Pollen types collected by *Tetragonisca angustula* (Hymenoptera: Apidae) in dry vegetation in Northeastern Brazil

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**Abstract.** Knowing the floral origin of the pollen collected by native bees in Neotropical ecosystems enables us to understand the dynamics of the interdependent relationships between the insects and the native flora. The objectives of this study were to: (i) investigate the spectrum of plant species from which pollen is collected by *Tetragonisca angustula* in caatinga (dry vegetation) areas in the semiarid region of Northeastern Brazil; (ii) identify pollen types that could be used as regional geographical markers and (iii) determine the pollen niche breadth and the pattern of use of floral resources by this stingless bee. In total, 23 samples of the pollen stored by *T. angustula* were collected monthly and subjected to acetolysis. Of the 45 pollen types identified that of *Prosopis juliflora* (Fabaceae) and *Solanum* (Solanaceae) were the most frequently represented in the samples and 19 were collected by the bees at both study sites. Species characteristic of caatinga, such as *Poincianella pyramidalis* (Tulasne) L.P. Queiroz and *Senna macranthera* (de Candolle ex Colladon) H.S. Irwin & Barneby, were among the pollen identified and they were used as indicators of the geographical origin of the samples. The average values of the pollen niche breadth (H') and equitability (J') indicate that *T. angustula* is a generalist and homogeneous forager of the floral resources in *caatinga*.

**INTRODUCTION**

Over the past several decades, deforestation, habitat loss and the introduction of exotic species have resulted in a decline in populations of native pollinators. This phenomenon has led the international scientific community to attempt to understand the implications of this decline for global ecosystems and propose mechanisms for reducing this decline (Allen-Wardell et al., 1998; International Pollinators Initiative, 1999; Biesmeijer et al., 2006; Freitas et al., 2009; Burkle & Alarcón, 2011; Giannini et al., 2012). As the environmental services provided by native bees in particular are important it is essential to understand those aspects of their biology, behaviour and ecology that can be used to develop conservation initiatives designed to maintain pollination (Allen-Wardell et al., 1998).

The pollinators are primarily female bees that collect pollen as the main protein source for themselves and their larvae (Michener, 2007). In the Neotropical region, however, it is difficult to carry out continuous and systematic studies of the flora that provide this protein diet for native bees. In Brazil, such studies have been ongoing for decades; however, they have not been done in all the biomes in Brazil (Barth, 2004; Borges et al., 2006).

Among the Brazilian biomes, the phytogeographic domain of *caatinga* is the largest gap in our botanical knowledge, largely because of the erroneous belief that this type of vegetation originated from the modification of another plant formation (Giulietti et al., 2004). According to Queiroz (2006), historically, the species diversity of the flora of the *caatinga* is considered to be low and include few endemics. It is suggested that the *caatinga* does not have an autochthonous flora and that most of its elements are derived from chaco and Atlantic rain forest (Rizzini, 1963; Andrade-Lima, 1981). These hypotheses were subsequently rejected (Prado & Gibbs, 1993). Recent studies indicate that the flora of the *caatinga* is surprisingly diverse (Giulietti et al., 2002, 2004, 2009; Albuquerque et al., 2012) and includes 4,320 species of angiosperms (Forzza et al., 2010), which have relationships with a range of pollinators including 187 species of native bees (Zanella & Martins, 2003). There is a mosaic of different ecosystems in the dry vegetation in the *caatinga*, which covers an area of approximately 800,000 km² and practically the entire semiarid climate region in Brazil, from the northeast through a region north of the State of Minas Gerais (Prado, 2003; Giulietti et al., 2004).

Among the Meliponini reported in the Brazilian *caatinga*, *Tetragonisca* sp. group *angustula* (Lateille, 1811) is one of the most well-known and studied groups. The geographical distribution of this group of stingless bees goes beyond the Brazilian territory, extending from Mexico to Argentina (Camargo & Pedro, 2012). Although the pollen collected by *T. angustula* has been studied in many different types of Brazilian vegetation
(Iwama & Melhem, 1979; Imperatriz-Fonseca et al., 1984; Carvalho et al., 1999; Morgado et al., 2011; Novais & Absy, 2013), few studies have been done in caatinga. Novais et al. (2006) performed a palynological study of the honey of this species collected in a hyperxerophytic caatinga region at the Canudos Biological Station in the northeastern micro-region of Bahia State. Recently, the pollen contents of honey from *T. angustula* colonies at Iaberaba and Ruy Barbosa in Bahia State were investigated (Novais et al., in press). However, there are no reports of palynological studies of the pollen stored by *T. angustula* foraging in caatinga vegetation.

Because there is no information on the floral spectrum foraged by *T. angustula* for collecting pollen in dry vegetation in the caatinga region of the Brazilian semiarid region we asked the following questions: (i) what pollen types reflect the flora visited by *T. angustula* in the caatinga of Bahia?; (ii) which of the various pollen types collected by this species can be used as regional geographical markers?, and (iii) does the pattern of use of pollen resources in the areas of caatinga investigated indicate a homogeneous or heterogeneous use of the local flora?

**MATERIAL AND METHODS**

**Study areas**

This study was carried out in the municipalities of Itaberaba (12°26′18.8″S, 40°13′12.7″W) and Ruy Barbosa (12°17′27.6″S, 40°30′03.0″W), which are located in the semiarid climate zone in the state of Bahia, Northeastern Brazil (Fig. 1). The annual precipitation in this area does not exceed 800 mm and the average temperature is approximately 25°C (EMBRAPA, 2013).

The most diverse plant families in the region include Fabaceae, Euphorbiaceae, Apocynaceae, Orchidaceae, Rubiaceae and Asteraceae, of which there are more than 120 recognized species (Cardoso & Queiroz, 2008). Depending on altitude, various plant formations can be distinguished locally, from caatinga with palm trees at lower altitudes (up to 450 m), to vegetation similar to savanna and areas of rupestrian fields (at approximately 900 m) (Cardoso & Queiroz, 2008).

**Sampling and laboratory processing**

Samples of pollen (23 in total) collected from stores in *T. angustula* colonies in the study areas were examined. At Itaberaba, samples were collected (*n* = 10) monthly during 2011, with the exception of July and November when there was a scarcity of pollen in the colonies. At Ruy Barbosa, the samples were collected monthly (*n* = 13) between October 2010 and December 2011, with the exception of June and November. Each month, the samples were collected from the same colony, mainly from the five pollen pots that had been sealed most recently by the bees, which were usually of a lighter colour than older pots. The material was collected from the food pots using pieces of disposable plastic straws (approximately 3 cm), which were packed in lidded pots, labelled and kept in a refrigerator until used in the laboratory procedures.

The protocol adopted was that of Novais & Absy (2013), which included drying the material in an oven at 40°C to stabilize the weight, dissolving in warm water and 95% ethanol, dehydrating in glacial acetic acid and acetolysis (Erdtman, 1960).

For each sample, three slides were prepared using glycerin gelatin and at least 500 pollen grains were quantified and identified using optical microscopy. The botanical identification of the pollen types was performed using catalogues (Roubik & Moreno, 1991; Silva, 2007) and the pollen library at the Laboratory of Plant Micromorphology of Bahia State University at Feira de Santana. Because each pollen type represents a morphological entity that potentially includes various species or genera (Joosten & de Klerk, 2002), the author was cited for each taxonomic entity referred to in the text.

After the microscopic analysis, the classes of occurrence (C.O.) were established based on the presence or absence of each pollen type in the samples as follows: *r* – rare (present in <10% of samples); *i* – infrequent (10–20%); *f* – frequent (21–50%); and *vf* – very frequent (>50%) (Jones & Bryant, 1996). These classes do not consider the number of pollen grains in each sample.

![Fig. 1. The map on the left indicates the location of Brazil (light grey) in South America (black). The map on the right shows the state of Bahia, indicating the study areas at Itaberaba (IT) and Ruy Barbosa (RB).](image-url)
This was calculated using the following equation:

\[ H' = - \sum p_i \ln p_i \]

In the above equation, \( p_i \) is the proportion of each pollen type \( i \) in the sample. This proportion is given by \( \frac{n_i}{N} \), where \( n_i \) is the number of grains of each pollen type \( i \) and \( N \) the total number of pollen grains in the sample.

Pielou’s equitability index \( (J’) \) (Pielou, 1977) was used to indicate the dynamics of the use of pollen resources by the bees. This was calculated using the following equation:

\[ J’ = \frac{H'}{H'_{\text{max}}} \]

In this equation, \( H'_{\text{max}} \) is the natural logarithm of the total number of different pollen types (\( S \)) in the sample; \( H'_{\text{max}} = \ln S \). The \( J’ \) index varies from 0 to 1. Values between 0 and 0.5 indicate a heterogeneous use of the local flora and those between 0.5 and 1 indicate a homogeneous use.

All analyses were performed using PAST (Palaeontological Statistics) software version 2.16 (Hammer et al., 2001).

**RESULTS**

Our data indicate that *T. angustula* used a large number of plant species as 45 pollen types are listed in the spectra (Tables 1 and 2). Of these 19 were common to both study areas, 10 were recorded only in the samples from Itabera and 16 only in the samples from Ruy Barbosa. In total, 29 pollen types belonging to 18 plant families were recorded in the samples from Itabera (Table 1) and 35 from 25 families in the samples from Ruy Barbosa (Table 2). Of the plant families the Fabaceae was the most frequently represented family, with eight morphologically different pollen types (Tables 1 and 2).
The main pollen types present in the samples collected at Itaberaba include the following: *Brosimum*, *Prosopis juliflora*, *Senna macranthera* and *Solanum* (Table 1). At Ruy Barbosa, the main types were *Brosimum*, *Heteropetys*, *Prosopis juliflora*, *Schinus*, *Senna macranthera* and *Solanum* (Table 2). These pollen types were present in more than 50% of the samples analyzed (“very frequent” class of occurrence) and greater than 10% in at least one of the samples. *T. angustula* very frequently collected pollen from *Prosopis juliflora* (Swartz) de Candolle. In addition to being present in all of the pollen samples analyzed from both study areas (*n* = 23), *P. juliflora* occurred with a frequency of greater than 10% in 20 of the samples (86.96%) (Tables 1 and 2). The second most represented pollen type was *Solanum*, which occurred in 90% of the samples from Itaberaba and 92.31% of those from Ruy Barbosa. The third most used pollen type at Itaberaba was *Senna macranthera*, which occurred in 80% of the sam-
samples from this municipality, followed by *Poincianella pyramidalis* (70%), *Mikania* (60%), *Brosimum* (60%), *Myrcia* (50%) and *Heteropterys* (50%). At Ruy Barbosa, *Brosimum* and *Schinus* each occurred in 84.62% of the samples, followed by *Senna* (76.92%) and *Heteropterys* (61.54%).

At Itaberaba, only the *Mikania* and *Poincianella pyramidalis* pollen types occurred at a frequency lower than 10% in more than 50% of the samples (Table 1). At Ruy Barbosa, the pollen types included in the “very frequent” class of occurrence demonstrated a greater than 10% frequency in at least one sample (Table 2). Although a large number of pollen types were recorded, less than 25% of these were found consistently at a high percentage (>10%) (Ramalho et al., 1985) in the *T. angustula* collections (Tables 1 and 2).

In 2011, eight pollen types were found in at least three consecutive months at Itaberaba, *Poincianella pyramidalis* (Jan–May), *Prosopis juliflora* (Jan–Dec), *Solanum* (Feb–Dec), *Senna macranthera* (Feb–Aug), *Zornia echinocarpa* (Apr–May), *Myrcia* (May–Aug), *Brosimum* (Jun–Dec) and *Mikania* (Sep–Dec). The occurrence of these pollen types in consecutive months was more evident in the second half of 2011 (Table 1).

At Ruy Barbosa, eight pollen types were also found in at least three consecutive months between Oct 2010 and Dec 2011: *Prosopis juliflora* (Oct 2010–Dec 2011), *Solanum* (Oct–Dec 2010 and Feb–Dec 2011), *Poincianella pyramidalis* (Dec 2010–Feb 2011), *Brosimum* (Jan–Oct 2011), *Geonoma* (Jan–Mar 2011), *Schinus* (Jan–Mar 2011), *Senna macranthera* (Feb–Aug 2011), *Zornia echinocarpa* (Apr–May 2011), *Myrcia* (May–Aug 2011), *Brosimum* (Jun–Dec 2011) and *Mikania* (Sep–Dec 2011). The occurrence of these pollen types in consecutive months was more evident in the second half of 2011 (Table 1).
In the months when the collections were performed at both study sites, the number of pollen types identified in the samples from Itaberaba each month was slightly lower than at Ruy Barbosa (Fig. 2, Tables 1 and 2). Furthermore, in these months, the number of pollen types recorded simultaneously in the spectra for the two areas varied from three (in Jan and Apr 2011) to eight (in Dec 2011) (an average of 4 ± 1.5 s.d., \(n = 9\)) (Fig. 2, Table 3).

In total, eleven pollen types were recorded in the samples of pollen collected by *Tetragonisca angustula* in the same months at Itaberaba and Ruy Barbosa, and include the families Anacardiaceae, Arecaceae, Asteraceae, Fabaceae, Malpighiaceae, Malvaceae, Moraceae, Myrtaceae, and Solanaceae (Table 3, Fig. 3).

The ecological analysis indicates that the average trophic niche breadth of *T. angustula* at Itaberaba was 1.10 ± 0.46 s.d. (\(n = 13\)). The month with the smallest niche breadth was Jan 2011 (\(H’ = 0.42\)) and the largest was Dec 2011 (\(H’ = 1.79\)) (Fig. 4, Table 1). At Ruy Barbosa, the average \(H’\) index was 1.11 ± 0.35 s.d. (\(n = 10\)), with minimum and maximum values recorded in Sep 2011 (\(H’ = 0.65\)) and Dec 2011 (\(H’ = 1.76\)), respectively (Fig. 4, Table 2).

The average equitability at Itaberaba was 0.50 ± 0.15 s.d. (\(n = 13\)), with the minimum and maximum recorded in Jan 2011 (\(J’ = 0.24\)) and Jun 2011 (\(J’ = 0.69\)), respectively (Fig. 4, Tables 1 and 2). At Ruy Barbosa, the

Fig. 3. Photomicrographs of some of the pollen types recorded in pollen stores of *Tetragonisca angustula* foraging in caatinga (seasonally dry tropical forest) vegetation in Northeastern Brazil. a – Brosimum, Moraceae; b – Geonoma, Arecaceae; c – Heteropterys, Malpighiaceae; d – Mikania, Asteraceae; e – Myrcia, Myrtaceae; f – Poincianella pyramidalis, Fabaceae; g – Prosopis juliflora, Fabaceae; h – *Senna macranthera*, Fabaceae; i – Solanum, Solanaceae. Bars = 10 µm.
average $J'$ value was $0.52 \pm 0.16$ s.d. ($n = 10$), with a minimum and maximum recorded in Sep 2011 ($J' = 0.31$) and Apr 2011 ($J' = 0.81$), respectively (Fig. 4, Tables 1 and 2).

The monthly $H'$ and $J'$ values for Itaberaba decreased or increased in parallel (Table 1). At Ruy Barbosa, however, this parallelism in behaviour was not consistent. In 2011, in Feb–Mar and Mar–Apr, opposite trends were recorded in the $H'$ and $J'$ values: the $H'$ index decreased when the $J'$ value increased and vice versa (Table 2).

**DISCUSSION AND CONCLUSIONS**

This study indicates that *T. angustula* collected pollen from a wide spectrum of flowers but especially from *Prospis juliflora* (Fabaceae) and *Solanum Linnaeus* spp. (Solanaceae), which makes up a major part of the pollen spectrum throughout most of the period studied. The consistent use of certain floral sources by the bees and other pollinators may reflect the ability of these insects to "learn" or "memorize" their floral characteristics, such as colour, odour, size, and patterns, which function as indicators of rewards, such as nectar, pollen or resin (Gonzalez et al., 1995; Chittka et al., 1999; Glover, 2007; Nicolson, 2007). Although the search for particular flower sources can often be inferred from pollen spectra, this fact does not necessarily imply that these species provide a greater amount of the resource than other species. Chittka et al. (1999) suggest that in the search for a preferred species, bees could neglect flowers of other species of plants that offer as much or more of the resource than the preferred plants. In addition, pollen is not always a reliable indicator of a nectar source because there are no in depth studies of how the availability of resources varies even within the same genus or of the floral and reproductive biology of a large number of species of plants in areas such as the *caatinga* (Iwama & Melhem, 1979; Machado & Lopes, 2004; Santos et al., 2010; Roubik & Moreno, 2013). Therefore, it is essential to carry out more field studies to ratify or rectify the inferences made based on pollen spectra (Roubik & Moreno, 2013).

The pollen grains in bee products provide clues to their geographical origin because the key plant species that characterize the region where they were produced can be identified (Maurizio, 1975; Louveaux et al., 1978). Giulietti et al. (2002) lists 318 species endemic to areas of *caatinga* in Northeastern Brazil. Of the genera identified in this study, 13 were represented in the pollen spectra recorded for Itaberaba and Ruy Barbosa, however, only a few of these can be considered to be geographical indicators of this type of vegetation, such as *Poincianella pyramidalis*, *Senna macranthera* and *Zornia echinocarpa* (Giulietti et al., 2004).

Queiroz (2008) reports that *Poincianella pyramidalis* (Tulasne) L.P. Queiroz var. *pyramidalis* is characteristic of areas of sensu stricto *caatinga*, where small-sized trees are predominant but do not form a continuous canopy, the trees and bushes exhibit xerophytic characteristics and the herbaceous plant layer is only present during the rainy season. *P. pyramidalis* var. *pyramidalis* is typical of the state of Bahia and adjacent regions of *caatinga* in Pernambuco and Alagoas States, mainly occurring in arboREAL *caatinga*. The majority of the insects visiting the flowers of *P. pyramidalis* seek nectar, although the volume of this resource per flower is limited, which may cause the visitor to forage from a larger number of flowers, thereby increasing the chance of effectively pollinating the species (Leite & Machado, 2009).

There are three different varieties of *Senna macranthera* (de Candolle ex Colladon) H.S. Irwin & Barneby occurring in areas of *caatinga* (Queiroz, 2008), of which only *S. macranthera* var. *micans* (Nees) H.S. Irwin & Barneby is reported by Cardoso and Queiroz (2008) in the region of the Serra do Orobó [Orobó Mountains] in the municipalities of Itaberaba and Ruy Barbosa. Pollen from the genus *Senna* Miller is recorded in various palynological studies carried out in areas of *caatinga* (Novais et al., 2009; Oliveira et al., 2010; Silva et al., 2012) and *S. macranthera* is an important source of pollen used by the bees in this type of vegetation (Maia-Silva et al., 2012). It is likely that *T. angustula* can collect the pollen remaining in the flowers of this species after their anthers have been shaken by the vibrations generated by larger bees, such as those of the genera *Xylocopa* Latreille and *Bombus* Latreille (Machado & Lopes, 2004; Maia-Silva et al., 2012).

Queiroz (2008) reports that *Zornia echinocarpa* (Moricand ex Meissner) Bentham occurs in *restinga* (coastal vegetation) and *caatinga* on sandy soil only in the state of Bahia. Cardoso & Queiroz (2008) confirm the presence of this species in the Serra do Orobó. Pollen from *Zornia echinocarpa* is among the dominant types in the pollen...
loads of *Apis mellifera* Linnaeus foraging in the *caatinga* region of Canudos, Bahia (Novais et al., 2010). In our study, *Zornia echinocarpa* pollen was present at a low level in the spectra. In contrast, pollen types indicating genera and species important for bees in the *caatinga* region are recorded, both in nectar [e.g., *Prosopis juliflora* (Fabaceae) and *Waltheria Linnaeus* (Malvaceae)] and pollen [e.g., *Solanum* (Solanaceae)] (Machado & Lopes, 2004; Santos et al., 2006; Maia-Silva et al., 2012). The *Prosopis juliflora* pollen type occurred most frequently in the samples studied. This species is not native to *caatinga*, however, it occurs spontaneously in various regions where agricultural activities have ceased (Queiroz, 2008). *Prosopis juliflora* may be pre-adapted to surviving in *caatinga* because of its rapid growth and resistance to drought (Sajjad et al., 2012). In the study areas, this species flowered practically throughout the entire year, constituting a continuous source of resources for local bees.

Other sources of pollen for bees in the study areas were the genera *Brossimum Swartz*, which is common in pollen spectra although predominantly an anemophilous species (Martins & Batalha, 2006), and *Virola Aublet* (Lena & Oliveira, 2006). Cesário & Gaglianone (2008) report that the simple flowers of *Schinus* Linnaeus are a source of pollen and nectar for various species of insect. *Heteropterys* Kunth, although considered to be a source of oil for Centridini bees in areas of *caatinga* (Aguiar et al., 2003), was also regularly visited during several months at Itaberaba and Ruy Barbosa and, therefore, is also a pollen source (Rocha-Filho et al., 2012).

Compared with that of other Meliponini, the trophic niche of *Tetragonisca angustula* is relatively small (H’ < 1.5) in the two areas studied (Carvalho et al., 1999; Oliveira et al., 2009; Ferreira et al., 2010). However, values close to those recorded in our study were obtained for *T. angustula* at other Brazilian locations. Novais & Absy (2013) investigated the pollen stored by *T. angustula* at Belterra (H’ = 1.04) and Santarém (H’ = 0.59) in the state of Pará, Carvalho et al. (1999) analyzed corbicular pollen stored by this species at Piracicaba, São Paulo (H’ = 2.70) and Morgado et al. (2011) investigated corbicular pollen stored by *T. angustula* foraging in the Ilha Grande, Rio de Janeiro (H’ = 0.43). It is likely that the small size of *T. angustula* determines the small size of its pollen niche in different ecosystems.

In general, the trends in H’ and J’ run parallel with one another. However, in some cases, this is not the case as was recorded from November to December 2010, at Ruy Barbosa. Although the value of H’ was the same (1.18) in both months, that of J’ increased (from 0.46 to 0.54), due to the decrease in the dominance of the pollen of *Prosopis juliflora*, which was the predominant pollen (71.01%) in November 2010. While the most dominant pollen type in December 2010 was *Solanum* pollen type (57.12%), but at a frequency lower than that recorded for *Prosopis juliflora* in the previous month. In contrast, in December 2010, the frequency of *P. juliflora* pollen was 17.97%. Thus, the co-dominance of the *Solanum* and *Prosopis juliflora* pollen types in December 2010 may explain the increase in uniformity recorded in the spectrum (J’).

Another example occurred from April to May 2011 when there was an increase in H’ and a decrease in J’. This decline was due to the dominance of *Solanum* pollen type (54.88%). In April 2010, *Prosopis juliflora* dominated the spectrum (54.91%), followed by *Solanum* (24.91%) and *Senna macranthera* (14.15%). In May 2010, the *Solanum* pollen type was followed by that of *Prosopis juliflora* (12.37%) and *Heteropterys* (11.67%). The lowest values of these last two types, compared to the frequencies of *Solanum* and *Senna macranthera* in April 2010, indicated a lower uniformity in the use of pollen resources in May 2010.

The uniform use of pollen sources was slightly lower at Itaberaba than at Ruy Barbosa. Carvalho et al. (1999) recorded that compared with *Apis mellifera* and three species of Meliponini of the genera *Nannotrigona* Cockerell, *Paratamona* Schwarz and *Plebeia* Schwarz, *T. angustula* foraged less uniformly at Piracicaba (São Paulo). The diversity of floral sources used by *Tetragonisca angustula*, together with the consistency exhibited by this species in the use of these resources, confirmed the generalist habit of this species. The small size of this insect may result in reduced energy expenditure (Imperatriz-Fonseca et al., 1984) per flower while foraging and probably per minute of flight, making various plants equally attractive for supplying food rewards. We may hypothesize that flowers offering a low quantity of resource (pollen or nectar) should be almost as attractive to small bees as those which offer high pollen/nectar rewards. This can be advantageous if they are competing with other foragers for the same resources.

Therefore, we conclude that *T. angustula* exploits a significant number of species of plants in the study areas, which confirms that this species has a polylectic foraging strategy. A number of species of plants can be used as indicators of *caatinga* vegetation, such as *Poincianella pyramidalis* and *Senna macranthera*. The pollen types *Prosopis juliflora* and *Solanum* were the most frequently recorded in the samples, demonstrating their prioritization as sources of pollen by *Tetragonisca angustula* at Itaberaba and Ruy Barbosa. The ecological analyses indicated a homogeneous use of floral resources by *T. angustula* in the areas of *caatinga* studied. High homogeneity implies low dominance of a single pollen type in the pollen spectrum. That the exotic *Prosopis juliflora* is such an important source of pollen for this native bee may have implications for the interactions between native and non-native species in this and other eco-regions that contain generalist species with broad niches.

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