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Antioxidant, antimicrobial, and cytotoxic properties of chitosan–PVA films functionalised with chitooligosaccharide and gallic acid for shelf-life extension of perishable foods

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Environment friendly, non-toxic, and functional packaging films are gaining interest worldwide for extending the shelf-life of perishable food items. Hence, this study explored the antioxidant, antimicrobial, and cytotoxicity patterns of earlier fabricated chitosan films containing polyvinyl alcohol (PVA), chitooligosaccharide (COS), and gallic acid (GA). In addition, the active properties of chitosan films were reconfirmed using tomatoes, green grapes, and pearl fish fillets as model perishable food products. Chitosan films containing COS and GA demonstrated strong DPPH ($99.48 \pm 0.24\%$) and ABTS ($98.06 \pm 0.91\%$) radical scavenging capacity and robust potentiality in ferric reducing activity (230.93 ± 1.42 , equivalent $\mu\text{M Fe}^{2+}$ per g sample). Additionally, the inhibitory rates of the fabricated films towards *Escherichia coli*, *Listeria innocua*, and *Saccharomyces cerevisiae* were $59.49 \pm 9.17\%$, $79.29 \pm 0.94\%$, and $79.55 \pm 8.45\%$, respectively. Furthermore, MTT assay exhibited that chitosan films were biocompatible with viability greater than 90% on HaCaT cells and 75% on Caco-2 cells. The fabricated films demonstrated non-cytotoxicity *in vitro* and show potential suitability for packaging applications. Moreover, the application of chitosan films on tomatoes and green grapes showed the lowest weight loss compared to the control film at room temperature ($23\text{ }^\circ\text{C}$) for up to 7 and 9 days, respectively. Additionally, these films also led to a reduction in total volatile basic nitrogen (TVBN) levels in the preservation of pearl fish fillets, indicating the potential for shelf-life extension.

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Sustainability spotlight

This research focuses on the development of functional, non-toxic, and sustainable active packaging films made from chitosan blended with polyvinyl alcohol, chitooligosaccharides, and gallic acid. The resulting active films demonstrated strong antioxidant properties against DPPH and ABTS radicals. Additionally, they displayed significant antimicrobial activity against *E. coli*, *L. innocua*, and *S. cerevisiae*, along with strong biocompatibility with HaCaT and Caco-2 cells. These films effectively reduced weight loss in tomatoes and grapes, as well as minimised total volatile basic nitrogen production during the preservation of pearl fish fillets, highlighting their potential as functional packaging solutions for extending the shelf life of perishable foods.

1 Introduction

Microbial contamination is a common phenomenon during the storage of perishable foods, leading to degraded food quality and a threat to consumers' health.¹ It is estimated that over 200 million tons of food could be wasted by 2050 due to improper

packaging and the short shelf-life of perishable food items.² Additionally, the use of petroleum-based materials in food packaging can lead to significant environmental pollution, posing a threat to both terrestrial and aquatic life.³ Moreover, the use of plastic in food packaging generates microplastics, which accumulate in living terrestrial and aquatic environments through the food chain and enter humans through bio-magnification.⁴ It is reported that microplastics have been detected in human blood, which can cause severe health risks.⁵ In this regard, using biodegradable and functional food packaging materials in the food industry may protect food by preventing contamination, reducing food waste, ensuring food safety, and contributing to the United Nations' Sustainable Development Goals (SDGs).^{6,7}

Several polysaccharides, including chitosan, pectin, starch, and alginate, are considered suitable for developing active food

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packaging materials.⁸ Among them, chitosan is gaining interest in functional packaging materials due to its film-forming capacity, antioxidant and antimicrobial activity, non-toxicity, biodegradability, and biocompatibility.^{9,10} Although chitosan has functional properties in the development of packaging films, it still has some drawbacks, including insufficient antioxidant and antimicrobial abilities, which greatly limit the application of neat chitosan in active food packaging.^{11,12} Additionally, the incorporation of polyvinyl alcohol (PVA) is increasing in the development of active chitosan films due to its excellent film-forming ability, biocompatibility, and improved flexibility, although it offers fewer functional properties. Therefore, chitosan derivatives and phenolic compounds are incorporated to fabricate functional chitosan films.¹³ Chitooligosaccharide (COS) and gallic acid are highly water-soluble and obtained from the enzymatic depolymerisation of chitosan and secondary metabolites in plant materials (*e.g.*, grapes, apples, blueberries, tea, *etc.*), respectively.¹⁴ Studies have reported that COS and gallic acid possess several functional activities, such as antioxidant, antimicrobial, and antiviral properties.^{15–17} The incorporation of COS in the development of active chitosan films, through conjugation with caffeic acid, showed potential antioxidant activity, particularly in scavenging DPPH radicals.¹⁸ Additionally, the combination of gallic acid with corn starch and pullulan in the production of functional films demonstrated both DPPH radical scavenging properties and antimicrobial activity against *Escherichia coli*, *Staphylococcus aureus*, and *Pseudomonas aeruginosa*.¹⁹ Hence, there has been growing interest in including COS and gallic acid in developing active food packaging materials.

In general, active food packaging materials come into direct contact with food and release bioactive compounds to extend product shelf life by reducing lipid oxidation and microbial growth.^{20,21} Even though using biomaterials in the development of packaging films is safe, the interaction of raw materials during the film's synthesis process could alter the packaging film's biocompatibility patterns.²² Therefore, the cytotoxicity patterns of packaging films need to be tested before practical application for extending food shelf-life. In addition, the packaging materials used in food applications should not release any toxic compounds that could harm human health, as stated by the Food Safety Act, 1991, in EC 1935/2004.²³ Thus, the main objective of the current study is to explore the antioxidant, antimicrobial, and cytotoxicity patterns of developed chitosan films containing polyvinyl alcohol, COS, and gallic acid.

2 Materials and methods

2.1 Materials

Chitosan (91% degree of deacetylation and 503 kDa molecular weight, M_w) was acquired from Weseta International in Shanghai, China. Chitooligosaccharide (COS) was synthesised following the method of Rajabi *et al.*²⁴ Polyvinyl alcohol ($\geq 99\%$ hydrolysed and 89 000–98 000 $g\ mol^{-1}$ M_w) and gallic acid (97.5% and 170 $g\ mol^{-1}$ M_w) were provided by Sigma-Aldrich, New Zealand. TPTZ (2,4,6-Tris(2-pyridyl)-s-triazine), M_w : 312.33, and $\geq 98\%$), ABTS (2,2'-azino-bis(3-ethylbenzothiazoline-6-sulfonic

acid, M_w : 548.68 $g\ mol^{-1}$, and $\geq 98\%$), and DPPH (2,2-diphenyl-1-picrylhydrazyl and molecular weight: 394.32) were obtained from Sigma-Aldrich, Switzerland, Canada, and Germany respectively. Potassium persulfate (M_w : 270.32 and $\geq 99\%$) was procured from Sigma-Aldrich. Methanol (CAS-No.: 67-56-1, $>99.8\%$, and M_w : 32.04) was supplied by Fisher Scientific, UK, Ltd, India. Glacial acetic acid (CAS No.: 64-19-7 and $>99.7\%$), sodium acetate trihydrate (M_w : 136.08, and 99% from Biolab (Aust) Ltd), ferrous sulphate (CAS No.: 7782-63-0 and 99%), iron(III) chloride hexahydrate (CAS No.: 10025-77-1, M_w : 270.30 $g\ mol^{-1}$, and 97%) were also obtained from Sigma-Aldrich, USA, for this study. *Escherichia coli* (ESR 916), *Saccharomyces cerevisiae* (ICMP 10067), and *Listeria innocua* (ESR 3024) were acquired from the Microbiological Lab culture collection of the Department of Food Science, University of Otago. Luria broth (LB), nutrient yeast peptone dextrose broth (YPD), and tryptic soy broth (TSB) were supplied by Becton, Dickinson, and Company, Le Pont de Claix, France. The Caco-2 cells (P6, human colon cells) and HaCaT cells (P38, human keratinocyte skin cells) were obtained from the American Type Culture Collection (ATCC, Manassas, VA, USA). DMEM (Dulbecco's modified Eagle's medium), FBS (fetal bovine serum), anti-anti (antibiotic-antimycotic), 0.25% trypsin-EDTA, DPBS (Dulbecco's phosphate buffered saline), calcein AM, and propidium iodide were obtained from Gibco (Life Technologies Corporation, Grand Island, USA). Additionally, ethylene tetrazolium bromide (3-(4,5-dimethyl-2-thiazolyl)-2,5-diphenyl-2H, MTT) was purchased from Thermo Fisher Scientific (Life Technologies Corporation, Eugene, USA), and dimethyl sulfoxide (CAS No.: 67-68-5, molecular weight: 78.13 $g\ mol^{-1}$, and $\geq 99.7\%$, DMSO) was procured from Sigma-Aldrich, USA. Potassium carbonate anhydrous (K_2CO_3 , M_w : 138.21, 99% assay) was purchased from M & B Laboratory Chemicals (May & Baker Ltd, Dagenham, England). Methyl red ($C_{15}H_{15}N_3O_2$), bromocresol green ($C_{21}H_{14}Br_4O_5S$), and phenolphthalein ($C_{20}H_{14}O_4$) were acquired from Hopkin & Williams Ltd, Chadwell Heath, Essex, England. Boric acid (H_3BO_3 , M_w : 61.83, 99.5% assay) was supplied by Ajax Chemicals, Auburn, Australia. Analytical grade ethanol (C_2H_5OH , 99.5%) was procured from the Department of Chemistry, University of Otago. Analytical reagent grade hydrochloric acid (HCl, M_w : 36.46, CAS-No.: 7647-01-0, and 37%) was supplied by Fisher Scientific, UK. Sodium hydroxide (NaOH, CAS-No.: 1310-73-2, M_w : 40, and $>98\%$ assay) was provided by Sigma-Aldrich, New Zealand. Fresh tomatoes, green grapes, and pearl fish fillets were purchased from the local supermarkets in Dunedin, New Zealand.

2.2 Chitosan film fabrication

The chitosan films were fabricated using the solvent casting method, following the process established by Bhowmik *et al.*²⁵ Briefly, 2% chitosan (CH) solution was obtained by mixing 0.4 g of CH in 20 mL of 2% acetic acid solution. After that, 0.2 g (50% based on CH) of polyvinyl alcohol (PVA), 0.02–0.04 g (5–10% based on CH) of chitooligosaccharide (COS), and 0.02–0.04 g (5–10% based on CH) of gallic acid (GA) were added and stirred for 30 min at room temperature (23 °C). Then, the film-forming solution was poured into square Petri dishes (10 cm × 10 cm) and kept for 72 h at 23 °C. The fabricated films were manually



separated from the Petri dishes. The film containing only chitosan was regarded as the CH film, followed by CP2 (CH + 0.2 g PVA), CP5 (CH + 0.2 g PVA + 0.02 g COS), CP6 (CH + 0.2 g PVA + 0.04 g COS), CP7 (CH + 0.2 g PVA + 0.02 g GA), CP8 (CH + 0.2 g PVA + 0.04 g GA), CP9 (CH + 0.2 g PVA + 0.02 g COS + 0.02 g GA), CP10 (CH + 0.2 g PVA + 0.04 g COS + 0.04 g GA), CP11 (CH + 0.2 g PVA + 0.04 g COS + 0.02 g GA) and CP12 (CH + 0.2 g PVA + 0.02 g COS + 0.02 g GA) films based on previous work (Table S1), where the physical, mechanical, structural, thermal, and biodegradation properties of the films were included.²⁵

2.3 Antioxidant activity of developed chitosan films

2.3.1 DPPH (2,2-diphenyl-1-picrylhydrazyl) assay. The DPPH assay, with some modifications, was used to measure the antioxidant properties of the fabricated chitosan films.²⁶ In short, 7.88 mg of DPPH powder was dissolved in 100 mL of 99.8% methanol to create 0.2 mM DPPH solution. After that, the film sample (5 mg) was mixed with 1.5 mL of DPPH solution and stored at 22 °C. A UV/Visible Spectrophotometer (Ultrospec 3300 Pro, Biochrom Ltd, Cambridge, England) was then used to test the mixture's absorbance at 517 nm across a range of time intervals (0.5–16 h). The scavenging capacity of DPPH free radicals was determined in triplicate, and the mean values are presented as per eqn (1).

$$\text{Scavenging ability (\%)} = \left[1 - \frac{(A_1 - A_2)}{A_0} \times 100 \right] \quad (1)$$

where A_0 is the absorbance of DPPH solution, A_1 is the absorbance of a film sample with DPPH solution, and A_2 is the absorbance of a film sample with methanol.

2.3.2 ABTS assay. The scavenging properties of ABTS free radicals of the developed chitosan films were evaluated using the methods of Božič *et al.*²⁷ and Riaz *et al.*²⁶ with some modifications. In summary, 384 mg of ABTS powder and 66 mg of potassium persulfate were separately dissolved in 100 mL of distilled water to create a 7 mM ABTS solution and 2.45 mM $K_2S_2O_8$ solution. After that, the same volumes of ABTS and $K_2S_2O_8$ solutions were mixed and left in the dark for 16 h at 22 °C. Subsequently, the combined solution was diluted 1 : 10 with a 50% methanol solution to create a workable solution. The absorbance of the working solution at 734 nm using a UV/Visible Spectrophotometer (Ultrospec 3300 Pro, Biochrom Ltd, Cambridge, England) was measured and found to be 0.96 (<1). After that, 1.5 mL of working solution and 5 mg of film sample were mixed, and the mixture was left in the dark. At various times (6 min–4 h), the absorbance at 734 nm was measured. Antioxidant ability was calculated in triplicate using eqn (2).

$$\text{Scavenging ability (\%)} = \left[1 - \frac{(A_s - A_m)}{A_w} \times 100 \right] \quad (2)$$

where A_w is the absorbance of a working solution, A_s is the absorbance of a film sample with a working solution, and A_c is the absorbance of a film sample with methanol.

2.3.3 FRAP assay. The ferric-reducing antioxidant power (FRAP) assay was conducted following the method of Gulzar *et al.*²⁸ with some modifications to measure the capacity of the

fabricated chitosan films to transform ferric iron (Fe^{3+}) to ferrous (Fe^{2+}) iron. Concisely, 300 mM acetate buffer (pH < 3.6) was prepared by dissolving $CH_3COONa \cdot 3H_2O$ in 100 mL ultrapure water containing 1.6 mL CH_3COOH , stored at 22 °C using an amber flask. Thereafter, a 10 mM HCl solution was prepared by mixing 40 μ L 37% HCl in 10 mL ultrapure water; after that, 10 mM 2,4,6-tris(2-pyridyl)-s-triazine (TPTZ) solution was prepared by adding 31 mg TPTZ in 10 mM HCl solution. A 20 mM $FeCl_3 \cdot 6H_2O$ solution was also prepared by dissolving 54.1 mg $FeCl_3 \cdot 6H_2O$ in 10 mL ultrapure water. Similarly, a 2000 μ M $FeSO_4 \cdot 7H_2O$ solution was made by mixing 55.6 mg $FeSO_4 \cdot 7H_2O$ in 100 mL ultrapure water. A 72 mL working solution was prepared by mixing 6 mL $FeCl_3 \cdot 6H_2O$ and 6 mL TPTZ with 60 mL acetate buffer. Then, 5 mg of film sample was mixed with 0.9 mL of working solution and incubated in a water bath (TECHNE, TE-10A Tempette, Total Lab Systems Ltd) at 37 °C for 15 min. The Fe^{2+} TPTZ complex was assessed by measuring absorbance at 593 nm and calculating its activity in triplicate by using the equation ($Y = 0.0008x + 0.1303$ and $R^2 = 0.99$) from the generated standard curve using $FeSO_4 \cdot 7H_2O$ (0–2000 μ M). The results were expressed as μ M Fe^{2+} equivalents per g of the film sample.

2.4 Antimicrobial properties of films

The antimicrobial activities and the integrity of microbial cell membranes of chitosan films were assessed towards Gram-positive bacteria (*Listeria innocua*), Gram-negative bacteria (*Escherichia coli*), and yeast (*Saccharomyces cerevisiae*) following the method of Bi *et al.*²⁹ and Sadiq *et al.*³⁰ with some modifications. *E. coli* and *L. innocua* were cultured in liquid Luria broth and tryptic soy broth media at 37 °C to the logarithmic phase, and *S. cerevisiae* was cultured in yeast peptone dextrose medium at 30 °C. After that, 0.2 mL microbial suspensions (with an initial absorbance of 1 at OD_{600}) of *E. coli*, *S. cerevisiae*, and *L. innocua* were diluted 50 times using 10 mL of LB, YPD, and TSB medium, respectively, and incubated with the films (6 mm \times 6 mm) with 0.2 mL liquid medium for 24 h. Then, the optical density at 600 nm was measured using a plate reader (CLARIOstarPlus, BMG LABTECH, Germany) at different time intervals (0–24 h), and the antimicrobial rate was analysed in triplicate using eqn (3).

$$\text{Antimicrobial rate (\%)} = (OD_1 - OD_2)/OD_1 \times 100 \quad (3)$$

where OD_1 is the microbial optical density of the control group (without a film), and OD_2 is the microbial optical density treated with developed films.

To assess the integrity of microbial cell membranes, 0.15 mL of microbial suspension was obtained after 24 hours, and it was centrifuged (1–16 K, SIGMA, Germany) at 11000 \times g for five minutes. Utilising 0.1 mL of the supernatant, the quantities of cellular constituents were assessed as proposed by Bi *et al.*²⁹ by determining the absorbance at 260 nm with a plate reader (CLARIOstarPlus, BMG LABTECH, Germany). The control group consisted of the microbial suspension supernatant that had not been film-treated.



2.5 Cell assay, viability, and proliferation assays

2.5.1 Cell assay. The frozen stocks of HaCaT cells (P38, human keratinocyte skin cells) and Caco-2 cells (P6, human colon cells) were removed from liquid nitrogen and warmed in a water bath at 37 °C and thawed for 50 seconds. Then, HaCaT and Caco-2 cells were grown in T-75 cm² flasks in complete DMEM (cDMEM) media containing 10% fetal bovine serum and 1% antibiotic–antimycotic (penicillin–streptomycin). Then, the culture flasks were kept in a humidified atmosphere of 95% air and 5% CO₂ using a cell incubator (MCO-19AIC (UV), CO₂ incubator, SANYO Electric Co., Ltd, Japan), maintained at 37 °C. The cDMEM was replaced every three days during the culture periods. Additionally, HaCaT and Caco-2 cells were passaged to maintain their proliferative state when they reached 70–90% confluence through trypsinisation. The cell counting was performed using Septer™ sensors (60 μm), EMD Millipore Corporation, Burlington, USA.

2.5.2 Cell viability assay. The toxicity and biocompatibility of developed chitosan films were evaluated using LIVE/DEAD® assay, which determines cytotoxicity/cell viability.^{31,32} Concisely, sterile film samples (6 mm × 6 mm) were immersed in cDMEM, and both the HaCaT and Caco-2 cells were seeded separately per well and incubated at 37 °C for 24 h, 48 h and 72 h. The cDMEM with cells but without a film was considered a control. The assay staining solution was prepared using 8 mL DPBS, 12 μL calcein AM, and 24 μL propidium iodide. After 24 h, 48 h, and 72 h, the medium was removed from the well plate, and 100 μL assay solution was added to each well and incubated at 37 °C for 10 min. After that, cells were observed under a fluorescence microscope (EVOS M5000, ThermoFisher, Waltham, USA) using EVOS™ M5000 software. The enzymatic conversion of calcein AM to calcein (excitation 494 nm, emission 517 nm) allowed identification of living cells.³³ The nucleic acids of cells with damaged cell membranes bound to propidium iodide indicate dead cells (excitation 528 nm and emission 617 nm). The number of living and dead cells was counted using Image J software (US National Institutes of Health) with a plugin.³⁴ The fluorescent images of each cell, captured at different culture periods, were converted to greyscale using ImageJ, resulting in monochromatic 16 bit images. This process transformed the pixels into individual data points. Next, we calculated the number of pixels representing dead (red) cells and live (green) cells. The percentage of live cells was then determined relative to the total number of cells in each image. To calculate the average value, three images of each cell from every culture period and each film were used, maintaining consistent settings applied across all images to ensure uniformity. The percentage of cell viability was calculated in triplicate using eqn (4).

$$\text{Cell viability (\%)} = \frac{\text{Number of live cells}}{\text{Total cell number}} \times 100 \quad (4)$$

2.5.3 Cell proliferation assay. The cell proliferation of HaCaT and Caco-2 cells was measured at 24 h, 48 h, and 72 h calorimetrically using ethylene tetrazolium bromide (3-(4,5-

dimethyl-2-thiazolyl)-2,5-diphenyl-2H, MTT) assay.³⁵ Briefly, 12 mM MTT stock solution was prepared by dissolving 50 mg MTT powder in 10 mL sterile DPBS and kept at –20 °C. The HaCaT and Caco-2 cells were seeded separately (5 × 10³ cells) per well containing chitosan film samples (6 mm × 6 mm) and incubated at 37 °C for 24 h, 48 h and 72 h in quadruplicate. After 24 h, 48 h, and 72 h, the medium was removed from the well plate, and 100 μL MTS reagent (earlier prepared 7 mL by dissolving 1.4 mL MTT solution with 5.6 mL cDMEM) was added to each well and kept in a humidified incubator for 4 h. Then, the medium was removed from the well plate, and 50 μL dimethyl sulfoxide (DMSO) was added to each well. DMSO containing HaCaT and Caco-2 cells was transferred to new well plates, and absorbance was measured at 540 nm using a plate reader (CLARIOstarPlus, BMG LABTECH, Germany). Standard curves (Fig. S1) were used to detect the unknown cell numbers of HaCaT and Caco-2 cells treated with films.

2.6 Evaluation of storage quality of tomato and green grape

The storage qualities of tomatoes and green grapes were assessed as model food products to investigate the active properties of the developed films (CH, CP9, and CP10). The prepared tomatoes (3.2–3.5 g) and green grapes (3.1–3.3 g) were kept in a Petri dish (3.5 cm × 3.5 cm) covered with a lid. The films (1.3 cm × 1.3 cm) were inserted inside the Petri dish, and the setups containing tomatoes and grapes were stored for 7 and 9 days, respectively, at room temperature (23 °C, relative humidity 46%). Tomatoes and green grapes in the Petri dish without a film were regarded as the control, and those without a lid and film were designated as non-packaging (NP). The Petri dish was weighed every day, and the weight was recorded. The weight loss of tomatoes and grapes was measured in triplicate using eqn (5).³⁶

$$\text{Weight loss (\%)} = (W_1 - W_2)/W_1 \times 100 \quad (5)$$

where W_1 is the initial weight of tomatoes and grapes in a Petri dish, and W_2 is the weight on the day of measurement.

2.7 Evaluation of storage quality of pearl fish fillet

The purchased pearl fish fillets were cut (20–25 g) and kept in a Petri dish with films (1.3 cm × 1.3 cm) separately at room temperature (23 °C, relative humidity 46%). The pH, drip loss, and total volatile basic nitrogen (TVBN) were measured daily in triplicate to detect the storage quality of pearl fish fillets. The pearl fish fillet without a film was considered the control, and the one without a film and lid was regarded as the NP group.

The pH of pearl fish fillet was detected following the method of Pang *et al.*³⁷ with some modifications. Five (5) grams of pearl fish fillet were vigorously homogenised in 45 mL of distilled water. After that, the pH of the pearl fish fillet filtrate was measured in triplicate using a pH meter (HI 5222, Hanna Instruments, USA). Additionally, the drip loss of packaged pearl fish fillets with films in a Petri dish was assessed using the weighing method.³⁶ The pearl fish fillets in a Petri dish were



weighed daily using an analytical balance, and eqn (6) was used to calculate the drip loss.

$$\text{Drip loss (\%)} = (W_1 - W_2)/W_1 \times 100 \quad (6)$$

where W_1 is the initial weight of pearl fish fillets in a Petri dish, and W_2 is the weight on the measurement day.

The TVBN content of pearl fish fillets was measured following the method of Lee *et al.*³⁸ with some modifications. Briefly, 5 g of ground pearl fish fillets were inserted in a 50 mL Falcon tube containing 45 mL distilled water and kept at room temperature (23 °C) for approximately 35 min. After that, Whatman No. 1 filter paper was used to filter the samples, and the supernatant was analysed. Before sample analysis, the Conway indicator solution was prepared by dissolving 66 mg of bromocresol green and 66 mg of methyl red in 100 mL of ethanol (99.5%). Additionally, a 0.01 N boric acid solution was prepared by dissolving 0.3 g boric acid in 150 mL of ethanol, and the volume was increased to 500 mL using 350 mL of distilled water. Also, a 50% potassium carbonate solution was made by dissolving 50 g of K_2CO_3 in 100 mL of distilled water. In the analysis of TVBN in pearl fish fillets, 1 mL of H_3BO_3 and 100 μ L of indicator solution were included in the inner part of the Conway unit, and 1 mL of supernatant and 1 mL of K_2CO_3 solution were added in the outer chamber of the Conway unit. Then, the Conway unit was kept at 37 °C for 2 h in an oven (THERMOTEC 2000, Lower Hutt, New Zealand). After that, 0.01 N HCl was incorporated into the inner part of the Conway unit until the formation of the pink colour. The TVBN content was calculated in triplicate using eqn (7).

$$\text{TVBN (mg/100 g)} = 0.14 \times (b - a) \times f \times d/W \times 100 \quad (7)$$

where b and a are the titration volumes (mL) of HCl for a sample and blank, respectively, f is the concentration of HCl (0.01 N), d is the dilution factor (45 mL), and W is the pearl fish fillet weight (5 g).

2.8 Statistical analysis

The statistical analysis used GraphPad Prism version 9.4.1 (GraphPad Software, Boston, USA). The obtained data from at least three replicates are presented as mean \pm standard deviation (SD). To look for significant differences ($P \leq 0.05$) in the generated films, cell viability, and total cell number, Tukey's test and ordinary one-way analysis of variance (ANOVA) were employed.

3 Results and discussion

3.1 Antioxidant activity of developed chitosan films

The antioxidant activity of fabricated chitosan films is presented in Fig. 1. In the DPPH scavenging assay, the fabricated films showed different activities during 0.5–16 h. Films CP8 (95.37 ± 0.57 – $95.51 \pm 0.56\%$), CP10 (91.37 ± 5.52 – $95.38 \pm 0.08\%$), and CP12 (88.30 ± 9.51 – $95.44 \pm 0.25\%$) exhibited the highest DPPH scavenging activities with greater consistency during the time periods, whereas CP7 (46.80 ± 1.44 – $94.12 \pm$

1.85%), CP9 (69.82 ± 5.48 – $96.30 \pm 0.04\%$), and CP11 (41.31 ± 6.14 – $99.48 \pm 0.24\%$) demonstrated linearity in different time intervals. CH (8.31 ± 0.43 – $13.13 \pm 4.26\%$), CP2 (6.88 ± 0.17 – $14.80 \pm 2.45\%$), CP5 (2.76 ± 1.79 – $7.16 \pm 3.13\%$), and CP6 (3.13 ± 0.21 – $8.24 \pm 2\%$) films presented the lowest percentage of DPPH scavenging activities. These research findings are consistent with Zhang *et al.*,³⁹ who reported that a neat chitosan film (15%) demonstrated the lowest DPPH scavenging capacity compared to a gallic acid-loaded film (50–83%), indicating that the addition of gallic acid in the chitosan film enhanced anti-oxidant activity due to the interaction between the native anti-oxidant properties of gallic acid moieties and the chitosan backbone. The addition of PVA and COS in the chitosan film did not improve the DPPH radical scavenging activity in the chitosan films. However, the inclusion of gallic acid and COS significantly enhanced the DPPH radical scavenging capacity of the chitosan–PVA films labelled CP9–CP12. These findings align with the research conducted by Yuan *et al.*,¹⁸ who elucidated that COS and caffeic acid-loaded chitosan films significantly increased scavenging ability (93.43%) compared to the control film (10%). Additionally, similar studies reported by Lee *et al.*⁴⁰ indicated that including gallic acid in the preparation of chitosan active films exhibited strong scavenging properties (95.7%) due to the presence of a phenolic hydroxyl group.

Similarly, in the ABTS scavenging assay, CP10 (97.85 ± 1.01 – $98.06 \pm 0.91\%$), CP12 (97.41 ± 0.69 – $97.57 \pm 0.53\%$), CP8 (94.12 ± 5.05 – $95.79 \pm 2.63\%$), and CP11 (94.34 ± 2.87 – $96.68 \pm 0.38\%$) showed the highest percentage of ABTS scavenging properties, whereas CP7 (84.23 ± 9.09 – $96.50 \pm 0.38\%$) and CP9 (85.60 ± 5.59 – $93.87 \pm 0.67\%$) demonstrated different activity with linearity in the experimental time, followed by CH (20.76 ± 7.30 – $80.99 \pm 1.28\%$), CP2 (12.80 ± 0.07 – $91.90 \pm 4.81\%$), CP5 (14.23 ± 10.02 – $90.94 \pm 4.81\%$), and CP6 (11.48 ± 8 – $87.06 \pm 12.72\%$). Similar findings were stated by Zhao *et al.*,¹¹ who described chitosan films (22%) as having the lowest ABTS scavenging activity, whereas adding gallic acid to chitosan films enhanced hydrogen supply capacity, improving ABTS radical scavenging capacity (85%).

Likewise, the highest FRAP (equivalent μ M Fe^{2+} per g sample) was obtained from the CP10 (230.93 ± 1.42) film, followed by CP12 (229.13 ± 4.20), CP8 (228.69 ± 0.49), CP9 (222.88 ± 4.76), CP7 (220.73 ± 1.36) and CP11 (218.11 ± 0.24), whereas the lowest FRAP was exhibited by CP6 (54.66 ± 1.11), CH (33.88 ± 1.79), CP5 (25.04 ± 0.06), and CP2 (24.43 ± 4.39) films. Adding COS and gallic acid to the chitosan–PVA film enhanced its ability to reduce Fe^{3+} to Fe^{2+} , potentially indicating synergistic effects. These findings align with those of Gulzar *et al.*,²⁸ who reported that adding COS and tannic acid to chitosan films demonstrated the highest FRAP activity, which depends on the amounts of COS and tannic acids in film development. A film with the capacity to convert ferric to ferrous ions and delay the production of free radicals might delay the lipid oxidation of foods and avoid oxidative stress.⁴¹ The highest FRAP activities by incorporating COS and gallic acid reconfirmed their active role in developing chitosan films with effective antioxidant properties.



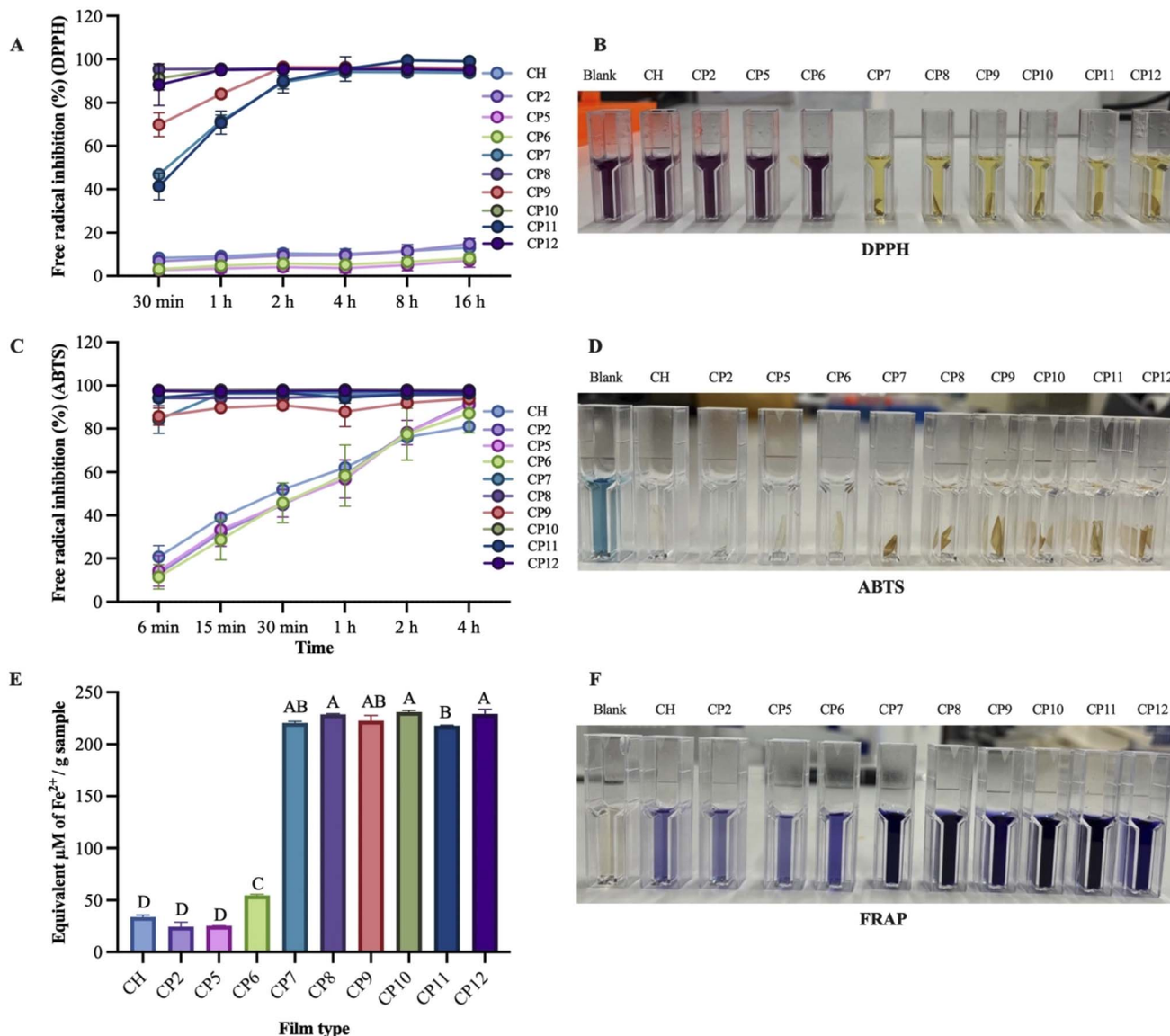


Fig. 1 Scavenging capacity of fabricated films towards DPPH (A), ABTS (C), and FRAP (E) radicals; colour changes of working solution observed during the experiment (B, D and F). FRAP: ferric reducing antioxidant power. Bars represent mean \pm SD with different letters (A–D) that are significantly different at $p < 0.05$, $n = 3$.

3.2 Antimicrobial properties of fabricated films

The antimicrobial activity of the developed chitosan film towards *Escherichia coli* (Gram-negative bacteria), *Listeria innocua* (Gram-positive bacteria), and *Saccharomyces cerevisiae* (yeast) is presented in Fig. 2. Overall, the COS and gallic acid loaded chitosan films showed higher antimicrobial activity and leakage of greater intracellular substances compared to the neat chitosan film. These findings suggested that adding COS and gallic acid effectively improved the antimicrobial properties of the chitosan film. The inclusion of COS and gallic acid in the chitosan film resulted in antimicrobial capacities of $59.49 \pm 9.17\%$ (CP10), $79.29 \pm 0.94\%$ (CP10), and $79.55 \pm 8.45\%$ (CP9). The findings of the study are in line with Zhao *et al.*,¹¹ who demonstrated that the addition of gallic acid improved

antimicrobial activity towards Gram-positive (*Staphylococcus aureus*) and Gram-negative (*Escherichia coli*) bacteria due to the synergistic effect of the positively charged amino group of chitosan and gallic acid, which disrupts the bacterial cell wall and results in bacteria death. Adding PVA and COS alone in the chitosan film did not significantly improve the antimicrobial properties of the chitosan film. These findings are consistent with those of Gulzar *et al.*²⁸ and Kanatt *et al.*,⁴² who reported that elevated concentrations of COS in the film failed to inhibit the growth of microbes (*Listeria monocytogenes* and *Escherichia coli*) effectively, and PVA in the chitosan film did not show any significant changes in microbial growth (*Staphylococcus aureus* and *Bacillus cereus*), respectively. However, combining COS and gallic acid in the chitosan film improved antimicrobial activity in the current study due to the synergistic interaction with



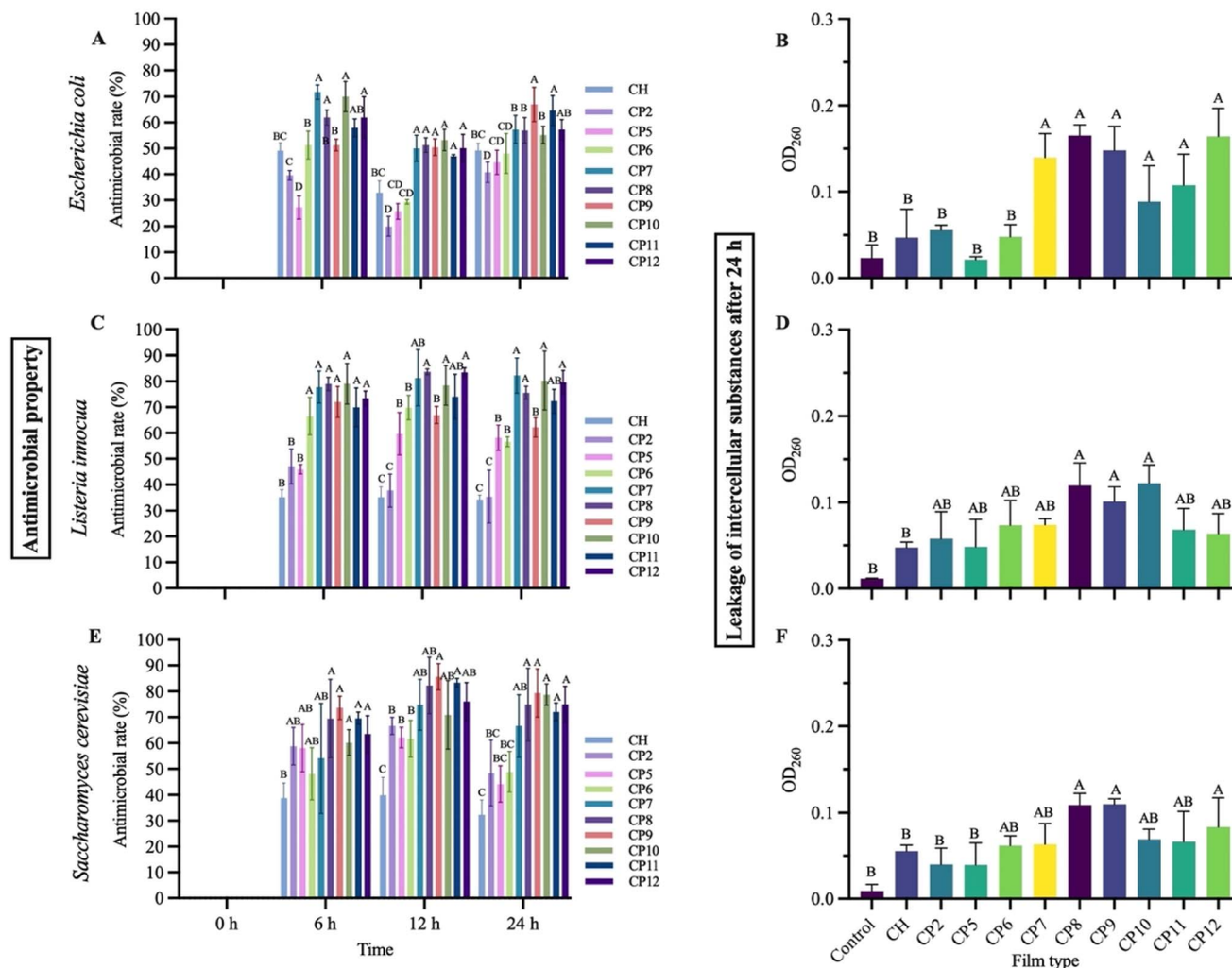


Fig. 2 Antimicrobial rate of fabricated chitosan films towards *Escherichia coli* (A), *Listeria innocua* (C), and *Saccharomyces cerevisiae* (E), and leakage of cell constituents of microbes after 24 h (B, D and F). Bars represent mean \pm SD with different letters (A–D) that are significantly different at $p < 0.05$, $n = 3$.

chitosan. Additionally, the release of cellular components is higher in COS and gallic acid-loaded films (CP7–CP12 (*Escherichia coli*), CP8–CP10 (*Listeria innocua*), and CP8, CP9, and CP12 (*Saccharomyces cerevisiae*)) than in the chitosan film. This could have happened due to the presence of an abundance of free amino groups, contributed by COS, in the film matrix, leading to electrostatic interactions with the phosphate groups of the microbial cell membrane to damage membrane integrity, resulting in the release of intercellular substances.^{29,43} Additionally, the potential interaction of GA with the chitosan film matrix caused membrane pores of microbes to destroy cell membranes and enhance their cell permeability, resulting in leakage of cell constituents.⁴⁴ Also, we hypothesise that the inclusion of COS and gallic acid in the chitosan film acts synergistically to improve membrane permeability and physical disruption of microbial cells, leading to death. These findings are consistent with those of Hou *et al.*,¹⁷ who reported that incorporating gallic acid and tannic acid into a cellulose/chitosan film matrix created an active film with enhanced

antimicrobial activity against *Escherichia coli* and *Staphylococcus aureus* due to their synergistic interactions. Therefore, the OD₂₆₀ values of COS and gallic acid-loaded chitosan films were higher than those of the neat chitosan film, reconfirming the enhancement of antimicrobial activity in the fabricated films of the current study.

3.3 Cytotoxicity patterns of chitosan films

3.3.1 Cell viability of HaCaT cells. The cell viability profile of HaCaT cells is presented in Fig. 3. The green and red fluorescence demonstrated live and dead cells, respectively. The higher proportion of live cells than dead cells confirmed the positive viability of HaCaT cells toward CH, CP9, and CP10 films. The HaCaT cells exhibited excellent viability, more than 90% in all tested films. There were no significant differences ($p > 0.05$) between the control and developed films at 24, 48, and 72 h, except for the CP10 film at 24 h. Following ISO 10993-5, a biological material is considered non-toxic if the cell survival rate is more than 70% during the culture period. Hence, the



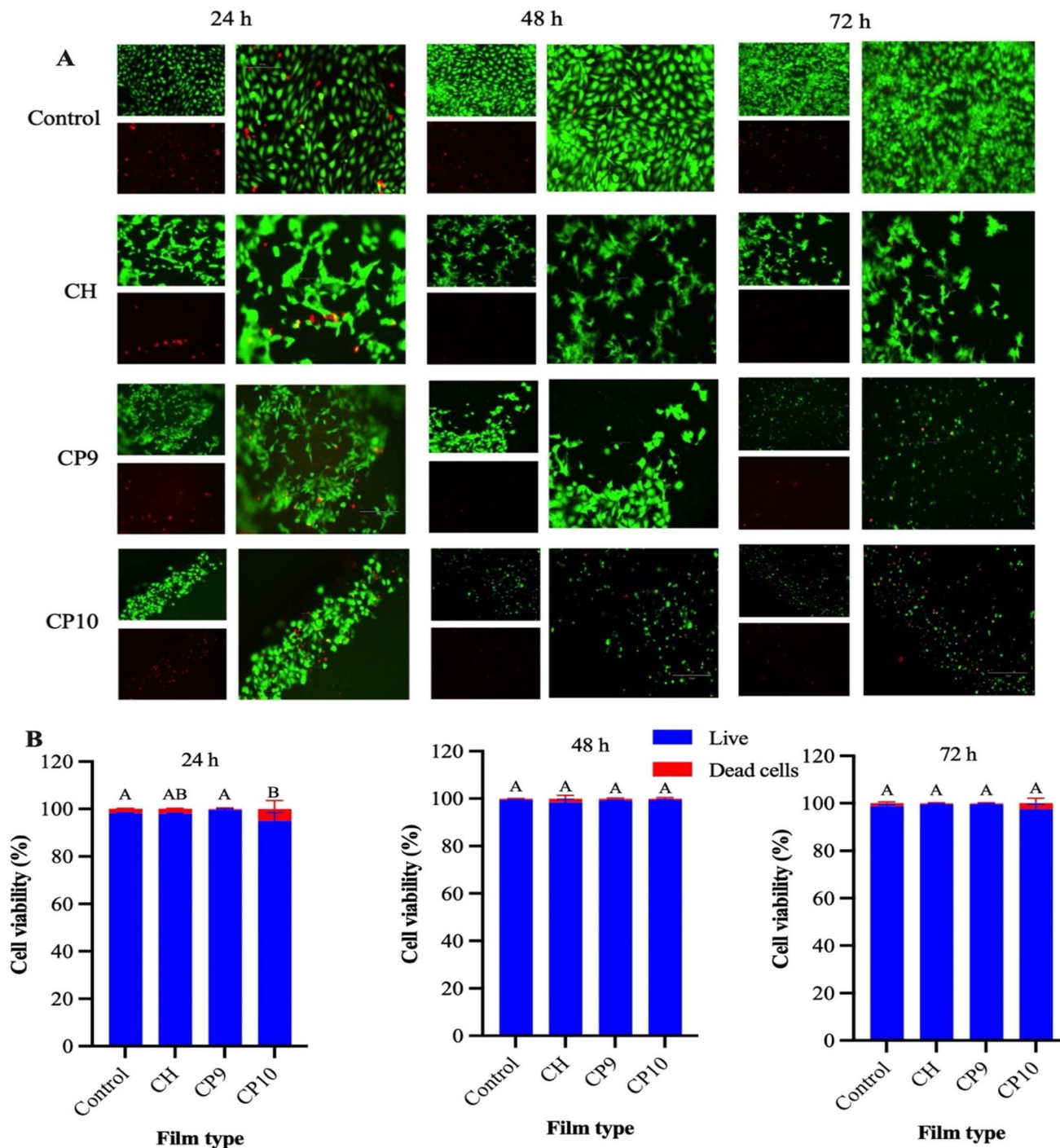


Fig. 3 Dye-based fluorescent microscopy images (A) and cell viability (B) of HaCaT cells after 24, 48, and 72 h of incubation with CH, CP9, and CP10 films. Bars represent mean \pm SD with different letters (A and B) that are significantly different at $p < 0.05$, $n = 3$.

film developed with chitosan, polyvinyl alcohol, COS, and gallic acid is considered a safe food packaging material. Yang *et al.*⁴⁵ fabricated a packaging film containing polyvinyl alcohol, hexamethylene guanidine, and gallic acid and assessed its cytotoxicity. Research findings of their studies showed that the developed film had good biocompatibility, and the cell viability of mouse fibroblast (L929) cells of the gallic acid-loaded film was greater than 70%. Riaz *et al.*⁴⁶ studied the safety of gallic

acid in the developing film containing agarose and showed more than 75% cell viability towards L929 cells. The current study findings demonstrated that adding GA to the chitosan film is non-toxic to HaCaT cells.

3.3.2 Cell proliferation of HaCaT cells. The cell proliferation of HaCaT cells is illustrated in Fig. 4. The results showed that the number of HaCaT cells increased from 24 h to 72 h, and no significant differences existed between the control and



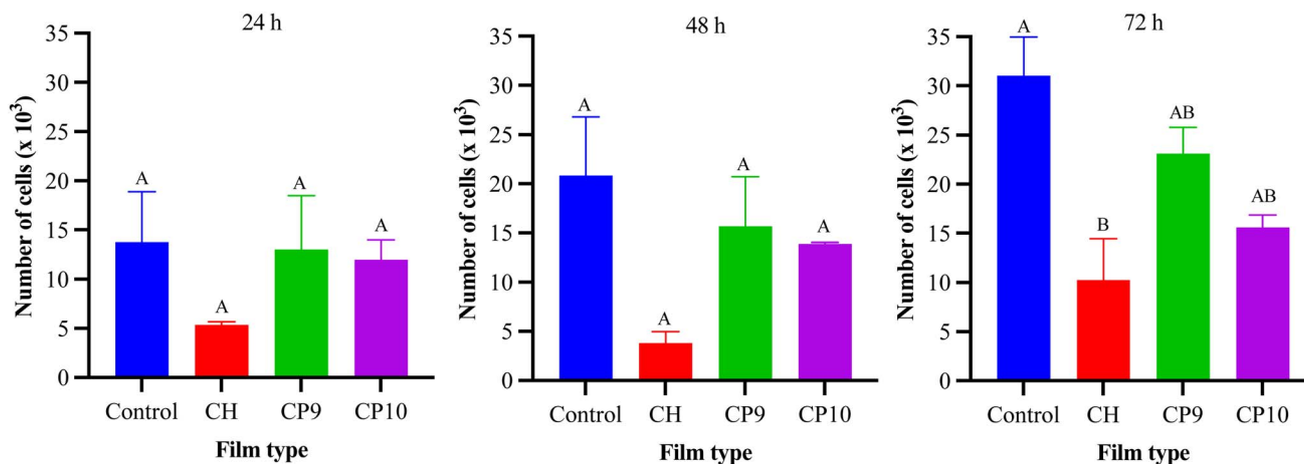


Fig. 4 Cell proliferation of HaCaT cells after 24, 48, and 72 h of incubation with CH, CP9, and CP10 films. Bars represent mean \pm SEM (standard error of the mean) with different letters (A and B) that are significantly different at $p < 0.05$, $n = 4$.

treated HaCaT cells after 24, 48, and 72 h, except for the CH film at 72 h. The results suggest that including PVA, COS, and gallic acid does not hinder the growth of cell numbers but rather increases HaCaT cell numbers, reconfirming the non-toxic properties of the developed films.

3.3.3 Cell viability of Caco-2 cells. The cell viability of Caco-2 cells is shown in Fig. 5. The green and red fluorescence showed live and dead cells, respectively. The higher number of live cells than dead cells in the graph confirmed the positive viability of Caco-2 cells toward CH, CP9, and CP10 films. The Caco-2 cells exhibited good viability, exceeding 75% at 24 h and 80% at 48 and 72 h in all tested films. Additionally, no significant differences ($p < 0.05$) exist between the control and developed films at 72 h, except for the CP10 film. The current research findings are consistent with Liu *et al.*,⁴⁷ who reported that adding gelatine and 3-phenyllactic acid in the development of the chitosan film demonstrated good biocompatibility on Caco-2 cells. The concentration of gallic acid in the chitosan film could affect the cell viability of Caco-2 cells; therefore, the percentage of live cells was significantly reduced in the CP10-treated film compared to the control, CH, and CP9 films. Forester & Waterhouse⁴⁸ reported that the increasing gallic acid concentration significantly inhibits the growth of Caco-2 cells. However, in the present study, all films, including CP10, showed more than 75% cell viability, revealing that the fabricated films are suitable for packaging.

3.3.4 Cell proliferation of Caco-2 cells. The cell proliferation of Caco-2 cells is shown in Fig. 6. The results depict that the number of HaCaT cells increased from 24 h to 72 h in the control compared to film-treated Caco-2 cells. However, no significant differences ($p > 0.05$) were observed between the control and film-treated Caco-2 cells at 24 and 48 h, except for the CH film at 48 h. At 72 h, Caco-2 cells treated with films exhibited significantly less proliferation than the control. The proliferation of Caco-2 cells depends on the viscosity and molecular weight of chitosan and the film-forming matrix.^{49,50} Additionally, the gallic acid content in the chitosan film could be attributed to minimising the proliferation of Caco-2 cells.

Some studies stated that gallic acid demonstrated anti-proliferative effects towards colorectal cancer cell models, including Caco-2 cells.^{48,51} The chitosan film containing gallic acid in DMEM media may generate H₂O₂, increasing cell auto-oxidation and reducing cell proliferation. These findings coincided with those of Mu & Kitts,⁵² who described that phenolic acid, especially gallic acid showed the greatest anti-proliferation properties against Caco-2 cells by generating H₂O₂ in DMEM media. Some researchers reported that gallic acid did not affect the proliferation of the normal cell lines compared to cancer cells.⁵³ Thus, fabricated films could be suitable as food packaging materials. While our current findings suggest suitability for packaging applications, we do not establish a safety margin for oral exposure at this stage. Therefore, the developed material cannot yet be classified as edible packaging, and comprehensive *in vivo* testing together with exposure-based risk assessment is required to confirm its safety for human consumption.

3.4 Application of films in tomato and green grape packaging

The appearance and weight loss of tomatoes under different pack-film conditions are presented in Fig. 7A and B. It was observed that the NP tomatoes gradually lost their appearance, and the surface became wrinkly due to the loss of turgidity in the cells. However, no microbial growth was seen in NP tomatoes compared to other film-packed groups. This observation could be explained by the fact that the NP samples, being without a lid and film, did not accumulate moisture droplets from the respiration of the tomatoes. The development of humidity inside the film-packed samples (control-CP10) created a suitable atmosphere for the growth of microbes. In the case of the control group, microbial growth was observed on day 2. In contrast, the development of microbes in the film-packed tomatoes with CH, CP9, and CP10 films was detected on days 3, 4, and 5, respectively. In other words, films such as CP10 delayed the onset of microbial growth. The current research



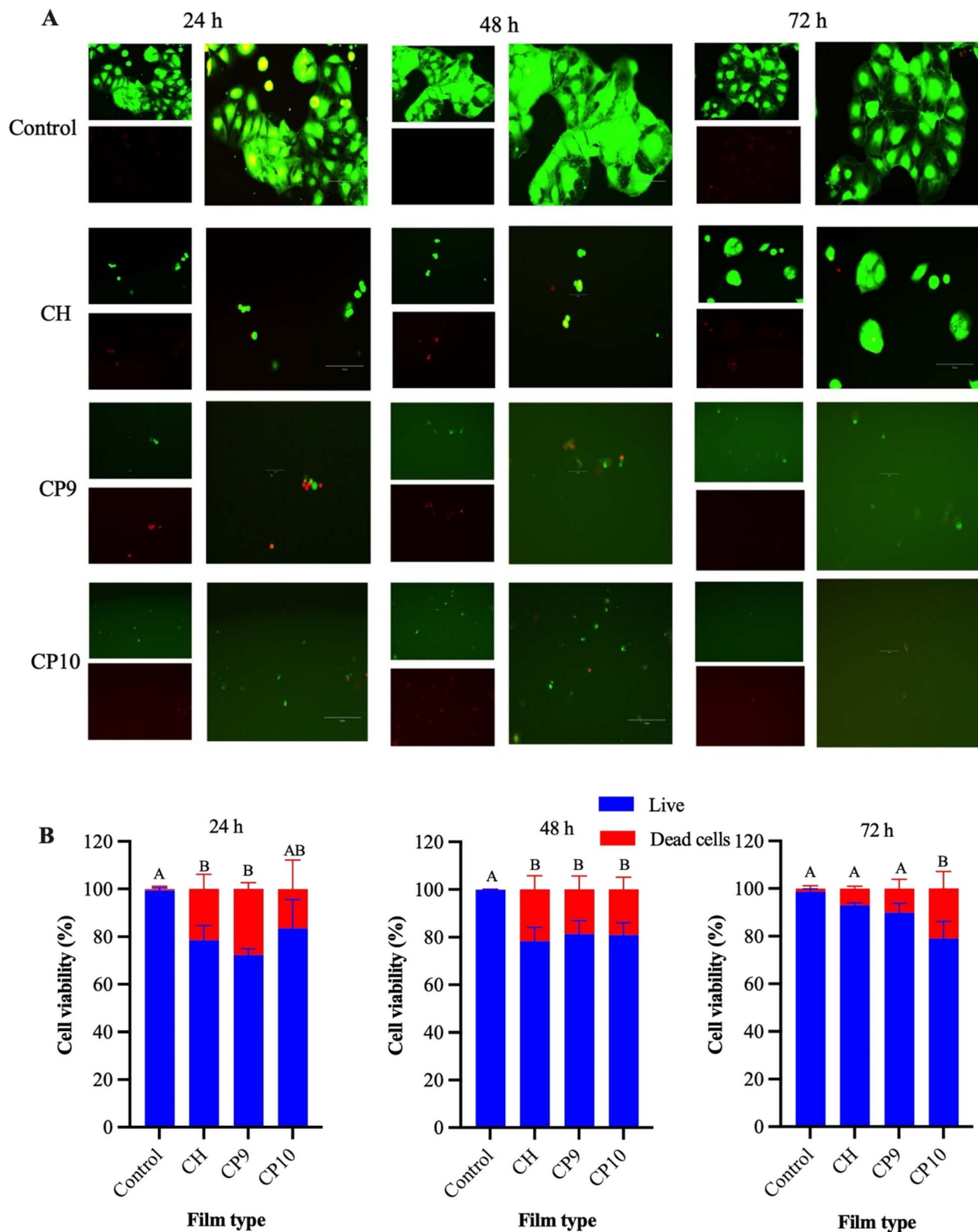


Fig. 5 Dye-based fluorescent microscopy images (A) and cell viability (B) of Caco-2 cells after 24, 48, and 72 h of incubation with CH, CP9, and CP10 films. Bars represent mean \pm SD with different letters (A and B) that are significantly different at $p < 0.05$, $n = 3$.

findings confirm the point that the fabricated films had antimicrobial ability that extended the shelf life of the tomatoes. On day 1, all the tomatoes (NK: $3.48 \pm 1.76\%$, control: $2.30 \pm$

0.38% , CH: $1.64 \pm 0.38\%$, CP9: $1.94 \pm 0.04\%$, CP10: $1.25 \pm 0.07\%$) showed more or less similar weight loss. However, by day 7, the NK group demonstrated the highest weight loss (37.5



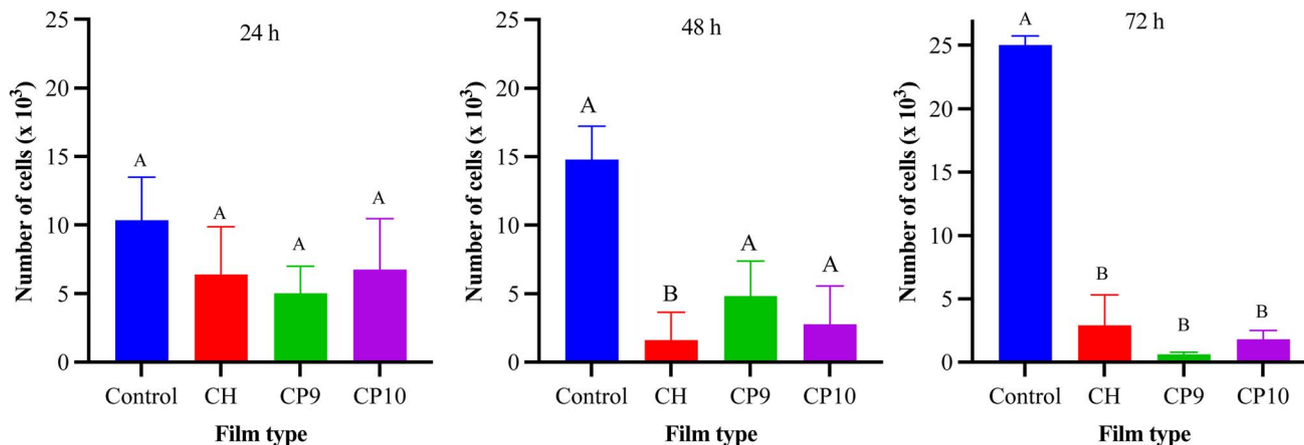


Fig. 6 Cell proliferation of Caco-2 cells after 24, 48, and 72 h of incubation with CH, CP9, and CP10 films. Bars represent mean \pm SEM with different letters (A and B) that are significantly different at $p < 0.05$, $n = 4$.

$\pm 1.32\%$), followed by the control ($16.57 \pm 1.11\%$), CH ($14.29 \pm 1.25\%$), CP9 ($13.56 \pm 1.5\%$), and CP10 ($9.63 \pm 0.55\%$). These findings are similar to those of Gasti *et al.*,⁵⁴ who reported unpacked green chillies demonstrating the highest weight loss compared to chitosan and gallic acid-loaded chitosan film-packed green chillies.

A similar observation was made for the grape samples. Non-packaged green grapes demonstrated the highest weight loss ($27.11 \pm 0.62\%$) and shrivelling during the storage periods compared to film-packed green grapes (control: $15.95 \pm 0.25\%$, CH: $14.77 \pm 1.2\%$, CP9: $13.1 \pm 0.94\%$, and CP10: $13.54 \pm 0.47\%$) (Fig. 7C and D). The current research findings align with those of Zhao *et al.*,¹¹ who observed the highest percentage of fresh-cut apple weight loss in an unpacked group compared to apple storage with the chitosan film and PVA-gallic acid incorporated chitosan film. Another study reported that non-packaged kiwifruit demonstrated shrinking and the highest weight loss compared to packing kiwifruit with the polylactic acid film containing chitosan and alizarin.³⁶ The growth of microbes was not observed with the naked eye in green grapes during the entire storage. This could be possible due to the presence of phenolic compounds, including tannins, anthocyanins, and flavonols, in grapes, which have preventive ability towards microbial growth.⁵⁵ Sun *et al.*⁵⁶ reported that unpacked grapes (*Vitis vinifera* L. Kyoho) demonstrated microbial growth and spoilage signs after 14 days of storage periods. In contrast, grapes packaged with chitosan enriched with montmorillonite and lauroyl arginate ethyl showed no microbial growth up to 20 days of storage periods.

3.5 Application of films in pearl fish fillet packaging

The appearance, drip loss, pH, and TVBN value of pearl fish fillets under different film-packaging conditions are presented in Fig. 8. It can be seen that the pearl fish fillets look shiny and show a 'fresh' colour on day 0 compared to day 2. Also, the shrinkage of pearl fish fillets was observed from day 2, especially in the NP group (Fig. 8A). There were significant differences in the drip loss of pearl fish fillets between the NP group and the film-packed group (Fig. 8B). The non-packaging and

control pearl fish fillets demonstrated 16.27% and 3.33% drip loss on day 2, respectively, whereas film-packed group showed below 3.22% drip loss. However, there are no significant differences in the drip loss of pearl fish fillets between the control and film-packed groups. This observation could be attributed to the fact that, in the current study, a small portion of the films was used, as compared to other studies where whole fish fillets were wrapped in the packaging film.^{36,57}

The pH of fresh pearl fish fillets was 6.7 ± 0.02 , and it can be seen from Fig. 8C that the pH of both the film-packed and NP groups of pearl fish fillets increased during storage. The pH of the non-packaging group pearl fish was increased to 8.92 ± 0.03 on day 2, followed by the control (8.82 ± 0.01), CH (8.76 ± 0.03), CP9 (8.64 ± 0.01), and CP10 (8.61 ± 0.01) films. The pH value of pearl fish fillets was significantly lower in the CP9 and CP10 film-packed fish compared to the NP, control, and CH packaging groups. The findings indicate that CP9 and CP10 films in packaging pearl fish could minimise the microorganism action and inhibit chemical deteriorative action, resulting in lower pH compared to other groups. The findings are consistent with the work of Yan *et al.*,⁵⁸ Amaregouda *et al.*,⁵⁹ and Wu *et al.*³⁶

The TVBN (mg/100 g) value of fresh pearl fish fillets was 5.04 ± 0.01 . The TVBN value of NP and film-packed pearl fish fillets was increased during storage (Fig. 8D). The TVBN value of NP (31.08 ± 1.45) and control (27.72 ± 2.52) pearl fish fillets was significantly higher than that of CH (23.51 ± 1.44), CP9 (24.36 ± 1.44), and CP10 (19.32 ± 1.46) film-packed fish fillets on day 1. However, the TVBN value increased to 57.96, 47.04, and 47.88 on day 2 in NP, control, and CH-packed pearl fish fillets, respectively, which is significantly higher than the changes observed in CP9 (45.36) and CP10 (41.16) films. The findings imply that fabricated films could minimise the decomposition of pearl fish fillets by diminishing antimicrobial action and lipid oxidation.⁶⁰ The findings in this study align with those of Chen *et al.*⁶¹ and Dong *et al.*,⁶² who developed active packaging films for hairtail fish and shrimp for shelf life extension and identified lowered TVBN compared to the control, respectively. Also, several research articles stated that fish is considered spoiled when the TVBN value exceeds 30 mg/100 g.^{63,64} In the



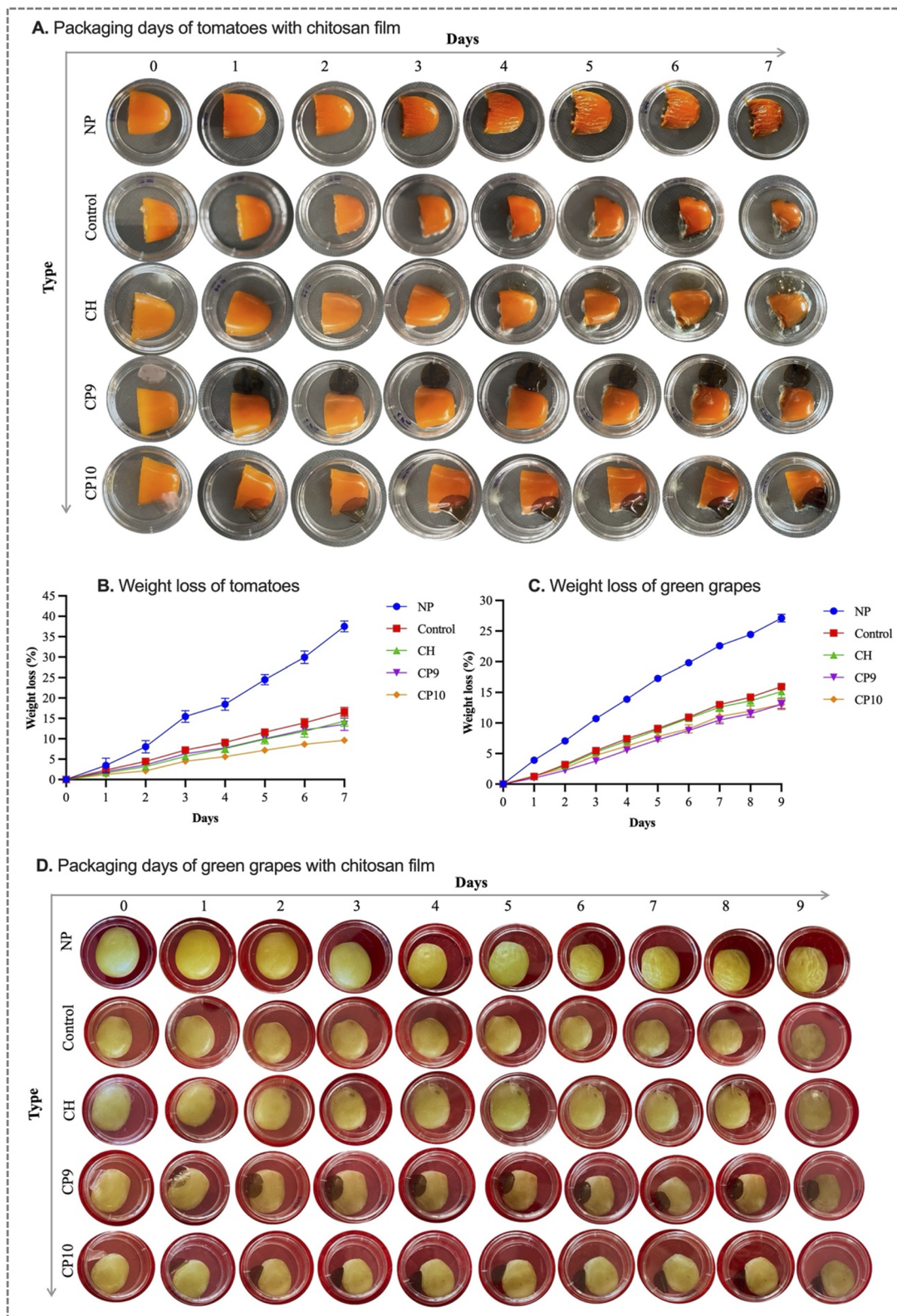


Fig. 7 Application of chitosan films (CH, CP9, and CP10) in freshness evaluation of tomatoes (A) and green grapes (D) stored at room temperature (23 °C) for 7 and 9 days, respectively, and their changes of weight loss (B and C). Non-packaging (NP): tomatoes or green grapes without a lid and film. Control: tomatoes or green grapes in a Petri dish without a film. CH: tomatoes or green grapes with a lid and chitosan film.



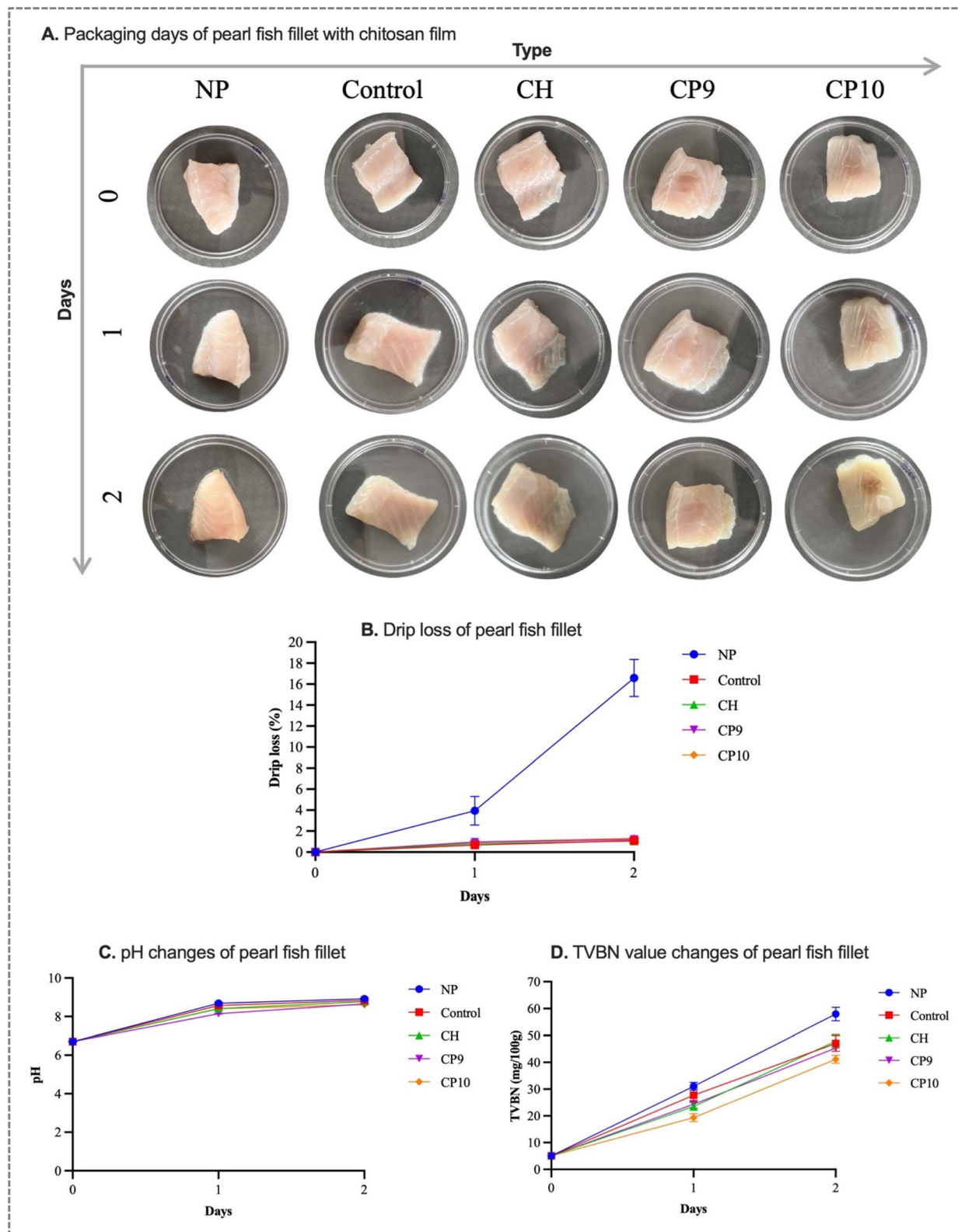


Fig. 8 Application of chitosan films in freshness evaluation of pearl fish fillet (A) stored at room temperature (23 °C) for two days, and their changes in drip loss (B), pH (C), and TVBN (D). Non-packaging (NP): pearl fish fillet without a lid and film, control: pearl fish fillet in the Petri dish without a film, and CH: pearl fish fillet with a lid and chitosan film.



current study, the TVBN value of pearl fish fillets was much lower in CP9 and CP10 film-packed fish on day 1; however, it exceeded the 30 mg/100 g cutoff value on day 2. These findings indicate that the films can potentially extend the shelf life of pearl fish fillets.

4 Conclusion

In the present research work, the functional properties of chitosan films containing PVA, COS, and gallic acid were investigated and evaluated. The COS and gallic acid-loaded chitosan-PVA films showed the highest scavenging capacity towards DPPH and ABTS radicals, and the highest FRAP was obtained from the film containing 10% COS and 10% gallic acid (CP10). In addition, the CP10 film demonstrated the highest antimicrobial efficacy towards *Escherichia coli* and *Listeria innocua*, and CP9 exhibited the highest leakage of cellular components towards *Saccharomyces cerevisiae*. Also, the highest cell viability of HaCaT and Caco-2 cells was observed on CH and CP9 films compared to CP10. Moreover, the application of CP9 and CP10 films on tomatoes and green grapes showed the lowest weight loss, and pearl fish fillets demonstrated the lowest pH and TVBN compared to CH, control, and non-packaging groups. Based on current research findings, chitosan films containing PVA, COS, and gallic acid could be considered functional and suitable packaging materials to extend the shelf life of perishable foods. Further studies may be necessary to examine the release profile of COS and gallic acid, focusing on their effects on cellular responses, microbial load data, and the quality indices of tomatoes and green grapes. Furthermore, a shelf ageing study of the packaging film could be conducted to evaluate the long-term stability of its functional properties during storage. This research could improve the understanding of the film's functional performance and the synergistic interactions between COS and gallic acid in practical food packaging applications.

Author contributions

Shuva Bhowmik: conceptualization, investigation, methodology, data curation, visualization, writing – original draft. Dominic Agyei: supervision, writing – review & editing. Azam Ali: supervision, conceptualization, project administration, writing – review & editing.

Conflicts of interest

The authors declare that they have no conflicts of interest.

Data availability

The authors affirm that the data supporting this study's findings are available within the research article and will also be shared upon request.

Supplementary information (SI) is available. See DOI: <https://doi.org/10.1039/d5fb00775e>.

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