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Reinforcement of biodegradable SiO₂NPs-modified cellulose-gelatin hydrogel films with antioxidant and antibacterial properties as potential food packaging composite

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Abstract

This study proposed an innovative approach to the development of sustainable and biodegradable food packaging materials by incorporating inexpensive nano-silica (SiO_2NP_s) in designed hydrogel (CSG) film employing biodegradable polymers: synthetic polymer polyvinyl alcohol (PVA), natural polymer - carboxymethyl cellulose (CMC) and protein-based bio-polymer -gelatine, and a commercial crosslinker, tetraethoxysilane (TEOS) through a conventional air-dry casting technique. The CSG hydrogel blends were modified with varying amounts of SiO₂NPs (0.05g, 0.1g, 0.15g and 0.2g) and compared with the blend without SiO₂NPs to determine the effect of SiO₂NPs loading through various characterisation techniques and applications including antioxidant and antimicrobial activity. Comprehensive characterizations of the CSG films revealed that CSG 0.1 (containing 0.1g SiO_2NPs) exhibited the most favourable functional properties, low crystallinity, high flexibility, suitable pore size, thermal stability, adequate tensile strength, elongation at the breaking point and maximum stability by swelling and diffusion test. The addition of SiO₂NPs consistently enhanced thermal and mechanical stability in all CSG films. Further, these CSG films were implemented for antioxidant test and antimicrobial activity against gram-positive Bacillus cereus and gram-negative Escherichia coli. SiO₂NPs integration significantly elevated the antioxidant capacity in all films, with CSG 0.1 showing $\sim 7\%$ improvement. The antimicrobial activity of SiO₂NPs-modified CSG films was also notable, with CSG 0.1 effectively inhibiting B. cereus by 1.2cm zone and E. coli by 0.5cm zone. A soil burial test was performed to pattern the biodegradability of CSG hydrogels. Therefore, the outstanding improvements in the intrinsic properties of CSG films, owing to SiO₂NPs modification, positioned these CSG hydrogels as promising candidates for advanced food packaging materials in various industries.

Keywords Nano-reinforcement · Hydrogel film formation · Biodegradability · Antibacterial activity · Antioxidant activity · Food packaging material

Abbreviations

CSG	Nano silica in designed hydrogel
PVA	Poly vinyl alcohol

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CMC	Carboxymethylcellulose
TEOS	Tetraethoxysilane
XRD	X-ray Diffraction technique
SEM	Scanning electron microscope
TGA	Thermogravimetric analysis
DTA	Differential Thermal analysis
UTS	Ultimate Tensile Strength
DCPIP	2,6-Dichlorophenolindophenol

Introduction

Exploration of sustainable food packaging materials is one of the high demand research areas globally due to the recent sustainable movement in packaging industries. Conventional packaging material containing plastics exhibits detrimental characteristics comprising hazardous chemicals and degradation into micro and nano-plastic causing serious contamination of soil and water bodies and also posing challenges for proper disposal/plastic waste management. Consequently, in the quest for more sustainable alternatives, materials such as textiles and paper have emerged as viable options for packaging though these options have numerous limitations such as easy penetration of liquid and less stability. On the other hand, glass or metal-based packaging is heavy-weight and requires high processing costs for production. Therefore, biodegradable polymer-based packaging materials are highly desired due to their advantageous characteristics (e.g., transparency, low weight, tear resistance, stiffness, the ability to be air and heat-sealed, and a high strength-to-weight ratio) to resolve environmental concerns as well as economical [1–4]. The food industries have also embraced the application of nanotechnology by utilizing engineered nanoparticles to enhance the quality of their products by enhancing the performance of polymers for food packaging applications by improving their mechanical strength, swelling behaviour, thermal characteristics, and biodegradability [4]. Among the advancements, hydrogels comprising polymers have emerged as a notable option due to their ability to absorb water and other liquids, reaching a maximum capacity of their weight [4, 5]. Therefore, this study proposed a novel SiO₂NPs in designed hydrogel (CSG) film as an active food packaging material.Biodegradable synthetic polymers PVA, along with natural polymer CMC played a significant role as important and biocompatible materials in food packaging industries for decades due to their sustainable and antimicrobial characteristics [6–8].CMC-containing carboxymethyl groups (-CH₂-COOH) are attached to certain hydroxyl (-OH) groups of the glucopyranose monomers comprising the cellulose backbone and serve as a viscosity modifier or thickener in food products and function as an emulsion stabilizer in various goods (including ice cream), non-food items such as toothpaste, laxatives, diet pills, water-based paints, detergents, textile sizing, reusable heat packs, and various paper products [6]. Gelatin, derived primarily from bovine collagen obtained from animal skin, bones, and connective tissues, results in a collection of peptides and proteins and is used as a gelling agent in various industries including food, beverages, pharmaceuticals, vitamin and medication capsules, photographic films and papers, and cosmetics [9, 10]. SiO₂NPs, also known as silica nanoparticles or nano-silica, possess favourable characteristics such as good biocompatibility, low toxicity, thermal stability, and ease of synthesis, making them promising for various biological applications and a high potential candidate for food packaging material [11, 12]. TEOS is the ethyl ester of orthosilicic acid, Si(OH)₄ and is predominantly used in the semiconductor as well as packaging industries as

a crosslinking agent in silicone polymers and as a precursor for SiO₂ and various applications such as coatings for carpets, production of aerogel as an additive in alcohol-based rocket fuels to reduce heat flow to the chamber wall in regeneratively cooled engines [13–15]. In this study, the preparation of CSG films utilising biodegradable polymers: synthetic polymer - PVA, natural polymer - CMC and protein-based biopolymer - gelatine, nano-compound - SiO₂NPs and a commercial crosslinker, TEOS were achieved through a simple conventional air-dry casting technique as sustainable and biodegradable food packaging material. This study aims to fabricate SiO₂NP-modified hydrogel-based CSG films as active food packaging materials addressing essential needs related to health, environment, and food industries. The objective of this research is to develop a novel polymeric film, SiO₂NP-modified CSG film for food packaging purposes which consists of non-toxic polymers and possesses biodegradable properties, allowing for easy decomposition. There have been very limited studies performed on engineered hydrogel composite for active food packaging in the literature so far. A related experimental study on engineered gelatine/ cellulose-silver nanoparticles (AgNPs) modified nanocomposite presented a high potential of antibacterial activities against E. coli and B. cereus as active food packaging material [2]. Compared to earlier studies, the material composition for the current study is unique, readily available and inexpensive compared to expensive materials like AgNPs, and carbon nanotubes (CNTs) [2, 16]. Also, the antibacterial (against E. coli and B. cereus) and antioxidant (against free radical of 2,6-dichlorophenolindophenol (DCPIP)) activities of the proposed active composite have been explored in this study. Then, the novelty of this study is to fabricate a new blend of composite material, SiO₂NP-modified CSG film containing gelatin-PVA-CMC-SiO2NPs-TEOS as active food packaging material and explore the improvement of intrinsic properties as well as antibacterial and antioxidant properties. The thermal and mechanical stability, swelling and diffusion characteristics and micro graphical orientation of this novel SiO₂NP-modified CSG film have been explored comprehensively in this study to determine whether this composite will be competitive for commercial packaging materials or not.

Materials and Methods

Materials

PVA (purity: >98%), CMC (purity: >98%), TEOS (purity: 98.5%), gelatin, SiO₂ NPs (purity: >99.98%), ethanol (100%), and 2, 2-dichloro phenol indophenol

(DCPIP) (100%), as vitamin C source namely ascorbic acid) were purchased from Sigma-Aldrich, United Kingdom.

Synthesis Methods of Hydrogel Film Preparation

The hydrogel film preparation has been used in this study is casting and blending method since it is well-established, user friendly, cost-effective method and all the control parameters can be maintained precisely throughout this technique [14]. To prepare the composite film, 0.1g of PVA was dissolved in 30mL of preheated distilled water through continuous stirring on a hot plate with a magnetic stirrer at a temperature of 60°C until complete dissolution. Similarly, 0.4g of gelatine and 0.6g of CMC were separately dissolved in 30mL and 40mL of distilled water, respectively, using continuous stirring on a hot plate at 60°C with a magnetic stirrer until complete dissolution. Subsequently, the PVA solution was added to the gelatine solution on the hot plate and mixed for 1h. After the 1h blending period, this PVA-gelatine solution was added to the CMC solution and blended for an additional 1h at 60°C. SiO₂NPs were dispersed in 30mL of distilled water and subjected to sonication using a digital ultrasonic cleaner for 0.5h at room temperature. The sonicated SiO₂NPs were then added to the PVA/CMC/ gelatine solution and mixed for 0.5h. Furthermore, 100µL of TEOS (dissolved in 20mL of ethanol (C₂H₅OH)) was added dropwise as a crosslinker, followed by stirring for 3h at 60°C. The prepared solution was poured into Petri dishes to obtain film samples, which were then dried at room temperature for the next 3h. The dried film samples were peeled and placed in a desiccator within polythene packets for further study. Additionally, a control sample of the film was synthesized for comparison purposes. The code and composition of the synthesized hydrogel base film samples are given in Table 1. The schematic representation of preparing hydrogel film is given in Fig. 1.

Table 1 The codes and compositions of PVA/CMC/gelatin films

Sample Codes	PVA(g)	CMC(g)	Gelatin(g)	SiO ₂ NPs(g)	TEOS(µL)
CSG	0.1	0.6	0.4	0	100
CSG 0.05	0.1	0.6	0.4	0.05	100
CSG 0.1	0.1	0.6	0.4	0.10	100
CSG 0.15	0.1	0.6	0.4	0.15	100
CSG 0.2	0.1	0.6	0.4	0.20	100

Characterizations

Fourier Transforms Infrared Spectroscopy (FTIR)

Fourier transform infrared spectrometry was used to investigate the functional groups of the hydrogels. The scanning was performed by Agilent Technologies Cary 630 FTIR with the range of wavelength from 4000-650cm⁻¹ and the resolution was 4.0cm⁻¹.

X-Ray Diffraction (XRD) Techniques

XRD analysis of hydrogel film samples was performed by an X-ray Diffraction machine (STOE, Japan) with scan step: 0.25, scan speed: 1°/minute and scan angle: 10-80 at 20. XRD analysis is useful for the estimation of the structure characterization of thin films. X-ray diffraction gives the material's molecular structure and estimates the dispersion of doped particles in the structure of the sample.

Scanning Electron Microscopy (SEM)

The samples were gold (Au) sputtered to make the samples conductive for the electron beam interaction. SEM analysis was performed to observe the surface morphology of synthesized hydrogel film samples. Scanning Electron Microscope (SEM, JSM, JEOL, Japan) was used to investigate the prepared dried films. Images were collected by using a scanning electron microscope at the magnification of 50 and 20µm.

Thermogravimetric/Differential Thermal Analysis (TG/DTA)

Thermogravimetric analysis was performed to estimate the thermal behaviour of the CSG hydrogel film series. The thermal decomposition activity of samples was investigated in an air environment with and heating rate of 10°C/min from 29 to 816°C with a TG/DTG instrument (Perkin Elmer Diamond TG/DTA instrument, Japan).

Ultimate Testing Machine (UTM)

The ultimate tensile strength of the designed nanohydrogel film series (CSG) was measured by a universal testing machine (AG-20KNXD Plus, Shimadzu). The hydrogel slices had a 2.5-inch length and breadth of 0.5 inches. The testing method was in accord with ASTM D412-98A. The gage length (1.60 inches) as well as the speed of the crosshead 5.0mm/min was set. Ultimate tensile strength was calculated in MPa simply by dividing the force (F) to split the film over the film sample's area (A). It also measured the percentage elongation at break (in % E) and estimated the average of at least three UTS determinations and percentage E.



Fig. 1 Schematic representation of CSG hydrogel film preparation

Swelling Test (ST)

In distilled water, the biodegradable hydrogel base film sample's swelling testing was examined. Cut the dried film sample (0.025g pre-weighted) into small pieces from a plastic petri dish and then load that dish with 80mL distilled water. The abundance of water weight of the film was measured after each 10min. The swollen sample was then placed in the water again till the establishment of equilibrium occurred. All experiments were repeated into triplets and the average results have been considered. By assigned relation, the swelling ratio for samples was calculated by Eq. (1).

Degree of Swelling =
$$\frac{Ws - Wd}{Wd}$$
 (1)

Where, $W_s = Swollen$ weight of the film and $W_d = Dry$ weight of the film

The swelling mechanism of the CSG series for water diffusion was evaluated by Eq. (2).

$$F = kt^n \tag{2}$$

Where F represents the fractional swelling at time t (min) that is calculated by the swelling ratio of W_t and W_{eq} . W_{eq}

and W_t are equilibrium swelling ratios at time t and k is the swelling rate constant, while n is the diffusion exponent that helps to find the release mechanism. The degree of swelling of the CSG series was used to analyze the values of n and k.

Applications of the Hydrogel Film

Antioxidant Activity

The standard solution of DCPIP was prepared by taking 0.024g of salt in a piece of aluminium foil in the dark and dissolved in 50mL C_2H_5OH . The prepared solution is then transferred to a 100mL volumetric flask and up to the mark the volume with absolute ethanol, this solution will be the standard solution of 1000µg/mL. The absorbance of the control was analyzed at 517nm. The standard antioxidant (Vitamin C) with different concentrations: i.e. 50,100,150,200,250,300 and 350µg/mL. The solutions were incubated for 30min at 25°C in the dark. The inhibition rate was measured at 517nm. All experiments were repeated into triplets and the average results have been considered. The antioxidant rate was determined by Eq. (3).

% Inhibition =
$$\left(\frac{Abs \ of \ control - Abs \ of \ sample}{Abs \ of \ control}\right) \times 100$$
(3)

The sample solutions of the hydrogels were prepared by considering a circular diameter of 3cm in 10mL of methanol (0.1mM in methanol) and shaking them in a water bath for 1h. 1mL of prepared hydrogel sample was mixed with 3mL of DCPIP. After 30min of keeping in the dark, the absorbance of all the standard (vitamin C) and hydrogel samples was measured at 517nm. The scavenging % was determined by Eq. (4) [17].

% Scavenging Activity =
$$\left(\frac{Abs \ of \ blank - Abs \ of \ sample}{Abs \ of \ blank}\right) \times 100$$
(4)

Antimicrobial Activity

The antibacterial activity was analyzed by the agar disc diffusion method. A nutritional medium for the development of bacterial strains was prepared to test the antibacterial properties of hydrogels. The agar solution was prepared following the standard method while agar was dissolved in distilled water was shaken thoroughly for proper mixing. The agar solution-containing bottle was covered with aluminium foil and autoclaved to sterilize the agar, ensuring that no other microorganisms except the selected one could thrive. The autoclave was performed at 120°C for 20min. After 20min of autoclave, once the temperature was down to 60°C, the agar solution bottle was placed under laminar flow. Following that, agar was placed into the plates, letting them cool down to set on the plates. Sanitized cotton swabs were used to spread a little bacterial culture across the plates. On the plates, two different bacterial strains, gram-positive B. cereus and gram-negative E. coli were grown in separate petri dishes with the use of a sterile cotton swab. Using sterile tweezers, five disc-shaped hydrogels were placed on the agar plates. The discs' diameter was maintained at 6mm. Plates were kept upside down at 37°C for around 24h while being maintained culture growth. The limiting area of the culture plates was determined by measuring them [18]. All experiments were repeated into triplets and the average results have been considered.

Soil Burial Test

The biodegradability of the designed nanohydrogel film series (CSG) was measured by a soil burial test [19]. The film samples were cut into 20x40mm size, weighed and then buried in soil which is in a container with a depth of 10cm. Well sprinkling water on the soil to maintain enough humidity and check with regular time intervals. The effect of soil and its' humidity was observed after a one-week interval of time and the CSG film was washed gently with distilled water to remove any soil from them. The CSG hydrogels were then dried and weighed. The samples were then buried again and the same procedure was repeated for 5 weeks and mass was taken after every week. All experiments were repeated into triplets and the average results have been considered. The biodegradation of the CSG nano hydrogel film samples was then calculated using Eq. (5).

$$Mass Loss (\%) = \frac{m_i - m_f}{m_i} \times 100$$
(5)

Where m_d is the dried weight of the polymer blend film and m_i is the initial weight of CSG hydrogel polymer blend film.

Results and Discussion

Spectral Studies of CSG Hydrogel

The presence of the carboxyl group (-COOH) in carboxymethyl cellulose presented in Fig. 2 showed a prominent absorption band at 1593 cm^{-1} . The stretching frequency of the -OH group is visible in the broad absorption band at 3284 cm^{-1} . At 2913 cm^{-1} , the C-H group's stretching vibration is visible. The vibration of -CH₂ scissoring at 1486 cm^{-1} and -OH bending at 1348.34 cm^{-1} , respectively, are ascribed to these bands. The CH-O-CH₂ band at 1046 cm^{-1} exhibits stretching [20]. The O-H stretching vibration of the polymer and water involved in hydrogen bonding is represented by the absorption peak of cellulose in the region of



Fig. 2 FTIR spectra of CSG hydrogel films

3114–3661cm⁻¹. Stretching modes for C-H can be seen in the area of FTIR spectra between 2831cm⁻¹ and 2989cm⁻¹. It was also noted that the region of about 1473cm⁻¹ was deformed by -CH₂. 1015cm⁻¹ has a range of pyranose rings. Because of their monosaccharide molecules containing pyranose rings, the range of 1015cm⁻¹ can be explained. Cellulose's C-H, C-O, and C-OH stretching infrared absorption spectroscopy in the 817-1166cm⁻¹ band [21].

PVA is associated with the stretching O-H from the intramolecular and intermolecular hydrogen bonds because of the broad bands between 3600-3000cm⁻¹. The vibrational band seen between 2857cm⁻¹ is due to the stretching of C-H from alkyl groups, and the peaks between 1750-1720cm⁻¹ are caused by the stretching of C-O from the acetate group still present in PVA [20]. Since its' monosaccharide contains pyranose rings, the range of 1015 cm⁻¹ can be explained. The existence of both Si-O-Si and Si-O-C was revealed by the asymmetric stretching bands in the 1105–1015cm⁻¹ range, which supported the crosslinking via silanol (Si-OH) groups. In the produced hydrogels, OH stretching of intra- and intermolecular hydrogen bonds was visible in a wide region between 3550-3150cm⁻¹. At 2921cm⁻¹, an alkyl group vibrational band (-CH stretching) was noticed. At 3710cm⁻¹, the Si-OH stretching vibration was observed [20].

Structural Studies of CSG Hydrogels

The X-ray diffraction (XRD) technique is important for analyzing the structural characteristics of the sample under examination. The technique of X-ray diffraction is also used to evaluate the dispersion of incorporated particles in film samples. Plotting an X-ray diffraction chart between two theta values and intensity in counts, where two thetas are taken across the x-axis and intensity along the y-axis. The X-ray wavelength is between 0.1-100 angstrom and the X-ray is considered to be electromagnetic radiation. Dispersion of SiO₂NPs was investigated by X-ray diffraction patterns in polymer structure. Results revealed in Fig. 3 that the CSG sample prepared with natural polymers, due to its amorphous behaviour indicated no peak in the pattern [22, 23]. The reinforcement of different amounts of SiO₂NPs (filler) in hydrogel film samples showed their crystalline behaviour because particles of SiO₂NPs tangled between the chains of the polymers and increasing the concentration of SiO₂NPs in samples increased their crystallinity. It was also observed that when the concentration of SiO₂NPs increased in film samples this increased the intensity at a fixed angle. Results showed samples CSG 0.15 and CSG 0.2 revealed high crystalline behaviour with maximum intensity and could not handle and carry the food item. Thus, sample CSG 0.1 with appropriate crystallinity and flexibility is preferred for food packaging film.



Fig. 3 Diffractogram results of CSG hydrogel films

Surface Morphology of CSG Hydrogel Films

Scanning electron microscopy (SEM) is important for studying the surface morphology of the CSG hydrogel films. SEM micrographs give information about pore size, polymeric structure as well and entanglement of particles (SiO₂NPs) with pores of hydrogel films. All the film samples consisted of the same polymeric concentration (PVA/CMC/gelatin) but different concentrations of SiO₂NPs. SEM micrographs showed that film sample CSG (without SiO₂NPs) exhibited prominent pores surface presented in Fig. 4(a, b). The surface of the film sample CSG 0.05 (with 0.05g SiO₂NPs) becomes a uniform reduction in pore size due to the presence of SiO₂NPs with nanoscale interaction with chains of polymers as in Fig. 4(c, d). As the concentration of SiO₂NPs (0.1g) was increased in film sample of CSG 0.1, then micrographs were observed along with evenly distributed SiO₂NPs with lessening the average pore size as shown in Fig. 4(e, f). Also, the further addition of $SiO_2NPs 0.15g$ and 0.2gin hydrogel film samples, CSG 0.15 and CSG 0.2, respectively, showed the large, deformed patches which presented the fractured surface a coarse appearance Fig. 4(g, h & i, j). SEM micrographs also revealed that increasing the concentration of SiO₂NPs in hydrogel films decreases the pore size of hydrogel films. It was clearly observed that the film sample, CSG 0.1 with favourable surface morphology, appropriate pore size and appropriate crystallinity can be preferred as a supportive format for food packages.

Thermal Analysis of CSG Hydrogel Films

TG thermograms in Fig. 5 revealed that the film samples with SiO_2NPs required more temperature for thermal



Fig. 4 Micrographs of CSG hydrogel films



Fig. 5 TGA results of CSG hydrogel films

degradation, so the incorporation of SiO₂NPs provided thermally stable films. The differential thermal analysis provides information about endothermic or exothermic changes that take place in the material under examination. In the present case, endothermic changes are revealed by prepared film samples with the addition of SiO₂NPs. The results of differential thermal analysis of prepared hydrogel film samples presented in Fig. 6 showed that each pattern revealed three endothermic peaks at approximately 345°C, 534°C and 615°C. All thermograms revealed that physical changes occurred in prepared film samples with the incorporation of SiO₂NPs.



Fig. 6 DTA results for prepared blended CSG hydrogel films

Mechanical Strength Analysis of CSG Hydrogel Films

Food packaging films demand durability to withstand stress during the packing and handling of food products. The concentration of SiO_2NPs in the prepared film samples affects the tensile strength significantly; it supports the polymer network in Fig. 7 and Table 2. It was observed that increasing the concentration of SiO_2NP exhibited improvement in the ultimate tensile strength of prepared film samples. This revealed that with the



Fig. 7 UTS results for prepared blended CSG hydrogel films

Sample codes	Ultimate tensile strength (MPa)	Elongation at break (%)
CSG	3.15	72
CSG 0.05	4.35	90
CSG 0.1	4.83	96
CSG 0.15	5.31	98
CSG 0.2	6.81	118

inclusion of the filler, a greater force was needed to break the film sample. Especially for the film sample with higher SiO_2NPs (filler) concentration showed the highest ultimate tensile strength.

Swelling and Diffusion Test of CSG Hydrogel Films

The distilled water was used for the measurement of the swelling activity of the prepared film samples. The swelling behaviour of film samples was estimated in the water until the equilibrium was reached. The film samples exhibited continuous increasing swelling activity with time Fig. 8. Results represented those increasing concentrations of SiO₂NPs in film samples CSG 0.05, CSG 0.1, CSG 0.15 and CSG 0.2 increased their swelling behavior. The equilibrium time of CSG, CSG 0.05, CSG 0.1, CSG 0.15 and CSG 0.2 was 70, 70, 100,70 and 80min, respectively. Results revealed hydrogel film CSG 0.1 is more stable than other samples, so hydrogel film CSG 0.1 is good for food packages.

The plot between ln F versus ln t is represented in Fig. 9 and the parameters of diffusion are elaborated in Table 3. CSG, CSG 0.05, CSG 0.15 and CSG 0.2 followed the Fickian



Fig. 8 Swelling test results for prepared blended CSG hydrogel films



Fig. 9 Diffusion responses for CSG hydrogels films

Table 3 Diffusion parameters of CSG hydrogel films

Parameters	CSG	CSG 0.05	CSG 0.1	CSG 0.15	CSG 0.2
n	0.23	1.30	0.80	0.31	0.104
Intercept	-1.40	-3.82	-2.54	-1.17	-0.24
K	0.24	0.021	0.07	0.31	0.78
Regression (R %)	92	85	83	99	96

diffusion phenomenon because the value of n < 0.5 meant there was an active water penetration due to a higher rate of relaxation than the rate of diffusion. CSG 0.1 showed non-Fickian diffusion behaviour as the value of n > 0.5 (0.80) which means, the water penetration rate was higher than the rate of polymeric chains relaxation, whereas CSG 0.05 showed a super case II diffusion mechanism as the value of n was greater than 1 (1.30).

Antioxidant Activity

Antioxidant activity of the CSG series was performed by simple method by using free radical of DCPIP. This antioxidant test is the most acknowledged method for assessing the free radical scavenging activity of hydrogel [24]. This method is extensively used to investigate the ability of hydrogels to behave as free radical scavengers/hydrogen donors and to assess antioxidant activity [25]. Antioxidant behavior is mainly based on the reduction behavior of DCPIP which gives maximum absorption at 517nm due to the odd number of electrons [26]. When the solution sample hydrogels react with DCPIP solution, the hydrogel sample turns paired in the presence of H-donor (which is the free radical scavenging activity of hydrogels) and it is reduced in the form of DCPIP-H. Due to reduction, the absorbance of hydrogels decreased as compared to the pure DCPIP [27]. The sample hydrogels of the CSG series displayed the increasing order of their antioxidant action [28]. The radicals that form DCPIP-H, mainly depend on the number of the electrons captured by the hydrogel H-donor [25, 29]. The intermolecular H-bonds also are formed among SiO₂NPs and other film production compounds which accelerate the anti-oxidant properties into the film [30]. CSG has lower antioxidant activity as compared to others, the addition of a strong antioxidant such as promises to demonstrate much greater improvement with the addition of SiO_2NPs [31, 32]. By increasing the amount of SiO₂NPs the hydrogels represented the increase in antioxidant activity as depicted in Fig. 10 and the absorbance measure is tabulated in Table 4 [33, 34].



Fig. 10 Antioxidant activity of blended CSG hydrogel films

Sample code	Absorbance (nm)
CSG	2.537
CSG 0.05	2.691
CSG 0.1	2.756
CSG 0.15	2.742
CSG 0.2	2.736

Antimicrobial Activity

To corroborate the drug release studies, antimicrobial activity was carried out to inter-relate the zone of inhibition of microbial growth with hydrogels. The study was performed by disc diffusion method by using the hydrogel sample of 6mm diameter against B. cereus and E. coli that was used as a bacterial model [35]. All the hydrogels exhibited the zone of inhibition of different centimetres. According to Table 5, every CSG hydrogel film inhibited its growth against E. coli and B. cereus. The empty areas on the agar plates with varying sizes demonstrated the effectiveness of the hydrogels for microbial resistance. The CSG hydrogel's inhibitory zone was minimal, but as the crosslinker concentration was increased, the CSG series displayed a reasonable amount of antimicrobial action. CSG 0.1 and CSG 1.5 both exhibited antimicrobial activity up to 0.5cm and 0.4cm in diameter respectively, but CSG and CSG 0.05 had inhibition zones of 0.2cm and 0.4cm, respectively presented in Fig. 11(a). On the other hand, CSG 0.1 and CSG 1.5 both exhibited antimicrobial activity up to 1.2 -1.5cm in diameter, respectively, but CSG and CSG 0.05 had inhibition zones of 0.4cm and 1.1cm, respectively presented in Fig. 11(b). SiO₂NPs-modified CSG films have significant influence on antibacterial activity due to its' characteristics such as hydrophobia, oleophobia, a cationic nature, resistance to acids, excellent flexibility, and resistance to abrasion and corrosion which inhibits adhesion and proliferation of microorganisms [36-39].

Soil Burial Response

Biodegradability of the designed CSG hydrogel films was observed using a soil burial environment having moisturized and nourished conditions. The 25°C temperature of the moisturized environment was observed which was

 Table 5
 Antibacterial activity (diameter of inhibition zone) of CSG hydrogel films

Bacteria type	CSG	CSG 0.05	CSG 0.1	CSG 0.15
	(cm)	(cm)	(cm)	(cm)
B. cereus	0.4	0.9	1.2	1.5
E. coli	0.2	0.4	0.5	0.4







Fig. 12 Effect of time on biodegradability of PVA/gelatin hydrogel films

satisfactory for the development of microorganisms' habitat that may play a dynamic part in the biodegradation of CSG hydrogel films. The CSG film samples were placed in the soil for 5 weeks and examined after each week. Figure 12 depicts the behaviour of different CSG films under the soil with a one-week time interval.

The neat CSG hydrogel films without reinforcement of SiO_2NPs exhibited more mass loss than any other CSGreinforced hydrogel films. Hence, it is apparent that mass loss declined as the amount of nano reinforcement was enhanced. Therefore, increasing the amount of nanofiller, the mass loss was rapid. Initially, the mass loss rate was lessened in the first one-week period and the trend is the same for all hydrogel films; but after that, the rate of biodegradation increased sharply. This phenomenon occured because of the polymeric nature of hydrogel film as the polymeric chain breakage started, the role of reinforced nano-reinforcements initiated acting and that provided the space and favourable environment for microorganisms to proliferate and initiate degradation of CSG hydrogel films which presented its' high potential to be used as a substitute packaging material.

Conclusions

The present study involved the preparation of novel food packaging films using gelatine, CMC, PVA, and SiO₂NPs crosslinked with TEOS. The addition of SiO₂NPs resulted in enhanced thermal and mechanical strength of the film samples. XRD analysis revealed distinct peaks corresponding to the presence of SiO₂NPs in the prepared films, with an increase in crystallinity observed with higher concentrations of SiO₂NPs at a fixed angle. SEM analysis indicated that the pore size of the hydrogel films decreased with increasing SiO₂NPs concentration. Among the film samples, CSG 0.1 exhibited favourable surface morphology and an appropriate pore size, leading to low weight and high specific surface area. This characteristic provides interesting opportunities for mechanical protection, thermal insulation, and adsorbing or releasing specific compounds that make it suitable for food packaging applications.TGA/DTA results demonstrated that the thermal stability of the prepared film samples improved with increasing SiO₂NPs concentration. UTS analysis revealed that the mechanical strength of the film samples was enhanced by higher concentrations of SiO₂NPs, but it displayed high crystallinity which is unfavourable for swelling characteristics of biodegradable hydrogels. Therefore, the film sample CSG 0.1, with an appropriate SiO₂NPs concentration of 0.1g, displayed 96% elongation at break and 4.8MPa ultimate tensile strength, offering flexibility with moderate crystallinity, making it a suitable supportive format for food packaging purposes. Furthermore, the swelling activity of the hydrogel films in water was highest for the CSG 0.1 film sample, attributed to the appropriate concentration of 0.1g of SiO₂NPs, which increased the film's swelling capacity. These findings highlight the suitability of CSG 0.1 for food packaging applications, considering its' favourable thermo-mechanical properties. Strong nature of biodegradability was observed with the utmost loading of nano reinforcement. To conclude, the film sample CSG 0.1 can be recommended for food packaging purposes due to its' desirable characteristics. The overall cost-effectiveness of this biofilm, market feasibility, environmental impacts, and comparative study with commercial synthetic plastic are highly recommended to be assessed in further studies.

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Data availability The data that support the findings of this study are available from the corresponding author, Nazia Hossain, upon reasonable request.

Declarations

Ethics approval and consent to participate The facts and views in the manuscript are solely ours, and we are totally responsible for authenticity, validity, and originality. We also declare that this manuscript is our original work, and we have not copied from anywhere else. There is no plagiarism in my manuscript.

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Competing interests The authors declare no conflict of interest.

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