A peer-reviewed version of this preprint was published in PeerJ on 14 March 2016.

View the peer-reviewed version (peerj.com/articles/1819), which is the preferred citable publication unless you specifically need to cite this preprint.

Rincón-Rabanales M, Vargas-López LI, Adriano-Anaya L, Vázquez-Ovando A, Salvador-Figueroa M, Ovando-Medina I. (2016) Reproductive biology of the biofuel plant *Jatropha curcas* in its center of origin. PeerJ 4:e1819
https://doi.org/10.7717/peerj.1819
Reproductive biology of the biofuel plant *Jatropha curcas* in its center of origin

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**Abstract**

In this work, we studied the main characteristics of flowering, reproductive system and diversity of pollinators for the biofuel plant *Jatropha curcas* (L.) in a site of tropical southeastern Mexico, within its center of origin. The plants were monoecious with inflorescences of unisexual flowers. The male flowers produced from 3062 to 5016 pollen grains (266-647 per anther). The plants produced fruits with both geitonogamy and xenogamy, although insect pollination significantly increased the number and quality of fruits. A high diversity of flower visiting insects (36 species) was found, of which nine were classified as efficient pollinators. The native stingless bees *Scaptotrigona mexicana* (Guérin-Meneville) and *Trigona (Tetragonisca) angustula* (Latreille) were the most frequent visitors and their presence coincided with the hours when the stigma was receptive. It is noteworthy that the female flowers open before the male flowers, favoring xenogamy, which may explain the high genetic variability reported in *J. curcas* for this region of the world.

**Keywords:** pollination, geitonogamy, xenogamy, stingless bees.
1 Introduction

*Jatropha curcas* (L.) (Euphorbiaceae), possibly native to Mexico and Central America [1,2], is considered the most promising non-edible plant for the production of biofuels. Many countries have established programs for its commercial cultivation [3]. In the Mesoamerican region, where the greatest genetic diversity of populations of *J. curcas* has been found [2,4-6], monocultures are being established, for example in Guatemala and the Mexican states of Chiapas and Michoacan [7].

When establishing new extensive crops, a multitude of factors must be taken into account, among which stands out reproductive biology [8], i.e., knowledge about flowering, phenological behavior, sexual system, and fruit and seed production.

Studies on floral biology and pollination ecology of *J. curcas* have been conducted, mainly in regions where this species is exotic, as in India [9-16]. Following from these studies, it is known that *J. curcas* is a monoecious species that presents geitonogamy and xenogamy; however, the first process, also called self-pollination, is prevalent. This could explain the low genetic diversity in Asian germplasm. It is possible that environment is affecting the prevalence of geitonogamy or xenogamy [17].

In the center of origin and diversity of *J. curcas* no such studies have been conducted, thus the sexual system and mode of pollination are unknown. Such knowledge could help design strategies to increase crop productivity; i.e. promoting efficient pollinators in commercial plantations. The aim of this study was to characterize the pollination process in *J. curcas* under field conditions at a site in southeastern tropical Mexico.

2 Material and Methods

2.1 Study site and biological material

The study was conducted within the living fences of a farm plot at Soconusco, Chiapas,
Southern Mexico (14.5036 N, 92.1704 W, 58 meters above sea level). In this zone the average annual temperature is 31°C, the average annual humidity is 80 %, and the average of rainfall is 2600 mm (Weather Station: 769043 MMTP of the Water National Commission, Mexico). The soil type was andosol with pH 5.7, 2.5 % of organic matter and 0.2 % of total nitrogen. Plants were selected based on their appearance (healthy and abundant foliage; approximately a 10 % of the plants of the fence had foliar damage) and location (in a sunny area), on a 600 m transect. The living fence was 10 years old and underwent annual pruning. According to people living in the surrounding communities, the studied plants are considered toxic, with occasional cases of children intoxication caused by seed ingestion.

2.2 Floral phenology

Flowering and fruiting dynamics were studied in 10 plants every 14 days during one year. We determined the average number of inflorescences per primary branch. In five inflorescences per plant, randomly selected, male and female flowers were counted. To estimate the time of flower anthesis and stigma receptivity, in other 10 plants, an inflorescence in the stage of flower buds was marked and observed daily (30 d) at intervals of every 10 min from 0700 h to 1200 h. To estimate the production of pollen and ovules, inflorescences were collected from other 10 plants with closed flowers, and 20 female flowers (FF) and 20 male flowers (FM) were randomly selected. Pollen grains were extracted from FM and mounted in glycerinated gelatin on a slide. Pollen was quantified using a stereomicroscope and the average number of pollen grains was estimated per anther and per flower. In FF the number of carpels and the number of ovules per carpel was counted. With these data the pollen to ovule ratio was estimated.

In order to comprehend the reproductive process, five pollination treatments were established: 1) geitonogamy or artificial pollination with pollen from the same inflorescence (GEI), 2) xenogamy (XEN) or artificial pollination with pollen from another plant, 3) apomixis (APO),
which was performed by removing the male flowers and placing non-toxic white glue
(Resistol®, Mexico) on stigma, 4) excluding pollinators (ExP), and 5) open pollination (OpP).

For each of the treatments 20 inflorescences were employed, one per plant (in total 20 plants
for this experiment), which were covered, except in OpP, with tulle mesh bags of 1 mm mesh
size. Fertilization was checked 14 days after pollination and the number of mature fruits per
treatment was quantified at 55 days.

2.3 Insect flower visitors and pollinators

To determine insect pollinators and visitors, observations were made on ten inflorescences
from independent plants, from 0800 h to 1700 at intervals of 10 min. For this, the time of
arrival at the flower was taken into account, the time they stayed, the resource used (nectar or
pollen), and movement among flowers of the same inflorescence and between inflorescences
of the same or another plant.

In addition insects visiting the flowers of J. curcas were collected on another living fence,
with similar characteristics, but distanced 500 m from the study site to avoid interference with
the previous experiments. For this part of the study, we sampled the area of influence of 100
plants. The insect collection was performed with entomological nets from 0600 h to 1800 h.
All insects captured were examined under stereoscope, dissected into head and thorax, and
identified using Ayala [18] and Michener [19] taxonomic keys. Moreover, the pollen adhered
to the legs of the insects was identified by microscopy.

The visiting insects were classified as: a) efficient pollinators, b) occasional pollinators, c)
accidental pollinators, or d) pillagers. We used the following criteria: 1) number of individuals
collected during different times of the day, 2) recurrence and time of visit to the male and
female flowers, 3) behavior observed on the flowers, and 4) presence of J. curcas pollen (pure
or mixed) on different parts of the body.

2.4 Statistical analysis
To understand differences between visitor groups in the frequency and time of visit to male and female flowers, the Chi-square test was applied. The number of fruits, and the quality of these in the different treatments were compared by analysis of variance (ANOVA) and Tukey test ($\alpha = 0.05$).

3 Results

3.1 Flowering and anthesis time

The plants studied had three flowering peaks (80%) in the months of April, May and September. In the period from December to February the plants showed no flowering. Subsequent to that, there were three periods of peak fruiting in the months of May, June and October. There were no fruits in the months from December to March.

It was found that, under the study conditions, *J. curcas* produces an average of 1.25 inflorescences per branch, and the number of female flowers per inflorescence was 1-11, with an average of 2.2. Meanwhile, the number of male flowers ranged from 35-198 flowers per inflorescence (mean: 106.7). It was found that the proportion of female/male flowers was 1:60. Pollen production by anther ranged from 266-647 pollen grains (mean: 475.1) and per flower was 3062-5016 pollen grains (mean: 4224.4). The proportion of pollen grains per ovule was 1408:1.

The male and female flowers begin to open at 0800 h, presenting the maximum aperture of female flowers (64.29%) and male (55.75%) at 0900 hours. The stigma was receptive at the time of 1000-1200 h.

3.2 Pattern of flower opening

Flowers regularly opened over an average period of 15 days (Figure 1). The pattern showed that the female flowers are the first to open and this process continues for eight days. Male flowers started opening two days after the female flowers, and lasted up to 13 days, with the
highest peaks between days 8 and 10.

3.3 Diversity of insect visitors

The variety of insect visitors to the flowers of *J. curcas* came from 36 species, which were grouped into four orders, 12 families and 16 genera as recognized (Table 1). Hymenoptera were the most diverse (75% of the species) and dominant (72.6% of the relative abundance) group, followed by Diptera (diversity of 19.4%; dominance of 26.3%).

We found three types of pollinators: a) accidental, including the fly Thachinidae sp. 2, Vespidae sp. 1, Vespidae sp. 2 and one species of Cerambicidae; b) occasional, comprising 14 species that included bees, ants, and wasps; and c) efficient, composed nine species, of which most were bees and two species of flies. The remaining insects were considered pillagers or nectar robbers (Table 1). Some of the individuals were carrying pure pollen loads of *J. curcas* in different parts of the body, while others carried *J. curcas* pollen mixed with other pollen types *[Ageratum aff. houstonianum* (Mill.), *Acacia aff. corninga* (L.) Willd., *Tridax aff. procumbens* (L.), *Zea mays* (L.)]. Pollen from other species represented less than 10% of the total loads (Table 2).

3.4 Frequency of visit

The activity of insects visiting the flowers of *J. curcas* was continuous, starting from 0600 h until shortly after 1800 h, showing a markedly bimodal behavior (Figure 2). The main activity peaks coincided with increased secretion of female flower nectar during daylight 0700-0900 h. The greatest wealth of insects was recorded at 0900 h and 1000 h (S=8), registering the highest peak at 1400 h (S=9) (Figure 2).

We found differences in the frequency and time of insects visiting the female flowers (*χ²* = 21.78, *p* < 0.01) and male (*χ²* = 39.69, *p* < 0.01). Insects observed in the marked panicles were *S. mexicana*, *T. (Trigona) fulviventris* (Guérin), *T. (Trigona) fuscipennis* (Friese), *Agapostemon nasutus* (Smith), *Augochlora (Augochlora) quiriguensis* (Cockerell),
Augochlora aurifera (Cockerell), Augochlora (Augochlora) smaragdina (Friese), Halictus (Seledonia) hesperus (Smith), Tachinidae sp. 1, Eristalis aff. persa (Williston), Camponotus (Mayr) and Vespidae sp. 1. Compared with other groups, the bees visited more flowers (43.2%) and stayed longer in resource foraging (38%). The second most important group was the Diptera with 39.3% (frequency) and 31% (time of visit). Specifically, bees visited a larger number of female flowers (55.6%) than male flowers (46.4%). Both bees and flies spent more time to visiting female flowers, while Vespidae sp. 1 had a preference for the male flowers. The Diptera foraged exclusively nectar, while bees collected nectar and pollen.

3.5 *Jatropha curcas* reproductive system

We found differences in the production and quality of fruits and seeds from the different reproductive systems of *J. curcas* (p < 0.001). The highest percentage of fruit set was recorded in free pollination treatments (OpP: 86.3 ± 2.2) and in xenogamy (XEN: 84.3 ± 6.3), being statistically equal between them; followed by the treatments with exclusion of pollinators (ExP: 18.1 ± 7.2) and geitonogamy (GEI: 16.2 ± 7.3). In the treatment of apomixis five fruits were formed, of which four were aborted and only one reached maturity (n = 55 female flowers). The only apomictic fruit had a high fresh weight, which contrasted with a low seed fresh weight, due to its thick endocarp.

Regarding the number of fruits that reached maturity, the OpP and XEN treatments were statistically superior to all other treatments (p < 0.001). The type of reproduction also influenced the quality of fruits and seeds, as the longest fruits were recorded in the OpP and XEN treatments (p < 0.001) and heaviest seeds occurred in OpP treatment (Table 3).

4 Discussion

Some aspects of the floral biology of *J. curcas* in the Mexican tropics, such as the proportion of male and female flowers (1:60), differ from those reported in studies conducted with
germplasm from other geographic areas. For example, Raju & Ezradanam [10] reported in India that an inflorescence could produce 1 to 5 female flowers and 25-93 male flowers, with a ratio of male flowers to female flowers of 1:29. Also, Pinto et al. [20] found 4-12 female flowers and 87-222 male flowers (relation: 1:20). This variable is not preserved and the differences depend on the genetic material, geographic region, climate, nutrition, time, and cultural practices, among other factors, which makes it a highly variable feature [11,13,20].

Pollen production per anther and per flower was higher than that reported in other studies, such as Bhattacharya et al. [11], who found that each flower produced 1617 pollen grains, with a ratio of pollen:ovule (P:O) of 539:1. In this regard, Cruden [21] mentions that the P:O ratio is an indicator of the reproductive system. In the case of *J. curcas* the P:O relation is very high, which could favor the xenogamy. On the other hand, there could be a compensatory mechanism of pollen loss caused by the constant arrival of insect visitors to the inflorescences, i.e., reflecting a low efficiency in pollen transfer.

The anthesis of male flowers and female flowers under the conditions of this study occurred at 0700 h, which coincides with the findings of Raju & Ezradanam [10], who reported that the flowers open daily between 0530-0630 h. Kaur et al. [16] reported that the male flowers open between 0600 h and 0700 h, while the female flowers open shortly after (0700-0800 h). The time it takes for the stigma to be receptive (1-2 h) is similar to that reported by Bhattacharya et al. [11]. The opening pattern of flowers in the morning was related to attracting insects, because the availability of resources (nectar and pollen) is significantly higher in the morning.

The flowering stage of the plants studied occurred from March to November, which was consistent with that reported by Sukarin et al. [9], who recorded two flowering peaks, one in May and one in November. Instead, Joker & Jepsen [22] observed that flowering occurs during the dry season with two flowering peaks, these authors mention that the plants bloom throughout the year in permanent wet conditions. While the fruiting stage occurred from April
to December, contrary to that reported by Toral et al. [23], who recorded that the fruits are produced in winter when the plant loses its leaves.

The wealth of insects foraging on flowers of *J. curcas* located in the southeastern area of the Mexican tropics was high (Table 2). However, of the 32 species of potential pollinators, not all were efficient as pollinators of *J. curcas*, because not all visited the flowers of both sexes, or they did not transport pollen on their body, or they did not coincide with the period of stigma receptivity. According to these observations, it was determined that an efficient pollinator for *J. curcas* was one that (i) visited several flowers of *J. curcas* during foraging [15], (ii) frequently shifted from one flower to another [15,24], (iii) transported specific pollen abundantly on its body, and (iv) was observed sliding some part of its body on the receptive stigma.

The most abundant insects were Hymenoptera and Diptera, which is consistent with some previous studies in other areas [25], but contrasts in the case of Diptera that have been cited as efficient pollinators of flowers of *J. curcas*, but which are not always present or are less diverse and abundant [10,11]. There is a high diversity and abundance of bees (19 species) and flies (9 species). Bees were generally small to medium size (5-10 mm) and particularly the native stingless bees were the most diverse, abundant, and with morphological characteristics correspondent to the floral syndromes of *J. curcas*. In general, the Diptera had body sizes 7-14 mm and abundant pilosity on the body, but their behavior does not allow cataloging them as efficient pollinators. Two exceptions were *Eristalis* sp. and Tachinidae sp. 1, which frequently visited male flowers looking for fresh nectar and transported pure pollen stuck onto their body for more than nine hours a day. They were observed simultaneously visiting female flowers for nectar during the period of maximum stigma receptivity facilitating xenogamy and geitonogamy, whereby they were classified as efficient pollinators. Our observations are consistent with studies by Raju & Ezradanam [10], who found that the
Diptera *Chrysomya megacephala* (Fabricius) was an efficient pollinator that promoted xenogamy and geitonogamy. Conversely, Rianti et al. [15] reported only *Eristalis tenax* (L.) as an infrequent visitor and not an efficient pollinator for *J. curcas* in West Java.

Native stingless bees *S. mexicana*, *T. (T.) angustula*, *T. (T.) fulviventris*, *T. (T.) fuscipennis*, *H. (S.) hesperus* and *A. nasutus* are of small body size (5-8 mm), with special structures for transporting pollen and a great quantity of specialized hairs [26]. It is possible that these characteristics enable them to efficiently perform the flow of pollen to the stigma of *J. curcas*, as has been observed in other species of bees [10,11,15,27].

In the present study, *A. mellifera* (L.) was infrequent in relation to other efficient pollinators (n = 3) and recorded only between 0800 and 0900 h. Additionally, two individuals transported *J. curcas* pollen in great abundance, in the head, chest and legs, while one of them transported pollen from different plant species. This result is contrary to most previous studies where *A. mellifera* has been registered as the most efficient pollinator in *J. curcas* [10-12,14-16,28].

We classify this species as occasional pollinator for *J. curcas* at this site in the Mexican tropics, and it can perform both geitonogamy and xenogamy.

The polylectic behavior of stingless bees has been reported in previous work, and this is due to the different sources of pollen that they forage for food. However, it has been reported that these bees can temporarily present an oligolectic strategy, taking advantage of a single source of food, as it occurs with *Cocos nucifera* (L.), *Mangifera indica* (L.), *Carica papaya* (L.), *Citrus limon* (L.) and *Capsicum annuum* (L.) [29]. Our data show that the efficient pollinating bees have oligolectic or monolectic behavior, at least during the season studied. Among those bees are included *S. mexicana* and *T. (T.) angustula*, which live in nests organized within tree trunks, have reduced foraging dispersion, and intensively use the floral resources available, for which they have a potential as inducers of pollination of extensive crops of *J. curcas*. 
The increased production of fruits (86.3%) was recorded in open-pollinated plants and in xenogamy treatment (84.3%), consistent with the findings of other authors [10,12,13]. However, a low fruit set (50-53%) is also reported in open-pollinated flowers [12]. Regarding the phenomenon of apomixis, we found a very low rate (2.5%), similar to that found by Santos et al. [30] in a semiarid region in Brazil (5%). In contrast, Bhattacharya et al. [12] and Kaur et al. [16] reported that apomixis might be responsible for the formation of more than 30% of the fruits. Chang-Wei et al. [13] reported a moderate effect of apomixis (12%).

The genetic diversity of *J. curcas* in this Mesoamerican area and particularly in this region of Mexico is high [1,4-6], and this may be largely due to the efficient pollination service by native stingless bees, to the strong trend for protogyny leading to pollination by xenogamy, and to the low prevalence of geitonogamy. Based in our results, strategies to improve the productivity of *J. curcas* in commercial plantations could be designed, for example increasing artificially the population of efficient pollinators as the stingless native bee *S. mexicana*.

**Acknowledgements**

We thank the Hermanos Ávalos Ranch, in Tapachula, Chiapas, for allowing us to conduct the study in living fences on their property. We are thankful with El Colegio de la Frontera Sur, for the support to identify Halictideae and stingless bees.

**References**

[1] Ovando-Medina I, Adriano-Anaya L, Vázquez-Ovando A, Ruiz-González S, Rincón-Rabanales M, Salvador-Figueroa M. Genetic diversity of *Jatropha curcas* in Southern Mexico. In: Bahadur B, Sujatha M, Carels N, editors. Jatropha, challenges for a new energy crop, Vol 2, New York: Springer; 2013, p. 219-50.

[2] Salvador-Figueroa M, Magaña-Ramos J, Vázquez-Ovando JA, Adriano-Anaya ML,
Ovando-Medina I. Genetic diversity and structure of *Jatropha curcas* L. in its centre of origin. Plant Genet Res 2014; doi:10.1017/S1479262114000550

[3] Renner A, Zelt T. Global market study on Jatropha. London: Gexsi LLP; 2008, p. 1-30

[4] Basha SD, Francis G, Makkar HPS, Becker K, Sujatha MA. Comparative study of biochemical traits and molecular markers for assessment of genetic relationships between *Jatropha curcas* L. germplasm from different countries. Plant Sci 2009;176:812-23.

[5] Ambrosi DG, Galla G, Purelli M, Barbi T, Fabbri A, Lucretti S, Sharbel TF, Barcaccia G. DNA markers and FCSS analyses shed light on the genetic diversity and reproductive strategy of *Jatropha curcas* L. Divers 2010; 5:810-36.

[6] Ovando-Medina I, Espinosa-García FJ, Núñez-Farfán JS, Salvador-Figueroa M. State of the art of genetic diversity research in *Jatropha curcas*. Sci Res Essays 2011;6:1709-19.

[7] Ovando I, Adriano L, Salvador M, Ruiz S, Vázquez A. Piñón (*Jatropha curcas*): bioenergía para el desarrollo de Chiapas. Biotecnol Agrop Biodivers Chiapas 2009;2:1-25.

[8] Silva C, Torezan H. Reproductive biology of tropical plants. In: Claro K, Oliveira P, Rico-Gray V, editors. Tropical biology and conservation management. Oxford; UNESCO 2008.

[9] Sukarin W, Yamada Y, Sakaguchi S. Characteristics of physic nut, *Jatropha curcas* L. as a new biomass crop in the tropics. Japan Agric Res Quart 1987;20:302-3.

[10] Raju SAJ, Ezradanam V. Pollination ecology and fruiting behaviour in a monoecious species, *Jatropha curcas* L. (Euphorbiaceae). Curr Sci India 2002;83:1395-98.

[11] Bhattacharya A, Datta K, Kumar SD. Floral biology, floral resources constraints and pollination limitation in *Jatropha curcas* L. Pak J Biol Sci 2005;8:456-60.

[12] Dhillon RS, Hooda MS, Handa AK, Ahlawat KS, Kumar Y, Subha SH, Singh N. Clonal propagation and reproductive biology of *Jatropha curcas* L. Indian J Agroforest 2006;8:18–27.
[13] Chang-Wei L, Kun L, You C, Yong-Yu S. Floral display and breeding system of Jatropha curcas L. Forest Stud China 2007a;9:114-19.

[14] Quin Y, Ping PD, Biao DZ, Liang W, Xiang SQ. Study on pollination biology of Jatropha curcas (Euphorbiaceae). J South China Agricult Univ 2007;28:62–6.

[15] Rianti P, Suryobroto B, Atmowidi T. Diversity and effectiveness of insect pollinators of Jatropha curcas L. (Euphorbiaceae). J Biosci 2010;17:38-42.

[16] Kaur K, Dhillon GPS, Gill R. Floral biology and breeding system of Jatropha curcas in north-western India. J Trop Forest Sci 2011;23:4-9

[17] Heller J. Physic nut, Jatropha curcas L. Promoting the conservation and use of underutilized and neglected crops. Rome: IPGRI; 1996, p. 10-11

[18] Ayala R. Revisión de las abejas sin aguijón de México (Hymenoptera: Apidae: Meliponini). Folia Entomol Mex 1999;106:1-123

[19] Michener CD. The bees of the world. in: Johns Hopkins Univ Press, editors. Baltimore; 2007. p. 122-832

[20] Pinto AC, Pimenta S, Oliveira B, Batista D, Oliveira H. Biologia floral e polinização artificial de pinhão-manso no norte de Minas Gerais. Pesq. Agropec. Bras.2009;44:1073-77.

[21] Cruden RW. Pollen–ovule ratios: conservative indicator of breeding systems in flowering plants. Evol 1997;31:32-46.

[22] Joker D, Jepsen J. Jatropha curcas L. Seed Leaflet 2003;83:1-2

[23] Toral OC, Iglesias JM, Montes OS, Sotolongo JA, García S, Torsti M. Jatropha curcas L., una especie arbórea con potencial energético en Cuba. Pastos Forrajes 2008;3:191-207.

[24] Free JB, Williams IH. The pollination of crops by bees. Apimondia Pub House: Bucharest; 1977, p. 1-14
[25] Luo CW, Huang ZY, Chen XM, Li K, Chen Y, Sun Y. Contribution of diurnal and nocturnal insects to the pollination of *Jatropha curcas* (Euphorbiaceae) in southwestern China. J Econ Entomol 2011;1:149-154

[26] Roubik DW Ecology and natural history of tropical bees. New York: Cambridge University Press; 1989, p. 43-70

[27] Atmowidi T, Rianti P, Sutrisna A. Pollination effectiveness of *Apis cerana* Fabricus and *Apis mellifera* Linnaeus in *Jatropha curcas* L. (Euphorbiaceae). Biotropica 2008;15:129-34

[28] Chang-Wei L, Kun L, You C, Yong-Yu S, Wen-Yun Y. Pollen viability, stigma receptivity and reproductive features of *J. curcas* L. (Euphorbiaceae). J. Northwest Plant 2007b;27:1994-2001.

[29] Martínez E, Cuadriello J, Ramírez E, Medina M, Sosa M, Melchor J. Foraging of *Nannotrigona testaceicornis, Trigona (Tetragonisca) angustula, Scaptotrigona mexicana* and *Plebeia* sp. in the Tacaná region, Chiapas, Mexico. Grana 1994;33(4-5):205-17.
Table 1. Potential pollinators of *Jatropha curcas* in the region of Soconusco, Southern Mexico

| Order      | Family  | Genus      | Species                  | Type of forage | Type of visitor | Relative abundance (%) |
|------------|---------|------------|--------------------------|----------------|------------------|------------------------|
| Hymenoptera| Apidae  | *Apis*     | *mellifera* Linneo       | 1, 2           | OP               | 1.1                    |
|            |         | *Trigona*  | *fulviventris* Guérin    | 1, 2           | EP               | 7.3                    |
|            |         | *Trigona*  | *fuscicarinella* Friese   | 1, 2           | EP               | 1.1                    |
|            |         | *Nannotrigona* | *perilampoides* Cresson | 1, 2           | OP               | 0.4                    |
|            |         | *Scaptotrigona* | *mexicana* Guérin-Meneville | 1, 2          | EP               | 30.5                   |
|            |         | *Tetragonisca* | *angustula* Lepeletier   | 1, 2           | EP               | 7.3                    |
|            |         | *Oxytrigona* | *mediorufa* Cockerell    | 1              | OP               | 0.4                    |
|            |         | *Melipona*  | *beechei* Bennett        | 1, 2           | OP               | 0.4                    |
|            |         | *Melipona*  | *solani* Cockerell       | 1, 2           | OP               | 0.4                    |
|            |         | *Ceratina*  | *capitosa* Smith         | 1, 2           | OP               | 0.4                    |
|            |         | *Tripeolus* | sp. Robertson            |                |                  | 0.4                    |
| Halictidae |         | *Agapostemon* | *nasutum* Smith         | 1, 2           | EP               | 7.3                    |
|            |         | *Augochlora* (Augochlora) | *quiriquensis* Cockerell | 1, 2           | OP               | 0.7                    |
|            |         | *Augochlora* (Oxystoglossella) | *aurifera* Cockerell   | 1, 2           | EP               | 0.7                    |
|            |         | *Augochlora* (Augochlora) | *smaragdina* Friese    | 1, 2           | OP               | 0.4                    |
|            |         | *Halicitus* (Halicitus) | *ligatus* Say          | 1, 2           | OP               | 0.4                    |
|            |         | *Halicitus* (Seladonia) | *hesperus* Smith       | 1, 2           | EP               | 9.1                    |
|            |         | *Lasioselis* (Dialictus) | sp. 1 Robertson        | 1, 2           | OP               | 0.4                    |
|            |         | *Lasioselis* (Dialictus) | sp. 2 Robertson        | 1, 2           | OP               | 0.4                    |
| Formicidae |         | *Camponotus* | sp. 1 Mayr               | 1              | OP, PI           | 0.4                    |
|            |         | *Crematogaster* | sp. 1 Lund              | 1              | OP, PI           | 0.4                    |
|            |         | *Crematogaster* | sp. 2 Lund              | 1              | OP, PI           | 0.4                    |
|            |         | *Dolichoderus* | sp. 1                  | 1              | PI               | 0.4                    |
|            |         | *Dolichoderus* | sp. 2                  | 1              | PI               | 0.4                    |
|            |         | *Formicididae* | sp. 3                  | 1              | PI               | 0.4                    |
|            |         | *Vespidae*  | sp. 1                   | 1              | AP, PI           | 0.4                    |
|            |         | *Vespidae*  | sp. 2                   | 1              | AP, PI           | 0.7                    |
|            |         | *Vespidae*  | sp. 1                   | 1              | PI               | 0.4                    |
| Diptera    |         | *Diptera*   | sp. 1                   | 1              | PI               | 0.4                    |
|            |         | *Syrphidae* | *Eristalis*             | 1              | EP               | 7.3                    |
|            |         | *Tachinidae* | sp. 1                  | 1              | EP               | 17.0                   |
|            |         | *Tachinidae* | sp. 2                  | 1              | AP, PI           | 0.4                    |
|            |         | *Syrphidae* | sp. 1                  | 1              | PI               | 0.4                    |
| Coleoptera |         | *Coleoptera* | *Cerambycidae*        | sp. 1          | AP, PI           | 0.7                    |
|            |         | *Hemiptera* | *Fulgoridae*           | sp. 1          | PI               | 0.4                    |

AP: accidental pollinator, EP: efficient pollinator, PI: pillager, OP: occasional pollinator. 1: nectar; 2: pollen.
| Species               | Individuals collected (n) | Type of pollen loads (Number and percentage) |
|-----------------------|---------------------------|---------------------------------------------|
|                       |                           | Pure loads       | Mixed loads      | Without loads    |
| Scaptotrigona mexicana| 84                        | 55 (65.5%)       | 17 (20.2%)       | 12 (14.3%)      |
| Tetragonisca angustula | 19                        | 14 (73.7%)       | 1 (5.3%)         | 4 (21.0%)       |
| Trigona fulviventris  | 19                        | 14 (73.7%)       | 2 (10.5%)        | 3 (15.7%)       |
| Trigona fuscipennis   | 3                         | 3 (100%)         | -                | -               |
| Halictus hesperus     | 25                        | 15 (60%)         | 3 (12%)          | 7 (28%)         |
| Agapostemon nasutum   | 19                        | 7 (36.9%)        | 2 (10.5%)        | 10 (52.6%)      |
| Tachinidae sp. 1      | 49                        | 31 (63.3%)       | 1 (2.0%)         | 17 (34.7%)      |
| Eristalis sp.         | 19                        | 14 (73.7%)       | 2 (5.3%)         | 4 (21.0%)       |
| Apis mellifera        | 3                         | 1 (33.3%)        | 1(33.3)%         | 1 (33.3%)       |
Table 3. Comparison of characteristics of fruits and seeds obtained from different pollination treatments in *Jatropha curcas* in the Soconusco region, Chiapas, Mexico

|                      | OpP  | XEN  | GEI  | ExP  | APO* | F**  | p     |
|----------------------|------|------|------|------|------|------|-------|
| Fruits per inflorescence (n) | 4.88 a | 4.20 a | 0.88 b | 1.00 b | 0.10 c | 21.02 | 0.001 |
| Fruit diameter (cm)    | 2.94 a | 3.04 a | 2.82 ab | 2.64 b | 2.9 | 2.98 | 0.035 |
| Fruit length (cm)      | 3.29 a | 3.26 ab | 2.98 bc | 2.85 c | 3.1 | 4.73 | 0.004 |
| Fruit fresh weight (g) | 12.90 a | 13.10 a | 12.42 a | 10.07 b | 13.86 | 4.48 | 0.005 |
| Seeds (n)              | 2.68 a | 2.77 a | 2.71 a | 2.16 b | 3.0 | 2.80 | 0.004 |
| Seed fresh weight (g)  | 1.65 a | 1.21 b | 1.30 b | 0.97 c | 1.21 | 20.95 | 0.001 |

OpP: open pollination, XEN: xenogamy, GEI: geitonogamy, ExP: excluding pollinators, APO: apomixis.

*Due to the reduced number of fruits, the apomixis treatment was not included in most of the ANOVA tests.

**One way ANOVA and Tukey tests were performed. Different superscripts letters in a row denote statistical differences.
**Figure 1.** Opening dynamic of female and male flowers in *Jatropha curcas* in the Soconusco region, Chiapas, Mexico.
Figure 2. Daily dynamics of insects visiting *Jatropha curcas* flowers in the Soconusco region, Southern Mexico.