The Effect of Some Antioxidants Compounds on Enzymatic and Nonenzymic Browning of Quince Juice During Thermal Treatment

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Abstract

Enzymatic browning by tyrosinase and nonenzymatic browning (Maillard reaction) are responsible for producing new compounds which contributes considerably to the undesirable of aroma, taste and colour in food. In this study, tyrosinase activity and browning intensity in quince juice were inhibited by treating them with antioxidant compounds vanillin, glutathione and melatonin as compared to the control. The maximum inhibition percentage of Tyrosinase were 83, 68.8 and 90.3 % and the maximum inhibitory effect of browning intensity were 13.9, 9.7 and 29.8% respectively.

For untreated quince juice which heated to 5 hours at 100 °C, the results showed increasing in A294, Browning intensity (A420) and reducing power, as well as decreasing in reducing sugar, free amino group, phenolic compounds contents and inhibition percentage of lipid peroxidation. The decreasing of A294, browning intensity and increasing in reducing power were showed after treating with (10mM) vanillin and (8mM) glutathione compared to control. Conversely to glutathione, the addition of vanillin to quince juice revealed increasing in inhibition percentage of lipid peroxidation, reducing sugar and phenolic contents and decreasing in free amino acid content compared to control. Inhibition Mode of tyrosinase by melatonin was non-competitive. The Km value remained constant (1.96mM), while Vmax reduced from 228.13 to 114.06 U.ml⁻¹.min⁻¹. Inhibition constant Ki value was 6.9 mM.

Keywords: Tyrosinase, maillard reaction, glutathione, melatonin, Quince

تأثير بعض المركبات المضادة للأكسدة على شدة الانسمرار الأنزيمي وغير الأنزيمي في عصير فاكهة السفرجل خلال المعاملة الحرارية

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الخلاصة

ان الانسمرار الأنزيمي الحاصل بواسطة انزيم التايروسينيز والانسمرار غير الأنزيمي (تفاعل ميلارد) هما المسؤولان عن إنتاج مركبات جديدة والتي تسهم إلى حد بعيد في تكون نكهه ولون وطعم غير مرغوب به في الأغذية. في هذه الدراسة ثبتت فعالية انزيم
**Introduction:**

Quince (Cydonia oblonga Miller) fruit is a valuable dietary product belongs to family Rosaceae, which contains healthy famous fruits such as apples and pears. On maturing, the abundant hairs which cover the peel were disappeared and this stage is extremely required for processing it into candy, jam, jelly and cakes (1). The fruit contains good quantity of water, carbohydrates, proteins, tannins, organic acids and crude fibers. The fruit is a worthy source of vitamins such as A and B and rich in vitamin C. It also contains potassium, calcium, phosphorous and has very low amounts of fat (2-5). The astringency and sweetness of quince by reason of it is rich in glucose, fructose and sorbitol and also in phytic, malic, citric and quinic acids (6-7). Quince fruits are described moderately higher quantity of amino acids (8), as well as antioxidant and phenolic compounds (9).

Quince has a number of important health benefits such as expectorant, carminative and anticancer (10), being also used for the treatment of cold, influenza, migraine (11) and conjunctivitis (12). It has hypoglycemic action, antimicrobial, anti-allergic, anti-inflammatory and antiulcer and act as a boost for heart and brain (13-16).

Two forms of browning are usually detected in fruits and vegetables, namely enzymatic and non-enzymatic browning. Enzymatic browning is produced via hydroxylated of monophenolic compounds to o-diphenols that were oxidized to o-quinones by polyphenoloxidases (also known as tyrosinase, phenolase, cresolase or catecholase). Non-enzymatic browning reactions include degradation of ascorbic acid, Lipid peroxidation and Sugar caramelization as well as reactions that are initiated by a
condensation of reducing sugars, with compounds which contain a free amino group, as amino acids, peptide and proteins. This reaction is also well-known as Maillard reaction (17-19), which contributes significantly to the undesirable appearance, aroma, taste and colour (20).

Likewise to reducing carbohydrates, ascorbic acid may also react with amino compounds to form brown pigments and accelerate Maillard browning (21-22). These reactions take place through, cookery, storage and processing of foods. But they may be desirable during manufacturing of baking of bread, tea, beer and coffee (23).

Recently, various strategies have been attempted for controlling Maillard reactions. Natural compounds such as polyphenols, amino acid, peptide and vitamins, as well as enzymes were used for targeting reactive sites, intermediates, or and products (24). This controlling is very important not only to improve flavour, odour and colour, but because of likely toxicity of the products. Maillard reactions can be inhibited using different systems of the reactants or removing one of the reacting materials (25), or by using chemical inhibitors (26). Aminoguanidine was one of the compounds that has the ability of inhibiting Maillard reaction (27) by trapping α-dicarbonyls mechanism (28-29).

Experimental

Preparation of juice:

Healthy quince fruits were harvested in November from Mosul, Iraq, and used in this research. Quince (80g) were washed, peeled, cut and blended with 100 ml phosphate buffer, filtered through muslin cloth and centrifuged at 3220 xg for 15 minutes respectively to get a clear juice (30).

Inhibitors preparation:

The concentrations (2 to 10mM intervals 2) of Vanillin (VAN), Glutathione (GSH), and Melatonine (MEL) were prepared and used as inhibitors for tyrosinase and MRPs.

Tyrosinase assay:

By using tyrosine as substrate, tyrosinase activity was spectrophotometrically measured at λ = 280 nm according to the Worthington manual Decker (1977) method. One unit of tyrosinase activity was defined as the amount of enzyme that caused a change in absorbance of 0.001 per minute at 25°C and pH = 6.8 (31).
The effect of inhibitors on tyrosinase activity and browning intensity in juice:

To four millilitre of clear juice, one milliliter of individual inhibitors was added. The mixtures were used to determine tyrosinase activity at 280nm and browning intensity at 420nm. A control container was kept without inhibitors. The inhibition concentration (IC₅₀) values of melatonin were determined by following equation: IC₅₀= 50-b/a. a and b are slope and intercept respectively of the straight line.

The effect of time on tyrosinase activity and browning intensity in juice:

Three sets of six containers were used. In each one, 4ml of juice and 1ml individual concentration of VAN (10mM), MEL (10mM) and GSH (8mM) were added successively and the sixth container was kept as a control. Tyrosinase activity and browning intensity at 420nm were measured at 0-120 minutes (30).

Inhibition type of tyrosinase by melatonin:

The inhibition mode of purified tyrosinase was planned using melatonin (5.9 mM). After incubating 0.5 ml of purified enzyme with 1 ml of inhibitor, the activity was followed using (0.6 -10 mM) of L-tyrosine (30).The calculation of inhibition constant (Ki) value was accomplished by the equation:

\[ K_{m'} = K_m (1 + I/K_i) \]

Preparation of Maillard reaction products (MRPs):

One millilitre individual above concentration of VAN and GSH was added to 4 ml of Juice. To screw-sealed tubes, the solution was conveyed, locked tightly and put in water bath at 100°C. After heating the samples were taken every one hour, instantly cooled and analysed. All samples systems were prepared in dublicate. A control container was kept without inhibitors and the first measured hour without heating.

Colourless intermediate and browning intensity measurement:

The absorbance at 294nm was measured to evaluat colourless intermediate. The browning intensity was estimated by monitoring the absorbance at 420 nm. using a spectrophotometer (Shimadzu 1800 UV Spectrophotometer, Double Beam, Japan) (33).
Reducing Sugars content estimation:

The amount of reducing sugars in juice was determined by Di-nitrosalicylic acid (DNSA). In tubes of 10 ml, 1 mL of sample and 1 ml of DNSA reagent are placed. Then tubes were heated at 100 °C, 5 minutes, cooled and completed with 8 mL of distilled water. The absorbance was read at 540 nm by using Cecil 1100 spectrophotometer. The sample was replaced by distilled water in blank solution. The standard curve was prepared by using glucose (10-100mg/ml) (34).

Free amino group content estimation:

The determined of free amino group content was achieved by adding 0.2 ml of ninhydrine reagent (2mg/ml in 20 mM acetic acid – acetate buffer pH 5) to 1 ml of the sample. The solutions were heated 10 minutes at 100°C and cooled. By using a Cecil 1100 spectrophotometer, the absorbance at 570 nm was measured. The preparation of standard curve was carried out by using lysine(5-50mg/ml) (35).

Total phenolic compounds content estimation:

Total phenolics were estimated according to method described by Anesini, Ferraro and Filip (2008)(36). 1ml of Cydonia oblonga Miller juice. was added to 5 mL 1N Folin-Ciocalteu reagent. After 5 minutes, 4 mL 7.5% Na₂CO₃ was added, and the reagent mixture was kept 60 minutes at room temperature. The quantification of phenolic compounds was spectrophotometrically determined by measuring the absorbance at 765 nm, a gallic acid (10-100 μg/mL) was used for a standard curve.

Reducing power estimation:

In glass tube 0.5mL of sample was mixed with 0.5 mL of pH 6.6 sodium phosphate buffer, (0.2 M) and 0.5 mL of K₃[Fe(CN)₆] (1%). Half mL of trichloroacetic acid (TCA) (10%) was added after incubating mixture at 50°C for 20 minutes then centrifuged 10 minutes at 805 xg. 0.5 ml of supernatant was treated with 0.5 ml of distilled water and 200 μL of 0.1% FeCl₃. The increasing in absorbance at 700 nm. represents reducing power (37).

The inhibition of lipid peroxidation:

Lecithin 10.0 mg/mL was prepared in 10 mM phosphate buffered saline, pH 7.4 by stirring with magnetic stirrer (solution A). 2 ml of concentrated hydrochloric acid ,15g of TCA and 0.37g of thiobarbituric acid were adjusted to 100 mL deionized water (solution B). In glass tube 1mL of solution A was added to 1mL of 0.4 mM ascorbic acid, 1mL of 0.4 mM FeCl₃ and 1mL of sample then
transferred to dark bath at 37°C for 1hour. From solution B 2mL was added, heated at 100°C for 15 minutes, cooled and centrifuged for 10 minutes at 1096 xg. The measure of absorbance at 532nm was investigated , and named as As. Deionized water was replaced for sample as blank (Ac). The inhibition percentage (%) = (Ac-As)/Ac×100 (38-39).

Results and Discussion

The effect of inhibitors on tyrosinase activity and browning intensity in juice:

Table (1, 2) explain the inhibition percentage of tyrosinase activity and browning intensity in quince juice treated with VAN, GSH and MEL by using different concentration and different time. The results indicate that the tyrosinase activity and browning intensity were reduced as compared to the control. The percentage of inhibition increased as the concentration of antibrowning agents increases (40). GSH and cinnamic acid were studied as anti-browning agent in apple juice. The browning reaction rate could be efficiently delayed due to the inhibition of PPO (31).

### Table 1: Inhibitory effect of VAN, GSH and MEL on tyrosinase activity and browning intensity

<table>
<thead>
<tr>
<th>Inhibitors conc. (mM)</th>
<th>VAN</th>
<th>GSH</th>
<th>MEL</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Inhibitory effect of browning %</td>
<td>Tyrosinase inhibition %</td>
<td>Inhibitory effect of browning %</td>
</tr>
<tr>
<td>Control</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>11.9</td>
<td>65.5</td>
<td>9</td>
</tr>
<tr>
<td>4</td>
<td>12.5</td>
<td>70.3</td>
<td>9</td>
</tr>
<tr>
<td>6</td>
<td>12.5</td>
<td>58.4</td>
<td>8.6</td>
</tr>
<tr>
<td>8</td>
<td>12.9</td>
<td>32.5</td>
<td>9.7</td>
</tr>
<tr>
<td>10</td>
<td>13.9</td>
<td>83.8</td>
<td>9</td>
</tr>
</tbody>
</table>

### Table 2: Time Effect on tyrosinase activity and Browning intensity using VAN, GSH and MEL

<table>
<thead>
<tr>
<th>Time (min.)</th>
<th>VAN (10mM)</th>
<th>GSH (8mM)</th>
<th>MEL (10mM)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Inhibitory effect of browning %</td>
<td>Tyrosinase inhibition %</td>
<td>Inhibitory effect of browning %</td>
</tr>
<tr>
<td>0</td>
<td>15.2</td>
<td>47.5</td>
<td>14.2</td>
</tr>
<tr>
<td>15</td>
<td>16.6</td>
<td>55.5</td>
<td>16.6</td>
</tr>
<tr>
<td>30</td>
<td>19.4</td>
<td>69.3</td>
<td>18</td>
</tr>
<tr>
<td>45</td>
<td>21.8</td>
<td>78.2</td>
<td>20.1</td>
</tr>
<tr>
<td>60</td>
<td>23.7</td>
<td>88</td>
<td>22.3</td>
</tr>
<tr>
<td>90</td>
<td>25.8</td>
<td>100</td>
<td>25.1</td>
</tr>
<tr>
<td>120</td>
<td>27.9</td>
<td>100</td>
<td>29.3</td>
</tr>
</tbody>
</table>
The inhibition Mode of tyrosinase by melatonin:

The inhibition mode was studied by using melatonin. Figure 8 explains the kinetic results as double reciprocal Lineweaver-Burk plots for melatonin (5.9 mM). The nature of the inhibition was non-competitive. The $K_m$ value was (1.96mM) calculated for melatonin that remained constant, while $V_{max}$ was reduced from 228.13 to 114.06 U.ml$^{-1}$.min$^{-1}$. Inhibition constant $K_i$ value was 6.9mM. Prior inhibition studies demonstrated that GSH is noncompetitive inhibitor of apple juice PPO respectively (31). Mushroom tyrosinase was noncompetitively inhibited by cinnamic acid with $K_i$ 1.994mM (41). On the other hand, Paudel and other (2019)(42) reported competitively inhibition of tyrosinase by bromophenol.

![Figure 8. Lineweaver–Burk plot of inhibition of purified Tyrosinase by melatonin](image)

Changes in UV$_{294}$ (A$_{UV_{294}}$) and browning intensity (A$_{420}$)

Maillard reaction is related with improvement of UV-absorbing intermediate complexes, before brown pigments generation. Several intermediate products may convert to ending brown complexes, but others intermediate products are still during heating. Ajandouz and others (2011)(43) indicate that the A$_{UV_{294}}$ and browning intensity (A$_{420}$) are reflected of intermediate and browning compounds respectively. A gradual increasing in AUV$_{294}$ was found with increasing heating time for untreated quince juice as depicted in (Figure 1). These results accord to increment in absorbance at 294 nm for glucose-glycine and chitosan -glucose mixtures with heat treatment (44-45)
Browning intensity (A_{294}) of control proportionally increased by heating time (Figure 2). Browning intensity (A_{294}) increasing for numerous model systems depending on sugar natures, intermediates and coloured polymers which are resulted (46). The decreasing of A_{UV \lambda_{max}} and browning intensity by treating with vanillin and glutathione was observed compared to control (Figure 1 and 2).
Changes in reducing sugar and free amino group content:

A reducing sugar and free amino acid contents of heated quince juice, are revealed in Figure 3 and 4. A markedly lower in reducing sugar but gradually in free amino group was found when heated to 5 h respectively. The lower in reducing sugar during heating of lysine-fructose, casein-glucose and chitosan-glucose systems was described (39,47,48). A continuous reduction in the amino group content was seen as the heating time increased (45). Treated quince juice with vanillin revealed an increase in the reducing sugar, whereas the decrease was observed with glutathione with increasing heating time compared to control (Figure 3).

![Figure 3. Changes in reducing sugar content during heating for various times](chart)

The results indicated the participation of the amino group and carbonyl group of sugar in the development of MRPs during heat treatment, as confirmed by the lower free amino groups and reducing sugars remaining upon heat treatment. Conversely to the reduced sugar, after treating of quince juice with vanillin the free amino acid content was diminished, while increased with glutathione treatment with increasing heating time compared to control (Figure 4).
The decrease in reducing sugar and free amino acid was in agreement with increment of $A_{\text{UV, max}}$ and browning intensity. This indicated that long heating catalyzed the interaction between amino groups, and reducing sugar through glycation process. The reaction rate of glycation was influenced by carbonyl electrophilicity (49-50).

**Changes in phenolic content:**

Phenolic constituents in the presence of oxygen in many edible plant products such as fresh fruits and vegetables were oxidized by polyphenol oxidases (PPO) or tyrosinases to produce o-quinones. The polymerization of o-quinones into complex brown pigments takes place in occurrence of amino acids and proteins (51). Furthermore, quinones formed in the initial browning reaction can contribute in coupled oxidation reactions, allowing them to oxidize additional polyphenols that cannot be directly enzymatically oxidized.

The results exhibit the decrease in phenolic content with increasing heating time. Treating the juice with vanillin and glutathione, phenolic content were increased and reduced compared to control respectively (Figure 5).
Changes in reducing power:

Figure 6 shows a proportionally slow increase in reducing power to end of heating time. Both vanillin and glutathione treatment were higher than control in reducing power absorbance. With increasing heating time, the reducing power of glucose-glycine (52), xylose-lysine (53) and glucose-lysine mixtures (52) was reported increased.
Changes in Inhibition of lipid peroxidation percentage:

Lipid peroxidation can be defined commonly as a process under which oxidants such as free radicals attack lipid containing unsaturated carbon-carbon bond to yield fatty acid hydroperoxide in presence of oxygen molecule. The unstable hydroperoxides is degraded easily to shorter chain hydrocarbons as aldehydes (45). The results in figure 7 shows the increasing inhibition of lipid peroxidation percentage evidently in the first hour of heating time and then decreased gradually with increasing heating time. Also that inhibition of lipid peroxidation percentage was incremented when treating the juice with vanillin, but it was decreased when treating it with glutathione compared to control.

![Figure 7. Changes in inhibition of lipid peroxidation during heating for various times](image)

Various efforts have been completed to control Maillard reactions in foods. For example, adding natural or artificial compounds to Maillard reactions eliminate one of the reactants, addition compounds, as N-acetylcysteine which inhibits reactions or generates colorless products (54-55), trapping of α-dicarbonyl compounds by activating hydroxyl-containing compounds, such as gallic acid (56-58). In addition to the Maillard inhibition has been effectively attained via modification of amines residues by acetylation shield lysine from further modification.(59).

Generally, browning initiation was prevented by antioxidants which react with oxygen. Antioxidants compounds also react with the intermediate products, thus breaking the chain reaction and melanin formation was prevented (60).
Conclusion:

The antioxidant properties of vanillin, glutathione and melatonin can contribute in controlling the millard reaction and inhibition of tyrosinase. Adding these compounds to foods during processing and storage may be beneficial for improving manufacture of fruit and vegetable. In some plant sources, these compounds have occurred as natural compounds, therefore more accepted as food ingredients than synthetically manufactured compounds.

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