Extraction and Determination of Anti-Oxidant Activity of Polyphenols from Carrot Pomace, and Their Use in Date Oat Bar

Hammad Naeem1*, Umar Niaz2 and Sarmad Sattar1

1National Institute of Food Science and Technology, Pakistan
2Department of Entomology, Pakistan

Received: November 27, 2018; Published: December 03, 2018

*Corresponding author: Hammad Naeem, National Institute of Food Science and Technology, Pakistan

Abstract

Carrot pomace has polyphenols in large amounts. Polyphenols play an important role in the prevention of degenerative diseases such as cancer and cardiovascular diseases. Carrot pomace goes to waste with all these useful compounds. Carrot pomace or its extract can be utilized in food products to combat food shortage. Extraction of molecules from biological materials by conventional techniques, such as simple maceration, is time consuming. Sonication breaks the cell membranes. It reduces considerably the extraction time and increasing the extract yield. These phenolics were extracted using conventional Maceration techniques & advance Sonication technique. The extracts were used in Functional date-oat bars. Bars were then tested for determination of total phenolics present in them. Total Polyphenol content of functional date bar having 1% extract were 3.160 mg GAE/g of extract. Antioxidant activity in terms of % inhibition of functional date bar having 1% extract is 26.70%. It is evident from the results that the extracts of carrot pomace contain significant number of polyphenols and anti-oxidant activity. The potential of reducing waste to combat the food security issues are high, as it is estimated that by 50% reduction in current food waste, the world would be saving 1314 trillion kcal per year. This represent a reduction of about 22% of the number of additional calories needed to feed the projected population by 2050.

Introduction

The carrot belongs to the family Apiaceae. The carrot originated in Asia. Carrots are particularly rich in carotene (pro-vitamin A) and also significant amount of polyphenols. Carrots are also known to have polyphenols and antioxidants. Carrots are consumed either fresh, as a salad crop, or cooked. Large quantities are also processed, either alone or in mixtures with other vegetables, by canning or freezing. Polyphenols are a group of chemical substances found in plants. Polyphenols are naturally present in plants. Polyphenols are not essential nutrients meaning that they are not required by the human body for sustaining life, but they can exert beneficial functions [1]. Fruits and vegetables are sources of polyphenols. Polyphenols play an important role in the prevention of degenerative diseases such as cancer and cardiovascular diseases. Polyphenols are antioxidants. Antioxidants are the substances that prevent oxidation. Antioxidants, polyphenols and carotenoids, may help protect cells from damage caused by free radicals. Extraction of molecules from biological materials by conventional techniques, such as simple maceration, is time consuming. The development of modern techniques such as extraction assisted by microwave or extraction assisted by ultrasound. Sonication breaks the cell membranes. It reduces considerably the extraction time and increasing the extract yield. The application of ultrasound disrupts the cell wall structures and accelerates diffusion through membranes; thus, the cell lyses and hence facilitates the release of cell contents. The potential of reducing waste to combat the food security issues are high, as it is estimated that by 50% reduction in current food waste, the world would be saving 1314 trillion kcal per year. This represent a reduction of about 22% of the number of additional calories needed to feed the projected population by 2050.
and vegetables suggest that most of these crop remains and by products can be utilized, recovered and converted into value added food products. Hence, value addition of these wastes through drying technology and extraction methods to dehydrated products and nutraceutical products, respectively could be an alternative market option for the food and the associated industries. We incorporated phenolic extract in date bars so that we can get benefit from their anti-oxidant properties. Date bars have high nutrition value. Consumer prefer date bars that are more tasted followed by proper textural features that could be obtained by equilibrium of ingredients. Antioxidant activity resulting from the presence of phenolic compounds in the bars is well established.

**Objectives**

a) To carry out proximate analysis of carrot pomace powder.

b) To determine the polyphenol content of carrot pomace extract.

c) To determine the Antioxidant activity of carrot pomace extract.

d) To produce functional date bars.

**Materials and Methods**

The research study was conducted at Food Sciences Research Institute (FSRI), National Agriculture Research Centre (NARC), Islamabad, Pakistan.

a. Selection of vegetables: Carrot vegetable were purchased from local vegetable market, Islamabad and transferred to FSRI, NARC.

b. Preparation of pomace powder: Vegetables were washed thoroughly, juice was extracted, and fresh weight of carrot pomace was taken on calibrated Top load balance. (Seedburo 8800A).

c. Drying of carrot pomace: Carrot pomace were sun dried for 10 hours and then carrot pomace dried in hot air oven (Memmet ULM-500) at 500°C for 48 hours until moisture content of carrot pomace was reduced to 10% or below. Dried carrot pomace was weighed on top load balance (Figure 1).

d. Grinding of Carrot Pomace: Dried carrot pomace was grounded to fine powder in a Cyclotech mill with a sieve size 0.5 mm and weighed.

e. Storage of pomace powder: The pomace powder was packed in air-tight zip bags and stored at refrigeration temperature 40°C until further studies (Figure 2).

**Determination of Moisture Content in Carrot Pomace Powder**

**Principle**

Sample is weighed and placed in hot air oven with set temperature at 130°C for 1 hour (60 minutes). The weight after drying is used to determine moisture content of the sample (Figure 3).
Apparatus:

i. Hot air oven (Memmert OVEN FML-13)
ii. Moisture dishes
iii. Desiccator
iv. Plastic spatula
v. Analytical weighing balance

Procedure:

Took two moisture dishes and labelled them. Take 2 g of carrot pomace powder in these moisture dishes. Note the weight of empty moisture dishes and exact weight of sample taken. Now cover the moisture dishes with lids and transferred them to hot air oven and now uncovered them [2,3]. Set the temperature at 130°C and note the time when temperature reached at 130°C. After 1 hour (60 minutes) turned off the oven and put the moisture dishes out from the oven with the help of gloves and covered the dishes with lids. Now place the moisture dishes in desiccator for 15-25 minutes (Figures 4-6). After, moisture dishes are cooled to room temperature. Remove the lids and weighed them again (Tables 1 & 2).

Table 1: Proximate analysis of carrot pomace powder. It is evident from table 2 that Carrot pomace powder has optimum composition as described by other researchers.

| Sr. No. | Proximate analysis | Composition (%) |
|---------|--------------------|-----------------|
| 1.      | Moisture           | 8.94            |
| 2.      | Ash                | 5.72            |
| 3.      | Fat                | 1.34            |
| 4.      | Fiber              | 16.43           |

Table 2: Yield of carrot pomace extract. The yield of carrot pomace extract is within range as described by earlier studies.

| Replication | Yield (%) |
|-------------|-----------|
| 1           | 24.14     |
| 2           | 25.67     |
| 3           | 26.09     |
| 4           | 25.88     |
| Mean± Std Dev. | 25.45±0.88 |

Calculation:

\[
\%_{\text{moisture}} = \frac{\text{InitialWeight} - \text{FinalWeight}}{\text{SampleWeight}} \times 100
\]

Determination of Ash in Carrot Pomace Powder

Principle:

At high temperature i.e. 550°C all organic compounds are burnt off. Minerals are less volatile than other food compounds and are not destroyed at this high temperature. The inorganic material left behind is called ash (minerals).

Apparatus:

a. Muffled furnace (CARBOLITE AAF 1100)
b. Crucibles with lids
c. Desiccator
d. Analytical weighing balance
Procedure:
Took clean crucibles at room temperature and weighed them without their lids. Now take 3g of carrot pomace powder. Note the weight accurately. Now place the lids on the crucibles and transferred them to the Muffled furnace [4]. Set the temperature at 550°C and let the sample incinerate for about 18 hours (overnight) until light whitish grey ash was obtained (Figure 7). Carefully remove the crucibles with the help of gloves and place them in Desiccator for 15-25 minutes. When the crucibles have reached room temperature weighed them accurately (Tables 3,4).

Calculation:
\[ \% Ash = \frac{\text{FinalWeight} - \text{CrucibleWeight}}{\text{sampleweight}} \times 100 \]

Determination of Crude Fiber Content in Carrot Pomace Powder

Principle:
Crude fiber is insoluble and combustible organic residue that remains after the sample has been treated under known conditions i.e. 0.25 N Sulphuric acid and 40.7% NaOH solution.

Materials and Apparatus:
A. Sulphuric acid 0.25 N
B. Sodium hydroxide 407 g/L (40.7%)
C. Distilled water
D. Crude fiber Digestion apparatus
E. Muslin cloth
F. Oven
G. Muffled furnace
H. Measuring cylinders
I. Beakers

Procedure:
Weighed an empty crucible. Weighed 2g of sample in crucible. Put the sample in beaker. Added 200mL Sulphuric acid. Boiled for 30 minutes. Turned off the heat and add 10ml of sodium hydroxide.

Table 3: Total polyphenols content of carrot pomace extract (ultrasound).

| Sample GQTL 1272 | Absorbance (nm) (Y) | Mg GAE/g of extract |
|------------------|---------------------|---------------------|
| Rep. 1           | 0.7646              | 17.96               |
| Rep. 2           | 0.6919              | 17.10               |
| Rep. 3           | 0.7358              | 17.89               |
| Rep. 4           | 1.2243              | 22.91               |
| Mean ± Std Dev.  | 18.97± 2.65         |

Table 4: Total polyphenols content of carrot pomace extract (Maceration). It is evident from table 4 & 5 that Extraction of total Polyphenol content obtained from Maceration technique was 6.11% higher than Ultrasound assisted extraction. Carrot pomace extract had high enough polyphenols content suitable for further study and preparation of functional foods.

| Sample GQTL 1272 | Absorbance (nm) (Y) | Mg GAE/g of extract |
|------------------|---------------------|---------------------|
| Rep. 1           | 0.7800              | 20.23               |
| Rep. 2           | 0.9522              | 18.21               |
| Rep. 3           | 1.0989              | 21.49               |
| Rep. 4           | 1.0099              | 20.6                |
| Mean ± Std Dev.  | 20.13± 1.38         |

Table 5: DPPH scavenging activity of polyphenols of carrot pomace extract (Ultrasound).

| Sample GQTL 1272 | Concentration(µg/mL) | Absorbance(nm) | Inhibition (%) |
|------------------|----------------------|----------------|---------------|
| Conc. 1          | 400                  | 0.8210         | 45.68         |
| Conc. 2          | 200                  | 0.8220         | 26.01         |
| Conc. 3          | 100                  | 0.8428         | 15.93         |
| Conc. 4          | 50                   | 0.8474         | 8.38          |

Table 6: DPPH Scavenging activity of polyphenols of carrot pomace extract (maceration). It is evident from table 6 & 7 that as the concentration of the Extract is decreased, the inhibition also decreases with slowed the decrease in antioxidant activity of carrot pomace extract.

| Sample GQTL 1272 | Concentration(µg/mL) | Absorbance(nm) | Inhibition (%) |
|------------------|----------------------|----------------|---------------|
| Conc. 1          | 400                  | 0.9154         | 47.22         |
| Conc. 2          | 200                  | 0.7472         | 28.12         |
| Conc. 3          | 100                  | 0.8159         | 15.02         |
| Conc. 4          | 50                   | 0.7921         | 9.56          |

Citation: Hammad N, Umar N, Sarmad S. Extraction and Determination of Anti-Oxidant Activity of Polyphenols from Carrot Pomace, and Their Use in Date Oat Bar. LOJ Med Sci 2(3)-2018. LOJMS.MS.ID.000138.
Boiled for another 30 minutes. Removed and filtered through muslin cloth (Tables 5 & 6). Washed the residue with hot distilled water to remove excess of alkali. Dried the crucible with residue at 130°C for 1 hour [5]. Let it cool and weighed (A). Ignited the residues at 600°C in muffled furnace overnight (Figure 8).

Cooled and weighed (W₂). 35-40 mL of solvent was added. Thimble and beaker were fixed in apparatus. Boiling position was set for 45 minutes. Rinsing position was set for 35 minutes (Figure 9). The solvent was collected by blocking the extraction outlet. Beaker with fat was dried at 105°C for 30 minutes and cooled in desiccator and weighed (W₂) (Tables 7 & 8).

Weight of crucible + Sample after drying (A)
Weight of crucible + Sample after ashing (B)

Calculations:

\[ \% \text{Crude fiber} = \frac{A - B}{\text{sample weight}} \times 100 \]

**Determination of Fat Content in Carrot Pomace Powder**

**Principle:**

The fat (triglycerides) can be determined by extracting the samples with suitable solvent e.g. hexane, petroleum ether etc. in a continuous extraction apparatus. Solvent is recollected, and remaining fat is oven dried and weighed.

**Material and Apparatus:**

a) Analytical balance
b) Buchi apparatus system (B-740)
c) Beakers
d) Cellulose Thimble
e) Desiccator
f) Oven
g) n hexane

**Procedure:**

3g of sample was weighed in to thimble and covered with tissue paper. Buchi classical beakers were dried at 105°C for 30 minutes.

![Figure 8: Crude fiber digestion apparatus.](image)

![Figure 9: Soxhlet apparatus.](image)

| Table 7: Proximate analysis of functional date bars. |
|-----------------------------------------------|
| Treatment        | Moisture% | Ash% | Fat% | Fiber% |
| T₀ (control)     | 17.11     | 19.95| 10.56| 1.81   |
| T₁ (1%)          | 17.07     | 20.42| 10.33| 1.66   |
| T₂ (3%)          | 17.00     | 21.04| 10.72| 1.47   |

| Table 8: Sensory Evaluation of functional date bar. It is clear from the table 13 that treatment T₂ (3% extract supplementation) was preferred over others and got the highest sensory score while control date bars were least liked and obtained the lowest sensory score. |
|-----------------------------------------------|
| Sample        | Color | Taste | Flavor | Texture | Overall acceptability |
| Control (T₀)  | 7.20  | 6.60  | 6.60   | 6.20    | 6.40                 |
| T₁ (1%)       | 7.00  | 6.80  | 7.00   | 6.40    | 6.80                 |
| T₂ (3%)       | 7.60  | 7.80  | 7.80   | 7.20    | 7.80                 |

**Calculation:**

\[ \% \text{Fat} = \frac{W₁ - W₂}{\text{sample weight}} \times 100 \]

**Extraction of Polyphenols**

**Ultrasound-Assisted Extraction**

**Principle:**

Sonicator produces ultrasonic waves which break the sample at molecular level. The solvent penetrates the sample and polyphenols are dissolved in the solvent.
Apparatus:

i. Sonicator
ii. Beakers
iii. Reagent bottles
iv. Glass funnels
v. Whatman filter paper 41
vi. Aluminum foils
vii. Round Bottom flasks
viii. Volumetric cylinder

Chemicals:

a) Ethanol (50% solution)
b) Methanol (50% solution)

Procedure:

Carefully weighed 3g carrot pomace powder and put it in each of the 4 reagent bottles (125mL). In 2 reagent bottles put 50% ethanol solution and in 2 reagent bottles was added 50% methanol solution. Sample to solvent ratio must be 1:20 [6]. Reagent bottles were closed tightly with their lids (Figure 10). Distill water was added in sonicator up to optimum level. Sonicator was turned on for 1 hour at 50°C. After 1-hour reagent bottles were taken out from sonicator and cooled to room temperature (Tables 9 & 10).

Table 9: Determination of Total Polyphenol Content of functional date bars. It is evident from table no. 9 that polyphenol content of functional date bar having 3% extract is more than the functional date bar having 1% extract.

| Sample          | Absorbance | mg GAE/g of extract |
|-----------------|------------|---------------------|
| T₀ (control)    | 0.4151     | 1.542               |
| T₁ (1%)         | 0.4746     | 3.160               |
| T₂ (3%)         | 0.5932     | 5.075               |

Filtration:

Procedure:

4 round bottom flasks were placed on their respective supports. Samples from sonicator were filtered into round bottom flasks using whatman filter paper no.41.

Solvent extraction:

Apparatus:

A. Rotary evaporator (BUCHI B-490)
B. Circulating chiller
C. Vacuum controller
D. Hot air oven
E. Beakers
F. Aluminum foil

Procedure:

Distilled water was added in water bath to optimum level. Water bath temperature was set at 50°C. Chiller was turned on and its temperature was set at 20°C. Pressure of vacuum controller was set at 120mbar. Round bottom flask was attached with condenser.
and rotator. Rotation was turned on. Solvent was evaporated and condensed in the condenser and collected in collection chamber. Solvent was evaporated until 10mL of sample remained. This procedure was repeated for each round bottom flask containing sample. Remaining extract of carrot pomace powder was collected in 30mL beakers (Figure 11). These beakers were placed in hot air oven at 45°C for 16 hours to completely evaporate remaining water. Each of the beaker was covered with aluminum foil and placed in refrigerator till further analysis.

**Maceration Extraction Technique:**

**Principle:**
Solution containing sample is placed in shaking water bath for 22 hours at 40°C. Shaking water bath agitates the sample thus, enhancing the extraction of polyphenols.

**Apparatus and Chemicals:**
- Shaking water bath
- Ethanol (50% solution)
- Methanol (50% solution)
- Carrot pomace powder
- Analytical Balance (Model 8800A)
- Conical Flasks
- Aluminum foil

**Procedure:**

4 conical flasks were taken and 3g of carrot pomace powder was weighed in each of them. 45mL of 50% ethanol solution was added in 2 conical flasks and 45mL of 50% methanol solution was added in remaining 2 flasks. Each of these conical flasks were covered with aluminum foils and were tightened by using rubber bands. Shaking water bath was filled with distilled water to optimum level [7]. Conical flasks were placed in shaking water bath for 22 hours at 40°C. Filtration, solvent extraction and pomace extract were stored by the same procedure described above for ultrasound-assisted extraction (Figure 12).

**Determination of Total Polyphenol Content**

The total polyphenol content of carrot pomace powder was determined by Folin-Ciocalteu method as explained by Singleton (1999). Gallic acid standard solutions were prepared at different concentrations (12.5, 25, 50, 100, 200, 400 µg/mL). Ethanolic solution of sample extract 10mg/mL was prepared for the analysis. 0.5mL ethanolic solution was mixed with 2.5mL of 10% folin-ciocalteu’s reagent dissolved in water and 2.5mL 7.5% Na2Co3. Blank was simultaneously prepared, containing 0.5mL ethanol, 2.5mL diluted Folin-ciocalteu reagent dissolved in water and 2.5mL 7.5% Na2Co3. Then the sample was incubated at 25°C for 30 minutes. The calibration curve was prepared from various concentrations of Gallic acid standard solutions (Figure 13). The absorbance was determined at 765nm with UV-visible Spectrophotometer. The total polyphenol contents were expressed as mg Gallic acid equivalent (GAE)/g of extract.

**Determination of Antioxidant Activity of Carrot Pomace Powder**

Anti-oxidant activity of carrot pomace extract was determined by DPPH (1, 1-diphenyl-2-picryl-hydrazyl) radical scavenging assay according to method of Brand William et al. (1995) with some modifications. The DPPH Stock Solution was prepared by dissolving 0.006g of DPPH with 25mL of methanol. Carrot pomace extract of concentrations 400µg/ml, 200µg/ml, 100µg/ml, 50µg/ml were prepared [8,9]. The working solution was obtained DPPH solution with methanol to obtain an absorbance of 0.9921 at 517nm using UV-visible spectrophotometer (Figure 14). 3mL from above mentioned working DPPH solution was mixed with 200µL of the samples at different concentrations (50-400µg/mL). The solution in the test tubes were shaken well and incubated in dark for 15 minutes at room temperature. Absorbance for all the test tubes was measured at 517nm with UV-visible spectrophotometer (Figure 15).
taken at 517nm. The Scavenging activity was estimated based on the percentage of DPPH radical.

**Preparation of Functional Date Bars**

**Ingredients:**

**A. For Control:**

i. Dates: 60%  
ii. Oats: 20%  
iii. Almonds: 16%  
iv. Coconut oil: 4%

**B. For T₁ (1%):**

i. Dates: 59%  
ii. Oats: 20%  
iii. Almonds: 16%  
iv. Coconut oil: 4%  
v. Extract: 1%

**C. For T₂ (3%):**

i. Dates: 57%  
ii. Oats: 20%  
iii. Almonds: 16%  
v. Extract: 3%

**Preparation:**

Almonds and oats were ground well in a grinder. Carrot extract were added at different ratio (1%, 3%). Dates were added into the grinder. Coconut oil was added into the grinder. All the ingredients were ground well. Transferred all the mixture into tray and placed the tray into refrigerator for 10 minutes. Removed tray out from refrigerator and cut the mixture into bars of 7.5cm length, 2.5cm width, 1cm height. Bars were packed in butter paper.

**Sensory Evaluation of Functional Date Bars:**
Sensory evaluation of these functional date bars was done for color, taste, flavor, texture and overall acceptability by a panel consisting of 5 judges male and female of different age groups and background. Sample were presented in successions and panelist were asked to rate evaluation variables according to 9-point Hedonic scale (1=dislike extremely and 9= like extremely) (Figures 15-19).

Proximate Analysis of Functional Date Bars:

The proximate analysis of functional date bars include moisture, ash, fat, crude fiber was carried out according to standard methods of AOAC (2010).

Determination of Total Polyphenol Content:

Polyphenol content of functional date bars were determined by the same method as determined from carrot pomace powder which has been described above.

Extraction of Polyphenols from Functional Date Bars:

Procedure for extraction of polyphenols from functional date bars is same as extraction of polyphenols in carrot pomace powder which has been described above.

Determination of Antioxidant Activity of Functional Date Bars:

Procedure for determination of antioxidant activity from functional date bars is same as determination of antioxidant activity from carrot pomace powder which has been described above.

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