) and dried extract was stored for 30 days at room temperature and used.

EtPTUAE cynaropicrin content was determined by HPLC A Dionex Ultimate 3000 system (Thermo Scientific, USA) equipped with a Diode Array detector was used. A Kinetex F5 2.6μm (4.6 × 150 mm) column (Phenomenex, USA) was employed at 30 °C, using a mobile phase of a mixture of water and acetonitrile (75:25 (v/v)) with a flow rate of 0.5 mL/min. All samples were pre-filtered with 0.22 μm pore size membrane filters ( Pall, USA).

2.4. Films development
Chitosan films, neat or with incorporated EtPTUAE extracts, were prepared by the solvent evaporation method and adapted from Ferreira et al. [25]. A Chitosan solution was prepared by adding 2% (w/v) in acetic acid aqueous solution 1% (v/v) with magnetic stirring at 200 rpm until the chitosan powder was completely dissolved. The polymer solution was vacuum filtrated twice using Miracloth® (Merck Millipore) as filter. Chitosan films doped with EtPTUAE extracts were also prepared. For that, EtPTUAE extracts on the proportion of 1, 5 and 10% (w/w) (cynaropicrin/chitosan) were firstly dissolved in 10 mL of ethanol and then added to a 25 mL filtered Chitosan solution. Chitosan and Chitosan with EtPTUAE forming solutions were centrifuged at 5000 rpm for 15 min to remove possible air bubbles. The solutions were then casted on polypropylene petri dishes (90 mm diameter) and dried at 40 °C for 48 h. Dried chitosan-based films were peeled off, weighted and equilibrated at 25 °C with different relative humidity ambiances, to further understand the influence of water on the final properties of the films. Defined saturated salt solutions were prepared in four different desiccators by mixing inorganic salts with distilled water, in order to obtain a specific relative humidity (RH) as described in Table 1.

2.5. Characterization of chitosan-based films
2.5.1. Scanning electron microscopy (SEM)
Surface and cross section images were used to detect morphology differences on chitosan films, with and without EtPTUAE extracts. The

<table>
<thead>
<tr>
<th>Salt</th>
<th>Relative humidity (RH%)</th>
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<tbody>
<tr>
<td>Potassium acetate</td>
<td>22.6</td>
</tr>
<tr>
<td>Sodium bromide</td>
<td>57.7</td>
</tr>
<tr>
<td>Sodium chloride</td>
<td>75.3</td>
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<tr>
<td>Potassium chloride</td>
<td>84.3</td>
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</table>
films surface and cross section images were obtained with different magnifications, respectively 1000× and 3000×, using an analytical Scanning Electron Microscope (Hitachi Ltd., Japan), equipped with a field-emission electron source of 15 kV. Prior to observation, films were cut with liquid nitrogen, mounted on a metal stub and coated with a thin layer film of gold-palladium.

2.5.2. Fourier transform infrared spectroscopy (FT-IR) analysis
For the analysis of possible interactions established between E2PUE extract components and chitosan, FT-IR spectra analysis was performed. FT-IR spectra were acquired using a Perkin Elmer Spectrum spectrometer (C105146, Perkin Elmer, France) equipped with Spectrum software (Perkin Elmer, France). Spectra were collected in the range of 4000-400 cm⁻¹ with 10 scans repetitions.

2.5.2. Swelling degree
In order to determine the swelling degree of the prepared films, samples were cut into 10 × 10 mm pieces, and thickness (e₀) was measured using a micrometer caliper (Mitutoyo, USA) with an accuracy of 0.1 μm. Films were immersed in phosphate buffered saline (PBS) solution at pH 7.4 during 24 h at 37 °C and, subsequently, the thickness (e₁) was measured [26]. The swelling volumetric degree (SW) was calculated using Eq. (1):

$$SW_{volumetric} = \frac{(e₁ - e₀)}{e₀} \times 100 \quad (1)$$

where SW_volumetric corresponds to volumetric swelling, e₀ and e₁ correspond to the initial and final film thickness, respectively.

2.5.4. Thermal properties
To evaluate the thermal stability of the films prepared, thermogravimetric analyses (TGA) were performed using a TGA Q50 analyzer (TA instruments, USA). The samples were heated at a constant rate of 10 K.min⁻¹, from room temperature up to 873 K, under argon atmosphere. The Universal Analysis software was used to determine the onset (T_onset) and decomposition (T_dec) temperatures.

2.5.5. Mechanical properties
Puncture tests were carried out using a TA.XTplus texture analyzer (Stable Micro Systems, UK) equipped with a 2 mm diameter cylindrical stainless steel probe. Samples were punctured through a hole (diameter of 10 mm), at a constant speed of 1 mm·s⁻¹.

Puncture stress (σ, Pa) was calculated according to the following equation:

$$σ = \frac{F}{S_c} \quad (2)$$

where F is the maximum force exerted by the probe (N) and S_c is the probe cross-sectional area (m²). In order to normalize results without the influence of membrane thickness, the puncture stress values were calculated (σ_on, MPa/mm) as follows:

$$σ_{on} = \frac{σ}{e_0} \quad (3)$$

2.5.6. Biological assays
2.5.6.1. Indirect cytotoxicity assays. Cytotoxicity assays were performed by an indirect method following the ISO 10993-5:2009 guidelines for biological evaluation of medical devices [27]. In order to collect film extract, pieces of 2 cm² were maintained in 5 mL of culture medium under controlled temperature (37 °C) and constant stirring rate (200 rpm) for 24 h. Extracts were then removed and diluted as follows: 100%, 75%, 50% and 25% using cell culture medium, for immediate use.

Bj5-ta human skin fibroblasts (American Type Culture Collection (ATCC)) were seeded at a density of 1.25 × 10⁴ cells/well on a 96-well tissue culture plate and incubated for 24 h at 37 °C in a humidified atmosphere with 5% CO₂ (Sanyo Electric Co, Japan). After this time, the culture medium was removed and cells were exposed to film extracts. Life control corresponded to cells simply incubated with culture medium (100% of the total volume). After 6 h and 24 h, cell metabolic assay was assessed by a MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphe-nyltetrazolium bromide, a tetrazole) assay, where 10% of the MTT reagent (0.5 mg/mL) dissolved in medium was added to each well, and cells were incubated at 37 °C for 2 h. The MTT solution was carefully aspirated, and formazan crystals were dissolved with a DMSO/ETOH (1:1 (v/v)) mixture. Absorbance was measured on a microplate reader (SpectraMax Plus, Molecular Devices, USA) at 570 nm. All the results were normalized relatively to life control, which was considered as 100% of viability. Triplicates were performed, in three independent experiments for each treatment, and the data were expressed as mean ± standard deviation. The viability results obtained for each film extract, at 6 and 24 h, were analyzed using the Prism version 6.00 for Windows (GraphPad Software, USA). Where differences did exist, the source of the differences at a p < 0.05 of significance level was identified by all pairwise multiple comparison procedure. The Tukey's test was used for pairwise comparisons.

2.5.6.2. IL6 (interleukin 6)-mediated skin inflammation. To determine the anti-inflammatory potential of film extracts, Bj5-ta cells were seeded at a density of 1.8 × 10⁵ cells/mL on a 12-well tissue culture plate for 24 h before the following procedure. Then, cells were stimulated with 5 ng/mL lipopolysaccharides (LPS) for 8 h. After stimulation, culture medium was removed and incubated cells were exposed to film extracts at 100% dilution, as well as to two E2PUE extracts with different concentration (1 μg/mL and 10 μg/mL) and two ciprofloxacin solutions with different concentrations (0.5 μg/mL and 5 μg/mL) for 16 h at 37 °C in a humidified atmosphere with 5% CO₂. Piroxicam, a known anti-inflammatory drug, at a concentration of 10 μg/mL, was used as negative control. A human ELISA kit was used to detect the IL-6 levels in accordance with the product manual. Triplicates were performed, in three independent experiments for each treatment, and the data was expressed as mean ± standard deviation. The IL-6 results obtained for each film extract, after 8 h of stimulation with LPS and after 16 h exposure to film extracts were analyzed using the Prism version 6.00 for Windows (GraphPad Software, USA). Where differences did exist, the source of the differences at a p < 0.05 of significance level was identified by all pairwise multiple comparison procedure. The Tukey's test was used for pairwise comparisons.

2.5.6.3. Acidine orange/propridium iodide staining assay. Bj5-ta cells were seeded at 1.8 × 10⁵ cells/mL on a 12-well tissue culture plate and left to stabilize overnight. The next day, the same conditions for IL-6 assays were replicated. After the respective incubations, cells were washed with PBS and incubated with a propidium iodide (PI) solution (50 μg/mL in PBS) for 15 min, after which they were incubated with an acidine orange (AO) solution (50 μg/mL in PBS) for another 15 min. In all incubation periods, cells were maintained in a humidified atmosphere at 37 °C, 5% CO₂ in the dark. Cells were then micrographed with an inverted fluorescent microscope (Olympus IX71), using FITC and TRITC filters.

2.5.6.4. Wound healing – scratch assay. Bj5-ta cells were seeded on an ibidi μ-slide 24 well (ibidi GmbH, Germany) at a density of 2.5 × 10⁵ cells/mL for wound healing assay and according to ibidi® instruction manual [28]. Cells were incubated in a humidified atmosphere at 37 °C, 5% CO₂, to allow cell adhesion and the formation of a confluent monolayer. Thereafter, ibidi culture inserts were removed, and cell monolayers with a cell-free gap of 500 μm ± 100 μm were gently washed with PBS, followed by addition of film extracts at a 100% dilution. Cells were then incubated for different time periods (0, 7 and 24 h). Cells
incubated in culture medium, within the same time periods, served as control. For each incubation period, the plate was placed under a phase-contrast microscope (Motic, China) equipped with a camera (Motic 2500, Motic, China) and images were acquired at 10× magnification with a computer-based microscopy imaging system (Motic Images Plus 2.0, Motic, China). Cell migration quantification was performed using Wound healing ACAS (Automated Cellular Analysis System) image analysis (Biodi GmbH, Germany). For each image, the gap area was measured at certain time intervals and compared to the initial gap area at time t = 0. Wound closure (expressed as a percentage) was calculated from the following equation:

\[
\text{Wound closure} (\%) = \left( \frac{W_0 - W_t}{W_0} \right) \times 100
\]

where, \(W_0\) and \(W_t\) correspond to the initial and time interval measured area, respectively.

3. Results and Discussion

Relative humidity may influence chitosan film properties. Based on this, chitosan and chitosan enriched with 1, 5 and 10% EtPUAE, were exposed to different relative humidity values until a constant mass was achieved. The films obtained were then characterized as described below.

3.1. Scanning electron microscopy (SEM)

Chitosan films obtained by solvent evaporation method were colorless, transparent and bright. Incorporation of EtPUAE extract at different chitin/chitosan ratios (1, 5 and 10% (w/w)), led to greenish but shiny films, as can be observed in Fig. 1 (a1 to d1).

Surface and cross section images of films obtained by SEM, at the four water activities under study, did not present significant differences between them, therefore, only surface and cross section images for an RH of 57.7% are presented in Fig. 1.

From the surface images obtained, it is possible to observe that the films prepared are dense and homogeneous, without visible cracks or defects, and no differences detectable between the chitosan film (Figure a2), and chitosan incorporated with EtPUAE extracts (1, 5 and 10% (w/w)) (Figures b2, c2 and d2, respectively). Regarding the cross-section images, the obtained results show that EtPUAE extracts are well dispersed in chitosan, presenting also a dense and homogeneous structure (Figures a3 to d3).

3.2. FT-IR

In order to understand the existence of possible molecular interactions between chitosan and EtPUAE extracts, a FT-IR analysis was performed to chitosan, chitosan +1% EtPUAE, chitosan +5% EtPUAE and chitosan +10% EtPUAE films, at a RH of 22.6%, 57.7%, 75.3% and 83.3%. The results obtained for films conditioned at a RH of 57.7% are depicted in Fig. 2. A similar pattern was observed for the four spectra obtained, although with some different transmittance intensities at certain peaks. Chitosan characteristic peaks corresponding to OH-stretching (3400-3300 cm⁻¹), N-H stretching in the amino group (3500-3300 cm⁻¹), C-H symmetric and asymmetric stretching (2900-2850 cm⁻¹), C-O stretching (amide I) (1650 cm⁻¹), N-H bending (amide II) (1539 cm⁻¹), C-N (amide III) (1323 cm⁻¹), asymmetric stretching of the C-O-C bridge (1153 cm⁻¹), and C-O stretching (1024 and 1066 cm⁻¹) were detected and are in accordance with previously reported results [29,30].

The addition of EtPUAE extracts did not reveal significant wavelength shifts, which may indicate the inexistence of covalent bonds established between the extracts and chitosan [31]. A new peak at 1746 cm⁻¹, characteristic of C=O stretching (6-lactone) [32], was observed as well as an intensity increase with an increasing extract concentration in the chitosan films (Fig. 2b). This peak, also identified on
Fig. 2. FT-IR spectra for chitosan, chitosan + 1% EtPUAE, chitosan + 5% EtPUAE, chitosan + 10% EtPUAE and EtPUAE at a RH of 57.7% where a) corresponds to full spectra and b) corresponds to detailed spectra on the sesquiterpene lactone peak wavelength zone (1700–1800 cm⁻¹).

Fig. 3. Chemical structures of cyanoprinin a) and chitosan b).

the EtPUAE spectra could be attributed to cyanoprinin, a sesquiterpene lactone identified as the major compound found on EtPUAE extracts [23].

For the chitosan film prepared with 10% (w/w) cyanoprinin, a decrease on the broad peak of -OH stretching (3400–3300 cm⁻¹) and/or -NH (1539 cm⁻¹) was observed and this may be due to possible binding interactions between cyanoprinin -OH groups (Fig. 3a) and -OH or -NH chitosan groups (Fig. 3b). Sun et al. [31] reported the same effect with hydrogen bonds established between polyphenols and chitosan, contributing to the interaction between these two compounds (Fig. 4).

When comparing the relative humidity effect on the chitosan film spectra it was possible to observe a decrease in characteristic peaks with reducing relative humidity. A low value for the -OH vibrational band can be attributed to a lower content of water molecules, whereas the increase of intensity from 57.7% to 84.3% of RH can be associated to increasing hydrogen bonding between chitosan chains, as explained by Sanchez et al 2017 [33].

When analyzing FT-IR spectra of chitosan +10% EtPUAE at 22.6 and 75.3% relative humidity and comparing it with neat chitosan FT-IR,

![Image of chemical structures and FT-IR spectra](image-url)

Fig. 4. FT-IR spectra for chitosan film conditioned at different relative humidity values: 22.6%, 57.7%, 75.3% and 84.3%.
respectively, Fig. 5a) and b), it is possible to observe that, at lower RH, the film with the extract presents an -OH vibrational band more intense than the film without extract. The higher intensity observed for the chitosan + 10% EPUAE film at low RH, comparatively to neat chitosan film, could be attributed to hydrogen bonding between extract and chitosan, which is in accordence with what was mentioned before. On the contrary, at higher RH, a similar effect cannot be observed because the higher water content overshadows the signal associated with hydrogen bonding.

3.3. Swelling

The creation of an optimal moist wound healing environment is critical to reduce the risk of complications on the surrounding skin and, based on this, wound dressings need to be able to absorb and retain exudates while maintaining an optimum level of moisture at the wound surface, which in turn must not spread to the surrounding skin [34]. Phosphate buffered saline (PBS) mimics for in vitro assessment, the osmolality and ion concentrations of the human body fluids. For this reason, PBS was selected for swelling experiments.

With respect to the swelling effect upon the different Woven/Chitosan ratios tested, it was possible to observe a general decrease of the film swelling capacity, with increasing EPUAE extract concentration (Table 2). The presence of hydrophobic compounds (e.g. cyanopiricin) reduces polymer hydrophilicity, decreasing its affinity to PBS, leading to a lower swelling ability as observed by several authors during the addition of lipidic fractions into chitosan films [30,35,38].

An increase of chitosan volumetric swelling was observed with an increase of the relative humidity, from 134% to 945% (p < 0.05) (Table 2).

As explained by some authors, a high relative humidity increases hydrophilic polymers moisture content, opening new binding sites for water, increasing its ability to PBS sorption, as observed for chitosan films [37]. The incorporation of EPUAE, due to its hydrophobic character, reverses this increase, leading to the lowest volumetric swelling effect. Considering an optimum exudate control on wound site, higher EPUAE content should be considered for light exudate wounds. Nevertheless, all films presented higher swelling degree comparing to the commercially available dressing, Tegaderm™ (6% after 48 h) [38].

3.4. Thermogravimetric analysis (TGA)

Thermogravimetric results for films of chitosan and chitosan loaded with EPUAE are represented in Fig. 6. For the chitosan film, a first mass loss was observed between 307 K and 428 K, corresponding to evaporation of water molecules present in the films [39]. Thermal decomposition of the chitosan film occurs between 400 K and 650 K, due to denaturation of the chitosan polymeric structure, which is in accordence with the literature [39–41].

In Table 3 and Fig. 6, it is possible to observe that, although T onset decreases with the increase of cyanopiricin concentration in films, such decrease is not significant, for the same relative humidity and for the different chitosan-based films under study. Exception made for the RH of 22.6%, where a higher percentage weight loss was observed for the chitosan film, which could be attributed to the lower moisture content, as observed from the swelling results.

The results obtained show that the thermal stability of chitosan films is not affected by the incorporation of EPUAE extracts and relative humidity. Considering the final application as wound dressing, the TGA analysis revealed that, at 37 °C, no weight loss was observed for the chitosan-based films, supporting their suitability for skin wound application.

3.5. Mechanical properties

The values of tension under puncture are represented on Table 4. The results show that, for the same relative humidity, the incorporation of EPUAE extracts promotes a decrease of the films tension upon rupture, although it is not observed a linear and statistically different decrease with the increase of the EPUAE film content (p < 0.05). As stress represents the maximum strength that films can support before breaking, the results indicate that films produced with the EPUAE

| Table 2: Volumetric swelling values obtained after 24 h immersion in a PBS solution, for chitosan, chitosan + 1% EPUAE, chitosan + 5% EPUAE, chitosan + 10% EPUAE, for films prepared under four different relative humidities of 22.6, 57.7, 75.3 and 84.3%.

<table>
<thead>
<tr>
<th>RH</th>
<th>Chitosan</th>
<th>Chitosan + 1% EPUAE</th>
<th>Chitosan + 5% EPUAE</th>
<th>Chitosan + 10% EPUAE</th>
</tr>
</thead>
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<tr>
<td>22.6%</td>
<td>134 ± 5%</td>
<td>151 ± 3%</td>
<td>103 ± 3%</td>
<td>76 ± 4%</td>
</tr>
<tr>
<td>57.7%</td>
<td>363 ± 25%</td>
<td>115 ± 9%</td>
<td>90 ± 6%</td>
<td>72 ± 12%</td>
</tr>
<tr>
<td>75.3%</td>
<td>453 ± 31%</td>
<td>94 ± 10%</td>
<td>62 ± 5%</td>
<td>46 ± 5%</td>
</tr>
<tr>
<td>84.3%</td>
<td>945 ± 30%</td>
<td>77 ± 8%</td>
<td>66 ± 4%</td>
<td>45 ± 6%</td>
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</table>

Average ± standard deviation estimated from three extracts analyzed in triplicate. Values with different superscripts, within the same column, are statistically different (p < 0.05).
extract incorporation present a lower mechanical resistance when compared with that produced only with chitosan. A similar behavior was observed by other authors, when adding carvacrol, α-tocopherol or apple polyphenols to a chitosan matrix [31,42,43]. It was proposed that the addition of hydrophobic compounds to the hydrophilic chitosan medium, induces the development of structural discontinuities in the highly packed chitosan matrix, lowering its ability to resist to mechanical solicitations [44]. The same behavior is envisaged to occur in the present work, considering the EPUAE hydrophobic behavior and the molecular interactions between EPUAE compounds and the chitosan polymer, which may weaken the polymer chain aggregation forces. Furthermore, films with the same extract content show a substantial decrease of puncture stress with an increase of the relative humidity. The lowest value (60.11 ± 5.66 MPa.mn⁻¹) was observed for the highest relative humidity (84.3%) and the highest EPUAE content (10%). This fact is attributed to the plastizer effect of sorbed water that reduces polymer-polymer interactions, lowering the films mechanical strength. However, the lower tensile strength obtained, did not prevent their use on skin healing since that according to some authors, the tensile strength of human skin tissue, varies between 5 and 30 MPa [45] although some authors have suggested that pre-stresses may be as great as 1 MPa [46].

### Table 3

<table>
<thead>
<tr>
<th></th>
<th>Chitosan</th>
<th>Chitosan + 1% EPUAE</th>
<th>Chitosan + 5% EPUAE</th>
<th>Chitosan + 10% EPUAE</th>
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<tr>
<td>T onset (K)</td>
<td>431</td>
<td>428</td>
<td>426</td>
<td>424</td>
</tr>
<tr>
<td>T dec (K)</td>
<td>538</td>
<td>532</td>
<td>532</td>
<td>542</td>
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</table>

**Table 3**

Onset temperature (*T*onset (K)) and decomposition temperature (*T*dec (K)) for the different films under study, chitosan, chitosan + 1% EPUAE, chitosan + 5% EPUAE and chitosan + 10% EPUAE and for the relative humidity values tested: 22.6, 57.7, 75.3 and 84.3.

### Table 4

<table>
<thead>
<tr>
<th>RH (%)</th>
<th>Tension (MPa.mn)</th>
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<tbody>
<tr>
<td></td>
<td>Chitosan + 1% EPUAE</td>
</tr>
<tr>
<td>22.6</td>
<td>215.55 ± 10.60*</td>
</tr>
<tr>
<td>57.7</td>
<td>117.00 ± 0.89*</td>
</tr>
<tr>
<td>75.3</td>
<td>115.32 ± 4.37*</td>
</tr>
<tr>
<td>84.3</td>
<td>86.59 ± 1.05*</td>
</tr>
</tbody>
</table>

**Average ± standard deviation estimated from three extracts analyzed in triplicate. Values with different superscripts, within the same column, are statistically different (p < 0.05).**
**Table 5**

<table>
<thead>
<tr>
<th>Film</th>
<th>Cell viability (%)</th>
<th>Cyn film extract (g/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>6h</td>
<td>24h</td>
</tr>
<tr>
<td>Chitosan</td>
<td>98.4 ± 4.6%</td>
<td>95.8 ± 7.1%</td>
</tr>
<tr>
<td>Chitosan + 1% EPIUAEX</td>
<td>95.7 ± 5.0%</td>
<td>88.5 ± 8.9%</td>
</tr>
<tr>
<td>Chitosan + 5% EPIUAEX</td>
<td>75.6 ± 5.7%</td>
<td>81.4 ± 8.9%</td>
</tr>
<tr>
<td>Chitosan + 10% EPIUAEX</td>
<td>51.1 ± 5.0%</td>
<td>34.1 ± 2.7%</td>
</tr>
<tr>
<td>EPIUAEX</td>
<td>26.1 ± 1.6</td>
<td>24.1 ± 1.1</td>
</tr>
<tr>
<td>Cynaropicrin</td>
<td>15.9 ± 1.4</td>
<td>9.7 ± 1.0</td>
</tr>
</tbody>
</table>

Average ± standard deviation estimated from three extracts analyzed in triplicate. Values with different superscripts, within the same column, are statistically different (p < 0.05).

### 3.6. Biological activity

#### 3.6.1. Cytotoxicity

Biocompatibility, including low or absence of cytotoxicity are key factors for a material to be considered suitable for use as medical device. According to ISO 10993-5:2009 guidelines for biological evaluation of medical devices, a reduction on cell viability higher than 30% is considered to present a cytotoxic effect [27]. Indirect cytotoxicity assay for chitosan films, with extracts at different dilution rates, were performed with telomerase-immortalized human normal skin fibroblasts (BJ-5-ta cell line) for incubation periods of 6 and 24 h, as presented in Table 5. According to the obtained results, after 6 h of exposure, the chitosan film led to a cell viability of 98.4 ± 4.6%, and the chitosan + 1% EPIUAEX film resulted in a cell viability of 95.7 ± 5.0%, with no statistical difference between them (p < 0.05). Although leading to a lower cell viability, according to the ISO 10993-5:2009 guidelines, the chitosan + 5% EPIUAEX film (75.51 ± 5.67) was not considered cytotoxic. On the other hand, with a cell viability of 51.09 ± 5.03%, the chitosan + 10% EPIUAEX film was considered cytotoxic. After 24 h, no statistical difference was observed between the values for the non-cytotoxic films (neat chitosan and chitosan + 1% EPIUAEX).

The IC₅₀ values at 6 and 24 h for the EPIUAEX extract and pure cyanaropicrin was determined. In addition, cyanaropicrin content in the film extracts used for the cytotoxicity experiments, was quantified. Analysis of the cyanaropicrin content, shows that films with incorporation of 1 or 5% (w/w) EPIUAEX extract contain respectively 0.177 and 2.96 μg/mL of cyanaropicrin, a concentration below its IC₅₀ (15.92 ± 1.45 μg/mL). Regarding the film of chitosan + 10% EPIUAEX, at 6 h of incubation, a significant cell viability reduction was observed (51.09 ± 5.03%). The final cyanaropicrin concentration of film extract was 17.73 μg/mL (chitosan + 10% EPIUAEX film) indicating that the observed cytotoxicity is most probably directly correlated with the cyanaropicrin concentration in the film.

#### 3.6.2. IL-6 (interleukin-6)-mediated skin inflammation

Continuous inflammation is a major cause of chronic wounds, and the expression modulation of pro-inflammatory interleukins is a suitable way to stop the inflammatory process, reducing epithelization, thus promoting a normal wound healing process [47]. IL-6 is one of the key pro-inflammatory factors produced by dermal fibroblasts in response to certain stimuli [48]. In this work, dermal fibroblasts (BJ-5-ta) were stimulated with lipopolysaccharide (LPS) and after this activation, which induces the IL-6 production, cells were exposed to the chitosan-based films under study. As controls, the EPIUAEX extract and a cyanaropicrin solution were directly applied to the stimulated BJ-5-ta cells.

Interestingly, the IL-6 levels were reduced on stimulated BJ-5-ta cells exposed to the film extracts (neat chitosan, chitosan + 1% EPIUAEX and chitosan + 5% EPIUAEX) (Fig. 7). A detailed analysis of results shows that, for the neat chitosan film, a reduction of IL-6 expression by LPS-stimulated fibroblasts was achieved, without statistical difference, in comparison with 10 ng/mL piroxicam. Incorporation of EPIUAEX into chitosan films seems to promote an anti-inflammatory effect as expressed by reduction of IL-6 levels, although without statistical difference comparing to neat chitosan (p > 0.05). The strongest effect was observed for...
Table 6
Cell viability for B16-ta LPS-stimulated cells after 16 h of exposure to films of chitosan + 5% EPUAE film compared to that of chitosan + 1% EPUAE, chitosan + 5% EPUAE, and EPUAE at 1 and 10 μg/mL, calculated from AO + PI assay.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Cell viability (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chitosan + 5% EPUAE</td>
<td>96.0 ± 0.2</td>
</tr>
<tr>
<td>Chitosan + 1% EPUAE</td>
<td>95.0 ± 0.2</td>
</tr>
<tr>
<td>Chitosan + 5% EPUAE</td>
<td>90.0 ± 3.2</td>
</tr>
<tr>
<td>EPUAE 1 μg/mL</td>
<td>96.0 ± 0.7</td>
</tr>
<tr>
<td>EPUAE 10 μg/mL</td>
<td>90.0 ± 3.6</td>
</tr>
<tr>
<td>Piroxicam</td>
<td>79.0 ± 4.1</td>
</tr>
</tbody>
</table>

the chitosan + 5% EPUAE film, with a reduction of ≈ 86% and 83% in the IL-6 expression by skin fibroblasts, comparatively to LPS and the neat chitosan film (p < 0.05), which is extremely promising for regulation of continuous inflammation in chronic wounds.

Further, the effect of the EPUAE extract and the cyanoapicrin solution was also evaluated, taking into account the cyanoapicrin levels quantified in chitosan-based films (Table 5). With a cyanoapicrin content of 51.7 mg/g extract, a higher reduction of IL-6 levels was observed with the EPUAE extract, when compared to the pure cyanoapicrin solution (in the same concentration range). Although also effective, the EPUAE extract at 10 μg/mL has a lower capacity to reduce IL-6 levels in stimulated cells when compared to the EPUAE 1 μg/mL (p < 0.05), which may be related to a slight increase in the material's toxicity with the higher extract load. Although EPUAE, as well as cyanoapicrin, can reduce IL-6 levels according to the results presented in Fig. 7, there are no differences in regard to this cytokine quantification for cells incubated with EPUAE extract (10 μg/mL) or cyanoapicrin (0.5 and 5 μg/mL, 1.44 and 14.4 μM, respectively) (p > 0.05). These results support a more potent anti-inflammatory effect for the EPUAE extract, comparatively to cyanoapicrin, underlying an interesting synergistic effect for the extract.

Taking into account the cytotoxicity associated to cyanoapicrin at 5 μg/mL (14.4 μM) (Table 5), a dual staining with acidine orange (AO) and propidium iodide (PI) was performed to distinguish viable cells from apoptotic/necrotic cells. Vander Berghe et al. [52] observed that necrosis is induced by several stimuli, such as tumor necrosis factor (TNF), anti-Fas (an apoptotic mediator) or double-stranded RNA, promote the secretion of the pro-inflammatory cytokine IL-6. According to our results, Table 6 and Fig. 8, the EPUAE extract at 10 μg/mL, and the cyanoapicrin solution at 5 μg/mL, induce a slight increase in cell death, as more PI (red) fluorescence is visible, comparatively to the lower tested concentrations. This could be related to necrosis promoted by high levels of the EPUAE extract and cyanoapicrin, mediated by increased levels of IL-6, as described by Vander Berghe et al.

![Fig. 8. Acidine orange (AO, green) and propidium iodide (PI, red) double staining fluorescent micrographs of LPS-stimulated B16-ta cells after 16 h of exposure to: (a) chitosan film, (b) chitosan + 1% EPUAE film, (c) chitosan + 5% EPUAE film, (d) EPUAE 1 μg/mL, (e) EPUAE 10 μg/mL, (f) positive control (LPS), (g) negative control (piroxicam) (h) unstimulated cells (life control). The images were captured in multiple times with a ×200 magnification.](image-url)
Regarding the chitosan-based films, a very marginal reduction in the number of viable cells (green fluorescence) was observed for the chitosan + 5% EtPUAE film, which may also be related to extract-related toxicity.

Although this is the first time that cyanaropicin and EtPUAE extracts are tested in LPS-stimulated 875-ta fibroblasts, cyanaropicin’s anti-inflammatory potential has been previously monitored by IL-6 expression downregulation in human keratinocytes. According to Takeda et al., cyanaropicin at 1 \( \mu \text{M} \) appeared to inhibit IL-6 and TNF-\( \alpha \) upregulation in UVB-treated normal human epidermal keratinocytes [48]. The results presented here show that EtPUAE extracts and loaded chitosan films reveal a similar IL-6-mediated anti-inflammatory activity in skin fibroblasts, which supports their applicability for chronic wound management.

### 3.6.3. Scratch assay

Although a stronger reduction in IL-6 levels of LPS-stimulated fibroblasts was observed for the film of chitosan + 5% EtPUAE, a decrease in cell viability was also evident, which indicates a potential toxicity issue.

Therefore, wound healing assays were only performed for films of chitosan and of chitosan + 1% EtPUAE.

Fibroblasts are responsible for most collagen and elastin synthesis as well as organization of the extracellular matrix (ECM) components [50]. Therefore, normal skin fibroblasts were selected for this study due to their need for determination of wound closure. Analysis of results reported on Table 7 and Fig. 9, shows that, after 7 h of incubation, films with extract did not affect fibroblasts migration with statistical significance (\( p < 0.05 \), comparatively to the sample control. On the other hand, after 24 h of incubation, a more effective wound closure was observed for the film with 1% EtPUAE (\( p < 0.05 \)), which suggests that a concentration of 0.177 \( \mu \text{g/mL} \) cyanaropicin can promote a faster wound healing.

### 4. Conclusions

In this work chitosan and chitosan films loaded with different EtPUAE extracts were prepared, conditioned at different relative humidity values and characterized in terms of structure, swelling, thermal and mechanical properties. In addition, preliminary assessment of the biological properties (cytotoxicity, IL-6 mediated skin inflammation and wound healing) were performed in order to evaluate the prospective use of the films developed as wound dressing materials.

SEM analysis of chitosan and chitosan-based films produced, revealed a homogeneous and dense structure, although after EtPUAE loading, to the naked eye, green and shiny films were obtained, in opposition to clear and translucent neat chitosan films. A decrease of the volumetric swelling capacity was observed with the EtPUAE loading, attributed to the extract hydrophobic character, which could be advantageous for wound dressings where an excessive fluid sorption is not necessary.

![Fig. 9. Microscopic images representing the in vitro scratch assay of chitosan-based films: 875-ta cells were incubated in the presence and absence of chitosan-based films and images were captured at 7 h and 24 h. The boundaries of the scratched wounds were determined by the yellow lines.](image-url)
References


