Improvement of nutritional and physicochemical proprieties of milk chocolates enriched with kale (Brassica oleracea var. acephala) and grape (Vitis vinifera)
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Abstract
Consumption of functional food has been growing recently, and products with healthier ingredients can influence the purchasing decisions. Studies have been shown that the development of chocolates with addition of functional ingredients can improve its nutritional properties. Considering it, the present study demonstrated that the addition of lyophilized kale and grape can modify physical, nutritional and physicochemical characteristics of the chocolates. The enriched milk chocolates showed higher values of dietary fibers and mineral content, which improved nutritional quality of the product when compared to plain milk chocolates. The results of the HPLC-DAD suggested the transference of phenolic compounds of kale and grape to the final enriched chocolates. The sensorial analysis indicated that differences among the formulated chocolates were indeed perceived and were well accepted. Thus, the addition of kale and grape in chocolates may be an alternative to improve the nutritional characteristics and the polyphenolic compound content in these chocolates.

Keywords: chocolate; fruits; vegetables; antioxidants.
Practical Application: The incorporation of fruits and vegetables in chocolate formulation has nutritional importance. The present study showed that addition of lyophilized kale and grape can modify the physical, nutritional and physical-chemical characteristics of the chocolates.

1 Introduction
The development of healthy formulations is a trend in the food market, according to “Brazil food trends 2020” (Madi et al., 2010). It involves products with lower caloric value, sugar content and trans fat, and more functional properties. Currently, consumers are looking for products which contribute to a healthy diet, with higher mineral and nutrient content and reduced values of fat and cholesterol.

Since past years, antioxidants have been in–depth explored due to their association with health benefits. It is known that the consumption of high amounts of different bioactive compounds is closely related to a lower risk of health disorders. The natural source of antioxidants includes a large number of vegetable and fruits (Oroian & Escrive, 2015). Moreover, antioxidant activity is commonly correlated with functional properties as an alternative to reduce or avoid non-degenerative diseases and precarious aging (Oroian & Escrive, 2015; Watson et al., 2013). For example, polyphenols are antioxidants that have promising chemical structures to prevent skin aging (Watson et al., 2013). Ali, Ismail, & Kersten (2014) also related the presence of these compounds with prevention of metabolic diseases and obesity.

Cacao has been extensively studied for its intake health benefit as the improvement of vascular function (Pereira et al., 2014) cardiovascular prevention (Grassi et al., 2015) and cognitive functions (Sokolov et al., 2013). In 1999, Vinson, Proch, & Zubik (Vinson et al., 1999) related the presence of the polyphenol content of cocoa to health benefits of chocolate due to differences in the polyphenol content in milk chocolate (52.2 ± 20.4 µmol/g), when compared to dark chocolate (126.0 ± 17.4 µmol/g), since there is a lower level of cocoa in milk chocolate. Pimentel et al. (2010) showed that the polyphenols in dark chocolate, with higher content of cocoa, are 21% higher than the dark chocolate with less cocoa content, which are 36% higher than milk chocolate and 55% higher than white chocolate.

Fruits and vegetables are also a very good sources of bioactive compounds, which can promote antioxidant actions in the body (Scalbert et al., 2005; Slavin & Lloyd, 2012). It is known that the its consumption is correlated to reduced risk of breast cancer (Eliassen et al., 2012); prevent obesity and diabetes (Mozaffarian, 2016) and the increase of HDL cholesterol (Sarriá et al., 2015). Moreover, dried fruits are also known to provide bioactive compounds to maintain human health balance (Chang et al., 2016).

Kale is an important vegetable of the Mediterranean diet and it is known for its antimicrobial proprieties and capacity for disease prevention. In its composition, nine phenolic compounds with high antioxidant capacity were already described (Ayaz et al., 2008; Murador et al., 2016). Murador et al. (2016) founded high carotenoid content in raw kale, as well as high total phenolic content (29.16 mg GAE/100g raw sample). Mageney et al. (2017) described the phenolic compounds on Brassica oleracea and identify major components as flavonol kampoferol and quecetin...
Also with a high phenolic content, grape consumption is recommended to prevent non-degenerative diseases and precocious aging. Grapes are sources of antioxidants since it are mainly composed by anthocyanins, flavonoids and resveratrol (Xia et al., 2010).

Thus, this study aimed to increase the nutritional value of milk chocolate by adding freeze-dried kale and grape into the formula.

2 Materials and methods

2.1 Lyophilization

Kale samples were purchased in a local Market in São Paulo city, Brazil. The leaves were cleaned, sanitized and chopped for storage in ultrafreezer for 24 hours at -88 °C.

Grapes also were purchased in a local market in São Paulo, Brasil. The variety of grape was Summer Royal without seed. Entire grapes were cleaned, sanitized and chopped for storage in ultrafreezer for 24 hours at -88 °C.

For the lyophilization process the Edward’s lyophilizer (Brazil) (Model: OS 3728) was used for 5 days. After that, kale leaves were crushed for 3 minutes in a termomixer Vorwerk (Brazil). The kale powder was stored in a desiccator to avoid humidity. Lyophilized grapes were cut for reducing particle size and also stored in a desiccator.

2.2 Production of chocolates

Chocolates were produced in a Ball Mill Machine WA-FA20 (Mazzetti, Italy). The process time was 2 hours at 45 °C. Chocolate formulation included: cocoa liquor, cocoa butter, soy lecithin, sugar, vanilla, milk powder and lyophilized kale or grape. Process and formulation are described in Figure 1.

Tempering was carried out in a temper (Universal, Brazil) and the samples were stored in an incubator with controlled humidity (60%) and temperature (20 °C).

2.3 Physicochemical evaluation

The physicochemical evaluation of the chocolates was based on (AOAC, 2005) for ash, moisture, protein and fat content. All analysis were carried out in triplicate.

Water activity was measure in equipment Novasina, model Labmaster (Novasina, Switzerland) in triplicate.

2.4 Texture, rheology and particle size

Particle size of chocolates was analyzed by using the digital micrometor Digimatic number 293 (Mitutoyo, USA), with five repetitions.

Rheology analysis was conducted on a rotacional rheometer (Rheotest 3.1, German) with thermostatic bath at 40 °C and probe S1. A ramp with shear rate of 200 s⁻¹ and time of 120 s were used. All the chocolate samples were analyzed in triplicate.

The texture was analyzed by using a texturerometer TA-XT2 (Stable Micro Systems, United Kingdom). Fracture properties were tested with a probe HDP/3PB at 25°, in triplicate. Parameters were: Speed pre-test and post-test: 2.0 mm/s; distance: 10 mm; charge cell: 25 kg; Trigger Force: 0.05 N; strength in compression – return to start. The data was collected by using the program “Texture Expert Exceed” – version 2.64 (Stable Micro Systems, United Kingdom), Sample: Chocolate bar with 9.0 cm x 2.5 cm x 1.3 cm (Lannes, 2008).

3 Antiradical activity

3.1 Extraction

The extraction method for all samples (chocolates, kale and grape) was conducted as suggested by (Genovese & Lannes, 2009). A sample of (0.5 g) was added to 20 mL of methanol 70%, using an Ultraturrax mixer (Marconi, Brazil) for one minute in speed 4, immersed in an ice bath. The extract was filtered through a paper filter and the resulting solution was stored in amber flask.

![Figure 1](image-url)

Figure 1. Schematic methodology of procedure on panel 1 and formulation on Panel 2, being A) Plain chocolate; B) chocolate with lyophilized kale C) chocolate with lyophilized grape.
3.2 Scavenging activity

Scavenging activity against the DPPH and ABTS radicals were conducted as suggested by Oliveira et al. (2014)

Stock DPPH was prepared in methanol and its final concentration was determined spectrophotometrically (ε515 nm = 1.25 x 104 L mol⁻¹ cm⁻¹). DPPH• stock solution was added to ethanol in a 10 mM absorbance quartz cell to final concentration of 80.0 µmol L⁻¹. The assay starts by adding the antiradical stock solution to a final volume of 3.0 mL. The absorbance at 515 nm was monitored for 30 minutes.

An ABTS stock aqueous solution was prepared and stored at 4 °C. For the oxidation of the ABTS 5 mL of the solution was used with 88 µL of potassium persulfate with concentration of 1.4 x 10⁻⁵ mol L⁻¹, which reacted for 16 hours, protected from the light. Resulting concentration was measured with a UV visible Spectrophotometer at 734 nm. ABTS•⁺ stock solution added to ethanol in a 10 mM absorbance quartz cell to a final concentration of 53.0 µmol L⁻¹. Reaction starts with the addition of antiradical stock solution to final volum of 3.0 mL. The absorbance at 734 nm was monitored for 30 minutes (Oliveira et al., 2014).

Trolox® (6-hydroxy-2,5,7,8-tetramethylcroman-2-carboxilic acid) (Sigma-Aldrich) stock solution was prepared with NaOH (0.01 mol L⁻¹) and ethanol.

All the reactions were monitored with a UV visible Spectrophotometer (Varian Cary 50 bio, USA) for 30 minutes and protected from the light.

The antiradical capacity was expressed in % trolox, which indicates the correlation between linear concentrations of Trolox and other antiradical compounds. Trolox is known as an analogous of vitamin E and is considered a proportion of the antiradical capacity for scavenging against the free radical.

3.3 HPLC-DAD

The extract obtained was rota-evaporated using a BUCHI equipment (Switzerland) with temperature and control rotation, and dissolved in methanol at a proportion of 1 mg/1 mL, and then processed in a SPE cartridge (Octadecyl C18/18%). Analyses were made in duplicate in HPLC-DAD (Agilent Technology model 1200 Infinity, USA) in an 5µm C₁₈ Ascentis column (250x4.6mm) (Supelco analytical, USA). The flow rate was of 1 mL/min and the volume of injection was 40 µL. The wave-length was 260, 280, 320 and 360 nm.

For analysis of phenolic compounds the mobile phase used was: eluent A) acidified water with 2% of acetic acid, and B) Acetic acid 0.5%: acetonitrile at a proportion of 50:50 (v/v). The gradient was: 0-20 min 10% of eluent B, 20-40 min 24% of eluent B, 40-60 min 30% of eluent B, 60-65 min 55% of eluent B, 65-70 min 70% of eluent B, 70-75 min 75% of eluent B and 75-80 100% of eluent B (Silva & Queiroz, 2016).

3.4 Polyphenolic compounds

The Folin-Ciocauteau method was used according to Singleton et al. (1999), with some modifications.

The standard curve was built with solutions of gallic acid (50,100,150 and 250 mg/L). For the reaction 20 µL of the extraction matter was used with 200 µL of Folin-Ciocauteau reagent and mixed; after 5 minutes 750 µL of sodium carbonate 20% was added and the volume was adjusted to 5 mL. The solution remained protected from the light.

The measurements were carried out using a UV visible Spectrophotometer (Spectrum meter Brazil) at 760 nm and repeated in triplicate.

3.5 Sensorial evaluation

The global acceptance test (texture, taste, appearance) using the hedonic scale with 9 points and purchase intention with 5 points was applied on 90 untrained panelists of the University of São Paulo, after approval of University Ethics Committee (n° 1.393.285).

Sensorial analyze was done in Sensorial Analysis Laboratory of Department of Biochemical-pharmaceutical Technology on Faculty of Pharmaceutical Science, in University of São Paulo.

Chocolates were prepared 24 hours before and storage at 20 °C with 60% of humidity, 5 g of samples were served in plastic plates, in monadic way, and panelists were advised to drink water between samples to avoid interferences.

3.6 Data analyses

All data analyses were carried out using with the software Statistica version 11 (StatSoft, EUA) for the Analysis of Variance (ANOVA) with a level of significance of p<0.05. The Tuckey’s method were done to check for significant differences among the groups (α=95%).

4 Results

4.1 Physicochemical analyses

Considering the main constituents of chocolates (Afoakwa, 2010), the nutritional composition of plain milk chocolates and chocolates with addition of kale and grape were evaluated and the results are presented in Table 1.

Table 1 shows that the addition of kale and grapes does not modify significantly (p<0.05) the moisture content and water activity. Also, the evaluation of the sugar content shows that there is no difference between plain chocolate and the chocolate with grape, even when reducing the sugar content in the formulation, which indicates that the sugar in the formulation can be replaced by lyophilized fruits such as grapes.

On the other hand, the addition of kale and grape in chocolates modify some of the nutritional contents, comparing to plain milk chocolate. The addition of kale increases significantly (p<0.05) soluble fibers, while the addition of grape raised insoluble fiber levels. These results corroborate with the review published
by Zhu et al. (2015) and with the USDA, which showed high levels of fibers in both grape and kale, respectively. Moreover, the enrichment of chocolate formulations with kale or grape have impacted the ash content (Table 1), causing a significant increase (p<0.05) of this value.

4.2 Rheology, texture and particle size

The addition of natural products on the chocolate could modify quality characteristics, like rheology, which may become it undesirable for consumption. The results indicated that the addition of kale and grape modify this parameter without impacting the sensory quality of this products.

The results of particle size, texture and rheological parameters can be found in Figure 1.

Chocolate with added grape shows an increase in particle size. Lyophilized grapes have high levels of sugar content which make them hygroscopic. For avoid humidity in chocolate samples, the grape were added after the process of production of chocolate. Thereafter, grapes did not pass throw refining stage, which can explain this increase in particle size. Despite desirable particle size of chocolate are between 20-30 µm (Afoakwa, 2010), sensorial analysis shows that the grape particles are desirable for some consumers.

The results show that there is a statistic difference (p<0.05) among all formulations related to particle size. The particle size is directly related to texture and yield values, and inversely related to Casson viscosity. Texture results indicated that the hardness of chocolate is modify with addition of grapes, with a statistic difference (p<0.05).

Rheology analysis reveal that plain milk chocolate shows the highest (p<0.05). Casson Viscosity, as well as the chocolate with added grape is the one with highest (p<0.05) yield value.

4.3 Sensory Evaluation

Sensorial analysis evaluated four parameters of all chocolate samples. The results for taste, texture and aspect are shown on Figure 2 and purchase intention on Figures 3 and 4.

Sensorial results showed that the difference in formulations affects the perception of consumers. The difference among chocolates were perceived by 92% of the panelists, which indicated that there were significant differences in sugar content, texture (of the chocolate with grape) and taste (among all chocolates). The chocolate with grape showed a higher variance in texture; 47.4% of the panelists indicated approval for "crunchy texture", while 52.6% of panelists disliked the product for the same reason.

In spite of these differences, all the chocolates were well accepted by consumers. This study showed a purchase intention of 61.1% for chocolate with kale, 66.6% for plain milk chocolate and 64.2% for chocolate with grape.

4.4 Antiradical Profile

The antiradical activity was evaluated by DPPH and ABTS method and the Folin Ciocântea method was used to found the total phenolic content. The antiradical capacity is shown in Table 2 and Table 3.

The results of lyophilized kale and lyophilized grape show a high antiradical activity against DPPH radical, which corroborate with phenolic content analysis. Cocoa also show high antiradical activity and phenolic content. This results indicated that the raw materials were promising for increase antiradical capacity in chocolate.

The analysis of antiradical activity against the DPPH radical show that plain chocolate and that with added grape are statistically different (p<0.05), while the chocolate with kale shows no difference in comparison with the other two samples. This indicates that the scavenging activity against DPPH was not increased with the addition of kale and at the same time it decreased with the addition of grape. Nevertheless, the antiradical capacity is comparable to beverages and foods known as having high values of antiradical capacity, such as mango (211 mg trolox/100 g) and wine (196.4 mg Trolox/100 g) (Floegel et al., 2011; Vijaya Kumar Reddy et al., 2010).

The addition of kale and grape did not interfere with respect to the antiradical capacity of milk chocolate, since no significant difference (p>0.05) among the samples was observed. Despite this, all the chocolates are considered as having high antiradical.

Table 1. Nutritional composition of A) Plain Milk Chocolate, B) Milk chocolate with kale, C) Milk Chocolate with Grape. Different letters means statistic difference (p<0.05).

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calories (kcal)</td>
<td>564.68</td>
<td>554.68</td>
<td>555.7</td>
</tr>
<tr>
<td>Moisture (g/100 g)</td>
<td>1.73±0.48a</td>
<td>2.07±0.29a</td>
<td>2.27±0.25a</td>
</tr>
<tr>
<td>Water Activity (Aw)</td>
<td>0.52±0.013a</td>
<td>0.50±0.01a</td>
<td>0.52±0.032a</td>
</tr>
<tr>
<td>Ash (g/100 g)</td>
<td>2.53±0.14a</td>
<td>2.68±0.054a</td>
<td>2.57±0.027b</td>
</tr>
<tr>
<td>Protein (g/100 g)</td>
<td>8.51±0.36a</td>
<td>8.52±0.37a</td>
<td>8.13±0.46a</td>
</tr>
<tr>
<td>Fat (g/100 g)</td>
<td>36.26±0.39a</td>
<td>35.06±0.38b</td>
<td>35.50±0.87a</td>
</tr>
<tr>
<td>Dietary Fibers Total (g/100 g)</td>
<td>7.89</td>
<td>8.59</td>
<td>8.05</td>
</tr>
<tr>
<td>Soluble Fibers (g/100 g)</td>
<td>3.18</td>
<td>4.18</td>
<td>3.16</td>
</tr>
<tr>
<td>Insoluble fibres (g/100 g)</td>
<td>4.71</td>
<td>4.4</td>
<td>4.88</td>
</tr>
<tr>
<td>Total sugar (g/100 g)</td>
<td>21.00±0.016</td>
<td>17.00±0.0083</td>
<td>21.00±0.015</td>
</tr>
<tr>
<td>Reduced sugars</td>
<td>7.4±0.0032</td>
<td>7.82±0.029</td>
<td>7.24±0.039</td>
</tr>
<tr>
<td>Non-reduced sugars (g/100 g)</td>
<td>13</td>
<td>9.21</td>
<td>13.45</td>
</tr>
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</table>
Figure 2. Results of (1) particle size, (2) Texture, (3) Casson Viscosity and (4) Yield Value for A) Milk chocolate B) Milk chocolate with added kale C) Milk chocolate with added grape. Different letters means statistic difference (p<0.05).

Figure 3. Sensorial evaluation results for 3 parameters: Taste, texture and aspect of A) Plain milk chocolate, B) Chocolate added kale C) Chocolate added grape. Values of means±SD for all samples.
Chocolate with *B. oleracea* and *V. vinifera*

The statistical analysis of quantification of polyphenolic compounds shows that there was no significance difference (p>0.05) between the milk chocolate and the chocolate with kale. However, when comparing the grape chocolate with the control, the study shows that there has been an increase of 50.44% in its value. This shows that the addition of kale did not modify the polyphenolic content and that the addition of grape did increase this content. However, the polyphenolic content of chocolate with kale and of plain chocolate are still high, being comparable to oranges (70 mg GAE/g) (Vijaya Kumar Reddy et al., 2010).

To confirm this hypothesis, the profile of phenolic compounds was carried out by HPLC-DAD (Figure 5).

The chromatogram analysis of milk chocolate with added kale, plain milk chocolate and lyophilized kale, shows that some compounds in lyophilized kale were transferred to the chocolate with added kale. The chromatograms of milk chocolate with added grape, plain milk chocolate and lyophilized grape show also that some of the grape peaks are transferred to the chocolate with added grape. Most of these compounds were previously identified in kale and grape which indicates that some of these compounds were transferred to the chocolates.
5 Discussion

The addition of lyophilized products aimed to improve the functional properties of chocolate and we showed here that it was able to modify some physicochemical characteristics of milk chocolate. The nutritional facts obtained here are similar to the values found by Afoakwa (2010) and some Brazilian national brands of milk chocolates. A low content of proteins was found for Afoakwa (2010) chocolate, 7 g/100 g, and also for three Brazilian brands (brand A- 6.8 g/100 g, brand B- 7.2 g/100 g and brand C- 6.5 g/100 g). Moreover, all the chocolates produced here had a higher lipid content than that of the Afoakwa (2010) study (33 g/100 g) and two of the Brazilian brands (Brand A-32.4 g/100 g, and Brand C- 32.5 g/100 g). Brand B showed higher lipid content than the chocolates of this present study (38.4%).

Mineral nutrients are essential for human health, helping vital cells, enzyme activation, hormonal secretion and better vitamin absorption. Elements like manganese, zinc and iron are important in the diet and improve the production of myoglobin and hemoglobin. The deficiency of iron affects a large part of the world population, so its supplementation is important (Gupta & Gupta, 2014). Here we demonstrate that the chocolate with kale that has added mineral content. According to Ayaz et al. (2006) kale has a high level of micronutrients like iron (72.6 μg g⁻¹), manganese (53.5 μg g⁻¹), zinc (9.4 μg g⁻¹), calcium (19.7 μg g⁻¹) and potassium (13.5 μg g⁻¹), which may explain why an increase of 8.6% of ash was found in the chocolate with this addition.

Results found in DPPH and ABTS assays indicated high profile of all produced chocolates as scavenging free radicals. The differences between these two parameters can be explain by reactivity of the two reagents used- DPPH radical reacts with more reactive antiradicals as the same as ABTS radical reacts with intermediary antiradicals (Oliveira et al., 2014). In spite of that, the results in this work also found that the addition of kale and grape, in these conditions, could not improve the antiradical activity of milk chocolate, which may can be explain by the interaction between proteins and polyphenolics. In fact, the difficulty to improve the antioxidant characteristics of chocolate with addition of fruits has been shown by other authors - (Todorovic et al., 2015) rated the polyphenolic profile of milk and dark chocolates produced in Serbia with addition of raspberry. Although no statistic difference between plain dark chocolate and dark chocolate with addition of raspberry, it was found that the antiradical activity against DPPH and ABTS radicals increase with the addition of raspberry. Belščak-Cvitanović et al. (2012) studies the addition of leaves of raspberry (Rubus ideus L.) aimed to increase phenolic compounds on milk and dark chocolate. Despite of the addition of lyophilized extract in all samples, the antiradical capacity as well as the polyphenolic content didn’t increase. It was proposed that sugar and milk interacts with catechins, resulting in lower values of antiradical activity.

Siebert (1999) studied the interaction between proteins and polyphenolic compounds. It is known that the benzene ring and the hydroxyl groups of phenolic compounds can interact with the amine or carboxyl groups of proteins. These bonds create a web that modifies the bioavailability of polyphenolic compounds. Arts et al. (2002) showed that the addition of proteins (casein) in tea, alters its antiradical capacity, resulting in a false result. Jakobek (2015) reviewed the interaction between the polyphenolic compounds with lipids, proteins and carbohydrates. According to the author, these interactions can alter the bioavailability and bio-accessibility of these compounds in the body. Jakobek (2015) explains that the phenolic compounds with high quantities

Figure 5. Chromatography profiles of A- plain chocolate (blue), kale (green), grape (purple) and chocolate with kale addition (orange) on C-18 reverse phase chromatography with detection at 280 nm. The arrows demonstrate the peaks correspondent to enrichment by kale or grape into the chocolate with additions of these two ingredients. B- plain chocolate (blue), grape (green), grape (purple) and chocolate with kale addition (orange) on C-18 reverse phase chromatography with detection at 350 nm. The arrows demonstrate the peaks correspondent to enrichment by kale or grape into the chocolate with additions of these two ingredients.
of hydroxyl groups and with high molecular weight, have the propensity to bind with the proteins. The phenolic compounds present in grape and kale have these chemical characteristics, which may explain the results obtained.

On the other hand, HPLC-DAD results show that some of the compounds present on kale and grape were might transfer to the enriched chocolates. When comparing the control chocolate with the chocolate with kale it is possible to perceive that there are some compounds added. The chromatography suggest the presence of derivate of benzoic acid and cinnamic acid in the chocolate with kale. Ayaz et al. (2008) identify the major phenolic compounds on kale as galic acid, vanillic acid, caffeic acid, ferulic acid, cummaric acid and sinapic acid, which are classified as hydroxycynamic acids and hydroxybenzoic acids. This study corroborates with the data found in the present work, which may explain the addition of these phenolic compounds on chocolate.

It was also observed the transfer of some components from grape to enriched-grape chocolate. Chromatography profiles suggest that these compounds are derivate of cinnamic acid, syringic acid and catequins. Several studies indicated the presence of phenolic compounds in grapes- Liang et al. (2012) found 36 phenolic compounds distributed in classes of anthocyanins, flavonols, hydroxycinamic acid, hydroxybenzoic acids and flavonoides. Ferreira et al. (2016) identify 24 polyphenols on mutant cultivar of Vitis vinifera which are manly phenolic acids, flavonols, stilbenes and anthocyanins. da Silva Padilha et al. (2017) found 20 phenolic compounds on Vitis vinifera and identify as major compounds syringic acid, peonidin, coumaric acid, catequins and epigallocatequins. Whereas this results, it is possible that some of these compounds are added to chocolate.

Catechins are mainly antioxidants of tea and they are associated with lower risk of chronic diseases, as well as the potential of inhibiting cancer cells and reducing the aging process (Braicu et al., 2013). Flavonols are also found in tea and have a high potential for reducing lipid peroxidation, acting as a cardiovascular protector (Heim et al., 2002). Hydrocynamic acids are studied for their antioxidant capacities, mainly because of their high bioactivity (Razzaghi-Asl et al., 2013) as well as hydroxybenzoic acids (Sevgi et al., 2015; Zhang et al., 2015).

We also evaluated physical proprieties of the enriched chocolates. Chocolate is described as a suspension of particles of sugar, milk and cocoa, disperse in cocoa butter. It is considering a non-Newtonian fluid, which means that the shear stress is not proportional to shear rate. Rheology determination of chocolates shows that this type of material has the characteristic to have yield stress and plastic viscosity. As well, it is known that the best way to describe rheological characteristics of chocolates is by the Casson model (Afoakwa et al., 2007; Lannes, 2008), which is adopted by OICC (Office International of Cocoa and Chocolate). The determination of rheological parameters of chocolate, particle size and texture are essential to quality control of these products, mainly because it have an effect on sensorial qualities (Afoakwa, 2010). Particle size higher than 30 µm impact on a gritty texture, as well as particles size next to 20 µm shows an soft texture and creamy mass (Afoakwa, 2010). Shear rate is related to processing of chocolate, as well as molding and coating and influence its structural and physical quality (Toker et al., 2016).

Low moisture in chocolate is an important factor of quality, whereas high values of moisture can cause sugar bloom, a defect that can occur during the shelf life of the product, modifying its characteristics of appearance and brightness (Afoakwa, 2010). Moreover, flow characteristics are also affected by the presence of moisture. Studies have shown that values above 3% of moisture can cause an increase of initial tension and viscosity (Afoakwa et al., 2007). The moisture content found for all three chocolates shows that this important quality characteristic is preserved with the addition of lyophilized fruit and leaves.

Fat content and moisture are modifiers of the characteristics structure of chocolate. Afoakwa (2010) identified that higher fat contents and lower particle size affected the firmness, consistency, cohesiveness and viscosity of chocolates. The hardness is related to the fat content and the type of fat used in the formulation. This can explain the results of hardness obtained in this study. The highest fat content (35%) found in these chocolates could result in higher hardness values. In addition, high fat content can reduce yield value and viscosity, which can also be noticed in this present study.

Rheology can also be affected by particle size characteristics – as texture. As Beckett (2011) reported, distribution of particles in chocolate influences viscosity; the higher the particle size, the lower the viscosity. Toker et al. (2016) show that the particle size is inversely related to yield stress and plastic viscosity and can affect the quality of the chocolates. The finds in the present study corroborate with this, since formulation A has the lowest particle size and a higher viscosity, while formulation C has a higher particle size and lower viscosity. Beckett (2011) reported that the values for chocolate viscosity are between 1-20 Pa.s. Also, the viscosity and yield values are related to distribution of particle size. Lower particle size results in more interaction among the particles, increasing superficial area and resulting in better flow (Afoakwa, 2010; Beckett, 2011; Konar et al., 2014); this relation was confirmed in this present study. The sensory evaluation shows that this difference on rheology and texture does not impact on sensorial perception and suggested that the product can be marketable.

6 Conclusion

Based on the exposed, it was found that it is possible to aggregate functional properties to milk chocolate. The addition of kale increased fiber and mineral content, as well as the addition of grape modified the structural properties of milk chocolate. The antiradical profile suggested that there was no increase of antiradical capacity with the enrichment; however, the chromatography profile of the phenolic compounds demonstrated that some compounds were transferred to the chocolates after the addition.

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References


