Chitosan/whey Protein (CWP) Edible Films Efficiency for Controlling Mould Growth and on Microbiological, Chemical and Sensory Properties During Storage of Göbek Kashar Cheese

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Abstract
The objective of present study was to evaluate the effects of the application of chitosan and chitosan/whey protein on the chemical, microbiological and organoleptic properties of Göbek Kashar cheese during ripening time (on 3rd, 30th, 60th and 90th d). Difference in microbiological and chemical changes between samples was found to be significant ($p<0.05$) during ripening period. Cheese samples with edible coating had statistically lower mould counts compared to the uncoated samples. Furthermore the highest and lowest mould counts were determined in control (4.20 Log CFU/g) and other samples (<1 Log CFU/g) at 60th and 90th d of storage. All samples exhibited higher levels of water soluble nitrogen and ripening index at the end of storage process. At the end of 90 day storage period, no significant differences in salt and fat values were observed among the cheeses studied. The edible coatings had a beneficial effect on the sensory quality of cheese samples. In the result of sensory analysis, while cheese C and the chitosan coated cheese samples were more preferred by the panelists, the chitosan/whey protein film-coated cheese samples received the lowest scores. This study shows coating suggests could be used to improve the quality of cheese during ripening time.

Key words: chitosan, chitosan/whey protein, Göbek Kashar, ripening

Introduction
As a type of cheese, Kashar is manufactured and consumed in an extended geography including Turkey, where the production types and techniques of Kashar may vary depending largely on regions, producers and the methods applied and it has gained great diversity. Among these delicious Kashar cheese types manufactured in certain regions and offering originality is the Göbek Kashar of Ardahan. This type of Kashar is usually produced from cow milk using traditional methods and consumed after a long ripening period changing from 3 mon to 1 year. Göbek Kashar appealing to the taste of local people and containing rich nutrition elements deserves to be the subject of the present study.

Growth of microorganisms on colonization of cheese surfaces is generally seen to be an important risk (Kousta et al., 2010) because such surfaces can allow ideal media for microbial growth by involving efficient amount of water and suitable pH conditions (Conte et al., 2013). Even though various antimicrobial applications can be done on the surfaces of some types of food using various techniques such as spraying, dipping or brushing in order to take under control microbial growth, such applications are seen to have confined effects (Ture et al., 2011) may be because the antimicrobial effects of the agents used are lost rapidly. The number of studies conducted on the maintenance of food safety and prolong shelf life has increased over the last years. In such studies, edible films to coat foods and enable antimicrobials to remain on the surface are taken into consideration because such films contain active ingredients which can be present at high concentrations where they are needed (Dos Santos Pires et al., 2008; Fajardo et al., 2010; Kristo et al., 2008; Ollé Resa et al., 2013; Ollé Resa et al., 2014; Ture et al., 2011). Furthermore the activities of edible coatings not only depend on the coating methods employed, but also the properties of the coating materials (type, amount, density, viscosity and surface tension) (Zhong et al., 2014).

Food coatings from edible films may contain lipids, proteins, carbohydrates or their combinations and are rep-

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ported to prolong food shelf-life and reduce the rate of hazardous occurrence in food products (Guilbert, 1986; Kester and Fennema, 1986; Krishna et al., 2012; Krochta and Johnston, 1997). From this point of view, the use of an edible film chitosan [poly-(1-4)-2-amino-2-deoxy-D-glucose], has been increasingly preferred in recent years. It is a hydrophilic biopolymer from alkaline N-deacetylation of chitin (Kurita, 2001) and exhibits broad-spectrum antimicrobial effects on fungi, bacteria and viruses (Coma et al., 2002; Liu et al., 2001; Rabea et al., 2003; Rhoades and Roller, 2000). Chitosan also can be seen in exoskeletons of crustaceans and insects and in the cell wall of some fungi (Mucoraceae) (Keser and Bilal, 2010) and serve as a protective additive due to its capability of holding fat and water, creating colour, improving durability and exhibiting antibacterial and antifungal characteristics (Gokmen and Gurbuz, 2011; Wang, 1992). In order to prolong shelf life, chitosan was treated on various cheese types as coating by various authors such as Mozzarella (Altieri et al., 2005), Emmental (Coma et al., 2002), Regional Saloio (Cerqueira et al., 2009) and Apulia spreadable cheese (Gammariello et al., 2008). Chitosan was used alone or as carrier of other natural antimicrobials, e.g., lysozyme (Duan et al., 2007), lysozyme and EDTA (Del Nobile et al., 2009).

To the best of authors’ knowledge, there has been no study dealing with the effect of chitosan on Göbek Kashar cheese. Therefore, the purpose of this study was to evaluate the effectiveness of using a chitosan and chitosan/whey protein (CWP) coating to prevent mould growth, and to investigate its effects on microbiological, physicochemical properties and proteolysis levels of Göbek Kashar cheese during ripening.

Materials and Methods

Materials

Cow milk was picked up from a livestock farm in Göle district, Ardahan province. Microbial rennet (1:15000) was purchased from Mayasan Company, Istanbul, Turkey. Chitosan used for coating Göbek Kashar cheeses was obtained from (Sigma-Aldrich Chemical, USA) and prepared as described by Baxter et al. (1992) with a degree of N-acetylation 9.0%. Spray-dried whey from bovine milk (product no. W1500, lot no. 81K0279), containing 11% w/w proteins (biuret) and 65% w/w lactose was purchased from Sigma Chemical Co. (USA).

Production of Göbek Kashar cheese

Since no literature could be found about Göbek Kashar cheese, the term “Göbek” was accepted to be used for this local type and it was prepared following the method composed of the experiences of elderly villagers and cheese producers in the province. Cheese samples were produced in a local dairy manufacturer (Alibey plant, Göle, Ardahan). Göbek Kashar cheese was produced traditionally from raw cow milk without starter culture. The raw cow milk (300 kg) was heated to 32°C and rennet diluted in water (1/10) was added for coagulation. After 30 min, the coagulum was cut into cubes and allowed to rest for 15 min. When pH reached 5.5, whey was removed from curd in the tank and the curd was taken to a vat. Boiling process was conducted at 83°C and 5.5% brine. Boiled cheese paste was cut into pieces in moulding machine at 75 to 83°C, desired weight and put into moulds. After resting in mould for 10 min, mould was removed and past expanded. After resting for an additional 10 min out of mould, Göbek Kashar cheese was taken on a wooden bench. Cheese samples were stored on benches for 10 hours at room temperature and then additional 10 hours at 10 to 15°C. Göbek Kashar cheese, a traditional cheese type, gets its name from its shape (göbek means belly in Turkish) and manufacturing process is given in Fig. 1.

Following the preripening process, cheese samples were divided into three groups; control (C; without edible films), chitosan (X; 0.8% w/v) and chitosan/whey protein (0.8% w/v chitosan and 2.4% w/v lyophilized milk whey; Y). Cheese samples were first analysed at the end of preripening process and before coating. Other analyses were carried out in 30 d intervals (i.e., on 3rd, 30th, 60th and 90th d) by duplicating all the analyses.

Preparation of chitosan/whey protein film forming solution

A CWP film forming solution, containing 0.8% w/v chitosan and 2.4% w/v lyophilized milk whey (11% WP), was prepared by mixing of chitosan dissolved in 0.1 N HCl and lyophilized milk whey dissolved in distilled water under continuous stirring. The pH of the chitosan-whey solution was adjusted to pH 5.0 with 0.1 N HCl, and the solution was filter-sterilized. Films were prepared by casting 32.5 mL of either chitosan (0.8% w/v) or CWP (0.8% w/v chitosan and 2.4% w/v lyophilized milk whey) solutions into polystyrene petri dishes (60×15 mm) and in the preparation of coating films was used method as in the study of Di Pierro et al. (2010).

Microbiological analysis

Dried coating solution was removed from each cheese
wheel using sterile gloves and the sample groups were subjected to microbiological analysis. For each sample, 11 g cheese was taken and diluted in 99 mL of 0.85% (wt/vol) sterile saline solution. After that, a Stomacher (Lab. Stomacher Blander 400 BA 7021, Swardmedical) was used to homogenise the samples in a sterile polyethylene bag for 1.5 min. Sterile 9 mL 0.85% (wt/vol) NaCl was used for dilution and the number of total aerobic mesophilic bacteria (TAMB; Merck, at 30±1°C for 72 h; Ozdemir and Sert, 1996); lactic acid bacteria (in MRS; Merck, at 30°C for 48 h in anaerobic conditions; Diliello, 1982); lactic acid bacteria (in M17; Merck, at 30°C for 48 h; Sert et al., 2007); coagulase-positive Staphylococcus Baird–Parker agar with egg yolk tellurite enrichment (Merck, at 37°C for 24 h; Ozdemir and Sert, 1996); coliforms (Violet Red Bile Agar, Oxoid, at 35±2°C for 48 h; Diliello, 1982) and moulds (Potato Dextrose Agar, Oxoid, at 25°C for 5 to 7 d; Koburger and Marth, 1984) were counted.

Chemical Analysis

After the removal of coating materials, each of cheese samples was fully shredded. Parameters of moisture, fat, salt and titratable acidity (by titration method with NaOH using phenolphthalein as an indicator) were measured conveniently with the method of Kurt et al. (2007). Kjeldahl method (International Dairy Federation, 1993) was used to determine total nitrogen (TN) while pH was measured using a pH-metre (WTW 340-1) as in the study of Savello et al. (1989).

Nitrogen fractions

Water-soluble nitrogen (WSN), 12% trichloroacetic acid-soluble nitrogen (TCA-SN) and pH 4.6-soluble- nitrogen in the percentage of cheese were determined in the aliquots of the same cheese extract prepared as described by Kuchroo and Fox (1982). A 20 g grated cheese sample was homogenised in 40 mL H2O for 2 min through an Ultra turrax blender (IKA, USA); stored at 40°C for 1 h; centrifuged at 3000 g for 30 min at 4°C; its fatty layer was removed and the supernatant was filtered with filter paper (Whatman 113). Twenty five mL extract prepared for WSN was taken at an equal volume of 24% (wv) and TCA was added for further fractionation of the nitrogenous compounds. The mixtures were incubated for 2 h at room temperature. Precipitates were filtered through filter paper (Polychroniadou et al., 1999). Contents of WSN, TCA-SN and pH 4.6-SN were determined by Kjeldahl method. The ripening index (RI) was determined using the formula WSN/ TN×100.

Sensory evaluation

Eight panellists experienced in the sensorial evaluation of Göbek Kashar cheese assessed the cheese samples on...
Counts of LAB in MRS were determined in cheese samples to range between 6.23 and 7.76 Log CFU/g which were found to be significantly lower ($p<0.05$) in control samples on 3rd and 30th days of ripening by closing as the values measured for coated samples. These results were found to be in convenience with the values found in previous studies (Yılmaz and Dagdemir, 2012). Di Pierro et al. (2011) found lactic acid bacteria (MRS) to be in the range of 5 to 6 Log CFU/g for the control on 14th d and for the chitosan/whey protein film-coated cheese samples on 30th d, which are lower than the findings in the present study. As can be seen in Fig. 2(c), the LAB (M17) values were found in cheese samples to be between 5.84 and 7.25 Log CFU/g. Yılmaz and Dagdemir (2012) found high results to be between 6.73 and 8.13 for 120 d of coated beeswax coated Kashar cheese samples. These results showed that coating with chitosan and CWP did not have any negative influences on the growth of necessary microorganisms for the maturation of cheese.

The treatment and storage processes were found to affect significantly the moulds ($p<0.05$) in cheese samples in the present study being between <1 and 4.15 Log CFU/g. Microbiological analyses showed that samples coated with chitosan and CWP exhibited a decrease on moulds compared to control after 90 d of storage while X represented higher mould counts than the sample Y at the end of ripening period. Similar results were reported by several studies (Sarroğlu and Oner, 2006; Yılmaz and Dagdemir, 2012). Mould counts were reported to be $10^2$ CFU/g for all samples in Yılmaz and Dagdemir (2012) while increasing for control sample from 2.75 to 4.60 Log CFU/g in the storage period and from 15th d of ripening. Sarroğlu and Oner (2006) stated that mould and yeast could not be counted in Na-caseinate film coated Kashar cheese samples from 60th d while in uncoated samples counting could not be done from 90th d. Fajardo et al. (2010) reported the counts of mould and yeast to be 4.53 and 6.06 Log (CFU/g), respectively and that on 27th d of storage, natamycin+chitosan coated cheese presented lower mould/yeast rate (4.95±0.27). Ramos et al. (2012) studied the efficacy of edible films produced from whey protein isolate, glyceroel and natamycin as antimicrobial agent. The authors showed through the viable cell counts assay that natamycin incorporated in the film displayed a cidal effect against *Y. lipolytica*. Balagué et al. (2013) observed no fungi in the cheese samples packaged with the active film on 26th d of storage at 4°C while there was fungi growth in control samples on 16th d of storage. Ollé Resa et al. (2014) studied the effectiveness of natamycin
against yeast in Port Salut cheese and exerted an initial fungicidal effect against *S. cerevisiae*. The barrier test performed in cheese with films Cas/N which means that the edible film exerted a fungicidal effect against moulds. Additionally coated materials prevented the contamination of the cheese by the microorganisms inoculated in the dipping.

Counts of coliform bacteria were found in cheese samples to be between <1 and 3.77 Log CFU/g in the present study while it was <1 Log CFU/g level in Yılmaz and Dagedmir (2012). Sarıoğlu and Oner (2006) stated that counts of coliform microorganisms could not be detected in Na-caseinate film coated Kashar cheese on 90th d while it was possible to count them in control samples on 60th d. *Staphylococcus aureus* count was under detectable level (2 Log CFU/g) in all samples during ripening which can be attributed to the scalding process applied in the production of Göbek Kashar cheese. Pranoto *et al.* (2005) determined the inhibitor effect of antimicrobial alginate film containing 0.4 % garlic oil on *S. aureus*. Torlak and Nizamoglu (2011) reported in Kashar cheese samples coated with renewable films that the counts of *S. aureus*...
decreased on 14th d compared to control group between 0.90-2.66 Log and all the film types exhibited antimicrobial effect at significant level compared to control group (p<0.05). Furthermore Mei et al. (2013) reported that the application of starch-chitosan film’s in the storage of Mongolian cheese for controlling microbial populations was effective.

**Chemical conclusion**

Chemical composition of the cheese samples is given in Table 1. Chemical changes in samples were found to be statistically significant (p<0.05) during ripening period on 90th d. Dry matter of the samples was found to change between 57.70 and 64.18%. This increase was higher for the cheese samples without coating (p<0.05). These results may show that the coating process with chitosan and CWP might have delayed moisture losses compared to control. Fajardo et al. (2010) reported the moisture content of coated Saloio cheese before storage to be significantly higher than that of uncoated cheese samples resulting mainly from the water content of the coating itself while this difference was valid at 4°C only for natamycin+chitosan coated cheese samples. Yıldırım et al. (2006) found that the dry matter content of Kashar cheese samples A (control), C (coating with casein solution), D (coating with casein solution containing natamycin) and E (dipping in natamycin solution) increased until the 60th d of storage (p<0.05) after which it did not change significantly (p>0.05).

In the present study, similar to the total dry matter content, the fat content in all the cheese samples increased with increasing ripening time (p<0.05). The lowest fat content was observed in sample Y (26.22%, w/w) while the highest in C (30.30%). In coated Kashar cheese samples, the lowest and highest fat rates were 31.37% and 43.25% while in control group they were 30.5% and 42.25% (Saroglu and Oner, 2006). Similar results were obtained by Yıldırım et al. (2006). Salt values were found in samples to be between 3.50 and 5.75%. Gulec et al. (2004) found the lowest and the highest salt rates of casein coated Kashar cheese samples and control to be 1.54% to 2.54% and 1.54% to 2.43%.

As can be seen in Table 1, the treatment and storage affected significantly pH values (p<0.05). Gulec et al. (2004) stated that pH ranged from 5.14 to 5.25 in casein coated and uncoated Kashar cheese samples on 90th of storage, which are in convenience with the present study. Lucera et al. (2014) determined pH of mozzarella cheese, monitored during the entire observation period, ranged between 6.50 and 6.30.

Acidity rates of the samples varied between 21.01 and 35.46 SH. Di Pierro et al. (2011) reported that titratable acidity of Ricotta cheese, coated with a chitosan/whey protein film, reached the same level as measured for the control sample at the end of storage.

In the present study, protein rates of the samples were between 25.12 and 28.06%. Gulec et al. (2004) reported the lowest and highest rates of protein in cheese samples and control Kashar samples to be 24.5 to 31.36% and 24.5 to 31.28% while Sarýoglu and Oner (2006) found this rate in Kashar samples and control group to be 27.70 to 30.5% and 27.30 to 31.28% respectively. WSN values increased during ripening period. Yilmaz and Dagdemir (2012) found WSN to be significantly higher in control.

| Cheese samples | Ripening time (d) | Fat (%) | Salt (%) | pH | Acidity SH (%) | Protein (%) | WSN (%) | Ripening index (%) | TCA-SN (%) | pH 4.6-SN (%) |
|---------------|------------------|---------|----------|----|----------------|-------------|---------|-------------------|------------|---------------|
| C             | 3                | 59.22±0.51a 27.30±0.32a 4.18±0.17b 5.85±0.03a 21.01±0.02b 26.74±0.03a 3.93±0.03a 14.71±0.12a | 4.54±0.30a 6.52±0.24a | 57.70±0.10b 22.32±0.02b 27.49±0.07b 3.42±0.03a 12.45±0.10b | 6.22±0.04b 7.55±0.35b |
|               | 30               | 60.70±0.78a 28.60±0.16a 4.70±0.25a 5.66±0.06a 23.27±0.06b 26.35±0.03a 4.10±0.02b 15.57±0.09a | 5.91±0.05b 6.97±0.18b | 59.42±0.19a 25.94±0.09b 25.87±0.55b 4.67±0.04a 18.07±0.20b | 6.81±0.15b 7.62±0.14b |
|               | 60               | 62.54±0.53a 29.71±0.20a 5.49±0.21b 5.49±0.06b 27.94±0.09a 25.87±0.55b 4.67±0.04a 18.07±0.20b | 6.81±0.15b 7.62±0.14b | 65.18±0.67a 25.32±0.23a 5.34±0.04a 21.11±0.12a | 7.62±0.20b 8.69±0.30b |
|               | 90               | 64.18±0.67a 30.30±0.19a 5.25±0.19a 5.34±0.04a 30.14±0.09a 25.32±0.23a 5.34±0.04a 21.11±0.12a | 7.62±0.20b 8.69±0.30b | 57.70±0.10b 22.32±0.02b 27.49±0.07b 3.42±0.03a 12.45±0.10b | 6.22±0.04b 7.55±0.35b |

*Mean values followed by different letters in the same column are significantly different (p<0.05).

Control (C) without edible films; X. chitosan (0.8% w/v); Y. chitosan/whey protein (CWP, 0.8% w/v chitosan and 2.4% w/v lyophilized milk whey).
cheese until day 120, followed by BW1 (single-layer coating). The ripening index value ranged from 14.71 to 21.11% for control sample and 12.45 to 19.95% for X and 14.10 to 18.66% for Y. Yıldırım et al. (2006) reported that the ripening index of Kashar cheese samples increased steadily until the 60th d of ripening. Gulec et al. (2004) found ripening index values of all samples were similar until 60th d while on 90th d, ripening index values of control increased more than coated samples. The coating method used significantly affected the ripening of cheese. As can be seen in Table 1, 12% TCA-SN content in the cheese samples increased with increasing ripening time ($p<0.05$). The values of 12% TCA-SN, expressed as %, showed a significant increase ($p<0.05$) during ripening process (Yılmaz and Dagdemir, 2012). Aydemir (2010) reported that the increase in WSN%TN and TCA%TN values was low when the Kashar cheese samples were ripened at 4±1°C following the pre-ripening. The pH 4.6-SN ranged from 6.52 to 8.69% for the control samples, 7.22 to 9.21% for X and 4.25 to 6.46% for Y was significant ($p<0.05$) during ripening time. The values of pH 4.6-SN were found to be 10.72-23.76% in the sample of Kashar cheese samples in Hayaloglu (2009).

**Sensory evaluation**

Results of the sensory evaluation of 90 day cheese samples on a scale from 1 (poor) to 9 (excellent) are shown in a radar plot in Fig. 3. A significant difference ($p<0.05$) was found to be between the samples for colour-appearance, taste, texture, odour and general acceptability. C sample was preferred the most by the panellists. Y sample was given the lowest scores by the panellists. Yıldırım et al. (2006) reported lower texture and taste scores of vacuum-packaged Kashar cheese compared to control and those coated with casein. Di Pierro et al. (2011) stated that compared to control, Ricotta cheese exhibited better texture conditions when it was coated with chitosan/whey protein film while no difference was found to be in visual appearance, texture, flavour and odour between uncoated and chitosan/whey protein film coated Ricotta cheese samples. Cetinkaya et al. (2004) reported significant differences between Kashar samples coated with beewax in terms of aroma, flavour, o colour, appearance and texture.

Coating of Göbek Kashar cheese with edible film composed of chitosan and chitosan/whey protein showed reducing effect on the microbial growth and extended the shelf-life of the products. It is indicated from the results that the use of chitosan/whey protein can suppress mould growth during the ripening time without any adverse effects on cheese quality. During the ripening process of cheese, the level of soluble nitrogen components determined the effect of the edible film coating on the rate and extent of proteolysis quantitatively in cheese. In terms of sensorial evaluation, panellists gave the highest scores especially to C group. This study may be important since it could indicate successfully through the coated samples with chitosan and chitosan/whey protein that the mentioned materials could be used as good coating materials in the manufacture of Göbek Kashar cheese and extended throughout this cheese production to improve quality.

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![Fig. 3. Changes in sensory characteristics of Göbek Kashar cheese after 90 d of storage.](image-url)
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