Phytochemical Profiling of Conventional and Supercritical Ginger Extract Based Baked Bars

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Abstract

Contemporary, nutraceuticals have attracted the consumers owing to their therapeutical potential aligned with metabolic arrays. In this amnibiance, ginger is a famous herb that has the ability to mitigate various health related disorders due to its unique photchemistry with special reference to gingerol and shagoal. To evaluate the health boosting ability of ginger, product namely ginger bars were prepared by the addition of 3% ginger conventional nutraceutical (CSE) as well as 0.3% of supercritical nutraceutical (SFE). The product was observed for color tonality in the form of L*, a*, b*. Chroma and Hue. The antioxidant potential of ginger bars was assessed by different antioxidant tests i.e. TPC, DPPH, Antioxidant activity, FRAP, ABTS and metal chelating ranges from 67.45 ± 2.29 to 112.28 ± 3.81 mg GAE/100g for TPC, 8.28 ± 0.28 to 30.72 ± 1.05% for DPPH, 13.27 ± 0.45 to 33.61 ± 1.14% for antioxidant activity, 22.15 ± 0.75 to 48.81 ± 1.66 µmole TE/g for FRAP assay, 5.94 ± 0.20 to 19.05 ± 0.65 µmole TE/g for ABTS and for metal chelating it varied from 16.41 ± 0.56 to 21.22 ± 0.72 by the addendum of ginger extracts. Furthermore the ginger bars were marked by hedonic response in terms of color, crispiness, taste, flavor and overall acceptability.

Keywords: Designer product; Physico-chemical analysis; TPC; DPPH; FRAP; ABTS; FRAP; Antioxidant activity; Metal chelating

Introduction

Diet along with its constituents contributes improved state of health other than reduced risk of diseases to enhance the quality of life. These concepts motivated to the addition of functional foods in routine diet that are health boosting foods processed with biologically active ingredient in precise quantity having both qualitative and quantitative influence on health. Hence, in modern age these healthy foods are important source in management and prevention of chronic disorders [1].

In present era, the consumption rate of designer foods is increasing day by day because of its health benefits beyond to nutritional value along with enhanced shelf life owing to the addition of antioxidants that lowers the process of rancidity [2-4]. A few epochs ago, the trend of cereal based food product moved towards designer foods by the addendum of phytoceuticals that improves the health stratum along with enhanced shelf life [2]. Innately, the marked chances resulted due to the oxidation reactions that transpire slowly during storage [3]. The food recipes that are modified by the supplementation of spices have improved stability against oxidation. These spices are more often used for oxidation stability to enhance shelf life in addition to providing flavor [5].

Baked products have been recognized as best vehicles for amalgamation of ginger, although there are positive effects on the physicochemical properties of baked products after the addition of ginger along with health benefits [6]. Baking is a complex process and results in manifold physical and biochemical effects including structure formation, taste development, color formation and synthesis of health promoting and health impairing constituents [7]. The bars are well known as a cradle of carbohydrates in food pyramid that ensures that a person is taking sufficient amount of nutrients in balance to require by the body. In formulation of bars, the ingredients provide its characteristics including color, flavor, taste, texture along with calories. Other parameters that have impact on attributes of bar are replacement of sugar and fat replacement over and above to addition of spices [8].

Incorporation of antioxidants such as bioactive ingredients in food products viz., baked bars have been grown rapidly because of improved health status awareness [9]. These natural moieties also act as mold inhibitors that delay the production and growth of mold on baked products and help in improved shelf life. The other method to get the interest of consumer is to develop the formation of chemical free product by replacing undesired ingredients augmented by antioxidants and enzymes [8].

Ginger owing to be a rich source of aromatic and pleasant flavoring properties is commonly used in the preparation of baked products, condiments and curries [10]. It has strong antioxidant potential that has been verified to be effectual in lipid oxidation inhibition as well as declining the level of oxidation in baked products. Although, nutraceutics from ginger have desirable characteristics such as being natural, non-GMO food and clean label ingredient as it can be labeled as a food ingredient in the label of food product [11].

Materials and Methods

Three types of bars were prepared using best treatment of each nutraceutical_{cse} and nutraceutical_{sfe} as described in AACC (2000) method no. 10-50D. The first (T1) contained nutraceutical_{cse} whilst other (T2) enriched with nutraceutical_{sfe} along with control (T0) for comparison purpose Table 1.

Physico-chemical analysis

The prepared bars were analyzed for the color, texture and...
antioxidant potential during the storage period. The color and texture parameters of bars were measured using the methods of Parn et al. [12].

**Color analysis**

The color analysis was performed by using CIE-Lab Color Meter (CIETLAB SPACE, Color Tech-PCM, USA). Prior to analysis, the colorimeter was calibrated using the zero and white calibration plates, respectively. Samples were also analyzed to find out their hue and chroma values.

\[
C* = \left[ (a*)^2 + (b*)^2 \right]^{1/2}
\]

\[
h = \tan^{-1} \left( \frac{b*}{a*} \right)
\]

**Texture analysis**

Texture analysis was performed using texture analyzer (single arm texture analyzer TA-XT Plus, Stable Micro Systems, Surrey, UK) with a load cell of 2 kg weight. A force versus time curve for a two-cycle compression was measured, with a disk probe of 35 mm diameter and a displacement speed of 10 mm/min. Built in software of the texture analyzer was used for analyzing the data generated.

**Antioxidant potential**

Antioxidant potential of ginger bar was determined by the protocols described by Sharma and Gujral [13].

**Total Phenolic Content (TPC)**

Total phenolic contents (TPC) in ginger bars extract were measured using Folin-Ciocalteu method that was based on the reduction of phosphotungstic acid to phosphotungstic blue and as result absorbance increased due to rise in number of aromatic phenolic groups. For the purpose, 50 µL of ginger bar extract was separately added to test tube containing 250 µL of Folin-Ciocalteu’s reagent, 750 µL of 20% sodium carbonate solution and volume was made up to 5 mL with distilled water. After two hours, absorbance was measured at 765 nm using UV/visible light spectrophotometer (CECIL CE7200) against control that has all reaction reagents except sample extract. Total polyphenols was estimated and values were verbalized as gallic acid equivalent (mg gallic acid/100 g).

Total phenolic compounds of each extract in gallic acid equivalents (GAE) was calculated by following formula:

\[
C = c \times \frac{V}{M}
\]

\[C = \text{Total phenolic contents (mg/g plant extract, in GAE)}\]
\[c = \text{Concentration of gallic acid (mg/mL)}\]
\[V = \text{Volume of extract (mL)}\]
\[M = \text{Weight of ginger extract (g)}\]

**Free Radical Scavenging Activity (DPPH assay)**

Sample solution of ginger bar extract was prepared by dissolving 0.025 mL of sample extract in 10 mL of respective solvent with 3 mL of freshly prepared DPPH solution in respective solvent that was mixed with 77 µL sample extract. Each sample was kept in dark place for about 15 minutes at room temperature and decrease in absorbance was measured at 517 nm on UV/visible light spectrophotometer. Similarly, blank sample absorbance having the same amount of solvent and DPPH solution except extract was prepared and absorbance was estimated at same wavelength on UV/visible light spectrophotometer. The free radical-scavenging activity of each ginger extract can be presented as percentage reduction in DPPH due to given amount of each extract.

\[
\text{Reduction of absorbance (\%)} = \left( \frac{AB - AA}{AB} \right) \times 100
\]

\[AB = \text{Absorbance of blank sample at } t = 0 \text{ minute}\]
\[AA = \text{Absorbance of tested extract solution at } t = 15 \text{ minutes}\]

**Antioxidant Activity (AA)**

Antioxidant activity of ginger bar extracts was based on coupled oxidation of β-carotene as well as linoleic acid. In this method, 2 mg of β-carotene was dissolved in 20 mL of chloroform. A 3 mL of aliquot was taken in flask containing 40 mg linoleic acid along with 400 mg Tween 20 and the mixture was then evaporated at 40°C for 10 min using rotary evaporator to remove chloroform. This mixture was diluted with 100 mL distilled water and was mixed properly by vortex mixer to prepare emulsion. 3 mL of β-carotene emulsion as well as 0.12 mL phenolic extracts were taken in test tubes and were thoroughly mixed. Afterward, test tubes were incubated at 50°C in a water bath for time duration of 30 minutes. Absorbance of each sample was measured at 470 nm on UV/visible light spectrophotometer. The degradation rate of the extracts was also calculated according to the first order kinetic reaction using following expression.

\[
\ln a = \ln b - \frac{AA}{t}
\]

\[a = \text{Initial absorbance on 470 nm at time zero}\]
\[b = \text{Absorbance on 470 nm after 30 min}\]
\[t = \text{Time in minutes}\]

The antioxidant activity was expressed as percentage inhibition (%) relative to the control by following equation.

\[
\text{AA (\%)} = \frac{\text{Degradation rate of control} - \text{Degradation rate of sample}}{\text{Degradation rate of control}} \times 100
\]

**Ferric Reducing Antioxidant Power (FRAP) assay**

The reducing power of ginger bar extracts was determined by measuring capability of extracts to reduce ferric tripyridyltriazine into blue colored ferrous that can be detected at 593 nm. FRAP reagent was prepared by mixing 25 mL acetate buffer (0.1 M at pH 3.6), 2.5 mL TPTZ (10 mM), and 2.5 mL ferric chloride (20 mM) and was incubated at 30°C for 10 minutes. To determine reducing power of ginger extract immediately 1.5 mL of FRAP reagent was mixed with 100 µL of ginger extract or standard and 100 µL of distilled water. Then absorbance was measured at 593 nm on UV/visible light spectrophotometer. A calibration curve was drawn using trolox (0-500 µmol/mL) and was expressed as µmol trolox equivalent per gram of sample.

**ABTS (2,2-Azino-Bis, 3-Ethylbenzothiazoline-6-Sulphonic Acid) Assay**

ABTS assay is a decolorizing method, the ABTS radical was freshly
prepared by adding 5 mL of a 4.9 mM potassium persulfate solution to 5 mL of a 14 mM ABTS solution and keeping the mixture in the dark for 16 hr. This solution was diluted further with respective solvent to yield an absorbance of 0.7 ± 0.02 at 734 nm and was used for antioxidant assay. The final reaction mixture (1 mL) comprised of 950 µL of ABTS solution and 50 µL of the extract or water was mixed for 30 seconds and allowed to stay for 5 min at ambient temperature. After the absorbance was recorded at 734 nm using a UV-visible spectrophotometer (Shimadzu UV-160A, Kyoto, Japan) and compared with the control ABTS solution. A calibration curve was made by using various concentration of Trolox (780-1000 µL/mL). ABTS radical scavenging activity was expressed as µmol trolox equivalent antioxidant capacity (TEAC) per gram of sample.

### Metal chelating potential

Ferrous ions chelating activity of extracts was estimated in which ginger bar extracts (0.1 mL) were added to a solution of 2 mM FeCl$_2$ (0.05 mL). The reaction was initiated by the addition of 5 mM ferrozine (0.1 mL) and 2.75 mL of distilled water. The mixture was shaken vigorously and left at room temperature for 10 min. The absorbance of the solution was then measured at 562 nm. The scavenging activity was calculated as follows:

$$\text{MC} \% = \frac{A_{\text{blank}} - A_{\text{sample}}}{A_{\text{blank}}} \times 100$$

Where,

- $A_{\text{blank}}$: absorbance of the control reaction
- $A_{\text{sample}}$: absorbance in the presence of plant extract

Samples were analyzed in triplicate.

### Hedonic response

The resultant bars were evaluated by a trained panel of judges using a nine point hedonic test scale. Attributes to be tested on the products included various quality parameters such as that of aroma, taste, color, texture, overall acceptability, which were based on a nine point hedonic test scale.

### Results and Discussion

#### Physico-chemical analysis of bars

The baked bars prepared by using conventional nutraceutical (3%) and nutraceutical supercritical (0.3% ginger extract) concentrations were analyzed for color tone, texture and antioxidant potential to assess the impact of treatment as well as 60 days storage interval.

### Color:

The mark of consumer acceptance of food product is their color. The color of ginger conventional extract is yellowish with the passage of time. As it was observed from present study that $L^*$ value for T$_0$ was maximum 38.68 ± 1.32 due to yellowish color of ginger conventional extract while minimum for T$_1$ (32.42 ± 1.10). Collaborative effect of treatment and storage has exposed that highest $L^*$ value was recorded in T$_3$ (37.68 ± 1.28) at beginning that increased to (39.52 ± 1.42) at end of storage. However, 60 days storage has increased $b^*$ values from 34.65 ± 1.10 to 36.44 ± 1.31. Values in Figure 1 revealed that addition of ginger conventional extract in $T_1$ and ginger supercritical extract in $T_2$ bars produced significant change in chroma value i.e. 39.25 ± 1.34 and 36.97 ± 1.33, respectively however, for $T_0$ it was minimum 33.15 ± 1.13 that changed as function of time from 35.51 ± 1.21 to 36.96 ± 1.33. For hue angle, values shown in Figure 1 changed from 77.34 ± 2.63 to 80.36 ± 2.89. So, we can interpret that with the increase in concentration of ginger conventional extract $L^*$ and $a^*$ values decreased while $b^*$, chroma and hue was increased.

### Color parameters:

The results of current research work are comparable with the findings of Abdel-Samie et al. [14] observed the color parameters of ginger enriched cookies and suggested that $L^*$ value of cookies was 65.3 ± 0.6 for control cookies prepared from wheat flour and 65.9 ± 0.9 to 59.9 ± 0.2 by the addition of various concentration of ginger. Similarly the $a^*$ was 8.9 ± 0.5 in control and changed to 7.1 ± 0.2 by the addition of ginger powder. In the case of $b^*$ values, it changed between 36.6 ± 0.2 to 37.5 ± 0.5 by the adjunct of ginger powder that was 38.6 ± 0.5 in control.

Furthermore, another group of scientists Ashoush and Gadallah, [15] prepared the biscuits by the augmentation of mango kernel peel and mango kernel powder along with control wheat flour biscuits. In their research work the color intensity of control wheat flour was 62.48 ± 1.90 for $L^*$ at start that decreased to 51.22 ± 0.27 at end of storage, 8.97 ± 0.93 for $a^*$ that decreased to 2.44 ± 0.31 while for $b^*$ value increased from 31.64 ± 0.39 to 33.69 ± 2.20 in the end of storage interval. The chroma intensity increased from 32.90 ± 0.43 to 34.04 ± 2.17 however for hue it changes from 70.62 ± 1.70 to 60.00 ± 0.30. Similarly, Haase et al. [7] concluded that by changing baking temperature from 180 to 240°C the $L^*$ values of baked products changes from 79.9 ± 4.27 to 72.7 ± 4.42.
Nonetheless, Sharma and Gujral, [13] who prepared wheat flour chapatties by the addition of barely flour. In their research work the L* of wheat flour chapatties decreased from 84.7 ± 0.3 to 80.4 ± 0.2 whilst the a* value decreased from 13.44 ± 0.53 to 12.41 ± 0.14 during storage. The L* value decreased due to the production of some melanoids that decrease L* value however the a* decreased because of the product of these intermediate maillard reaction compounds that moves it towards greenish shade and bluish shade instead of yellow in the case of b*. Although the decrease in L* and a* are due to the baking in which brown pigments produced during baking [16]. Moreover, Pasqalone et al. [17] prepared the wheat flour biscuits by the enrichment of grape mac and concluded that the L*, a* and b* values of biscuits was 41.14 ± 4.20, 9.74 ± 2.14 and 32.18 ± 0.40, respectively.

Texture: Figure 2 has represented the effect of treatment as well storage on hardness of bars. It was observed that hardness for T5 (3% ginger conventional extract) was minimum 0.62 ± 0.02 in contrast to T2 (0.3% supercritical extract) 0.65 ± 0.02 and T0 (control) was maximum 0.80 ± 0.03 kg force whilst after the cookies as the force required breaking it. They depicted that the force gradually decreased from 1st day to 60th day of storage. The L* value decreased due to the production of some melanoids that decreases L* value however the a* decreased because of the product of some melanoids that moves it towards greenish shade and bluish shade instead of yellow in the case of b*. Despite the decrease in L* and a* are due to the baking in which brown pigments produced during baking. The L*, a* and b* values of biscuits was 41.14 ± 4.20, 9.74 ± 2.14 and 32.18 ± 0.40, respectively.

Antioxidant activity of bars

Total phenolic content: In baked products the main problem is the reactance that reduces the attention of people. So, some attempt was employed in present research work by incorporating ginger powder and its extract in bars having bioactive moieties of ginger that were examined for their antioxidant perspective during four days storage. It is cleared from means Table 2 TPC of bars that T2 (0.3% ginger supercritical extract) 0.65 ± 0.02 and T0 (control) 0.72 ± 0.02 kg force. Interactive effect of treatment as well as storage has depicted that hardness of bars containing 3% ginger conventional extract (T2) was less effect from 0 to 60 days storage period as compared to supercritical treatment. Storage of bars caused significant decrease in hardness from 0.75 ± 0.03 to 0.56 ± 0.02 kg. It is clear from the Figure 2 that the bars became soft with the passage of time due to which the force gradually decreased from 1st day to 60th day of storage.

Table 2: Effect of treatments and storage on TPC (mg GAE/100 g) of bars.

Table 3: Effect of treatments and storage on DPPH (%) of bars.

DPPH: It is mostly used to assess the antioxidant potential that valued the antioxidant indices through free radical scavenging. Means for DPPH Table 3 demonstrated that free radical scavenging activity of T5 was maximum 30.72 ± 1.05% followed by T2 and T0 16.88 ± 0.58% and 8.28 ± 0.28% respectively. Throughout storage interval the DPPH assay increased from 17.64 ± 0.60 to 23.56 ± 0.85% whilst, maximum increase was observed in bars prepared by ginger supercritical extract ranging from 29.38 ± 1.00 to 31.56 ± 1.14%.

Antioxidant activity (AA): Mean antioxidant potential Table 4 regarding three treatments i.e. T0 (control), T1 (3% conventional ginger extract) and T2 (0.3% supercritical ginger extract) has revealed maximum activity (33.61 ± 1.14%) was observed in T2 followed by T1 and T0 16.88 ± 0.58% and 8.28 ± 0.28% respectively. Throughout storage interval the antioxidant activity (AA) has also influenced ß-carotene bleaching rate of each treatment that was highest at 60th day 23.56 ± 0.85% and lowest on first day i.e. 21.45 ± 0.73%.

Ferric Reducing Antioxidant Potential (FRAP)

Results Table 5 have illustrated that T2 extract has maximum ferric reducing power 48.81 ± 1.66 that was low in T0 35.60 ± 1.21 μmole trolox equivalents/g ginger bar and 22.15 ± 0.75 in control bars. In the same way, significant effect was noted in storage time factor for each

![Texture](image-url)
treatment that was higher 36.34 ± 1.24 at 0 day while lower 34.64 ± 1.25 μmole trolox equivalents/g/bar at 60th day.

**ABTS assay**

From means Table 6, it was observed that maximum ABTS value was recorded in T2 19.05 ± 0.65 followed by T1 11.29 ± 0.38 and lowest was in control 5.94 ± 0.20 μmole trolox equivalents/g. Furthermore, it was also predicted as function of storage duration that maximum ABTS value 25.7% was observed at 60th day while minimum 17.66 ± 0.60% at 0 day.

**Metal chelating potential**

Values for effect of solvent and time Table 7 have shown highest chelating potential in T2 21.22 ± 0.72% followed by 17.88 ± 0.61% in T1 and 16.41 ± 0.56% in T0. Storage time also affected chelating potential as maximum amount 19.36 ± 0.70% was observed at 60th day while lowest 11.65 ± 0.42 μmole trolox equivalents/g.

**Antioxidant potential**

The results of current research were in harmony with the findings of Abdel-Samie [14] with his colleagues evaluated the effect of ginger as antioxidant on the dough mixing properties and quality of cookies and concluded that the total phenolic content of control cookies that were prepared by wheat flour alone were 78.5 ± 1.1 mg GAE/100 g of cookies that increased from 90.8 ± 0.8 to 109.8 ± 2.7 mg GAE/100 g of cookies by the gradually supplementation of ginger. Similarly, the antioxidant assay of ginger based cookies increased from 45.8 ± 1.8 to 64.6 ± 1.0% by increasing the concentration of ginger that was 41.0 ± 0.6% in control cookies.

Furthermore, Ashoush and Gadallah [15] prepared wheat flour biscuits ad concluded that the total phenolic contents on wheat flour were 1.59 ± 0.05 mg GAE/g of wheat flour biscuit that increased to 7.08 ± 0.07 mg GAE/g by the addition of mango kernel powder as well as the DPPH assay of control wheat flour biscuits were 26.13 ± 0.05% that increased to 91.57 ± 0.11% by the enrichment of mango kernel powder. At the same moment, Zhu et al. [18] assessed the antioxidant potential of defatted wheat germ and resulted that the total phenolic content in wheat germ was 14.63 ± 0.04 mg GAE/g and DPPH assay was 75%. For ABTS radical scavenging the value was 9.37 ± 0.05 mg/ml as IC50 β carotene based antioxidant activity was 35.90% and for metal chelating potential the value of wheat germ was 25.7%.

Additionally, an alternative group of researchers, Haase et al. [7] evaluated the ABTS and FRAP assay of wheat flour biscuits and clinched that after baking the ABTS assay of wheat flour based biscuits were 7.12 ± 2.06 to 7.68 ± 1.91 mmol TE/kg wheat flour though for FRAP assay the value was 3.10 ± 0.98 to 3.84 ± 1.01 mmol TE/kg wheat flour that varied by changing the baking temperature from 210°C to 240°C. Moreover, Ahmad et al. [19] who prepared tiger nut enriched biscuits and assessed for nutritional and sensory aspects. They concluded that control biscuits without tiger nut supplementation have the total phenolic content of 2.11 mg/G of wheat flour and DPPH assay of 6.51% that gradually increased by the supplementation of tiger nut flour.

Another group of scientist Sharma and Gujral [13] prepared the wheat chapatties by the addition of barely flour and concluded that total phenolic content of wheat flour based chapatties were 2062 ± 36 µg/g in flour which increased by the incorporation of barely flour but decreased during baking (2016 ± 22 µg/g of chapattie) due to the decomposition of molecules at higher temperature beyond to 80°C. Similarly, the antioxidant of wheat chapatties was 16.1 ± 1% in the start that increased up to 30.6 ± 0.4% during storage. During the processing of baking the antioxidant activity of baked products increased as compared to flour due to the maillard reaction that takes place in the availability of sugars and proteins. Some dark compounds normally brown colored are produced due to the thermal processing of baked products. These melanoidins (brown pigments) are briefly known to possess antioxidant properties [14]. Likewise, they determined the metal chelating power of wheat chapatties that was 27.4 ± 0.5% and increased to 30.9 ± 1.0% during baking and storage. In the meanwhile, they observed reducing power of wheat flour chapatties that was 29.1 ± 1.2 μmole ascorbic acid at the start and decreased slightly during storage.

Recently, Parn et al. [12] evaluated the antioxidant potential of wheat based fruit bars by utilizing date paste and concluded that the total phenolic content of bar ranges in 240.33 ± 6.35 to 224.33 ± 1.15 mg GAE/100 g although, the DPPH scavenging varied from 30.69 ± 1.06 to 32.75 ± 0.46.

**Sensory evaluation of bars**

For sensory evaluation, bars were ranked using 9 point hedonic

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**Table 5:** Effect of treatments and storage on FRAP (μmole TE/g) of bars.

| Storage intervals (days) | Treatments | Means |
|-------------------------|------------|-------|
|                         | T0         | 5.94 ± 0.20c |
|                         | T1         | 11.29 ± 0.38b |
|                         | T2         | 19.05 ± 0.65a |
| 0                       | T0         | 6.37 ± 0.22 |
|                         | T1         | 11.72 ± 0.40 |
|                         | T2         | 19.43 ± 0.66 |
| 15                      | T0         | 6.14 ± 0.20 |
|                         | T1         | 11.59 ± 0.37 |
|                         | T2         | 19.26 ± 0.62 |
| 30                      | T0         | 5.96 ± 0.18 |
|                         | T1         | 11.26 ± 0.34 |
|                         | T2         | 19.07 ± 0.57 |
| 45                      | T0         | 5.72 ± 0.22 |
|                         | T1         | 11.03 ± 0.42 |
|                         | T2         | 18.85 ± 0.72 |
| 60                      | T0         | 5.49 ± 0.20 |
|                         | T1         | 10.85 ± 0.39 |
|                         | T2         | 18.62 ± 0.67 |
| Means                   |             | 22.15 ± 0.75 |

**Table 6:** Effect of treatments and storage on ABTS (μmole TE/g) of bars.

| Storage intervals (days) | Treatments | Means |
|-------------------------|------------|-------|
|                         | T0         | 5.94 ± 0.20c |
|                         | T1         | 11.29 ± 0.38b |
|                         | T2         | 19.05 ± 0.65a |
| 0                       | T0         | 6.37 ± 0.22 |
|                         | T1         | 11.72 ± 0.40 |
|                         | T2         | 19.43 ± 0.66 |
| 15                      | T0         | 6.14 ± 0.20 |
|                         | T1         | 11.59 ± 0.37 |
|                         | T2         | 19.26 ± 0.62 |
| 30                      | T0         | 5.96 ± 0.18 |
|                         | T1         | 11.26 ± 0.34 |
|                         | T2         | 19.07 ± 0.57 |
| 45                      | T0         | 5.72 ± 0.22 |
|                         | T1         | 11.03 ± 0.42 |
|                         | T2         | 18.85 ± 0.72 |
| 60                      | T0         | 5.49 ± 0.20 |
|                         | T1         | 10.85 ± 0.39 |
|                         | T2         | 18.62 ± 0.67 |
| Means                   |             | 22.15 ± 0.75 |

**Table 7:** Effect of treatments and storage on metal chelating potential (%) of bars.
scale for their color, flavor, taste, crispiness and overall acceptability. Color being the most important character is the key of success of any product. If color does not affect then consumer would not like to even taste it. Means color marks for outcome of treatment has Figure 3 elucidated non-significant effect on color of bars; maximum 7.27 ± 0.25 were assigned to T2 (3% ginger extract) followed by T1 (7.26 ± 0.25) while minimum 7.23 ± 0.25 to T0 (0.3% ginger nutraceutical). Color scores for bars significantly decreased as a function of storage from 7.43 ± 0.25 to 7.07 ± 0.25 during sixty days. It is obvious from the Figure 3 that the color was approximately same for all the treatments and hormonally the score of color decreased with time. Flavor is one of the characteristics which make product liked or disliked by the consumers. The flavor showed various results for treatments as well as storage intervals. It is obvious from the Figure 3 that T0 got higher marks for flavor 7.33 ± 0.25 in contrast to T2 (7.05 ± 0.24) and T1 (6.66 ± 0.23) as ginger extracts in T1 and T2 caused pleasant flavor in bars. Figure 3 showed the flavor of supercritical extract based bars got maximum marks among all the treatments followed by conventional extract based and control bars. Similarly, storage study has revealed that flavor of bars also changed significantly ranging from 7.27 ± 0.25 to 6.75 ± 0.24. Values for taste (Figure 3) showed that maximum score for taste was assigned to T1 (7.29 ± 0.25) while minimum to T2 (6.97 ± 0.24). Likewise, storage also decreased the taste marks from 7.38 ± 0.25 to 6.95 ± 0.25. Crispiness specifies the crusty expertise of the food products. Same as the flavor and taste of nutraceutical, extract based bars was best from all three treatments as depicted by Figure 3. For bars crispiness Figure 4, maximum scores 7.41 ± 0.25 was noted for T1 while minimum for T1, T2 7.25 ± 0.25 and 7.15 ± 0.24, correspondingly. Storage intervals also showed significant reduction from 7.42 ± 0.25 to 6.98 ± 0.25 in bars crispiness. Figure 4 proved that the control bars have good crispiness as compared to nutraceutical based bars. In view of the overall acceptability (Figure 4), T2 was considered best with allocated marks 7.34 ± 0.25, whereas T1, at the lower level with marks 7.04 ± 0.24. Overall acceptability also decreased with time from 7.47 ± 0.25 to 6.98 ± 0.25 during sixty days storage of bars however remained highest for T1. Making an allowance for hedonic scale response, Figure 4 concluded that the bars containing 0.3% ginger supercritical extract were rated higher marks.

The results of current research work were in accordance to the finding of Abdel-Samie et al. [14] who observed the sensory profile of ginger based cookies. They concluded that the appearance color score of control cookies was 8.0 ± 1.21 that changed from 8.0 ± 0.9 to 7.5 ± 1.1 by the addition of ginger. Similarly the texture of ginger based cookies was marked as 7.3 ± 1.3 to 6.9 ± 1.1 however it was 7.4 ± 1.7 for control cookies. Furthermore, for the flavor grades were 7.5 ± 1.4 for control and 7.0 ± 1.7 to 6.4 ± 1.4 after the augmentation of ginger powder. Nonetheless, for overall acceptability best count was 8.1 ± 0.8 to 7.0 ± 1.5 for ginger cookies that was 7.4 ± 1.4 for control cookies.

Similarly, Oluwamuokomi et al. [20] prepared wheat-cassava composite biscuits by the addition of soy flour. In their research work they concluded that the crispiness of wheat flour biscuits was marked as 8.0 while taste, aroma, shape, color and overall acceptability was scored as 8.0, 7.5, 6.0, 7.0 and 8.5 correspondingly.

**Conclusion**

During research work, two types of nutraceutical bars were prepared after supplementing with ginger enriched fractions against control. In the case of bars the T1 contained 3% ginger nutraceutical and T2 supplemented with 0.3% ginger supercritical extract. The treatments and storage exhibited significant variations in color tonality and texture that decreased from 61.02 ± 2.07 at 0 day to 56.80 ± 2.04 at 60th day for L*, 7.75 ± 0.26 to 6.15 ± 0.22 for a* however the values of b*, chroma and hue increased during storage from 34.65 ± 1.18 to 36.44 ± 1.31 for b*, 35.51 ± 1.21 to 36.96 ± 1.33 for chroma and for hue the value was 77.34 ± 2.63 to 80.36 ± 2.89. Among antioxidant perspectives, T2 showed maximum values for all tests such as TPC (112.28 ± 3.81 mg GAE/100g), DPPH (30.72 ± 1.05%), antioxidant activity (33.61 ± 1.14), FRAP (48.81 ± 1.66 µ mole TE/g), ABTS (19.05 ± 0.65 µ mole TE/g) and metal chelating (21.22 ± 0.72%). T2 was followed by T1 with values 87.12 ± 2.96 mg GAE/100g, 21.45 ± 0.73% and metal chelating (21.12 ± 0.72%). T0 was followed by T1 with values 79.56 ± 2.96 mg GAE/100g, T30 16.88 ± 0.71% antioxidant activity, 35.60 ± 1.21 µ mole TE/g FRAP, 11.29 ± 0.38 µ mole TE/g ABTS and 11.29 ± 0.38% for metal chelating. During storage the antioxidant potential decreased from 92.54 ± 3.15 to 84.97 ± 3.06 mg GAE/100 g in TPC, 36.34 ± 1.24 to 34.64 ± 1.25 µ mole TE/g for FRAP and 12.51 ± 0.43 to 11.65 ± 0. µ mole TE/g for ABTS although, in DPPH it increased from 17.64 ± 0.60 to 19.56 ± 0.70%, 21.45 ± 0.73 to 23.56 ± 0.86% antioxidant activity and metal chelating potential increased from 17.66 ± 0.60 to 19.36 ± 0.70 %. Hedonic response was also assessed using 9-point hedonic scale for the estimation of color, flavor, crispiness, taste and overall acceptability of ginger bars. The maximum scores for color was 7.27 ± 0.25 (T0), 7.33 ± 0.28 (T1) for flavor, 7.29 ± 0.25 for taste (T1), 7.25 ± 0.25 for control and 7.34 ± 0.25 (T2) for overall acceptability.

**Acknowledgement**

This work was carried out under Pak-US Science and Technology project for establishment of Functional and Nutraceutical Research Section at University of
Agriculture, Faisalabad-Pakistan. The financial and technical assistance under the framework of this project is highly acknowledged.

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