Polyphenols, Ascorbic Acid and Carotenoids Contents and Antioxidant Properties of Habanero Pepper (Capsicum chinense) Fruit

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ABSTRACT

Their high bioactive compounds content and importance as dietary antioxidants has increased interest in Capsicum fruit. The fruit of seven Capsicum chinense Jacq. var. habanero genotypes grown in Yucatan, Mexico, were analyzed to quantify their phenolic compounds, carotenoids and ascorbic acid contents, and to measure their free radical scavenging (ABTS assay) and antioxidant activities (β-carotene/linoleic acid assay). Phenolics (20.54 to 20.75 mg/100 g sample), carotenoids (1.00 to 1.26 mg/100 g sample) and ascorbic acid contents (187.24 to 281.73 mg/100 g sample) varied between genotypes. Trolox equivalent antioxidant capacity (TEAC) ranged from 1.55 to 3.23 mM/mg sample. During the 120-min decolorization trial, antioxidant capacity decreased over time in the studied genotypes. Values ranged from 36% to 57% β-carotene bleaching during the first 30 minutes. Fruit from all seven studied genotypes are good antioxidant sources and hold promise as natural ingredients in functional foods.

Keywords: Capsicum chinense Jacq; Phenolic Compounds; Carotenoids Content; Ascorbic Acid Content; Antioxidant Activities

1. Introduction

Antioxidants are of interest because they reduce free radicals and reactive oxygen species (ROS) in the organism. ROS and other free radicals are generated as part of normal cellular metabolism and in response to environmental factors such as ultraviolet irradiation [1]. Accumulation of these highly reactive molecules in cells can damage cellular components such as lipids, membranes, nucleic acids and proteins. This oxidative stress can directly and/or indirectly lead to human diseases such as cardiovascular disease and cancer. The human body is equipped with an antioxidant defense system that deactivates these highly reactive free radicals [2]. One of the most practical ways of managing free radical activity in the body is through diet. Dietary antioxidants may play an important role in protecting against cell damage caused by free radicals by acting as radical scavengers, reducing agents and quenchers of singlet oxygen formation, and by forming complexes with pro-oxidant metals [3].

Carotenoids impart orange and red colors in vegetables. The carotenoids in peppers include capsanthin and carotene. The yellow-orange color of peppers is formed by α- and β-carotene, zeaxanthin, lutein and β-cryptoxanthin [4]. Beta-carotene is a hydrocarbon carotenoid found widely in the chloroplasts of higher plants, and exercises pro-vitamin A and powerful antioxidant activities. Carotenoids’ cancer-preventive activities have been associated with their antioxidant properties [5]. Perera and Yen [6] reported that consumption of carotenoid-rich foods reduces the incidence of several disorders such as cancers, cardiovascular diseases, age-related macular degeneration, cataracts, diseases related to compromised immune function, and other degenerative diseases.

Plant phenols include simple phenols, flavonoids, anthocyanins, lignans and lignins, stilbenes and tannins. Phenols function as antioxidants with properties similar to vitamins C and E, and β-carotene, and have received extensive study. A wide variety of spice-derived phenolic
compounds, such as capsaicin, possess potent antmuta-
genic and anticarcinogenic properties [7]. Phenolic comp-
ounds cannot be produced by the human organism and are
acquired mainly through diet. Knowledge about the
nutritional and therapeutic role of dietary phenolic anti-
oxidants is essential for development of functional foods,
which are conventional foods with augmented health be-
niefs [2].

Ascorbic acid is a required human nutrient, and func-
tions primarily as an antioxidant in biological systems,
preventing common degenerative processes. Testing of
ascorbic acid derivatives on cancer cells showed ascorbic
acid esters to have promising anticancer activity [8]. As-
corbic acid as found in most fruits and vegetables also
protects against heart disease, high cholesterol, high blood
pressure and cancer [5].

Capsicum species have been domesticated since the
advent of agriculture and have formed part of the human
diet since at least 7500 BCE. Along with beans (Phaseo-
lus spp.), corn (Zea mays L.) and cucurbits (Cucurbita-
_ceae), Capsicum species were among the first plants cul-
tivated in the Americas. Indigenous peoples in the Ame-
ricas began growing chilies between 5200 and 3400 BCE,
making them some of the oldest cultivated crops in the
Americas. Ancient farmers began the selection process
with wild Capsicum varieties, leading eventually to the
myriad varieties known today. A member of the family
Solanaceae, the Capsicum genus includes five species commonly recognized as domesticated: C. annuum, C.
baccatum, C. chinense, C. frutescens and C. pubescens.
There are also approximately 20 documented wild spe-
cies [9].

Mexico is one of the principal centers of origin and
domestication of the genus Capsicum. Some variants in
Mexico are known and used primarily in certain regions.
For example, five variants are common in Yucatan state:
Habanero (C. chinense, Jacq.); Ya’xik (green pepper);
Xcatik (sweet pepper); Chowak, Picopaloma, Sukurre (C.
annuum L.), grown in cornfields and backyard gardens;
and Ma’xik (C. annuum var. aviculare Dierb. D’Arcy &
Eshbaugh), grown in a semi-wild state in backyards [10].
Habanero pepper fruits known to be an excellent source of phytochemicals, including vitamins A and C, phenolic
compounds, flavonoids and carotenoids [5]. Hot chili pep-
ers are widely consumed in Mexico, with an average
annual per capita intake of approximately 7 to 9 kg, mak-
ing it the second most consumed vegetable after toma-
toes [11]. Such a widely consumed vegetable with vari-
ous potential health benefits merits characterization of its
bioactive compounds content to better understand their
possible applications and increase consumer awareness
of these benefits. The present study objective was to quan-
tify polyphenols, carotenoids and ascorbic acid levels,
and measure free radical scavenging activity using an

2. Materials and Methods

2.1. Capsicum chinense Jacq. Genotypes

Capsicum chinense Jacq. Genotypes were donated by the
germ plasm bank of the Instituto Nacional de Investiga-
ciones Forestales, Agrícolas y Pecuarias (INIFAP) of Me-
xico from the September 2011 harvest in Yucatán, Mex-
ico. Material was selected based on mature fruit color 82
days after transplantation. Selected colors and materials
were orange (Orange and L-184); red (Red and L-149);
yellow (L-110 and L-36); and brown (L-37). Fruit from
the selected genotypes were refrigerated during 12 hours
before their use. After that fruits were ground in a blen-
der at a 1:2 (fruit:distilled water) ratio. Samples of the
fruit were frozen and dried at ~45°C and 133 × 10⁻³ mbar
in a lyophilizer (Labconco Free Zone 4.5, Kansas City,
MI, USA).

2.2. Capsicum chinense Jacq. Extracts

Extracts were produced by first suspending 1 g (d.b.) lyo-
philized sample in 20 mL methanol aqueous solution at
80% (v/v) and stirring for 3 h at room temperature. The
suspension was centrifuged at 2500 × g for 15 min (Mis-
stral 3000i, Curtin Matheson Sci.), the supernatant separa-
ted and a second extraction done. The two supernatants
were mixed and filtered through No. 41 Whatman paper.

2.3. Total Polyphenols

Total polyphenols concentration in all samples was quan-
tified by the Folin-Ciocalteau method, according to Sin-
gleton et al. [12]. The reaction mixture consisted of 500
mL sample in 4.5 mL water to which a solution of 200
mL Folin-Ciocalteau and 500 mL saturated Na₂CO₃ was
added. This mixture was stirred and 4.3 mL distilled wa-
ter added. After 1 h, absorbance was read at 765 nm (Ther-
mospectronic Genesis 10 uv, Madison, WI, USA). Total
polyphenol content was calculated as Trolox equivalents.
Trolox standard solutions were used for constructing the
 calibration curve. Total phenols content was expressed as
mg Trolox equivalents per 100 g of sample.

2.4. Total Carotenoids

Total carotenoids concentration was quantified according
to Rodriguez-Amaya and Kimura [13]. Briefly, 1 g fresh
sample was homogenized in 20 mL acetone and the su-
pernatant decanted. This process was repeated until at-
taining complete removal of all pigments. The sample
was filtered and washed with 30 mL acetone, the acetone

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evaporated and the dry sample dissolved in 60 mL petroleum ether. The resulting solution was filtered, transferred quantitatively to a 100 mL volumetric flask, and volume completed with petroleum ether. Of this solution, 2 mL were placed in a test tube with 8 mL petroleum ether. Absorbance was read at 475 nm (Thermospectronic Genesis 10 uv, Madison, Wi, USA) and concentration calculated with a β-carotene curve.

2.5. Ascorbic Acid

Ascorbic acid concentration in all samples was quantified using the official AOAC titrimetric method [14]. Briefly, 2 mL of 3% metaphosphoric acid-8% acetic acid sample extracts were titrated with indophenol solution (25% 2,6-dichlorophenol and 2% NaHCO3 in water) until a light but distinct rose pink color appeared and persisted for more than 5 seconds. Indophenol concentration was expressed as mg of ascorbic acid equivalent to 1 mL of reagent. Ascorbic acid content was expressed as mg of indophenol equivalents per 100 g of sample.

2.6. ABTS Decolorization Assay

Antioxidant activity was analyzed following Pukalskas et al. [15]. The ABTS** radical cation was produced by reacting ABTS with potassium per sulfate. To prepare this solution, 4.0908 g NaCl, 0.1347 g KH2PO4, 0.7098 g Na2HPO4, and 0.0749 g KCl dissolved in 500 mL ultrapure water. If pH was lower than 7.4, it was adjusted with NaOH. A 70 mM K2S4O8 solution in ultrapure water was prepared. ABTS radical cation was produced by reacting 10 mL ABTS stock solution with 40 mL K2S4O8 solution and allowing the mixture to stand in darkness at room temperature for 16 - 17 h before use. The radical was stable in this form for more than 2 days when stored in darkness at room temperature.

Antioxidant compound content in all samples was analyzed by diluting the ABTS** solution with PBS to an absorbance of 0.800 ± 0.03 AU at 734 nm. After adding 990 mL diluted ABTS** solution (A734 nm = 0.800 ± 0.03) to 10 mL antioxidant compound or Trolox standard (final concentration 0.5 - 3.5 mM) in PBS, absorbance was read at ambient temperature exactly 6 min after initial mixing. All analyses were run in triplicate. The percentage decrease in absorbance at 734 nm was calculated and plotted as a function of the Trolox concentration for the standard reference data. The radical scavenging activity of the tested samples, expressed as inhibition percentage (%I), was calculated with the equation:

\[ %I = \left( \frac{A_b - A_i}{A_b} \right) \times 100 \]

where \( A_b \) is absorbance of the blank sample (\( t = 0 \)), and \( A_i \) is absorbance of the sample with antioxidant after 6 min.

The Trolox equivalent antioxidant coefficient (TEAC) was quantified by a regression analysis of percent inhibition versus Trolox concentration using the following formula:

\[ \text{TEAC} = \left( \frac{\%I - b}{m} \right) \]

where \( b \) is the intersection and \( m \) is the slope.

2.7. β-Carotene Decolorization Assay

Antioxidant activity was quantified according to Miller [16]. The assay is based on determining the decolorization exhibited by β-carotene in response to the action of linoleic acid oxidation products. In a test tube, 1 mL 0.2 mg/mL β-carotene in chloroform was added to a mixture of 20 mg linoleic acid and 200 mg Tween 20. The chloroform was evaporated and 50 mL bidistilled water added. Aliquots (2.5 mL) of this mixture were added to tubes containing 100 mL sample. Water was used as a reference blank and 2,6-di-t-tert-butyl-4-methyl phenol (BHT) dissolved in 96% ethanol was used as a positive control. Samples were incubated at 50°C, aliquots taken at 30, 60, 90 and 120 min, and absorbance measured at 470 nm (Thermospectronic Genesis 10 uv, Madison, WI, USA).

2.8. Statistical Analysis

All results were analyzed using descriptive statistics with a central tendency and dispersion measures. One-way ANOVAs were run to evaluate polyphenols, carotenoids and ascorbic acid contents, as well as ABTS and β-carotene decolorization assay results. A LSD multiple range test was applied to determine differences between treatments. All analyses were done according to Montgomery [17] and processed with the Statgraphics Plus version 5.1 software.

3. Results and Discussion

Hot peppers are known to be a good source of phenolic compounds such as aspolyphenols, carotenoids and ascorbic acid. These phytochemicals exhibit high antioxidant activity, and their consumption has been linked to a decreased risk of developing chronic and degenerative diseases. We studied Capsicum chinense Jacq var. Habanero pepper fruit, one of the most widely consumed hot peppers in southeast Mexico. Total polyphenols, total carotenoids and ascorbic acid contents, as well as antioxidant activity (TEAC and β-carotene assay), were quantified in seven C. chinense Jacq genotypes selected based on mature fruit color 82 days after transplantation.
3.1. Total Polyphenols

Among phytochemicals, polyphenols are particularly interesting for their free radical scavenging properties and in vivo biological activities. Epidemiological studies have indicated a possible association between dietary polyphenols intake and risk of coronary heart disease and cancer [18]. Peppers (Capsicum sp.) are known for being rich in micronutrients and bioactive compounds (e.g. phenolic compounds), and are recognized as beneficial to health. Total polyphenols content in the studied genotypes ranged from 20.54 to 20.75 mg/100 g sample (Figure 1). Content was higher (p < 0.05) in red (L-149 and red) and brown (L-37) genotypes, and did not differ (p > 0.05) among yellow and orange genotypes.

Polyphenols are secondary metabolites widely distributed in plants [11]. Levels vary widely during growth and maturation and contribute to fruit pungency, bitterness, flavor and color [9]. The more intensely colored genotypes studied here had the highest polyphenols content, which agrees with previous reports of high phenolics content in red genotypes. Zhang and Hamauzu [19] examined phenolic compounds content in bell peppers (Capsicum annum L.) of different colors, and reported that phenolics content varied depending on color (9.2 - 15.4 mg GAE/100 g fresh weight). In phenolic extracts, green pepper had a higher (p < 0.05) phenolics content than red and yellow peppers, while in methanol extracts red pepper had a higher (p < 0.05) phenolics content than red and yellow peppers. The present results for Habanero genotypes are comparable to reported total phenolics content in Chiltepin (4.85 mg/g GAE) and Habanero fruits (5.92 mg/g GAE) [11].

Phenolic compounds are an important group of secondary metabolites synthesized by plants as an adaptation toiotic and abiotic stress conditions (infection, wounding, water stress, cold stress, high visible light) [2]. In recent years, phenolic compounds have attracted increasing research interest because they are potentially powerful antioxidants that can protect the human organism from free radicals, formation of which is associated with normal natural metabolism of aerobic cells. The antiradical activity of flavonoids and phenolics is based on the redox properties of their hydroxy groups and the structural relationships between different parts of their chemical structure. Phenolic compounds cannot be produced by the human body, and thus must be acquired through the diet. Understanding the nutritional and therapeutic roles of dietary phenolic antioxidants is essential to developing functional foods, which are conventional foods with augmented health benefits [2]. In a study of total phenols content in mature C. chinense fruits from 63 accessions originally acquired from Belize, Brazil, Colombia, Ecuador, Mexico, Peru, Puerto Rico and the United States, an accession from Mexico (PI-438648) was found to contain the highest total phenols content (349 μg g⁻¹ fresh fruit) [5].

A phenols content double this has been reported for C. chinense Habanero (759.12 mg/100 g fresh weight) [20]. In a study of chemical composition and biological activities in two C. annum cultivars (acuminatum small and cerasiferum) common in the Mediterranean diet, C. annum var. acuminatum extract had a phenol content (970.2 mg/100 g) twice that of C. annum var. cerasiferum (426.2 mg/100 g) [21]. These levels are higher than the total polyphenols contents reported for fresh red (296 mg/100 g), yellow (284 mg/100 g) and green (215 mg/100 g) peppers [22]. Hot peppers are clearly a good source of polyphenols, but compositional studies are needed to determine how variables such as cultivar, season and pre- and post-harvest conditions may affect pepper fruit chemical composition.

Figure 1. Polyphenols content in C. chinense Jacq. Genotypes. Data are presented as means (n = 3). *Different superscript letters indicate statistical difference (p < 0.05).

3.2. Total Carotenoids

Carotenoids are fat-soluble antioxidants found in many fruits and vegetables and are required for human epithelial cellular differentiation [19]. Carotenoids content in the studied Habanero genotypes ranged from 1.00 to 1.26 mg/100 g sample (Figure 2). No differences (p > 0.05) were identified between the highest carotenoid levels in L-36, Orange, L-184 and L-37. Variation in carotenoids content in different colored peppers has also been reported in bell peppers (Capsicum annum L.), where yellow varieties had the highest levels and green peppers the lowest [19].

Carotenoids are terpenoid compounds formed by the condensation of eight isoprene units. Specific chemical groups present at the chain terminus determine carotenoid chromophore properties and allow their classification into two families based on color: red and yellow/orange [23]. Capsicum fruit owe their intense coloring to carotenoid pigments [24], which coincides with the high carotenoid content observed here in all seven stud-
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3.3. Ascorbic Acid

Peppers have the highest ascorbic acid content of the vegetables; for example, consumption of 100 g fresh weight of peppers provides 100% - 200% of the recommended dietary allowance of ascorbic acid. Ascorbic acid content in the seven studied Habanero genotypes ranged from 187.24 to 281.73 mg/100 g sample (Figure 3). The Orange genotype had the highest level and the L-149 the lowest. Similar variation has been reported for bell peppers (*Capsicum annum* L.), in which red peppers had the highest ascorbic acid level, and green peppers the lowest [19]. The high ascorbic acid content of peppers is one of their primary nutritional qualities. Factors such as genotype, environment and fruit maturity affect levels of ascorbic acid and other nutritional compounds.

Ascorbic acid levels were high in all the studied genotypes, which coincides with reported increases in ascorbic acid content as pepper fruit matures; highest content is reached at full maturity [26]. Carotenoids, ascorbic acid, flavonoids, phenolic acids and other chemical constituent concentrations increase as peppers reach maturity [24]. In a study of several pepper cultivars, ascorbic acid was found to increase as color developed in some cultivars, but remained unchanged or diminished in others; black, purple, and white pepper fruit contained lower ascorbic acid levels compared to green, yellow, red, brown

mg/100 g dry weight for var. Ancho and var. Mulato. Differences in carotenoids content are probably due to the influence of genotype and maturity stages.

Carotenoids are known to play an important role in preventing oxidative damage, which is caused by free radicals in age-related diseases such as cancer. Most current carotenoids research is focused on their probable function as lipid antioxidants, which can protect against oxidation and other destructive processes mediated by singlet oxygen and free radicals, although more specific immune system effects are also receiving attention [21].

![Figure 2. Carotenoids content in *C. chinense* Jacq. Genotypes. Data are presented as means (n = 3). **Different superscript letters indicate statistical difference (p < 0.05).](image)

![Figure 3. Acid ascorbic content in *C. chinense* Jacq. Genotypes. Data are presented as means (n = 3). **Different superscript letters indicate statistical difference (p < 0.05).](image)
and orange peppers. High ascorbic acid levels have also been reported for pickled Serrano (584 mg ascorbic acid/100 g dry weight), fresh Serrano (1385 mg ascorbic acid/100 g dry weight), pickled Jalapeño (794 mg ascorbic acid/100 g dry weight), chipotle (694 mg ascorbic acid/100 g dry weight) and fresh Jalapeño peppers from three growing regions in the state of Chihuahua, Mexico: Meoqui (1696 mg ascorbic acid/100 g dry weight); Flores Magón (1185 mg ascorbic acid/100 g dry weight); and Ascención (2153 mg ascorbic acid/100 g dry weight) [27]. These authors concluded that the high variability in ascorbic acid content among Jalapeño samples may have been due to differences in growing conditions, maturity and particularly post harvest handling. In the study of the mature fruit of 63 C. chinense accessions by Antonious et al. [5], they reported that accessions PI-152452 (Brazil) and PI-360726 (Ecuador) contained the highest ascorbic acid concentrations (1.2 and 1.1 mg·g⁻¹ fresh fruit, respectively). These accessions were identified as potential candidates for mass production of antioxidants with health-promoting properties. Over all, C. chinense Jacq. is clearly an alternative source of phytochemicals such as ascorbic acid, a potent antioxidant.

3.4. Antioxidant Activity

3.4.1. ABTS Decolorization Assay

Trolox equivalent antioxidant capacity (TEAC) (mM/mg sample) differed (p < 0.05) between the genotypes, ranging from 1.55 to 3.23 mM/mg sample (Figure 4). The TEAC assay is based on scavenging of the ABTS⁺ radical cation by the antioxidants present in a sample. This radical is bluish-green in color, with maximum absorbance at 734 nm [28]. When antioxidant compounds are present in the reaction medium, they quench the free radical, causing loss of color and a consequent reduction in absorbance that corresponds quantitatively to antioxidant concentration. TEAC values have also been reported for pickled Serrano (3988 mmol Trolox equivalent/100 g sample), fresh Serrano (4787 mmol Trolox equivalent/100 g sample), pickled Jalapeño (2776 mmol Trolox equivalent/100 g sample), chipotle (2864 mmol Trolox equivalent/100 g sample) and fresh Jalapeño peppers from three growing regions in the state of Chihuahua, Mexico: Meoqui (5541 mmol Trolox equivalent/100 g sample); Flores Magón (3513 mmol Trolox equivalent/100 g sample); and Ascención (5211 mmol Trolox equivalent/100 g sample) [27]. In their study of Chiltepin and Habanero varieties, Rodriguez-Maturino et al. [11] suggested that differences in total antioxidant activity in peppers may be attributed to causes such as fertilization, fruit maturity and temperature. Their results showed no correlation between total phenolic compounds and antioxidant capacity in Chiltepin (11.10 mg of GAE/g of dry weight) and Habanero (8.20 mg of GAE/g of dry weight) fruit, which may indicate that antioxidant capacity could be affected by the presence of phytochemicals such as carotenoids.

Scavenging ability (%) also varied significantly (p < 0.05) between the genotypes, ranging from 94.98% to 44.46% (Figure 5). Scavenging ability values exhibited the same behavior as the TEAC values: the highest values were in the L-36 and Red genotypes, and the lowest in L-37. Higher antioxidant capacity in yellow, orange and red genotypes agrees with observations made by Guillerro et al. [24] that antioxidant capacity in C. annuum genotypes is comparable to that of commercial antioxidants, with the highest activity in orange, red and yellow genotypes.

3.4.2. β-Carotene Decolorization Assay

β-carotene decolorization is an effective antioxidant activity assay. In the absence of an antioxidant, β-carotene undergoes rapid discoloration since the free linoleic acid radical attacks the β-carotene molecule, which loses its double bonds and consequently its characteristic orange color. During the 120-min decolorization trial, antioxidant capacity decreased over time in the studied genotypes (Figure 6). Values ranged from 36% to 57% β-carotene decolorization.
tene bleeding during the first 30 minutes, with L-36 having the highest percentage and L-149 the lowest. After 120 minutes, values ranged from 11% to 32%. Overall, L-149 was the most stable antioxidant since its capacity decreased only 33% after 120 minutes.

Phenolic compounds, carotenoids and ascorbic acid can contribute to antioxidant activity in fruits, vegetables and grain products [29]. Free radical scavenging is one of the known mechanisms by which antioxidants inhibit the lipid oxidation caused by free radicals. The high polyphenols, carotenoids and ascorbic acid contents observed here in all the genotypes suggest that their observed antioxidant activities are largely due to their antioxidant content. The present results provide useful and important data for researchers to augment the antioxidant capacity and functional value of Habanero pepper in the food and nutraceutical industries.

4. Conclusion

Fruit from the seven studied Habanero chili genotypes is apparently good sources of antioxidants, including ascorbic acid, carotenoids and polyphenols. Habanero fruit can be consumed directly or used in extracts to increase nutritional value in different foods and diets. Development and consumption of Habanero pepper genotypes with high antioxidant activity may help in decreasing the incidence of certain types of diseases in humans.

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