- 1 Original article Biological control
- 2 Natural biocontrol of fruit flies by parasitoids in indigenous hosts: a
- 3 perspective for the control of populations within and outside the orchards.
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Abstract

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Fruit flies (Diptera Tephritidae) are a major pest to fruit production in Africa. Most efforts to control these pests are focused directly on the orchards where fruit damage has been recorded. However, fruit flies are generally able to develop on alternative indigenous fruits from which they continuously recolonize the orchards, challenging the ability to efficiently control these pests. In the natural and semi-natural environments, parasitic Hymenoptera remain the primary means of keeping populations of fruit flies at a reasonable level, but little is known about the diversity, the distribution and the ability of these parasitoids to effectively control populations. In this study, we explored the diversity and abundance of parasitoids of fruit fly pests in indigenous and exotic cultivated fruits of two separate regions of South Africa: North-East (Limpopo and Mpumalanga provinces) and South-West (Western Cape Province). A total of 16 fruit species were collected and incubated to obtain emergence of fruit flies and their associated parasitoids. Ceratitis capitata (Wiedemann), Ceratitis rosa sensu lato Karsch, Ceratitis cosyra (Walker) and Bactrocera dorsalis (Hendel) were reared from the incubated fruit. In the Western Cape Province, only C. capitata and C. rosa sensu lato were recorded. Our results show an important contrast in the distribution of fruit fly parasitoid species between regions and between indigenous fruit types. Fopius ceratitivorus Wharton and Psyttalia humilis (Silvestri) (Hymenoptera: Braconidae) dominated parasitism in the north-eastern part of the country, while fruit fly parasitoids were almost absent on fruit flies in the Western Cape Province. Parasitism rate of fruit flies ranged from 0 to ca 50% depending on fruit species and locality within a region. The exact drivers of this variation in parasitism rate remain unclear, but smaller fruits generally seem to allow better control of infesting fruit flies by parasitoids than larger ones. The fruits of some indigenous trees (*Berchemia discolor* (Klotzsch) Hemsl.; *Syzygium cordatum* Hochst. ex Krauss and *Garcinia livingstonei* T. Anderson) are hosts of high densities of both fruit flies and their parasitoids. These species are good candidates for the application of augmentoria for the mass production and release of biological control agents.

Key words

Natural biological control, Fruit flies, Tephritidae, Alternative hosts, Sub-Saharan Africa

1. Introduction

Fruit-infesting fruit flies (Diptera, Tephritidae) are pests occurring worldwide that cause substantial damage to fruit production (Ekesi et al., 2016). In addition to the losses directly caused by the development of the larval stages in the fruits, numerous fruit fly species have a quarantine status and can strongly impact the export market of the country in which they occur (José et al., 2013). In Sub-Saharan Africa, the horticultural sector is affected by serious losses caused by native species (*Ceratitis capitata* (Wiedemann), *Ceratitis cosyra* (Walker), *Ceratits rosa* Karsch and *Ceratits quilicii* De Meyer, Mwatawala & Virgilio), but also alien species (*Bactrocera dorsalis* (Hendel), *Bactocera zonata* (Saunders)) that have invaded, established and spread rapidly through this region (De Meyer et al., 2007; Khamis et al., 2009; De Villiers et al., 2016; Manrakhan, Venter & Hattingh, 2015). Across many African countries, the total financial losses due to these pests are substantial and absence of management can lead to the complete loss of production (Ekesi et al., 2016).

In Africa, the traditional management of fruit flies using pesticides is challenged by the strict residue levels applied in export markets, in the larger context of the recognized detrimental effect of these compounds on human and ecosystem health (Bourguet and Guillemaud, 2016). The management practice has therefore evolved toward alternative methods such as bait application technique (BAT) using bio-pesticides, sterile insect technique (SIT) or orchard sanitation methods (OS) (Klungness et al., 2005; Ekesi and Billah, 2007; Manrakhan and Addison, 2014; Ekesi et al., 2016). Among OS methods, the augmentoria consist in the collecting of infested fruits and their storage in a container with a mesh retaining the fruit flies and allowing the biocontrol agents (parasitic hymenoptera) to emerge (Klungness et al., 2005). This simple approach contributes to control the population density of pest fruit flies and is usually used in combination with other control methods. All these methods are mostly deployed in the infested orchards or in the direct vicinity due to the cost and human resources needed for their application. However, fruit-infesting Tephritidae are polyphagous and

they are able to establish population reservoirs in neighbouring indigenous fruits (De Meyer et al., 2002; Copland et al., 2009; Aluja et al. 2014; Threon et al., 2017; Grové et al., 2017). Due to the dispersal ability of the adult stage (Moxley et al., 2017), these reservoir populations can spill over into orchards, resulting in limited efficiency of the above mentioned methods. Therefore, sustainable management of these pests also requires an understanding of the drivers affecting their abundance in natural and semi natural environments, within the framework of an area-wide integrated pest management (IPM) approach (Mau et al., 2007).

In natural and semi natural environments, fruit fly populations are naturally controlled by co-evolved parasitic Hymenoptera mainly belonging to the subfamily Opiinae (Wharton and Gilstrap, 1983). The Afrotropical region features a wide range of opiine species associated with tephritids, and an important and sustained effort has been made to characterize this diversity for biocontrol purposes (Wharton and Gilstrap, 1983; Wharton, 1999; Wharton, 2009; Wharton & Yoder, 2018). In addition to native species, several alien species were introduced in an attempt to improve the control of populations of fruit fly in orchards and crops, sometimes with remarkable success (see Ekesi et al., 2016 for a review). These species are koinobiont endoparasitoids which lay their eggs in the flies' eggs or first instar larvae and emerge as adults from the puparium, leading to the death of the host fly (Rousse et al., 2005).

There is little information, however, on how efficiently the Opiinae and other parasitoids are controlling the populations of fruit flies in natural and semi natural environments in Africa. Surveys were conducted only on wild olives in southern Africa to estimate the natural control of populations of *Bactrocera oleae* (Rossi) by parasitoids in non-crop habitat (Neuenschwander, 1982; Mkize et al., 2008). For most fruit-infesting flies, there is lack of data on how the parasitoids are controlling these populations in alternative indigenous fruits. As a result, it is currently unclear if indigenous fruits are reservoirs only for the fly population or for both the flies and parasitoids, and to what extent and for what reason. This information is crucial because it will determine how habitats should be managed to improve the control of fruit fly populations at a regional scale (Landi et al., 2000).

In this study, we (i) explored the diversity and distribution of parasitoids of fruit infesting flies in South Africa, and we (ii) made a preliminary estimation of the ability of these parasitoids to control fruit flies in wild indigenous or exotic ornamental and cultivated fruits. The potential use of indigenous fruit trees as reservoirs of parasitoids for the application of augmentative techniques is discussed.

2. Materials and Methods

2.1. Study area and fruit sampling

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Fruits were sampled between 2013 and 2018 in two separate and climatically different regions located in the North-East (Mpumalanga and Limpopo Provinces) and the South-West (Western Cape Province) of South Africa. The Western Cape province (WC) has a Mediterranean-type climate characterized by winter rainfalls, with cold, wet winters and hot, dry summers (ARC, 2014) that is suitable for the production of deciduous fruits. In contrast, the provinces of Mpumalanga and Limpopo (M&L) are characterised by a subtropical climate with summer rainfall and hot, wet summers and cold, dry winters (Government SA, 2018) which is more suitable for the production of tropical fruits. Fruits were collected in 18 localities, 5 in M&L and 13 in WC (Table 1). Exotic cultivated fruits were sampled in abandoned, organic orchards or home gardens where no chemicals, potentially affecting wasp populations, were sprayed (including neurotoxic bio-pesticides such as Spinosad). Fruits were collected on the trees at the appropriate time of maturation, and from the ground to maximize the range of parasitoids recorded (Eitam and Vargas, 2007). Several collecting events (hereafter called replicates) were conducted for each fruit type in different places and/or time of the year. Between 1 and 10 sampling replicates were carried out for each fruit type. A total of 16 fruit types were sampled, with 7 being wild indigenous, 3 exotic ornamental, and 7 exotic cultivated on a commercial scale (Table 1). In the WC Province, the natural vegetation surrounding orchards is of the Fynbos type (characterised by proteoid, ericoid and restioid plants), which is a very dry and woody vegetation. Apart from wild olives, that are not focal hosts for fruit infesting Ceratitis spp., there was no alternative wild hosts available for these species. Therefore, only exotic cultivated and ornamental fruits (Mespilus sp.) could be collected in this province. After collection, fruits were counted, weighed and placed in plastic boxes with sterile sand at the bottom to allow the larvae to emerge and pupate. Only one layer of fruit was placed in each box to limit excessive liquid accumulation that would be detrimental for the emergence of flies and their parasitoids. Plastic boxes were placed in incubators consisting of a Perspex box of 1x1 meter. The incubator was covered by black material except at the base of the collecting tube placed on top, in order to allow emerging insects to go into it. Incubators were maintained in at 25°C in a climatic chamber for a period of 3-4 weeks (photoperiod 16:8, L:D). All emerging flies and parasitoids were collected every two days, killed in a freezer and/or directly preserved in 96% ethanol before they

2.2. Species identification

were identified and counted.

Species identification of fruit flies based on morphology was made according to Virgilo et al. (2014) and De Meyer et al. (2016). Ceratitis rosa and C. quilicii are reported as C. rosa sensu lato, because some samples were collected and identified before the description of C. quilicii (De Meyer et al., 2016). The species of parasitoids were identified following Wharton & Yoder (2018), Wharton and Gilstrap (1983) and Wharton (1999). Some species in the subfamily Opiinae comprise species complexes (Rugman-Jones et al. 2009), therefore we also sequenced the standard barcode fragment of the mitochondrial cytochrome oxydase I (COI, Hebert et al., 2003) in order to cross-validate morphological identifications. DNA was extracted from a leg or a fragment of the abdomen of specimens using a DNeasy Blood & Tissue kit (Qiagen, Hilden, Germany), according to the manufacturer's protocol. PCR amplification were performed using the standard primers for barcoding (mitochondrial cytochrome c oxidase subunit I) invertebrates: LCO1490: 5'-GGTCAACAAATCATAAAGATATTGG-3' and TAAACTTCAGGGTGACCAAAAAATCA-3' (Folmer et al., 1994). PCR reactions were carried out on a Mastercycler® Nexus (Eppendorf, Hamburg, Germany) in a volume of 10 μL PCR mix containing 5 μl of Multiplex Master Mix (Qiagen, Hilden, Germany), 0.8 μl primers (Forward and Reverse at 2 μM) and 2 μ l of DNA. The PCR conditions were as follows: initial DNA denaturation at 94°C for 15 minutes, followed by 10 cycles of 30 s each at 94°C, 1 min at 60-50°C (touchdown of -1°C per cycle) and 30 s at 70°C, followed by 30 cycles of 30 s at 94°C, 1 min at 55°C, and 1 min at 72°C with a final extension of 20 min at 72°C. The PCR products were sequenced by Eurofins Genomics (http://www.eurofinsgenomics.eu/). Barcode sequences were aligned using CodonCode Aligner V.3.7.1. (CodonCode Corporation, Centerville, MA, USA), and checked to identify the presence of pseudogenes using standard detection methods (Haran et al., 2015). Each sequence was then aligned with reference barcode sequences of parasitoids of Fruit fly (Rugman-Jones et al. 2009; Haran et al. 2018). All voucher specimens were mounted on card or placed in 96% ethanol and were deposited in the South African Museum collections (SAMC, Cape Town). Additional material was deposited at CBGP, Montpellier, France (doi.org/10.15454/D6XAKL).

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2.3. Fly infestation and parasitism rates

The fly infestation rate was considered as the mean number of fruit flies emerging from 1 kg of fruit across replicates. Parasitism rate of fruit flies for a specific fruit type was computed as the ratio of parasitoids over the total number of flies and parasitoids (each representing one parasitized fruit fly) emerging from this fruit across sites. Parasitism rates were estimated only for solitary parasitoids in which a single adult parasitoid emerged from a fly puparium, and for parasitoid taxa where we were confident that they were associated with Tephritidae. A reliable estimation of parasitism rate is

usually conducted on a large number of replicates of fruit samples from a single locality over several years. This allows controlling the natural seasonal fluctuations of the parasitoid populations. Due to the exploratory nature of this study, and the limited number of spatial and temporal replicates, only an estimation of parasitism rate is provided. Therefore, the standard deviation of the mean parasitism rates were not computed, and only higher and lower values of parasitism rates from single replicates are reported.

3. Results

A total of four fruit fly species were recorded on all incubated fruits: *Ceratitis capitata, C. rosa sensu lato, C. cosyra* and *B. dorsalis,* with only the first two species found in the Western Cape province of South Africa. The infestation rates of fruit fly were higher in the M&L provinces than in the WC province, with an average number of flies of 213.36 and 2.72 per kg of fruit, respectively. Five species of parasitic Hymenoptera emerged from incubated fruits: two species of Opiinae (Braconidae): *Fopius ceratitivorus* Wharton, 1999 and *Psyttalia humilis* (Silvestri, 1913), one species of Alysiinae (Braconidae): *Asobara* sp. Förster, 1862, one species of *Aganaspis* Lin, 1987 (Figitidae), and *Tetrastichus giffardii* Silvestri, 1913 (Eulophidae). The identification of *P. humilis* was confirmed by the barcode sequences of specimens from South Africa that showed 99% similarity with a Genbank sequence available for this species (accession number: EU761026.1).

The parasitoid *Aganaspis* sp. was the only species recorded in both WC and M&L provinces of South Africa. The opiine species *F. certitivorus* and *P. humilis* and the Alysiine *Asobara* sp. were only recorded in the M&L provinces, while *T. giffardii* was only found in the South-West. No representatives of Opiinae were found in infested fruit samples of the Western Cape Province. Within the North-Eastern sites, *F. ceratitivorus* was only collected around Nelspruit (Mpumalanga province), while *P. humilis* was collected in both Mpumalanga and Limpopo provinces (Levubu, Letsitele town, Nelspruit). Across all fruit samples, these two species never co-occurred in the same batch of fruit. Mean parasitism rate of fruit flies by Opiinae ranged from 0% (*Carissa spinarum* L., *Psidium gujava* L. and *Sclerocarya birrea* (A.Rich.) Hochst.) to 38.5 % (*Syzygium cordatum* Hochst. ex Krauss) among fruit kinds (Table 2). Estimations of parasitism rate reached 54 % in a single replicate of fruit incubation of *Syzygium jambos* (L.) Alston. *Psyttalia humilis* and *F. certitivorus* showed variations in estimated parasitism rate, both sometimes not recovered or recorded at low level (mean of 2.65% and 5.96%, respectively), or reaching higher rates (mean of 22.38% and 38.5 %, respectively) among fruits collected.

Except for *Syzygium jambos* and *Carissa spinarum*, the smaller fruits (average weight of individual fruit below 10 g) allowed a higher parasitism rate than the larger ones (mean weight above 10 g),

with an estimated mean parasitism rate of 21.46% and 8.42%, respectively. The two opine *P. humilis* and *F. ceratitivorus* emerged together with all four species of fruit flies recorded in the study (Table 2).

4. Discussion

The environment surrounding an orchard is critical for the management of populations of polyphagous fruit flies. Indigenous hosts are known to be potential reservoirs of fly populations and their associated parasitoids in Central America (Ovruski et al. 2005, Aluja et al. 2014, Montoya et al. 2016). But, little quantitative investigation of parasitoids of fruit flies in indigenous hosts has been conducted in Africa in general (but see Copeland et al., 2009; Moxley et al., 2017). This study provides the first insights into the potential for wasp parasitoids to effectively control populations of fruit flies out of orchards in South Africa.

Among all the species of parasitoids that emerged from the fruits collected, only *Fopius ceratitivorus*, *Psyttalia humilis* and *Tetrastichus giffardii* are confirmed fruit fly parasitoids. Apart from a few exceptions (Ovruski et al. 2005; Tormos et al., 2013), the species belonging to the subfamily Alysiinae and the family Figitidae emerging from fruits are presently considered as parasitoids of Drosphilidae that are co-infesting the fruits (Wharton & Yoder, 2018). The two species belonging to these two groups (*Asobara* sp. and *Aganaspis* sp. respectively) will not be considered for further discussion.

The three parasitoid species we found on fruit flies contrast with the species assemblage obtained by Manrakhan et al. (2010) on *Ceratitis* species attacking berries of *Coffea* species in South Africa (Mpumalanga Province). In this study, we found a large representation of *P. humilis. Psytallia perproxima*, *Bracon celer* and *Tetrastichus giffardianus* were not recovered in this study. In contrast, *F. ceratitivorus* and *T. giffardi* were not reported on *Ceratitis* spp. attacking berries of *Coffea* species while they were found, sometimes at a high rate, in the present study. This difference in species composition might be due to differences in sampling localities, where parasitoid faunal assemblages may vary at a local scale, or due to a difference in attraction of parasitoids between infested fruits and berries of *Coffea* species.

4.1. A contrasted distribution pattern of parasitism of fruit flies in South Africa

This study highlighted a remarkable difference of parasitism rate of fruit flies by their parasitoids among two climatically different regions of South Africa. The two major Tephritidae obtained on various fruits (*Ceratitis capitata* and *C. rosa* s. l.) are two native species widely distributed and abundant across South Africa (De Villiers et al., 2013; Karsten et al., 2015; De Meyer et al., 2016).

While in the north-eastern areas, *C. capitata* and *C. rosa* s.l were parasitised by the dominant opiine parasitoïds (*F. ceratitivorus* and *P. humilis*), in the South-West regions these parasitoids were largely absent in the fruit sampled where these two fruit fly species were reared. Indeed, among the total of 3430 fruits belonging to 7 species infested by fruit flies collected in the WC province, not a single opiine specimen emerged. The eulophid *T. giffardi* was the only parasitoid emerging from *Ceratitis* spp. in this province and it was recorded only once on Medlar (*Mespilus* sp.).

Species of fruit flies, including *C. capitata* and *Bactrocera* sp. are known to have large climatic niches (Weldon et al., 2018) that may be wider than their parasitoids used in biocontrol (Lane et al., 2018) even at reduced scale (Etiam et al. 2004). It can be hypothesized, for *F. ceratitivorus* in particular, that the climatic conditions in the Western Cape are not appropriate for this species to develop on its host. Indeed, the genus *Fopius* is mainly distributed in the tropics (Wharton, 1999; Wharton and Yoder, 2018). The temperate winter-rainfall climatic conditions found in the Cape region are probably not suitable for the persistence of *Fopius* species in this area.

The case of *P. humilis* is different. This species is widely distributed in South Africa, from the North Eastern part of the country, the southern coast (Jh, pers obs), to the WC province where it has been mostly recorded from *Bactrocera oleae* attacking olives (Wharton and Yoder, 2018). Even though this species is present across the two sampled localities, it is found in much higher abundance on a variety of fruit in the North East, whereas the WC populations attcking olive fly do not seem to be able to shift effectively onto *Ceratitis* species attacking cultivated and ornamental fruit sampled in this province. Further investigations are needed to explore whether populations of *P. humilis* are specialists on a specific host-fly or fruit type in South Africa.

Fopius ceratitivorus and P. humilis never co-occurred in the same batch of fruit, while being distributed in the same regions. This result suggests a competitive exclusion of the two species, in line with the study of Wang et al. (2008) that reported at individual level a systematic exclusion of Psyttalia species by F. ceratitivorus when sharing the same host. It is therefore possible that at local scale and for specific fruit types, the early acting species F. ceratitivorus (oviposition in the eggs of the host) can completely supress the population of a later attacking species such as P. humilis (oviposition in larvae of the host). More generally, the absence of co-occurrence of the two species in a batch of fruit suggests contrasted performances of the two species that are commentary for the control of fruit flies in indigenous hosts.

4.2. Fruit flies are unequally controlled by parasitoids in indigenous fruits

Fruit flies encountered in this study are using a large range of indigenous fruits to complete their life cycle in tropical Africa (Copeland et al., 2009; Grové et al., 2017). Our results show a contrasting control efficacy by their associated parasitoids across fruit type in South Africa. In fruit of some families: Apocynaceae, Anacardiaceae and some Myrtaceae, opiine parasitoids seem unable to control fruit flies, suggesting that some indigenous or ornamental trees can play a role of sole fruit fly multiplier (Aluja et al. 2014). In contrast, these parasitoids were found attacking fruit flies at a substantial rate in fruit of other families: Clusiaceae, Rhamnaceae and some Myrtaceae. The exact drivers of this variation is difficult to determine as they may encompass a large range of parameters. Parasitoids usually follow the phenology of their hosts, track the larvae using plant kairomones, or oviposition marks of the flies. It is interesting to note that several fruit types that displayed either no parasitism, or a rather high parasitism rate of fruit flies were collected in the same localities. Therefore, these variations cannot be due to local absence of some parasitoids species. This observation rather suggests that some fruits might not be attractive for the opiine parasitoids, or alternatively their physical structure may prevent the parasitoids from efficiently locating the immature stages of fruit flies for oviposition.

Over all fruits sampled, the smaller fruits generally allowed a higher parasitism rate than the larger ones. Other studies showed similar results in North America (Wang et al. 2009a, b) or tended to confirm this trend in South Africa, with similar parasitism rates of fruit flies recorded from small fruits such as *Coffea* berries and wild olives (Mkize et al., 2008; Manrakhan et al., 2010). This observation is in line with the oviposition strategy of the genus *Psyttalia* that lays its eggs in the larval stages of fruit flies. In this case host larvae might be difficult to reach with their ovipositor in large fleshy fruits. However, this explanation does not apply to the genus *Fopius* that target the eggs of fruit flies at the surface of the fruits directly after the oviposition of fruit flies. It can also be hypothesized that opiine parasitoids are laying through oviposition marks on fruits, as has been documented in other braconid species (Stelinski et al., 2010). In this case, numerous small fruits would allow more oviposition events than a few large ones for an equal amount of resource reward for fruit fly larvae.

Three fruit tree species (*Berchemia discolor*, *Garcinia livingstonei* and *Syzygium cordatum*) showed a high infestation by fruit flies (about 800, 100 and 60 flies per kg of fruit respectively) that were themselves strongly parasitized by opiine species (about 22, 34 and 40 % of estimated parasitism rate respectively). The presence of these trees around orchards is notable, as they may play an attractant role for fruit flies, and hence for the parasitoids controlling them resulting in the development of a relatively high reservoir population of parasitoid wasps in an adjacent habitat. They would therefore contribute to the reduction of natural populations of fruit flies as described by

Aluja et al. (2014). In practice, however, the parasitism rate estimated still suggests that these trees would release substantial volumes of non-controlled fruit flies near the orchards. Therefore, it seem more appropriate to combine these trees with augmentative techniques, such as the harvesting of fruits and their placement in augmentauria or simply the placement of appropriate nettings under these in order to allow an important augmentative release of parasitoids together with a strong reduction of fruit fly populations. As an illustration, our results show that in some localities, one kg of the Brown Ivory fruits (*Berchemia discolor*) placed in an augmentoria can release about 300 parasitoid specimens while retaining about 800 fruit fly individuals. Other estimations reached 20,000 parasitoids per tree (Aluja et al. 2014). This approach for production of parasitoids may be more cost effective than traditional laboratory mass rearing for field release. More generally, these trees can potentially be used as "decoy trees", attracting and retaining fruit flies out of the orchard, killing them and releasing parasitoids using an augmentoria derived technique. This approach has economic potential as it could be used to limit both orchard infestations and to reduce wild populations of fruit flies.

5. Conclusion

This study showed a contrasting distribution of the natural biocontrol agents of fruit flies across two climatically disparate regions in South Africa, with a higher diversity and higher efficacy of parasitoids in the sub-tropical summer-rainfall part of the country in comparison to the area with a temperate, winter-rainfall climate. This survey also suggests that some indigenous trees are host for the flies and their parasitoids while some species seem to host only non-controlled populations of fruit flies. This suggests that indigenous trees around orchards could be selected and potentially developed to contribute to the reduction of fruit fly populations in natural and semi-natural environment using augmentative methods. More investigation is needed across the rest of Africa in order to determine the contributory role played by indigenous trees towards pest fruit fly control. More generally, a better characterization of the underlying drivers affecting biocontrol efficiency of fruit flies by the opiine parasitoids is needed, such as adaptation or preference of the parasitoid wasps for specific host flies, host plants, or climatic conditions.

Table captions

Table 1. Details of fruits samples incubated from the Limpopo, Mpumalanga and Western-Cape provinces of South Africa. **Dates**: Months and Years of collecting of fruits; **Repl.**: number of sampling

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replicates for each fruit kind in one locality; Avg fruit: mean number of fruit collected per replicate; Avg weight: Mean total fruit weight (Kg); AVg flies: mean number of flies obtained; Avg flies/ kg: mean number of flies per Kg of fruit incubated. Pres. Parasit: presence of confirmed parasitoids of Fruit flies in the fruit sample. Fruits with a * refer to exotic cultivated species and with a ** to exotic ornamental species. Italic values indicates results obtained for simple replicates. Table 2. Parasitism rate of fruit fly infesting indigenous fruits in South Africa. T: total number of fruit collected among replicates, W: total weight of fruit collected (g), Sz: average fruit weight (g), %: parasitism rate (%) among replicates, R- & R+: lower and higher parasitism rate obtain in single replicates. The fruit fly recorded from each fruit reared are reported. Acknowledgements We acknowledge the students Monique James and Bianca Stead (Stellenbosch University) for their help with the sampling and the rearing of Fruit flies and parasitoids. We thank the Western Cape Nature Conservation Board (South Africa for authorization to collect specimens (Permits N° CN44-30-4229). We also thank Aruna Manrakhan (Citrus Research International) for interesting discussions on early versions of the manuscript. This research was conducted for the project Opi-SYST, funded by the program Agreenskills (UE) and Hortgro STONE and Hortgro POME (South Africa). **Competing interests** Declarations of interest: none. References Aluja, M., Sivinski, J., Van Driesche, R. Anzures-Dadda, A., Guillén, L. 2014. Pest management through tropical tree conservation. Biodivers. Conserv.23, 831–853. https://doi.org/10.1007/s10531-014-0636-3 Bourguet, D., Guillemaud, T. 2016. The Hidden and External Costs of Pesticide Use. In E. Lichtfouse (ed.), Sustainable Agriculture Reviews, Sust. Agr. Rev. 19. DOI 10.1007/978-3-319-26777-7_2 Copland, R.S., Wharton, R.A., Luke, Q., De Meyer, M., Zenz, N., Machera, P., Okumu, M. 2006. Geographic Distribution, Host Fruit, and Parasitoids of African Fruit Fly Pests Ceratitis anonae, Ceratitis cosyra, Ceratitis fasciventris, and Ceratitis rosa (Diptera: Tephritidae) in Kenya. Ann. Entomol. Soc. Am. 99, 261–279.

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| Province of SA | locality | coordinates | fruit species | Dates | repl. | Avg. fruit | Avg. weight (kg) | Avg. flies | Avg. flies/kg | Pres. Parasit |
|----------------|------------------|------------------------------------|---|----------------------------|-------|---------------|---------------------|---------------|------------------|------------------|
| Limpopo | Letsitele | 23°39'24.10''S 30°40'15.47''E | Carissa spinarum L. Simple spine num-num | February 2014-2015 | 2 | 81.5 | 0.189 | 122 | 645.50 | |
| | Levubu | _ | Ekebergia capensis Sparrm. Cape-ash | February 2014 | 1 | 328 | 0.42 | 125 | 297.62 | Χ |
| | | - | <i>Psidium guajava</i> L. Guava* | October 2013 | 1 | 40 | 3.43 | 111 | 32.36 | |
| | Morebeng | _ | <i>Psidium guajava</i> L. Guava* | May 2015 | 1 | 30 | 1.22 | 121 | 99.18 | |
| | Thohoyandou | - | Sclerocarya birrea (A.Rich.) H Marula | | 2 | 52.5 | 1.07 | 14 | 13.08 | Χ |
| | Letsitele | 23°39'24.10''S 30°40'15.47''E | Berchemia discolor (Klotzsch, Brown ivory | | 3 | 441.67 | 0.36 | 283.67 | 787.97 | |
| Mpumalanga | Mbombela | 25° 27'06.24''S | Psidium cattleianum Afzel. ex | February-March-April | 5 | 63.6 | 0.23 | 61.4 | 266.96 | Х |
| | | 30° 58'09.66''E 25° 27'06.24''S | Strawberry Guava** Syzygium jambos L. | 2014-2016 December | 3 | 40.33 | 1.32 | 53 | 40.15 | Х |
| | | 30° 58'09.66''E 25° 27'06.24''S | Rose apple** Syzygium cordatum Hochst. | | 2 | 125 | 0.2319 | 14 | 60.37 | Х |
| | | 30° 58'09.66''E 25° 27'06.24''S | Water berry Sclerocarya birrea (A.Rich.) H | 2014 February | 1 | 50 | 1.13 | 129 | 114.16 | |
| | | 30° 58'09.66''E | Marula <i>Garcinia livingstonei</i> T.Ande | 2015 | | | | | | |
| | | 25° 27'06.24''S 30° 58'09.66''E | African mangosteen | 2013-2014 | 2 | 55.5 | 0.44 | 44.5 | 101.14 | Х |
| | | 25° 27'06.24''S 30° 58'09.66''E | <i>Cordyla africana</i> Lour. Wild mango | February 2016 | 1 | 20 | 1.08 | 110 | 101.85 | Х |
| Western Cape | Tulbagh | 33° 16'21.00''S 19° 10'08.40''E | Prunus domestica L. Plum* | January-April 2016-2017 | 2 | 11 | 0.5745 | 4.5 | 7.83 | |
| | Stellenbosch | 33° 55'19.68''S 18°48'4374''E | | January-April 2016-2017 | 4 | 50 | 2.87 | 14.5 | 5.05 | |
| | Ceres | - | | January-April 2016-2017 | 2 | 55 | 6.4 | 0 | 0.00 | |
| | Somerset West | 34°02'56.58''S 18°50'25.72''E | | February 2018 | 2 | 583 | 11.47 | 11 | 0.96 | |
| | Tulbagh | 33° 16'21.00''S 19° 10'08.40''E | <i>Prunus persica</i> L. Peach* | January-April 2016-2017 | 2 | 8.5 | 0.33 | 0.5 | 1.52 | |
| | Ceres | _ | | January-April 2016-2017 | 2 | 51 | 6.055 | 1.5 | 0.25 | |
| | Stellenbosch | 33°56'21.63''S 18°51'00.79''E | | February 2018 | 1 | 69 | 5.24 | 18 | 3.44 | |
| | Robertson | 33° 48'15.60''S 19° 53'00.60''E | | March 2016-2017-2018 | 2 | 103 | 16.085 | 12 | 0.75 | |
| | Stellenbosch | 33°55'18.20''S 18°55'41.57''E | <i>Malus domestica</i> B. Apple* | January-April 2016-2017 | 2 | 52.5 | 5.6 | 1.5 | 0.27 | |
| | Stellenbosch | 33°56'51.84''S 18°51'42.30''E | | January-April 2016-2017 | 2 | 50 | 2.75 | 0 | 0.00 | |
| | Stellenbosch | 33°55'18.20''S 18°55'41.57''E | <i>Pyrus communis</i> L. Pear* | January-April 2016-2017 | 2 | 50 | 4.65 | 44 | 9.46 | |
| | Stellenbosch | 33°56'51.84''S 18°51'42.30''E | | January-April 2016-2017 | 2 | 75 | 5.35 | 22 | 4.11 | |
| | Stellenbosch | 33°58'11.23''S 18°47'37.35''E | | February 2018 | 1 | 300 | 24.18 | 128 | 5.29 | |
| | Stellenbosch | 33°55'24.82''S 18°52'22.36''E | | March 2018 | 2 | 190 | 10.45 | 58 | 5.55 | |
| | Tulbagh | 33° 16'21.00''S 19° 10'08.40''E | | January-April 2016-2017 | 2 | 35 | 5.18 | 3.5 | 0.68 | |
| | Riebeeck Kasteel | 33°23'12.00''S 18°54'00.00''E | <i>Vitis</i> sp. Table grape* | January-April 2016-2017 | 1 | 50 | 4 | 1 | 0.25 | |
| | De Doorns | 33°28'28.80''S 19°39'35.40''E | | January-April 2016-2017 | 1 | 50 | 1.7 | 0 | 0.00 | |
| | Stellenbosch | 33°56'51.84''S 18°51'42.30''E | Citrus limon L. Citrus * | January-April 2016-2017 | 1 | 130 | 3.6 | 2 | 0.56 | |
| | Stellenbosch | 33°56'15.59''S 18°51'24.09''E | <i>Mespilus</i> sp. Medlar** | February 2018 | 1 | 103 | 2.11 | 12 | 5.69 | |

| | | | | | | | | | doji | | | |
|-----------------------|----------------------|------|--------|-------|------------------|-------|-------|-------|------|----|---|-----------|
| Species | Species | Т | W(g) | Sz(g) | Parasitoid sp | % | R - | R+ | ن. | ر. | 5 | <u>8.</u> |
| Berchemia discolor | Brown ivory | 1325 | 1069.4 | 0.80 | P. humilis | 22.37 | 14.02 | 40.32 | Χ | | | |
| Carissa spinarum | Simple spine num-num | 163 | 378.0 | 2.31 | | 0 | 0 | 0 | Χ | | | |
| Cordyla africana | Wild mango | 20 | 1079.1 | 53.95 | P. humilis | 2.65 | na | na | | Χ | Χ | Χ |
| Ekebergia capensis | Cape-ash | 328 | 421.0 | 1.28 | P. humilis | 6.02 | na | na | Χ | | | |
| Garcinia livingstonei | African mangosteen | 111 | 873.5 | 7.87 | F. ceratitivorus | 34.41 | 18.82 | 50.00 | Χ | Χ | | |
| Psidium cattleianum | Strawberry Guava | 318 | 1146.3 | 3.60 | F. ceratitivorus | 5.96 | 0 | 13.33 | Χ | Χ | Χ | Χ |
| Psidium guajava | Guava | 70 | 4649.8 | 66.42 | | 0 | 0 | 0 | Χ | Χ | Χ | Χ |
| Sclerocarya birrea | Marula | 212 | 4523.3 | 21.33 | | 0 | 0 | 0 | | | Χ | |
| Syzygium cordatum | Water berry | 250 | 463.8 | 1.86 | F. ceratitivorus | 38.54 | 33.33 | 43.75 | Χ | Χ | | |
| Syzygium jambos | Rose apple | 121 | 3940.7 | 32.57 | F. ceratitivorus | 17.98 | 0 | 53.93 | | Χ | | |