Original article

Novel Clotrimazole and Vitis vinifera loaded chitosan nanoparticles: Antifungal and wound healing efficiencies

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A B S T R A C T

Chitosan integrated nanoparticles of clotrimazole and Egyptian Vitis vinifera juice extract was evaluated in order to maximize the antifungal activity and reduce the gross side effects. In the present study Egyptian Thompson Seedless Vitis vinifera and Clotrimazole (Cz) loaded chitosan nanoparticles (NCs/VJ/Cz) showed a promising antifungal effect with average inhibition zone diameters of 74 and 72 mm against Candida albicans and Aspergillus niger respectively. NCs/VJ/Cz ointment was examined by experimental rats with wounded skin fungal infection. Data proved the ability of NCs/VJ/Cz to gradually release the drugs in a sustained manner with complete wound healing after 7 days administration. As a conclusion NCs/VJ/Cz ointment can be used as a novel anti-dermatophytic agent with high wound healing capacity.

1. Introduction

Dermatophytosis are keratinophylic fungi that affect skin, hair, and nails superficially. Cryptococcus neoformans, Candida and Aspergillus species have been found to be the most common fungi causing systemic mycosis which was considered as a cause of high morbidity and mortality rates (Fuentefria et al., 2018). In order to overcome dematophytic infection, imidazoles (e.g., bifonazole, clotrimazole, miconazole) were the most commonly used antifungal drugs for the treatment of superficial cutaneous mycoses infections. Precautionary measures must be taken in the application of all fungal drugs with fungistatic mechanism of action due to high toxicity, insufficient bioavailability and the development of resistance by fungal pathogens or emerging innate resistance among fungal species (Cavaleiro et al., 2015). Search for a novel antifungal drug will be necessary for the future, natural products are important source for new therapeutic agents of different strategies that can be applied to improve the antifungal drugs.

Grapes (Vitis vinifera) are cultivated especially on the Mediterranean area (Falah-Tafti, 2010). Vitis vinifera juice extract facilitates and accelerate skin wound healing with antioxidant, antimicrobial, antiviral and anti-cancer activities (Nassiri-Asl & Hosseinzadeh, 2009). On the other hand, Chitosan is a polysaccharide (deacetylated from chitin) which has unique properties offering many industrial and biomedical applications. Chitosan nanoparticles may increase the solubility and enhance the bioavailability of different active agents, with the ability of modulating a drug targeted nano-system (Leonida et al., 2018). Chitosan nanoparticles are among biodegradable polymers which were studied extensively as delivery systems with controlled release of active ingredients (Divya & Jisha, 2018). Chitosan nanoparticles have the ability to cross different biological barriers and deliver the drug in a controlled release manner to the target site, improve the drugs bioavailability, drugs pharmacokinetics modification and to protect the loaded drugs (Divya & Jisha, 2018). The small nanoparticles size also enhance the interfacial interaction with the target cell membrane caused by the endocytosis effect (Ghadi et al., 2014; Reina et al., 2019). These properties favored the chitosan nanoparticles with several clinical potentialities to concur the
increasing incidence of microbial resistance (Landriscina et al., 2015).

The present study aimed to synthesize a novel drug derived from the combination between clotrimazole and Thompson seedless Vitis vinifera juice extract loaded chitosan nanoparticles against infectious skin mycosis.

2. Material and methods

2.1. Vitis vinifera (Grape) samples

Vitis vinifera samples namely: Thompson (VJ) and Flame seedless (FJ) Vitis vinifera were collected during September 2019, from Alexandria, Egypt. The samples were cleaned and rinsed with sterile water then air dried.

2.2. Microorganisms

Some dermatophytes namely: Aspergillus niger, Aspergillus flavus, Fusarium oxysporum, Candida albicans, Mucor hiemalis and a standard strain Candida albicans ATCC 14,053 were kindly identified and provided by Assiut University Mycological Center (AUMC) (Mousafa, 2006).

2.3. Seed culture preparations

Spore suspension and yeast seed culture were prepared according to CLSI guidelines (CLSI, 2017).

2.4. Preparation of the Vitis vinifera juice extract

Vitis vinifera samples were macerated and extracted using an alcoholic solvent (water/ethanol: 20/80 v/v) at 60 °C for 1 hr then filtered through a Whatman filter paper No. 1 (Katalinic et al., 2013).

2.5. Antifungal activity of Vitis vinifera juice extract

Antifungal activity of all the prepared extracts was evaluated by disc-diffusion method using Potato dextrose agar (PDA) plates (Oliveira et al., 2013).

2.6. Combination between grape juice extracts and the commonly used antifungals

Clotrimazole (Cz), and Fluconazole (Flz) discs were loaded with 25 μl of Vitis vinifera juice extracts one at a time and was applied on the surface of the PDA fungal inoculated plates (Nirmala & Narendhirakannan, 2011). Further evaluation of the combination efficacy was done using minimum inhibitory concentration (MIC), minimum fungicidal concentration (MFC) and checkerboard dilution technique (Foucquier & Guedj, 2015).

2.7. Chemical analysis of Thompson Seedless Vitis vinifera juice extract (VJ)

2.7.1. GC–MS analysis

Thompson Seedless Vitis vinifera juice extract (VJ) was subjected to GC–MS analysis using capillary column SBP5 (30 m × 0.32 mm × 0.25 μm) installed in a Hewlett-Packard 5890 Series II instrument (Nirmala & Narendhirakannan, 2011). GC–MS analysis was done at National Institute of Oceanography and Fisheries, Alexandria, Egypt.

2.8. Nanoparticles preparation

2.8.1. Nano-chitosan preparation (NCs)

Nano-chitosan (NCs) was prepared using ionic gelation method according to Elnaggar et al. (2020) using sodium TPP solution (0.2% w/v in deionized water).

2.8.2. Nano-chitosan /Thompson Seedless Vitis vinifera juice extract (NCs/VJ)

NCs encapsulated with VJ (20 mg) were prepared using incorporation method according to Ong et al. (2017).

2.8.3. Nano-chitosan/Clotrimazole (NCs/Cz)

NCs encapsulated with Cz (10 mg) was prepared using incorporation method according to Koukaras et al. (2012).

2.8.4. Nanochitosan /Thompson Seedless Vitis vinifera /Clotrimazole (NCs/VJ/Cz)

NCs encapsulated with Cz (10 mg) and VJ (20 mg) were prepared by mixing Cz and VJ with the prepared TPP solution under stirring. The mixture was added drop-wise into two different chitosan concentrations (0.25 and 0.5%) one at a time to reach higher loading efficiency. The prepared nanoparticles were stored at 4°C in sterile Falcon tubes after centrifugation and lyophilization for further investigations.

2.9. Antifungal activity of the prepared nanoformulae

The prepared Nano-formulae (NCs, NCs/VJ, NCs/Cz and NCs/VJ/Cz) were evaluated using disc-diffusion method; MIC, and MFC (Nina et al., 2015). Candida albicans treated cells with the most promising formula were examined using transmission electron microscopy (TEM) using a Transmission Electron Microscope (JEM-100 CX Joel) at the Electron Microscope Unit, Faculty of Science, Alexandria University, Egypt. While A. niger treated hyphae was examined using light microscope (Ribes et al., 2017).

2.10. Test for antioxidant (DPPH scavenging) activity

The DPPH (2,2- diphenyl-1-picryl-hydrazyl) scavenging effect was calculated using the following equation:

\[
\text{Scavenging percentage} = \frac{\text{ADPPH} - \text{AS}}{\text{ADPPH}} \times 100
\]

where AS is the absorbance of the solution when the sample under test has been added at a particular concentration and ADPPH is the absorbance of the DPPH solution (Xie et al., 2015).

2.11. Characterization of the most promising nanoformula

The ultrastructure, size and shape of the prepared nanoformulae were examined using TEM (JEM-100 CX Joel) (Vinodhini et al., 2017). FTIR (FTIR-BRUKER, wave number ranged from 4000 cm\(^{-1}\) to 450 cm\(^{-1}\)) was used to investigate the nanoparticles functional groups and X-Ray diffraction (XRD-BRUKER AXS, Cu Kα radiation source (λ = 0.154 nm), at scan rate = 10/ min) of the most promising nanoformula were analyzed (Elnaggar et al., 2020). However, the particle size (PS), polydispersity index (PDI) and Zeta potential of the most promising Nano-formula was determined by dynamic light scattering (DLS) technique (Malvern Zetasizer) (Elnaggar et al., 2020).

The entrapment efficiency of the most promising nanoformula was investigated according to El-Refaie et al. (2015). The entrapment efficiency percentage (EE%) was calculated using the following equation:
Finally, the release of Clotrimazole and Thompson seedless Vitis vinifera juice extract were determined using dialysis bag (2.5 cm in diameter) at different time intervals (0.5, 1, 2, 3, 4, 5 and 6 hrs) according to El-Refaie et al. (2015). Cz and VJ release% were calculated and drug concentration was measured with UV absorbance at 237 and 280 nm respectively (Bonazzi et al., 1998; Xiao et al., 2018).

2.12. Ointment preparation

Four grams of gelatin powder was dissolved in 100 ml distilled water at 40 °C for 30 min. After complete homogenization, the gelatin solution was incorporated with 500 mg/ml of each VJ loaded chitosan nanoparticles, Cz loaded chitosan nanoparticles and the most potent formula (NCs/VJ/Cz) one at a time and mixed with constant stirring for 1 h at 4 °C (Jridi et al., 2017).

2.13. Formula evaluation using animal model

2.13.1. Ex vivo permeation studies

2.13.1.1. Preparation of skin samples. Ethics Committee at the Faculty of Science, Alexandria University, Egypt has approved the present experiment (ALEXU IACUC Institutional Animal Care and Use Committee on research (AU04190413302). The animals were kept under standard laboratory conditions and veterinary supervision with access to water and food ad libitum. Thiopental solution (40 mg/kg) was injected intraperitoneally for rats anesthetization (IACUC Faculty and Staff, 2016). Shaving razor was used for the removal of the rats’ dorsal hair carefully (Butani et al., 2014). The hypodermis part of the shaved skin was removed with scalpel (Aggarwal, 2006) which was then washed with saline and stored in refrigerator (−20 °C) until use (Sahoo et al., 2014).

2.13.1.2. Skin permeation studies. Franz diffusion cells (surface of 3.14 cm² and volume = 9 ml) were used in the ex vivo release studies of Egyptian Thompson seedless Vitis vinifera (VJ) and Clotrimazole (Cz) from the prepared ointment (NCs/VJ/Cz ointment). 9 ml of sodium phosphate buffer (SPB) of pH 5.5 was poured into the Franz diffusion cells’ receptor compartment at 32 °C ± 0.5 °C (Kumar et al., 2014). The excised skin was placed between the receptor compartment and the donor of the Franz diffusion cell (Song et al., 2014). The ointment (0.5 g) was applied uniformly on the excised skin. Then at different time intervals (2, 4, 6, 8, 10, 12, and 24 h) a 2 ml aliquot was withdrawn (replaced with freshly prepared SPB medium) and filtered. Each aliquot was analyzed using UV–vis spectrophotometer at 237 and 280 nm respectively (Bonazzi et al., 1998; Xiao et al., 2018) to calculate the percentage of cumulative Cz and VJ drug permeated through the rats’ excised skin.

2.13.1.3. Skin retention study. After ending the permeation studies the excised skin was separated and washed with methanol then the excised skin was vortexed in SPB. For effective extraction of the retained VJ and Cz, each skin sample was immersed in methanol for 24 hrs under shaking condition (100 rpm at 37 °C) (Song et al., 2012; Butani et al., 2014). Each sample was then sonicated and centrifuged (8000 rpm) for 20 min (Verma et al., 2014). The supernatant was analyzed with a UV method at 237 and 280 nm respectively (Bonazzi et al., 1998; Xiao et al., 2018) and the placebo nanoparticles was used as a blank (Ge et al., 2014). The percentage of Cz and VJ deposition were calculated and represented as average ± SD.

2.13.2. In vivo studies

2.13.2.1. In vivo skin retention study. In vivo skin retention study was carried out according to Aggarwal & Goindi (2013). Male rats (Rattus norvegicus albinus) weighing 195 ± 30 g were anesthetized using intraperitoneal injection using 10 mg/kg xylazine and 80–100 mg/kg hydrochloride (Gupta & Vyas, 2012; Altuntas et al., 2014). Shaving razor was used to shave the rats’ dorsal hair carefully (Butani et al., 2014). Half (0.5) g of the prepared ointment of NCs/VJ/Cz was applied on the shaved area. The animals were sacrificed after 5 min, then after 2, 4, 6, and 24 hrs. The shaved dorsal skin was excised and washed with methanol. Skin preparation was done as described in the ex vivo drug retention study to assess the deposition of VJ and Cz using UV–vis spectrophotometer at 237 and 280 nm respectively (Bonazzi et al., 1998; Xiao et al., 2018).

2.13.2.2. In vivo antifungal activity of NCs/VJ/Cz ointment. In vivo antifungal activity study of NCs/VJ/Cz ointment was done following Gupta and Vyas (2012).

2.13.2.2.1. Animal preparation. Fifty-five male albino rats (Rattus norvegicus albinus), 4 months old with average body weight of 180 ± 30 g were assigned to eleven groups each including 5 rats and kept in a cage under conventional conditions of temperature and humidity in separate cages in average temperature (25 ± 2 °C) inside an adequately ventilated room (Elsheikh et al., 2012). Dorsal inter-scapular region of each albino rat was disinfected with ethanol (70%) and shaved one day prior to infection. All experiment was done under aseptic conditions. A circular wound was created by excising the skin carefully with a 15-mm biopsy punch; circular wounded areas were left open until the end of the experiment (Tramontina, 2002). Aliquots of 1x10⁸-10⁹ A. niger spore suspension was injected intradermally in the wounded area (Qushawy et al., 2018).

However, treatment regimen began 72 hrs post-fungal infection where test formulations were topicaly applied once daily (0.5 g/day) for three consecutive days. The rats were divided into eleven groups according to their treatment regimen as follows:

- **Group I**: Assigned as negative control with neither fungal infection nor treatment regimen.
- **Group II**: Assigned as positive control, infected with A. niger strain with no treatment regimen.
- **Group III**: A. niger Infected rats treated with gelatin solution only.
- **Group IV**: A. niger Infected rats treated with NCs placebo only.
- **Group V**: A. niger Infected rats treated with Cz only.
- **Group VI**: A. niger Infected rats treated with VJ only.
- **Group VII**: A. niger Infected rats treated with the combination of Cz and VJ.
- **Group VIII**: A. niger Infected rats treated with ointment I (NCs/VJ: gelatin (1:1 w/v)).
- **Group IX**: A. niger Infected rats treated with ointment II (NCs/Cz: gelatin (1:1 w/v)).
- **Group X**: A. niger Infected rats treated with ointment III (NCs/VJ/Cz: gelatin (1:1 w/v)).
- **Group XI**: Assigned as the treatment control (ointment III) with no infection (toxicity assessment).

Rats were checked twice daily during the experimental period to ensure no adverse reactions were observed. This is the first work to emphasis the effect of NCs/VJ/Cz conjugated nanoparticles as a treatment for wounded skin fungal infection in rats.
2.13.2.2.2. Histological studies. The excised healed skin tissue of each rat group was fixed in 10% formalin to allow good fixative penetration for histological examination. Sections (5 μm thick) were stained with Ehrlich’s hematoxylin and counterstained with eosin (H&E). The selected sections were examined with light microscope and photographed (Arizmendi et al., 2014).

2.14. Hemocompatibility test
Estimation of hemocompatibility of the most promising formula was performed through Hemolysis studies using fresh human blood, collected in EDTA tube and then diluted with normal saline solution (2 ml blood + 2.5 ml normal saline) according to Basak et al. (2018). The hemolysis percentage was calculated according to following equation:

\[
\text{%Hemolysis} = \frac{OD_{test} - OD_{negative}}{OD_{positive} - OD_{negative}} \times 100
\]

According to Basak et al. (2018), if the hemolysis % was less than 5% then the tested drug was considered highly hemocompatible. If the hemolysis % was within 10 , the tested drug was considered hemocompatible. On the other hand, if the hemolysis %was more than 20, the drug was reported as non hemocompatible.

2.15. Statistical analyses
All experiments were performed in triplicates and results were expressed as Mean ± Standard Deviation. Statistical analysis was performed by SPSS 16.0.

3. Results

3.1. Antifungal activity of Egyptian grape juice extract

Data in Table 1 showed that Flame Seedless *Vitis vinifera* juice extract (FJ) reported higher antifungal activity than Clotrimazole (Cz) and Thompson Seedless *Vitis vinifera* juice extract (VJ). It was revealed that *A. niger* and *Candida albicans* were the most resistant strains against the tested agents with inhibition zone diameter of 12.12 and 10.37 mm respectively.

3.2. Combination between *Vitis vinifera* juice extract and some antifungal agents

Data in Table 2 revealed that the antifungal effect of Cz/VJ showed significantly higher antifungal activity (Inhibition Zone diameter: 61.16 mm) than other combinations against the tested dermatophytes. Data revealed that Flame Seedless *Vitis vinifera* juice extract showed antagonistic effect with the tested antifungal agents while Thompson Seedless *Vitis vinifera* juice extract (VJ) showed a promising synergistic effect. VJ was selected for further studies. The combined antifungal activity of VJ and Cz were further evaluated by the checkerboard dilution technique. It was revealed that VJ/Cz had a synergistic effect with FICI equaled 0.5 against both *A. niger* and *Candida albicans*.

3.3. GC/MS analysis of Thompson Seedless *Vitis vinifera* juice extract

Thompson seedless *Vitis vinifera* juice extract was prepared and analyzed using GC–MS (Fig. 1). hydroxymethyl furfural, 5,3-Hydroxy-2,3-dihydromaltol, and ethyl-2-hydroxymethylimidazolide (7, 13 and 80% respectively) had been identified on referring to the corresponding acquisition time. Other compounds were detected in lower concentrations (e.g. eugenol 4% and α-terpineol 0.01).

3.4. The antifungal effect of the synthesized nanoparticles

Data in Table 3 revealed that NCs (0.25%)/VJ/Cz was the most promising formula with inhibition zone diameter of 72 and 74 mm against *A. niger* and *Candida albicans* respectively and therefore it was used for further studies. Other tested chitosan concentrations gave inferior results.

3.5. Determination of MIC, MFC and MIC index of the most promising formula

Table 4 revealed that the MIC that led to the maximum inhibition was reported with NCs/VJ/Cz nanoformula (2 μg/ml). According to the MIC index values, it was revealed that VJ alone had a Fungistic effect but after the formula incorporation the effect was converted to Fungicidal effect due to the interaction between VJ, Cz and NCs.

3.6. Microscopic examination of fungal treated cells

It was observed that the formula was precipitated and adsorbed on the cell surface leading to cell deformation followed by cell membrane disruption, releasing the cellular contents and became as a ghost cell (Fig. 2a and b). On the other hand, Light microscopic examination of *A. niger* treated cells showed deformed filamentous forming ghost shape (Fig. 2c and d).

3.7. Antioxidant activity of the prepared formula

Data in Table 5 revealed that the DPPH activity% of NCs/VJ/Cz was 97.5% with IC50 2.7 mg/ml.

3.8. Characterization of the synthesized nanoparticles

3.8.1. Transmission electron microscopic examinations
TEM images (Fig. 3) illustrated that the prepared NCs had a vesicle size between 216 and 263 nm, upon the addition of VJ the vesicle size was 59–124 nm. On the other hand, NCs/Cz showed relatively low vesicle size (67–75 nm) while NCs/VJ/Cz had a vesicle size between 50 and 89 nm. The prepared nanoparticles had a denser color in the core.

Table 1
Antifungal activity of Egyptian Seedless *Vitis vinifera* juice extract and some some pharmaceutical products.

| Organism                  | Average inhibition zone diameter (mm) ± Standard deviation |
|---------------------------|------------------------------------------------------------|
|                           | Thompson Seedless (VJ) | Flame Seedless (FJ) | Clotrimazole (Cz) | Fluconazole (Flz) |
| *A. niger*                | 13.6 ± 0.25          | 13.6 ± 0.28        | 6.1 ± 0.02        | 15.2 ± 0.55 |
| *A. flavus*               | 15.6 ± 0.28          | 14.2 ± 0.72        | 10.0 ± 0.10       | 17.8 ± 0.67 |
| *Fusarium oxysporum*      | 17.8 ± 0.28          | 26.6 ± 0.28        | 11.0 ± 0.11       | 17.7 ± 0.71 |
| *Candida albicans* ATCC   | 14.9 ± 0.46          | 17.2 ± 0.50        | 11.0 ± 0.11       | 16.4 ± 0.68 |
| *Candida albicans*        | 12.6 ± 0.76          | 11.4 ± 0.5         | 6.0 ± 0.1         | 11.5 ± 0.67 |
| *Mucor hiemalis*          | 17.0 ± 0             | 21.0 ± 0.59        | 11.2 ± 0.1        | ± 0.66     |
3.8.2. Fourier transform infrared spectroscopic (FTIR) analysis and X-Ray diffraction

FTIR spectral details of NCs/VJ/Cz (Fig. 4a) showed absorption band obtained at 3433.9 cm⁻¹/C₀ attributed to a collective –NH and OH group resulted in a broadened band due to the physical interactions with TPP when converting chitosan to a Nano particulate form. The peak at 1730.15 cm⁻¹/C₀ was assigned to the C = O and C–O which may be explained by the functionalization of eugenol and 5,3-Hydroxy-2,3-dihydromaltol on the prepared chitosan nanoparticles. Data in Fig. 4b showed the X-Ray diffraction patterns of NCs loaded with VJ and Cz showed two peaks at 2θ = 27 indicating a shift from the pure chitosan peaks around 20 revealed the increased amorphous nature.

3.8.3. Determination of entrapment efficiency, particle size, PDI and Zeta potential

The physical characteristics of NCs/VJ/Cz formula (Table 6) revealed that the mean diameter, PDI value and Zeta potential were 35.4 nm, 0.248 and + 31 mV respectively indicating homogeneous size distribution with good stability. The entrapment efficiency of VJ/Cz into the chitosan vesicles was found to be 94.7%.

3.8.4. Drug release study

NCs/VJ/Cz was able to gradually release the drug load by time. After 2 hrs, the cumulative release of VJ from the Nano carrier was 47.9 ± 1.9% compared to 35 ± 1.5% for Cz (Fig. 5). The initial burst release might be attributed to the diffusion of Cz and VJ that were localized at or close to the surface of the prepared nanoparticles.

3.9. Formula evaluation using animal model

3.9.1. Skin permeation study

Ex vivo permeation studies of Egyptian Thompson Seedless *Vitis vinifera* (VJ) and Clotrimazole (Cz) from the loaded Chitosan nanoparticles (NCs/VJ/Cz) ointment were tested. The ex vivo release pattern of VJ and Cz was shown in Fig. 6. Ex vivo permeation of Egyptian Thompson Seedless *Vitis vinifera* (VJ) and Clotrimazole (Cz) was enhanced upon its incorporation into the ointment showing almost a 1.5-fold permeation increase.

3.8.2. Fourier transform infrared spectroscopic (FTIR) analysis and X-Ray diffraction

- **Table 2** Combination study of different Egyptian Seedless *Vitis vinifera* juice extracts with some pharmaceutical products

| Organism          | Average inhibition zone diameter (mm) ± Standard deviation |
|-------------------|----------------------------------------------------------|
|                   | A. Niger | A. Flavus | Fusarium oxysporum | Candida albicans ATCC | Candida albicans | Mucor hiemalis |
| Fluconazole/VJ    | 20.0 ± 0.10 | 30.0 ± 0.20 | 30.0 ± 0.10 | 30.0 ± 0.10 | 21.0 ± 0.10 | 25.0 ± 0.15 |
| Clotrimazole/VJ   | 15.0 ± 0.10 | 23.0 ± 0.15 | 22.0 ± 0.20 | 20.0 ± 0.10 | 16.0 ± 0.20 | 20.0 ± 0.20 |
| Fluconazole/Cz    | 29.0 ± 0.15 | 35.0 ± 0.10 | 37.0 ± 0.10 | 31.0 ± 0.10 | 25.0 ± 0.15 | 40.0 ± 0.15 |
| Clotrimazole/Cz   | 60.0 ± 0.20 | 74.0 ± 0.30 | 60.0 ± 0.20 | 77.0 ± 0.20 | 66.0 ± 0.15 | 30.0 ± 0.20 |

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3.9.2. Skin retention study
Skin retention study was performed to prove the sustained locality effect of the prepared ointment into the skin. Skin samples were harvested after ending the skin permeation study to assess the VJ and Cz deposition in the skin. VJ and Cz deposition percentage in the skin was higher than the permeated percentage. Illustration of the comparison between VJ and Cz permeated and deposited percentages was shown in Table 7.

3.9.3. In vivo studies
3.9.3.1. In vivo skin deposition study. Data represented in Table 8 revealed that the VJ and Cz exhibited rapid penetration into the skin. After 5 min application of the novel formulated ointment, 69.2 ± 0.01 and 76.9 ± 0.7 μg of VJ and Cz were deposited respectively into the skin. The novel formulated ointment afterward successfully released the drugs with sustained manner through time. It is important to mention the differences between the results of ex vivo and in vivo skin retention studies which can be explained by the blood circulation to the skin which helps in drug elimination and lowering the deposition in the skin. Hence, a hemocompatibility study was done to confirm the formula safety upon topical application.

3.10. Histological and cytotoxicity studies
It is of considerable interest that animals from experimental groups did not exhibit any case of mortality. Wound contraction showed gradual reduction of wound area in 6 days and complete healing after 7 days (Fig. 7). Examination of light micrographs of vertical skin sections of healthy wounded rats after healing showed normal histological architecture (Fig. 8a). While fungal infected wounded skin, showed that the epidermal keratinocytes not completely differentiated into layers with severe inflammation (Fig. 8b). On the other hand, the light micrograph of fungal infected wounded skin treated with the formula (NCs/VJ/Cz) ointment showed more defined feature of healing. Epidermis was completely regenerated with its derivatives (Fig. 8c). Finally, the light micrograph of the group treated with NCs/VJ/Cz ointment showed normal histological architecture which proved the formula safety with no sign of inflammation, irritation or any observed side effects (Fig. 8d). Other groups were added to supplementary file (From S1Fig to S8 Fig).

3.11. Hemocompatibility test
The limit prescribed by Brazilian law and international standards was 1% for spontaneous hemolysis. In this study, the hemolysis activity of the promising formula (NCs/VJ/Cz) was proved to be highly hemocompatible (0.3%).

4. Discussion
Dermatophytic infections were usually treated with imidazoles (e.g., bifonazole, clotrimazole, miconazole) however some precautionary measures must be taken due to their high toxicity, insufficient bioavailability and the development of resistant fungal pathogens (Cavaleiro et al., 2015). Novel antifungal drugs using
natural products has been the last resort in treating dermatophytic infections. Grapes (*Vitis vinifera*) are known to facilitate and accelerate skin wound healing with antioxidant, antimicrobial, antiviral and anti-cancer activities (Nassiri-Asl & Hosseinzadeh, 2009). In the present work, *Vitis vinifera* reported higher antifungal activity than clotrimazole (Cz). This can be explained by the high amounts of polyphenols in *Vitis vinifera* (VJ) that may play a major role in the observed antifungal effects (Fraternale et al., 2015). Houillé et al. (2014) revealed that two dimethoxy-resveratrol derivatives (3,4-dimethoxy-resveratrol and 3,5-dimethoxyresveratrol) displayed interesting antifungal activities with minimum inhibitory concentration (MIC) values of 28–37 µg/ml against *Candida* species. Consequently, the observed antifungal activity of VJ in the present study can be reverted to the presence of dimethoxy-resveratrol derivatives. Kuruc & Čonková (2017) tested the in vitro antifungal activity of an ethanolic extract of *Vitis vinifera* tendrils (TVV) and it was revealed that the MIC values was from 250 to 300 ppm. Simonetti et al. (2019) evaluated the antifungal activity of *Vitis vinifera* extracts and it was revealed that the MIC

### Table 6

| Formula used | Average size (nm) | Potential | PDI | EE% |
|--------------|-------------------|-----------|-----|-----|
| NCs/VJ/Cz    | 35.40             | +31.00    | 0.24| 94.70|

### Table 8

| Time (hrs) | Deposition (µg) |
|------------|-----------------|
|            | VJ              | Cz              |
| 0.08       | 69.2 ± 0.01     | 76.9 ± 0.70     |
| 2          | 60.7 ± 1.00     | 70.1 ± 1.50     |
| 4          | 54.1 ± 1.10     | 51.9 ± 0.30     |
| 6          | 43.9 ± 0.50     | 38.3 ± 1.00     |
| 24         | 28.4 ± 1.50     | 19.7 ± 2.40     |

### Table 7

| Formula used | VJ release percentage* | Cz release percentage* | VJ permeation percentage* | Cz permeation percentage* | VJ retention percentage** | Cz retention percentage** |
|--------------|-------------------------|------------------------|---------------------------|---------------------------|---------------------------|---------------------------|
| NCs/VJ/Cz    | 59.4 ± 0.2              | 34.8 ± 0.3             | 1.2 ± 0.4                 | 2.4 ± 0.02                | 3.0 ± 0.07                | 5.6 ± 0.1                 |
| VJ¹          | 92.2 ± 0.01             |                        | 1.2 ± 0.4                 |                          |                           |                           |
| Cz²          | 81.9 ± 0.05             |                        |                            |                           |                           |                           |

*: Dialysis bag, **: Skin, -: Not tested, ¹: VJ control and ²: Cz control
values ranged from 53.58 to 214.31 μg/ml against Candida spp. and from 43.54 to 133.02 μg/ml against dermatophytes. Kuruc & Čonková (2017) revealed that the efficacy of antimycotics can be increased by a proper combination between commercial drugs and natural substances. The combined antifungal activity of VJ and Cz were evaluated by checkerboard dilution technique which revealed a synergistic effect between VJ and Cz against both A. niger and Candida albicans. VJ/Cz loaded chitosan nanoparticles had the highest antifungal activity with inhibition zone diameter of 72 and 74 mm against A. niger and Candida albicans respectively. The antifungal mechanism of action of the prepared nano-formula was assessed by electron microscopic study which revealed the nanoparticles precipitated and adsorbed on the fungal cell surface leading to membrane disruption and releasing of the cellular contents. The antifungal activity against fungal mat was due to the diffusion of charged nanoparticles onto the cell wall of the fungal species which eventually causes cell death (Ing et al., 2012). Chitosan nanoparticles has a metal chelating and DNA binding activities that inhibit mRNA synthesis which can attribute to the observed antimicrobial activity (Divya & Jisha, 2018). Klis et al. (2007) reported higher resistance activity of A. niger strains after chitosan nanoparticles treatment which can be explained by the fact that A. niger cell wall contains 10% of chitin (the non-deacetylated form of chitosan).

In the present study, the antioxidant activity of NCs/VJ/Cz was 97.5% with IC50 reached 2.7 mg/ml. De la Cerda-Carrasco et al. (2015) tested the antioxidant activity of white Vitis vinifera juice extracts which presented higher antioxidant capacities and higher contents of total phenols and total proanthocyanidins compared with red Vitis vinifera juice extracts. Huang & Li (2014) tested the DPPH and ROS radical scavenging activity of Chitosan/fucoidian nanoparticles which showed potent antioxidant effect. Yen et al. (2008) explained that the observed antioxidant activity of chitosan was due to the iron chelating ability and hydroxyl radical scavenging activity.

The formulated NCs/VJ/Cz had a denser color in the core, which might be due to the higher phosphorous content of TPP in the cross-linked core of the nanoparticles (Abdelkader et al., 2017). FTIR spectral details of NCs/VJ/Cz (Fig. 4a) showed absorption band obtained at 3433.9 cm⁻¹ attributed to the physical interactions with TPP when converting chitosan to a Nano particulate form (Abdelkader et al., 2017). Fig. 4b showed the X-Ray diffraction patterns of NCs loaded with VJ and Cz which showed two peaks at 2θ = 27 indicating a shift from the pure chitosan peaks around 20 revealing the increased amorphous nature of chitosan after crosslinking with sodium tripolyphosphate (Abdelkader et al., 2017). Sharma and Sharma (2013) reported that terbinafine (an antifungal agent) was effectively encapsulated in chitosan nanoparticles (77.8%).

Ex vivo and in vivo evaluation of skin retention, permeation and wound repair potentiality were examined. NCs/VJ/Cz formula was able to gradually release the drug by time. The initial burst release might be attributed to the diffusion of Cz molecules and VJ that were localized at or close to the surface of the nanoparticles. The sustained release then occurred due to the diffusion of the drug through the cross-linked Cs matrix (Ing et al., 2012). The drug flux through a membrane (dialysis bag) increases as the thermodynamic activity of the drug increases (Hemmati, 2015; Abdelkader et al., 2017). The thermodynamic activity of a drug can be increased by decreasing the drug solubility in the carrier or by increasing the drug concentration (Higuchi, 1960). Nanoformulations of drugs can enhance the drug penetration into the skin through topical application (Madheswaran et al., 2013). Ex vivo permeation studies of Egyptian Thompson Seedless Vitis vinifera (VJ) and Clotrimazole (Cz) from the loaded Chitosan nanoparticles (NCs/VJ/Cz) ointment revealed the enhanced drug permeation. Its incorporation into ointment reached almost a 1.5-fold permeation increase. This can be explained on the basis of the physicochemical characteristics of the prepared Nanoformula (Madheswaran et al., 2013). Both Nano-chitosan and gelatin were known for their skin penetration effect (Lou et al., 2013). In addition, the particle size of NCs/VJ/Cz could influence the ex vivo permeation of VJ and Cz (Madheswaran et al., 2013). Whereas, the smaller particle size (less than 100 nm) means large
surface area which allow the drug to deeply penetrate the skin (Stoye et al., 1998). Similarly, Modi et al. (2013) prepared a ketoconazole loaded chitosan nanoparticle which showed potent sustained release over time due to highly absorption in the gastric mucosa.

The drug deposition was enhanced significantly due to high similarity between gelatin (gelation material of the prepared ointment) and the collagen present in the skin which allows the drug delivery and diffusion through the skin (Gosenca et al., 2013). Aggarwal & Goindi (2013) suggested that the rapid penetration of the drug has a significant effect in choosing the suitable skin-targeting formulation. Hence, in the in vivo skin deposition study through different time intervals was done. Data represented in Table 8 revealed that the Vj and Cz exhibited rapid penetration into the skin.

It is of considerable interest that animals from all the experimental groups did not exhibit any case of mortality. Wound contraction showed gradual reduction of wound area in 6 days and complete healing after 7 days (Fig. 7). Hemmati (2015) tested topical application with Thompson seedless grape juice extract in a eucerin base (2% w/w) and it was proved to be highly hemocompatible (0.3%). Brown et al. (2010) revealed that resveratrol (active constituent of Vitis vinifera seed extract) at doses of 0.5 and 1 g was completely safe and that adverse gastrointestinal effects appeared with doses of 2.5 and 5 g.

Conclusion

Chitosan nanoparticles loaded with Thompson seedless Vitis vinifera juice extract (NCs/Vj/Cz) ointment was prepared and the ex vivo and in vivo evaluation of skin retention, permeation and wound repair potentiality were examined by experimental skin fungal infected rats. Data revealed that the formula was able to gradually release the drugs in a sustained manner with tissue repairing and complete wound healing effect after 7 days. Hence, Chlormizalone combination with Thompson Seedless Vitis vinifera juice extract (Vj) loaded on chitosan nanoparticles can be used as novel anti-dermatophytic agent with potent wound healing capacity.

Financial & competing interests’ disclosure

The authors have no relevant affiliations or financial involvement with any organization or entity with a financial interest in or conflict with the subject matter or materials discussed in the manuscript. This includes employment, consultancies, honoraria, stock ownership or options, expert testimony, grants or patents received or pending, or royalties.

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Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.sjbs.2021.10.041.
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