Expanding horizons of active packaging: Design of consumer-controlled release systems helps risk management of susceptible individuals

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Abstract

The objective of this study was the development of zein based antilisterial films that contain a consumer-controlled and pH-triggered release mechanism for lysozyme (LYS). For this purpose, composite films were formed by mixing hydrophobic zein with hydrophilic soy protein isolate (SPI) or lentil protein isolate (LPI). Active property of films was formed by maintaining 30 to 50% of total LYS in free form (LYSfree). On the other hand, the pH-triggered release mechanism was formed by exploiting attractive charge-charge interactions between LYS (pI: 11.4) and SPI or LPI (pI values = 4.5), and binding remaining LYS (LYSbound) in film matrix. The pH-triggered release mechanism of composite films worked in buffers that had pHs varying between 4.3 and 7.3. The composite films bound majority of LYS between pH 5.3 and 7.3, but they released LYS at pH < 4.5. The pH-triggered release of LYS was achieved with zein-LPI films activated by acidification in packed cold-stored beef, lamb and smoked salmon. The zein-SPI films performed pH-triggered LYS release in packed food with the exception of packed smoked salmon. The LYSfree and LYSbound released from pH-triggered films showed antimicrobial activity on Listeria innocua. Consumer-controlled release mechanisms enable increasing antimicrobial stress over pathogens during transfer from market to home and cold storage at home. Thus, such films could initiate personalized packaged food to help risk management of susceptible individuals.

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1. Introduction

The increasing threats of foodborne outbreaks related to raw, minimally processed or ready-to-eat food products have boosted the research in the field of active packaging technologies such as antimicrobial, antioxidant, and bioactive packaging (Cagri, Ustunol, & Ryser, 2001; Lu, Zhu, Li, & Chen, 2015; Lynch, Tauxe, & Hedberg, 2009). Antimicrobial packaging is the most promising active packaging technology since it could be used as an effective hurdle to increase safety and/or shelf-life of food (Appendini & Hotchkiss, 2002; Gennadios, Hanna, & Kurth, 1997; Han, 2000; Ouattara, Simard, Piette, Bégin, & Holley, 2000; Quintavalla & Vicini, 2002). Antimicrobial packaging targets mainly the food surface on which microbiological changes occur most intensively (Appendini & Hotchkiss, 2002). Thus, it uses lower amounts of antimicrobials than adding antimicrobials into bulk of food. The use of natural antimicrobial compounds in edible films attracts a particular interest since health concerns and environmental problems originating from chemical food additives and plastics have been increasing continuously (Cha & Chinnan, 2004; Han, 2003; Pérez-Pérez, Regalado-González, Rodríguez-Rodríguez, Barbosa-Rodríguez, & Villaseñor-Ortega, 2006; Suppakul, Miltz, Sonneveld, & Bigger, 2003).

Hen egg white LYS is one of the most potential candidates for antimicrobial packaging since (1) it has a Generally Recognized as Safe (GRAS) status, (2) it shows good stability and activity in different films and food systems under refrigeration (Mecitoglu et al., 2006; Unalan, Korel, & Yemencioglu, 2011), and (3) it has been tested extensively in major edible film materials (Cha, Choi, Chinnan, & Park, 2002; Park, Daeschel, & Zhao, 2004; Bower, Avena-Bustillos, Olsen, McHugh, & Bechtel, 2006; Mecitoglu et al., 2006; Joerger, 2007; Mendes de Souza, Fernández, López-Carballoc, Gavara, & Hernández-Munoz, 2010). The antimicrobial mechanism of LYS originates from its lytic activity that causes splitting of the bonds between N-acetyl muramic acid and N-acetylgalactosamine of the peptidoglycan in Gram-positive bacterial cell wall (Appendini & Hotchkiss, 2000; Quintavalla, Vicini, & Bigger, 2004).

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Hotchkiss, 1997; Shah, 2000). Some Gram-positive pathogenic bacteria such as Staphylococcus aureus show extreme resistance against LYS (Sudagdan & Yemencioglu, 2012), but LYS shows high potency against the critical bacterial pathogen Listeria monocytogenes (Duan, Park, Daeschel, & Zhao, 2007). It is essentially important to control contamination and growth of L. monocytogenes in food since this bacterial pathogen regularly causes deadly infections in pregnant women, and elderly and immunosuppressed people (Vazquez-Boland et al., 2001; Álvarez-Ordóñez, Leong, Hickey, Beaufort, & Jordan, 2015). Antimicrobial packaging could be used as a highly effective hurdle to prevent the growth of L. monocytogenes in risky foods that contaminate easily with this pathogen and enable its growth. For example, Min, Rumsey, and Krochta (2008) successfully employed LYS in whey protein films to control L. monocytogenes growth on smoked salmon. Unalan, Arcan, Kørel, and Yemencioglu (2013) employed LYS containing zein composite films to prevent the growth of L. monocytogenes in fresh cheese. In these studies it was aimed to maximize free LYS in the films by minimizing interactions and bonding between LYS and edible film matrix. The presence of free LYS ensures the initiation of enzyme release from films onto food surface and provides an antimicrobial activity.

Boyaci, Korel, and Yemencioglu (2016) have developed LYS containing whey protein based edible films with a novel concept named activate-at-home type packaging (ActiHome packaging). These authors employed this concept in preventing listerial growth in cold-smoked salmon. ActiHome packaging is a novel concept which means that packaging material is activated by consumers to form a hurdle for the development of critical pathogenic bacteria in food kept at home in refrigerators until consumption (Boyaci et al., 2016). ActiHome packaging could be an alternative technology to reduce the risk of food poisoning from L. monocytogenes whose growth cannot be controlled by traditional packaging methods such as vacuum packaging (VP) (Duffes, 1999; Gram, 2001). Some studies have shown that it is possible to control L. monocytogenes growth in risky food by modified atmosphere packaging (MAP) containing elevated levels of CO₂ (100% CO₂) (Michaelsen, Sebranek, & Dickson, 2006; Rutherford et al., 2007). However, the protective effect of MAP is lost after opening the package, and remaining food kept for later consumption becomes quite risky for the development of L. monocytogenes (Buchanan & Klawitter, 1990; Davies, 1997; Tsigarida, Skandamis, & Nychas, 2000). In ActiHome packaging concept, edible films containing both soluble and bound antimicrobials are used for packaging or coating food products. Release of soluble antimicrobials provides antimicrobial effect at the food surface during transportation, storage, and marketing (Fig. 1A). The release of bound antimicrobials starts after activation of the film by the consumer immediately after purchasing the product at the market (Fig. 1B). Alternatively, the consumer could activate the packaging at home. This is reasonable when a protective packaging (MAP or VP) is opened at home shortly after purchase, and only part of the food is consumed (Fig. 1C). The first generation of ActiHome packaging materials developed by Boyaci et al. (2016) from whey protein were quite successful to bind and immobilize positively charged LYS (pl at 11.4) onto negatively charged whey protein (pl between 4.4 and 5.4) films in the pH range of smoked salmon (between pH 6.0 and 6.3). However, due to the excessive negative charges of whey protein film matrix, these films bound and immobilized all incorporated LYS effectively. Thus, they lacked free LYS and were not active unless their pH-triggered release mechanism was activated by the consumer by wetting the film with acidicified lemon juice. In the present study, zein based composite packaging materials have been developed with both active and activate-at-home type properties. Zein is a protein obtained as a byproduct during corn processing by the bioethanol and oil industries (Selling, Woods, Sessa, & Biswas, 2008; Xu, Reddy, & Yang, 2007). It is a great candidate as an edible film-forming agent since it has excellent film-forming ability and good solubility in solvents like ethanol. Zein is a highly hydrophobic protein since it contains very limited number of hydrophilic amino acids that could form interactions with incorporated hydrophilic antimicrobial agents. Thus, protein-based antimicrobial agents like LYS remain mainly in free and soluble form when they are incorporated into zein films (Mecitoglu et al., 2006; Gűçbilmez, Yemencioglu & Arslanoglu, 2007).

Therefore, composites of hydrophobic zein were formed by using suitable amounts of LPI or SPI that contain negatively charged groups (pl value ≈ 4.5) to bind part of the positively charged LYS. This strategy allows maintaining desired amounts of LYS in free soluble form while putting sufficient amounts of LYS in bound form within the LPI or SPI fraction in the film matrix. In this work, the presence of LYSfree and the good working of the proposed pH-triggered release mechanism for LYSbound in the composite films was proved by conducting extensive release tests in buffers and on selected food samples. Sliced beef, lamb meat, and smoked salmon were selected as food samples since they have been reported to pose risk of listeriosis (Dalgaard & Jørgensen, 1998; Vermeulen, Devlieghere, De Loy-Hendrickx, & Uyttendaele, 2011; Di Cicco et al., 2012; D’Ostuni, Tristezza, Giorgi, Rampino, Grieco, & Perrotta, 2016; Smith et al., 2011). Antimicrobial activity of films originated from their LYSfree and LYSbound was also demonstrated on Listeria innocua. This work presented the second generation of ActiHome packaging materials that could be used both as a traditional antimicrobial film and as an advanced additional hurdle controlled by the consumer.

2. Materials and methods

2.1. Materials

Corn zein and LYS were obtained from Sigma Chem. Co. (St. Louis, MO, USA). Lentil (Cultivar Alidayı) used to produce LPI, was provided by the General Directorate of Agricultural Research in Ankara, Turkey. Soybean (non-GMO) used to produce SPI, and beef, lamb meat, and cold-smoked salmon were purchased from local supermarkets in İzmir, Turkey. The salmon sample was cold-smoked and vacuum packed, and it contains 3.5% salt (product information). Fresh lamb and beef samples were from leg parts with the fascia layer on their surface. Bacterial strain of Listeria innocua (NRRL B-33314) used in antimicrobial tests was provided by United States Department of Agriculture, Microbial Genomics and Bioprocessing Research Unit, Peoria, Illinois (USA).

2.2. Preparation of lentil and soy protein isolates

LPI and SPI were extracted from dry lentil and soy beans. To remove lipids and phenolic compounds, lentil and soy seeds were first processed to acetone powder (AP) according to the method given by Arcan and Yemencioglu (2007). The APs were stored at -18 °C until they were used for protein extraction. Protein extraction was conducted by mixing 50 g of AP with 200 mL of distilled water, adjusting pH of the mixture to 9.0 with 1 N NaOH, and applying continuous stirring for 30 min at room temperature. To remove insoluble debris the extract was then centrifuged at 11,000 x g for 30 min at 4 °C. The supernatant containing solubilized protein was then separated, and its pH was adjusted to 4.5 with 1 N acetic acid solution to precipitate the proteins. The precipitated proteins were then collected by centrifugation at 11,000 x g for 30 min at 4 °C, and they were dissolved again in 150 mL distilled water. The solubilized protein was then precipitated for the second time as described above by bringing extract pH...
antilisterial activity during transportation, marketing and storage; B and C: activate-at-home type property originating from LYS-bound release of following activation provides additional antilisterial activity based on different scenarios). (Heidolph, Germany, rotor F

Dawson (1998). Briefly, 1.4 g zein was dissolved in 8.2 mL ethanol (96%) by mixing slowly with a magnetic stirrer for 25 min. Glycerol (0.4 mL) was added into the solution as plasticizer. The film-forming solution was then heated to boiling point and boiled for 5 min for denaturation of zein proteins. After cooling to room temperature, LPI or SPI at 130 or 390 mg/g of zein (1.5 or 4.5 mg/cm² of dried film), and LYS at 61 mg/g of zein (0.7 mg/cm²) were added into film-forming solution. The solution was then homogenized at 10,000 rpm for 4 min using a homogenizer-disperser (Heidolph, Germany, rotor Φ = 6.6 mm tip). After that, 4.3 g of film-forming solution was spread onto 8.5 cm × 8.5 cm glass plates, previously cleaned with ethanol. Casted films were dried at 25 °C for 20±2 h.

2.3. Preparation of films

Zein-based composite films were prepared by modifying the standard zein film-making method described in Padgett, Han, and Dawson (1998). To 4.5, collecting precipitated protein by centrifugation, and dissolving precipitate in distilled water. The pH of protein solution was then adjusted to 7.0 with 1 N NaOH, and the solution was lyophilized using a freeze dryer (Labconco, Kansas City, KS, USA). The lyophilized LPI and SPI were stored at -18 °C until they were used in film preparation.

2.4. Determination of LYS activity

LYS activity was measured spectrophotometrically at 660 nm by using Shimadzu (Model 2450, Japan) spectrophotometer equipped with a constant temperature cell holder at 30 °C. The reaction mixture was prepared by mixing 0.1 mL enzyme containing solution (incubated at 30 °C for 1 min) and 2.4 mL Micrococcus lyso- dexticus suspension (at 30 °C) prepared in 0.05 M Na-phosphate buffer at pH 7.0. The reaction mixture was mixed with a vortex, and the decrease in absorbance was monitored for 120 s. Enzyme activity was calculated from the slope of the initial portion of absorbance vs. time curve, and it was expressed as Units released per cm² of films. One Unit was defined as 0.001 change in absorbance in 1 min. Average of three measurements was used in calculations. The activity of LYS solution determined by this method was 74548 U/mg of solution.

2.5. Test of designed pH-triggered LYS release mechanism in buffers at different pH

The working of the designed pH-triggered release mechanism of films was tested in buffers at different pH values. This test shows not only the activity of LYSfree, but also shows the activity of LYS-bound immobilized by charge-charge interactions. The release profiles of LYS at different pH was evaluated by incubation of films (zein, zein–LPI or zein–SPI films containing 0.7 mg/cm² LYS) at 4 °C for 24 h in series of buffers (50 mL of Na-phosphate buffers at pH 7.3, 6.3, 5.3 or Na-phosphate–HCl buffer at pH 4.3) and monitoring their released LYS activities. The films cut into 16 cm² (4 cm × 4 cm) pieces were sequentially kept firstly in buffer at pH 7.3 for 24 h, secondly in buffer at pH 6.3 for 24 h, thirdly in buffer at pH 5.3 for 24 h, and lastly in buffer at pH 4.3 for 24 h. The LYS activity was measured as described in section 2.4 by taking 0.1 mL samples periodically and conducting three measurements (3×0.1 mL) at each incubation time. Calculations were corrected by considering activities removed from the aqueous media during sampling. The total activity of released LYS from a film was calculated by determining the sum of each activity at pH 7.3, 6.3, 5.3 and 4.3 within 96 h (4 × 24 h). The total activity of released LYS from control zein film was accepted as 100%. All other activities were reported as percentage in respect to total activity of released LYS from control zein film. The results were expressed on histograms by plotting released activity (%) from each type of film within 24 h at different buffers.
2.6. Test of designed pH-triggered LYS release mechanism in different foods

The pH-triggered release mechanism of films was also tested on slices of beef, lamb meat, and cold smoked salmon. Briefly, zein, zein-LPI (4.5 mg/cm² LPI) or zein-SPI (4.5 mg/cm² SPI) films (2 cm × 2 cm) containing 0.7 mg/cm² LYS were placed on one side of the food slices (4–4.5 g of beef and lamb meat, 1.4–1.5 g of salmon). The films on beef and lamb meats were in contact with the fascia/epimysium layer of the muscle. The coated samples were then wrapped with a plastic stretch film, and then with an aluminum foil to increase film-food contact. The samples were then cold-stored at 4 °C for 2 days to cause release of LYSfree in films. At the end of 2 days, the packs were opened, and 0.2 mL of 10% (v/v) lactic acid solution was pipetted onto surfaces of the films to activate films. The acidification aimed to drop film pH ≤ 4.5, a suitable pH to initiate release of bound LYS in the film matrix. The coated samples were then packed again (with plastic stretch films and aluminum foil) and cold-stored at 4 °C for 5 days (total 7 days) to monitor release of LYS by acidification. A control of each film group was also used in packaging of different food, but these groups were cold-stored without acidification. During cold-storage, some packs were taken every 24 h, and assayed for LYS activity after enzyme extraction. For extraction of LYS, the films were first removed from the food surface. Food sample was then homogenized for 1 min in 10 mL of buffer (0.1 M PBS at pH 6.0) using a blender (Waring with 30 mL jar, USA, Model: 7011HS). The homogenate was then clarified by centrifugation at 11,000 × g for 10 min at 4 °C. LYS activity was determined by using 0.1 mL of supernatant in triplicate (3 × 0.1 mL) as described in section 2.4. The activity of LYS released into sample was then calculated and expressed as Units released per cm² of the films.

2.7. Antimicrobial activity of films

The antimicrobial activity of films was determined by the modification of the classical zone-inhibition assay, using Listeria innocua (NRRL B-33314) as test microorganism. The inoculum was prepared in peptone water (0.1%), using a 24 h culture of L. innocua growth on nutrient agar incubated at 37 °C. The cell concentration was set to 1.0 McFarland unit (3 × 10⁸ CFU/mL). Sixteen discs from each film were cut by a sterile cork-borer, and the discs (13 mm in diameter) were placed on agar plates inoculated by L. innocua. Petri dishes were incubated at 37 °C for 24 h, and the area of the clear zones formed around the discs was measured and calculated using a digital micrometer. These results originated from LYSfree in the films. After the measurement of zones, the discs on the agar plates were carefully removed with a sterile pincher and transferred to surfaces of newly inoculated fresh agar plates. Each type of film discs were then separated into two groups. One of the two disc groups was then acidified to activate pH-triggered LYS release by pipetting 10 μL of 0.5 M sterile Na-acetate buffer at pH 4.3 onto film surfaces. The other groups were monitored as control, and they were not acidified. All the plates were then incubated at 37 °C for 24 h, and the area of the clear zones formed around transferred discs was calculated after measuring the zone diameters using a digital micrometer. The results from acidified films originated mainly from residual soluble LYSfree left in the films following first incubation, and LYSbound liberated by acidification applied before initiation of second incubation.

2.8. Mechanical and morphological properties of films

Mechanical properties of films were evaluated by measuring tensile strength, elongation at break and elastic modulus of films using the TA.XT-2 texture analyzer (Stable Microsystems, Godalming, UK) according to ASTM Standard Method D-882-02 (ASTM, 2002). Films were conditioned in an environmental chamber at 25 °C and 50% RH for 24 h. For mechanical tests, films were cut into 8 mm-wide and 80 mm-length strips. The initial grip distance was 50 mm, and the crosshead speed was 50 mm/min. At least seven replicates of each film were tested. Tensile properties were calculated from the plot of stress versus strain.

The film morphologies were determined by obtaining their cross-sectional photographs using a scanning electron microscope (SEM) (Philips XL 30S FEG, FEI Company, Eindhoven, Netherlands). Film strips with 8 mm width were first broken in liquid nitrogen, and then they were coated by gold palladium for 1 min in a Magnetron Sputter Coating Instrument (Emitech K550X, Quorum Technologies Inc., UK) before photographed in the SEM. The average thicknesses of films were measured from the 5000× magnified micrographs. The film thicknesses were measured from SEM cross-sectional views by using Scandium software (Olympus Soft Imaging Solutions GmbH, Münster, Germany).

2.9. Statistical analysis

Statistical analysis was performed by using MINITAB® release 14 (Minitab Inc., State College, Pa., U.S.A.). Data were analyzed by using one-way analysis of variance (ANOVA). Significance was accepted at p < 0.05.

3. Results and discussion

3.1. Test of designed pH-triggered release mechanism in different buffers

LYS release profiles of zein, zein-LPI and zein-SPI films in different buffers are presented in Figs. 2 and 3. Tests conducted at pH 7.3 for 24 h clearly showed the release of LYSfree from different films. Due to the limited number of negatively charged groups in zein (Shukla & Cheryan, 2001; Argos, Pedersen, Marks, & Larkins, 1982), 80 to 90% of total activity for LYS released from zein control films was due to LYSfree released from films at pH 7.3. It is clear that the addition of LPI or SPI at 1.5 mg/cm² did not considerably affect the activity of LYSfree released from zein-LPI and zein-SPI films at pH 7.3. In contrast, the increase of LPI and SPI from 1.5 to 4.5 mg/cm² caused almost 48% and 58% lower LYSfree release from zein-LPI and zein-SPI films than related zein controls at pH 7.3.

![Fig. 2. Activities of LYS released from control film (1) and LPI film (1.5 or 4.5 mg/cm²) at each of buffers at pH 7.3, 6.3, 5.3 and 4.3.](image)
respectively. These results suggested that the negative charge formed at pH 7.3 in composite films containing SPI or LPI at 4.5 mg/cm² caused binding of a significant portion of positively charged LYS. Moreover, it is important to report that the sum of activities for LYS released from zein-LPI film at pH 6.3 and 5.3 was almost 5-fold higher than that released from zein-SPI film at the same pH values. The recent study of Aydemir and Yemencioglu (2013) showed that pl for lentil protein fractions varied in a wide range between pH 4.5 and 6.5. Thus, it seemed that the reduced negative charge intensity of some lentil protein fractions in the zein-LPI film matrix at pH 6.3 and 5.3 caused the release of part of the LYSbound from zein-LPI films. On the other hand, the transfer of zein-SPI and zein-LPI films from buffer at pH 5.3 to buffer at pH 4.3 had initiated release of considerable amounts of LYS from films. It was clear that the reduced pH caused loss of negative charges on the majority of LPI and SPI fractions in the composite film matrix, and this liberated a significant portion of LYSbound from the films. Activity for released LYSbound at pH 4.3 from composite films with 4.5 mg/cm² LPI and SPI was almost 30% and 61% of total activity for released LYS from these films at different pH values (at pH 7.3, 6.3, 5.3 and 4.3 within 96 h), respectively. The higher LYSbound of zein-SPI films than zein-LPI films at pH 4.3 suggested that the SPI is particularly rich in protein fractions with low pl value (<pI 5.3). Thus, LYSbound in zein-SPI films released as a bulky fraction only when pH dropped from 5.3 to 4.3, slightly below their reported pl of 4.9 (Okubo, Waldrop, Iacobucci, & Myers, 1975). In contrast, LPI contains protein fractions with pl between 4.3 and 6.3. Thus, LYSbound in zein-LPI films released gradually as pH dropped from 7.3 to 4.3. The results of release tests also showed that the inherently charged groups in zein control films also had a limited LYS binding capacity. The sum of activities for LYS released from zein control films at pH 5.3 and 4.3, (pH values below pl of zein, 6.2), changed from 7 to 13% of the total activity for LYS released from these films at different pH values (at pH 7.3, 6.3, 5.3 and 4.3 within 96 h).

The total activity of LYS released into buffers at different pH values within 96 h also helped calculation of yield for films based on released enzyme activity (Yield of films for LYS activity = (Total activity of LYS released from films/Activity of LYS incorporated into films) x 100). The average LYS activity yield for different control zein films (n = 2) was 91%. On the other hand, LYS activity yields for composite films with 1.5 and 4.5 mg/cm² LPI were 105% and 89%, respectively. The yield over 100% for the composite film with LPI at 1.5 mg/cm² suggested some slight heterogeneity in LYS distribution of this film (a 4 cm × 4 cm piece from a 8.5 cm × 8.5 cm cast film). The higher yield of this composite film than the others also suggested the reduced entrapment of LYS in this film. It seemed that the distribution of hydrophilic LPI molecules within composite film matrix reduced the fraction of LYS physically entrapped within the hydrophobic zein clusters. In contrast, similar activity yields of composite film with 4.5 mg/cm² LPI and control film suggested some LYS entrapment by aggregated LPI and/or LPI–zein complexes. On the other hand, LYS activity yields of composite films with 1.5 and 4.5 mg/cm² SPI were 77% and 74%, respectively. These results suggested significantly higher capacity of SPI than that of LPI to cause physical entrapment of LYS within zein film matrix.

3.2. Test of designed pH-triggered release mechanism in different packed foods

3.2.1. LYS release profiles before acidification of films (24th and 48th h of cold storage)

Results of release tests conducted with slices of lamb meat, beef, and smoked salmon packed with developed films, and cold stored at 4 °C are presented in Fig. 4. The release of LYSfree from different films onto lamb meat before activation conducted by acidification of films was seen in Fig. 4A and C. The activity of LYS released onto lamb meat from zein films is almost 3 and 2.4-fold higher than those released from zein-LPI and zein-SPI films at the end of 48 h, respectively. This result was expected since previous release tests conducted in buffers proved the lower amounts of LYSfree in composite films than in zein control film.

The activity of LYS released onto beef samples from zein films at the end of 48 h is also almost 3.3 and 3.5-fold higher than that released from zein-LPI and zein-SPI films, respectively. Initial release profiles of LYS from zein films onto beef and lamb samples were similar (Fig. 4D). Zein-LPI films also showed similar initial LYS release profiles for beef and lamb samples in the first 48 h (Fig. 4E). However, it is important to note that zein and zein-LPI films showed almost 33% and 28% higher LYS activity on beef than that on lamb meat within 48 h. On the other hand, the initial LYS release profiles of zein-SPI films followed a different profile in lamb and beef samples (Fig. 4F). Release of LYS from zein-SPI films onto lamb meat occurred rapidly in the first 24 h, but instability of enzyme in lamb meat caused a moderate drop (-20%) in its LYS activity at 48 h. In contrast, LYS release from zein-SPI films onto beef occurred slowly, but it increased continuously and reached almost the same activity with lamb meat at the end of 48 h.

Activity measurements suggested that all films released much more LYS on smoked salmon than that on lamb meat and beef within shorter time periods (Fig. 4G and I). For example, LYS activities measured for smoked salmon packed with zein, zein-LPI and zein-SPI films within 24 h were 1.5–6.4-fold higher than those of beef and lamb meat. It seemed that the fascia/epimysium layer at the surfaces of lamb meat and beef acted as a barrier for the film swelling, and this slowed down the diffusion of LYS from films onto these samples. In contrast, films placed on salmon slices might have swelled rapidly since these samples lacked such a continuous protective layer. It is also interesting to report that zein and zein-LPI films released almost the same amount of LYSfree onto the salmon surface at the end of 24 h. This result suggested that the release of LYSbound initiated (triggered) in zein-LPI films before acidification. It seemed that the salt used in salmon curing (~8% (w/w)) caused the destabilization of attractive charge-charge interactions between LYSbound and zein-LPI film matrix, and this caused release of some enzyme from films. The destabilization of electrostatic interactions between a protein and a charged matrix in presence of salts is a well-known scientific truth, and this is frequently exploited to extract enzyme bound by electrostatic interactions in tissues.
(Yemenicioglu & Cemeroglu, 1999). The same principle is also used to elute enzyme and protein from ion-exchange columns during purification (Yemenicioglu, Ozkan, & Cemeroglu, 1998). In contrast, a considerable solubilization of LYS\textsubscript{bound} was not observed in smoked salmon packed with zein-SPI films. Thus, the salmon samples packed with zein-SPI films showed 1.8–1.9-fold lower LYS activity than those packed with zein and zein-LPI films at the end of 24 h. This could be related to the more powerful (extensive) attractive charge-charge interactions of LYS with zein-SPI film matrix than zein-LPI film matrix. In fact, the findings obtained with release tests conducted in buffer at different pH supported that the LYS attached more tightly on zein-SPI release tests conducted in buffer at different pH supported that the LYS/LYS activities of acidified and non-acidified samples were monitored for their LYS activities. At this stage, the release test became highly complex since there are multiple dependent variables [LYS release rate, loss in LYS activity due to instability, and activity of LYS released (LYS\textsubscript{free} and/or LYS\textsubscript{bound} in acidified/non-acidified samples)] against the independent variables (storage time and acidification). Lamb meat samples packed with non-acidified and acidified zein films showed a sharp reduction in

**Fig. 4. Activities of released LYS from acidified and non-acidified zein, zein-LPI and zein-SPI films applied on different food (A, B, C: Lamb meat, D, E, F: Beef, G, H, I: Cold-smoked salmon; Note-1: LYS\textsubscript{free} release was expected at 0th, 24th and 48th hours since film acidification was applied at the end of 48th h; note-2: LYS\textsubscript{free} and/or LYS\textsubscript{bound} release was expected at 72nd, 96th, 120th and 168th h; note-3: storage times (h) with statistically significant differences (p < 0.05) between LYS activities of acidified and non-acidified samples: A: 72, 168; B: 96, 168; C: 72, 96, 168; D: 72, 96, 120, 168; E: 72, 120, 168; F: 72, 96, 120, 168; G: 72, 96; H: 72, 96, 168; I: 96).**
their LYS activities after 48th h of cold storage. However, LYS activity in lamb meat packed with non-acidified zein films showed much more dramatic reduction (76%) than those packed with acidified zein films (54%) at the end of 168 h. These findings clearly showed the high instability of LYS in lamb meat. Moreover, the results also suggested that the pH-triggered release of LYSbound by inherently charged groups in zein control film might improve LYS activities of the product. On the other hand, the activation of zein-LPI and zein-SPI films by acidiﬁcation at the end of 48th h caused a gradual increase (started at 72nd and peaked at 96th h) and a sharp increase (at 96th h) in LYS activities of packed lamb meat, respectively. At similar acidiﬁcation conditions, 24 h earlier initiation of LYS activity increase in lamb samples coated with zein-LPI films suggested a better response of pH-triggering mechanism in these ﬁlms than that in zein-SPI ﬁlms. LYS activities of lamb meats packed with both acididiﬁed composite ﬁlms showed a slight reduction in LYS activity after 96 h. However, it is important to report that LYS activities of lamb samples packed with acididiﬁed composite ﬁlms at the end of 168h of cold storage were almost 2-fold higher than that for lamb meat packed with non-acididiﬁed control zein ﬁlms. Thus, it appeared that the acidiﬁcation to exploit pH-controlled release properties of composite ﬁlms was a more beneﬁcial delivery strategy than classical active packaging conducted with non-acididiﬁed zein ﬁlms. On the other hand, it should be reported that the activity of LYS released onto lamb meat from acididiﬁed composite ﬁlms at 168h were not considerably higher (almost 10%) than that from acididiﬁed zein ﬁlms. Thus, it is clear that the composite ﬁlms should be prepared with higher levels of LYS and LPI or SPI to increase amounts of their LYSfree and LYSbound fractions.

On the other hand, results for LYS activity measurements with cold-stored beef samples after acidiﬁcation followed a slightly different pattern than those of lamb samples. First of all, the pH-triggered release mechanism in acididiﬁed zein control ﬁlms showed its beneﬁt only at the 72nd h of cold storage by causing a peak LYS activity at 21600 U/cm². In contrast, the acididiﬁed zein-LPI ﬁlms caused a gradual increase in LYS activities of beef samples. It is important to note that in beef packed with acididiﬁed zein-LPI ﬁlms a peak LYS activity (almost 21000 U/cm²) was reached at the 120th h, two days later than that observed for beef samples packed with acididiﬁed zein ﬁlms. This result showed the possibility of achieving a sustained LYS release by using pH-triggered release mechanism in beef packed with zein-LPI ﬁlms. The pH-triggered release mechanism also worked on beef samples packed with zein-SPI composite ﬁlms. However, the release of LYS from acididiﬁed zein-SPI ﬁlms onto beef samples made a peak at the 96th h. Moreover, the maximal activity reached for beef samples packed with acididiﬁed zein-SPI ﬁlms was almost 30% lower than those reached for beef samples packed with acididiﬁed zein and acididiﬁed zein-LPI ﬁlms. This result showed that the response of pH-triggered release mechanism of zein-LPI ﬁlm on the beef surface is better than that in zein-SPI ﬁlm.

On the other hand, the overall results clearly showed that the maximum LYS activities and ﬁnal LYS activity levels reached in packed beef were higher than those of lamb meat. It seemed that differences in biochemical stability of lamb meat and beef affected the stability of LYS. The proteases (e.g. calpains, cathepsins, calpastatins and Ca-dependent proteases) play a central role in biochemical changes in meat. However, the activity and stability of proteases as well as level of protease regulating inhibitors could be variable in meats of lamb and cattle (Morton, Bickerstaffe, Kent, Dransﬁeld, & Keeley, 1999; Oualli & Talmant, 1990; Koochmarie, Whipple, Kretchman, Crouse, & Mersmann, 1990).

Release of LYS from acididiﬁed and non-acididiﬁed control ﬁlms on smoked salmon followed a quite similar proﬁle and showed a gradual increase up to 120th h of cold storage. Salmon samples packed with acididiﬁed zein ﬁlms showed signiﬁcantly higher LYS activities (1.2–1.3-fold) than those packed with non-acididiﬁed zein ﬁlms at 72nd and 96th h of storage (p < 0.05). On the other hand, some more signiﬁcant differences were observed between LYS activities of salmon samples packed with acididiﬁed and non-acididiﬁed zein-LPI ﬁlms. The activities of LYS released on salmon packed with acididiﬁed zein-LPI ﬁlms were signiﬁcantly higher (1.4–1.8-fold) than those of salmon packed with non-acididiﬁed zein-LPI ﬁlms at the 72nd, 96th and 120th h of cold storage (p < 0.05). However, it should also be reported that signiﬁcant reductions were observed in LYS activities of salmon packed with acididiﬁed (-23%) and non-acididiﬁed (-43%) zein-LPI ﬁlms after 96th and 120th h of cold storage, respectively. This result suggested the exhaustion of LYS reservoirs in zein-LPI ﬁlms at the later periods of cold storage. In contrast, the LYS activity of acididiﬁed and non-acididiﬁed zein-SPI ﬁlms increased continuously up to the 168th h of cold storage. At the 96th h of cold storage, salmon packed with acididiﬁed zein-SPI ﬁlms showed signiﬁcantly higher (1.6-fold) LYS activity than that packed with non-acididiﬁed zein-SPI ﬁlms (p < 0.05). However, LYS release proﬁles of acididiﬁed and non-acididiﬁed zein-SPI ﬁlms applied on salmon samples were quite similar at the 120th and 168th h of cold storage (p > 0.05). These ﬁndings suggested that at the later stages of cold-storage the LYSbound in non-acididiﬁed zein-SPI ﬁlms liberated spontaneously from SPI fraction in the ﬁlm matrix and released onto smoked salmon. Thus, it is once more proved that the cured products that contain salts which are capable to destabilize ionic interactions between LYS and ﬁlm matrix might interfere with the pH-controlled release system. It is important to note that the non-acididiﬁed zein-LPI ﬁlm suffered from spontaneous release of LYSbound at the beginning of storage (within the ﬁrst 48 h) while the same problem was observed in zein-SPI ﬁlms at the later stages of cold storage (>120 h). Thus, it is clear that the zein-SPI ﬁlm is a better option than zein-LPI ﬁlm when target food is rich in salts. Further studies are needed to optimize SPI content of ﬁlms for cured samples and test alternative proteins that could form more resistant ionic interactions with LYS.

3.3. Antimicrobial activity of ﬁlms

The antimicrobial activity of developed zein, zein-LPI, and zein-SPI ﬁlms were tested against non-pathogenic Listeria innocua which is used as an indicator for the critical pathogenic bacteria L. monocytogenes (Francis & Bernie, 1997). The zone inhibition tests were conducted ﬁrst with non-acididiﬁed ﬁlms to prove the active properties of ﬁlms originated from LYSfree (Table 1). The zones determined for non-acididiﬁed ﬁlms within 24 h incubation period were almost 2-fold larger than those for non-acididiﬁed zein-LPI and zein-SPI ﬁlms that gave quite similar zone areas (p > 0.05). This result was expected since the majority of the LYS in zein ﬁlms was free while composite ﬁlms obtained with 4.5 mg/cm² SPI or LPI
contained lower activities of LYS free at slightly acidic to neutral pH values (between pH 5.3 and 7.3). At the end of first 24 h incubation on agars, the films were aseptically transferred onto new agars inoculated with fresh L. innocua culture. Half of the films were acidified while the remaining films were not acidified. The formation of clear zones in non-acidified zein and non-acidified composite films at the end of second 24 h incubation period in fresh medium clearly showed the presence of considerable amounts of residual LYS free in the films. It should also be reported that the zone areas of non-acidified zein and composite films at the end of second incubation period were not significantly different from each other (p > 0.05). However, it is important to report that the zone areas of acidified zein-SPI and zein-LPI composite films at the end of second incubation were almost 1.4-fold higher than those of the non-acidified composite films. The acidified composite control films lacking LYS did not form clear zones. Thus, these results clearly indicated any considerable changes in classical highly porous nature of zein films to a dense one by incorporation of SPI and LPI contents) and secondary incubation period suggested the lack of any considerable antimicrobial potential for LYS bound in these films.

3.4. Mechanical and morphological properties of films

Mechanical properties of zein and composite films were evaluated by determining their tensile strength, elongation at break and Young’s modulus values (Table 2). Although the incorporation of LYS into zein films caused a significant reduction in their Young’s modulus, it caused no significant changes in tensile strength and elongation of films (p > 0.05). The SEM cross-section photos (Fig. 5A and B) of zein control and LYS containing zein films did not also indicate any considerable changes in classical highly porous nature of zein films (Arcan, Boyaci, & Yemenicioğlu, 2017). These results suggested the lack of any considerable interaction and networking between hydrophobic zein film matrix and hydrophilic LYS. The incorporation of 4.5 mg/cm² LPI or SPI into zein films with LYS did not cause a significant change in tensile strength of LYS containing zein films (p < 0.05). However, addition of LPI and SPI at 4.5 mg/cm² caused significant increases and reductions in elongation and Young’s modulus values of composite films (p < 0.05). The SEM pictures of film cross-sections suggested the change of highly porous nature of zein films to a dense one by incorporation of SPI and LPI into films (Fig. 5C and D). Such dramatic morphological changes in composite films suggested interaction of hydrophobic zein with LPI and SPI that are amphiphilic proteins with good surface active properties (Aydemir & Yemenicioğlu, 2013). The interactions could change the typical orientation of zein molecules in film that has been defined as a meshwork which is composed of doughnut structures formed by asymmetric rods joined to each other (Guo, Liu, An, Li, & Hu, 2005). The interaction of zein with LPI or SPI could also reduce the hydrophobic interactions among asymmetric zein rods. Thus, the porous nature of films has turned to denser one. These results suggested that the zein based composites could be an opportunity to reduce the classical brittleness problem of zein films.

During food applications both control zein film and composite films showed good affinity on food surfaces and maintained their integrity if they are not disturbed (see supplementary file 1). However, at the later stages of cold-storage (e.g. at 96th h for beef) a gentle attempt to unpeel zein control films from food surfaces caused local disintegration in acidified and nonacidified films (see supplementary file 2). Superficial local cracks also appeared in zein-LPI and zein-SPI films, but these films (acidified or non-acidified) peeled off from food surfaces without disintegration. These results suggested a certain improvement in mechanical properties of composite films. However, more detailed studies are needed to determine long-term effects of SPI and LPI on mechanical and morphological properties of zein films during processing and storage.

4. Conclusions

The results of this work clearly showed the possibility of designing antimicrobial composite films with active and consumer-controlled activate-at-home-type antimicrobial properties. The release tests conducted with zein-SPI and zein-LPI films in different buffers and in different foods clearly showed the presence of free and bound LYS fractions in the composite films. The timing of release for LYS bound could be controlled in lamb meat and beef simply by acidification of composite films placed on food surface. However, the designed pH-triggered release mechanism did not work successfully in smoked salmon due to the liberation of LYS bound from non-acidified films in presence of salt used in curing. Further studies are needed to optimize film compositions (LYS, SPI, and LPI contents) and fix the amount of free and bound enzyme fractions in the films. Moreover, an industrial design of a special packaging is also needed to develop practical commercial applications for the developed delivery method. For example, integration of a simple acidification mechanism into packaging could enable activation of films in closed packages. This provides an additional antimicrobial effect to VP or MAP during transportation from market to home and enables extra safety for closed or opened packed food kept at home until consumption. The development of ActiHome packaging concept might expand the horizons of standard antimicrobial packaging by involving conscious consumers in the active packaging process. This helps development of value-added personalized food products that help risk management for susceptible people such as pregnant women, elderly and immunosuppressed people.

<table>
<thead>
<tr>
<th>Incorporated concentration (mg/cm²)</th>
<th>Thickness (µm)</th>
<th>Tensile Strength (MPa)</th>
<th>Elongation (%)</th>
<th>Young’s modulus (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LYS</td>
<td>135.72 ± 6.2</td>
<td>6.40 ± 0.9a</td>
<td>1.36 ± 0.1f</td>
<td>470.29 ± 38.6a</td>
</tr>
<tr>
<td>LPI</td>
<td>126.09 ± 3.3</td>
<td>5.62 ± 0.3b</td>
<td>1.52 ± 0.1c</td>
<td>370.14 ± 30.5b</td>
</tr>
<tr>
<td>SPI</td>
<td>119.94 ± 2.9</td>
<td>5.34 ± 0.5b</td>
<td>3.08 ± 0.5a</td>
<td>138.75 ± 23.1c</td>
</tr>
</tbody>
</table>

**Note:** different letter at each column indicate statistically significant changes at p < 0.05.

Table 2: Mechanical properties of zein, zein-LPI and zein-SPI films.
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Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.foodhyd.2017.12.038.

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