Comparison of total phenol content and antioxidant activity of herbal infusions with added Stevia rebaudiana Bertoni

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Abstract
Antioxidant capacity (AC) was determined by the ABTS method and DPPH, and total phenol content (TPC) in dehydrated plant material, in infusions and in residues (plant material after preparing the infusion) of white, black, red, green, stevia, lemon grass and chamomile teas to which stevia leaves were added or not added; addition of processed stevia powder was also tested. Three independent experiments were set up: with dehydrated plant material, with infusions and with residues. For the case of dehydrated plant material, white tea had the highest TPC (10813.5 mg GAE/100g) and AC by the ABTS method (1183.3 µM TE/g) and DPPH method (1525.0 µM TE/g). In infusions, black tea had higher TPC (180.82 µg GAE/ml) and AC by the methods ABTS and DPPH (0.6114 and 2.5983 µM TE/ml, respectively). In residues, TPC was higher in white tea, while green tea had the highest AC values. AC of dehydrated plant material increased when stevia leaves were added, according to the DPPH and ABTS methods, but not in residues by ABTS. Addition of stevia leaves in infusions increased AC in white, lemon grass, chamomile and stevia teas by the methods ABTS and in spearmint, black, red, and green teas with the DPPH method.

Keywords: Camellia sinensis; Matricaria chamomilla; Cymbopogon citratus and Mentha spicata.

Practical Application: Addition of stevia leaves in most of the herbal infusions tested increased TPC and AC by DPPH.

1 Introduction

An infusion is a beverage obtained from parts (dry leaves, flowers and fruits) of the shoot of several herbs or aromatic plants. Worldwide, most of the infusions are prepared from two types of manufactured teas: black and green teas (Konieczynski et al., 2017), which are obtained from Camellia sinensis leaves, as are white and red teas. However, the growing season, geographic region and methods of fermentation create much variation in the composition and characteristics of the tea (Zhao et al., 2011) in addition to the variation associated with different stages of maturity of the leaves (Kosińska & Andlauer, 2014). Based on the method of fermentation, teas have been classified as non-fermented (white and green tea), semi-fermented (red tea) and fermented (black tea) (Camargo et al., 2016). Today, consumption of infusions has increased because of their beneficial properties, which are attributed to the presence of phenolic compounds (Cleverdon et al., 2018). These compounds have higher antioxidant capacity than fruits and vegetables and are even more potent than vitamin C, E and carotenoids (Wiseman et al., 1997). Because of these characteristics, teas are often used as remedies to cure diverse organic disorders. In this respect, Hernández-Saaavedra et al. (2013) report that herbal infusions can help reduce obesity, insulin intolerance and systemic inflammation. In addition, dehydrated stevia leaves can be added to infusions to improve their characteristics (Shevchenko et al., 2013). Because stevia has therapeutic values such as antioxidant, antimicrobials, antidiabetics, antihypertensive, cardiotonic; also; it has stevioside and rebaudioside, which give it an intense sweet flavor; it is good source of protein, crude fibre, minerals and amino acids (Joseph & George, 2019; Ahmed & Mukta, 2017). Processed stevia is up to 200-300 times sweeter than sucrose (Ghaheri et al., 2018), it is considered the best sugar substitute (Joseph & George, 2019). Although stevia extracts are being used only as sweeteners, aqueous extract of the dehydrated leaves have numerous functional properties (Ahmed & Mukta, 2017). It is believed that commercial infusions sweetened with stevia are a viable option that can contribute to the health of people who consume them. The aim of this study was to compare TPC and AC in dehydrated plant material, infusions and residues of white, black, red, green, spearmint, stevia, lemon grass and chamomile teas with or not addition stevia leaves and processed stevia.

2 Materials and methods
The study was conducted in the plant physiology laboratory of the CUSUR-UDG. Eight different plant materials were used: white, black, red and green tea (Camellia sinensis), spearmint (Mentha spicata), lemon grass (Cymbopogon citratus), chamomile (Matricaria chamomilla) and Stevia rebaudiana B. Three independent experiments were set up: 1) plant material before infusion, 2) infusion, and 3) plant material after infusion (considered...
residue). In each experiment, a two-factorial analysis was used in which one factor was the plant material (white, black, red and green tea, spearmint, lemon grass, chamomile and stevia); the other factor was the addition of 0.25 g dehydrated ground stevia leaves (ASL), commercial processed powdered stevia (PS) (Member’s Mark) and without stevia (WSL). In experiments 1 and 3, only ASL and WSL were considered, and in experiment 2 ASL, PS and WSL were studied. The commercial brand of chamomile, spearmint, green tea and lemon grass was Vit te® and black, white and red tea were Euro te®. Stevia leaves were produced in a greenhouse, dried in an oven at 50 °C (Binder series FD) and ground in a mill (KRUPS GX4100).

2.1 Preparation of infusions

To prepare the infusions, the technique described by Muñoz-Velázquez et al. (2012) was used with slight modifications. One gram of each of the eight plant materials were used (White, black, red and green teas, spearmint, lemon grass, chamomile and stevia); each infusion was replicated four times. The plant materials were sweetened by adding 1) one envelope of processed powdered stevia, 2) 0.25 g dried ground stevia leaves and 3) no sweetener. Filter paper was used to wrap the plant material and place it in the infuser; 250 ml water at 90 °C was poured into a cup and the infuser was introduced and left for 5 min. After this time the infuser was withdrawn and left to drain on a support for 5 min. The plant material was then taken out of the infusers, and both the infusion and the residues were left to cool to room temperature. Later, 45 ml of the infusion of each replication was stored at -20 °C until quantification. In the same way, the plant residues were stored at -20 °C until extraction and later quantification of total phenol content and antioxidant capacity.

2.2 Sample extraction

Sample extractions were carried out following Saura-Calixto, (1998), with some modifications. Of the eight dehydrated plant materials, ASL and WSL, 0.5 g were used, and 2 g of the moist plant material after use in the infusion (residue), which is equivalent to about 0.5 dehydrated plant material, was adjusted to real weight at calculation. Both dehydrated and moist (residues) plant material ASL and WSL were placed in a 50 ml plastic Corning tube with a stopper. Each material was replicated four times. The substrates were weighed on a scale (Santorius Te2145), and 4 ml acidified (pH 2) methanol was added. The tubes were covered with aluminum foil and shaken for 1 h in a Benchmark Orbi Shaker. After shaking, they were centrifuged at 1792 g for 15 min at room temperature with a SIGMA 3-16 KL centrifuge. The supernatant was deposited in 10 ml flask and covered with aluminum foil and kept 4 °C during the extraction process. Again 4 ml of 70% acetone was added to the plant material, it was shaken and centrifuged as in the previous step. The supernatant was deposited in the same flask where the supernatant of the first step was deposited, and 10 mL were gauged with acidified methanol (pH 2). Four aliquots of 1.5 mL of this extraction were placed in Eppendorf tubes and stored at -20 °C until quantification.

2.3 Quantification of total phenols

Total phenols content were quantified at a micro-scale, as proposed by Arnous et al. (2002), with modifications. A calibration curve was constructed from a standard solution of gallic acid (GA) using concentrations of 0-500 µg/mL. Ten µL of the extracts were placed together with 790 µL distilled water, 150 µL sodium bicarbonate at 20% and 50 µL of Folin reagent. The aliquots were left to stand for 1 h protected from light. Each sample was loaded in triplicate in plates and absorbance was read at 760 nm in a Thermo Scientific Multiskan Go 1510 spectrophotometer. The results were expressed in mg gallic acid per 100g of extract (mg GA/100g dry base) for dehydrated plant material and residues and in µg GA/mL for infusions.

2.4 Determination of antioxidants by DPPH and ABTS

Antioxidant capacity was determined by the ABTS method, following Prior et al. (2005). A trolox calibration curve was used (0 to 0.8 mM). An ABTS mother solution (7 mM) and potassium persulfate (2.4 mM) were placed in a precipitation beaker with 10 mL of distilled water and immediately covered to protect it from light. It was maintained at room temperature for 16 h in a magnetic stirrer (Dagger Hotplate/Stirrer). Then, was adjusted with 100% methanol to an absorbance of 0.7. Each sample was loaded in triplicate, depositing 20 µL of the sample, plus 200 µL ABTS. Absorbance was read at 734 nm in the Multiskan Go up to 30 min of reaction. The results in AC were expressed in µM Trolox equivalents (TE)/g, dry base for dehydrated plant material and residues and in mM TE/mL for infusions.

A DPPH solution (150µM) was prepared. A trolox calibration curve was used (0 to 0.8mM), and each sample was loaded in triplicate. Twenty µL of sample (blank and the curve) were deposited plus 200 µL DPPH. Absorbance was read at 520 nm in the Multiskan Go, reaction of up to 30 min.

An analysis of variance was performed and afterward a Tukey comparison of means test (α=0.05) with the software Statistical Analysis System version 9.1.3 (SAS Institute, 2007).

3 Results

3.1 Dehydrated plant material

Total phenol content in dehydrated plant material

Average values of total phenol contents in dehydrated plant material showed that white tea had the highest TPC with 10813.5 mg GA/100g dry base, followed by black tea, green tea and stevia with 6759.5, 5539.9 and 4208.4 mg GA/100g, respectively. The rest of the plant materials had lower values and were statistically equal (Figure 1a). Independent analysis of each of the materials revealed that white tea WSL was statistically superior with a value of 11750 mg GA/100g, followed by white tea ASL with 9877 mg GA/100g. Stevia, black tea and green tea had intermediate values, and the plant materials ASL and WSL had lowest values and were statistically equal. No differences were found in averages between plant materials without stevia leaves and plant materials with added stevia leaves (Table 1).
Figure 1. Total Phenol Content (TPC), Antioxidant capacity (AC) by the ABTS and DPPH methods of dehydrated plant material, infusions and residue.

Table 1. Total Phenol Content (TPC, mg GA/100g), Antioxidant capacity (AC, µmol TE/g) by the method ABTS and DPPH of dehydrated plant material and residues without stevia leaf (WSL) and added with stevia leaves (ASL).

<table>
<thead>
<tr>
<th>Dehydrated plant material</th>
<th>TPC</th>
<th>ABTS</th>
<th>DPPH</th>
<th>TPC</th>
<th>ABTS</th>
<th>DPPH</th>
</tr>
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<tr>
<td>White tea WSL</td>
<td>11750</td>
<td>1014</td>
<td>1653.5</td>
<td>13211</td>
<td>971.6</td>
<td>513.7</td>
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<td>White tea ASL</td>
<td>9877</td>
<td>1351.6</td>
<td>1396.5</td>
<td>10583</td>
<td>620.6</td>
<td>168.6</td>
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<tr>
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<td>180.7</td>
<td>99.3</td>
<td>2264</td>
<td>134.0</td>
<td>78.0</td>
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<tr>
<td>Spearmint ASL</td>
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<td>203.9</td>
<td>407.7</td>
<td>3354</td>
<td>171.5</td>
<td>132.0</td>
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<tr>
<td>Lemon G. WSL</td>
<td>959</td>
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<td>57.3</td>
<td>1464</td>
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<td>7.3</td>
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<tr>
<td>Lemon G. ASL</td>
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<td>154.8</td>
<td>95.1</td>
<td>2338</td>
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<td>1566</td>
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<tr>
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<td>85.1</td>
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<td>2618</td>
<td>128.1</td>
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<td>741.2</td>
<td>6719</td>
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<td>2796</td>
<td>204.2</td>
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<td>144.2</td>
<td>185.6</td>
<td>3443</td>
<td>199.7</td>
<td>104.5</td>
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<td>Stevia WSL</td>
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<td>628.0</td>
<td>5161</td>
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<td>640.4</td>
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<tr>
<td>Stevia ASL</td>
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<td>581.1</td>
<td>726.2</td>
<td>5416</td>
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<td>1466.5</td>
<td>6155</td>
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<td>515.16</td>
<td>587.7</td>
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<td>636.9a</td>
<td>5089a</td>
<td>3664b</td>
<td>459.2a</td>
</tr>
</tbody>
</table>

Means with the same letter in column are not significantly different (Tukey, p<0.05).
Dehydrated plant material antioxidant capacity by the ABTS method

Average values obtained by the ABTS method for AC of dehydrated plant materials show that white and black tea are statistically equal and superior (1183.3 and 1179.4 µM TE/g, respectively), followed by green tea with 1053.2 µM TE/g. The lowest values were found in chamomile (Figure 1b). Independent analysis of each plant material revealed that white tea ASL had the highest AC with 1351.6 µM TE/g, followed by black tea ASL with 1323.8 µM TE/g. In contrast, chamomile WSL had the statistically lowest values with 45.7 µM TE/g. Adding stevia leaves to any of the dehydrated plant material increased average AC values, from 515.16 µM TE/g to 612.3 µM TE/g (Table 1).

Antioxidant capacity of dehydrated plant material by the DPPH method

By the DPPH method, the average AC values of plant material was statistically equal and superior to white tea with a value of 1525.0 µM TE/g, followed by green tea with 1474.7 µM TE/g, while again chamomile had the lowest values with 46.0 µM TE/g (Figure 1c). Independent analysis of each dehydrated plant material showed that, white WSL was that with the highest antioxidant capacity with 1653.5 µM TE/g, followed by green tea ASL with a value of 1483.1 µM TE/g; chamomile WSL had the statistically lowest value, 32.2 µM TE/g. Analysis of average values of the group of plant materials WSL compared with plant materials ASL showed that adding stevia leaves resulted in statistically higher values of antioxidant capacity, from 587.7 to 636.9 µM TE/g (Table 1).

3.2 Infusions

Total phenol content in infusions.

Total phenol content was statistically highest in the infusion of black tea, 180.82 µg GAE/ml, followed by stevia, red tea, spearmint, white tea and green tea with values of 94.61, 80.20, 66.69, 59.33 and 58.48 µg GAE/ml, respectively. The infusions that showed the lowest values were chamomile and lemon grass (Figure 1d). Independent analysis of each of the infusions showed that black tea ASL was statistically superior to 279.17 µg GAE/ml, followed by black tea WSL and PS with values of 142.17 and 121.13, µg GAE/ml, respectively. The infusion of stevia, red tea, green tea, spearmint and white tea with the different sweeteners had intermediate values, and the infusions with the lowest values were of lemon grass and chamomile. TPC averages was statistically higher in the infusions with added stevia leaves with a value of 90.28, µg GAE/ml than the infusions with processed stevia and without stevia leaves. This means that addition dehydrated stevia leaves increases AC, while adding processed stevia does not contribute to incrementing AC of the infusions (Table 2).
Antioxidant capacity of infusions by the ABTS method

The DPPH method for average AC of WSL, ASL and PS of each infusion resulted in that black tea and stevia were statistically equal and superior to the rest of the teas with values of 2.6053 and 2.5983 μM TE/ml, respectively, followed by red tea, white tea, green tea, spearmint, chamomile and lemon grass with values of 1.9351, 1.7611, 1.6290, 1.3708, 0.5422 and 0.3723 μM TE/ml respectively (Figure 1f). Analyzed independently, green tea ASL obtained the highest statistical value, 2.8790 μM TE/ml, followed by stevia ASL and black tea ASL, with values of 2.8382 and 2.8296 μM TE/ml, respectively. All the infusions with added dehydrated stevia leaves had higher antioxidant capacity than infusions WSL or PS, except the infusion of spearmint. In general, both ABTS and DPPH methods showed the highest TPC and AC in black tea and the infusions with the lowest values were chamomile and lemon grass. The DPPH method, comparing only averages of the infusions WSL, PS and ASL, showed that AC had the same trend as TPC in that the infusions with added stevia leaves showed higher AC than those to which stevia leaves were not added or those with added processed stevia (Table 2).

3.3 Residues
Total phenol content in residues

Total phenol contents was statistically superior in white tea with 11897 mg GAE/100g, followed by black and green teas with values of 7663 and 6198, mg GAE/100g respectively; the rest of the residues were statistically lower than those mentioned (Figure 1g). The TPC of the residues of white tea WSL, 13211 mg GAE/100g, was statistically higher, followed by white tea ASL, 10583 mg GAE/100g. In contrast, lemon grass tea had the lowest value (1464 mg GAE/100g). No differences were found when we compared the average of all the residues without stevia leaves with all the residues with added stevia leaves (Table 1).

Antioxidant capacity of residues by the DPPH method

Green tea had the statistically highest AC by ABTS with 916.3μM TE/g, followed by black tea and white tea, with values of 816.2 and 796.1 μM TE/g, respectively, while chamomile again had the lowest values, 96.7 μM TE/g (Figure 1h). Individually, black tea residues WSL had the highest AC with 1070.7 μM TE/g, followed by green tea WSL and white tea WSL with values of 1031.3 and 971.6 μM TE/g, respectively. The lowest values, which were statistically equal, were those of lemon grass WSL and chamomile WSL, with values of 64.3 and 70.3 μM TE/g respectively. The comparison of all the residues without stevia leaves with all the plant materials with added stevia leaves showed that those to which stevia leaves were not added were statistically superior with a value of 511.5μM TE/g (Table 1).

4 Discussion
4.1 Plant material

Dehydrated plant tissue of white tea had the highest average values of both total phenol content (TPC) and antioxidant capacity (AC) by the ABTS and DPPH methods, even though green, black, red and white teas are from the same plant (Camellia sinensis). The results obtained for TPC are attributable to the process and maturity of the teas. White is from young leaf buds, it is the least processed and is not fermented. Green tea is from fresh mature leaves, like black tea, but the difference is that black tea is fermented (Kosińska & Andlauer, 2014), while red tea undergoes two fermentations. In this respect, Kim et al., (2011) reports that the TPC and AC were higher in leaves of less processed teas, suggesting that fermentation decreases AC and can result in low potential health benefits. Zhao et al. (2011) also reported higher content of phenolic compounds in powdered white tea (74.7346 mg/g) than in green tea (27.9767 mg/g) and green Pu-erh (46.5586 mg/g). Another explanation of why white tea has higher TPC and AC is that shoots and young leaves have higher TPC and AC than mature leaves (Chan et al., 2007).

In our study, we found that dehydrated stevia had average values of 38.56 mg GAE/100g, which were lower than those reported by González et al. (2014), who studied the stevia...
TPC was not different when stevia leaves were added. However, in AC, adding stevia leaves increased by both DPPH and ABTS. This agrees with other authors who report that stevia has considerable AC (Periche et al., 2014) that increases AC of other plant materials when mixed (Benítez & Pérez, 2016; Shevchenko et al., 2013).

### 4.2 Infusions

Worldwide, the infusions are an important source of phenolic compounds, and represent a class of bioactive molecules that are closely associated with a variety of health benefits (Cleverdon et al., 2018). Our results for TPC in infusions coincide with Gorjanović et al. (2012), who also report that black tea is that with the highest TPC. In contrast, Atoui et al. (2005), (Deetae et al., 2012), Konieczynski et al. (2017) and (Moraes-de-Souza et al., 2008) report higher TPC in green tea than in black tea. Our results are numerically different and lower than those reported by Muñoz-Velázquez et al. (2012). However, the trend is the same since they report values for chamomile of 61.84 to 69.28, lemon grass 69.91 to 75.66, spearmint 150.80 to 231.85, arnica 173.31, boldo 312.71 and green tea 1628.05 µg GAE/ml. Likewise, Atoui et al. (2005) report lower TPC for chamomile and high for black tea and Chinese green tea (106, 847 and 1216 mg GAE/cup, respectively). Moraes-de-Souza et al. (2008) also report high TPC for green and black teas than for chamomile, coinciding with our study. On the other hand, Camargo et al. (2016) report higher TPC in infusions of white tea (85.36 µg GAE/mL) than in green, red and black teas, with 76, 45.47 and 43.34 µg GAE/mL, respectively. Moraes-de-Souza et al. (2008) and Gorjanović et al. (2012) report that TPC can be different in different studies because of the way the teas are prepared (processing of the plant, concentration, time and temperature of the infusion), herb (species, part used, development stage), characteristics of production (soil, climate, stress) and methods of analysis, among other factors.

The stevia infusion had a high TPC in our study, surpassed only by black tea. Other authors, such as Shukla et al. (2012) have also reported high TPC in aqueous and ethanol extracts of stevia, obtaining 56.74 and 61.50 mg GA/g, respectively. Likewise, Deetae et al. (2012) reports that stevia is outstanding in TPC, like black and green teas, among 18 infusions evaluated. In this respect, Periche et al. (2014) mentions that because stevia is a natural non-caloric sweetener with beneficial properties, considerable antioxidants and amino acids, its consumption as an infusion is increasing. We found that the infusion of black tea was that with the highest AC, according to both ABTS and DPPH. These results agree with Camargo et al. (2016), who also found the highest AC in black tea when compared with red, green and white teas using the same methodologies. In our study, stevia obtained higher average values than black tea and even had higher AC by the DPPH method than green tea. This agrees with Deetae et al. (2012), who conclude that stevia has properties similar to black and green teas. In this respect, Shukla et al. (2012) concludes that stevia is able to eliminate free radicals. Also, Periche et al. (2014) report that it is a source of natural antioxidants that can benefit health.

The results of AC obtained in our study by the DPPH method in chamomile and spearmint infusions are similar to those reported by Muñoz-Velázquez et al. (2012), who report that the AC of chamomile is 0.52-0.60, 0.63-0.70 for lemon grass, spearmint 1.28-2.04, arnica 1.47, boldo 3.61, and green tea 4.90 µM TE/ml, although, the values we obtained for green tea were lower. However, in both studies the same tendency was found: chamomile and lemon grass have the lowest values, spearmint intermediate values and green tea the highest values. Moreover, Atoui et al. (2005) report low levels for chamomile (0.17 TE) and high levels for black and green teas (0.54 and 0.57 TE, respectively). Moraes-de-Souza et al. (2008), however, report that infusions of chamomile, green tea, and black tea have a high capacity for oxidative inhibition.

The high values found in TPC and AC by the DPPH method when stevia leaves were incorporated into the infusions agree with Benítez & Pérez (2016), who studied the variation in antioxidant capacity and phenolic compounds of aromatic herbs during the process to obtain a filtrate with stevioside as a natural sweetener. Benítez & Pérez (2016) and Shevchenko et al. (2013) found that aromatic herbs and Camelia sinensis respectively had increased antioxidant capacity when stevia leaves were added. It is important to highlight that the addition of processed powdered stevia does not increase antioxidant capacity or total phenol contents, thus serving only as a sweetener. In contrast, dehydrated stevia leaves, besides sweetening, also increases TPC and AC as evaluated by the DPPH method.

### 4.3 Residues

Although it has been demonstrated that most of the phenolic compounds (80-90%) are released in the first five minutes, the release time varies among teas. For example, most of the phenolic compounds in green tea are released in the first 2-3 min (Cleverdon et al., 2018). However, our results show clearly that, even after infusion, the residues still contain a large quantity of TPC and AC, and the availability of TPC increases in all of the evaluated materials, making it feasible to reuse the residues for another infusion.

### 5 Conclusions

The addition dehydrated stevia leaves increases AC in dry material and infusions of white tea, lemon grass, chamomile, stevia by both methods, and of spearmint, black tea, red tea, and green tea by DPPH method. But, the addition of commercial processed powdered stevia does not increases AC or TPC in infusions. The infusion process were not enough to deplete the initial contents of AC and TPC once the dry material was used.

### References

