Investigation of Galactomannan/deacetylated Chitosan Nanocomposite Films and their Anti-bacterial Properties

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Abstract

A little packaging is a wonderful thing, a lot of packaging is a nightmare, particularly when landfills around the world threaten to engulf our living space. The topic of edible packaging is still of interest to the food industry and other organization funding research to solve packaging dilemmas. In this research, galactomannan (GM) was used as raw material and deacetylated chitosan (DE-ChN) was used as strengthening modifier to prepare GM based packaging films. The chemical structure of the composite film was analyzed with SEM and FTIR. The properties influence for films of different DE-ChN content were studied. The obtained GM/DE-ChN nanocomposite films showed superior hydrophobicity and high tensile strength. The nanocomposite films against *Escherichia coli*, *Bacillus subtilis*, *Staphylococcus aureus* and *Streptococcus pneumoniae* showed great antibacterial properties. Moreover, the GM and GM/DE-ChN nanocomposite film showed no toxicity to RAW264.7 macrophage cells. The final obtained GM/DE-ChN packaging film provides a foundation for the potentials for futural plastic packaging alternatives.

Introduction

Consumers’ growing awareness of healthy lifestyles has prompted people to explore new technologies that can extend the shelf life of food without using preservatives. Due to its ability to improve the quality of foods, edible films and coatings are specially considered in food preservation. Today, traditional bio-based polymers from biomass have been gradually applied to food packaging films and coatings materials, and the commercialization of biopolymer film and coating has become more profound (Galus and Kadziński 2015). Natural substrates such as polysaccharides, proteins, and lipids can be used to make edible films and coatings (Gutiérrez et al. 2015). The changes in the mechanical properties and barrier properties of the main components in the biopolymer matrix have aroused people's interest in composite structures, which enables people to explore the complementary advantages of each component and minimize its shortcomings (Lee et al. 2004).

Polysaccharides such as starch, cellulose, chitosan, pullulan and Tara gum are some of the most normally used materials for producing edible films (Wu et al. 2012; Xiao et al. 2012). Galactomannan (GM) is a representative polysaccharide, GM derived from the *Sesbania cannabina* seed endosperm is a natural renewable polysaccharide, which can effectively reduce the use of non-renewable petroleum resources when preparing food packaging materials. At the same time, it can be used as a safe edible film to reduce consumers’ concerns about the safety of food packaging. However, the single GM film has poor mechanical properties, and is brittle, which is not conducive to its application. It is often necessary to add hydroxyl groups and polyols to improve the tensile strength, flexibility, etc. of the film, avoiding the surface breakage of the film after preparation, and making the film surface smooth (Li et al. 2019). Glycerin, polyvinyl alcohol and polysaccharides containing hydroxyl groups are commonly used plasticizers. These plasticizers are similar to polysaccharides because of their molecular structure. Hydrogen bonding will occur between shared hydroxyl groups, but different plasticizer molecules differences in structure, size, C, O element content and spatial configuration lead to different film-forming properties (Antoniou et al. 2014).
As a deacetylation product of natural polysaccharide chitin, deacetylated chitosan (DE-ChN) has many excellent physical and chemical properties, such as biocompatibility, biodegradability, non-toxicity, multi-reactivity, low irritation, broad-spectrum antibacterial properties, etc. (Verlee et al. 2017; Ye et al. 2018). There have been a large number of literatures reported that chitosan has broad-spectrum antibacterial properties, and has a good inhibitory effect on a variety of bacteria, fungi, and even some viruses. Park et al. found that the fresh-keeping time of strawberries covered with chitosan film was prolonged and the water loss was significantly reduced (Park et al. 2005). In addition, chitosan also has broad application prospects in the biomedical industry.

The antibacterial properties of DE-ChN are closely related to its degree of deacetylation, molecular weight, solvent and pH of the system. The surface of DE-ChN is rich in amino groups, which is the key to the antibacterial properties of DE-ChN. At present, a large number of scholars have studied the influencing factors of chitosan's antibacterial properties (Sudarshan et al. 1992; Kong et al. 2010). Researchers have found that the source (Chien et al. 2016), deacetylation degree, molecular mass, and concentration of chitosan have a great influence on the antibacterial effect (No et al. 2002; Zheng and Zhu 2003). At the same time, the pH, water content, external factors such as solvents also have a significant impact on the antibacterial ability of chitosan.

In this paper, different amount of DE-ChN were added into GM to prepare GM/DE-ChN nanocomposite films. The effects of different concentration of DE-ChN on the nanocomposite film's mechanical properties, hydrophobicity, thermal stability and four common antibacterial properties were investigated. The final prepared GM based functional food packaging film provides a foundation for the potentials for futural plastic packaging alternaties. The prepared functional food packaging film based on GM can be used as a plastic packaging alternative.

**Experimental**

**Materials**

*Sesbania cannabina seed endosperm* was purchased from Lianyungang city (Jiangsu Province, China). Chitin with a degree of deacetylation of 6.9% was purified from crab shells (*Eriocheir sinensis*) as described in the Liu et al work (Liu et al. 2016). All other chemicals were of analytical grade. *E. coli* (CGMCC 1.8723), *B. subtilis* (CGMCC 1.3358), *S. aureus* (CGMCC 1.2465) and *S. pneumoniae* (CGMCC 1.8722) were purchased from China General Microbiological Culture Collection Center (CGMCC) for the antibacterial test.

**Preparation of GM**

In order to separate GM, the *Sesbania cannabina seed endosperm* was swelled in distilled water for 24 hours, and then mechanically crushed using a juicer. The resulting slurry was stirred at 50 °C for 24 h, and then centrifuged to separate the suspended solids. From the supernatant GM solution, GM was
precipitated with 95% (v/v) ethanol (The volume ratio of GM and ethanol was 1:3) and washed with additional ethanol of 95% (v/v). The precipitated and washed GM was then freeze-dried to yield a powder after the ethanol was evaporated under a fume hood. Finally, the average molecular weight (Mw) of GM was calculated by gel permeation chromatography to be 430,000 Daltons (GPC, Agilent Technologies, Santa Clara, CA, USA), the extracted GM sample consisted of 91.43 ± 0.36 % galactomannan was determined by high performance liquid chromatography (HPLC, ICS 3000, Dionex, USA). The molar ratio of mannose to galactose was determined to be 2.10. These results indicated that the extracted GM was a biomacromolecular with higher purity and high molecular weight, and thus is an ideal matrix for making various materials.

**Preparation of DE-ChN**

Partially deacetylated chitin was prepared as described in a previous report (Fan et al. 2010). Purified chitin was suspended in a (30%wt) NaOH solution and heated at 90°C for 4 h, partially deacetylated chitin with degree of deacetylation of about 25% was prepared successfully, after deacetylation, the sample was collected and washed with distilled water until the supernatant reached a neutral pH. Then the sample was freeze-dried.

To prepare DE-ChN, the sample after freeze-dried was dispersed in distilled water at a concentration of 0.4% (w/v), and the pH was adjusted to 3 using acetic acid under constant stirring; then, the suspension was homogenized at 10,000 rpm for 30 s by a homogenizer (T25, IKA, Germany) and the sonication procedure was conducted at 500 W for 5 min (an interval of 3 s) by an ultra-sonicator (VCX500, USA) and repeated for 5 times; finally the dispersion was centrifuged, and the supernatant was collected as the DE-ChN. For further use, DE-ChN dispersions with 0.5% (w/v) were prepared by dilution or concentration using rotary evaporation.

**Preparation of GM/DE-ChN nanocomposite films**

To prepare GM/DE-ChN nanocomposite films, the GM solution (10 g/L) was first obtained at 50°C with 3 h constant stirring. The GM/DE-ChN nanocomposite films were prepared by ultrasonic dispersion and solution casting. A series of DE-ChN with mass fractions of 20%, 40% and 60% (according to the dry weight of GM) were gradually added to the GM solution under stirring. After ultrasonic defoaming treatment, the above mixture was poured into a polytetrafluoroethylene mold diameter of 9 cm and dried in an oven at 40° C and the nanocomposites films were obtained. All films were stored at 25°C and 50% relative humidity (RH) for at least 48 h before measurement. The composite films were named GM, GM/20% DE-ChN, GM/40% DE-ChN, GM/60% DE-ChN and DE-ChN according to the amount of DE-ChN (0–60%), respectively.

**Characterization of GM/DE-ChN nanocomposite films**
The micro-morphology of nanocomposite films was observed by scanning electron microscopy (SEM) using a Quanta 200 (FEI, American) microscope. The cross section of the films was obtained by applying liquid nitrogen freeze-cracking composite films. To prepare the sample for imaging, the film was pasted onto a row of staples with the cross section facing upwards. The fixed film was then coated with a 20 nm gold layer before recording SEM images.

The light transmittance spectra of nanocomposite films were detected in the range of 200–900 nm using a UV–vis spectrophotometer (Ultrospec 2100, Amersham Bioscience).

The FT-IR spectra of the original GM film, DE-ChN film and GM/DE-ChN nanocomposite film samples were scanned from 400 to 4000 cm\(^{-1}\) in a Nicolet Antaris FT-IR apparatus under dry air at 30 °C.

The mechanical properties (tensile strength and elongation) of GM/DE-ChN nanocomposite films were measured with a TRAPPEZIUM X type tester (Shimadzu Corporation, Japan). Each tested film was cut into 10 x 50 mm specimens. The tensile strength at break of GM/DE-ChN nanocomposite film was measured with a sensor equipped with a load of 500 N. The span length and testing speed were 30 mm and 1 mm/min, respectively. Five specimens of each film were tested and the mean values of the results were calculated.

Thermogravimetric analysis (TGA) of GM/DE-ChN nanocomposite films was carried out with a TGA/209F1 (Libra, German) instrument with a sample weight of approximately 5–15 mg and a heating rate of 10 °C min\(^{-1}\) from 30 °C to 800 °C under flowing nitrogen.

The hydrophobicity of the prepared films was evaluated by recording the water contact angle using an Attension Theta contact angle system apparatus (Biolin Scientific, Sweden). During the measurement, a 4 µL water droplet was dropped on the surface of nanocomposite film and the contact angle was calculated from the average of three parallel samples.

**Cytotoxicity evaluation of GM/DE-ChN nanocomposite films**

Cytotoxicity evaluation of GM/DE-ChN nanocomposite films were carried out in vitro using RAW 264.6 cells (Mouse monocyte macrophages). First, both a GM solution, DE-ChN solution and an GM/DE-ChN solution (concentrations of 25–400 µg/mL) were added to a macrophage medium and cultured with the cells at 37 °C for 24 h in 96-well flat-bottom plate (2×10\(^4\) cells/well). Then, each solution (100 µL) was added to a 10-fold dilution CCK-8 reagent at room temperature in the dark for 1 h. The absorbance of the macrophages cultured in the GM and GM/DE-ChN solutions was determined at 450 nm.

**Antibacterial performance**

To investigate the germicidal ability of the GM/DE-ChN nanocomposites films, a shake flask method was used to test against *B. subtilis*, *E. coli*, *S. aureus* and *S. pneumoniae*. After cultivation in liquid lysogeny
broth (LB) medium (containing 10 g/L peptone, 5 g/L yeast extract, and 10 g/L sodium chloride) for 12 h at 37°C, the microorganisms were diluted with the LB liquid culture medium to obtain a bacterial suspension with approximately $1.0 \times 10^6$ CFU/mL concentration. According to Xu et al work (Xu et al. 2019), 0.01 g composites film was immersed in 10 mL of bacterial suspension adjusted pH 5.5. The solution was then cultured in a shaker stirring at 150 rpm and at 37°C for 6 h. Then, a 100 µL of 7-fold serial dilutions was pipetted into agar plates. The mixture was incubated in incubator at 37°C for 12 h and the number of bacteria was obtained using the colony forming count method.

**Statistical analysis**

Statistical analysis was performed using a commercial SPSS program (SPSS 20.00, SPSS INC., Chicago, Illinois, USA). The data were expressed as mean ± standard deviation (SD) and $p < 0.05$ was taken as the minimal level of significance.

**Results And Discussion**

**Morphology and UV absorption of GM/DE-ChN nanocomposite films**

DE-ChN has been widely reported in the preparation of composite film, which endows the composite film excellent chemical and physical properties. Light transmittance is an auxiliary means to evaluate the compatibility of polymers. If the compatibility between DE-ChN and GM molecules in the GM/DE-ChN nanocomposite film is poor, then light reflection or scattering will occur at the phase interface between them, the light transmittance of the film is reduced. The film prepared by mechanically mixing had a smooth surface and higher transparency due to its high level of uniformity (Gennadios et al. 1998).

Digital images of the composite film showed the transparency of the film as can be seen from Fig. 1. The original GM film was slight matt, and the light transmittance of the GM film compounded with DE-ChN increased from 73% (original GM film) to 86% (GM/60% DE-ChN film). Both films had no UV absorption. Figure. 2a’ ~ e’ displayed the SEM images of the cross-section of GM/DE-ChN nanocomposite films with different content of DE-ChN addition. All the cross-section SEM images of GM/DE-ChN nanocomposite film showed a multilayer structure, just like the physical structure of natural mother pearl shellfish, so the composite film had good physical and chemical properties (Wang et al. 2017).

**FT-IR analysis of GM/DE-ChN nanocomposite films**

To investigate the effect of DE-ChN on the structure of the GM films, the chemical bonds in the original GM film, DE-ChN film and the GM/DE-ChN nanocomposite film were surveyed using ATR-FTIR. When a H atom was connected to O, F and other atoms with a small radius and strong electronegativity, the electron cloud moved and exposed the H atom nucleus, which was easily electrostatically attracted to other atoms with lone pairs of electrons, forming a strong dipolar effect, this was the hydrogen bond, which would
affect the infrared spectrum. From Figure 3, it can be seen that the FTIR spectra range of 3310–3350 cm\(^{-1}\) indicated the -OH and the -NH stretching vibration. The -CH stretching vibration was attributed to the range of 2800–3000 cm\(^{-1}\). The typical amino bond of chitosan wavenumber appeared in the range of 1620–1720 cm\(^{-1}\) and 1520–1580 cm\(^{-1}\) (1500 cm\(^{-1}\), 1655 cm\(^{-1}\) and1630 cm\(^{-1}\)) due to the amine and (Martins et al. 2012; Li et al. 2013; Darbasi et al. 2017). The ATR-FTIR spectra contained intense bands at 1647 cm\(^{-1}\) (C = O), 1016 cm\(^{-1}\) (C-O-C and C-OH on the pyran ring) (William Kemp 2002). The 1380 cm\(^{-1}\) was the absorption peak of -CH\(_3\) symmetric deformation vibration and -CH bending vibration (Lawrie et al. 2007), which proved that chitosan was successfully compounded on GM (Jóźwiak et al. 2017).

**Mechanical properties of GM/DE-ChN nanocomposite films**

The mechanical properties of the original GM film, DE-ChN film and GM/DE-ChN nanocomposite film materials were shown in Table 1. Compared with the original GM film, the tensile strength values of GM/DE-ChN nanocomposite films improved significantly. This was because chitosan has higher mechanical properties than that of GM due to its linear characteristics. The interaction mechanism of DE-ChN and GM in nanocomposite films as can be seen from Figure 4. The addition of polysaccharide enhanced the hydrogen bonding force between GM molecules, thereby increasing the tensile strength of the GM/DE-ChN nanocomposite film (Mollah et al. 2016). According to the data in Table 1, when the addition amount of DE-ChN was 60%, the maximum tensile strength of GM/DE-ChN nanocomposite film was 107 MPa, which was 343% higher than the original GM film. Similar results were reported by Pinotti et al. (Pinotti et al. 2007) who indicated the reduction in methylcellulose film flexibility with increasing chitosan concentration. Khan et al. (Khan et al. 2012) reported that chitosan incorporation (5%- 36% wt) in methylcellulose-based films significantly improved the tensile strength of films with the reduction of viscoelasticity. The 50% chitosan containing starch-based films was considered as the optimum because the films had good strength (47 MPa). Moreover, the appearance of the films was quite transparent. As the content of chitosan increased, the hydrogen bonding force between polysaccharide molecules increased, and the tensile strength of the corresponding film also increased.

### Table 1

| Sample          | Tensile strength (MPa) | Elongation at break (%) | Young’s module (GPa) |
|-----------------|------------------------|-------------------------|----------------------|
| GM              | 31.2 ± 5.2             | 7.5 ± 3.9               | 1.8 ± 0.6            |
| DE-ChN          | 110.1 ± 6.0            | 4.1 ± 0.3               | 4.1 ± 0.4            |
| GM/20%DE-ChN    | 80.6 ± 4.1             | 4.6 ± 0.7               | 3.2 ± 0.4            |
| GM/40%DE-ChN    | 103.0 ± 2.9            | 6.6 ± 1.4               | 4.1 ± 0.4            |
| GM/60%DE-ChN    | 107.1 ± 9.8            | 3.3 ± 0.9               | 4.0 ± 0.5            |
TGA and DTG analysis of GM/DE-ChN nanocomposite films

The thermal stability of the GM/DE-ChN nanocomposite films with different DE-ChN addition amounts were shown in Fig. 5. The weight loss of the composite films was divided into the following three stages which can be observed from Fig. 5A. The first weight loss started from 40 °C to 150 °C. This part was mainly the physical removal of water in the GM and DE-ChN composite film. The next step started at about 250°C, and the GM began to decompose at high temperature (Pang et al. 2015). The ash content of GM film, GM/20% DE-ChN film, GM/40% DE-ChN film, GM/60% DE-ChN film and DE-ChN film were 16.3%, 20.5%, 22.9%, 27.1% and 20.1%, respectively. The maximum degradation temperature was 298.3 °C, 293.8 °C, 288.4 °C, 286.6 °C and 268.7 °C, respectively. It could be seen from Fig. 4 that the change trend of the residual mass fraction was GM/60% DE-ChN > GM/40% DE-ChN > GM/20% DE-ChN > DE-ChN > GM. From the derivative thermogravimetric analysis curve (DTG) of the composite films, it could be seen that the original GM film, the original DE-ChN film and the GM/DE-ChN nanocomposite film had different endothermic peaks in the DTG curve. The endothermic peak of the GM composite film with different addition amounts of DE-ChN had shifted. This characteristic due to the different degradation temperature of the material. The degradation temperature of the original GM film was about 300 °C, while the DE-ChN/GM composite film had a second endothermic peak at 400 °C (Yao et al. 2006a). For the original DE-ChN film, the maximum degradation temperature started at about 400 °C, so the DE-ChN/GM composite film had a third peak, but the original GM film did not contain DE-ChN without this peak. The results showed that the GM composite film added with DE-ChN still maintained the thermal stability of the original GM film.

Water contact angle determination of GM/DE-ChN nanocomposite films

Figure. 6 showed the effects of DE-ChN addition on the water contact angle of composite films. The water contact angle of GM film was about 73°. DE-ChN could significantly improve the water contact angle of composite films. This was because the molecular chain of chitosan was semi-rigid and contained a large number of hydroxyl and amino groups. The hydroxy and hydroxy groups, the hydroxy groups and the amino groups or the amino groups and the amino groups were paired with each other to form a zipper-like hydrogen bond, which bind the polar groups in the film, and there were fewer polar groups on the film surface, resulting in the surface water contact angle of the GM/DE-ChN nanocomposite film increased (Liu et al. 2010). GM/20% DE-ChN nanocomposite films had the significant maximum water contact angle of 119° (P < 0.001), which were increased by about 63%. That may be related to the high content of hydroxyl and amino groups in the film. Compared with other samples with DE-ChN content, more hydroxyl and amino groups in the film were paired with each other, forming more zipper-like hydrogen bonds, and more polar groups of the film were bound inside, thereby reducing the polar groups on the surface of the film, resulting in a high-water contact angle of the film sample.
Cytotoxicity evaluation of GM/DE-ChN nanocomposite films

In order to evaluate the safety of the prepared films, in vitro cytotoxicity tests were carried out for GM and GM/DE-ChN nanocomposite solutions by culturing with RAW 264.7 macrophages cells. Each sample was added to the macrophage medium at the concentration of 25–800 µg/mL and CK was used as a control blank (Zhou et al. 2018). The effects of different DE-ChN addition amount of GM composite solution on the toxicity of macrophages was shown in Fig. 7. On the whole, the effect of GM/DE-ChN nanocomposite solution on macrophage activity increased in a concentration-dependent manner and then decreased. Chitosan was a linear polymer of N-acetyl-D glucosamine and deacetylglucosamine. It had certain characteristics of glycosaminoglycan and hyaluronic acid, and had certain biological activities (Tian and Ye 2012). Chitosan can promote human monocytes to produce TGF-inhibitor and platelet derived growth factor (TGF-inhibitor) (Yao et al. 2006b). In addition, chitosan treatment could also induce the expression of major histocompatibility complex (MHC) mannose receptor and other activation markers. It has been reported in the literature that polysaccharides enhance the expression of arginase, thereby enhancing the activity of arginase metabolic pathways, and stimulating the proliferation of macrophages (Bhardwaj et al. 2020).

Antibacterial performance

The sterilization property against *E. coli*, *B. subtilis*, *S. aureus* and *S. pneumoniae* of composite films was determined by the colony counting method. The bacterial suspension was inoculated in the solid medium and the numbers of total viable counts were visualized in Figure. 8 ~ 11. The bacterial colony of *E. coli*, *B. subtilis*, *S. aureus* and *S. pneumoniae* were almost fully grown on the agar plates after the treatment of original GM film, but less observed in the presence of GM/DE-ChN nanocomposite films. The antibacterial effects of the original GM solution and the GM/DE-ChN nanocomposite solution on *E. coli* are shown in Fig. 8. As the amount of DE-ChN added in GM increased, the antibacterial effect of GM/DE-ChN nanocomposite solution on *E. coli* also increased. The inhibitory rate of GM/40% DE-ChN film to *E. coli* was 7%, and the inhibitory rate of GM/60% DE-ChN film to *E. coli* was 33%, and the antibacterial effect was significant (P < 0.01). The antibacterial effect of the film on *B. subtilis* was shown in Fig. 9. As the amount of DE-ChN added in GM increased, the antibacterial effect of DE-ChN/GM composite solution on *B. subtilis* also increased. GM/20% DE-ChN, GM/40% DE-ChN, GM/60% DE-ChN and original DE-ChN films had an inhibitory rate of 99% against *B. subtilis*, the antibacterial effect was particularly significant (P < 0.001). Figure 10 showed the antibacterial effect of the film on *S. aureus*, the GM/20% DE-ChN film had an inhibitory rate of 13% against *S. aureus*, GM/40% DE-ChN and GM/60% film had an inhibitory rate of 46% and 47% against *S. aureus*, and the inhibitory rate of DE-ChN film against *S. aureus* was 57%, the antibacterial effect was particularly significant (P < 0.001). The antibacterial effect of the composite film on *S. pneumoniae* was shown in Fig. 11. The bacteriostatic rate of GM/20% DE-ChN against *S. pneumoniae* was 29%, GM/40% DE-ChN, GM/60% DE-ChN and the inhibitory rates of DE-ChN against *S. pneumoniae* were significantly 88%, 92% and 94%, respectively (P < 0.001). This was because the amino group in chitosan was cat ionized at pH 5.5, then the interaction between chitosan and bacteria was
activated (Raafat et al. 2008). The permeability of the bacterial cell membrane changes, causing internal osmotic imbalance, leading to the leakage of electrolytes such as K⁺ and other low-molecular-weight protein components in the cell, and finally leading to the apoptosis of microorganisms (Liu et al. 2004). The different antibacterial activities of the films against gram-positive bacteria and gram-negative bacteria might be due to the different cell wall structures of the two bacteria (Jing et al. 2007). The cell wall of gram-positive bacteria was composed of polypeptidoglycan, and chitosan could easily pass through the peptidoglycan network and directly act on the cell membrane. However, the cell wall of gram-negative bacteria contained not only an inner membrane composed of peptidoglycan and lipopolysaccharide, but also an outer membrane composed of lipoproteins and phospholipids. The outer membrane of gram-negative bacteria could act as an effective external barrier for macromolecules, which might prevent the macromolecule chitosan from reaching the plasma membrane. Therefore, chitosan showed a stronger antibacterial effect on gram-positive bacteria than gram-negative bacteria.

Conclusions

The GM/DE-ChN nanocomposite films showed great antibacterial properties for the prohibition of *E. coli*, *B. subtilis*, *S. aureus* and *S. pneumoniae*. When the additional amount of DE-ChN was 60% into GM, the antibacterial rate of GM/DE-ChN nanocomposite film against *E. coli*, *B. subtilis*, *S. aureus* and *S. pneumoniae* reached 33%, 99%, 47% and 92%. The obtained GM/DE-ChN nanocomposite films had improved tensile strength (107 MPa), which was 343% higher than the original GM film. The GM/DE-ChN nanocomposite films had excellent hydrophobicity of 107°, which was 147% higher than the original GM film. Moreover, the GM/DE-ChN nanocomposite film showed no toxicity to macrophage cells. The above conclusions indicated that the biocompatible DE-ChN had significant antibacterial properties against common bacteria. The physical and chemical properties of GM films were improved. Therefore, combining DE-ChN with renewable and sustainable packaging materials to prepare packaging materials with antibacterial properties will provide a new method for food packaging without harmful additives.

Declarations

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Author contribution

Wanying Liu: Conceptualization, Methodology, Formal analysis, Writing-Original Draft, Writing-Review & Editing.

Ling Zhe: Visualization
Compliance with ethical standards

Conflict of interest

The authors declare no competing financial interest.

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Figures

Figure 1

UV transmittance of GM/DE-ChN nanocomposite films
Figure 2

Digital (a ~ e) and SEM (a’ ~ e’) images of the cross-section of GM/DE-ChN nanocomposite films: (a) Original GM film; (b) GM/20%DE-ChN film; (c) GM/40%DE-ChN film; (d) GM/60%DE-ChN film; (e) Original DE-ChN film
Figure 3

FT-IR spectra of GM/DE-ChN nanocomposite films

Figure 4
The hypothetical interaction mechanism of DE-ChN and GM in nanocomposite films.

Figure 5

TGA (A) and DTG (B) analysis for GM/DE-ChN nanocomposite films.

Figure 6
The water contact angle of GM/DE-ChN nanocomposite films (P<0.05 *, P<0.01 **, P<0.001 ***)

Figure 7

Effects of the GM and GM/DE-ChN nanocomposite films on macrophage activity (P<0.05 *, P<0.01 **, P<0.001 ***)

Figure 7

Effects of the GM and GM/DE-ChN nanocomposite films on macrophage activity (P<0.05 *, P<0.01 **, P<0.001 ***)
**Figure 8**

Antibacterial effect of treating E. coli with nanocomposite films (Blank control group, a GM film, b GM/20%DE-ChN film, c GM/40%DE-ChN film, d GM/60%DE-ChN film, e DE-ChN film) (P<0.05 *, P<0.01 **, P<0.001 ***)

**Figure 9**

Antibacterial effect of treating B. subtilis with nanocomposite films (Blank control group, a GM film, b GM/20%DE-ChN film, c GM/40%DE-ChN film, d GM/60%DE-ChN film, e DE-ChN film) (P<0.05 *, P<0.01 **, P<0.001 ***
Figure 10

Antibacterial effect of treating S. aureus with nanocomposite films (Blank control group, a GM film, b GM/20%DE-ChN film, c GM/40%DE-ChN film, d GM/60%DE-ChN film, e DE-ChN film) (P<0.05 *, P<0.01 **, P<0.001 ***)

Figure 11

Antibacterial effect of treating S. pneumoniae with nanocomposite films (Blank control group, a GM film, b GM/20%DE-ChN film, c GM/40%DE-ChN film, d GM/60%DE-ChN film, e DE-ChN film) (P<0.05 *, P<0.01 **, P<0.001 ***)
