EFFECTS OF *TOMATO ZONATE SPOT VIRUS* INFECTION ON THE DEVELOPMENT AND REPRODUCTION OF ITS VECTOR *FRANKLINIELLA OCCIDENTALIS* (THYSANOPTERA: THRIPIDAE)

Xue Zheng, Jie Zhang, Yongdui Chen, Jiahong Dong* and Zhongkai Zhang* Biotechnology and Germplasm Resource Institute, Yunnan Academy of Agricultural Sciences, Yunnan Province Key Laboratory of Agricultural Biotechnology, Kunming 650223, China

Corresponding authors; E-mail Addresses: dongjhn@126.com; zhangzhongkai99@126.com

ABSTRACT

The western flower thrips, *Frankliniella occidentalis* (Pergande) (Thysanoptera: Thripidae), is the main vector of *Tomato Zonate Spot Virus* (TZSV) in Yunnan, China. We investigated the life history parameters of *F. occidentalis* on tomato and pepper leaves either with or without TZSV infection. The total duration of the immature stages of *F. occidentalis* reared on TZSV-infected leaves was significantly shorter than the total duration of those reared on uninfected leaves. Also the survival rates of the instars and prepupae on TZSV-infected tomato and pepper leaf disks were significantly higher than those on uninfected tomato and pepper leaf disks. The *F. occidentalis* populations reared on TZSV-infected tomato and pepper leaf disks respectively increased 11.97- and 10.64-fold in 1 generation, while those reared on uninfected tomato and pepper leaf disks increased only 8.10- and 6.45-fold, respectively. These results demonstrated that TZSV infection improved the fitness and host suitability of its vector, *F. occidentalis*. Also our findings suggest that TZSV-infection is likely to induce larger field populations of *F. occidentalis*, thereby increasing the probability of TZSV transmission.

Key Words: *Tomato zonate spot virus* (TZSV), western flower thrips, development, reproduction

Resumen

Los trips occidental de las flores, *Frankliniella occidentalis*, (Pergande) (Thysanoptera: Thripidae), es el principal vector del Virus de la Mancha Zonada del Tomate (VMZT) en Yunnan, China. Se investigó los parámetros del ciclo de vida de *F. occidentalis* sobre hojas de tomate y pimiento con o sin infección por VMZT. La duración total de las etapas inmaduras de *F. occidentalis* criadas en las hojas infectadas por el VMZT fue significativamente más corta que la duración total de las criadas en las hojas no infectadas. También, las tasas de sobrevivencia de los instares y prepupas sobre discos de hojas de tomate y pimiento infectadas por VMZT fueron significativamente más altas que las tasas sobre discos de hojas de tomate y pimiento no infectadas. Las poblaciones del *F. occidentalis* criadas sobre discos de hojas de tomate y pimienta infectados por VMZT, incrementaron 11.97 y 10.64 veces, respectivamente en 1 generación, mientras que las criadas sobre discos de hojas de tomate y pimienta no infectadas incrementaron sólo 8.10 y 6.45 veces, respectivamente. Estos resultados demostraron que la infección por VMZT mejoró el estado físico e idoneidad de su vector, *F. occidentalis*. También nuestros hallazgos sugieren que es probable que la infección por VMZT induzca poblaciones de campo de *F. occidentalis* más grandes y con ello incrementando la probabilidad de transmisión de VMZT.

Palabras Clave: Virus de la Mancha Zonada del Tomate (VMZT) , trips occidental de las flores, desarrollo, reproducción

Tospoviruses have received international attention because they have caused major crop losses throughout the world in recent years (Dong et al. 2013; Pappu et al. 2009; Puangmalai et al. 2013; Seepiban et al. 2011). Parrella et al. (2003) reported that TSWV can infect more than 1,090 plant species and is one of the most economically important plant viruses in the world. *Tomato zonate spot virus* (TZSV) is a new member of the ge-

nus *Tospovirus* and family *Bunyaviridae*, it was first reported in Yunnan province in China (Dong et al. 2008, 2010). Like other tospoviruses, the genome of TZSV is comprised 3 molecules of negative-sense or ambisense single-stranded RNAs (ssRNAs) that encode 4 structural proteins and 2 non-structural proteins (Dong et al. 2008). *Tomato spotted wilt virus* (TSWV) is the dominant species of the genus *Tospovirus* in Yunnan province,

and this species has been causing substantial yield losses in vegetable, tobacco and other cash crops in recent years. Dong et al. (2010) reported that the host plant range of TZSV includes about 25 plant species belonging to 7 families, including economically crops and weeds such as tomato (*Solanum lycopersicum* L.; Solanales: Solanaceae), chili pepper (*Capsicum annuum* L*.*; Solanales: Solanaceae), *Rumex dentatus* L.; Caryophyllales: Polygonaceae) and *Bidens pilosa* L.; Asterales: Asteraceae).

It has long been known that most species of tospoviruses are transmitted by thrips. For example, the type species TSWV is transmitted by *Frankliniella occidentalis* (Pergande) (Thysanoptera: Thripidae), the western flower thrips, in a persistent propagative manner (Ullman et al. 1997, 2002). Our previous work showed that *F. occidentalis* is also the main vector of TZSV in Yunnan (Dong et al. 2008; Zheng et al. 2013). *Franliniella occidentalis* is a destructive pest that was first reported in China at the beginning of the 20th-century (Zhang et al. 2003). The short reproductive cycle and high fecundity of this insect contribute to its success as a destructive pest. This, together with its extremely wide host range, broad geographical distributions and capacity to transmit many tospoviruses, makes *F. occidentalis* difficult to control by current strategies. This insect therefore has become a major constraint on the production of numerous economically important crops (Morse & Hoddle 2006).

Some studies have indicated that upon infection of *Sogatella furcifera* (Horváth) (Hemiptera: Delphacidae) with *Southern rice black-streaked dwarf virus* (SRBSDV), or the infection of *F. occidentalis* with *Tomato spotted wilt virus,* the fitness of both of these vectors was improved (Maris et al. 2004; Zhang et al. 2014). Although the effects of some other plant viruses, such as *Tomato mottle virus* (ToMoV) and *Barley yellow dwarf virus* (BYDV), on their vectors were studied (McKenzie 2002; Fiebig et al. 2004), little is known about the effect of TZSV infection on its vector, *F. occidentalis*. The present study investigated the impact of TZSV infection on the life cycle, reproductive success and population trend index of *F. occidentalis* in 2 host species, tomato (*S. lycopersicum*) and pepper (*C. annuum*).

Materials and Methods

Virus Source, Host Plant and Vector

TZSV inoculum was collected in 2012 from TZSV-infected tomato plants. This material was tested by RT-PCR to confirm infection in the field at Yuanmou, Yunnan Province, and the inoculum was kept frozen at -80 °C. Tomato and pepper were seeded in pot mixtures in controlled climate chambers at 25 ± 1 °C and 16:8 h L:D, and finally

in the dark at 25 °C for 24 h prior to inoculation. Four-week old plants with at least 2 true leaves were mechanically inoculated with either TZSVinfected or uninfected leaf tissue that had been steeped in the inoculation buffer (0.05 M of sodium phosphate buffer, pH 7.0). Then the leaves were rinsed with water, and the plants were maintained in the climate chambers. Thirty days after TZSV inoculation, some leaf tissues from inoculated plants were tested by RT-PCR to confirm infection. After confirmation, leaves from plants with confirmed infection were used for the experiments.

Frankliniella occidentalis thrips were obtained from healthy tomato plants from Mile, Yunnan province of China and the *F. occidentalis* colony was maintained on green bean pods (*Phaseolus vulgaris* L.; Fabales: Fabaceae) at 25 ± 1 °C and 16:8 h L:D for more than 1 yr in the Institute of Biotechnology and Germplasm Resources, Yunnan Academy of Agricultural Sciences, Kunming, China.

Development of *F. occidentalis* Fed on TZSV-Infected or Uninfected Tomato or Pepper

We used the method of Zhang et al. (2007) to assess the effects of TZSV-infection on the development *F. occidentalis*. Briefly, 20 newly emerged adult females were collected, divided into groups of 5 and each group was placed in a Petri dish (12 cm diam) containing 25 mm-diam leaf disks, with or without TZSV-infection, over moistened filter papers. The adults were removed after they had oviposited on the leaf disks during 12 h. The Petri dishes, each with a moist filter paper, were then placed in the climate chambers and maintained at 25 ± 1 °C and 16:8 h L:D for investigations on thrips development. After hatching, 40 larvae were transferred individually into new Petri dishes (7 cm diam) containing TZSV-infected or uninfected tomato and pepper leaf disks. Each Petri dish was sealed with parafilm to prevent *F. occidentalis* from escaping. The leaf disks were changed daily. Growth of *F. occidentalis* in each Petri dish was monitored daily until the emergence of adults. Duration of each developmental stage of *F. occidentalis* was recorded for all treatments. All dead individuals of any developmental stage were excluded when calculating the average developmental time for a specific stage. Each of the experiments was repeated 5 times.

Life Table Parameters of *F. occidentalis* Fed on TZSV-Infected or Uninfected Tomato or Pepper

One hundred neonates were collected randomly as founders of the experimental population and reared on the leaf disks as described above. When the neonates developed into 2nd instars and pu-

pae, the survival rates representing those developed into the 2nd instar (Su1) or those that had developed into the prepupae-to-pupae (Su2) were determined. The emerged males and females were then counted daily to determine the emergence rate (Er) and percentage of females (Fr). The leaf disks were inspected thoroughly for the numbers of non-hatched eggs. Fecundity (Fy) was determined based on the average number of eggs produced by the females. Hatchability (Hy) was calculated as the total number of neonates over the total number of neonates plus the number of non-hatched eggs.

A new TZSV-Infected or Uninfected leaf disk was replaced every 2 days in each Petri dish, as appropriate, and the number of eggs laid on the previous leaf disk was counted by Nikon SMZ1500 stereomicroscope (100×). The experiment was repeated 5 times. The population growth index (I) was calculated as follows:

 $Nt = No \times Su1 \times Su2 \times Er \times Fr \times Fy \times Hy$ $I = Nt / No$

Where No is the number of neonates used in the initial population and Nt is the number of individuals in the next-generation population.

Data Analysis

All data were analyzed using the SPSS®, Version 11.5. Tukey-Kramer method $(P < 0.05)$ was used to determine significant differences between treatments, and all factors were standardized using the arcsine square-root transformation.

RESULTS

Development of *F. occidentalis* Fed on TZSV-Infected or Uninfected Tomato or Pepper

The results in Table 1 show the duration time of each stage for *F. occidentalis* reared on tomato and pepper leaves with or without TZSV infection. Although the durations of development (days) of the eggs, 1st instars and pupae were slightly shorter for *F. occidentalis* reared on TZSV-infected than on the uninfected tomato or pepper leaf disks, these differences were not statistically significant. However, the durations of development of 2nd instars and total development from egg to adult of *F. occidentalis* reared both on TZSV-infected tomato or pepper leaf disks were significantly shorter $(P < 0.05)$ than those reared on the uninfected leaves.

Life Table Parameters of *F. occidentalis* Fed on TZSV-Infected or Uninfected Tomato or Pepper

The survival rates from neonates to 2nd instars (Su1) of *F. occidentalis* reared on TZSV-infected tomato and pepper leaf disks (Table 2) were significantly higher than those reared on uninfected tomato and pepper leaf disks. Likewise the survival rates from prepupae to pupae (Su2) of *F. occidentalis* reared on TZSV-infected tomato and pepper leaf disks were significantly higher than those reared on uninfected tomato and pepper leaf disks. Although the emergence rates, female ratios and fecundities of *F. occidentalis* reared on TZSV-infected tomato and pepper leaf disks were higher than those observed for *F. occidentalis* reared on their uninfected counterparts, there were no significant differences between the 2 treatments (TZSV-infected and uninfected) as well as between the 2 host plants.

Frankliniella occidentalis populations reared on TZSV-infected tomato and pepper leaf disks in the laboratory increased 11.97- and 10.64-fold, respectively in 1 generation (Table 2), but those reared on uninfected tomato and pepper leaf disks increased only 8.10- and 6.45-fold, respectively.

Discussion and Conclusions

Knowledge of how a virus impacts fitness of its insect vector is a key element in understanding outbreaks of a virus disease. To date, the effects of several viruses, such as *Rice dwarf virus* (RDV),

Table 1. Average duration (days) of each developmental stage of *Frankliniella occidentalis* reared on TZSV-infected or uninfected tomato or pepper leaf disks.

Stage of F. occidentalis	Tomato		Pepper	
	TZSV-infected	Uninfected	TZSV-infected	Uninfected
Egg	2.72 ± 0.14 a	3.02 ± 0.26 a	2.27 ± 0.04 b	2.29 ± 0.04 b
1st instar larva	2.65 ± 0.22 a	2.78 ± 0.06 a	2.28 ± 0.05 b	2.36 ± 0.11 ab
2nd instar larva	3.04 ± 0.11 a	3.44 ± 0.15 b	3.83 ± 0.09 c	4.32 ± 0.24 d
Prepupa	1.05 ± 0.15 a	1.18 ± 0.10 a	1.13 ± 0.08 a	1.57 ± 0.20 b
Pupa	2.25 ± 0.08 a	2.46 ± 0.24 a	1.42 ± 0.16 b	1.98 ± 0.08 b
Total immature stage	11.71 ± 0.20 a	12.88 ± 0.15 b	10.93 ± 0.03 c	12.52 ± 0.38 b

Each value represents the mean ± standard error (SE) of data from 5 replicates. Values in the same row with different letters are significantly different at *P* < 0.05 level (Tukey-Kramer method)

Table 2. Life table parameters of *Frankliniella occidentalis* reared on leaf disks from TZSV-infected and uninfected tomato and pepper.

No, number of individuals in the initial population; Su1, survival rate from neonate to 2nd instar; Su2, survival rate from prepupa to pupa; Er, emergence rate; Fr, female ratio; Fy, fecundity (eggs per female); Hy, hatchability; N, predicted number of offspring; I, population trend index (folds of increase compared to the number in the preceding generation. Each value represents the mean \pm SE of data from 5 replicates. Values in the same row with different letters are significantly different at $P < 0.05$ level (Tukey-Kramer method).

Tomato mottle virus (ToMoV), *Chrysanthemum stem necrosis virus* (CSNV), TSWV and *Barley yellow dwarf virus* (BYDV), on their transmission vectors have been studied (Nakasuji & Kiritani 1970; McKenzie 2002; Fiebig et al. 2004; Maris et al. 2004;).

TZSV transmitted by *F. occidentalis* alters the quality of the host plant, which could influence the performance of *F. occidentalis*, including its longevity, growth, fecundity, etc. (Brodbeck et al. 2002; Scott Brown et al. 2002; Scheirs et al. 2003; Abe et al. 2012; Shrestha et al. 2012; Jacobson & Kennedy 2013; Miray & Mehmet 2013; Ogada et al. 2013; Okuda et al. 2013). For example, Ogada et al. (2013) reported that host plants infected with TSWV could improve longevity and survival of the vector, *F. occidentalis*. Viral infection can alter a plant's suitability for further infection through alteration of the vector's preference and performance, because viral infection reduces plant nutrient levels or induces plant defense mechanisms that, in turn, offset insect reproduction (Harris et al. 2001; Messina et al. 2002). In a series of experiments, the direct and indirect (through the host plant) effects of plant viruses on the vectors have been quantified. For example, it was reported that the durations of development and survival of *F. occidentalis* juveniles were positively affected by being reared on TSWV-infected plant leaves, and plants infected with TSWV were more attractive than uninfected plants to *F. occidentalis* (Bautista et al. 1995; Maris et al. 2004; Belliure et al. 2005; Abe et al. 2012; Ogada et al. 2013). On the other hand TSWV, *Impatiens necrotic spot virus* (INSV), which is closely related to TZSV in the genus *Tospovirus*, was found to negatively impact *F. occidentalis* (De Angelis et al. 1993). In the current study, we showed that *F. occidentalis* was able to complete its life cycle

on the tomato and pepper leaves with or without TZSV-infection. Although the durations of development of the egg, 1st instar, prepupa and pupa were shorter for *F. occidentalis* reared on TZSVinfected tomato and pepper leaves than those reared on uninfected leaves, these differences were not significant. The durations of development of 2nd instars and the total immature stage of *F. occidentalis* reared on TZSV-infected tomato and pepper leaves were, however, significantly shorter than those reared on uninfected leaves. These results are consistent with those reported for TSWV by Maris et al. (2004) and suggest that TZSV-infected tomato and pepper are more suitable hosts for *F. occidentalis* development than their uninfected counterparts.

Life-table studies are fundamental to population ecology. The life table gives the most comprehensive description of survivorship, reproduction, etc, of a population and provides basic data on population growth parameters. In the present study, we compared *F. occidentalis* reared on uninfected tomato and pepper leaves with *F. occidentalis* reared on TZSV-infected tomato and pepper leaves, and we found that the latter had a slightly greater female ratios, fecundities, egg hatchabilities, and emergence rates, but these differences were not statistically significant. However, the survival rates of *F. occidentalis* from neonate to 2nd instar (Su1) and from prepupa to pupa (Su2) reared on TZSV-infected tomato and pepper leaves were significantly higher than those reared on uninfected leaves.

TZSV has spread across many parts of Yunnan Province in large part because of the reproductive success of its vector. In the current study, lower fecundities were observed in *F. occidentalis* reared on uninfected tomato and pepper leaves. Moreover, the population trend indexes (I) of *F. occidentalis* reared on TZSV-infected plants were considerably higher than those reared on uninfected leaves. It is known that virus infection may improve the availability of vital nutrients such as free amino acids in infected host plants (Ajayi 1986). Shrestha et al. (2012) have reported that the increased oviposition of viruliferous *F. fusca* could have been influenced by the availability of increased concentrations of free amino acids in TSWV-infected plants. Thus, in the present study, the population trend index variations might be correlated with changes in host nutrition values upon TZSV infection, and changes in the life-table parameters are highly important with respect to abundance of viruliferous vectors in the field. It is noteworthy that life-table parameters often vary with different environmental, host species and other factors. Clearly the results presented in this report provide valuable information relevant to better understanding the progress of TZSV epidemics, and to the development of strategies and tactics for combating TZSV and *F. occidentalis*.

Frankliniella occidentalis may also benefit from viral infection, because TZSV-infected tomato and pepper plants are more suitable hosts for the development and reproduction of *F. occidentalis*. van Lenteren & Noldus (1990) reported that faster developmental rate and higher fecundity of an insect species on a given host plant indicate better suitability of the host plant. We conclude that TZSV infection improves the suitability of tomato and pepper for TZSV's vector, *F. occidentalis*. In summary, our findings suggest that TZSV-infected tomato and pepper increased in fitness and reproductive success of *F. occidentalis*, thereby increasing the probability of greater vector populations and greater TZSV transmission.

ENDNOTES

The study was supported by National Natural Science Foundation of China (31060237, 31360430), Chinese Postdoctoral Science Foundation (2013M531992), Science and Technology Program of Yunnan Province (2012CH007). Xue Zheng and Jie Zhang are co-first authors because they contributed equally to this study.

References Cited

- Abe, H., Tomitaka, Y., Shimoda, T., Seo, S., Sakurai, T., Kugimiya, S., Tsuda, S., and Kobayashi, M. 2012. Antagonistic plant defense system regulated by phytohormones assists interactions among vector insect, thrips and a tospovirus. Plant Cell Physiol. 53: 204-212.
- Ajayi, O. 1986. The effect of *Barley yellow dwarf virus* on the amino acid composition of spring wheat. Ann. Appl. Biol. 108: 145-149.
- Bautista, R. C., Mau, R. F. L., Cho, J. J., and Custer, D. M. 1995. Potential of tomato spotted wilt tospovirus plant hosts in Hawaii as virus reservoirs for

transmission by *Frankliniella occidentalis* (Thysanoptera: Thripidae). Phytopathology. 85: 953-958.

- Belliure, B., Janssen, A., Maris, P. C., Peters, D., and Sabelis, M. W. 2005. Herbivore arthropods benefit from vectoring plant viruses. Ecol. Lett. 8: 70-79.
- Brodbeck, B. V., Funderburk, J., Stavisky, J., Andersen, P. C., and Hulshof, J. 2002. Recent advances in the nutritional ecology of Thysanoptera, or the lack thereof, pp. 145-153 *In* R. Marullo and L. Mound [eds.], Thrips and Tospoviruses: Proc. 7th Intl. Symp. on Thysanoptera. Australian Natl. Insect Collection (ANIC), Canberra.
- De Angelis, J. D., Sether, D. M., and Rossignol, P. A. 1993. Survival, development, and reproduction in western flower thrips (Thysanoptera. Thripidae) exposed to *Impatiens necrotic spot virus*. Environ. Entomol. 22: 1308-1312.
- Dong, J. H., Cheng, X. F., Yin, Y. Y., Fang, Q., Ding, M., Li, T. T., Zhang, L. Z., Su, X. X., McBeath, J. H., and Zhang, Z. K. 2008. Characterization of *Tomato zonate spot virus,* a new tospovirus species in China. Arch. Virol. 153: 855-864.
- Dong, J. H., Zhang, Z. K., Yin, Y. Y., Cheng, X. F., Ding, M., and Fang, Q. 2010. Natural host ranges of *Tomato zonate spot virus* in Yunnan. J. Insect Sci. 10: 12-13.
- Dong, J. H., Yin, Y. Y., Fang, Q., and Zhang, Z. K. 2013. A new tospovirus causing chlorotic ringspot on Hippeastrum sp. in China. Virus Genes 46: 567-570.
- Fiebig, M., Plehling, H. M., and Borgemeister, C. 2004. *Barley yellow dwarf virus*, wheat, and *Sitobion avenae*: a case of trilateral interactions. Entomol. Exp. Appl. 110: 11-21.
- Harris, M. O., Sandanayaka, M., and Griffin, A. 2001. Oviposition preferences of the Hessian fly and their consequences for the survival and reproductive potential of offspring. Ecol. Entomol. 26: 473-486.
- Jacobson, A. L., and Kennedy, G. G. 2013. Specific Insect-Virus Interactions Are Responsible for Variation in Competency of Different *Thrips tabaci* Isolines to Transmit Different *Tomato Spotted Wilt Virus* Isolates. Plos One. 8: e54567.
- Maris, P. C., Joosten, N. N., Goldvach, R. W., and Peters, D. 2004. *Tomato spotted wilt virus* infection improves host suitability for its vector *Frankliniella occidentalis*. Phytopathology 97: 706-711.
- McKenzie, C. L. 2002. Effect of *Tomato mottle virus* (ToMoV) on *Bemisia tabaci* biotype B (Homoptera: Aleyrodidae) oviposition and adult survivorship on healthy tomato. Florida Entomol. 85: 367-368.
- Messina, F. J., Taylor, R., and Karren, M. E. 2002. Divergent responses of two cereal aphids to previous infestation of their host plant. Entomol. Exp. Appl. 103: 43-50.
- Miray, A. S., and Mehmet, A. S. 2013. Spread of *Tomato spotted wilt virus* from an internal virus source by thrips species in Samsun, Turkey. Phytoparasitica 41: 159-168.
- MORSE, J. G., AND HODDLE, M. S. 2006. Invasion biology of thrips. Annu. Rev. Entomol. 51: 67-89.
- Nakasuji, F., and Kiritani, K. 1970. Effects of *Rice dwarf virus* upon its vector, *Nephotettix cincticeps* Uhler (Hemiptera: Deltocephalidae), and its significance for changes in relative abundance of infected individuals among vector populations. Appl. Entomol. Zool. 5: 1-12.
- Ogada, P. A., Maiss, E., and Poehling, H. M. 2013. Influence of *Tomato spotted wilt virus* on performance

and behavior of western flower thrips (*Frankliniella occidentalis*). J. Appl. Entomol. 137: 488-489.

- Okuda, S., Okuda, M., Matsuura, S, Okazaki, S., and Iwai, H. 2013. Competence of *Frankliniella occidentalis* and *Frankliniella intonsa strains* as vectors for *Chrysanthemum stem necrosis virus*. European J. Plant. Pathol. 136: 355-362.
- Pappu, H. R., Jones, R. A. C., and Jain, R. K. 2009. Global status of tospovirus epidemics in diverse cropping systems: successes achieved and challenges ahead. Virus Res. 141: 219-236.
- Parrella, G., Gognalons, P., Gebre-Selassie, K., Vovlas, C., and Marchoux, G. 2003. An update of the host range of *Tomato spotted wilt virus*. J. Plant Pathol. 85: 227-264.
- Puangmalai, P., Potapohn, N., Akarapisarn, A., and Pascha, H. J. 2013. Inheritance of *Tomato necrotic ring viru*s resistance in *Capsicum annuum*. J. Agric. Sci. 5: 129-133.
- Scheirs, J., De Bruyn, L., and Verhagen, R. 2003. Host nutritive quality and host plant choice in two grass miners: primary roles for primary compounds? J. Chem. Ecol. 29: 1373-1389.
- Scott Brown, A. S., Simmonds, M. S. J., and Blaney, W. M. 2002. Relationship between nutritional composition of plant species and infestation levels of thrips. J. Chem. Ecol. 28: 2399-2409.
- Seepiban, C., Gajanandana, O., Attathom, T., and Attathom, S. 2011. *Tomato ring spot virus*, a new of tospovirus isolated in Thailand. Arch. Virol. 156, 263-274.
- Shrestha, A., srinivasan, R., Riley, D. G., and Cul-BREATH, A. K. 2012. Direct and indirect effects of a thrips-transmitted *Tospovirus* on the preference and

fitness of its vector, *Frankliniella fusca*. Entomol. Exp. Appl. 145: 260-271.

- Ullman, D. E., Sherwood, J. L., and German, T. L. 1997. Thrips as vectors of plant pathogens, pp. 539- 565. *In* T. Lewis [ed.], Thrips as Crop Pests. CAB Intl., Wallingford, United Kingdom.
- Ullman, D. E., Medeiros, R. B., Campbell, L. R., Whitfield, A. E., Sherwood, J. L., and German, T. L. 2002. Thrips as vectors of tospoviruses. Adv. Bot. Res. 36: 113-140.
- van Lenteren, J. C., and Noldus, L. P. J. J. 1990. Whitefly-plant relationships: behavioural and ecological aspects, pp. 47-89 *In* D. Gerling [ed.], Whiteflies: their bionomics, pest status and management. Intercept, Andover, UK.
- Zhang, J., Zheng, X., Chen, Y. D., Hu, J., Dong, J. H., Su, X. X., and Zhang, Z. K. 2014. *Southern rice blackstreaked dwarf virus* Infection Improves Host Suitability for Its Insect Vector, *Sogatella furcifera* (Hemiptera: Delphacidae). J.Econ. Entomol. 107: 92-97.
- Zhang, Y. J., Wu, Q. J., Xu, B. Y., and Zhu, G. R. 2003. Dangerous invasive species – the western flower thrips caused damage in Beijing. Plant Prot*.* 29: 58- 59. (in Chinese with English summary)
- Zhang, Z. J., Wu, Q. J., Li, X. F., Zhang, Y. J., Xu, B. Y., and Zhu, G. R. 2007. Life history of western flower thrips, *Frankliniella occidentalis* (Thysan., Thripae), on five different vegetable leaves. J. Appl. Entomol. 131: 347-354.
- Zheng, X., Liu, C. M., Li, H. G., Zhang, J., Dong, J. H., AND Zhang, Z. K. 2013. Investigation of thrips, host plants of Tospoviruses in Honghe of Yunnan Province. China Plant Prot. 33: 41-44. (in Chinese with English summary)