The Effect of Honey Supplementation on Plasma Levels of Short Chain Fatty Acids in Healthy Infants

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Abstract

Background: Honey is a sweet and flavorful natural product, which is consumed for its high nutritive value and for its positive effects on human health, including antioxidant, antimicrobial, anti-inflammatory, wound healing, antitumor, immunomodulator, prebiotic and probiotic properties. Honey is produced by honey bees from plant nectars, plant secretions and excretions of plant-sucking insects. Concerning its nutrient profile, it represents an interesting source of natural macro- and micro-nutrients, consisting of a saturated solution of fructose, glucose, and to Fructo-Oligosaccharides (FOS) that can serve as prebiotics, but also of a wide range of minor constituents, especially phenolic compounds.

Aim of Study: Evaluation of the effect of honey supplementation on plasma levels of short chain fatty acids in healthy infants and on the anthropometric measurements.

Subjects and Methods: This study was a single arm prospective interventional study. Twenty healthy infants aged 1 to 2 years were consecutively recruited from the Outpatient Pediatric Clinic of Ain Shams University, Egypt during the period from January, 2020 to February, 2020. Each infant received 2g honey/kg/day for 8 weeks. The plasma level of SCFAS and the anthropometric measurements were compared before and after honey intervention.

Results: There was a statistically significant increase in plasma level of SCFAS (formic, acetic and butyric) after honey intervention in the studied group (p-value <0.05). There was also a statistically significant increase in the body weight (p-value 0.001) and OFC (p-value 0.03 1). Honey consumption did not produce any adverse effect in the studied group for 8 weeks.

Conclusion: 8 weeks of honey consumption resulted in increased plasma SCFAS level (formic, acetic and butyric) and increase in the body weight and OFC.

Key Words: Honey – Short chain fatty acids – Occiput frontal circumference.

Introduction

HONEY is a natural substance produced by honey bees. It has both nutritive and health benefits. The positive effects of honey on health are due to its anti-oxidant, anti-microbial, anti-inflammatory and wound healing effects [1].

Honey has to be employed as a dietary adjunct. In this respect, it acts as a prebiotic, which is defined as a non-digestible food ingredient that beneficially affects the host by selectively stimulating the growth and/or activity of a limited number of bacteria (bifid bacteria and lactobacilli) in the intestine [2].

On the account of the nutritional value (303 kcal/100g honey) and fast absorption of its carbohydrate, honey is a food suitable for humans of every age [3]. Simply, when orally consumed, its carbohydrates are easily digested and quickly transported into the blood and can be utilized for energy requirements by the human body. The positive effects of honey in infant diet are attributed to effects on the digestion process. One possible cause is the well-established effect of oligosaccharides on B. bifidus [4].

Honey has been used as nutritional and medical product since the earliest times [5,6].

Honey contains more than 180 substances, including amino acids, vitamins, minerals and carbohydrate constituents as fructose and glucose in addition to oligosaccharides [7]. Oligosaccharides are provided as fermentable prebiotic substances [8]. To selectively modulate the gut microbial balance in favour of probiotic Lactobacilli and bifid bacteria, thus improving the host metabolic function [9,10].
Clinical trials try to manipulate the microbiome in the effort to prevent or treat infancy and childhood diseases with a focus on probiotics, prebiotics and symbiotics. Functional foods are foods that affect the component of the gut micro biota, which lead to generation of SCFAs [11].

Aim of the work:

Evaluation of the effect of honey supplementation on plasma levels of short chain fatty acids and on the anthropometric measurements in healthy infants.

Subjects and Methods

This study was a single arm prospective interventional study that was conducted on twenty healthy infants aged 1 to 2 years. The sex ratio was 1:1. The infants were consecutively recruited from the Outpatient Pediatric Clinic of Ain Shams University, Egypt during the period from January, 2020 to February, 2020.

The study was approved by the Local ethical committee, and an informed consent was obtained from the mother or care giver of each infant before enrolment into the study.

Plasma level of SCFAS were measured and the anthropometric measurements were compared before and after honey intervention including Body Weight (BW) which was measured to the nearest 10g by using a standard digital scale, Body Length (BL) which was also measured to the nearest 1cm, weight for height which more accurately assesses body build and distinguishes wasting (acute malnutrition) from stunting (chronic malnutrition), Body Mass Index which was calculated as the following equation (BMI = Weight (kg)/length (m)²), head circumference (the measurement was approximated to the nearest 0.1cm) and Mid-Upper Arm Circumference (MUAC) which was recorded to the nearest 1mm.

All anthropometric measures were plotted on WHO, 2006, data were entered on WHO anthropometric software.

Honey intervention: The honey used in this study was a raw, unprocessed clover honey, collected from Al-Mahala-Gharbia Governorate, Egypt, and directly supplied by a beekeeper. Physicochemical analysis of the honey was done in the Chemical Analysis Laboratory of Honey Bee Products, Beekeeping Research Center, Plant Protection Research Institute, Agriculture Research Center, Giza, Egypt. The honey had a pH of 3.7; moisture content of 18.8%; electrical conductivity of 0.27 mS/cm; and a carbohydrate content of 78.4g/100g, with a fructose to glucose ratio of 1.2:0.8, respectively, and a non-reducing sugar content of 3.4g/100g. The Hydroxymethylfurfuraldehyde (HMF) content was 1.6mg/kg. Values of HMF less than 15mg/kg indicate fresh honey not exposed to heat [12]. Microscopic examination of samples from honey confirmed the presence of pollen grains, which were mainly of clover (Trifolium alexandrinum). The honey was also tested for the presence of Clostridium botulinum spores before use (no spores were detected). Examination of honey for C. botulinum spores was done by centrifugation and filtration of the supernatant, followed by culture on cooked meat [13]. Each participant received honey in an oral dose of 2g/Kg/day for 8 weeks. The calculated dose of honey was dissolved in water in a ratio of 1 to 3 respectively and ingested before breakfast. Dissolving honey in water enhances the antimicrobial effect of honey [14]. The total caloric value of honey was subtracted from the total daily caloric intake. Each participant was provided by 7 glass containers each week. Each container contained the calculated honey dose to be dissolved in water just before ingestion. The caregivers instructed not to give their infants any additional honey doses during the study.

Plasma level of short chain fatty acids (Acetic acid, butyric acid and formic acid) were measured before and after honey intervention using HPLC (Agilent technologies 1100 series, with a quaternary pump (G131A model) according to the method described previously by Miwa & Yamamoto [15] and modified by Hussein et al., [16] and Youness et al., [17].

Statistical analysis:

Data were collected, revised, coded and entered to the Statistical Package for Social Science (IBM SPSS) version 23. The quantitative data were presented as mean ± standard deviation and ranges when their distribution was parametric (normal) while non-normally distributed variables (non-parametric data) were presented as median with Inter-Quartile Range (IQR). Also qualitative variables were presented as number and percentages. The comparison between groups with qualitative data was done by using Chi-square test and Fisher exact test instead of Chi-square test only when the expected count in any cell less than 5. The comparison between two groups with quantitative data and non-parametric distribution was done by using Mann-Whitney Test. The comparison between two paired groups with quantitative data and parametric distribution was done by using paired t-test while
with non-parametric data the comparison was done by using Wilcoxon Rank test. Spearman correlation coefficients were used to assess the correlation between two quantitative parameters in the same group. The confidence interval was set to 95% and the margin of error accepted was set to 5%. So, the \( p \)-value was considered significant as the following: \( p > 0.05 \): Non significant. \( p < 0.05 \): Significant. \( p < 0.01 \): Highly significant.

### Results

There was a statistically significant increase in plasma level of SCFAS (formic, acetic and butyric) after honey intervention in the studied group \( (p\text{-value} < 0.05) \) as shown in (Table 1).

There was also a statistically significant increase in the body weight \( (p\text{-value} 0.001) \) and OFC \( (p\text{-value} 0.031) \) as shown in (Table 2).

#### Table (1): Comparison between serum SCFAS level before and after honey intervention in the studied group.

<table>
<thead>
<tr>
<th>SCFAS Type</th>
<th>Median (IQR)</th>
<th>Range</th>
<th>Test value</th>
<th>( p )-value</th>
<th>Sig.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Formic (microg/ml):</td>
<td>15.44 (9.98-22.92)</td>
<td>9.08-45.31</td>
<td>17.23 (11.43-30.35)</td>
<td>39 (–6-73)</td>
<td>( -2.170 )</td>
</tr>
<tr>
<td>Acetic (microg/ml):</td>
<td>11.52 (6.27-15.35)</td>
<td>3.82-30.23</td>
<td>24.34 (17.5-67.06)</td>
<td>189 (–3-425)</td>
<td>( -2.280 )</td>
</tr>
<tr>
<td>Butyric (microg/ml):</td>
<td>22.38 (18.11-47.27)</td>
<td>11.93-197.67</td>
<td>29.43 (16.3-83.7)</td>
<td>53 (12-183)</td>
<td>( -2.121 )</td>
</tr>
</tbody>
</table>

\( p \)-value >0.05: Non significant. \( p \)-value <0.05: Significant. \( p \)-value <0.01: Highly significant.

#### Table (2): Comparison between actual values after honey supplementation and expected values for age of anthropometric measurements.

<table>
<thead>
<tr>
<th>Measurement</th>
<th>Actual value</th>
<th>Expected value</th>
<th>Test value</th>
<th>( p )-value</th>
<th>Sig.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wt in kg:</td>
<td>Mean ± SD</td>
<td>10.58±1.35</td>
<td>10.28±1.37</td>
<td>3.929*</td>
<td>0.001 HS</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>8.8-13.3</td>
<td>8.3-13</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lt in cm:</td>
<td>Mean ± SD</td>
<td>79.80±7.44</td>
<td>80.80±3.20</td>
<td>–0.783</td>
<td>0.443 NS</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>73-98.5</td>
<td>75-86</td>
<td></td>
<td></td>
</tr>
<tr>
<td>OFC:</td>
<td>Mean ± SD</td>
<td>46.95±1.99</td>
<td>46.27±1.09</td>
<td>2.250*</td>
<td>0.037 S</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>43-49</td>
<td>44.4-48</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\( p \)-value >0.05: Non significant. \( p \)-value <0.05: Significant. \( p \)-value <0.01: Highly significant. *: Paired t-test.

Honey consumption did not produce any adverse effect in the studied group for 8 weeks.

### Discussion

In the present study, 8 weeks consumption in twenty healthy infants resulted in such positive effect as increasing plasma level of SCFAS (formic, acetic and butyric) through providing to gut microbiota otherwise indigestible polysaccharides, oligosaccharides, and proteins to fermentate to SCFAS which can be incorporated in glucose-and lipid synthesis and can function as signaling molecules in several metabolic and endocrine processes [18] and providing also a major source of energy and nutrients [19].

Honey, having 0.75% of fructo oligosaccharides in its constitution, may act as a prebiotic. Moreover, the detrimental effects of bile salts on Bifidobacterium spp, a ubiquitous inhabitant of the human gastrointestinal tract, can be overcome by the action of fructooligosaccharides and their monomeric derivatives [20].

Honey contains many oligosaccharides and low molecular weight polysaccharides that can resist degradation by host enzymes, and thus remain available as nutrient source for the intestinal micro flora thereby acknowledged as a prebiotic material [21].

When prebiotics enter the lower part of the gastrointestinal tract, they stimulate the growth and/or activity of health-promoting bacteria in the colon. It has been demonstrated that these bacteria generally including lactobacilli and bifidobacteria species protect the host by competing with bacterial and fungal pathogens for availability of nutrients and space and also modulating the immune system [21].

In most of the studies reported, honey has shown to support the growth of the probiotics when incubated in optimum conditions with milk (including reconstituted or fermented) or selective
growth media. Furthermore, inhibitory action was demonstrated against the pathogens and other intestinal microbes [22-24].

Similar to the positive effect of honey on study of Haddadin [25] to evaluating effect of honey on the growth and metabolism of two Bacterial Species (Bifidobacterium infantis and Lactobacillus acidophilus—both of human intestinal origin) showed that honey beneficially influenced the growth and metabolism [production of Short-Chain Fatty Acids (SCFA)] of these two organisms and it might be reasonable to assume that honey ingested by a consumer would have a similar effect on the native populations of these species in the lower intestine.

To our knowledge, the present study may be the first study correlated between honey supplementation and serum SCFAS levels in healthy infants.

There was also a statistically significant increase in the body weight and OFC as honey supplementation increased GET (gastric emptying time) patients with positive effect on the improvement in the anthropometric measurements and serum albumin [26].

Aly [27] found that Infants who received honey gained more weight than controls, which can be explained by the increased caloric intake in association with honey consumption.

Rao [28] demonstrated increased weight gain in infants receiving prebiotics, whereas Mugambi [29], did not show any benefit of adding prebiotic combinations of Galacto-Oligosaccharides (GOS) and FOS or FOS alone on weight gain. It is known that honey has both pre-and probiotic effects [30].

Evidence of the role of SCFA in appetite regulation has recently appeared in a study using selective modulation of colonic propionate in humans which demonstrated that propionate appears to induce short-term appetite regulation [31].

**Conclusion:**

8 weeks of honey consumption resulted in increased plasma SCFAS level (formic, acetic and butyric) and increase in the body weight and OFC.

**Acknowledgments:**

We thank very much all children and their parents who agreed to participate in this study. We also thank Dr. Seid J, Professor of Medical Biochemistry at National Research Center who measured plasma level of SCFAS.

**References**


تأثير إعطاء عسل النحل على مستوى الأحماض الدهنية قصيرة السلسلة في بلازما الأطفال الرضع الأصحاء

حيث أن الهدف من الدراسة معرفة إعطاء عسل النحل على مستوى الأحماض الدهنية قصيرة السلسلة في بلازما الأطفال الرضع الأصحاء وعلى قياسات الأطفال الجسمانية المختلفة.

وهذه الدراسة عبرة عن دراسة متصلة تفاعلية حيث جرت على 20 طفل من الأطفال يتراوح أعمارهم من سنة إلى سنتين ومن كلا الجنسين المنحدرين من عيادة الخارجية في مستشفى الأطفال مستشفيات جامعة عين شمس، مصر، وتراوحت فترة الدراسة شهرين (8 أسابيع). وتم الحصول على الموافقة من أحد الوالدين على الاقل قبل عمر الدراسة على الأطفال. وتم خضوع جميع الأطفال إلى ما يلي:

1. تجميع بيانات من الأطفال المشاركين فيما يتعلق بالعديد الذي، أي حالة مرضية تتعارض مع إمتصاص الفذاء أو عدم الإستقامة من بشكل مباشر، التأثير غير أن امتصاص الماء أو مزيج. وفحص البذور الكامل والشراب لجميع أذى الجسم وقياسات النمو بما في ذلك قياسات نسبة الوزن والطول والوزن بالنسبة للعمر والطول بالنسبة للعمر والوزن بالنسبة للعمر والوزن بالنسبة للعمر ومستوى الدهون ومستوى منتصف النزاف قبل.

ويعتبر عسل النحل.

نتناول كل طفيلة جرعة من العسل تبلغ 2 مل/كم/يوم عن طريق الفم لمدة 8 أسابيع، وتم إدراة الجرعة المحسوبة من العسل على ماء بنسبة 1:1 على التوالي. وذلك قبل تناولها مباشرة قبل الطقف قبل الإفطار حيث إن إبذاء العسل في الماء سوف يقلل فاعلية العسل كمضاد للUIKitوريBAL. ويتناسب في ضغط جرعة العسل. يتم إدراة جرعة العسل وفقًا لكل على حدة والإستقالة بها بعداً عن الضوء في وعاء مغلق جيداً لحين استخدامه وتم إعطاء الطفل 7 جرعات لكل إسبوع وكان هناك توصية بعد جرعة العسل إضافية خلال فترة الدراسة وخصم السعرات الحرارية للطفل اليومية من جميع السعرات الحرارية اليومية للطفل. حيث إن العسل المستخدم في هذه الدراسة هو عسل نحل خام غير معالج. ضغيفة الغربة، مصر، ثم تناولها مباشرة من قبل مربي النحل وتم إختياره بحلاً من جرائم.

في هذه الدراسة فيما يتعلق بالقياسات الإحصائية، بعد دورا التحليل عن تناول العسل كمكمل غذائي، أظهرت النتائج تقدم زيادة ذات

دالة إحصائية عالية بقيمة 2.000 في الوزن ودالة إحصائية بقيمة = 7.720 في محيط الورس عند مقارنتها بالقيم الفعلية لوزن ومحيط الورس بالنسبة للفحص بعد تناول العسل. فيما يتعلق بمختلف الأحماض الدهنية قصيرة السلسلة في الدم في دراستنا، وجد أن هناك لفرق بين

دالة إحصائية بعد تناول العسل كمكمل غذائي بقيمة إحصائية = 24.000 22.000 23.000 21.000 20.000 مع الببتيريك والأسوتيك والفورميز بالتبادل.