Natural Rubber Films Incorporated with Red Propolis and Silver Nanoparticles Aimed for Occlusive Dressing Application

Charlene Silvestrin Celi Garcia a, Ana Elisa Dotta Maddalozzo a, Paulo Miguel Celi Garcia a, c, Cristian Padilha Fontoura b, Melissa Machado Rodrigues b, Marcelo Giovanela b, João Antonio Pêgas Henriques b, Cesar Aguzzoli b, Janaina da Silva Crespo b, Mariana Roesch-Ely a*

aUniversidade de Caxias do Sul, Instituto de Biotecnologia, Caxias do Sul, RS, Brasil.
bUniversidade de Caxias do Sul, Área do Conhecimento de Ciências Exatas e Engenharias, Rua Francisco Getúlio Vargas, 1130, 95070-560, Caxias do Sul, RS, Brasil.
cSociedade Brasileira de Cirurgia Plástica (SBCP), Rua Mato Grosso, 306, Higienópolis, 01239-040, São Paulo, SP, Brasil.

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In this work, NRL films were enriched with red propolis and silver nanoparticles with the aim of obtaining dressings with bactericide properties. To improve the properties of these materials, plasma treatment was also applied to the films. Cytotoxic effects were evaluated by MTT and agar diffusion assays on secondary and stem cells. Determination of cell behavior on the films were assessed by EDS and FEG-SEM. In general, the plasma modified NRL films presented hydrophilic character, favorable condition for biomedical applications. RBS results showed that silver is probably in nanoparticulate form. MTT assays revealed that non-cytotoxic effects were observed for all the samples extracts. Either the plasma treatment or silver deposition were effective in sterilizing the samples and modifying their surfaces. It was demonstrated in this in vitro study that ADSCs associated with NRL films enriched with propolis should be considered promising for application in the healing of occlusive wounds.

Keywords: natural rubber latex films; plasma treatment; wound dressing; red propolis; silver nanoparticles.

1. Introduction

Burnings represent a severe risk to human health, being the third cause of accidental death on a global scale 1. The lack of skin tissue coverage increases vulnerability to infections and allows microorganism proliferation. As a consequence, functions related to skin protection against infections, lesions and trauma become compromised 2. In this scenario, biomaterials applied as wound dressings guarantee the integrity of the tissue and its protective function.

Natural rubber latex (NRL) has attracted much attention amongst biomaterials for activating biological healing responses 3, such as angiogenic activity, which is the ability of healing burns through the promotion of blood vessel formation 4. Due to this property, it can be used to aid people who suffer from diabetes and the accompanying hardships of wound healing 5.

A special concern related to burns is the predisposition to secondary infection processes, which is responsible for about 75% of registered deaths amongst burnt patients 6. The use of substances that have antibacterial properties in association with latex minimizes infections, like the ones caused by Staphylococcus aureus, commonly found in burnings 7,8.

Literature has reported the antibacterial activity of red propolis due to its complex and varying chemical composition 9 which includes flavonoids and aromatic acids, as well as different types of terpenoids 10,11. It is constituted of exudates from floral shoots and buds of various plants, presenting many benefits to human health, like antioxidant, anti-inflammatory and regenerating properties 9-12. Red propolis (RP) was chosen to be studied in this work, as it is a product that can be only found in Brazil, so it represents a native resource that can help in public health. Brazil is considered one of the greatest propolis producers in the world due to the large diversity of ecosystems and to the favorable climate 13,14. Also, there are not many studies about RP in comparison to green propolis (the most commercialized type). Besides differences in the chemical composition of both products, the literature reported that RP presents more toxic activity than the green sort 15.

The association of latex films to silver guarantees further bactericide character and low toxicity to human cells 16,17. Silver as a topical antimicrobial agent has been featured in wound dressing applications 18-20, being useful in the treatment of acute wounds, burns or surgical wounds and effectively accelerating the healing process 17. Silver is effective against Escherichia coli, one of the most difficult
microorganisms to treat and found in burning injuries. This type of bacteria has become multi-resistant to antibiotics, the reason why silver is a good option to be used\textsuperscript{21,22}. When in nanoparticle or ion form, silver has antimicrobial effects against Gram-positive and Gram-negative bacteria, so it is largely employed in biomedical industry\textsuperscript{23,24}.

Besides protection against microorganisms\textsuperscript{25}, a desirable feature for occlusive dressing materials is avoidance of fibroblast adhesion, so the removal of the dressing from the wound will not damage the newly generated tissue\textsuperscript{26,27}. In addition, a dressing should be hydrophilic, because it will accelerate scarring process as wound-released exudates are absorbed without compromising the moist environment\textsuperscript{28}. Literature has pointed that surfaces of low hydrophilicity encourages growth of bacteria, compromising healing\textsuperscript{29}.

Aiming to enhance hydrophilicity, avoidance of cell adhesion and bactericide activity, different processes in this study such as silver nanoparticles (AgNPs) deposition, incorporation of RP and plasma treatment were tested and compared trough the latex samples as promising products. Plasma treatment is a vastly used technique for modifying all classes of materials, sterile packaging and biomedicine-aimed products, but it is efficiency has not yet been exploited thoroughly in wound dressing applications\textsuperscript{30}. Apart from that, another advantage of plasma-assisted processes is that the modification takes place solely on the surface being treated, while bulk properties, such as mechanical ones, are maintained\textsuperscript{31}. In this case, the elastic properties of latex would be maintained and remain ideal for wound dressing, as the limbs could move freely without breaking the polymeric membrane.

In polymers, plasma-assisted processes are used to modify surface properties of great importance in biomaterials\textsuperscript{32}, such as wettability, permeability and biocompatibility. When a polymeric material is submitted to plasma treatment, it interacts with the ionized gas environment and generates new functional groups, which are responsible for modifying the aforementioned properties, as plasma active species interact with molecules on the polymer’s surface\textsuperscript{33}. Apart from benefiting from many biomedically relevant physicochemical properties, plasma treatment can effectively sterilize the treated surface\textsuperscript{34}, reason to why the authors chose to use this technique. These treatments are also eco-friendly, so they do not damage the environment.

For the treatment of chronic injuries, the association of autologous stem cells and biomaterials is recommended\textsuperscript{35}. The application of autologous adipose derived stem cell (ADSCs) with biocompatible materials can be a promising strategy to assist burn patients who require a large graft area to cover extensive sites and who are limited by the availability of donor skin\textsuperscript{36,37}. Some advantages of working with ADSCs are the large amount of these cells found in our body, the facility to be collected with little risk of morbidity to the donor, presence of plasticity, multipotentiality and capability to differentiate into epithelial tissue, including dermis and epidermis\textsuperscript{38}.

In this context, the main goal of this work was to produce NRL films for a possible application as wound dressings. Physico-chemical characterization and biological assays were also carried out to investigate surface properties and the biocompatibility of these films during contact with steam cells and secondary fibroblasts.

2. Materials and Methods

2.1. Natural rubber latex (NRL)

NRL used to produce the matrix of the films was acquired by Du Latex (São Paulo, Brazil) and is derived from trees of the same clone. A centrifugation process was performed to decrease the amount of allergenic proteins by stabilizing sulfur and ammonium hydroxide. The elastomer belonged to lot LA032618, with a pH of 10.4.

2.2. Red propolis (RP)

RP from Alagoas State (Northeast region of Brazil) was used in this study. RP was stored in a dry, light exposure-free environment and frozen at –20 °C until the extract was prepared. RP samples were milled into a thin powder and 50 g were mixed into 500 mL of EtOH-H\textsubscript{2}O 70%-30% (v/v). This mixture was brought to mechanical stirring for a period of 24 h at room temperature and then filtered. The solvent was later evaporated at 37 °C and freeze-dried. At the end of this procedure, RP was kept at –20 °C until analyses were performed.

2.3. Natural rubber latex films

NRL films were prepared by shedding 40 mL of latex in a Petri dish, which was inserted on a furnace (Q317M – Quimis, Diadema, Brazil) for 24 h at 65 °C, until complete dry. Biological properties are favorable when heat treatment on NRL is conducted between 65–85 °C. At this temperature range, rubber particles coagulate faster during the evaporation of latex byproducts, thus supplying a greater presence of phospholipids and proteins not bonded to rubber particles. These proteins are responsible for angiogenic activity in latex\textsuperscript{39}, so biological properties are favored.

2.4. Natural rubber latex films containing red propolis (NRL-RP)

RP was diluted in 2 mL of EtOH-H\textsubscript{2}O 50% (v/v), and then added to 38 mL of NRL. The mixture remained on a furnace (Q317M – Quimis, Diadema, Brazil) for 24 h at 65 °C until complete dry. Figure 1 displays a flow chart of how RP was prepared until incorporation into NRL.

2.5. Plasma treatment

The NRL and NRL-RP films were inserted into a vacuum chamber (homemade), which was evacuated up to 3.0 × 10\textsuperscript{-2} mbar. The gases used for plasma discharge were oxygen (Linde, Brazil, purity of 99.9992%) and hydrogen (Linde, Brazil, purity of 99.9992%) in a 1:1 ratio. Work pressure of 1.0 mbar was used. To generate the electric potential, a 13.56 MHz radiofrequency source (Tokyo Hy-Power RF-300, Japan) was applied with 30 W of power. The process was carried out for 10 min. It is important to highlight that input power must be kept at the minimum possible value, once temperature may ignite
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Degradation on NRL. High frequency glow discharges, operating at 13.56 MHz and low pressures (≤ 100 Pa), are usually adequate for general polymeric films due to excellent uniformity (i.e., plasma permeation) in the treatment and absence of hot arcs, which may harm the material\textsuperscript{39}. The films produced in this process were NRL-Pla and NRL-RP-Pla.

2.6. Silver deposition

Silver deposition on the NRL and NRL-RP films was carried out by magnetron sputtering technique. A solid silver target (99.9% purity) distanced 6.0 cm from samples was used. The base pressure was of 3.0 × 10\textsuperscript{-7} mbar and argon (Linde, Brazil, 99.9992%) plasma was ignited. NRL films containing silver (NRL-Ag) and silver + RP (NRL-RP-Ag) were produced. The optimized parameters used in the silver deposition were shown in Table 1.

To form nanoparticles instead of a continuous film, the time and power of the process was set as low. In addition, the treatment took place at room temperature, so that the silver does not diffuse into the matrix and has difficulty in being leached.

| Experimental parameter | Value          |
|------------------------|----------------|
| Work pressure          | 2.0 × 10\textsuperscript{-7} mbar |
| DC power               | 20 W           |
| Argon flow             | 5.0 cm\textsuperscript{3}/min |
| Temperature            | 25 °C          |
| Deposition time         | 3.0 s          |

2.7. Physico-chemical characterization

2.7.1. Attenuated total reflectance Fourier transform infrared spectroscopy (ATR-FTIR)

The physico-chemical characterization of the NRL and NRL-RP films was carried out by analyzing their functional groups through ATR-FTIR using a spectrophotometer equipment (model 400, Perkin-Elmer, USA). The spectra were registered in transmittance mode using an interval of wavelengths between 4000 and 500 cm\textsuperscript{-1}.

2.7.2. Elemental quantitative analysis (RBS)

Elemental quantitative analysis was done by Rutherford backscattering spectrometry (RBS). A 3.0 MV Tandem ion accelerator with a monoenergetic ion beam of He\textsuperscript{+} of 2.0 MeV and backscattering detection angle of 165° was used. Through this analysis, it was possible to obtain the concentration of deposited silver on the produced NRL-Ag films.

2.7.3. Water contact angle (WCA)

Through sessile drop technique, it was possible to measure the contact angle to determine the wettability of the films (NRL, NRL-Ag, NRL-RP, NRL-RP-Ag, NRL-Pla) in a goniometer (model 300, SEO Phoenix, South Korea).

Three drops were tensioned with distilled water, each with 10 measurements at room temperature. Shelf life was estimated along a period of 5 months to evaluate the maintenance of obtained wettability as function of time. The samples were stored at room temperature/light inside a Petri dish to avoid humidity.

2.8. Biological assays

Assays were performed on all the samples (NRL, NRL-Ag, NRL-RP, NRL-RP-Ag, NRL-Pla and NRL-RP-Pla).

Figure 1. Flow chart for the methodology adopted in the preparation of NRL-RP films.
2.8.1 Evaluation of the efficiency of the plasma sterilization process

The evaluation of the effectiveness of the plasma sterilization process was performed by submerging samples in the thioglycolate broth (Biomerieux, St, Lombard, USA) medium during 14 days. After this period, the appearance of turbidity in the thioglycolate medium was observed in case of contamination. Thus, the samples that showed no turbidity of the medium were classified as effectively sterilized.

2.8.2 Isolation of adipose tissue-derived stem cells

This study was approved by the Committee of Ethics in Research of the University of Caxias do Sul (Rio Grande do Sul State, Brazil) under the number 1.230.848 and is found in the system Comitê de Ética em Pesquisa/Comissão Nacional de Ética em Pesquisa (CEP/CONEP), Plataforma Brasil, on a national and unified basis of registers for research involving human beings, belonging to the Ministry of Health under the protocol CAAE: 46199015.2.0000.534. The cell isolation was based on the protocol described by Zuk et al.40, with some modifications.

In the ZuK protocol, collagenase I is used. In our isolation of stem cells from adipose tissue, collagenase II was used. The adipose tissue was consecutively washed with 1× PBS and submitted to the enzymatic process of complete tissue digestion with 1.0 mg/mL of collagenase II (Sigma Aldrich, St. Louis, Missouri, USA) in 1× PBS for 30 min in a water bath, with shaking every 5 min. After, the adipose tissue was centrifuged for 10 min at 1400 rpm, the supernatant was removed and stromal vascular fraction (SVF) was isolated. Dulbecco’s modified eagle medium (DMEM) supplemented with 10% of fetal bovine serum (FBS) and 1% of penicillin/streptomycin (P/S) was added to the inactivation of the enzymatic digestion process and centrifugation was performed at 1000 rpm for 10 min. The pellet was resuspended in DMEM/F12 supplemented and seeded in culture bottle and kept in a humidified incubator at 37 °C with 5% of CO₂.

In addition, stem cells were characterized by the methodology described by Dominic et al.41, with modifications; the cell differentiation in vitro for bone, cartilage and adipose tissue was proceeded using a cell differentiation proper medium.

2.8.3 Cytotoxicity assays

The cytotoxicity assays were carried out using both mouse fibroblasts L929 cells and ADSCs. To determine the cytotoxicity and the biocompatibility of the films, standardized methods were applied according to ISO 10993-0542 and ISO 10993-143. To obtain an extract solution to be used upon cells in 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay, the DMEM culture medium supplemented with 10% of FBS and 1% of P/S was put into contact with the samples at a 0.1 g/mL concentration during 24 h at 37 °C in 5% of CO₂.

2.8.4 Indirect MTT cytotoxic assays

Cytotoxicity was evaluated by indirect MTT assays. This method is based on the reduction of MTT to formazan crystals by mitochondrial dehydrogenases44. L929 and ADSCs were seeded at a density of 5.0 × 10⁴ cells/mL in 100 μL of DMEM culture medium supplemented with 10% of FBS and 1% of P/S. After obtaining a confluence of 70–80%, cells were treated with the extract solution obtained as previously reported from the immersion of the samples in culture medium for 1, 2 and 7 days at 37 °C and 5% of CO₂. For negative control, a DMEM medium (10% of FBS and 1% of P/S) was used and for positive control, 5% of dimethylsulfoxide (DMSO). The medium was removed and 1.0 mg/mL of MTT in a medium free from FBS and P/S was added to the wells of microplates. The plates were incubated at 37 °C during 2 h in a humidified atmosphere with 5% of CO₂. Subsequently, MTT solution was removed and formazan crystals were dissolved in 100 μL of DMSO. Spectrophotometric reading at 570 nm was carried out using a microplate reader (Max 190 Spectra, Molecular Devices, USA) and results were expressed as percentage of cell viability. Absorbance of negative control represented 100% viability and values of treated cells were calculated as a percentage of control.

Statistical significance was evaluated through t-test and One-way analysis of variance (ANOVA) with the single-step multiple comparison procedure (Tukey) to evaluate the statistical differences in case of normal distribution. In this work, we used six different samples and three different times, so the purpose of applying ANOVA was to perform the statistical test to check if there is a difference in the distribution of a measure between the groups. Significance was accepted for p-value inferior to 0.05, using Statistical Package for the Social Sciences (SPSS, version 19.0) for Windows.

2.8.5 Agar overlay assay

Cytotoxicity was evaluated by the indirect method of the neutral red dye assay with transformation of lysosomes from viable cells45. L929 and ADSCs were seeded into 6-well plates at a density of 1.0 × 10⁴ cells per well in a DMEM culture medium supplemented with 10% of FBS and 1% of P/S. After 24 h, the culture medium was removed and 1.0 mL of the covering medium was added to each well. The covering medium was composed by 1.8% of agar with neutral red dye at 0.01% and concentrated MEM 2X, in equal amounts.

Then, the sterilized samples were put at the center of each well in an incubator at 37 °C, with 5% of CO₂ for 24 h. The samples were evaluated macro and microscopically (Table 2) by observing halo formation around the samples, which is correlated to their cytotoxic features.

2.8.6 Determination of topographical distribution and morphological patterns of L929 and ADSCs deposited on the films

For determining properties such as adhesion, topographical distribution and morphology, L929 and ADSCs were seeded into 6-well plates at a density of 5.0 × 10⁴ cells/mL during 1, 2, and 7 days. Exactly 2000 μL of DMEM culture medium supplemented with 10% of FBS and 1% of P/S was used over the samples for 1, 2 and 7 days. Following this, cells were fixed with 3% glutaraldehyde solution in PBS (v/v) for 15 min and dehydrated with increase ethanol grade 30, 50, 70, 90 and 100% (v/v) during 10 min for each concentration. The samples were kept inside a desiccator until analysis. A
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Field emission gun equipped scanning electron microscope (FEG-SEM) (Mira 3, Tescan, Czech Republic) was used in these assessments. The images were made in the treated side of the samples. The working distance was between 15 and 20 mm and the accelerating voltage was either 10 or 15 kV.

2.8.7 Elemental analysis through energy dispersive spectroscopy (EDS)

The elemental analysis of the films was obtained through a microscope (Mira Tescan 3), equipped with an energy dispersive spectrometer (EDS) of the silicon drift detector (SDD) type. The images were obtained from the treated side of the samples. The working distance was between 15 and 20 mm and the accelerating voltage was 15 kV.

3. Results and Discussion

3.1. Attenuated total reflectance Fourier transform infrared spectroscopy (ATR-FTIR)

Figure 2 shows the ATR-FTIR spectra for NRL and NRL-RP films. As can be observed, both spectra are very similar, as already seen on the literature\(^\text{46}\). This is an indicative that the functional groups on NRL and RP are almost the same. New bands were not seen on the NRL-RP spectrum in comparison with the NRL spectrum, so no chemical interaction between latex and red propolis took place. Physical interaction between NRL and RP, by other hand, is desired, once it allows the gradual and easier release of RP compounds into the wound environment\(^\text{47}\). In order to study the functional groups present in the NRL matrix, which will be the same for all films produced and, therefore, has great importance for this study, the wavelengths related to its compounds were analyzed. The attribution of the main bands related to ATR-FTIR spectrum of NRL is listed in Table 3. Absorption bands for the characteristic vibrational modes of cis-1,4-polyisoprene are highlighted in bold.

Because they have carbonyl groups (aldehydes, esters, carboxylic acids, amides and ketones), the region between 1800 and 600 cm\(^{-1}\) is of greatest interest. These functional groups are mostly related to latex proteins, which are associated to possible regenerative process. It is noted that there is no indication of sulfur presence, which is used in latex during manufacturing for stabilization\(^\text{38}\). This fact indicates that the amount of this element is relatively low, which favors biomedical use.

Adsorption bands that characterize RP are mostly present in 1646, 1451, 1249 and 1043 cm\(^{-1}\), concurring with literature reports\(^\text{39,41}\). These bands are originated from the presence of aromatic rings and flavonoids, which are the main responsible agents in bactericidal and antioxidant properties of propolis\(^\text{52}\).

3.2. Elemental quantitative analysis (RBS)

RBS analysis on NRL-Ag film gave an overall insight on elemental composition. The obtained spectrum shows a peak that refers to carbon (mostly present in cis-1,4-polyisoprene), besides nitrogen and oxygen, which are related to proteins and humidity (Figure 3). This is in full consonance with ATR-FTIR spectrum, previously displayed.
Moreover, a narrow and intense peak attributed to silver was observed, indicating that this element is present at the film surface. This appearance is also an indication that silver is in the nanoparticulate form, not forming a film, where the peak would appear as a plateau, as shown in the literature. The areal amount of deposited silver calculated through the spectrum was $1.6 \times 10^{16}$ atoms/cm$^2$ or 2.87 μg/cm$^2$. The literature supports that different products consisting in wound dressings can contain silver nanoparticles concentrations between 1.0 μg/cm$^2$ and 1.34 mg/cm$^2$. This way, the silver concentration found in this work would fit into the lower limit within the range of these traditional products, reducing the risk of cytotoxicity to human cells.

Another peak circled in blue (around 1300 keV) could be related to elements such as calcium and potassium, both of which are absorbed by the soil, taking part in the biosynthesis of natural latex.

### 3.3 Water contact angle (WCA)

The contact angle measurements for NRL, NRL-Pla, NRL-Ag, NRL-RP and NRL-RP-Ag films are represented in Table 4.

![Figure 3. RBS spectrum of NRL-Ag film.](image)

Silver-containing films (NRL-Ag and NRL-RP-Ag) also provided a decrease in the contact angle, falling in hydrophilic range as expected, once incorporation of AgNPs is responsible for rising the wettability in a matrix, besides providing the bactericidal effect. RP, hydrophilic itself, is also an enhancer of wettability when incorporated. This result indicates that NRL-Ag, NRL-RP, NRL-RP-Ag and NRL-Pla films could be used as dressings, since hydrophilicity is correlated with improvement in both scarring and tissue regeneration.

Furthermore, it is important to assess shelf life, in terms of wettability of the samples, as it will dictate for how long they retain a favorable condition for biomedical application, i.e., the maintenance of their hydrophilicity. There is a tendency in contact angle increasing as function of time. This happens due to the aging effect, which depends on the temperature, polymer type and storage humidity. Opposed to polymers predominantly crystalline, which have a more rigid structure and have a lower aging rate, amorphous polymers have greater mobility in their polymeric chains and are more likely to aging effect. After some time of storage, hydrophilicity of the plasma treated films is reduced due to the reorientation of functional groups formed on the surface toward bulk material and the diffusion of these groups on the polymeric matrix.

The samples containing silver also present a decrease in the contact angle. This phenomenon is expected because the AgNPs tend to form aggregates, promoting roughness effects that affect the wettability. To investigate this effect, contact angles were measured throughout five months. The results are found in Table 5.

This assessment highlights the period that the samples will retain their hydrophilic character for 4 to 5 months. NRL-RP-Ag samples should be preferably used by the end of the final month after their manufacturing. On the other hand, NRL-RP did not present significant changes in contact angle along the evaluated time, indicating it could be used over a longer period.

### 3.4. Evaluation of the efficiency of the plasma sterilization process

Although simple, the thioglycolate broth test can serve as a support for the sterility control of several materials. As already explained, to evaluate the effectiveness of the sterilization process of plasma treatment and silver deposition by magnetron sputtering in a vacuum chamber, the films were immersed in medium of thioglycolate broth for 14 days. After this period, the appearance or not of turbidity in the thioglycolate broth...
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was evaluated. In the NRL films with no plasma treatment or silver deposition (NRL and NRL-RP), the medium was cloudy (Figure 4a), evidencing bacterial contamination. In this case, the samples would require a further sterilization process, such as ethylene oxide. Plasma treatment or silver deposition, on the other hand, were effective to eliminate microorganisms (Figure 4b), where no turbidity of the broth was seen (NRL-Pla, NRL-RP-Pla, NRL-Ag and NRL-RP-Ag).

Many of the usual sterilization processes may cause modifications in the films chemical structure, interfering directly in their properties. High temperature, as seen in autoclave, may degrade the biomaterial, while the ethylene oxide alters surface chemistry, allowing the formation of toxic residues and undesirable byproducts.

Due to their flexibility regarding material size and shape, plasma-based treatments are gaining more visibility. Cold plasma method is efficient in sterilizing biomedical devices and offers several advantages, one of them is a clean process happening in a wide range of temperatures, including room temperature. The biocide effect is possible due to the formation of reactive species that attack nucleophilic sites in the microorganisms, inactivating them. Therefore, the treatments applied on the films can be effective as a sterilization process, minimizing costs and time in industrial processes.

3.5. Cell culture assays

3.5.1. Isolation of adipose tissue-derived stem cells

The adipose tissue was obtained from a liposuction procedure and submitted to enzymatic digestion for the purpose of isolating stem cells. After 4 days, the cultures were monitored in an inverted microscope, through which a great amount of adhered cells in expansion were observed, corroborating the findings of Dominic et al., showing an adhesion to plastic and expansion when in culture.

In addition, characterization through flow cytometry analysis, which garnered evidence that ADSCs exhibited little or no expression for all hematopoietic cells markers were performed. It is also expected that cells do not express in vitro markers that have the ability to differentiate into osteoblasts, adipocytes and chondroblasts, a fact also evidenced in the analysis (Table 6). Mesenchymal stem cells derived from adipose tissue were also assessed regarding its differentiation capacity in various types of mesenchymal lineages. All induced ADSCs samples presented the ability to differentiate into osteogenic, adipogenic and chondrogenic tissues.

ADSCs are a source of mesenchymal stromal cells and, when used in grafting, have released factors that stimulate angiogenesis, accelerating the wound healing process. Lotfi et al. showed that autologous ADSCs transplantation, increases vessel density, type I collagen and growth factor secretion, healing the wound. The application of ADSCs associated to biocompatible materials has shown favorable results in injured places, being suggested as a possible therapy for the treatment of skin wounds.

3.2.2. Indirect MTT cytotoxic assays

To evaluate the cytotoxic effect of the films, an extract was initially prepared referring to each condition according to the described methodology of standards ISO 10933-5 and ISO 10993-12, and these were put into contact with...
L929 cell line recommended by international standards and ADSCs. MTT assay results are represented in Figure 5.

According to ISO 10993-5, a material triggers cytotoxic effects when a reduction in cell viability is shown to be superior to 30% after 24 h of exposition. Comparing cell viability for these films with the international standards classification for medical devices, the assays revealed that none of the samples present cytotoxic effect for L929 and ADSCs (Figure 5a and 5b).

After 1 and 2 days of incubation, it was verified that the viability of the extracts was superior in comparison to 1 day (69.79%) and 2 days (59.29%) positive control for L929 cells, and 1 day (79.08%) and 2 days (51.49%) for ADSCs, showing statistical differences related to exposure time in both cells. Our findings concur well with the findings of Watthanaphanit and Rujiravanit in cell viability assay with L929 cells, which demonstrated that the percentage of cell viability for all samples and all concentrations of the extract with NRL was higher than 70%, thus ensuring its applicability accordingly to safety standards.

Another aspect than can be emphasized in the cell cytotoxicity assay is the assessment taking into account the exposure time of L929 and ADSCs for 7 days; a greater reduction in viability was observed for films: (1) NRL (83.58% and 72.21%); (2) NRL-Pla (76.34% and 78.58%); (3) NRL-RP (75.77% and 83.72%); (4) NRL-Ag (75.51% and 84.50%); (5) NRL-RP-Ag (83.87% and 79.84%). On the seventh day, for both L929 and ADSC cells, we were able to observe a static similarity; however, according to the indirect cytotoxicity assay, no statistical difference was observed between the treatments using primary and cell lineage culture.

Films not showing cytotoxic effect even after a week of exposition in the MTT assay, according to ISO 10993-5, with viability superior to positive control (42.26% and 27.61%). According to Pichayakorn et al., studying NRL with peel-off masks, it was observed that they can be applied to pharmaceutical products and cosmetic formulations, as they do not present toxic effect and irritability when used in humans. Zancanela et al. showed that NRL films with incorporation of RP presented antifungal activity against Candida albicans and potentially act on wound healing process. In another in vitro study, the author adds that the incorporation of RP into NRL films did not demonstrate cytotoxic activity in rat fibroblast cells (3T3), by the contrary, it maintained its mechanical characteristics and could act as a physical barrier against the penetration of infectious agents.

For silver containing membranes (NRL-Ag and NRL-RP-Ag), the cell viability was above the minimum threshold of 70%, indicating no toxic behavior. The amount of silver for the produced membranes (previously discussed in the elemental quantitative analysis section) Table 5. Cell surface markers observed through flow cytometry analysis used for the identification of stem cells derived from adipose tissue and its percentage.

| Cell surface markers | Result (%) |
|----------------------|------------|
| **Positive**         |            |
| CD29                 | 98.7 ± 1.06|
| CD44                 | 92.7 ± 2.65|
| CD73                 | 98.3 ± 0.96|
| CD90                 | 98.7 ± 1.39|
| CD105                | 97.1 ± 2.17|
| **Negative**         |            |
| CD14                 | 0.4 ± 0.61 |
| CD34                 | 0.1 ± 0.12 |
| CD45                 | 0.1 ± 0.06 |
| CD184                | 0.1 ± 0.10 |
| HLA-DR               | 0.5 ± 0.87 |
| STRO1                | 2.2 ± 1.99 |

Figure 5. Cytotoxic effect obtained by the MTT indirect test according to ISO 10993-5-2009 and ISO 10993-12-2004 of the extracts compared to the negative (DMEM medium, 10% of FBS and 1% of P/S and 5% of DMSO) and positive control (DMEM medium, 10% of SFB, 1% of P/S and 5% of DMF) on the viability of (a) L929 and ADSCs (b), treated for 1, 2 and 7 days. NegCont: negative control; PosCont: positive control; NRL, NRL-Pla, NRL-RP, NRL-RP-Pla, NRL-Ag, NRL-RP-Ag films. * Letters (a, b, c, d, e, f, g, h and i) correspond to the statistically significant differences using the ANOVA-Tukey test (p ≤ 0.05).
is not sufficient to generate toxic effects on the cell lines, especially once they are on the low range of silver concentration for wound dressing applications.

Thus, it was observed that standard L929 cells did not differ from the ADSC isolated in this study. The prospects of performing autologous ADSC grafts in burns associated with the topical use of a potent antimicrobial, anti-inflammatory and regenerating agent based on RP extracts could be advantageous with occlusive dressings with natural latex rubber.

3.2.3. Agar overlay assay

Neutral red dye has been used as an indicator of qualitative cytotoxicity in primary cultures and other cell lineages. It is characterized by accumulating lysosomes in the viable cells that incorporated this vital dye, slightly cationic. This reagent penetrates cell membrane by non-ionic passive diffusion and concentrate on cells lysosomes, where it fixates through electrostatic bonding to anionic groups, including hydrophobic phosphates of the lysosomal matrix. This assay measures a direct contact and evaluates cytotoxicity simulating material in direct skin contact over a long period.

No evidence of cytotoxic-related halo formation surrounding the films was observed. Conversely, a microscopic assay showed some cell alteration, such as damage to cell morphology and neutral red dye distaining, indicating possible cell lysis. Such findings were accentuated in L929 cell lineage. However, the findings for L929 and ADSCs were classified under moderate cytotoxicity to agar overlay assay, due to the fact that they presented a distinct zone of 1.0 cm from the sample. It is worth to stress that although the ADSCs presented a smaller discoloration zone than that of L929, a decrease in cell intensity in red stain for viable cells was observed in comparison to its counterpart positive control (Figures 6 and 7).

3.2.4. Determination of adhesion, topographical distribution and morphological patterns of L929 and ADSCs deposited on the films

Adhesion, topographical distribution and morphology of both L929 and ADSCs deposited over the aforementioned films were evaluated. It could be observed that while the first and second days showed similar cell proliferation, in the seventh day NRL-RP, NRL-Ag and NRL-RP-Ag presented a small increase over L929 cell adhesion and proliferation (Figures 8 and 9).

In addition, analogies between obtained results in the contact angle measurements were stated: a reduction in water contact angle is inversely proportional to cell adhesion and spreading over the material surface. Similar results were found by Tong et al., in which the behavior of 3T3 mouse fibroblast adhesion on diamond films and as a drawn conclusion, adhesion and proliferation for these cells is preferentially favored over hydrophilic surfaces, due to the favoring of protein adsorption, such as fibronectin and vitronectin that have a fundamental role in cell adhesion.

For Liu et al., the wettability parameter is also one of the most important when it comes to cell response on materials. It is possible that the cells have a preference for hydrophilic surfaces. However, in ATR-FTIR analyses, functional groups with nonpolar characteristics were observed, thus the presence of nonpolar functional groups could plausibly explain the little cell adsorption and their morphological aspect, round-shaped and retracted cytoplasm, not expressing their fibroblastic characteristics.

The influence of roughness and porosity is equally important. In this study, surfaces of the films containing RP, treated with plasma and silver presented visually perceived rough surfaces and more hydrophilic surface, which may be favorable to cell adhesion. The possibilities to modify the hydrophilic-hydrophobic characteristics make NRL a candidate for tissue engineering.

It should be mentioned, however, that cell adhesion is a complex process, affected by different factors, including particular cellular metabolism, surface properties of the host biomaterial and biological environment. Regarding a biomaterial, roughness, hydrophobicity, surface energy, chemical composition and porosity are crucial factors when expecting a certain cell response.

3.2.5. Elemental analysis through energy dispersive spectroscopy (EDS)

Literature points out that NRL constitutes the cytoplasm of the lactiferous cells of the vegetal tissue on the tree cambium region. Nonetheless, the substances present in latex distribute themselves into three phases (mass/latex mass): (a) rubber particles (35%): elastomer, phospholipids and polymerase enzymes; (b) aqueous phase (55%): water, proteins and electrolytes; and (c) lutoid phase (10%): proteins, lipids and carotenes.

EDS analyses for NRL-RP film revealed the presence of the following elements: point (1) oxygen, gold and potassium; point (2) oxygen, gold, potassium, calcium and phosphor; and point (3) oxygen and gold (Figure 10a). In addition, NRL film showed the presence of the following chemical elements: point (4) calcium, oxygen and gold; point (5) calcium, oxygen, phosphor, potassium and sulphur; and point (6) calcium, oxygen and gold (Figure 10b). These results confirm several deposited particles, especially calcium, phosphorus and potassium, which are well-known functional materials for tissue regeneration. Gold, on other hand, is frequently used for recovery of samples prior to the analysis on the microscope.

Therefore, d’Auzac also pointed that the aqueous medium of latex is charged with mineral salts, such as K+, Na+, Mg2+ e Ca2+. No nitrogenous groups resulted from the ammonia-based latex obtaining process were verified. This finding is important, once ammonia is known for its volatility and irritability for both respiratory tracts and skin pathways.
Figure 6. Overlay assay with L929 cells. Visualization on inverted microscope (magnification of 40 ×). Measurements of discoloration zones (DZ) carried out on ImageJ, with scale calibration (0.1 cm).

Figure 7. Overlay assay with ADSCs. Visualization on inverted microscope (magnification of 40 ×). Measurements of discoloration zones (DZ) carried out on ImageJ, with scale calibration (0.1 cm).
Figure 8. FEG-SEM micrographs (500 ×). NRL; NRL-Pla; NRL-RP; NRL-RP-Pla; NRL-Ag; NRL-RP-Ag after 1, 2 and 7 days of L929 cellular incubation.
Figure 9. FEG-SEM micrographs (500 ×). NRL; NRL-Pla; NRL-RP; NRL-RP-Pla; NRL-Ag; NRL-RP-Ag after 1, 2 and 7 days ADSCs cellular incubation.
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4. Conclusions

From the carried-out analyses, it is possible to conclude that plasma treatment associated with silver deposition and RP incorporation on the surfaces of the NRL films modify their wettability, providing hydrophilic features, which in turn make them more suitable for biomedical application. Also, it was confirmed that these films could be stored and used up to several months after production, as hydrophobicity becomes a dominant feature on their surfaces over time.

Furthermore, the ATR-FTIR spectra showed that RP was physically incorporated into the NRL matrix, being a favorable condition to its release into the wound environment. The amount of deposited silver was calculated through RBS spectrum, indicating also that the metal is possibly found in the form of nanoparticles, and not in a film structure.

The results also allowed the association of factors like contact angle reduction and cellular dispersion over the films. It is possible that cells prefer hydrophilic surfaces, however, the presence of nonpolar functional groups most likely explains the little adsorption and the morphological aspect of the cells, which lost their fibroblast morphological characteristics. In this context, it was verified that the RP-containing samples and those that received plasma treatment and silver deposition presented greater cell adherence. In addition, these processes were effective in eliminating microorganisms, thus sterilizing the samples.

The immortalized L929 cell line and ADSCs collected from patients showed similar cytotoxicity patterns. Overall, the direct cell viability assays pointed out to moderate cytotoxicity and the indirect assays presented a cell viability percentage for all films and all extract concentrations above 70%, demonstrating international safety standards.

The properties of the in vitro studies materials were satisfactory, once their aimed application is related to a wound dressing, avoiding cellular adhesion, i.e., not promoting cellular tear off from the affected site. Besides, it is desired that the films present hydrophilic features to drain excess exudate formed from the wound, while not being toxic. However, it must be highlighted that these findings instigate further research so that the films can be applied as protective and regenerating occlusive skin dressings.

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Figure 10. Elemental analysis through energy dispersive spectroscopy (EDS) micrographs: (a) NRL-RP and (b) NRL.
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