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Physiological responses and nutritional qualities of tomato fruits to chitosan coating during postharvest storage

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ABSTRACT

An experiment was conducted at the Department of Agricultural Chemistry, Bangladesh Agricultural University, Mymensingh, during the period from February to September, 2017 to evaluate the effect of chitosan coating on physiological responses and nutritional qualities of tomato fruits at postharvest storage. There were four treatments of chitosan viz. control, 0.1, 0.2 and 0.3% solution in two storage conditions viz. 4°C and room temperature. The tomato fruit samples were collected at 10, 20, 30 and 50 days after postharvest storage to assess physiological parameters viz. shelf life and weight loss as well as to determine lycopene and mineral constituents viz. Ca, Mg, P, S, Na and K. The mean weight loss of tomato fruits were 0.64, 1.28, 1.59 and 2.28% at 4°C, while it was 0.88, 1.84, 2.60 and 4.80% at room temperature at 10, 20, 30 and 50 days after postharvest storage, respectively. The shelf life of tomato fruits ranged between 58.3-100.0, 50.0-100.0, 33.3-75.0 and 16.7-66.8% at 4°C, while the ranges were 66.8-100.0, 50.0-100.0, 33.3-75.0 and 0.0-41.8% at room temperature at 10, 20, 30 and 50 days after postharvest storage, respectively. As regards to weight loss and shelf life, the study results inferred that chitosan coating with 0.2% solution is useful at postharvest storage of fruits. The study results revealed that storage conditions (4°C and room temperature) did not affect on nutrient contents of tomato fruits but significantly reduced lycopene content at refrigerated condition. But the effect of chitosan coating on different nutrient contents of tomato fruits at different days after postharvest storage were highly significant at both conditions. Finally, the study results concluded that 0.2% chitosan based coatings in tomato fruits proved to extend the shelf life by decreasing the decay incidence and weight loss, and refrigerated condition is better than that of room temperature.

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Keywords: Chitosan coating, postharvest storage, tomato, nutritional quality

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1. INTRODUCTION

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Tomato (*Lycopersicon esculentum*) is one of the most important supplementary sources of minerals, phenolics and vitamins in human diet. The estimated annual production of tomato in Bangladesh was 385 thousand metric tons in 2017-2018 fiscal year [1], which is not enough to meet up local demand for the country, thus Bangladesh government has been imported several thousand metric tons from foreign countries in every year. Tomato is highly perishable, it encounters several problems in its transportation, storage and marketing [2]. Hence, postharvest losses make its production in most parts of the world unprofitable. According to Rehman et al. [3] postharvest losses in tomatoes can be as high as 25-42% globally. Thousands of tons of vegetables and fruits go to waste annually in Bangladesh due to a lack of sufficient technologies and knowledge on postharvest handling, packaging, storage and transportation. Bangladesh Bureau of Statistics report showed that postharvest

29 loss of tomato was 27.64% while the national level loss of tomato was 64252 tons in 2015-
30 2016 [4].

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32 Chitosan is commercially produced from shells of crabs, shrimp and lobsters, and coastal
33 areas of Bangladesh produce huge amount of shrimps. Thus the raw materials for chitosan
34 production is abundant in Bangladesh, which has a wide scope of use in agricultural field. In
35 the meantime, Department of Agricultural Chemistry of Bangladesh Agricultural University
36 (BAU) has extracted chitosan from shells of crabs and shrimp using local techniques.
37 Chitosan is soluble in dilute organic acids, and its coating is non-toxic and safe, and could
38 theoretically be used as a preservative for coating fruits [5]. Chitosan exhibits antifungal
39 activity against several fungi [6]. Meanwhile, it has been well documented that chitosan has
40 broad-spectrum antimicrobial activity [7, 8] and *in vivo* studies showed that chitosan
41 treatment could control or delay postharvest decay of fruits and vegetables [9]. Owing to lack
42 of information on appropriate postharvest treatments, packaging, temperature, etc. the
43 tomato fruits not only lose their quality like consumer acceptability, nutrient status of fruits,
44 and financial income to producers but also encounter a substantial postharvest loss.
45 Considering the facts stated above, this study was undertaken to assess the physiological
46 effects of chitosan application at postharvest storage, and to determine nutritional qualities of
47 tomato fruits at different stages of storage.

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49 **2. MATERIAL AND METHODS**

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51 **2.1 Collection and Screening of Tomato Fruits**

52 To conduct this experiment 15.0 kg of fully matured and partially ripen tomato fruits (cv.
53 Ruposhi) were collected from farmer's field and immediately brought to the laboratory of the
54 Department of Agricultural Chemistry, BAU, Mymensingh. After collection, tomato fruits were
55 screened on the basis of their uniformity in shape, size and level of maturity (colour). Almost
56 similar shape, size and matured fruits were selected for the experiment. Damaged and
57 disease infected fruits were removed at the beginning.

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59 **2.2 Treatments of Chitosan**

60 Chitosan used in the experiments was collected from the Department of Agricultural
61 Chemistry, BAU, Mymensingh, which has been extracted from shells of shrimp. There were
62 4 (four) treatments of chitosan used for the experiment viz. T0 (control/ no chitosan), T1
63 (0.10% chitosan solution), T2 (0.20% chitosan solution) and T3 (0.30% chitosan solution).

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65 **2.3 Preparation of Chitosan Coating Solutions**

66 To prepare 1.0 L of 0.10, 0.20 and 0.30% chitosan solutions, at first exactly 1.0, 2.0 and 3.0
67 gm of chitosan, respectively were dissolved in three different beakers containing about 25
68 mL of glacial acetic acid. Then the content was shaken well until chitosan dissolved
69 completely. After then dissolved chitosan solution was transferred into a litre volumetric flask
70 containing about 800 mL of distilled water and shaken well. Finally, the volume was made up
71 to the mark with distilled water. Acid solution without chitosan was used as control. The pH
72 of the solution was adjusted to 5.0 with 0.1 M NaOH solution.

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74 **2.4 Postharvest Application of Chitosan**

75 Previously selected 7-8 tomato fruits were dipped for 30 seconds in each treatment of
76 chitosan, and same number of fruits were also dipped similarly in the distilled water having
77 pH 5.0 (control). All treated fruits were allowed to air dried for 1 hr at 20°C. One group was
78 regarded as a replicate, and there were three replications and two conditions (room temp.
79 and refrigerated temp.) for the experiment. Thus, there were 24 (4×3×2) groups of tomato

80 fruits in this experiment. The treated and control fruits were packaged in zip-lock bags, to
81 maintain the relative humidity (RH) about 90-95%, and finally, the samples were stored at
82 room (20⁰C) and refrigerated (4⁰C) temperature.

84 **2.5 Data Recorded at Postharvest Storage**

85 Data on shelf life and weight loss of tomato fruits were measured and recorded at 10, 20, 30
86 and 50 days after storage. One tomato fruit from each replication was also collected
87 randomly at the same interval for chemical analyses.

88 **2.6 Nutritional Quality of Tomato Fruits**

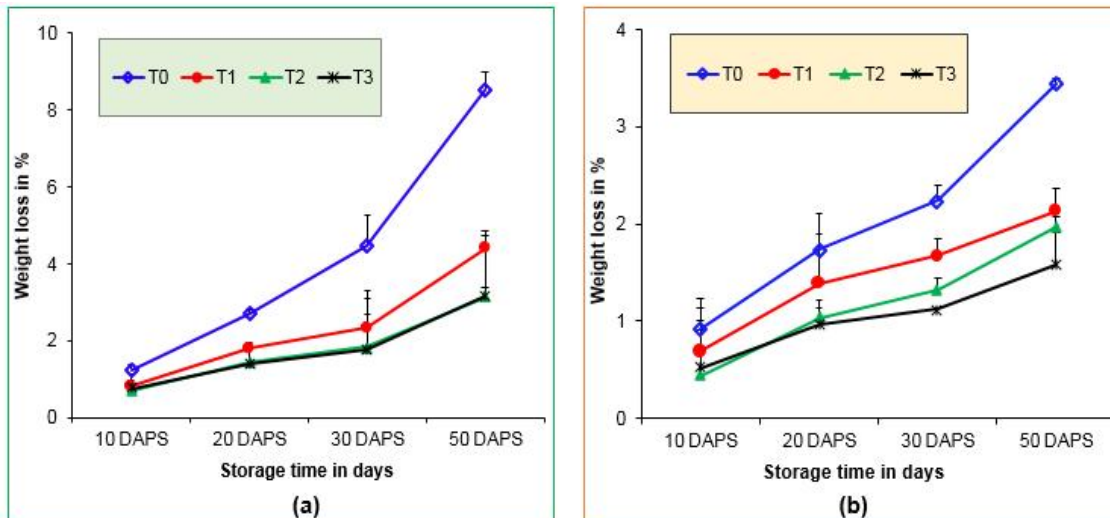
89 One tomato fruit sample from each replication was collected at 0 (fresh), 10, 20, 30 and 50
90 days interval for the determination lycopene and nutrient contents (Ca, Mg, P, K, Na and S).
91 Lycopene is responsible for the red colour of tomato. The carotenoids in the sample are
92 extracted in acetone and then taken up in petroleum ether following the method described by
93 Sadasivam and Manickam [10]. To determine different nutrient elements, collected fruit
94 samples were cut into small pieces using a sharp stainless steel knife and dried in an electric
95 oven at 50⁰C temperature for about 72 hrs. Then the samples were ground by a grinding mill
96 and used to prepare tomato fruit extract by wet oxidation method using di-acid mixture as
97 described by Singh et al. [11]. Among the nutrient elements, Ca and Mg were determined by
98 titrimetrically, P and S were measured by spectrophotometrically, and Na and K were
99 estimated by flame photometrically as mentioned by Singh et al. [11].

100 **3. RESULTS AND DISCUSSION**

101 **3.1 Weight Loss of Tomato Fruits**

102 Weight losses of tomato fruits in storage at 4⁰C (in refrigerator) and room temperature are
103 presented in Fig. 1. At 4⁰C temperature, the ranges of weight loss of tomato fruits were 0.44-
104 0.92, 0.97-1.74, 1.13-2.24 and 1.58-3.45% at 10, 20, 30 and 50 days after postharvest
105 storage (DAPS), respectively. It is apparent from Fig. 1 that the rate of weight loss was
106 higher in control treatment with the storage time at both temperature. While postharvest
107 chitosan coating treatment significantly decreased weight loss with increasing
108 concentrations. But there was very little difference in weight loss of tomato fruits at different
109 storage time between the treatments T2 and T3. The study results inferred that chitosan
110 coating with T3 (0.3% solution) is the best to retard water loss of tomato fruits in storage at
111 4⁰C temperature.

112 At room temperature, the ranges of weight loss of tomato fruits were 0.70-1.24, 1.40-2.70,
113 1.75-4.48 and 3.12-8.54% at 10, 20, 30 and 50 DAPS, respectively. Present study revealed
114 that the weight losses of tomato fruits were almost twice at different storage time, when they
115 were stored at room condition. Finally, the study results inferred that chitosan coating may
116 be used to prevent water loss of tomato fruits at postharvest storage and refrigerated
117 condition is better than that of room temperature. Similar observation was also reported by
118 Meng et al. [12] in case of table grape fruit stored at 20⁰ and 0⁰C temperature. Chien et al.
119 [13] also reported that coating of citrus fruits with low molecular weight chitosan significantly
120 decrease weight loss. They also stated that postharvest water retention prevents rapid
121 deterioration by shriveling of fruits and before shriveling becomes apparent, postharvest
122 water loss may also alter metabolism and, in some instances, accelerate fruit ripening.
123 Therefore, reducing water loss from fruit during storage or ripening helps to maintain the
124 quality of fruit.



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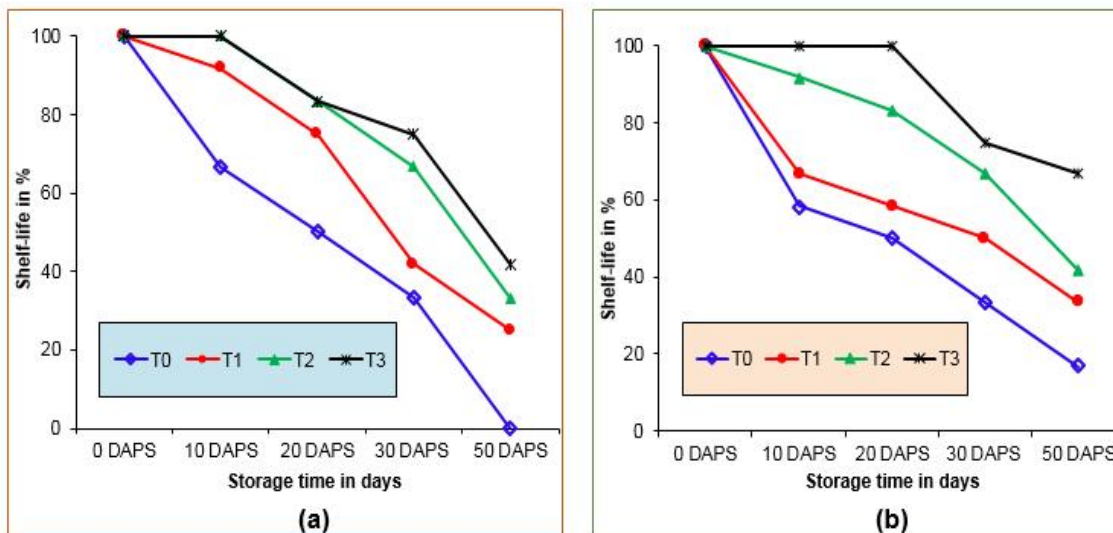
Fig. 1 Effects of different doses of chitosan coating on weight loss (in %) of tomato fruits at different days after post-harvest storage (DAPS) at room temperature (a) and 4°C temperature (b).

3.2 Shelf life of Tomato Fruits at Storage

138 Shelf lives of tomato fruits in storage at 4°C (in refrigerator) and room temperature are
139 presented in Fig. 2. At 4°C temperature, the ranges of shelf life of tomato fruits were 58.3-
140 100.0, 50.0-100.0, 33.3-75.0 and 16.7-66.8% at 10, 20, 30 and 50 DAPS, respectively. It is
141 apparent from Fig. 2 that the shelf life of tomato fruits decreased significantly in control
142 treatment with the storage time at both conditions. But postharvest chitosan coating
143 treatment significantly increased shelf life of tomato fruits with increasing concentrations. It is
144 also prominent from Fig. 2 that the treatment T3 (0.3% chitosan solution) could maintain
145 shelf life of tomato fruits 100% upto 20 days after storage. Furthermore, the shelf lives of
146 tomato fruits at storage were 75 and 66.8% at 30 and 50 days, respectively with the same
147 treatment. So, T3 treatment can be used for long time storage of tomato fruits at postharvest
148 storage at 4°C temperature.

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At room temperature, the ranges of shelf lives of tomato fruits were 66.8-100.0, 50.0-100.0,
33.3-75.0 and 0.0-41.8% at 10, 20, 30 and 50 DAPS, respectively. Present study results
revealed that there was no significant difference for shelf life of tomato fruits in between the
treatments T2 and T3. So, it can be inferred from this study that chitosan coating may be
used to extend shelf life of tomato fruits at postharvest storage and refrigerated condition is
better than that of room temperature, which might be due to controlling effect of chitosan on
postharvest diseases of tomato fruits caused by different organisms. Similar observation was
also reported by Liu et al. [14] and they stated that chitosan at 0.5 and 1% could significantly
decrease gray mould and blue mould caused by *Botrytis cinerea* and *Penicillium expansum*
in tomato fruit stored at 25 and 2°C temperature, respectively. Furthermore, Romanazzi et
al. [15] reported that chitosan application had shown promising disease control, at both
preharvest and postharvest stages. According to their report, chitosan showed a dual mode
of action on the pathogen and on the plant, as it reduces the growth of decay-causing fungi
and food borne pathogens and induces resistance responses in the host tissues.



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Fig. 2 Effects of different doses of chitosan coating on shelf-life (in %) of tomato fruits at different days after post-harvest storage (DAPS) at room temperature (a) and 4°C temperature (b).

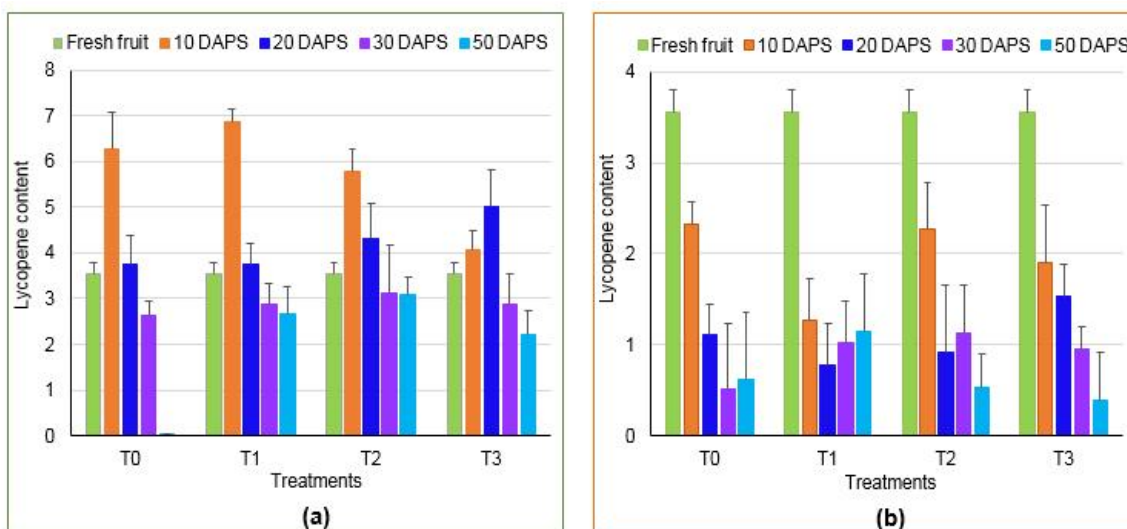
3.3 Lycopene Content of Tomato Fruits

172 Lycopene is one kind of carotenoids responsible for the red colour of tomato. The amount of
173 lycopene in tomato fruits at postharvest storage at 4°C (in refrigerator) and room
174 temperature are presented in Fig. 3. Epidemiological, as well as cell culture and animal
175 studies suggest that lycopene and the consumption of lycopene containing foods may
176 reduce cancer or cardiovascular disease risk [16]. At room temperature, the amount of
177 lycopene present in tomato fruits ranged between 4.07-6.86, 3.76-5.01, 2.64-3.12 and 0.0-
178 3.08 mg in 100 gm tomato fruits at 10, 20, 30 and 50 DAPS, respectively. The amounts of
179 lycopene were higher compared to fresh tomato (3.55 mg in 100 gm tomato fruits) at 10 and
180 20 DAPS, which might be due to extend physiological process during postharvest storage at
181 room temperature.

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183 At 4°C temperature, the amount of lycopene present in tomato fruits ranged between 1.27-
184 2.32, 0.78-1.54, 0.51-1.14 and 0.39-1.15 mg in 100 gm tomato fruits at 10, 20, 30 and 50
185 DAPS, respectively. These amounts were smaller compared to fresh tomato (3.55 mg in 100
186 gm tomato fruits), which might be due to low temperature during postharvest storage (4°C).
187 It is evident from Fig. 3 that coating of chitosan at different doses did not affect on the
188 lycopene content of tomato fruits at both temperatures. However, present study revealed
189 that in most cases, the amount of lycopene in tomato fruits decreased with postharvest
190 storage time. After bringing the fruit from room temperature to refrigerator temperature, the
191 abundance of most volatiles was greatly reduced within 3 to 5 hrs [17]. Exposure to storage
192 temperatures below 13°C may induce significant chilling injury in tomato fruit. Severity of
193 chilling injury is dependent on the length of the exposure to cold temperature as well as on
194 the ripening stage of the tomato fruit [18, 19]. They also stated that refrigerator storage at
195 around 4-6°C temperature may cause a severe alteration in fruit quality of tomato including
196 fruit discolouration and lycopene degradation. Following prolonged storage at chilling
197 temperature, a decrease in lycopene content was observed due to a decreased synthesis
198 and/or an increased breakdown. However, present study revealed that in most cases, the
199 amount of lycopene in tomato fruits decreased and/or remained unchanged with postharvest
200 storage time. Lycopene in fresh tomato fruits occurs essentially in the all-trans configuration.

201 The main causes of tomato lycopene degradation during processing are isomerization and
 202 oxidation [20]. Isomerization converts all-trans isomers to cis-isomers due to additional
 203 energy input and results in an unstable, energy-rich station.
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 206 **Fig. 3 Effects of different doses of chitosan coating on lycopene content (mg in 100**
 207 **gm sample) in tomato fruits at different days after post-harvest storage (DAPS)**
 208 **at room temperature (a) and 4°C temperature (b).**
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210 211 3.4 Nutrient Contents of Tomato Fruits

212 3.4.1 Calcium (Ca) content

213 Effect of chitosan application on Ca content of tomato fruits at different days after
 214 postharvest storage at both temperatures were highly significant at 1% level of probability
 215 (Tables 1 and 2). At refrigerated condition, the highest amounts of Ca were recorded from
 216 10, 30 and 50 DAPS at T3 (0.387%), T2 (0.514%) and T3 (0.518%) treatments, respectively.
 217 But the lowest amounts of Ca were found from 10 and 50 DAPS at T1 treatment and 30
 218 DAPS at control treatment. On the other hand, at room temperature, the maximum amounts
 219 of Ca were recorded from 10, 30 and 50 DAPS at T3 (0.421%), T2 (0.340%) and T2
 220 (0.624%) treatments, respectively. Instead, the minimum amounts of Ca were found from
 221 control treatments at different DAPS at room temperature. The amounts of Ca in tomato
 222 fruits at different DAPS both at 4°C and room temperatures were comparatively higher than
 223 the fresh tomato fruits (Tables 1 and 2). So, in context of Ca, it may be inferred that the
 224 treatment T2 (chitosan application at 0.2% solution) can be recommend for postharvest
 225 storage of tomato fruits. It is also evident from the present study that storage condition (4°C
 226 and room temperature) did not affect Ca content in postharvest storage of tomato fruits. Paul
 227 and Shaha [21] obtained 27.0±1.2 mg% Ca in tomato fruits collected from the northern
 228 region of Bangladesh. According to Parvin et al. [22], the tomato variety *Roma VF* contained
 229 0.32 to 0.69% Ca, which is almost at par with the present study.

230 **Table 1 Effects of different doses of chitosan coating on nutrient elements (Ca, Mg, P, S, Na and K) of tomato fruits at different**
 231 **days after post-harvest storage (DAPS) at 4⁰C temperature**

Treatments	Ca (%)			Mg (%)			P (%)			S (%)			Na (%)			K (%)		
	10 DAPS	30 DAPS	50 DAPS	10 DAPS	30 DAPS	50 DAPS	10 DAPS	30 DAPS	50 DAPS	10 DAPS	30 DAPS	50 DAPS	10 DAPS	30 DAPS	50 DAPS	10 DAPS	30 DAPS	50 DAPS
T ₀	0.312b	0.233d	0.402c	0.070c	0.070c	0.129c	0.004b	0.007b	0.001c	0.145b	0.226a	0.191c	0.214c	0.243c	0.230c	0.273c	0.372b	0.365c
T ₁	0.297b	0.463b	0.367d	0.094b	0.048d	0.192b	0.002c	0.008b	0.005b	0.152b	0.199b	0.215b	0.286ab	0.282b	0.248c	0.225d	0.328c	0.398b
T ₂	0.386a	0.514a	0.495b	0.181a	0.165a	0.224a	0.010a	0.012a	0.005b	0.187a	0.202b	0.261a	0.305a	0.209d	0.307a	0.404a	0.307c	0.432a
T ₃	0.387a	0.321c	0.518a	0.094b	0.145b	0.139c	0.004b	0.007b	0.007a	0.202a	0.239a	0.168d	0.268b	0.310a	0.281b	0.343b	0.404a	0.244d
LSD _{0.05}	0.0197	0.0146	0.0178	0.0103	0.0168	0.0119	0.0008	0.0020	0.0013	0.0168	0.0188	0.0157	0.0197	0.0157	0.0188	0.103	0.231	0.215
Level of significance	**	**	**	**	**	**	**	**	**	**	**	**	**	**	**	**	**	**
CV (%)	2.98	2.01	2.10	4.69	8.60	3.92	10.48	12.11	17.21	5.27	4.53	4.04	3.97	3.17	3.82	1.85	3.52	3.11
Average content in fresh fruit	0.273 ± 0.036			0.066 ± 0.018			0.003 ± 0.0002			0.158 ± 0.017			0.234 ± 0.021			0.381 ± 0.063		

232 ** = Significant at 1% level of probability

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237 **Table 2 Effects of different doses of chitosan coating on nutrient elements (Ca, Mg, P, S, Na and K) of tomato fruits at different**
 238 **days after post-harvest storage (DAPS) at room (25⁰C) temperature**

Treatments	Ca (%)			Mg (%)			P (%)			S (%)			Na (%)			K (%)		
	10 DAPS	30 DAPS	50 DAPS	10 DAPS	30 DAPS	50 DAPS	10 DAPS	30 DAPS	50 DAPS	10 DAPS	30 DAPS	50 DAPS	10 DAPS	30 DAPS	50 DAPS	10 DAPS	30 DAPS	50 DAPS
T ₀	0.233c	0.295b	0.318c	0.093c	0.179c	0.071c	0.008b	0.001c	0.008	0.221b	0.198c	0.262b	0.309b	0.259b	0.356b	0.296c	0.379ab	0.393b
T ₁	0.269b	0.296b	0.478b	0.094c	0.179c	0.093b	0.008b	0.002b	0.008	0.161c	0.239b	0.228c	0.251d	0.267ab	0.384a	0.294c	0.344c	0.365c
T ₂	0.266b	0.340a	0.624a	0.207b	0.252a	0.072c	0.009ab	0.008a	0.008	0.253a	0.291a	0.358a	0.353a	0.266ab	0.393a	0.422a	0.359bc	0.435a
T ₃	0.421a	0.308b	0.441b	0.235a	0.210b	0.176a	0.011a	0.002b	0.007	0.202b	0.229b	0.289b	0.272c	0.281a	0.394a	0.312b	0.395a	0.418a
LSD _{0.05}	0.0198	0.0197	0.0963	0.0168	0.0084	0.0133	0.0021	0.0006	0.0017	0.027	0.013	0.029	0.017	0.018	0.025	0.119	0.231	0.238
Level of significance	**	**	**	**	**	**	**	**	ns	**	**	**	**	**	**	**	**	**
CV (%)	3.51	3.42	11.01	5.72	2.24	6.58	12.37	9.56	12.19	6.73	2.84	5.38	3.04	3.51	3.40	2.01	3.34	3.14
Average content in fresh fruit	0.273 ± 0.036			0.066 ± 0.018			0.003 ± 0.0002			0.158 ± 0.017			0.234 ± 0.021			0.381 ± 0.063		

239 ** = Significant at 1% level of probability; ns = not significant

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241 **3.4.2 Magnesium (Mg) content**

242 Effect of different doses of chitosan coating on Mg content of tomato fruits at different DAPS
243 at 4°C and room temperatures are presented in Tables 1 and 2, respectively. Mg contents of
244 tomato fruits were highly significant at 1% level of probability at both conditions. At 4°C
245 temperature, the highest amounts of Mg were 0.181, 0.165 and 0.224% from 10, 30 and 50
246 DAPS, respectively at T2 treatment (0.2% chitosan solution). Alternatively, the lowest
247 amounts of Mg were recorded from 10, 30 and 50 DAPS at control treatment. Present study
248 results found that the higher doses of chitosan solution (T3 = 0.3% solution) at refrigerated
249 condition reduces the amount of Mg in tomato fruits at different DAPS.

250 In case of room temperature, the maximum amounts of Mg were recorded from 10, 30 and
251 50 DAPS at T3 (0.235%), T2 (0.252%) and T3 (0.176%) treatments, respectively.
252 Alternatively, the minimum amounts of Mg were found from control treatments at different
253 DAPS, which were statistically similar with T1 treatments of 10 and 30 DAPS. The amounts
254 of Mg in tomato fruits at different days after postharvest storage both at 4°C and room
255 temperatures were comparatively higher than the fresh tomato fruits (Tables 1 and 2). So, in
256 context of Mg, it may be inferred that the treatment T2 (chitosan application at 0.2% solution)
257 can be recommended for postharvest storage of tomato fruits. It is also evident from the
258 present study that storage condition (4°C and room temperature) did not affect Mg content in
259 postharvest storage of tomato fruits. Paul and Shaha [21] reported 17.0±1.8 mg% Mg in
260 tomato fruits, while Olaniyi et al. [23] found 0.222% Mg. Similarly, Cole et al. [24] reported
261 that tomato fruits contained 0.167% (DW) Mg and these results are almost at par with the
262 present study.

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264 **3.4.3 Phosphorus (P) content**

265 There were highly significant difference at 1% level of probability among the treatments of
266 chitosan coating on P content of tomato fruits at different DAPS at both temperatures, but at
267 room temperature, P content at 50 DAPS was insignificant (Tables 1 and 2). At 4°C
268 temperature, the highest amounts of P were 0.01, 0.012 and 0.007%, which were obtained
269 from 10, 30 and 50 DAPS, respectively at T2 and T3 treatments. Instead, the lowest
270 amounts of P were recorded from 10, 30 and 50 DAPS at T1 and control treatments. On the
271 other hand, at room temperature the maximum amounts of P were recorded from 10, 30 and
272 50 DAPS were 0.011% (T3), 0.008% (T2) and 0.008% (T0-T2), respectively, while the
273 minimum amounts of P were found from control treatments at DAPS. The amounts of P in
274 tomato fruits at different DAPS both at 4°C and room temperatures were comparatively
275 higher than the fresh tomato fruits (Tables 1 and 2). So, in context of P, it may be inferred
276 that the treatment T2 (chitosan application at 0.2% solution) can be recommended for
277 postharvest storage of tomato fruits. It is also evident from the present study that storage
278 condition (4°C and room temperature) did not affect P content in postharvest storage of
279 tomato fruits. Paul and Shaha [21] reported 28.0±1.8 mg% P in tomato fruits collected from
280 the northern region of Bangladesh. But Kadiri et al. [25] reported 1.02±0.01 mg kg⁻¹ P in
281 tomato fruits, which was almost similar to the present study.

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283 **3.4.4 Sulphur (S) content**

284 Effect of chitosan coating on S content of tomato fruits at different DAPS at both
285 temperatures were significant at 1% level of probability (Tables 1 and 2). In case of
286 refrigerated condition, the highest amounts of S were recorded from 10, 30 and 50 DAPS at
287 T3 (0.202%), T3 (0.239%) and T2 (0.261%) treatments, respectively, while the lowest
288 amounts of S were obtained from 10 and 50 DAPS at control treatment and 30 DAPS at T1
289 treatment. On the other hand, at room temperature, the maximum amounts of S were
290 recorded from 10, 30 and 50 DAPS and the contents were 0.253, 0.291 and 0.358%,
291 respectively which all were obtained from T2 (0.2% chitosan solution) treatment.

292 Alternatively, the minimum amounts of S were obtained from 10 and 50 DAPS at T1
293 treatment and 30 DAPS at control treatment. The mean amounts of S in tomato fruits at
294 different days after postharvest storage at room temperatures were almost similar to the
295 fresh tomato fruits but the amounts were little smaller at different DAPS at 4^oC (Tables 1 and
296 2). However, in context of S, it may be inferred that the treatment T2 (chitosan application at
297 0.2% solution) can be recommend for postharvest storage of tomato fruits. It is also evident
298 from the present study that refrigerated condition (4^oC) reduced S content in postharvest
299 storage of tomato fruits compared to room temperature. According to Mukta et al. [26], the
300 content of S in tomato fruits varied from 0.05 to 0.39%, which is almost at par with the
301 present study.

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3.4.5 Sodium (Na) content

304 There were highly significant difference among the treatments of chitosan coating on Na
305 content of tomato fruits at different DAPS at both temperatures (Tables 1 and 2). In case of
306 refrigerated condition, the highest amounts of Na were 0.305, 0.310 and 0.307%, which
307 obtained from 10, 30 and 50 DAPS, respectively at T2 and T3 treatments, while the lowest
308 amounts of Na were recorded from 10, 30 and 50 DAPS at control treatment. On the
309 contrary, at room temperature, the maximum amounts of Na were recorded from 10, 30 and
310 50 DAPS at T2 (0.353%), T3 (0.281%) and T3 (0.394%) treatments, respectively. But the
311 both treatments of T1 and T2 were statistically similar with T3 at 30 and 50 DAPS. However,
312 the minimum amounts of Na were found from control treatments at 30 and 50 DAPS. The
313 amounts of Na in tomato fruits at different DAPS both at 4^oC and room temperatures were
314 comparatively higher than the fresh tomato fruits (Tables 1 and 2). So, in context of Na, it
315 may be inferred that the treatment T2 (chitosan application at 0.2% solution) can be
316 recommended for postharvest storage of tomato fruits. Paul and Shaha [21] reported 5.5±0.9
317 mg% Na in tomato fruits collected from the northern region of Bangladesh, while Kadiri et al.
318 [25] found 7.73±0.9 mg kg⁻¹ Na. However, Na concentration obtained by this study was
319 greater than the reports stated above.

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3.4.6 Potassium (K) content

322 Effect of chitosan coating on K content of tomato fruits at different DAPS at both
323 temperatures were significant at 1% level of probability (Tables 1 and 2). At 4^oC
324 temperature, the highest amounts of K were recorded from 10, 30 and 50 DAPS at T2
325 (0.404%), T3 (0.404%) and T2 (0.432%) treatments, respectively, while the lowest amounts
326 of K were obtained from 10, 30 and 50 DAPS at T1, T2 and control treatments, respectively.
327 At room temperature, the maximum amounts of K were recorded from 10, 30 and 50 DAPS
328 at T2 (0.422%), T3 (0.395%) and T2 (0.435%) treatments, respectively, while the minimum
329 amounts of K were obtained from 10, 30 and 50 DAPS at T1 treatment. The mean amounts
330 of K in tomato fruits at different DAPS at both temperatures were almost similar to the fresh
331 tomato fruits (Tables 1 and 2). However, it is evident from the study results that tomato is a
332 good source of K and the treatment T2 (chitosan application at 0.2% solution) can be
333 recommend for postharvest storage of tomato fruits. According to Olaniyi et al. [23], the
334 tomato variety *Roma VF* contained 0.148% K. On the other hand, Mukta et al. [26] stated
335 that the K content in tomato fruits varied from 0.76 to 0.90%, which is almost twice than the
336 present study.

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4. CONCLUSION

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340 Chitosan coating of different doses did not affect on the lycopene content of tomato fruits at
341 both temperatures. But storage conditions (4^oC and room temperature) showed remarkable
342 affect on lycopene content of tomato fruits. Particularly, at 4^oC temperature, the amount of
343 lycopene reduced significantly compared to fresh tomato. On the contrary, storage

344 conditions did not show any remarkable change in nutrient contents of tomato fruits, but the
345 effect of chitosan coating on different nutrient contents of tomato fruits at different days after
346 postharvest storage at both temperatures were highly significant. The study results revealed
347 that postharvest chitosan coating treatment significantly decreased weight loss with
348 increasing concentrations at both 4°C and room temperatures. The rate of weight loss in
349 tomato fruits was higher in control treatment with the postharvest storage time at both
350 conditions. However, it worth mentioning that the weight losses of tomato fruits were almost
351 twice at different postharvest storage time, when they were stored at room temperature. The
352 shelf life of tomato fruits decreased significantly in control treatment with the postharvest
353 storage time at both 4°C and room temperatures. Present study results revealed that there
354 was no significant difference for shelf life of tomato fruits in between the treatments T2 and
355 T3. So, it can be inferred from this study that chitosan coating with 0.2% solution may be
356 used to prevent weight loss and to extend shelf life of tomato fruits at postharvest storage,
357 and refrigerated condition is better than that of room temperature. Finally, the study results
358 concluded that chitosan coatings have potential for extending shelf life, improving storability,
359 and enhancing some nutritional qualities of tomato fruits. At the same time, consumer
360 acceptance of such coated fruits and vegetables will also have to investigate in future.

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364 **COMPETING INTERESTS**

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Authors have declared that no competing interests exist.”

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