Enzyme activity of raw honey harvested from different localities of Kannad region, Aurangabad District (M. S.), India

Waykar Bhalchandra, Mahesh A Joshi and Nilesh Jawalkar

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Abstract
The present study deals with the enzyme activity of honey samples harvested from three different locations in the Kannad region of Aurangabad district (M. S.), India. Diastase and invertase activity was analyzed by using Schade et al., (1958) and Siegenthaler (1977) method respectively. Results clearly indicate that the honey harvested from agricultural and forest areas show the highest enzyme activity than the roadside area. The values of enzymes in honey were varied from location to location. This study also clearly demonstrates thin viat honey harvested from the Kannad region was fresh and unheated because the honey samples have more enzyme activity. The values of diastase and invertase were within the quality regulation limits proposed by Codex standards.

Keywords: Raw honey, diastase, invertase, enzyme activity, honey quality, Kannad region

Introduction
Honey is a sweet viscous food made by honeybees from the sugary secretions of plants, such as nectar by regurgitation, enzymatic activity and water evaporation as well as stored in wax structures called honeycombs [1]. Honey contains small amounts of different enzymes and the most important ones being diastase (α- and β-amylase), invertase (α-glucosidase), glucose oxidase, catalase and acid phosphatase, which comes from the nectar sources, salivary fluids and the pharyngeal gland secretions of the honeybee. The enzyme content in honey is one of the characteristics which make honey or its products different from other sweeteners. The biochemical composition of honey varies greatly and it mainly depends on the floral, regional and climatic conditions. Because of the great variation, a lot of research has been carried out to classify and identify the origin of honey in relation to its physical and biological properties. To our knowledge, the very limited study was carried out on the relationship between the biochemical (enzymes) and nutritional components of honey. Often, the major concern of honey consumers, regardless of honey origin, is the quality of honey. This relationship is very important for the food processing industry, particularly for those industries using honey as an ingredient in their food products [2].

The enzymes are closely related to the nutritional content and honey freshness. Even though enzymes are present in very small amounts, they have a significant effect on the quality of honey. This is because of the enzymes would significantly affect the protein content, free amino acid profile and acidity of honey samples. Mostly, enzymes found in honey samples were secreted from bee salivary fluids namely oxidases, catalases and amylases [3]. These enzymes would break down complex sugars into simple sugars such as fructose and glucose. Because of fermentation, the simple sugars might also be further catalyzed into alcohol and acetic acid under an appropriate amount of moisture content at the right temperature. Besides affecting the pH value, the activity of enzymes might change the flavour and aroma of honey after fermentation [2]. Since enzymes are present in trace amounts, many studies are likely to focus on sugar composition to determine honey origin [4]. Besides as macronutrient, the composition of monosaccharide and disaccharides as well as their ratios could be used to determine the degree of honey maturity. Usually, ripe honey samples have lower disaccharides such as sucrose and maltose content than those from honey harvested at an earlier stage. This is because most of the disaccharides have been converted into monosaccharides by the action of enzymes. Hence, the predominant sugars and their ratios are crucial parameters for honey characterization [5].
1) Diastase
It is a common name for the enzyme α-amylase. Its activity indicates the honey quality, which is used to determine if honey has been extensively heated during processing. It is found in the nectar and also added by the honey bee during the collection and ripening of nectar [8]. The diastase content of fresh, unheated honey is known to vary over a wide range. Any type of honey possesses several kinds of enzymes that play both nutritional and analytical roles in the product. It is the most critical honey enzyme capable of breaking down glycosidic linkages in oligo- and polysaccharides. The activity of this enzyme decreases with the time of storage and that of heating. This starch-digesting enzyme of honey is used as a quality indicator of honey because of its sensitivity to heat treatment [6].

2) Invertase
It is an enzyme which is widely distributed among plants and microorganisms and that catalyzes the hydrolysis of the disaccharide sucrose into glucose and fructose. Invertase (α-glucosidase) is an enzyme that is produced in the hypopharyngeal glands of the honeybee and which is a catalyst for one of the most important reactions in the transformation of nectar (sucrose) into honey (glucose and fructose) [9]. It is more sensitive than diastase to thermal treatment and storage of honey. Therefore invertase is the best parameter for the characterization of thermal treatment and storage time [8]. Hence, invertase activity is the quality indicator for the thermal treatment of honey and was used as a freshness indicator [9]. Therefore, the present study determines invertase and diastase activity of blended raw honey samples harvested from three different bee species from three different localities of the Kannad region of Aurangabad district (M. S.) India were indicated.

Materials and methods

Study area: The total area of Aurangabad district is about 10.07 lakh hector. Out of which 8.12 lakh hecter is under agriculture and 0.12 lakh is under forest area. Geographically, Kannad taluka of Aurangabad district is located at 20º 27’ N and 75º 13’ E. The average altitude of this area is 633 meter above sea level. Honey samples were collected from three different locations of Kannad taluka of Aurangabad district.

Collection of samples: Honey samples were collected from three bee species (Apis florea, Apis cerana indica and Apis dorsata) from three different locations of the Kannad region during October 2015 to September 2016. Total 23 different honey samples were collected: 9 from an agricultural area, 9 from roadside area and 5 from the forest area. Area wise honey samples were blended in equal quantity (100g each) and honey samples were put in airtight sterilized plastic containers. They were labeled, brought to the laboratory and stored at 0 - 4º C until analysis.

Determination of Diastase and Invertase: The diastase and invertase activity of honey samples was determined according to the procedure of Schade et al. (1958) and Siegenthaler (1977), respectively.

The calculation for Diastase Activity
The classical method for the determination of diastase activity is the method of Schade et al., (1958). There was a perfect correlation (r = 0.987) between the two measurements. Linear regression of y (diastase number) against x (ΔA420) yielded the following relation:

\[ DN = 28.2 \times \Delta A_{420} + 2.64 \]

The calculation for Invertase Activity
The amount of p-nitrophenol in μM produced during the test corresponds exactly to the amount of substrate in μ utilized. Therefore, the honey invertase activity can be calculated from the absorbance measured at 400 nm and is indicated in Invertase Number (IN).

\[ IN = 21.64 \times \Delta A_{400} \]

Results and Discussion

In the present study the diastase and invertase activity were determined in blended raw honey samples harvested from three different locations of Kannad taluka of Aurangabad district and obtained results were presented in the Table No.1 and Fig. No. 1 and 2.

The enzymes are important honey quality parameter and biological activity indicators. Honey naturally preserves small amounts of enzymes that have a huge impact on human life processes. Enzymes like invertase and diastase are very sensitive to heat and storage, they are also acts as freshness indicator of honey [9].

1) Diastase activity in honey
Enzyme diastase (amylases) breaks down starch into simple sugars. The activity of diastase in honey is affected by storage and is sensitive to temperature, which is used as an indicator of storage time/freshness and controls during the processing of the honey. Therefore proper heating and storage is of utmost importance to retain the market value of honey. The obtained results in our study for the diastase activity were summarized in table no. 1. The mean values of diastase number in the blended honey samples harvested from different locations of the Kannad region were within the range 5.35-7.91 (DN). Result showed that the mean diastase numbers in honey obtained from forest area are higher than in agricultural areas and roadside areas.

There was no significant difference in the values of diastase activity from species to species and location to location. All the collected honey samples were within the imposed limit of Codex [12, 13]. Many researchers carried out similar investigations. Yilmaz and Kufrevioolu, (2001) reported the mean value of diastase was 14.6 (DN) from Eastern Antonia [14]. Terrab et al., (2002) reported the diastase activity of Moroccan unifloral honey ranges between 0.18 to 236 Gº [15]. Serrano et al., (2004) determined the chemical and physical parameters of Andalusian honey and found the values ranges from 1.47 to 49.42 (DN) [16], Guler et al., (2007) studied the biochemical properties of honey to discriminate between pure and adulterate honey from Turkey and reported a mean value of 16.50 (DN) [17]. Serrano et al., (2007) reported diastase activity of Andalusian honey in the range of 3.99 to 49.42 (DN) [18], Silva et al., (2009) reported the values of diastase ranges from 3 to 38 (DN) from the Luso region (Portugal) [19]. Aloisi (2010) observed the diastase activity of honey from Chubut (Argentinian Patagonia) ranges between 3.90 to 39.28 Gothe units [20]. Estevinho et al., (2012) assessed the diastase value in the
range of 13.9 to 16.4 (DN) with a mean value of 15.3 (DN) from the Tras-Os-Montes regions of Portugal [21]. Buba et al., (2013) determined the diastase activity of honey samples from North-East Nigeria ranges between 8.00 to 13.00 (DN) [22]. Iftikhar et al., (2014) mentioned the diastase activity of local and imported brands of honey samples available in the Rawalpindi and Islamabad markets Pakistan ranged from 0.00 to 17.0 (DN) [23]. Chakir et al., (2016) reported the diastase activity of some honey produced from different plants in Morocco ranges between 4.30 to 29.60 (DN) [24]. Silva et al., (2017) observed the diastase activity of Portuguese honey from the Castelo Branco region in the range of 5.2 to 15.8 (DN) [25]. Goncalves et al., (2018) reported the diastase activity of selected Portuguese commercial monofloral honey samples ranges between 6.4-13.3 (DN) [26]. Bouhlali et al., (2019) determined the diastase activity of eleven monofloral honey samples produced in Morocco in the range of 7.40 to 29.29 (DN) [27]. Sajid et al., (2020) comparatively studied the diastase activity of fresh and branded honey from Pakistan in the range of 26.97 to 43.46 (DN) in fresh honey and 5.95 to 10.35 (DN) in branded honey [28].

Invertase activity in honey

Invertase activity is the quality indicator for the thermal treatment of honey and is used as a freshness indicator [9]. Invertase activity was determined in blended raw honey samples from three different localities in the Kannad region, and obtained results were summarized in table no. 1. The mean values of invertase activity in the blended honey samples harvested from the study area were within the range of 7.89-18.4 (IN). The mean invertase values in honey obtained from forest areas are higher than roadside areas as well as an agricultural areas. The variability in enzyme activity found in the different honey types was probably due to the nectar collection period (consequently the physiological stage of the colony); abundance of nectar flow and its sugar content (a high flow of concentrated nectar leads to lower enzyme content); the age of the bees (the glands of honeybees produce more digestive enzymes when they become a forager); pollen consumption, etc. [3]. Invertase activity has a great natural variation; its use has been proven in honey quality control.

In our study the values of invertase activity clearly demonstrate the significant difference from location to location. The obtained values of invertase activity for all the collected honey samples were within the imposed Codex limit [12, 13]. The enzyme invertase is more sensitive to storage as compared to the diastase in thermal treatment and storage of honey. Therefore invertase is a better parameter for the characterization of thermal treatment and storage time [9]. Many authors have expressed different opinions about the propriety of those parameters. Bonvhei et al., (2000) reported invertase activity in fresh and processed honey and recorded the values of invertase ranging from 4.04 to 46.2 (IN) [29]. Vorlova and Pridal, (2002) determined the invertase and diastase activity in the honey of Czech Provenience and reported the invertase values ranging from 0.8 to 25.9 (IN) with a mean value of 15.7 (IN) [30]. Serrano et al., (2004) studied the chemical and physical parameters of Andalusian honey and found the invertase number (IN) ranges from 0.20 to 50.5 (IN) [16]. Serrano et al., (2007) reported diastase activity of Andalusian honey in the range of 1.2 to 36.8 (IN) [18]. Silva et al., (2017) observed invertase activity of Portuguese honey from the Castelo Branco region in the range of 12.8 to 37.4 (IN) [25]. Boussaïd et al., (2018) reported the invertase activity of six Tunisian honey samples from various floral origins ranges between 46.25 to 184.68 IU (International Units) [31]. Sajid et al., (2020) comparatively studied the invertase activity of fresh and branded honey from Pakistan in the range of 58.55 to 81.9 (IN) in fresh honey and 3.10 to 9.66 (IN) in branded honey [28].

<table>
<thead>
<tr>
<th>Site of Collection</th>
<th>Diastase Activity (DN)</th>
<th>Invertase Activity (IN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agricultural Area</td>
<td>6.81±0.072</td>
<td>12.46±0.082</td>
</tr>
<tr>
<td>Forest Area</td>
<td>7.91±0.064</td>
<td>18.45±0.086</td>
</tr>
<tr>
<td>Road Side Area</td>
<td>5.35±0.041</td>
<td>7.89±0.063</td>
</tr>
<tr>
<td>Codex Alimentarius Standards, 1998 and 2019.</td>
<td>≤8</td>
<td>6.5-17.7</td>
</tr>
</tbody>
</table>

± indicates the Standard Deviation

Table 1: Invertase and diastase activity in blended raw honey of three bee species harvested from three different locations in the Kannad region of Aurangabad district.

Location Map

Fig: Map showing the locations of honey sample collection.
Conclusion
Overall we conclude that the honey harvested from agricultural and forest areas in the Kannad region shows higher invertase and diastase activity than roadside areas according to limits proposed by Codex Alimentarius Standards. Hence, the honey harvested from agricultural and forest areas has the highest nutritional value than the roadside area. It indicates honey freshness and the honey was unheated, which has great nutritional importance in the human diet.

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References


