Enhanced Antibacterial and Antioxidant Properties of Chitosan Films Blended with Gallic Acid and Incorporated with Thymol Silver Nanoparticles

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Abstract

In the present study, an attempt has been made to build and evaluate Chitosan+Glycerol/Gallic acid/Thymol-silver nanoparticles or chitosan blended (C+G/GA/T-SNPs) film to significantly improve antioxidant and antibacterial activity for accelerated wound healing. Methanolic Gallic acid is used for the first time in antibacterial chitosan control (C+G) films. All developed films, compounds was Thymol and Gallic acid and their synthesized Thymol silver nanoparticles (T-SNPs) and Gallic acid silver nanoparticles (GA-SNPs) were characterized by Fourier-transform infrared spectroscopy (FT-IR), Scanning electron microscopy (SEM), X-Ray diffraction (XRD), Zeta potential (ZP), Dynamic light scattering (DLS), and UV-Vis spectroscopy. T-SNPs and GA-SNPs are rod and spherical in shape and were sufficient to reduce, capped, and stabilize. T-SNPs and GA-SNPs were measured Dynamic Light scattering and found to be 123.2 nm and 121.1 nm with surface charges of -19.7 and -20.3 respectively. The incorporation of methanolic Gallic acid and T-SNPs into chitosan films, as predicted, effectively enhanced antioxidant and antimicrobial activity. The antimicrobial activity of Thymol, T-SNPs and C+G/GA/T-SNPs film showed more zone of inhibition than Gallic acid, GA-SNPs and C+G film. The elasticity, texture and folding endurance of the C+G film and C+G/GA/T-SNPs films have been substantially improved. The ecological quality of the generated C+G and C+G/GA/T-SNPs film was determined by the assessment of soil degradation and water degradation parameters. These findings lead to the conclusion that the C+G/GA/T-SNPs film produced with Gallic acid and T-SNPs can improve wound healing.

Keywords: Chitosan, Gallic Acid, Thymol Silver Nanoparticles, Antibacterial and Antioxidant Properties.

1. INTRODUCTION

Damage to the integrity of the skin leads to the formation of a wound that can occur anywhere in the body and adapts to spontaneous wound healing (Sabine et al. 2017; Wang et al. 2020). The largest external organ in the human body, the skin, plays a crucial role in protecting the body from mechanical damage and microbial invasion. As a consequence, wound healing is the most important process for shielding the human body from serious injury.

In general, wound healing occurs in four phases; (1) haemostatic stage, (2) inflammatory response stage, (3) cell proliferation stage, and (4) collagen fibre remodeling stage. During the haemostatic transition, platelets, and fibrin coagulate successively to form thrombi, causing the skin wound to contract and preventing further blood loss. The inflammatory stage is characterized by the invasion of immune cells to provide optimal conditions for the eventual spread of fibroblasts. Consequently, cell proliferation occurs, where fibroblasts divide rapidly in response to various growth factors and secrete collagen fibers. These collagen fibers grow and mature to complete the wound healing process. In the event that wounds lead to infection could lead to serious problems, thus care for wound healing is essential (Wang et al. 2020).

Chitosan, a deacetylated descendant of chitin, is widely used in the manufacture of biodegradable, biocompatible wound healing material. Chemically chitosan consists of (1, 4)-linked 2-amino-deoxy-b-D-glucan. It has a wide range of applications in pharmacy, medicine, food, agriculture, textile products, hygiene, and water treatment due to improved mechanical, chemical, and biological properties. In addition, in order to improve the mechanical, hydrophobic, antimicrobial, and antioxidant properties of chitosan film, various active compounds have been complexed into chitosan film (Kaya et al. 2019).

Gallic acid is a phenolic compound chemically referred to as 3,4,5 trihydroxybenzoic acid. It is a crystalline solid, slightly colorless, or slightly yellow with a molecular weight of 170.12 g/mol and a molecular formula of C₇H₆O₆. The melting point of Gallic acid is 210°C which decays between 235° and 240°C, which generates carbon dioxide and carbon monoxide (Fernandes et al. 2016). Gallic acid has been
reported to stimulate various biological activities, such as antibacterial, anti-fungal, anti-viral, anti-inflammatory, anti-oxidant, anti-cancer, anti-diabetic, etc., which have potential in the food, medical, pharmaceutical, biotechnological, and cosmetic industries (Nayeen et al. 2016).

Thymol, chemically known as 2-isopropyl-5-methylphenol, is a white crystalline monoterpene phenol that gives thyme a strong flavor, a pleasant aromatic fragrance and strong antiseptic properties. Thymol is used as an antibacterial, antioxidant and also as a surgical disinfectant, a general disinfectant and anaesthetic agent (Marchese et al. 2016; Meeren et al. 2017). The pharmacophoric properties of Thymol are the phenolic hydroxyl group, in its chemical structure, which delivers defence-free radicals and increases the endogenous antioxidant.

Considering the different useful properties of the biopolymers Chitosan, Gallic acid, Thymol silver nanoparticles (T-SNPs), the purpose of this study was to produce wound healing chitosan blended (C+G/GA/T-SNP) film, consisting of methanolic Gallic acid (antioxidant and wound healing) and incorporated T-SNPs (antibacterial). Synthesized C+G/GA/T-SNP film have been characterized physicochemically by solubility assay, UV spectrophotometer, Fourier Transform Infra-Red Spectroscopy (FTIR), Zeta Potential (ZP), X-Ray Diffraction (XRD), and Dynamic Light Scattering (DSL). In addition, essential properties such as biodegradability and water solubility are evaluated. Furthermore, the synthesized T-SNPs, GA-SNPs, and Thymol and Gallic acid have been used for analysis of antimicrobial, antioxidant properties. In particular, C+G/GA/T-SNPs films have been tested for biological potential including antimicrobial, antioxidant, and wound healing activities.

2. MATERIALS AND METHODS

2.1. Materials

2.1.1. The solvents required for solubility test. The Acetic acid, Chloroform, Acetonitrile, Acetone, Formaldehyde, Toluene, Tert butyl alcohol, Ethyl ether, Methanol, 100% alcohol, Tween 20, Dimethyl sulfoxide, N, N Dimethyl formamide, and double distilled water were of analytical grade (Sigma-Aldrich Chemicals).

2.1.2. The requirement for silver nanoparticles formulation. 250 mL flask, double distilled water, 100 mL measuring cylinder, Silver nitrate (AgNO₃) was purchased by Sigma-Aldrich.

2.1.3. The reagent used for Antioxidant activity. 2,2-Diphenyl-1-Picrylhydrazyl (DPPH, Sigma-Aldrich Chemicals), Ascorbic acid (ACS Chemicals), and Methanol (Sigma-Aldrich Chemicals).

2.1.4. The reagent used for Antimicrobial analysis. Chitosan (medium molecular weight and 85% deacetylation degree, Cas No. 440877) was purchased from SIGMA-Aldrich, glycerol (C₃H₈O₃), Gallic Acid (HiMedia), Thymol (S. D. Fine Chem. Limited), and Silver Nitrate (AgNO₃) (Sigma-Aldrich Chemicals).

2.1.5. The media and bacterial culture used in Antimicrobial assay. Nutrient Agar with composition as follows peptone (5.000 gms/Litre), sodium chloride (5.000 gms/Litre), HM peptone B# (1.500 gms/Litre), Yeast extract (1.500 gms/Litre), Agar (15.000 gms/Litre) and final pH (at 25°C) 7.4±0.2. The bacterial culture such as Escherichia coli, Klebsiella pneumoniae, Staphylococcus aureus, and Sporosarcina Pasteurii were gift from the Consultancy, Rajiv Gandhi Biotechnology Centre, Rashtrasant Tukadoji Maharaj Nagpur University, L.T.T Premises, Nagpur, Maharashtra, India.

2.1.6 Materials for Chitosan films production. Chitosan and Gallic acid were purchased from Himedia and acetic acid (CH₃COOH), glycerol (C₃H₈O₃), and methanol (CH₃OH) were supplied from Sigma-Aldrich Chemicals.

2.2. Solubility test of Thymol and Gallic acid

The solubility of Thymol and Gallic acid was undertaken in various organic solvents. Analyzes were performed using 2 mg of Thymol and Gallic acid separately in the test tube and 5 mL of solvents were introduced and blended (Acetic acid, Chloroform, Acetonitrile, Acetone, Formaldehyde, Toluene, Tert-butyl alcohol, Ethyl ether, Methanol, 100% alcohol, Tween 20, Dimethyl sulfoxide, and N, N Dimethyl formamide) (Lade et al. 2014). Further, after a few minutes, the solubility and time needed for the solubility of Thymol and Gallic acid in various solvents were documented.

2.3. Synthesis of silver nanoparticles of Thymol and Gallic acid using silver nitrate (AgNO₃)

Silver nanoparticles using Thymol and Gallic acid were synthesized by combining 95 mL of 1 mM silver nitrate (AgNO₃) with 5 mL of 1% Thymol and Gallic acid separately. The reaction was performed in the presence of sunlight (March 2020; temperature 25-30°C) and after 5 minutes, 1 N, NaOH solution was added to the mixture solution to maintain alkaline 8.5-9 pH (Lade et al. 2018).

2.4. Characterization of synthesized T-SNPs and GA-SNPs

The bio reduction of silver nitrate using 1% Thymol and Gallic acid was monitored at various times from 0 min to 24 hr and visual examination for change in colour of the solution was observed. The T-SNPs and GA-SNPs were centrifuged (Spectrofuge 7M) at 13,000 rpm for 15 min to forms colloidal pellets at the bottom. The synthesized T-SNPs and GA-SNPs were used for characterization purposes.

2.5. UV-Visible spectroscopy (UV-VIS)

The synthesis of T-SNPs and GA-SNPs was studied using Shimadzu 1800 UV-Vis spectrophotometer was used for recording the UV-vis spectra using a 10 mm path length quartz cuvette (Chandra et al. 2018). The instrument was set for base line correction and auto zero using methanol. Measuring the UV-Visible spectra at a resolution of 200-800 nm.

2.6. Fourier transforms infrared spectroscopy (FTIR) analysis

The FTIR analysis of the Thymol, Gallic acid, T-SNPs, and GA-SNPs were carried out to identify the primary functional groups of the compounds. The FTIR measurements were carried out using an Agilent Technology Cary 630 FTIR spectrophotometer by employing the KBr disc technique. The FTIR spectrum of the formulation was recorded at a resolution of 4 cm-1 in the transmission mode (4000–440 cm-1) (Chandra et al. 2018).

2.7. Zeta potential (ZP) analysis

The surface charge on T-SNPs and GA-SNPs determined using zetasizer (nano ZS, Malvern instrument Ltd., UK) at 25 °C. The zeta potential determines the colloidal stability of particles and surface charge over nanoparticles. The T-SNPs and GA-SNPs are subjected to Brownian motions due to thermal energy that produces Rayleigh scattering (Chandra 2018; lade et al. 2018). Liquid samples of the nanoparticles (5 mL) were diluted with double distilled water (50 mL) using NaCl as suspending electrolyte solution (2 x10-2 M
NaCl]. Connect a 1 ml pre-filled syringe with the sample to a cell port and point the cell down (ports down), then slowly inject the sample to the middle of the circuit at the bottom of the cell that it has formed and make sure no bubbles form in the cell. Return the cell in the vertical position (ports up) and continue to inject the sample from the syringe. Fill to the most extreme level of the cell (complete volume 0.8 ml or 0.75 mL relying upon the cell). Then, at that point, make sure that the electrodes are completely immersed in the liquid and there are no bubbles in the cell. Remove the syringe and cap the filled cells with its two caps (Caputo et al. 2015).

2.8. Dynamic light scattering (DLS) analysis

The size and size distribution of particles in the colloids T-SNPs and GA-SNPs were measured using a Nano Zetasizer system (Malvern Instruments). Measurement parameters were as follows: a laser wavelength of 633 nm (He-Ne), a scattering angle of 173° (fixed-without changing possibility), a measurement temperature of 25°C, a medium viscosity of 0.8872 mPa·s and a medium refractive index of 1.330, and material refractive index of 0.200 (Tomaszewksa et al. 2013). Prior to DLS estimation, the colloid was gone through a 0.2 µm polyvinylidene fluoride (PVDF) layer. The sample was loaded into quartz cuvette and five estimations were performed, for which the mean outcome was recorded. DLS studies were done in two modes: general purpose mode (with normal resolution) and multiple narrow modes (with high resolution).

2.9. X-ray diffraction (XRD) analysis

The crystal structures of T-SNPs and GA-SNPs were measured by a Rigaku® X-ray diffractometer (RINT2000, Tokyo, Japan) at a voltage of 40 kV and 15 mA. The scattered radiation was detected in the angular range 2θ = 5 - 80° with a scanning speed of 5°/min (Cui et al. 2018).

2.10. Scanning electron microscopy (SEM) analysis

The morphology of T-SNPs and GA-SNPs were used for scanning electron microscopic analysis using Jeol 6390LA/OXFORD XMX N with accelerated voltage of 20 kV in DST-SAIF Cochin, India (Lade et al. 2018).}

2.11. Antioxidant activity

2.11.1. DPPH (2,2′-diphenyl-1-picrylhydrazyl) radical scavenging assay. UV-visible spectrometry was applied to evaluate the scavenging rate of 2,2-diphenyl-1-picrylhydrazyl (DPPH) radicals of Thymol, T-SNPs, Gallic acid, GA-SNPs, C+G film, and C+G/GA/T-SNPs film. Briefly, 10 mg of each sample was dissolved in 10 mL of methanol to give a concentration of 1 mg/mL. Then prepared sample concentrations of 5, 10, 15, 25 and 50 µg/mL and make up the volume of each sample in methanol then add 75 µL of 1mM DPPH. The mixtures were incubated for 30 min in dark room condition and then measured at 517 nm via a Shimadzu UV-1800 UV spectrophotometer and IC50 (Inhibitory concentration to scavenge 50% free radicals) is also determined. Lower the absorbance of the reaction mixture indicates higher free radicals scavenging activity. IC50 value denotes the concentration of sample required to scavenge 50% of the DPPH free radicals (Bhalodia et al. 2011). The capability to scavenge the DPPH radicals was calculated using the following equation. The investigation was repeated three times and the antioxidant activities for each example were processed with the following formula:

\[
A \text{ control} - A \text{ sample} \\
A \text{ control} \\
\times 100
\]

Where,

A control = Absorbance of DPPH Solution.
A sample = Absorbance of DPPH along with a different concentration of samples.

IC50 was calculated from equation of line obtained by plotting a graph of concentration versus % inhibition

2.12. Antimicrobial activity

Antimicrobial activity of Gallic acid, Thymol, T-SNPs and GA-SNPs effectively against Escherichia coli, Klebsiella pneumoniae, Staphylococcus aureus, and Soporosarcina Pasteurii were evaluated using agar well diffusion method.

2.12.1. Agar well diffusion method. Agar well diffusion method is broadly used to assess antimicrobial activity. The agar plate surface inoculated by spreading a volume of the microbial inoculums over the entire nutrient agar plate surface. Then, a hole with a diameter of 4 to 6 mm is punched aseptically with a sterile cork borer or tip, and a volume (30 µL) of the antimicrobial agent or Thymol and Gallic acid solution at desired concentration is introduced into the well. Then, Nutrient agar plates are incubated at 37°C for 24 hours (Balouiri et al. 2016).

2.13. Chitosan films production

The two different types of C+G films were produced, one of which were incorporated with Gallic acid methanolic extract and T-SNPs while the one was used as a control without incorporating any extract. Briefly, 1 g of chitosan was added to 100 mL of 1% acetic acid to make chitosan solution. The mixture was stirred for 24 h via magnetic stirrer. Then, 1 g of Gallic acid was added to 5 mL methanol and was vortex (REMI CM-101, Vortex mixer) for 5 min. Then 1 mL of methanol mixture and 1 mL liquid T-SNPs was incorporated with 50 mL of chitosan solution and 200 µL of glycerol as plasticizer (Kaya et al. 2018). All solutions were homogenized at 2600 rpm for 5 min with a homogenizer (Heidolph, Silent Crusher M). Each solution was poured into a plastic Petri plate and the Petri plates were kept at 30°C for 48 h. Finally, the films were collected from Petri plate and stored for further analysis.

2.14. Characterization of chitosan control and chitosan blended film

2.14.1. Visual appearance. The Visual appearance C+G film and C+G/GA/T-SNPs film were observed for colour and physical form or appearance (Naik et al. 2019).

2.14.2. Determination of thickness. The thickness of each film was measured using screw guage (thickness tester) at different position of the film and the average was calculated (Naik et al. 2019).

2.14.3. Folding endurance. The folding endurance is expressed as the number of folds in films (number of times the film is folded at the same place, either to break the strip/ film or to develop visible cracks). The C+G film and C+G/GA/T-SNPs film was folded in the centre, between the fingers and the thumb and then opened. This was termed as one folding. The cycle was repeated till the film showed rupture or breaks in centre of film or folded up to 100 times which is considered satisfactory to reveal good film properties (Naik et al. 2019). The total numbers of folding were named as folding endurance value. This test was carried out on all the C+G film and C+G/GA/T-SNPs film for three times and the mean of three readings were recorded.

2.14.4. Biodegradability of film. Biodegradability was tried by using two unique strategies i.e., water solubility and soil
degradation (Kaya et al. 2018). Both of the techniques were repeated thrice. Solubility in water and soil was determined by using the following equation:

\[ WL(\%) = \frac{\text{weight loss}}{\text{initial weight}} \times 100 \]

### 2.1.4.1 Solubility in water

For testing the films solubility in water, the films sample was cut into small pieces (2 × 2 cm) and the initial dry weights were measured for all the two samples. Next, 20 mL of water was added to the films and were kept at room temperature for 72 h. After 72 h of immersion of the film samples; they were dried at 60°C and were weight again.

### 2.1.4.2 Soil degradability

Small pieces of the dry films (2 × 2 cm) were scaled and then put into the enriching soil. Around 10 mL of water was added to the samples day by day and following 15–30 days the final weights were measured (Kaya et al. 2018).

### 2.1.5 Surface pH

The C+G film and C+G/CA/T-SNPs film were left to swell for 1 hour on the surface of the agar plate, prepared by dissolving 2% w/v agar in warmed double distilled water with constant stirring and poured into the petridish to solidify at room temperature. The pH paper was placed on the surface of the swollen film and the pH was determined (Naik et al. 2019). The mean of three readings was recorded.

### 2.1.6 Fourier transforms infrared (FTIR) analysis

The FTIR analysis of the pure chitosan, C+G film and C+G/CA/T-SNPs film was carried out to identify the primary functional groups of the compounds. The FTIR measurements were carried out using an Agilent Technology Cary 630 FTIR spectrophotometer. The FTIR spectrum of the formulation was recorded at a resolution of 4 cm⁻¹ in the transmission mode (4000–4400 cm⁻¹) (Chandra et al. 2018).

### 2.1.7. X-ray diffraction analysis

The crystal structures of C+G film and C+G/CA/T-SNPs film were measured by a Rigaku® X-ray diffractometer (RINT2000, Tokyo, Japan) at a voltage of 40 kV and 15 mA. The scattered radiation was detected in the angular range 2θ = 5–80° with a scanning speed of 5°/min (Cui et al. 2018).

### 2.1.8. SEM analysis

The morphology of C+G film and C+G/CA/T-SNPs film were used for scanning electron microscopy analysis using jeol 6390LA/OXFORD XMX N with accelerated voltage of 20 kV in DST-SAIF Cochin, India (Lade et al. 2018).

### 2.1.9. Antimicrobial activity

Antimicrobial activity of C+G film and C+G/CA/T-SNPs film effectively against Eschericchia coli, Klebsiella pneumoniae, Staphylococcus aureus, and Sporosarcina Pasteurii were evaluated using agar disc diffusion method.

### 2.1.9.1 Agar disc diffusion method

Agar disc diffusion method is widely used to evaluate the antimicrobial activity. The agar plate surface inoculated by spreading a volume of the microbial inoculums over the entire nutrient agar plate surface. After 30 mins the C+G film and C+G/CA/T-SNPs film (2×2 cm) were placed on the semi dried inoculated plates using sterile forceps and press slightly in the centre of the film using the forcep. Then, agar plates are incubated at 37°C for 24 hours. The antimicrobial agent diffuses in the agar medium and inhibits the growth of microbial strain tested (Naik et al. 2019).

### 3. Results and Discussion

The Present study investigated the antimicrobial and antioxidant potential of Thymol, gallic acid, T-SNPs, and GA-SNPs. To validate, the solubility test of Gallic acid and Thymol in different solvents at the room temperature and characterized through the instruments such as UV-Visible Spectroscopy, Fourier Transform Infra-Red Spectroscopy, Zeta potential, and Dynamic Light Scattering. Further, the antimicrobial activity of Thymol and Gallic acid in different concentration was studied against four different bacterial cultures (E. coli, Klebsiella, S. aureus, and P. Pasteurii). Finally a C+G film and C+G/CA/T-SNPs film is formulated and antimicrobial and antioxidant potential was evaluated against gram positive and gram negative bacteria.

### 3.1 Solubility analysis of Thymol and Gallic acid

The solubility of Thymol and Gallic acid were recorded in different types of solvent and distilled water. The Thymol was completely soluble in Acetic acid, Chloroform, Acetonitrile, Acetone, Formaldehyde, Tolune, Tert-butyl alcohol, Ethyl ether, Methanol, 100% alcohol, Tween20, Dimethyl sulfoxide, and N, N Dimethyl formamide. However, Thymol was slightly soluble in distilled water. Similarly, it was observed that the Gallic acid was completely soluble in Acetone, Formaldehyde, Methanol, 100% alcohol, Dimethyl sulfoxide, and N, N Dimethyl formamide in a minute and Gallic acid totally insoluble in Tween 20, Ethyl ether, Chloroform, and Toluen. Gallic acid incompletely soluble in Acetic acid, Acetonitrile, Tert-Butyl alcohol, and Distilled water. Yet, it was observed that after 14 hours Gallic acid was soluble in Acetonitrile, Tert-Butyl alcohol, and Distilled water. In the study of Lade et al. (2018) they have also performed similar kind of solubility test however, in that case they have used plant extract but in this case Thymol and Gallic acid pure compound were used. The solubility of compound assists to determine its utility in various drug and polymers complexation for pharmaceutical industries applications.

### 3.2. Synthesis and visual characterization of T-SNPs and GA-SNPs

The silver nanoparticles have been synthesized successfully using Gallic acid and Thymol has been synthesized by mixing 95 mL of 1 mM silver nitrate (AgNO₃) with the 5 mL of 1% Gallic acid and Thymol independently under sun light conditions. Similarly, Lade and Patel et al. (2017) have used sunlight conditions for silver nanoparticles synthesis. In contrast, Li et al. (2015) have prepared silver nanoparticles by Gallic acid. They have used 2 mL of AgNO₃ (20 mM) which was reduced with 2 mL of Gallic acid (10 mM) at room temperature. Also, there are no reports of silver nanoparticles synthesis using Thymol under sunlight conditions. The change in colour of solution from transparent to dark brown and black colour was observed and confirmed the formulation of T-SNPs and GA-SNPs. The synthesis of T-SNPs and GA-SNPs are confirmed preliminary by observing change in colour from colourless, became dark brown colour of Thymol and black colour of Gallic acid. Similar change in colour for silver nanoparticles synthesis was observed by Lade and Patel et al. (2017).

### 3.3. UV-Visible spectroscopy analysis of T-SNPs and GA-SNPs

The UV-Visible absorption spectrum of T-SNPs and GA-SNPs is showed in Fig. 1. It is seen from the Fig. 1, the 1% Thymol and 1% Gallic acid are sufficient to act as reducing agents to form silver nanoparticles. The formation of T-SNPs with absorbance of 0.7 A and GA-SNPs with absorbance of 0.6 A
using Thymol and Gallic acid showed the well defined absorbance band at 415 nm and 410 nm. The work of Li et al. (2015) suggested that the green synthesis of Gallic acid-coated silver nanoparticles UV-vis absorption spectrum of GA-SNPs shows typical and narrow absorption peak at approximately 400 nm. However, the Tonello et al. (2018) have reported the UV-visible absorption spectra of 0.25 mM Thymol at 550 nm with absorbance of 0.7 A.

3.4. FT-IR analysis of Thymol, T-SNPs, Gallic acid and GA-SNPs

3.4.1. FT-IR analysis Thymol and T-SNPs. In order to conclude the molecular interactions in the functional groups present in Thymol and silver AgNO₃ to form T-SNPs, the FT-IR analysis was carried out. The FTIR spectra of Thymol and T-SNPs are shown in Fig. 2 in the spectrum of Thymol, the Characteristic FTIR spectra were recorded with the functional groups as C=O Stretch (Halo Compound) 1058.675 cm⁻¹, C=N Stretch (Amine) 1241.722 cm⁻¹, C-O Stretch (Aromatic ester) 1285.521 cm⁻¹, O-H Stretch (Alcohol) 1517.732 cm⁻¹, N-O Stretch (Nitro compound) 1561.336 cm⁻¹, N-H Bending (Amine) 1584.503 cm⁻¹, C=C Stretch (α, β unsaturated ketone group) 1619.383 cm⁻¹, C-H Bending (aromatic compound) 1742.610 cm⁻¹, N=N=N Stretch (Azide compound) 2106.625 cm⁻¹, O-H Stretch (Carboxylic acid) 2869.304 cm⁻¹.

As seen from the spectrum of T-SNPs, structural changes occurred during the silver nanoparticles synthesis. The intermolecular interactions were observed in C=N Stretch (1058.675 cm⁻¹) and conjugation to strong C-F Stretch (1016.018 cm⁻¹). Also, the C=N Stretch (1156.54 cm⁻¹) is shifted to C-F Stretch (1113.295 cm⁻¹). The shifting of O-H Stretch (1419.674 cm⁻¹) from the Thymol was recorded to O-H Bending (1404.368 cm⁻¹) in synthesized silver nanoparticles. A complete transformation of band was observed in C=O Stretch and C-O Stretch Alkene (1638.056 cm⁻¹). The peak absorption at N=N=N Stretch Azide (2106.625 cm⁻¹) was changed to N=C=S Stretch Isothiocyanate (2010.053 cm⁻¹). The peak shift in O-H Stretch 2603.778 cm⁻¹ and 3173.427 cm⁻¹ to C=O Stretch (2205.04 cm⁻¹), and C-H Stretch (3324.369 cm⁻¹) was observed for T-SNPs.

3.4.2. FT-IR analysis Gallic acid and GA-SNPs. FTIR spectroscopy was used in order to characterize Gallic acid and GA-SNPs. As can be seen in Fig. 2, C-Cl Stretch are observed around 732.969- 789.011 cm⁻¹ are the Halo Compound of Gallic acid, C-N Stretch at 1017.853 cm⁻¹-1235.695 cm⁻¹ are associated with Amine group, additionally the characteristics band of O-H Bending are observed at 1305.573 cm⁻¹ are for Phenol group. The C-H Bending at 1438.510 cm⁻¹ to 1535.999 cm⁻¹ are specific for Alkane functional group in Gallic acid. Moreover, the C=O Stretch was recorded at 1611.774 cm⁻¹ corresponds to α, β Unsaturated ketone.

Chandra et al. (2018) suggested the absorption peak recorded at 1685, 1579, 1571 and 1481 cm⁻¹ was assigned for vibration stretch of C=C bonds in an aromatic group. Various peaks observed in 1313 – 1020 cm⁻¹ regions corresponded to C-O stretch vibration bond and bending vibration of O-H bond of phenol Gallic acid (GA).

The C-H Bending stretches from 1694.253 cm⁻¹ to 1993.963 cm⁻¹ are specific for Aromatic compound. C=O Stretch at wavelength 2117.161 cm⁻¹ is for Alkylene. The O-H Stretch observed at 2652.804 to 3063.549 cm⁻¹ belongs to Carboxylic acid and 3268.788 to 3469.614 cm⁻¹ is specific for Alcohol.

As seen from the spectrum of GA-SNPs, C=C stretching vibration band is observed around 1017.853 cm⁻¹, indication of the Amine groups of Gallic acid and is shifted to C-F Stretch at 1016.107 cm⁻¹ specific for Alkyl halide in silver nanoparticles. Furthermore, C=C Stretch stretching vibration band moved slightly to higher wave numbers, from 1611.774 cm⁻¹ for Gallic acid to 1637.501 cm⁻¹ for the silver nanoparticles. This kind of shift of such bands indicates physical interactions between Gallic acid and silver nitrate. Additionally, the C-H Bending (1914.176 cm⁻¹) belongs to Aromatic compound which are modified into C=C=C Stretch (1918.218 cm⁻¹) as allene, suggesting intermolecular interaction among themselves. Moreover, the C=O Stretch (2117.161 cm⁻¹) are modified to N=C=S Stretch (2083.241 cm⁻¹) as isothiocyanate. The peak shifted in O-H Stretch 3268.788 cm⁻¹, and 3343.388 cm⁻¹ to O-H Stretch (3266.524 cm⁻¹) and (3310.658 cm⁻¹) was observed for GA-SNPs.

The Park et al. (2016) indicates the appearance of broad band’s at 3422 cm⁻¹ for GA-AgNPs confirmed the existence of intermolecular hydrogen bonds of phenolic groups on the surface of the nanoparticles and Chandra et al. (2018) shows a broadband peak between 3552 and 2500 cm⁻¹ and the narrow peak at 1772 cm⁻¹ due to the stretching vibration of O-H (alcohol) group and a C-O, which confirms the presence of carboxyl group in the Gallic acid. This carboxyl group binds to the surface of the AgNPs. These bonds wholly disappeared after the nanoparticles synthesis.

Figure 1: The UV-visible absorption spectra of Thymol silver nanoparticles (T-SNPs) and Gallic acid silver nanoparticles (GA-SNPs)
3.5. Zeta potential analysis of T-SNPs and GA-SNPs

The surface charge, zeta deviation and conductivity on the synthesized T-SNPs and GA-SNPs were recorded using the Zeta Potential. The Zeta potential results are presented in Fig. 3. As is seen from the Fig. 3, average zeta potential, zeta deviation and conductivity value was found to be -19.7 mV, 9.70 mV, and 0.105 (mS/cm) for T-SNPs. However, the average zeta potential, zeta deviation and conductivity value for GA-SNPs was found to be -20.3 mV, 5.66 mV and 0.115 (mS/cm). Although, the negative values -19.7 and -20.3 detected as a more regular stability of T-SNPs and GA-SNPs respectively. In the study of Li et al. (2015) it is reported that the Gallic acid-coated silver nanoparticles had a negative zeta potential distribution of −10.5 mV and were spherical in shape. The hydrodynamic diameter of GA-AgNPs evaluated by DLS was 17.6 ± 5.3 nm.

3.6. Dynamic light scattering analysis of T-SNPs and GA-SNPs

The size and size distribution of particles of colloids T-SNPs and GA-SNPs were measured by Dynamic Light scattering (see Fig. 3) and found to be 123.2 nm and 121.1 nm respectively. Li et al. (2015) reported the hydrodynamic diameter of GA-AgNPs evaluated by Dynamic Light scattering analysis was 17.6 ± 5.3 nm and Abed et al. (2019) reported diameter of Thymol AgNP’s 152 nm.
3.7. X-ray diffraction of T-SNPs and GA-SNPs

The exact nature of the silver particles can be presumed from XRD patterns of the T-SNPs synthesized using Thymol. The XRD pattern of the T-SNPs (see Fig. 4) showed four intense peaks in the whole spectrum of 2 Ө values ranging from 20 to 80. The 20 peaks observed at 38.08°, 44.23°, 64.44°, and 77.37° corresponds to (1 1 1), (2 0 0), (2 2 0), and (3 1 1) reflection planes of an fcc lattice of silver (ICSD No. 98-018-0878), respectively (Kumar et al. 2017a, b) suggesting AgNO₃ complete reduction by Thymol. The 38.08° peak is showing high intensity due to preferred crystal orientation and the remaining peaks shows low intensity due to crystal arrangements in random manner. Generally, the broadening of peaks in the XRD patterns of samples signifies smaller particle size. From the XRD patterns, it indicates that the T-SNPs powders have large crystalline domain sizes in the mentioned concentration and correspond to pure Ag metal with face-centered cubic (fcc) symmetry. The other unassigned peaks could be due to the crystallization (bcc) of bioorganic phase that occurs on the surface of the nanoparticles.

Similarly, the XRD pattern of the GA-SNPs (see Fig. 4) showed four 20 peaks at 38.08°, 44.23°, 64.44°, and 77.37° that corresponds to (1 1 1), (2 0 0), (2 2 0), and (3 1 1) reflection planes of a fcc lattice of silver (ICSD No. 98-018-0878), respectively (Kumar et al. 2017a, b) suggesting AgNO₃ complete reduction by Gallic acid. Therefore, an XRD pattern concludes that the GA-SNPs powders have large crystalline domain sizes and correspond to pure Ag metal with fcc symmetry. The Crystallite size Dp of GA-SNPs and T-SNPs are shown in Table 1 and Table 2 using Scherer’s formula for Crystallite size Dp = K λ / (B cosθ). The average crystallite size from a sharp peak is estimated by using the Scherer’s formula. The average crystallite size for GA-SNPs was found to be 25.20 nm and T-SNPs were found to be 28.31 nm.

Figure 3: (a) Zeta potential and (b) Dynamic light scattering of Thymol silver nanoparticles (T-SNPs) and Gallic acid silver nanoparticles (GA-SNPs)

Figure 4: X-ray diffraction pattern of Thymol silver nanoparticles (T-SNPs) and Gallic acid silver nanoparticles (GA-SNPs) which corresponds for Bragg’s reflections planes of the metallic silver nanoparticles
Table 1: Shows the Crystallite size \( D_p \) of Gallic acid Silver nanoparticles.

| Sr. No. | Peak position 20 (°) | FWHM \( B_{\text{size}} \) (°) | \( D_p \) (nm) | \( D_p \) Average (nm) |
|---------|----------------------|-------------------------------|--------------|------------------|
| 1       | 22.74                | 0.26                          | 32.58        |                  |
| 2       | 29.198               | 0.265                         | 32.38        |                  |
| 3       | 35.73                | 0.50                          | 17.45        |                  |
| 4       | 37.861               | 0.458                         | 19.17        |                  |
| 5       | 39.14                | 0.44                          | 20.03        | 25.20            |
| 6       | 44.036               | 0.62                          | 14.45        |                  |
| 7       | 47.38                | 0.18                          | 50.38        |                  |
| 8       | 48.27                | 0.30                          | 30.33        |                  |
| 9       | 57.32                | 0.52                          | 18.20        |                  |
| 10      | 64.279               | 0.449                         | 21.84        |                  |
| 11      | 77.185               | 0.519                         | 20.47        |                  |

Table 2: Shows the Crystallite size \( D_p \) of Thymol Silver nanoparticles

| Sr. No. | Peak position 20 (°) | FWHM \( B_{\text{size}} \) (°) | \( D_p \) (nm) | \( D_p \) Average (nm) |
|---------|----------------------|-------------------------------|--------------|------------------|
| 1       | 23.09                | 0.26                          | 32.60        |                  |
| 2       | 29.402               | 0.249                         | 34.48        |                  |
| 3       | 35.93                | 0.46                          | 18.98        |                  |
| 4       | 38.081               | 0.452                         | 19.43        |                  |
| 5       | 39.40                | 0.31                          | 28.45        | 28.31            |
| 6       | 44.231               | 0.61                          | 14.69        |                  |
| 7       | 47.607               | 0.34                          | 26.69        |                  |
| 8       | 48.53                | 0.26                          | 35.03        |                  |
| 9       | 57.49                | 0.30                          | 31.57        |                  |
| 10      | 64.442               | 0.467                         | 21.02        |                  |
| 11      | 72.85                | 0.18                          | 57.33        |                  |
| 12      | 77.375               | 0.545                         | 19.52        |                  |

### 3.8 SEM analysis of T-SNPs and GA-SNPs

The SEM analysis of T-SNPs and GA-SNPs are presented in Fig. 5. The shape of T-SNPs is rod that ranges from 85.37 nm to 97.07 nm. However, GA-SNPs are spherical in shape and size ranges from 77.75 nm to 125.79 nm. Li et al. (2015) reported the GA-AgNPs were spherical in shape and nearly mono dispersed and the average size of GA-AgNPs was computed as 16.8 ± 4.8 nm. However, Abed et al. (2019) prepared the AgNPs using thymol and the thymol AgNPs spherical in shape.

Figure 5: Shows scanning electron microscopy micrograph of Thymol silver nanoparticles (T-SNPs) and Gallic acid silver nanoparticles (GA-SNPs)
3.9. Antioxidant activity of Thymol, T-SNPs, Gallic acid, and GA-SNPs

The significant antioxidant potential of Thymol, T-SNPs, Gallic acid, and GA-SNPs was evaluated by DPPH radical scavenging assay and the ascorbic acid was used as the standard drug. The IC50 values of Ascorbic acid, Thymol, T-SNPs, Gallic acid, and GA-SNPs was determined to be 19.47 µg/mL, 173.5 µg/mL, 112.1 µg/mL, 2.09 µg/mL, and 15.19 µg/mL respectively. The graphs showed percentage inhibition and DPPH free radical scavenging activity of Ascorbic acid, could be plotted and compared with Thymol, T-SNPs, Gallic acid, and GA-SNPs as shown in Table 3.

In present examination, the decrease capacity of DPPH radical was determined by the decrease in its absorbance at 517 nm, which is induced by different antioxidants. The decrease in absorbance of DPPH radical caused by antioxidants because of the reaction between antioxidant molecules and radical progress which results in the scavenging of the radical by hydrogen donation. It can be declared that T-SNPs showed a more effective hydrogen donating capacity (IC50 - 112.1 µg/mL) than pure Thymol compound. However, reverse was observed in case of Gallic acid, the pure Gallic acid showed more effective hydrogen donating capacity (IC50 - 2.09 µg/mL) than GA-SNPs. Moreover, it can be concluded that pure Gallic acid more effective hydrogen donating capacity (IC50 - 2.09 µg/mL) than pure standard Ascorbic acid (IC50 - 19.47 µg/mL).

In the study of Bhadoriya et al. (2012) it was reported that isolated Gallic acid showed significant in vitro free radical scavenging activity; at the concentration level of 20 µg/mL-1, 61.45±0.44 % and 52.94±0.67 % inhibition of free radical was observed for ABTS and DPPH assay respectively. While 95.22±0.71 % and 91.99±0.59 % inhibition of free radical was observed for ABTS and DPPH assay respectively at the concentration of 50µg/mL-1. The Esmaili et al. (2011) has reported the antioxidant activity of Thymol using stable radical scavenging 2, 2’-Diphenyl-1-picrylhydrazyl (DPPH). The value of IC50 in this test was equal to 36.1±0.18 which showed good effects.

3.10. Antimicrobial activity of Thymol and Gallic acid

The antimicrobial activity of Thymol and Gallic acid at specific concentrations were used to check the zone of inhibition values against E. coli, Klebsiella, S. aureus, and S. pasteurii. The Fig. 6 and 7 showed the antimicrobial activity of Thymol against S. aureus, S. pasteurii, E. coli and Klebsiella. It was observed that the Thymol showed the maximum zone of inhibition of 12 mm, 14 mm, and 14 mm against the bacterium Klebsiella at the concentration of thymol 200 mg/ml, 400 mg/ml, and 600 mg/ml. In 800 mg/ml and 1000 mg/ml concentration of thymol showed the zone of inhibition of 16 mm and 17 mm against E. coli pathogen. Thymol shows the more antimicrobial activity than Gallic acid and when increases the concentration of Thymol and Gallic acid increased the zone of inhibition. The concentration of Gallic acid 200 mg/ml and 400 mg/ml showed the maximum zone of inhibition of 4 mm and 6 mm against the S. pasteurii and 600 mg/ml concentration showed 8 mm zone against S. aureus. The E. coli and S. aureus showed zone of inhibition of 8 mm and 11 mm at concentration 800 mg/ml and 1000 mg/ml.

A report from Olasupo et al. (2003) revealed the antibacterial effect of thymol with the lowest minimum inhibitory concentration (MIC) values of 1.0 mmol/L (S. typhimurium) and 1.2 mmol/L (Escherichia coli). Thymol (200 mg/ml) could inhibit the growth of E. coli by inducing the permeabilization and depolarization of the cytoplasmic membrane (Xu et al. 2008).

3.11. Antimicrobial activity of T-SNPs and GA-SNPs

The T-SNPs and GA-SNPs at specific concentrations (60, 70, 80, 90, and 100 µl) were used to check the zone of inhibition values against E. coli, Klebsiella, S. aureus, and S. pasteurii. The zone of inhibition values for T-SNPs and GA-SNPs against E. coli, Klebsiella, S. aureus, and S. pasteurii are presented in Fig. 8 and 9. The T-SNPs showed the maximum zone of inhibition of 4 mm, 6 mm, 8 mm, 10 mm, and 12 mm against the E. coli pathogen at the T-SNPs concentration of 60 µl, 70 µl, 80 µl, 90 µl and 100 µl. The T-SNPs all showed the zone of inhibition in E. coli other than Klebsiella, S. aureus, and S. pasteurii pathogens.

The GA-SNPs concentration 60 µl and 70 µl showed the maximum zone of inhibition of 2 mm and 4 mm against S. aureus and E.coli. The 80 µl, 90 µl and 100 µl concentration of GA-SNPs showed maximum zone of 5 mm, 6 mm and 8 mm against S. aureus pathogen. The T-SNPs show the more antimicrobial activity than GA-SNPs.

Similar to our work, Li et al. (2015) reported the antimicrobial activity of GA-AgNPs against two bacterial strains, Escherichia coli (E. coli) and Staphylococcus aureus (S. aureus), and one fungus, Candida albicans (C. albicans), with minimum inhibitory concentrations of 6, 30 and 24 µg/mL completely inhibited the growth of E. coli, S. aureus and C. albicans, respectively.

Figure 6: Graphical representation of antimicrobial activity of Thymol and Gallic acid in concentration of 0, 200, 400, 600, 800, and 1000 mg/mL against each of the four bacterial pathogens (S. aureus, S. pasteurii, E.coli, and Klebsiella)
Figure 7: Agar well diffusion technique demonstrates the zone of inhibition of Thymol and Gallic acid in concentration of 0, 200, 400, 600, 800, and 1000 mg/mL against each of the four bacterial pathogens (S. aureus, S. pasteurii, E.coli, and Klebsiella).

Figure 8: Graphical representation of antimicrobial activity of Thymol silver nanoparticles (T-SNPs) and Gallic acid silver nanoparticles (GA-SNPs) in concentration of 60, 70, 80, 90, and 100 µL against S. aureus, S. pasteurii, E.coli, and Klebsiella.

Figure 9: Agar well diffusion technique demonstrates zone of inhibition of Thymol silver nanoparticles (T-SNPs) and Gallic acid silver nanoparticles (GA-SNPs) in concentration of 60, 70, 80, 90, and 100 µL against S. aureus, S. pasteurii, E.coli, and Klebsiella.
Table 3: DPPH radical scavenging activity of Ascorbic acid, Thymol, T-SNPs, Gallic acid, and GA-SNPs at various concentrations

| Sample       | Concentration in µg/mL | % Inhibition | IC50 value in µg/mL |
|--------------|------------------------|--------------|---------------------|
| Ascorbic acid| 5                      | 25.0         |                     |
|              | 10                     | 27.7         |                     |
|              | 15                     | 54.8         | 19.47               |
|              | 25                     | 58.7         |                     |
|              | 50                     | 96.1         |                     |
| Thymol       | 5                      | 2.26         |                     |
|              | 10                     | 4.53         |                     |
|              | 15                     | 7.31         | 173.5               |
|              | 25                     | 10.30        |                     |
|              | 50                     | 15.15        |                     |
| T-SNPs       | 5                      | 13.29        |                     |
|              | 10                     | 15.56        | 112.1               |
|              | 15                     | 20.20        |                     |
|              | 25                     | 22.37        |                     |
|              | 50                     | 28.76        |                     |
| Gallic acid  | 5                      | 32.47        |                     |
|              | 10                     | 57.42        |                     |
|              | 15                     | 82.78        | 2.09                |
|              | 25                     | 93.60        |                     |
|              | 50                     | 97.57        |                     |
| GA-SNPs      | 5                      | 18.55        |                     |
|              | 10                     | 37.11        |                     |
|              | 15                     | 59.69        | 15.19               |
|              | 25                     | 87.42        |                     |
|              | 50                     | 91.95        |                     |

3.12. Visual appearance of films

Visual appearance and colour of the film as is shown in the Fig. 10. The C+G film was appeared as clear, glossy, smooth, and without any tents. Whereas, the visual appearance of C+G/GA/T-SNPs film was observed as light brown colour. Kaya et al. (2018) have reports also, explained the same phenomena, that adding plant extracts to chitosan matrix and changed the film’s colour.

Figure 10: Visual appearance of the chitosan control (C+G) film and the formulated chitosan blended (C+G/GA/T-SNPs) film
3.13. Determination of Thickness

Thickness of C+G film and C+G/GA/T-SNPs film was measured at six different points and average thickness with standard deviation was calculated. The thicknesses of C+G film and C+G/GA/T-SNPs film are 0.1 mm and 0.12 mm. The data of film thickness indicates that there was slight difference in thickness within the formulations.

3.14. Folding endurance

Folding endurance value for C+G film was found to be 32 folds. It indicates that C+G film had not much satisfactory film flexibility. In contrast, C+G/GA/T-SNPs film was found to be more than 200 folds that indicate film had good flexibility in C+G/GA/T-SNPs film.

3.15. Biodegradability of film

Fig. 11 shows the water solubility of the C+G film was recorded as 20% in 72 h. Incorparations of Gallic acid extract and T-SNPs into the films demonstrated decrease the film solubility as 12% for C+G/GA/T-SNPs film. The strong intermolecular interactions between Gallic acid and T-SNPs and chitosan matrix resulted in lower degradation values into the C+G/GA/T-SNPs film. Lower solubility rate for the C+G/GA/T-SNPs film was probably due to the faster diffusion of Gallic acid and T-SNPs in water.

Fig. 12 shows that the soil degradation rates in 15 days were recorded as 100% for C+G film and 45% for C+G/GA/T-SNPs film. Then after 30 days C+G/GA/T-SNPs film was 100% Soil degradation. The C+G film exhibited higher soil degradability; however, lower soil degradability was observed for the C+G/GA/T-SNPs film. Kaya et al. (2018) reported that the water solubility of the chitosan control film was recorded as 24% in 48 h and Soil degradation rates in 15 days were recorded as 89.04% for chitosan control film.

Figure 11: Biodegradability of films in water (a) chitosan control (C+G) film, (b) chitosan blended (C+G/GA/T-SNPs) film, and (c) chitosan blended (C+G/GA/T-SNPs) film shows the diffusion of Gallic acid and T-SNPs in water

Figure 12: Biodegradability of chitosan control (C+G) film and chitosan blended (C+G/GA/T-SNPs) film in soil.
3.16. Surface pH

Surface pH of the chitosan control (C+G) film and chitosan blended (C+G/GA/T-SNPs) film were found to have pH between 6-7. Hence the chances of irritation are very less.

3.17. FTIR analysis

Fig. 13 shows the FTIR analysis of the pure chitosan, C+G film and C+G/GA/T-SNPs film was carried out to identify the primary functional groups of the compounds. The pure chitosan FTIR spectrum is very wavy and zigzag. Also, it consists of large numbers of peaks. The FTIR spectrums of C+G film suggest that there is good molecular interaction between Chitosan and Glycerol. Moreover, Glycerol acts as good plasterer to chitosan as it reduces the wavy and zigzag to smooth in flow. The C+G/GA/T-SNPs film FTIR spectrum shows that the functional group between 1800-1200 could have taken part for bending bonds, bonding bonds, modifications and interactions to form the stable film. The Fig. 13 shows the FTIR spectra of Chitosan pure, chitosan control C+G film and C+G/GA/T-SNPs film.

![FTIR spectra](image1)

**Figure 13:** FTIR spectra of the pure chitosan, chitosan control (C+G) film, and chitosan blended (C+G/GA/T-SNPs) film were carried out to identify the primary functional groups of the compounds.

It can be seen, that the FTIR of chitosan suggest the functional groups as C=C Bending for Alkene (893.893 cm⁻¹), C-N Stretch for Amine (1021.901 cm⁻¹, 1060.042 cm⁻¹, 1149.553 cm⁻¹), C-O Stretch for Aromatic ester (1308.695 cm⁻¹), O-H bending for Phenol (1373.769 cm⁻¹, 1420.104 cm⁻¹), C≡C Stretch for Alkene (1645.897 cm⁻¹), C=C=C Stretch is Allene (1916.665 cm⁻¹, 1950.513 cm⁻¹, 1998.658 cm⁻¹), the C≡C Stretch at 2104.814 cm⁻¹, 2181.125 cm⁻¹ indicates Alkyne, C≡N Stretch is for Aliphatic primary amine group at 3359.641 cm⁻¹.

It can be seen, that the FTIR C+G film and C+G/GA/T-SNPs film indicates molecular interaction between functional group as C-H Bending for Alkene (922.770 cm⁻¹), S=O Stretch of Sulfoxide (1033.014 cm⁻¹), C-N Stretch for Amine (1244.148 cm⁻¹), N≡C≡S Stretch for Isothiocyanate (2005.769 cm⁻¹), and C-H Stretch for Alkane (2933.718 cm⁻¹), O-H Stretch Alcohol (3265.019) between Glycerol and chitosan. The interface at the C-O Stretch which is specific for Tertiary alcohol (1152.707 cm⁻¹) is modified to C-N Stretch known for Amine group (1155.370 cm⁻¹). The functional group S=O Stretch for Sulfone (1323.122 cm⁻¹) may be displaced by C-N Stretch Aromatic amine (1334.076 cm⁻¹). In specific, O-H Bending which is Phenol (1379.746 cm⁻¹) is modified to N-O Stretch Nitro compound.
The peak shifting was observed in C=C Stretch of Cyclic alkene (1566.704 cm⁻¹) to C=C Stretch (1535.832 cm⁻¹). The peak shifting was observed in C=C=C Stretch of Allene (1924.540 cm⁻¹) to C≡C ≡C Stretch (1914.544 cm⁻¹). Moreover, the C≡C≡C Stretch Allene (1804.119 cm⁻¹) is transformed to N=C=S Stretch Isothiocyanate (2004.119 cm⁻¹). The complete transformation of N≡C≡S Stretch Isothiocyanate (2114.509 cm⁻¹) to C≡C Stretch Alkyne (2207.818 cm⁻¹) and C-H Stretch Alkane (2888.438 cm⁻¹) into O=C=O Stretch Carbon dioxide (2349.779 cm⁻¹) was observed from spectrum.

Kaya et al. (2018) were recorded chitosan peaks as –OH bonds (3288.10 cm⁻¹), C–H aliphatic stretching vibration (2873.78 cm⁻¹), amide I (1651.51 cm⁻¹), amide II (1557.79 cm⁻¹), amide III (1374.07 cm⁻¹) and C–O stretching vibration (1025.51 cm⁻¹). The spectrum of chitosan film, structural changes occurred during the film formation. Addition of the glycerol to the film matrix (chitosan, acetic acid, and water), affected the intermolecular interactions resulted in expand in the OH bond and increase in the intensity of amide I, II, and III bands.

### 3.18. X-Ray diffraction

The crystalline structures of the C+G film and C+G/GA/T-SNPs film were characterized by X-ray diffraction (XRD) see Fig. 14. Both the films have observed similar diffraction patterns, with a broad peak at 21.66° and the peak is higher than 20 2-theta (deg). Indication of peak to peak in curve suggests the formation of phase or phases in the composite during polymerization process. Compressive strain to lattice structure, lattice size 20 2-theta (deg) increases. The pattern shows a complete broadening in all 2 theta values as it is nano material, the peaks are broadened. The silver nanoparticles must be embedded inside of the C+G film. Cui et al. (2018) have reported x-ray diffraction patterns of the chitosan film has three characteristic absorption peaks at 2θ = 10.22°, 19.9°, and 20.5°.

### 3.19. SEM analysis

The surface characterization of C+G film and C+G/GA/T-SNPs film are presented in Fig. 15. As is seen from the figure, surface morphology of C+G film was observed as small pores present in film and T-SNP are seen embedded in C+G/GA/T-SNPs film which is visible and are present in surface of film. Kaya et al. (2018) reported that the chitosan-control was observed as smooth, compact, and without any crack or pores.

### 3.20. Antioxidant activity

The significant antioxidant potential of C+G film and C+G/GA/T-SNPs film was evaluated by DPPH radical scavenging assay and the ascorbic acid was used as the standard drug. The IC50 values of C+G film and C+G/GA/T-SNPs film were determined to be 32.85 µg/mL and 27.12 µg/mL respectively. The graphs showed percentage inhibition and DPPH free radical scavenging activity of C+G film and C+G/GA/T-SNPs film as shown in Fig. 16 (a and b) and Table 4. Kaya et al. (2018) reported that the antioxidant activities of chitosan control film were measured as 37.21±2.07%.

**Figure 14:** The X-ray diffraction pattern of chitosan control (C+G) film and chitosan blended (C+G/GA/T-SNPs) film

**Figure 15:** Shows scanning electron microscopy micrograph of chitosan control (C+G) film and chitosan blended (C+G/GA/T-SNPs) film
Figure 16: Shows the DPPH free radical scavenging activity of (a) chitosan control (C+G) film and (b) chitosan blended (C+G/GA/T-SNPs) film at various concentrations.

Table 4: DPPH radical scavenging activity of C+G film and C+G/GA/T-SNPs film

| Sample                  | Concentration in µg/mL | % Inhibition | IC50 value in µg/mL |
|-------------------------|------------------------|--------------|---------------------|
| C+G film                | 5                      | 10.23        |                     |
|                         | 10                     | 19.18        |                     |
|                         | 15                     | 33.42        | 32.85               |
|                         | 25                     | 46.68        |                     |
|                         | 50                     | 67.40        |                     |
| C+G/GA/T-SNPs film      | 5                      | 14.22        |                     |
|                         | 10                     | 25.56        |                     |
|                         | 15                     | 41.18        | 27.12               |
|                         | 25                     | 54.56        |                     |
|                         | 50                     | 75.11        |                     |

3.21. In-vitro antimicrobial activity

The zone of inhibition values for C+G film and C+G/GA/T-SNPs film against *E. coli*, *Klebsiella*, *S. aureus*, and *S. pasteurii* are presented in Fig. 17 and 18. The zone of inhibition of the C+G/GA/T-SNPs film is found to be very high as compared with the C+G film. In specific, C+G film shows ZOI of 6 mm for *S. aureus*, 4 mm for *S. pasteurii*, 5 mm for *E. coli*, and 4 mm for *Klebsiella*. However, C+G/GA/T-SNPs film shows the zone of inhibition of 14 mm for *S. aureus*, 18 mm for *S. pasteurii*, 13 mm for *E. coli* and 11 mm for *Klebsiella*. It is also noted that the C+G/GA/T-SNPs film shows more inhibition against Gram Positive Bacteria (*S. aureus* and *S. pasteurii*) than Gram Negative Bacteria (*E. coli* and *Klebsiella*). Kaya et al. (2018) reported that chitosan control film showed lower inhibition zone than blended films. The incorporating of the plant extract into chitosan films provided higher antimicrobial activity. The chitosan blend films with the plant extract were found to be effective against especially Gram-negative bacteria such as *P. microbilis*, *P. vulgaris*, *P. Aeruginosa*, and *E. coli*.

Figure 17: Graphical representation of antimicrobial activity of chitosan control (C+G) film and chitosan blended (C+G/GA/T-SNPs) film against each of the four bacterial pathogens (*S. aureus*, *S. pasteurii*, *E. coli*, and *Klebsiella*)
4. CONCLUSION

This study explored potential aspects of Thymol, Gallic acid and T-SNPs, GA-SNPs. The T-SNPs and GA-SNPs are synthesized successfully using Thymol and Gallic acid under sunlight condition at pH 9 and 8.5. The UV-spectroscopic analysis confirms the T-SNPs and GA-SNPs at 415 and 410 nm. The FTIR determines the involvement of functional groups that have the potential to reduce and stabilized the T-SNPs and GA-SNPs. The surface charge of T-SNPs is -19.7 mV and GA-SNPs are -20.3 mV. The IC50 values of Thymol, T-SNPs, Gallic acid, and GA-SNPs conclude that T-SNPs showed a more effective hydrogen donating capacity than pure Thymol compound. The pure Gallic acid showed more effective hydrogen donating capacity than GA-SNPs. The antimicrobial potential of Thymol and Gallic acid against E.coli, Klebsiella, S. aureus, and S. pasteurii. Thymol and Gallic acid showed high antibacterial activity against Gram-positive (S. aureus and S. Pasteurii) and Gram-negative (E. coli and Klebsiella) pathogens. Higher concentration Thymol and Gallic acid resulted in an increase in the zone of inhibition. Thymol and T-SNPs showed the more antimicrobial activity than Gallic acid and GA-SNPs against E. coli, Klebsiella, S. aureus, and S. pasteurii pathogens. Thus T-SNPs could an alternative in pharmaceutical industry for the treatment of skin infection and anesthetic for medical filed. The pure Gallic acid and T-SNPs with strong antioxidant activity may be good alternative for the developing the wound healing dressing material in pharmaceutical industry. Further, the Gallic acid (highly antioxidant) and T-SNPs (intense antimicrobial) and were transferred into C+G films. As a result, high antioxidantive and antimicrobial C+G/GA/T-SNPs films (against four pathogens) were successfully produced and well characterized with commonly used analytical techniques. Additionally, the glycerol enhanced the plasticity of chitosan and the Gallic acid enhance the biodegradability time of the C+G/GA/T-SNP films.

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Conflict of interest

The authors declare that they have no conflict of interest in the publication.

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