Antibacterial activity and physicochemical characteristics of honey from Mato Grosso’s Amazon, Cerrado and Pantanal, Brazil

The objective of this report was to evaluate the physicochemical characteristics and the antibacterial activity of honey produced for human consumption in some counties in Amazon, Cerrado and Pantanal of Mato Grosso, Brazil. Honey samples were collected in 6 municipalities, totaling 18 samples, 3 samples per municipality. The physicochemical characteristics evaluated were moisture, reducing sugars, apparent sucrose, glucose and fructose by HPLC system, ashes, insoluble solids, free acidity, HMF, diastase activity, Lund, Fiehe and Lugol reactions, electrical conductivity and color. The content of total phenolic and flavonoids were quantified using standard curve of gallic acid and quercetin. The antibacterial activity of honey was evaluated by disk diffusion to five bacteria. Fifteen out of eighteen samples were reproved in some of the physicochemical characteristics. Honey’s average content of total phenolic was 35.68 mg GAE/100 g of honey. The content of total phenolic and flavonoids showed positive correlation with the color of the honeys. The studied honeys did not present antibacterial activity, but the phenolic and flavonoid contents originate potential antioxidant and prebiotic for the darkest honeys. Mato Grosso has potential to the production of the floral honey, in need of adequacy on the quality of the honey to the physicochemical characteristic’s patterns.

Keywords: Flavonoids; Phenolic; Color.

Antividade antibacteriana e características físico-químicas do mel da Amazônia, Cerrado e Pantanal de Mato Grosso, Brasil

O objetivo deste trabalho é avaliar as características físico-químicas e a atividade antibacteriana do mel produzido para consumo humano em alguns municípios da Amazônia, Cerrado e Pantanal de Mato Grosso, Brasil. Amostras de mel foram coletadas em 6 municípios, totalizando 18 amostras, 3 amostras por município. As características físico-químicas avaliadas foram: umidade, açúcares redutores, sacarose aparente, glicose e frutose pelo sistema HPLC, cinzas, sólidos insolúveis, ácide livre, HMF, atividade diastásica, reações de Lund, Fiehe e Lugol, condutividade elétrica e cor. O teor de fenólicos totais e flavonóides foi quantificado usando curva padrão de ácido gálico e quercetina. A atividade antibacteriana do mel foi avaliada por disco difusão a cinco bactérias. Quinze das dezoito amostras foram reprovas em algumas das características físico-químicas. O teor médio de fenólico total dos meis foi de 35,68 mg GAE / 100 g de mel. O teor de fenólicos totais e flavonóides apresentou correlação positiva com a cor dos meis. Os meios estudados não apresentaram atividade antibacteriana, mas os teores de fenólicos e flavonóides originaram potencial antioxidante e prebiótico para os meios mais escuros. Mato Grosso tem potencial para a produção do mel floral, necessitando de adequação na qualidade do mel aos padrões das características físico-químicas.

Keywords: Flavonóides; Fenólico; Cor.
INTRODUCTION

The production and quality of honey are positively influenced by the richness of species, number of flowers and close to native forests (SANDE et al., 2009; ALVES et al., 2011), which makes Mato Grosso, a potential state to beekeeping production in Brazil.

The state consists of three biomes of high biodiversity: Amazon (53.6% of the territory), Cerrado (39.6%) and Pantanal (6.79%) (MATO GROSSO, 2013). Nevertheless, Mato Grosso occupies the 16th place in the country’s honey production, with a total of 379 tons/year (IBGE, 2013). These is because of the lack of political and economic projects, structure associations and enable beekeepers (DALLEMOLE et al., 2010).

The floral honey bee *Apis mellifera* L. is a food product produced from the nectar of flowers (AZEREDO et al., 2003), consisting of supersaturated solution of sugars, primarily fructose and glucose and other minority compounds such as proteins, enzymes, amino acids, organic acids, lipids, vitamins, phenolic acids, flavonoids, carotenoids and minerals giving the honey nutritional property and functional activity (ALVAREZ-SUAREZ et al., 2010; KHALIL et al., 2011).

Interest in natural products with functional activity has grown in recent years, which has generated a growing demand for bee products (BALTRUŠAITYTĖ et al., 2007), due to their nutritional and therapeutic properties (SANT’ANA et al., 2012).

The honey ability to inhibit the growth of several Gram positive and Gram negative comes possibly from their flavonoids and phenolic compounds (ALVAREZ-SUAREZ et al., 2010; ESCUREDO et al., 2012), which are produced by secondary metabolism plants and are responsible also for the antioxidant effect (SILICI et al., 2010). The darker honeys have higher concentrations of secondary compounds such as phenolics, flavonoids and carotenoids and consequent greater antimicrobial activity (ALVAREZ-SUAREZ et al., 2010).

The Manuka Honey from New Zealand, monofloral *Leptospermum scoparium*, own evidenced antimicrobial activity. Bacteria are not able to manifest resistance to natural product and even strains multiresistant to antibiotics and drugs may be susceptible to this honey (BLAIR et al., 2009; COOPER et al., 2010). To combat multidrug-resistant bacteria used the synergism between antibiotics and honey and have found that an amount of honey added to the antibiotic enhances its effect (JENKINS et al., 2012).

The physicochemical, sensorial and microbiological characteristics determine the honey quality. The physicochemical analysis can verify the authenticity of the product and detect tampering or the presence of artificial components, guaranteeing food safety for consumers (ABADIO FINCO et al., 2010; BELAY et al., 2013).

The authenticity of honey is defined by legislation of country of origin (CAC, 2001), which, in Brazil follow the Normative Instruction nº 11, of October 20th. 2000, elaborated by the Ministry of Agriculture, Livestock and Supply (BRASIL, 2000). Nevertheless, the labeling of honey has not always met the requirements of Brazilian law (BERA et al., 2005).

The Brazilian honey has been considered in good physicochemical quality, with some exceptions. The Brazilian honey from Bahia was in accordance with the legislation (LACERDA et al., 2010), but 50% of honey
samples from Tocantins were disapproved in relation to legislation (ABADIO FINCO et al., 2010). Excellent quality was found in 87% of samples of organic honey from the Paraná River islands (ALVES et al., 2011). The honey in Ceará was considered with good quality to the physicochemical characteristics, except to HMF content (ALMEIDA-MURADIAN et al., 2013).

The increase in honey production can be achieved with the appreciation of the use and consumption of the product, through the proof of their therapeutic and biological activity. Furthermore, there is a need to develop new classes of antibiotic based on natural products which microorganisms do not exhibit resistance.

The objective of this report was to evaluate the physicochemical characteristics and the antibacterial activity of honey produced for human consumption in some counties in Amazon, Cerrado and Pantanal of Mato Grosso, Brazil.

MATERIAL AND METHODS

Study area and collecting honey

Honey samples from A. mellifera bee were collected in six municipalities of Mato Grosso, Brazil (Figure 1). A total of 18 samples were collected during the study, 3 samples per municipality.

The municipalities were grouped within the biome that had predominance of territorial footprint (Table 1) using AutoCAD 2013 for this. This criterion was used because four of the six cities studied are occupied by two or more biomes (Figure 1). Thus, honey samples were collected in the Amazon biome in the municipalities of Alta Floresta and Marcelândia in the Cerrado in Comodoro and Nossa Senhora do Livramento and the Pantanal in Cáceres and Poconé.

Figure 1: Mato Grosso State municipalities selected to collect honey spread across biomes
These cities were selected because they are the largest state producers (FERREIRA, 2014; IBGE, 2013), or for being part of a Local Productive Arrangement (CBA, 2013).

Amazon occupies an area of 480,215 km² in Mato Grosso, characterized by different physiognomic aspects, with presence of palm trees and lianas, humid climate without dry season, with rainfall well distributed throughout the year and high temperatures (IBGE, 2013).

Cerrado occupies an area of 354,823 km² in Mato Grosso, characterized by vegetation of Savannah, with more than 6500 cataloged plants; the weather is warm, semi-humid and has two distinct seasons: rainy summer and dry winter (IBGE, 2013).

Pantanal occupies a territory of 60,885 km² in the state, has opened vegetation like the Cerrado, but shows parts of rainforests and extensions of the Amazon ecosystem. Its location in a depression favors seasonal flooding during the rainy summer, when the flood (IBGE, 2013).

Table 1. Predominant biome in the cities due to territorial footprint of Amazon, Cerrado and Pantanal biome in the six municipalities selected for the study.

<table>
<thead>
<tr>
<th>Municipalities</th>
<th>Amazon</th>
<th>Cerrado</th>
<th>Pantanal</th>
<th>Predominance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alta Floresta</td>
<td>100</td>
<td>0.0</td>
<td>0.0</td>
<td>Amazon</td>
</tr>
<tr>
<td>Cáceres</td>
<td>5.93</td>
<td>8.97</td>
<td>85.10</td>
<td>Pantanal</td>
</tr>
<tr>
<td>Comodoro</td>
<td>43.88</td>
<td>56.12</td>
<td>0.0</td>
<td>Cerrado</td>
</tr>
<tr>
<td>Marcelândia</td>
<td>100</td>
<td>0.0</td>
<td>0.0</td>
<td>Amazon</td>
</tr>
<tr>
<td>Nossa Senhora do Livramento</td>
<td>0.0</td>
<td>63.95</td>
<td>36.05</td>
<td>Cerrado</td>
</tr>
<tr>
<td>Poconé</td>
<td>0.0</td>
<td>15.74</td>
<td>84.26</td>
<td>Pantanal</td>
</tr>
</tbody>
</table>

Honeys used in the study were from the 2012/2013 crop, harvested between July and October, acquired directly with the beekeepers and samples stored in amber bottles under refrigeration.

Microorganisms

The antibacterial activity was evaluated against bacterial strains from the American Type Culture Collection (ATCC): Staphylococcus aureus (ATCC 25923), Streptococcus pyogenes (ATCC NEWP 0015), Escherichia coli (ATCC 25922), Salmonella typhimurium (ATCC NEWP 0028) and Shigella flexneri (ATCC NEWP 0122).

Physicochemical characteristics

The determination of phenolic and flavonoid content, reducing sugars, apparent sucrose, insoluble solids, pH, free acidity, diastase activity, electrical conductivity, ashes and the reaction of Lugol were carried out in the Laboratory of the Center for Studies in Beekeeping (CETApis), Campus Cáceres at the University of Mato Grosso (UNEMAT).

The determination of moisture, color, HMF, Fiehe and Lund reactions, glucose and fructose were performed in Food Analysis Laboratory in the Department of Food and Experimental Nutrition, Faculty of Pharmaceutical Science School, University of São Paulo (FCF/ USP).

Moisture of honey was determined by refractive method with the aid of Abbe refractometer and Chataway Table (AOAC, 1990). Reducing sugars and apparent sucrose were quantitated by titration of Fehling
Solutions A and B (CAC, 2015). Insoluble solids content was determined by the weight of the residue remaining wash sample (CAC, 2015).

Electrical conductivity was determined with the aid of conductivity, taking account of the fact that the higher the ash content and acidity in honey, the higher the resulting conductivity (VORWOHL et al., 1964). Ash content was calculated by gravimetric method to quantify the organic substances not subjected to high temperatures in a muffle which volatilization (CAC, 2015).

Free acidity was determined by simple titration of honey solution with a solution of 0.05 N sodium hydroxide at pHmeter until pH 8.5 (AOAC, 1990). The optimal pH range of honey is between 3.2 and 4.5 (BERA, 2004). For the quantitative determination of hydroxymethylfurfural (HMF) it was used a high Diastase activity was based on the Schade procedure, which incubate a buffered starch solution using spectrophotometric method (CAC, 2015).

Honey color analysis was performed by colorimetry using the colorimeter "C 221 Honey color analyzer" brand Hanna. The result obtained on the device was compared to the Pfund scale in mm (ALMEIDA-MURADIAN et al., 2013; LANARA, 1981).

Lund reaction was based on natural protein precipitation of honey by tannic acid (IAL, 2005; ALMEIDA-MURADIAN et al., 2008). Fiehe qualitative reaction indicate the presence of HMF. Lugol colorimetric reaction indicate the addition of commercial glucose in honey (IAL, 2005; ALMEIDA-MURADIAN et al., 2008). Analysis of glucose and fructose were performed by HPLC (BOGDANOV ET AL., 1997).

**Total phenolic and flavonoid content**

For the determination of total phenolic content, it was used the Folin-Ciocalteu method (SINGLETON et al., 1985). An aliquot of 100 µL of aqueous solution of honey to 0.2 g/mL was mixed with 0.5 ml of Folin-Ciocalteu and 1.5 mL of sodium carbonate 20% w/v and the volume was completed with distilled water to 10 mL. The absorbance was read at 765 nm. The assay was performed in triplicate, with the average in mg of gallic acid equivalents (GAE)/100 g of honey. The reading of the data was extrapolated into a standard analytical curve of gallic acid (r=0.99).

Total flavonoid content was determined using adapted methods (MEDA et al., 2005), with an aliquot of 3 mL of honey solution in methanol: water (1:1) at a concentration of 500 mg/ml was mixed with 3 mL of 2% methanolic solution of hydrated aluminum chloride (Vetec). After 30 minutes rest, the absorbances were read at a wavelength of 415 nm against the blank consisting of 3 mL of methanol. The data were extrapolated into a standard analytical curve of quercetin in methanol (r=0.99).

**Antibacterial activity**

The determination of the antibacterial activity of honey was performed using the disk diffusion method (MDD), established as standard by the National Committee for Clinical Laboratory Standards (NCCLS, 2003), and the minimum active dilution (MAD) was estimated.

The honeys were tested at concentrations of 5%, 12.5%, 25%, 50%, 75% (w/v) and pure honey. The
microorganisms grown on Mueller Hinton Agar (*Staphylococcus aureus* and *Escherichia coli*), Mueller Hinton Agar supplemented with 5% defibrinated sheep blood in anaerobic environment produced by the anaerobic system Anaerobac - Probac Brazil (*Streptococcus pyogenes*) and Salmonella-Shigella Agar (*Salmonella thypimurim* and *Shigella flexneri*) poured into Petri dishes (NCCLS, 2005).

A suspension of the microorganism to $10^6$ McFarland was spread on the plates (NCCLS, 2005), and immediately thereafter, 10 µL of honey solutions were added on paper discs.

As a positive control it was used commercial discs of antibiotics Oxacillin (1 µg) and Amoxicillin (10 µg) for *Staphylococcus aureus* and *Streptococcus pyogenes* (SCALABRIN et al., 2003; CRUVINEL et al., 2011), Amoxicillin and Ciprofloxacin (5 µg) for *Escherichia coli* (ZANATTA et al., 2004), Ciprofloxacin and Azithromycin (15 µg) for *Salmonella typhimurium*, and Amoxicillin and Azithromycin for *Shigella flexneri* (TESSMANN et al., 2008; MESQUITA et al., 2009).

An artificial honey solution (3 g of sucrose, 35 g of fructose and 45 g of glucose in 17 g of sterile deionized water) was tested to determine whether the inhibitory effects linked to content sugar of honey (MERCÊS et al., 2013).

The reading of the antibacterial activity was performed after 24 and 48 h incubation at a temperature of 37 °C (OSTROSKY et al., 2008). The antibacterial activity was measured by the diameter (mm) of inhibition zone (halo) around the disks, using a digital caliper (FERRONATTO et al., 2007).

**Data analysis**

The physicochemical characteristics were analyzed using descriptive statistics, with the mean and standard deviation of the traits.

Total phenolic and flavonoids content between honeys from different geographical origins was tested with analysis of variance (ANOVA) for geographical origin with six levels the statistical model composed by total phenolic content as the dependent variable and the origin as an independent variable. The statistical model was simplified to three levels, testing the geographic origin for the three biomes (Amazon, Cerrado and Pantanal) as an independent variable, at 5% significance level. The same statistical model was used for total flavonoids.

The contrast test was used to group the geographical origin levels from the average of total phenolics and flavonoids of honey ($p>0.05$). Total phenolics data were transformed to logarithm and total flavonoids to square root to adjust the distribution Normal errors.

To test the relationship of the color of honey with phenolic and flavonoid content, data were submitted to Spearman correlation analysis to be an analysis that does not relate cause and effect of the variables included in the statistical model, removing the dependency between them and facilitating the interpretation of the relationship. The statistical model for correlation analysis was used to compare the color intensity of honey (mm) and the phenolic content. The same statistical model was used to flavonoids. The Spearman correlation is suitable for distribution of non-normal errors.

The minimum active dilution of each geographical origin honey was calculated by linear regression,
a statistical model formed by the zone of bacterial inhibition as the dependent variable and honey concentrations as an independent variable.

The minimum active dilution (MAD) honey for *Staphylococcus aureus* between honey from different geographical origins were tested using analysis of variance (ANOVA), the Kruskal-Wallis type. This analysis was used because the data was not normal. The statistical model has the MAD as the dependent variable and the source municipalities as an independent variable at 95% significance level.

The minimum active dilution of honey for *Escherichia coli* and *Salmonella typhimurium* between honeys of different geographical origins were tested with analysis of variance (ANOVA) to geographical origin in six levels in the statistical model composed of MAD as the dependent variable and municipalities’ source as an independent variable. Means were compared using Tukey mean test with the minimum level of statistical significance of $p<0.05$. The *Escherichia coli* data were transformed to square roots for adjusting the errors from Normal distribution.

The inhibition zones of antibiotics on bacteria were described as the mean and standard deviation of repetitions.

The Shapiro-Wilk test was used to check the fit of the distribution Normal errors. The analyses were performed with R software (R version 3.0.3, 2014).

**RESULTS AND DISCUSSION**

**Physicochemical characteristics**

Moisture content of evaluated honey in this study ranged from 13.6 to 16.8 %, averaging 15.2 % (Table 2), all samples are in accordance with the pattern of 20 % recommended by the Brazilian and international legislation (BRASIL, 2000; CAC, 2001). These results are like those found for honeys from São Paulo, who had moisture content between 16.68 to 18.13% (CANO et al., 2001; PONTARA et al., 2012).

Mato Grosso honey samples had moisture content exceeding 18 % (FERREIRA, 2014; LONGO, 2013), unlike the present study, which indicates that beekeepers have adequate management, improving the quality of Mato Grosso honey. The moisture content was related with honey harvest period (ALVAREZ-SUAREZ et al., 2010). Honey from São Paulo was collected during the dry season as well as samples of this work.

The concentration of reducing sugars from Mato Grosso honeys ranged from 66.25 to 89.23 % with an average of 74.39% (Table 2), all samples are in accordance with the standard stablished by the Brazilian legislation (minimum 65%) (BRASIL, 2000), and international recommendation (minimum 60%) (CAC, 2001). These results are close to those observed in honey produced in native vegetation sites in Paraná (67.13 to 72.04%) (ALVES et al., 2011), and in honey produced in Ethiopia (from 67.79 to 74.62%) (BELAY et al., 2013). The sugars are the major components of honey, including about 95% by dry weight of the honey (KÜÇÜK et al., 2007).
Table 2. Physicochemical characteristics of *Apis mellifera* honey from Amazon, Cerrado and Pantanal of Mato Grosso, Brazil, 2014, expressed as mean and standard deviation (SD), according to the national and international quality standards

<table>
<thead>
<tr>
<th>Physicochemical characteristics</th>
<th>Mean ± SD (N=18)</th>
<th>BRAZIL&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Codex&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture (%)            </td>
<td>15.2 ± 0.95</td>
<td>Max. 20</td>
<td>Max. 20</td>
</tr>
<tr>
<td>Reducing sugars (%)</td>
<td>74.39 ± 6.01</td>
<td>Min. 65</td>
<td>Min. 60</td>
</tr>
<tr>
<td>Apparent sucrose (%)</td>
<td>2.41 ± 3.18</td>
<td>Max. 6</td>
<td>Max. 5</td>
</tr>
<tr>
<td>Insoluble solids (%)</td>
<td>0.10 ± 0.07</td>
<td>Max. 0.1</td>
<td>Max. 0.1</td>
</tr>
<tr>
<td>Electrical conductivity (μS.cm&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>433.67 ± 193.6</td>
<td>-</td>
<td>Max. 800</td>
</tr>
<tr>
<td>Ashes (%)</td>
<td>0.24 ± 0.20</td>
<td>Max. 0.6</td>
<td>-</td>
</tr>
<tr>
<td>Free acidity (mEq.kg&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>38.27 ± 11.69</td>
<td>Max. 40</td>
<td>Max. 50</td>
</tr>
<tr>
<td>HMF (mg.kg&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>34.69 ± 35.05</td>
<td>Max. 60</td>
<td>Max. 80</td>
</tr>
<tr>
<td>Diastase (Un. Goethe/g of honey)</td>
<td>50.65 ± 3.55</td>
<td>Min. 3</td>
<td>Min. 8</td>
</tr>
<tr>
<td>Fiehe reaction (positive)</td>
<td>11 samples</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Lund reaction (mL)</td>
<td>0.81 ± 0.45</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Lugol reaction (positive)</td>
<td>0 samples</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Glucose (%)</td>
<td>39.09 ± 3.82</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Fructose (%)</td>
<td>48.07 ± 4.08</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Legend: ( - ): not required standard; HMF: hydroxymethylfurfural; Max.: maximum; Min.: minimum; SD: standard deviation.

<sup>a</sup>Value recommended by Brazilian legislation (BRASIL, 2000).

<sup>b</sup>Value recommended by the Codex Alimentarius (CAC, 2001).

The apparent sucrose showed values between zero and 12.15%, with an average of 2.41% (Table 2). Only one sample was out of the standard of the Brazilian legislation (BRASIL, 2000), which establishes a maximum of 6% and three samples were not in accordance with the maximum of 5% from international recommendation (CAC, 2001). The higher levels of apparent sucrose (1.98 to 6.22%) was found in honey samples from Turkey and attributed this to the fact that bees possibly have been fed with sugar syrup in place of nectar (KAHRAMAN et al., 2010), which was not what occurred in this study in Mato Grosso.

The increase of sucrose in honey are due to early crop, when honeycombs are not operculate or even due to addition of sucrose commercial honey (GOMES et al., 2010). This management feature should provide guidance to beekeepers as good beekeeping practices for quality honey production.

Insoluble solid content of the studied honey varied from 0.0085 to 0.24% (Table 2) and six samples were out of the Brazilian standard (BRASIL, 2000; CAC, 2001), which limits to a maximum of 0.1%. This result was like honey from Ethiopia who found levels of up to 0.38% of insoluble solids (BELAY et al., 2013). The presence of impurities in honey (wax, pollen, honeycomb debris, dirt and bees’ particles) may be attributed to the lack of knowledge or lack of quality in the product extraction, processing and packaging (BELAY et al., 2013).

Electrical conductivity varied 147 to 901 μS.cm<sup>-1</sup>, with an average of 433.67 μS.cm<sup>-1</sup> (Table 2) and only one sample was out of the international standard which recommends a maximum of 800 μS.cm<sup>-1</sup> (CAC, 2001). Electrical conductivity is not present in the Brazilian legislation (BRASIL, 2000). Similar values (340 and 1040 μS.cm<sup>-1</sup>) was found for electrical conductivity of Tocantins honey samples (ABADIO FINCO et al., 2010), and this variation related to the pH, acidity, and ash content present in honey.

Apparent sucrose content and electrical conductivity were two characteristics that presented more samples out of the standard for previous study for honeys from Mato Grosso (FERREIRA, 2014; LONGO, 2013), different from the present study which only one sample was out of the standard. This shows that
beekeepers are paying attention to achieving the harvest at the right time, avoiding pH extremely acid and high levels of apparent sucrose.

Ashes average content of the honey was 0.24%, with values from 0.02 to 0.87% (Table 2). Two of the samples were out of the recommended by the Brazilian legislation (maximum of 0.6% for floral honey samples) (BRASIL, 2000). The ashes average content allows detecting some irregularities in honey, especially the lack of hygiene and/or filtration after harvesting the honey by beekeepers (VILHENA et al., 1999). The monofloral honey *Eucalyptus* from São Paulo had more ashes compared with honey of other floral sources (BARTH et al., 2013; FELSNER et al., 2004), indicating that this characteristic may be related to the botanical origin.

Free acidity level of honeys ranged from 23.13 to 59.81 mEq/kg (Table 2), with eight samples outside of the Brazilian standard (maximum 40 mEq/kg) (BRASIL, 2000), but only four samples were out of the international standard (maximum 50 mEq/kg) (CAC, 2001). Some authors reported that besides the polinic source, incorrect storage or early harvest can modify the acidity of honey, which can be improved with the use of good practice of beekeeping management (GOMES et al., 2010). Moreover, the environment with high temperature and high humidity promotes the acidification reaction of honey (ALMEIDA-MURADIAN et al., 2007).

The use of honey house together with the number of hives are the most important beekeeping practices to ensure the physicochemical properties of honey, and the number of hives expresses the professionalism of the beekeeper from Baixada Cuiabana and the Pantanal of Mato Grosso, Brazil (LONGO, 2013).

HMF content presented an average of 34.69 mg.kg$^{-1}$, ranging from 2.56 to 127.28 mg.kg$^{-1}$, with four samples in disagreement with the Brazilian standard (maximum 60 mg/kg) (BRASIL, 2000), and two samples out of the international recommendation (maximum 80 mg/kg) (CAC, 2001). The fresh honey does not contain HMF. However, if the honey is subjected to high temperatures (BOGDANOV et al., 2004), inadequate storage conditions or long storage time it begins to show high levels of HMF (SANTOS et al., 2010). In Mato Grosso, predominantly high temperatures throughout the year, it demands greater care with honey conservation.

All samples presented HMF values were in accordance with the Brazilian legislation (minimum of 3 mg.kg$^{-1}$) (BRASIL, 2000), and international recommendation (minimum of 8 mg.kg$^{-1}$) (CAC, 2001). For diastase value, it was obtained an average of 50.65 Un. Goethe/g of honey (Table 2). The honey Cuba was within the international standard (13.4 to 33.4 Un. Goethe/g of honey) (ALVAREZ-SUAREZ et al., 2010), and state that this characteristic indicates that honey has quality.

Eleven of the 18 samples showed positive reaction Fiehe. The results of qualitative analysis of HMF to Tocantins honey showed that 25% of the samples showed a positive response to Fiehe reaction (ABADIO FINCO et al., 2010), indicating poor quality of these honeys for possible tampering or overheating. The results of qualitative HMF of this study were confirmed by quantitative analysis, which found four samples in disagreement with the Brazilian standard and two with the international standard, although all agree
regarding diastase values, indicating possible overheating of some samples.

For Lund reaction, five of the 18 samples were not in accordance with the recommendation of 0.6 to 3 mL (ALMEIDA-MURADIAN et al., 2008; IAL, 2005), with a deposit below 0.6 mL. In honeys from Tocantins, eight samples exceeding the standard value from Lund reaction, suggesting loss or addition of protein substances in product during the processing (COOPER et al., 2010).

All samples showed negative Lugol reaction. The same result was found in honey from Ceará (PAULINO et al., 2009). This test indicates qualitatively the possible adulteration of the honey with commercial glucose.

The honey content of the two sugars in total was found up to 70–95%. The honey samples content in glucose and fructose was determined by HPLC (32.45–43.98% and 36.76–51.64%, respectively).

Honey color analysis showed a great variation, extra white (17 mm on the Pfund scale) to amber (100 mm), but most of the samples (44%) were predominated light amber color (50 to 84 mm).

pH average was 3.9, ranging from 3.6 to 4.4, similar with Tocantins’ honey which presented pH between 3.35 and 4.50 (ABADIO FINCO et al., 2010).

Content of total phenolic and flavonoid

For total phenolic, Amazon honeys studied in this work showed 15.0 to 41.5 mg GAE/100 g of honey (x=31.33), Cerrado honeys 17.0 to 59.5 mg of EAG/100 g of honey (x=34.08) and Pantanal honeys 14.0 to 122.5 mg GAE/100 g of honey (x=47.92) (Table 3), with no significant difference between the three studied biomes (p=0.64).

Total phenolic content in honey was similar between the source of the municipalities of Alta Floresta, Cáceres and Comodoros (x=26.67), which was grouped by the contrast test (p=0.68), the municipalities of Marcelândia, Nossa Senhora do Livramento and Poconé had similar content (x=48.89) and formed another group (p=0.24).

The Santa Catarina honey was found content of phenolic compounds between 36.37 and 71.69 mg GAE/100 g of from (RIZELIO, 2011). The honey from other countries and different floral origins, and found total phenolic content from 32.59 to 114.74 mg GAE/100 g of honey (NCCLS, 2003), while a publication related to honey from Pantanal of Mato Grosso do Sul identified levels of total phenolic compounds with an average of 61.52 and 222.03 mg GAE/100 g of honey, depending on the place of harvest (BERTOLDI et al., 2012). Those results were considerably higher than those of honeys evaluated in this study (BERTOLDI et al., 2012; MEDA et al., 2005).

Table 3. Levels of total phenolics and flavonoids present in honey of Apis mellifera, from Mato Grosso, Brazil, 2014

<table>
<thead>
<tr>
<th>Biome honey source</th>
<th>Total phenolics * (mg de GAE/100 g of honey)</th>
<th>Total flavonoids * (mg de QE/100 g of honey)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pantanal</td>
<td>47.92 ± 37.96</td>
<td>0.26 ± 0.04</td>
</tr>
<tr>
<td>Cerrado</td>
<td>34.08 ± 15.03</td>
<td>0.28 ± 0.09</td>
</tr>
<tr>
<td>Amazon</td>
<td>31.33 ± 11.18</td>
<td>0.27 ± 0.06</td>
</tr>
</tbody>
</table>

Legend: GAE: gallic acid equivalent; QE: equivalent quercetin.

*Means in columns do not differ at 5% by F test. N = 18.
For total flavonoid content, it was obtained an average of 0.27 mg QE/100 g of honey for Amazon samples, with the highest concentration of 0.36 and less than 0.19 mg QE/100 g of honey. The average value obtained for Cerrado honeys was 0.28 mg QE/100 g of honey, ranging from 0.21 to 0.47 mg QE/100 g of honey. Pantanal honeys presented an average of 0.26 mg QE/100 g of honey, ranging from 0.20 to 0.34 mg QE/100 g of honey (Table 3). These results did not differ significantly between the three studied biomes (p=0.89).

The flavonoid content in honey was similar between the source of the municipalities of Alta Floresta, Cáceres and Comodoro (x=0.24), which was grouped by the contrast test (p=0.69). The municipalities of Marcelândia, Nossa Senhora do Livramento and Poconé had similar content (x=0.30) and formed another group (p=0.73).

Flavonoid content found in the studied honey are lower than those honey of various types, from different countries, and found levels between 0.17 and 8.35 mg QE/100 g of honey (MEDA et al., 2005).

The phenolic content and total flavonoids were grouped into two levels as the geographical origin of the municipalities. The botanical source of phenolic compounds and flavonoids may be the same, so the grouped municipalities should be similar for the availability of floral resources.

These results suggested that the geographical origin of municipalities, not the biome of origin, influences the content of these metabolites, however there is similarity between the municipalities of Alta Floresta, Cáceres and Comodoro and between Marcelândia, Nossa Senhora do Livramento and Poconé.

The phenolic and flavonoids are responsible for the honey bioactivity. They have antioxidant, antimicrobial, antiviral and anticarcinogenic capacity, generating a beneficial effect on human health, and awakening interest in identification and quantification (KÜÇÜK et al., 2007; RIZELIO, 2011).

The honey color variation depends on the combination of many factors, including the content of phenolic acids and flavonoids (ALVAREZ-SUAREZ et al., 2010; BERETTA et al., 2005). The honey production methods and agricultural practices can also influence the color (ALMEIDA-MURADIAN et al., 2013).

Total phenolic content was positively correlated with the color of honey (r=0.735, p<0.01, Table 4, Figure 2), corroborating the work of Alvarez-Suarez et al. (2010), which found higher total phenolic contents for the amber honey, and lower level for extra white honey. The lighter honey Portugal the phenolic content was lower than in the darkest honey (FERREIRA et al., 2009).

The amber honey (darker color) had higher levels of total flavonoids (0.47 mg QE/100 g of honey), while extra white honey (lighter color) had the lowest content of flavonoids (0.19 QE mg/100 g of honey). This was confirmed by the positive correlation between color and the quantity of total flavonoids (r=0.891, p<0.01, Table 4, Figure 3).

A positive correlation between total flavonoid content and the color of honeys from Cuba was observed, as the amber honey showed a high flavonoid content and white extra honey very low content. The ratio of increase in the color intensity of honey with the concentration of total phenolics, and flavonoids (MEDA et al., 2005).

A significant positive correlation was found between phenolic content and total flavonoid (r=0.764,
p<0.01), as well as Cuban honey that the increase of the intensity of color appears to be related to the increased concentration of total flavonoids and phenolic compounds (ALVAREZ-SUAREZ et al., 2010). Thus, the present study results seem to confirm that the contents of these compounds can directly affect the color of honey.

Table 4. Spearman correlation between the color and the total phenolic and flavonoids content of *Apis mellifera* honey, Mato Grosso, Brazil, 2014

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Color</th>
<th>Phenolic</th>
<th>Flavonoid</th>
</tr>
</thead>
<tbody>
<tr>
<td>Color</td>
<td>1.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phenolic</td>
<td>0.73**</td>
<td>1.00</td>
<td></td>
</tr>
<tr>
<td>Flavonoid</td>
<td>0.89**</td>
<td>0.76*</td>
<td>1.00</td>
</tr>
</tbody>
</table>

Legend: *p<0.05, **p<0.01. N = 18

Figure 2. Relationship of color (in mm Pfund scale) and total phenolic content (mg GAE/100 g of honey) of honey from the Amazon, Cerrado and Pantanal of Mato Grosso, Brazil, 2014. (Spearman correlation: r=0.73, p<0.01)

Figure 3: Relationship of color (in mm Pfund scale) and total flavonoid content (mg QE/100 g of honey) of Brazilian honey from the Amazon, Cerrado and Pantanal area of Mato Grosso, 2014. (Spearman correlation: r=0.89, p<0.01)
Antibacterial activity

The antibacterial activity of Mato Grosso honey studied in this work presented minimum active dilution for the bacteria *Staphylococcus aureus*, *Escherichia coli* and *Salmonella typhimurium* of 44.58% w/v (Table 5). The minimum active dilution for each bacterium did not differ between geographical origins of honey (Table 5). Honey showed no antibacterial activity against strains *Streptococcus pyogenes* and *Shigella flexneri*, with no inhibition zone formation around the discs with treatment.

The minimum active dilution of 44.58% w/v was considered not effective for *Staphylococcus aureus* compared the Cuban honeys with MAD 4.02% v/v (deemed sensitive) (ALVAREZ-SUAREZ et al., 2010), and MAD 7.96 and 8.02% v/v for *Escherichia coli* and *Bacillus subtilis*, respectively (considered moderate sensitive).

The estimated *Staphylococcus aureus* zone of inhibition that for every mm was required a concentration of 15.5% w/v in Greek honey (VOIDAROU et al., 2011), concentration three times lower than the concentration found for Mato Grosso honey. In another study, the minimum active dilution of Manuka honey and pasture honey against *Staphylococcus* coagulase-negative isolates from New Zealand patients ranged from 2.7 to 5.0% v/v (FRENCH et al., 2005), concentration nine times lower than that presented by Mato Grosso honey.

The artificial honey did not form inhibition zone for any of the tested microorganisms, Manuka honey and Ulmo honey inhibited the microorganisms tested more extensively than the artificial honey (SHERLOCK et al., 2010). The idea that sugar is solely responsible for the honey antibacterial activity was not validated (HANNAN et al., 2009), and claim that the antibacterial effect of natural honey is not connected only with the high osmolarity, because in their study, the artificial honey inhibited the bacteria tested at higher concentrations than the natural honey.

<table>
<thead>
<tr>
<th>Bacteria</th>
<th>MAD of honey (mean ± SD)**</th>
<th>R²</th>
<th>p*</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Staphylococcus aureus</em></td>
<td>44.58 ± 141.42</td>
<td>0.67</td>
<td>0.69**</td>
</tr>
<tr>
<td><em>Escherichia coli</em></td>
<td>44.58 ± 99.84</td>
<td>0.84</td>
<td>0.98**</td>
</tr>
<tr>
<td><em>Salmonella typhimurium</em></td>
<td>44.58 ± 321.29</td>
<td>0.96</td>
<td>1.00**</td>
</tr>
</tbody>
</table>

**Minimum active dilution of honey in % w/v, resulting in a 1 mm zone of inhibition.

*ANOVA for comparison of the probability of minimum active dilution of honey between geographical origins by Kruskal-Wallis test for *Staphylococcus aureus* and the F test for *Escherichia coli* and *Salmonella typhimurium*. N=18.

Bacteria were sensitive to commercially antibiotics, with inhibition zone average of 33.26 mm for *Staphylococcus aureus*, 27.27 mm for *Streptococcus pyogenes*, 22.61 mm for *Escherichia coli*, 22.09 mm for *Salmonella typhimurium* and 24.88 mm for *Shigella flexneri*.

The antimicrobial activity of Cuban honey was related to differences in the profile of phenolic and flavonoids compounds (quercetin and gallic acid, respectively), because the honey that showed higher phenolic and flavonoid content showed higher antimicrobial activity (ALVAREZ-SUAREZ et al., 2010). The high antimicrobial activity of honey from Portugal was associated to the high content of phenolics and flavonoids (quercetin and caffeic acid, respectively) (MIGUEL et al., 2013).

The honey produced in other countries may have different antimicrobial activity of honey produced
in Brazil, more specifically of Mato Grosso honey, because there can be great variation in the profile of therapeutic components (phenolics and flavonoids) depending on their geographical origin (ESTEVINHO et al., 2012; MOLAN, 2002).

Honey from Amazon, Cerrado and Pantanal of Mato Grosso need to be studied in search of other different phenolic compounds of gallic acid investigated in this work, such as protocatechuic acid, p-hydroxybenzoic acid, caffeic acid, chlorogenic acid, vanillic acid, p-coumaric acid, benzoic acid, cinnamic acid and ellagic acid, identified in Portuguese honey (ESTEVINHO et al., 2008). Other flavonoids different from quercetin should be investigated in Mato Grosso honey, such as naringenin, kaempferol, apigenin, pinocembrine and chrysin, identified in Portuguese honey (ESTEVINHO et al., 2008). These phenolic and flavonoids compounds may be associated with antimicrobial activity attributed to honey (ALJADI et al., 2003; YAO et al., 2003).

The methodological limitations of this study were due to antimicrobial activity by disk diffusion and not by microdilution broth, which is stronger in the search for pharmacological interest, and the representativeness of the six municipalities in Mato Grosso state in relation to the universe one hundred forty-one municipalities.

CONCLUSION

Mato Grosso has potential to floral honey production, even if the physicochemical quality of the product needs to be adequate, especially in relation to apparent sucrose, insoluble solids, ashes and free acidity.

The darker honey has a higher total phenolic and flavonoid content, although they are not indicated in the control of *Staphylococcus aureus*, *Streptococcus pyogenes*, *Escherichia coli*, *Salmonella typhimurium* and *Shigella flexneri*.

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