Development of activate-at-home-type edible antimicrobial films: An example pH-triggering mechanism formed for smoked salmon slices using lysozyme in whey protein films

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**A B S T R A C T**

The aim of the present study was to develop the first generation of a novel natural antimicrobial packaging system which could be activated by consumers at home. For this purpose, antimicrobial films having a pH-triggering mechanism which could be activated by acidification were developed by exploiting isoelectric properties and pH induced charge–charge interactions between whey proteins (WP) and lysozyme (LYS). The release tests of films in buffers and on coated smoked salmon slices at 4°C showed the immobilization of positively charged LYS by the negatively charged WP films at pHs ≥ 5.5. The LYS release initiated when pH was reduced below 5.5 where WP started to gain positive charges. The amount of LYS released and LYS release rate increased as pH reduced gradually from 5.5 to 3.0. The preparation of composite films of WP with beeswax (BW) and oleic acid (OLE) increased the total released LYS from films, WP, WP-OLE and WP-BW films showed antimicrobial activity against *Listeria innocua* in laboratory media. The smoked salmon slices coated by activated WP containing WP-OLE films showed significantly lower (almost 0.6 decimal) *L. innocua* counts than controls after 1 week at 4°C. This study clearly showed that the antimicrobial films could be activated by consumers to improve safety of opened packed food stored in home type refrigerators.

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1. Introduction

The rising threat of foodborne outbreaks related to raw and minimally processed products has caused a great interest in functional antimicrobial packaging materials (Cagri, Ustunol, & Ryser, 2001; Lu, Zhu, Li, & Chen, 2015; Lynch, Tauxe, & Hedberg, 2009). Antimicrobial packaging is a promising area since it might inhibit or suppress the growth of pathogenic or spoilage microorganisms on food surface by using minimum amounts of antimicrobials (Appendini & Hotchkiss, 2002; Gennadios, Hanna, & Kurth, 1997; Han, 2000; Ouattara, Simard, Piotte, Bégin, & Holley, 2000; Quintavalla & Vicini, 2002). The incorporation of chemical compounds into plastic films is a simple and effective method to develop antimicrobial packaging. However, the growing health concerns of the consumers against chemicals, and environmental problems originating from plastics cause a great interest in using natural antimicrobial compounds and biopolymers in antimicrobial packaging (Cha & Chinnan, 2004; Han, 2003; Pérez-Pérez, Regaldo-González, Rodríguez-Rodríguez, Barbosa-Rodríguez, & Villaseñor-Ortega, 2006; Suppakul, Miltz, Sonneveld, & Bigger, 2003).

Different natural antimicrobials including antimicrobial enzymes, bacteriocins, essential oils and phenolic compounds can be employed in active packaging (Cha & Chinnan, 2004; Cooksey, 2005; Joerger, 2007; Mastromatteo, Mastromatteo, Conte, & Del Nobile, 2010). Lysozyme (LYS) obtained from hen egg white is one of the most potential candidates for antimicrobial packaging since it has a Generally Recognized as Safe (GRAS) status. Also, it shows good stability and activity in different films and food systems under refrigerated storage temperatures (Mecitoglu et al., 2006; Unalan, Korel, & Yenemiçioğlu, 2011). LYS has been tested extensively in different edible materials including zein, gelatin, soy protein, carrageenan, whey protein, chitosan, alginate and pullulan (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-Carballe, Gavara, & Hernández-Muñoz, 2010).

The LYS shows antimicrobial activity mainly on Gram-positive...
bacteria by splitting the bonds between N-acetylmuramic acid and N-acetylglucosamine of the peptidoglycan in their cell walls (Appendini & Hotchkiss, 1997; Shah, 2000). Thus, the application of LYS in active packaging targets mainly the inhibition of critical Gram-positive pathogenic bacteria such as Listeria monocytogenes (Duan, Park, Daeschel, & Zhao, 2007). The prevention of food contamination by L. monocytogenes, and application of hurdles to prevent listerial growth in risky food are critically important since this bacterium may cause deadly infections (Alvarez-Ordóñez, Leong, Hickey, Beaufort, & Jordan, 2015). It is strongly suggested that individuals highly susceptible to Listeria, such as pregnant women, old people and the immunosuppressed people should be particularly careful during the consumption of risky food (Vazquez-Boland et al., 2001). The smoked salmon is a typical L. monocytogenes risk food (Daigaard & Jørgensen, 1998; Di Cicco et al., 2012; Vermeulen, Devlieghere, De Loy-Hendrickx, & Uyttendaele, 2011). However, the struggle against listeriosis in smoked salmon is really challenging since this bacteria could be isolated from different zones of smoked-fish processing plants, and it can grow even at the refrigeration temperatures with or without the presence of oxygen (Dass, Abu-Ghamam, Antony-Babu, & Cummins, 2010). Thus, vacuum packaging (VP) is not considered as an effective way to prevent this critical pathogen (Duffes, 1999; Gram, 2001). On the other hand, recent studies have shown that it is possible to control L. monocytogenes growth in cold stored smoked salmon slices by MAP containing elevated levels of CO2 (70% CO2/30% N2) (Nilsson, Huss, & Gram, 1997). However, the protective effect of MAP is lost after opening of the package at home, and the remaining food kept for later consumption becomes quite risky for the development of L. monocytogenes (Buchanan & Klawitter, 1990; Davies, 1997; Tsigarida, Skandamis, & Nychas, 2000).

The antimicrobial packaging could be used as a highly effective hurdle to prevent the growth of L. monocytogenes not only in smoked salmon but also in other risky foods. For example, Min, Rumsey, and Krochta (2008) successfully employed LYS incorporated whey protein (WP) films to control L. monocytogenes growth on cold stored smoked salmon. In that study, Min et al. (2008) adjusted the pH of WP film solution during film making to eliminate charge-charge interactions between LYS and WP films, and to ensure the initiation of free enzyme’s release from films following food application. On the other hand, Unalan, Arcan, Korel, and Yemencioğlu (2013) employed LYS containing zein composite films having controlled release properties to prevent the growth of L. monocytogenes in fresh cheese.

The aim of the present work is to develop the first generation of an antimicrobial edible film which can be activated by the consumers. Such a packaging concept which could be named as activate-at-home-type packaging (ActiHome packaging) is novel, and it could be applied by the consumer as a hurdle step for the preservation of VP or MAP food after opening of the packaging and ending of the preservation effect of packaging atmosphere. The ActiHome packaging could reduce poisoning from opened VP or MAP food left for later consumption. It is proposed that the precast ActiHome packaging films containing immobilized antimicrobials might be placed onto top and/or bottom surface and between layers of food or they could be applied as a coating on food surface by the producers. It is desirable that the immobilized antimicrobials at the film surface or within the film provide some antimicrobial effect at the food surface during transportation, storage and marketing. However, it is more important for an ActiHome type packaging to maintain a significant portion of the incorporated antimicrobial during transportation, storage and marketing within the film, but to start to release this antimicrobial after activation of film by the consumer at home. The WP films could be successfully employed as ActiHome type films due to their suitable pI between 4.4 and 5.4 (Etzel, 2004) and their importance as an agro-industrial waste. The use of LYS in WP film system has a good potential since pl of WPs enables binding of this antimicrobial enzyme (positively charged in food systems due to its high pl at 11.4) by the negatively charged WP film matrix when it was employed for packaging of risky food at pH close to neutrality. The WP-lys film system forms a perfect pH-triggering mechanism since it starts to release positively charged LYS by acidification that causes drop of pH below 5.0. Thus, in the present study, antimicrobial films having a pH-triggering mechanism which could be activated by acidification of consumer at home were developed by exploiting isoelectric properties and pH induced charge-charge interactions between whey protein (WP) and LYS. The release properties of developed WP films were also modified and improved by the preparation of composites of WP with oleic acid (ODE) and beeswax (BW), respectively. The designed films were tested for their release properties and antimicrobial effects in buffer systems and in a real packaging application conducted with smoked salmon. The present work is the first study planning the participation of consumer in active packaging and reducing incidence of deadly infections by critical pathogens in opened VP and MAP food stored at home for later consumption.

2. Materials and methods

2.1. Materials

Commercial LYS, Micrococcus lysodeicticus, oleic acid (90%) and beeswax were obtained from Sigma Chem. Co. (St. Louis, MO, USA). Whey protein isolate (BIPRO®, 97.8% protein) was kindly donated by Davisco Foods International, Inc. (MN, USA). Glycerol was obtained from Merck (Darmstadt, Germany). Bacterial strain of Listeria innocua (NRRL B-33314) used in antimicrobial zone tests was provided by United States Department of Agriculture, Microbial Genomics and Bioprocessing Research Unit, Peoria, Illinois. Cold-smoked salmon used in food release tests was obtained from a supermarket in Izmir (Turkey).

2.2. Film making

WP films were prepared by the slight modification of the method presented in McHugh and Krochta (1994). Briefly, 10% (w/w) of WP solution was prepared in distilled water, using magnetic stirrer. Glycerol (at 60% of WP) was added as plasticizer, and the solution was stirrer for 5 min. The film making solution was then incubated in a water bath at 90 °C for 30 min for protein denaturation, and cooled in ice water bath for 5 min. After that, LYS (7.87 mg/g film solution) was added into medium, and using a homogenizer—disperser (Heidolph, Germany, rotor Ψ = 6.6 mm tip) the mixture was homogenized at 10,000 rpm for 4 min. The five gram portions of film forming solution was then cast onto plastic Petri dishes (8.5 cm inner diameter), and dried for 24 h in a controlled test cabinet at 50% relative humidity and at 25 °C. This procedure was used to obtain standard WP films. The same procedure was also followed to obtain composite films of WP with oleic acid (WP-OLE). The oleic acid (at 5% of WP) was added after 5 min cooling of the heated film making solution. The LYS was added finally after weighting of film solution. The homogenization, casting and drying steps of WP-OLE films were applied as described above for the standard WP film. However, the sequence of the steps of the method was slightly modified to obtain composites of WP with beeswax (WP-BW). The beeswax (at 30% of WP) was added in hot film making solutions immediately after heating at 90 °C for 30 min. The LYS was added finally after weighting of film solution. The homogenization at 10,000 rpm for 4 min was applied to hot film making solution to enable melting and dispersion of the
beeswax. The cooling, casting and drying steps for WP-BW films were applied as described above for the standard WP film.

2.3. LYS activity

LYS activity was measured spectrophotometrically at 660 nm by using Shimadzu (Model 2450, Japan) spectrophotometer equipped with a constant temperature cell holder set to 30 °C. Reaction mixture was prepared by mixing 0.1 ml enzyme containing solution (incubated at 30 °C for 1 min), and 2.4 ml M. lysodeicticus suspension (at 30 °C) prepared in pH 7.0 Na-phosphate buffer. After mixing, the decrease in absorbance was monitored for 2 min, and enzyme activity was calculated from the slope of the initial portion of absorbance vs. time curve. Average of three measurements was used in calculations, and enzyme activity was expressed as Units per cm² of films. One Unit was defined as 0.001 changes in absorbance within 1 min.

2.4. Release profiles of films placed in different buffer systems

The LYS release tests were conducted by incubation of WP, WP-OLE and WP-BW films at 4 °C in different buffers between pH 3.0 and 6.0. The buffers at pH 4.5, 5.0, 5.5 and 6.0 were obtained by using 0.05 M Na-phosphate while, the buffers at pH 3.0 and 4.0 were obtained by 0.05 M Na-acetate-acetic acid buffer. Briefly, films (1/4 of each film, area: 14.2 cm²) were placed into glass Petri dishes containing 50 mL of buffer solution. The dishes were kept at 4 °C in an incubator and shaken with an orbital shaker working at 80 rpm. The release tests of each film were conducted until equilibrium was reached for the release of LYS or an insignificant increase was observed in LYS release. The LYS activity was monitored by taking 0.3 ml (3 × 0.1 ml) aliquots from the release test solution at different time intervals. The enzyme activities in collected aliquots were determined spectrophotometrically as presented in Section 2.3. All calculations were corrected by considering the activity removed by collected aliquots during sampling. The total LYS activity released from each film corresponded to maximum units released per cm² of the films (U/cm²) at the equilibrium. All activity measurements were conducted for three times. The release curves were formed by plotting the calculated released activities (U/cm²) vs. time (days). The initial release rates of LYS were determined from the slope of the initial linear portion of release curve. The release rates were expressed as U/cm²/h.

The protein release from different films was determined by designing separate release tests at the same conditions described for LYS above. However, the sampling (3 × 0.1 ml) was done only at the end of release test at 72 h. The protein contents of the samples were determined with the Bradford method by mixing 0.1 ml portions of samples with 1 ml of Bradford reagent. After 60 min of incubation, the absorbance of samples was measured at 595 nm. Bovine serum albumin was used as standard and protein solubilized from films were expressed as mg/g film dry weight.

2.5. Release profiles of films placed on smoked salmon slices

For these release tests, 13 mm diameter discs of WP, WP-OLE and WP-BW films were placed on one surface of 13 mm diameter discs obtained from smoked salmon slices (average thickness: 4.02 ± 0.4 mm; average weight: 0.62 ± 0.05 g). The film and salmon discs were prepared at aseptic conditions, using a sterile cork-borer. The coated salmon discs were first wrapped with a stretch food contact. The packed coated salmon discs were then stored at 4 °C for 24 h. At the end of 24 h cold storage, the packs were opened, and 0.2 ml 5% (w/v) citric acid solution prepared in clarified lemon juice was pipetted onto surfaces of film discs to trigger release of LYS. The samples were then packed again with the stretch film and aluminium foil, and further cold stored for 96 h. The LYS activity released from films to salmons was monitored during cold storage. For determination of the activity salmon discs were homogenized with 10 mL of deionized water for 2 min, using a homogenizer (IKA, DI18 Basic, Germany). The homogenates were centrifuged at 11,000 rpm for 20 min, and LYS activity was determined in the clear supernatants for three times (3 × 0.1 ml) as described in Section 2.3.

2.6. Antimicrobial activity of films in laboratory media

The antimicrobial activities of WP, WP-OLE and WP-BW films were determined by the classical zone- inhibition assay, using L. innocua (NRRL B-33314) as test microorganism. The inoculums were prepared in pepton 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 × 10⁶ CFU/mL). Fifteen discs were cut from each film by a cork-borer under aseptic conditions and three discs were placed onto surfaces of each inoculated agar. Petri dishes were then incubated at 37 °C for 48 h, and the area of clear zones formed around the discs was calculated by using a digital micrometer.

2.7. Antimicrobial activity of films on coated smoked salmon slices

The L. innocua was activated by transferring one loop of frozen culture (−80 °C) to 9 mL brain heart infusion broth (BHIB) (Merck, Darmstadt, Germany) and incubating at 37 °C for 24 h. One-ml aliquot from the active culture was transferred to the BHIB, and incubated at 4 °C for an additional 24 h for adaptation of its growth at cold storage temperature. After dilution with 0.1% pepton water (Merck, Darmstadt, Germany) an inoculum of 1 × 10⁷ CFU/mL was obtained for the inoculation of food sample. The surfaces of the smoked salmon slices (Size: 4 × 4 cm, Weight: 10 g) were inoculated with 0.125 mL of inoculum, and the inoculum was spread with a sterile plastic spreader. The inoculated slices were kept in a sterile cabinet for 10 min for absorption of the inoculum, and then antimicrobial films at the same size were placed carefully on their surfaces. The films were then acidified for triggering the release of LYS by pipetting 0.5 mL 5% (w/v) citric acid solution prepared in clarified lemon juice onto their surface and spreading this solution with a sterile plastic spreader. Controls were prepared similarly without acidification. Each coated smoked salmon slice was then wrapped with a plastic film and then with an aluminium foil. The five different groups used in active packaging were as follows; (1) Uncoated samples, (2) Samples coated with WP-OLE films, (3) Samples coated with WP-OLE films containing LYS (0.7 mg/cm²), (4) Samples coated with WP-OLE films, and acidified, (5) Samples coated with WP-OLE films containing LYS (0.7 mg/cm²) and acidified. The samples were stored at 4 °C, and enumerated for their L. innocua counts periodically, at 0th, 1st, 3rd, 5th and 7th days of cold-storage. During bacterial counts each salmon slice (10 g) was placed into a stomacher bag containing 90 mL sterile 0.1% peptone water, and homogenized for 60 s by using a stomacher (BagMixer® 400, Interscience, France). The serial decimal dilutions were prepared from this homogenate, and appropriate dilutions (0.1 mL) were spread plated onto Oxford Listeria Selective Agar (Merck, Darmstadt, Germany) containing Oxford Listeria Selective Supplement (Merck, Darmstadt, Germany). The plates were incubated at 37 °C for 48 h, and small black colonies with halos on the plates were enumerated. The counts were performed in triplicate plates. The three separate samples for each of the five different groups were analysed at each sampling day.
2.8. Mechanical properties of films

Tensile strength at break, elongation at break and elastic modulus were determined, using a Texture Analyser TA-XT2 (Stable Microsystems, Godalming, UK) according to ASTM Standard Method D-882-02 (ASTM, 2002). 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.

2.9. Moisture contents of films

The moisture contents of films were determined by the vacuum oven drying method applied at 70 °C 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.

3. Results and discussion

3.1. pH controlled LYS release profiles of different films in buffer systems

3.1.1. Release profiles of WP films

The LYS release profiles of WP films in buffers with different pHs at 4 °C were presented in Fig. 1. The pI of major WP fractions changed between 4.4 and 5.4 (Etzel, 2004), while pI of LYS is at 11.4 (Product information sheet, Sigma–Aldrich, USA). Thus, effective binding of positively charged LYS by the negatively charged WP occurred, and the films did not release any considerable enzyme activity at pH 5.5 and 6.0. The films incubated at pH 5.0, which is close to the pI of protein fractions forming WP, released LYS slowly. However, the total amount of enzyme activity released from these films at the equilibrium reached within 7 days incubation was low and corresponded only 22% of LYS activity incorporated into films (14,776 U/cm²) (Table 1). In contrast, the films incubated at pH 4.5, 4.0 and 3.0 released almost 2, 3 and 5 fold higher total LYS activity at their equilibrium than films incubated at pH 5.0, respectively. It is important to report that the activity of LYS released from WP films at pH 3.0 was almost equal to the LYS activity incorporated into films (activity released corresponded to 102% of activity incorporated). These data clearly showed the roles of attractive and repulsive charge–charge interactions formed among ionisable groups of LYS and WP in binding and release of LYS. It is clear that the pH drop caused loss of attractive charge–charge interactions between positively charged LYS and WP film matrix which started to gain positive charges while its negative charges disappeared. Moreover, the LYS release rates of films increased as pH of the release medium was reduced. In fact, it is worth to note that the LYS release rate of films incubated at pH 3.0 is 4.2 and 7.7 fold higher than those incubated at pH 4.5 and 4.0, respectively. It appeared

3.10. Statistical analysis

Statistical analysis was performed by using MINITAB® release 17 (Minitab Inc., State College, Pa., U.S.A.). LYS activities, antimicrobial inhibition zones, microbial counts and mechanical properties were analysed by using one-way analysis of variance (ANOVA). Significance threshold was p < 0.05.

Table 1

<table>
<thead>
<tr>
<th>Film composition</th>
<th>LYS (mg/cm²)</th>
<th>OLE (%)</th>
<th>pH</th>
<th>Initial lysozyme release rate (U/cm²/h)</th>
<th>Total released lysozyme activity (U/cm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.7</td>
<td></td>
<td></td>
<td>3.0</td>
<td>62,676 (0–1)</td>
<td>69,138 ± 9663*(102%)*</td>
</tr>
<tr>
<td>0.7</td>
<td></td>
<td></td>
<td>4.0</td>
<td>8066.8 (0–3)</td>
<td>47,019 ± 7552*(70%)*</td>
</tr>
<tr>
<td>0.7</td>
<td></td>
<td></td>
<td>4.5</td>
<td>1496.5 (0–3)</td>
<td>35,021 ± 853*(52%)*</td>
</tr>
<tr>
<td>0.7</td>
<td></td>
<td></td>
<td>5.0</td>
<td>1513 (0–3)</td>
<td>14,776 ± 770*(22%)*</td>
</tr>
<tr>
<td>0.7</td>
<td></td>
<td></td>
<td>5.5</td>
<td>No considerable release</td>
<td></td>
</tr>
<tr>
<td>0.7</td>
<td></td>
<td></td>
<td>6.0</td>
<td>No considerable release</td>
<td></td>
</tr>
<tr>
<td>0.7</td>
<td>9</td>
<td></td>
<td>4.0</td>
<td>9875.3 (0–3)</td>
<td>51,391 ± 1983*(76%)*</td>
</tr>
<tr>
<td>0.7</td>
<td>9</td>
<td></td>
<td>4.5</td>
<td>3221.9 (0–3)</td>
<td>74,184 ± 6730*(110%)*</td>
</tr>
<tr>
<td>0.7</td>
<td>9</td>
<td></td>
<td>5.0</td>
<td>1918 (0–3)</td>
<td>23,817 ± 647f(35%)*</td>
</tr>
<tr>
<td>0.7</td>
<td></td>
<td></td>
<td>5.5</td>
<td>No considerable release</td>
<td></td>
</tr>
<tr>
<td>0.7</td>
<td>9</td>
<td></td>
<td>30</td>
<td>11,288 (0–3)</td>
<td>56,811 ± 954*(84%)*</td>
</tr>
<tr>
<td>0.7</td>
<td>9</td>
<td></td>
<td>4.5</td>
<td>3004.5 (0–3)</td>
<td>40,709 ± 3592*(60%)*</td>
</tr>
<tr>
<td>0.7</td>
<td>9</td>
<td></td>
<td>5.0</td>
<td>1705 (0–3)</td>
<td>20,821 ± 1215*(31%)*</td>
</tr>
<tr>
<td>0.7</td>
<td></td>
<td></td>
<td>5.5</td>
<td>No considerable release</td>
<td></td>
</tr>
</tbody>
</table>

*a–* Different letters in each column show significant differences at p < 0.05.

*b* Concentrations of beeswax and oleic acid as % of WP (w/w).

*c* Time periods (h) of data used in best fit. R² value varies between 0.73 and 1.

*d* Time (days) at which the equilibrium was reached for LYS release.

*e* Proportion of released LYS activity incorporated in the film.
that the acceleration of LYS release by pH drop is due to the increased repulsion between the positively charged enzyme molecules, and WP which net positive charges increased as pH was reduced below its pI. These results clearly showed the possibility of controlling release rate and total amount of LYS released from WP films by changing the pH of release medium. This finding forms the basis of designing ActiHome type antimicrobial edible films which can be activated by the consumer simply by acidification of film placed into the package (on food surface, on bottom of food or between food layers) by the producer.

3.1.2. Release profiles of WP-BW and WP-OLE films

The incorporation of different fatty acids and waxes into films, and preparation of composite structures is frequently employed to modify morphologies, release profiles, permeability characteristics and mechanical properties of edible films (Arcan & Yemencioglu, 2013; Fabra, Talens, & Chiralt, 2008). In this work, the composites of WP with OLE and BW were prepared to create chemical and physical modifications in WP film matrix and change its LYS release profiles. The incorporation of OLE into WP films caused the formation of a protein-fatty acid emulsion and this turned transparent WP films to slightly opaque (Fig. 2A and B). On the other hand, the incorporation of BW caused the formation of numerous microscopic wax particles within the films and this tuned transparent WP films to highly opaque (cloudy) (Fig. 2C).

The release profiles of WP films and their composites between pH 4.0 and 5.5 are seen in Fig. 3A–D. Similar to the results obtained for WP films, the charge–charge attractions between positively charged LYS and negatively charged WP film matrix at pH 5.5 prevented the release of enzyme from WP-OLE and WP-BW composite films (Fig. 3A). However, the reduction of pH to 5.0 initiated release of the LYS from composite films. The release curves
obtained at pH 5.0 and 4.5 (Fig. 3B and C) clearly showed the higher LYS release rates and total amounts of released LYS from composite films at the equilibrium than WP films. On the other hand, similar LYS release rates and total released LYS activities were observed for all films at pH 4.0 (Fig. 3D). This finding suggested the dominance of repulsive charge–charge forces in LYS release from all films at pH 4.0. It is important to note that the total amount of LYS released from WP-OLE films at pH 4.5 was 1.8 and 2.1 fold higher than those of WP-BW and WP films, respectively. It is interesting to report that the released activity from WP-OLE films at pH 4.5 corresponded to 110% of calculated incorporated LYS activity in these films. The 10% higher activity of released LYS activity than that of calculated incorporated LYS activity could be simply due to slight heterogeneities in film thickness or in enzyme distribution within the films (quarter of round cast films were cut and used in release tests). Thus, this result practically means that almost all LYS released from WP-OLE films at pH 4.5. The results of release tests for WP films showed that these films should be incubated at pH 3.0 to release all their LYS. Thus, it is clear that the use of WP-OLE composites in food application at pH 4.5 instead of WP films could be more beneficial to increase LYS release on food surface.

The protein release profiles of WP, WP-OLE, and WP-BW control (films lacking LYS) films at the end of 72 h incubation at 4 °C in different buffers were also presented in Fig. 4. The films showed quite similar protein solubility between pH 4.0 and 6.5. The protein solubility of films was at insignificant levels (<1% of film dry weight) at pH 4.0, 4.5 and 5.0 (a range that is critical for release of LYS from films). A slight increase was observed in protein solubility above pH 5.0, but the maximum solubility was still below 3.7% of dry film weight even at pH 6.5. The films used in the current study were obtained after effective heat denaturation of whey protein (see Section 2.2) at 90 °C for 30 min. Thus, they contain very little amount of soluble protein at the pH range (pH 4.5 and 5.0) designed for activation of films by pH triggering. These results showed that the higher LYS release from WP-OLE and WP-BW films than WP films was not due to increased protein erosion from matrices of composite films. The WP that forms the film matrix is directly responsible for charge–charge binding (due to attraction) and release (due to repulsion) of LYS from the films when pH was changed. Due to their composite structures, the WP-OLE (9% of WP was changed with OLE) and WP-BW (30% of WP was changed with BW) films contain lower amounts of WP than the WP films. Thus, the reduced WP content of composite films and declined number of charge–charge interactions between LYS and composite film matrix might be a factor effective on increased LYS release from WP-OLE and WP-BW films. This factor should be more effective on release profiles of WP-BW films than those of WP-OLE films, since 30% of WP was changed with BW in these composite films. On the other hand, in WP-OLE films only 9% of WP was changed with OLE. Thus, there should be some other factors affecting the higher amounts of LYS release from WP-OLE films than WP-BW films at pH = 4.5. It appeared that the emulsion formed between WP and OLE might have caused a shift in the pKa of WP by masking some of the charged groups of this protein. The release of highest amount of LYS from WP-BW films occurred at pH 4.5. Thus, it is possible that the effects of emulsion induced changes (like masking) in charges of WP-OLE film matrix were maximized at pH 4.5, and this triggered

Table 2

Antimicrobial activity of WP, WP-OLE and WP-BW films on L. innocua in laboratory media with and without acidification.

<table>
<thead>
<tr>
<th>Film composition</th>
<th>LYS (mg/cm²)</th>
<th>OLE (%)</th>
<th>BW (%)</th>
<th>Acification</th>
<th>Average area of clear zones (mm²)</th>
<th>Acification</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>24 h</td>
<td>48 h</td>
</tr>
<tr>
<td>----</td>
<td>----</td>
<td>----</td>
<td>----</td>
<td>----</td>
<td>No clear zone</td>
<td>No clear zone</td>
</tr>
<tr>
<td>0.7</td>
<td>----</td>
<td>----</td>
<td>----</td>
<td>+</td>
<td>51.7 ± 8.5d</td>
<td>63.5 ± 13.9</td>
</tr>
<tr>
<td>0.7</td>
<td>--</td>
<td>9</td>
<td>----</td>
<td>+</td>
<td>101.1 ± 15.3c</td>
<td>84.6 ± 22.5</td>
</tr>
<tr>
<td>0.7</td>
<td>9</td>
<td>----</td>
<td>----</td>
<td>+</td>
<td>121.5 ± 14.9b</td>
<td>93.9 ± 17.4</td>
</tr>
<tr>
<td>0.7</td>
<td>9</td>
<td>----</td>
<td>30</td>
<td>+</td>
<td>141.6 ± 26.4d</td>
<td>124.3 ± 32.7</td>
</tr>
<tr>
<td>0.7</td>
<td>--</td>
<td>30</td>
<td>----</td>
<td>+</td>
<td>54.5 ± 7.4d</td>
<td>55.6 ± 11.0</td>
</tr>
<tr>
<td>0.7</td>
<td>--</td>
<td>30</td>
<td>----</td>
<td>+</td>
<td>58.5 ± 21.2d</td>
<td>88.11 ± 55.0</td>
</tr>
</tbody>
</table>

* Different letters within each incubation time show significant differences at p < 0.05.
* Concentrations of beeswax and oleic acid as % of WP (w/w).
* 5% (w/v) citric acid solution in lemon juice.
extensive LYS release from films due to minimized attractive charge–charge interactions between LYS and film matrix.

3.2. pH controlled LYS release profiles of films in coated cold stored smoked salmon slices

The designed pH-controlled release system was tested on WP, WP-BW and WP-OLE films coated smoked salmon slices cold stored at 4 °C. The film surfaces of half of the packed samples were acidified to test working of the pH-controlled triggering mechanism designed to initiate LYS release to salmon slices while the other half of the packed samples were cold stored without acidification. Fig. 5 shows the lack of any considerable LYS release from the films in non-acidified salmon controls that average pH value is almost 6.3. This result confirms the stability of attractive charge–charge interactions between LYS and WP film matrix at the initial pH of target food. In contrast, the release of LYS from films to food surface initiated in all samples following acidification that dropped pH of salmon slices from 6.3 to 6.4 to 4.5–4.6. There are no statistically significant differences among LYS activity of salmon slices packed with acidified WP, WP-OLE and WP-BW films up to 48 h of cold storage (p > 0.05). However, LYS activities released on salmon slices packed with acidified WP-OLE and WP-BW composite films were 37% and 33% higher than those released on salmon slices packed with standard WP films (p < 0.05) at the end of 72 h cold storage, respectively. Different from the results obtained in buffer systems, the difference between total released LYS activities of acidified WP-OLE and WP-BW coated salmon slices were not statistically significant (p > 0.05). It should also be reported that LYS activities released on salmon slices from cm² of WP, WP-OLE and WP-BW films within 72 h at pH 4.5 were 24%, 50% and 13% lower than those released from these films in buffer at pH 4.5 at equilibrium, respectively. It is possible that the reduced LYS activity release onto salmon slices was due to more limited swelling of films on salmon surfaces than in the buffers. However, quantitative LYS determination is needed to prove this hypothesis due to some potential changes in LYS activity by active proteases in salmon and by complex formation with salmon constituents. On the other hand, these results clearly showed the working of the designed pH-controlled release system in cold stored smoked salmon. Moreover, the beneficial effect of using composite WP-OLE and WP-BW films in increasing LYS delivery was also clearly demonstrated.

3.3. Antimicrobial activity of films

3.3.1. Antimicrobial activity of films in classical zone-inhibition test

Antimicrobial tests in this part targeted determining the effects of composite film making on antilisterial activity of films. Table 2 shows the antimicrobial properties of WP films and its composites following 24 h and 48 h incubation periods at 37 °C. During antimicrobial tests the control films (acidified or non-acidified) did not form any clear zone areas, but the formation of turbid zones very close to film disc borders (almost 1 mm) was observed around control WP films. This result suggested a limited inherent antimicrobial activity of films against L. innocua, possibly due to the presence of some residual natural antimicrobial milk components like lactoferrin or antimicrobial peptides in the commercial WP preparation. The limited antilisterial effect of films obtained from the same commercial WP product was also reported by Unalan, Ucar, Arcan, Kolel & Yemencioglu, (2011). However, it should also be reported that some extensive bacterial growth was also observed at the bottom of control films (Fig. 6). On the other hand, both acidified and non-acidified WP, WP-OLE and WP-BW films containing LYS caused formation of clear zones on L. innocua. The acidification of films caused significant increases in average area of clear zones formed by WP and WP-OLE films containing LYS (p < 0.05). In contrast, the acidification of films and the activation of LYS release did not cause any significant increases in the antimicrobial activity of LYS containing WP-BW films. It is also worth to report that the non-acidified LYS containing WP and WP-BW films showed similar antimicrobial activity while acidified LYS containing WP-BW films showed lower antimicrobial activity than acidified LYS containing WP films. These results suggested the change of LYS release profiles of WP-BW films at 37 °C within 24 h. It appeared that the modifications in the wax component, such as softening and melting at 37 °C which is essential to apply for obtaining bacterial growth during incubation caused the migration of wax components within the film matrix to film surface, and this formed a hydrophobic barrier for diffusion of LYS to agar. On the other hand, for both acidified and non-acidified films the highest antimicrobial activity was observed for WP-OLE films. This finding compared well with the release test experiments conducted in buffers and showed the release of highest amounts of LYS from WP-OLE films.

3.3.2. Antimicrobial activity of films on cold stored coated smoked salmon slices

The WP-OLE film, which showed superior performance than the

![Image](image-url)

**Table 3**

Antimicrobial activity of WP and WP-OLE films on L. innocua inoculated smoked salmon slices cold stored at 4 °C.

<table>
<thead>
<tr>
<th>L. innocua counts during storage at 4 °C (log CFU/g)</th>
<th>Day 0</th>
<th>Day 1</th>
<th>Day 3</th>
<th>Day 5</th>
<th>Day 7</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uncoated</td>
<td>4.98± 0.08&lt;sup&gt;A&lt;/sup&gt;</td>
<td>4.82± 0.10&lt;sup&gt;B&lt;/sup&gt;</td>
<td>4.82± 0.10&lt;sup&gt;B&lt;/sup&gt;</td>
<td>4.80± 0.11&lt;sup&gt;B&lt;/sup&gt;</td>
<td>4.83± 0.13&lt;sup&gt;B&lt;/sup&gt;</td>
</tr>
<tr>
<td>WP-OLE (control)</td>
<td>4.92± 0.10&lt;sup&gt;A&lt;/sup&gt;</td>
<td>4.72± 0.10&lt;sup&gt;C&lt;/sup&gt;</td>
<td>4.73± 0.08&lt;sup&gt;B&lt;/sup&gt;</td>
<td>4.78± 0.09&lt;sup&gt;B&lt;/sup&gt;</td>
<td>4.84± 0.11&lt;sup&gt;B&lt;/sup&gt;</td>
</tr>
<tr>
<td>WP-OLE + LYS</td>
<td>4.92± 0.08&lt;sup&gt;A&lt;/sup&gt;</td>
<td>4.68± 0.07&lt;sup&gt;B&lt;/sup&gt;</td>
<td>4.69± 0.15&lt;sup&gt;B&lt;/sup&gt;</td>
<td>4.74± 0.12&lt;sup&gt;B&lt;/sup&gt;</td>
<td>4.49± 0.12&lt;sup&gt;C&lt;/sup&gt;</td>
</tr>
<tr>
<td>WP-OLE (acidified)</td>
<td>4.97± 0.04&lt;sup&gt;A&lt;/sup&gt;</td>
<td>4.84± 0.11&lt;sup&gt;B&lt;/sup&gt;</td>
<td>4.77± 0.13&lt;sup&gt;B&lt;/sup&gt;</td>
<td>4.80± 0.12&lt;sup&gt;B&lt;/sup&gt;</td>
<td>4.57± 0.09&lt;sup&gt;B&lt;/sup&gt;</td>
</tr>
<tr>
<td>WP-OLE + LYS (acidified)</td>
<td>4.97± 0.07&lt;sup&gt;A&lt;/sup&gt;</td>
<td>4.49± 0.13&lt;sup&gt;B&lt;/sup&gt;</td>
<td>4.45± 0.13&lt;sup&gt;B&lt;/sup&gt;</td>
<td>4.27± 0.10&lt;sup&gt;B&lt;/sup&gt;</td>
<td>4.24± 0.15&lt;sup&gt;C&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>*<sup>A</sup>-<sup>B</sup> Different letters within each storage time show significant differences at p < 0.05.</sup>
WP-BW film in release tests and antimicrobial tests was selected for the food application. The acidification conducted with 5% (w/v) citric acid solution prepared in clarified lemon juice brought the salmon slice pH from 6.3–6.4 to 4.5–4.6, a critical pH level which caused maximum LYS release from WP-OLE composites after pH triggering. The degree of acidification in salmon slices by this procedure was found acceptable to 10 trained panelists that conducted a preliminary sensory panel (results were not given). The procedure was found acceptable to 10 trained panellists that conducted a preliminary sensory panel (results were not given). The results of listerial counts during cold storage of smoked salmon slices coated with different films are presented in Table 3. The samples did not show any significant differences in L. innocua counts at 0th day. However, the salmon slices coated with acidified LYS containing WP-OLE films showed significantly lower (p < 0.05) L. innocua counts than all the other samples (uncoated, coated with WP-OLE, LYS containing WP-OLE (non-acidiﬁed) or WP-OLE (acidiﬁed) ﬁlms) on the 1st, 3rd, 5th and 7th days of cold storage. It should also be reported that the L. innocua count of salmon slices packed with acidified LYS containing WP-OLE ﬁlms was almost 0.6 decimals lower than those of uncoated salmon slices and WP-OLE coated salmon slices at the end of 1 week cold storage. The results in the ﬁrst ﬁve days also clearly showed the lack of any considerable antimicrobial effect of non-acidiﬁed LYS containing WP-OLE ﬁlms and acidiﬁed LYS lacking WP-OLE ﬁlms. The results of the present study and the recent ﬁndings of Min et al., (2008) that successfully inhibited the growth of L. monocytogenes in cold stored salmon slices with LYS containing WP ﬁlms clearly show the good potential of employing LYS containing edible WP ﬁlms as an effective antilisterial hurdle.

3.4. Mechanical properties of ﬁlms

The results showing ﬁlm thickness, tensile strength at break, elongation at break and Young’s modulus of WP ﬁlms are presented in Table 4. The incorporation of LYS and OLE into WP ﬁlms did not cause a signiﬁcant change in tensile strengths of ﬁlms (p > 0.05), but a signiﬁcant reduction (p < 0.05) occurred in tensile strength of ﬁlms by incorporation of BW. This result was expected due to local disruptions of ﬁlm matrix continuity with wax aggregates. A similar reduction of ﬁlm tensile strength was also previously reported by Fabra et al. (2008) who incorporated BW into Na-caseinate ﬁlms. The results also showed signiﬁcant increases in ﬁlm elongations by incorporation of LYS and OLE due to the plasticizing effects of indicated ﬁlm components. The plasticizing effect of OLE was also reported by Arcan and Yemenciaoglu (2013) in zein ﬁlms, but these authors did not determine any plasticizing effect of LYS in zein ﬁlm system. In contrast, the incorporation of BW into WP ﬁlms did not have any effect on ﬁlm elongation. On the other hand, it should also be reported that the incorporation of LYS and OLE caused a signiﬁcant reduction in Young’s modulus values of WP ﬁlms while addition of BW into WP ﬁlm matrix did not cause any signiﬁcant change in this parameter. The overall results of mechanical tests clearly showed the presence of some modiﬁcations in mechanical properties of WP ﬁlm by incorporation of different ingredients, but the observed mechanical changes were not dramatic, and did not interfere with the applicability of the developed ﬁlms.

4. Conclusions

The results of this work clearly showed the possibility of designing activate-at-home- type antimicrobial packaging materials, using LYS, OLE and WP ﬁlms. The LYS release mechanism and antimicrobial activities of the designed ﬁlms based simply on acidification of the ﬁlm was successfully tested in laboratory model systems and on coated acidiﬁed smoked salmon slices stored at the standard cold storage temperature of 4 °C. This study showed the possibility of employing antimicrobial edible ﬁlms with pH controlled release properties to increase the safety of remained food left after opening of the original vacuum or modiﬁed atmosphere packaging at home. Further food applications are needed to optimize the inhibitory concentrations of LYS against L. monocytogenes at different conditions and determine the effects of antimicrobial packaging on different pathogenic bacteria.

Acknowledgements

We thank the Centres for Materials Research, and Biotechnology and Bioengineering at Izmir Institute of Technology (Izmir, Turkey) for providing their facilities to conduct some microscopic and antimicrobial studies of ﬁlms used in this study, respectively.

References


Table 4
Mechanical properties of WP, WP-OLE and WP-BW ﬁlms.

<table>
<thead>
<tr>
<th>LYS (mg/cm²)</th>
<th>OLE(%)</th>
<th>BW(%)</th>
<th>Film thickness (µm)</th>
<th>Tensile strength (MPa)</th>
<th>Elongation at break (%)</th>
<th>Young’s modulus (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>–</td>
<td>–</td>
<td>–</td>
<td>69 ± 0.4</td>
<td>2.9 ± 0.02</td>
<td>14.7 ± 0.3</td>
<td>17 ± 1.3</td>
</tr>
<tr>
<td>0.7</td>
<td>–</td>
<td>–</td>
<td>71 ± 0.7</td>
<td>2.8 ± 0.44</td>
<td>37.7 ± 12</td>
<td>7.6 ± 2.1</td>
</tr>
<tr>
<td>0.7</td>
<td>9</td>
<td>–</td>
<td>76 ± 1.2</td>
<td>2.8 ± 0.26</td>
<td>30.1 ± 11.2</td>
<td>10.7 ± 5.0</td>
</tr>
<tr>
<td>0.7</td>
<td>–</td>
<td>30</td>
<td>103 ± 5.2</td>
<td>2.1 ± 0.14</td>
<td>14.1 ± 3.5</td>
<td>15.6 ± 3.3</td>
</tr>
</tbody>
</table>

Note: Different letters in each column show signiﬁcant difference at p < 0.05. Concentrations of beeswax and oleic acid as % of WP (w/w).