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CRYPTIC INVASION OF THE EXOTIC *BEMISIA TABACI* BIOTYPE Q OCCURRED WIDESPREAD IN SHANDONG PROVINCE OF CHINA

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ABSTRACT

Bemisia tabaci (Gennadius) is an important agricultural pest worldwide. The pest is a species complex composed of numerous biotypes, among which biotypes B and Q are the 2 most invasive and widely distributed. Our previous study found that the ratio of the biotype Q has been increasing and displacement of biotypes B by Q has been occurring on cotton and eggplant in Shandong Province of China during the past several years. To determine whether biotype Q has been increasing on other hosts and possible displacement of biotypes B and Q on cultivated and wild host species near cotton or eggplant fields in 7 locations of Shandong Province during 2005-2008 with cleavage amplified polymorphic sequence (CAPS) of the *mtCOI* (mitochondrial cytochrome oxidase subunit I) marker. This research showed biotype Q has been occurring in the province as a whole. The displacement of biotypes B by Q has been increasing on all kinds of host plants and the displacement of biotypes B by Q has been occurring in the province as a whole. The displacement of biotypes B by Q has been increasing on all kinds of host plants and the displacement of biotypes B by Q has been occurring in the province as a whole. The displacement of biotypes B by Q has been occurring in the province as a whole.

Key Words: Bemisia tabaci, biotype Q, mitochondrial cytochrome oxidase marker

RESUMEN

Bemisia tabaci (Gennadius) es una plaga importante a la agricultura mundial. La especie es un complejo de especies compuesta de biotipos numerosos, entre ellos los biotipos B y Q son los 2 mas invasivos y distribuidos ampliamente. Nuestro estudio anterior encontró que la proporción de biotipo Q ha ido incrementando y el desplazamiento del biotipo B por el Q ha ido ocurriendo sobre algodón y barenjena en la Provincia de Shangdon en China durante varios de los años pasados. Para determinar si el biotipo Q se ha ido incrementando sobre otros hospederos y si el posible desplazamiento de biotipos ha ido ocurriendo en toda la provincia, tambien muestreamos los biotipos B y Q de B. tabaci sobre especies de hospederos cultivados y silvestres cerca de campos de algodón y berenjena en 7 localidades en la Provincia de Shandong durante 2005-2008 usando el metodo escisión de la secuencia polimórfica amplificado (ESPA) del marcador subunidad I de *mtCOI* (oxidasa citocromo mitocondrial). Esta investigación mostró que el biotipo Q se ha ido incrementando sobre toda clase de plantas hospederos y el desplazamiento de biotipo B por el Q ha ido ocurriendo en toda la provincia. El mecanismo del desplazamiento debe ser investigado y este conocimiento puede guiar la aplicación de insecticidas o un ajuste de los cultivos para controlar la plaga efectivamente.

Bemisia tabaci (Gennadius) is an important agricultural pest worldwide. It damages crops through direct feeding and vectors many plant viruses. The pest has been considered as a species complex that includes many genetic groups that are morphologically indistinguishable. Some are labeled as biotypes or host races because of differences in host range, geographical distribution, transmission ability of plant virus, and other biological characteristics. Biotype B is believed to originate from the Middle East-Northern Africa and have spread into many countries or regions over the past 2 decades. Biotype Q may have originated in the Mediterranean countries and circumstantial data shows that this biotype has been introduced into many non-Mediterranean countries or regions during the past several years (Chu et al. 2005; Ueda 2006; Brown et al. 2007).

Bemesia tabaci outbreaks in the mid-1990s in both Southern China and Northern China and subsequent research showed that the whitefly occurring in most of China was biotype B (Wu et al. 2002). In 2003, biotype Q was found in Kunming of Yunnan Province and then subsequently found in Beijing and Henan (Chu et al. 2006). Many populations of B. tabaci in Shandong Province, one of the most important agricultural provinces of China, have proved to be biotype B, but biotype Q was discovered in 2006 by use of mitochondrial cytochrome oxidase subunit I (mt-COI) sequence (Chu et al. 2007). It is important to monitor the spreading and the density of the biotype Q because it possesses greater resistance to many insecticides than biotype B in many countries (Dennehy et al. 2005; Horowitz et al. 2005). The biotypes of B. tabaci on cotton and eggplant in 6 locations within Shandong Province was determined with mtCOI sequences and biotype B-specific primers (Chu et al. 2010), which showed that the ratio of the biotype Q has been increasing and displacement of biotypes B by Q has been occurring during the past 4 years (2005-2008) on these two crops. To determine whether biotype Q has been increasing on other hosts and possible displacement of biotypes has been occurring in the province as a whole, we further surveyed *B. tabaci* biotypes on cultivated and wild host species near cotton or eggplant fields in 7 locations of Shandong Province during 2005-2008.

MATERIALS AND METHODS

Bemisia tabaci biotype was determined with the cleaved amplified polymorphic sequences (CAPS) of *mtCOI* amplified with new primers (C1-J-2195/R-BQ-2819). Adult whiteflies were collected from different plants including crops and weeds in 7 locations, DeZhou, ZiBo, Shou-Guang, JiNan, LiaoCheng, LinYi and ZaoZhuang in Shandong Province during 2005-2008 (Table 1). The adults were placed in tubes with 95% ethanol and stored at -20°C. Individual adults were ground and DNA was extracted. The

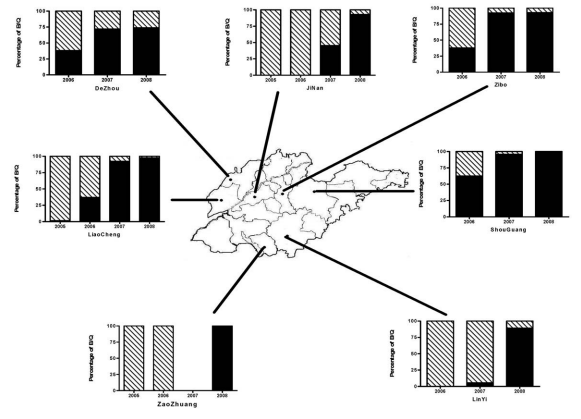


Fig. 1. Composition changes of *B. tabaci* biotypes B and Q in Shandong Province during 2005-2008 \blacksquare , biotype Q; \blacksquare , biotype B.

Location	Year	Host plant	Total individuals	Number of biotype		Percentage of biotype (%)	
				В	Q	В	Q
DeZhou	2006	Cotton	31	23	8	74.2	25.8
		Eggplant	61	34	27	55.7	44.3
		All host plants	92	57	35	62.0	38.0
	2007	Tomato	21	7	14	33.3	66.7
		Cotton	71	17	54	23.9	76.1
		Eggplant	45	7	38	15.6	84.4
		Zucchini	37	18	19	48.6	51.4
		All host plants	174	49	125	28.2	71.8
	2008	Cotton	27	0	27	0.0	100.0
		Japanese hop	30	0	30	0.0	100.0
		Eggplant	30	0	30	0.0	100.0
		All host plants	87	0	87	0.0	100.0
ZiBo	2006	Cotton	45	32	13	71.1	28.9
		Morning glory	20	0	20	0.0	100.0
		Eggplant	25	23	2	92.0	8.0
		All host plants	90	55	35	61.1	38.9
	2007	Cotton	75	2	73	2.7	97.3
		Eggplant	48	7	41	14.6	85.4
		All host plants	123	9	114	7.3	92.7
	2008	Japanese hop	28	0	28	0.0	100.0
		Cotton	29	0	29	0.0	100.0
		Eggplant	30	6	24	20.0	80.0
		All host plants	87	6	81	6.9	93.1
ShouGuang	2006	Chinese cabbage	60	23	37	38.3	61.7
		Rosebush	17	16	1	94.1	5.9
		Eggplant	30	1	29	3.3	96.7
		All host plants	107	40	67	37.4	62.6
	2007	Tomato	49	0	49	0.0	100.0
		Cucumber	24	0	24	0.0	100.0
		Pepper	25	0	25	0.0	100.0
		Japanese hop	66	1	65	1.5	98.5
		Cotton	98	13	85	13.3	86.7
		Pumpkin	62	0	62	0.0	100.0
		Eggplant	57	3	54	5.3	94.7
		Sweet pepper	24	0	24	0.0	100.0
		Chinese cabbage	41	1	40	2.4	97.6
		All host plants	446	18	428	4.0	96.0
	2008	Japanese hop	29	0	29	0.0	100.0
		Cotton	25	0	25	0.0	100.0
		Eggplant	30	0	30	0.0	100.0
		All host plants	84	0	84	0.0	100.0
JiNan	2005	Cotton	24	24	0	100.0	0.0
	2006	Eggplant	28	28	0	100.0	0.0
		Cotton	31	31	0	100.0	0.0
		All host plants	59	59	0	100.0	0.0
	2007	Cotton	47	18	29	38.3	61.7
	2008	Japanese hop	24	1	23	4.2	95.8
		Cotton	29	0	29	0.0	100.0
		Eggplant	19	4	15	21.1	78.9
		All host plants	72	5	67	6.9	93.1
LiaoCheng	2005	Cucumber	33	33	0	100.0	0.0
0		Cotton	39	38	1	97.4	2.6
		All host plants	72	71	1	98.6	1.4

TABLE 1. DATA ON WHITEFLY COLLECTION AND BIOTYPE DETERMINATION BASED ON THE MARKERS.

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Location	Year	Host plant	Total individuals	Number of biotype		Percentage of biotype (%)	
				В	Q	В	Q
	2006	Winter squash	27	23	4	85.2	14.8
		Cotton	28	13	15	46.4	53.6
		Morning glory	11	10	1	90.9	9.1
		Eggplant	26	13	13	50.0	50.0
		Cucumber	24	14	10	58.3	41.7
		All host plants	116	73	43	62.9	37.1
	2007	Cotton	54	0	54	0.0	100.0
		Eggplant	51	8	43	15.7	84.3
		All host plants	105	8	97	7.6	92.4
	2008	Japanese hop	30	1	29	3.3	96.7
		Cotton	30	0	30	0.0	100.0
		Eggplant	26	1	25	3.8	96.2
		All host plants	86	2	84	2.3	97.7
LinYi	2006	Cotton	28	28	0	100.0	0.0
		Eggplant	29	29	0	100.0	0.0
		Cucumber	8	8	0	100.0	0.0
		All host plants	65	65	0	100.0	0.0
	2007	Cucumber	13	11	2	84.6	15.4
		Cotton	21	21	0	100.0	0.0
		Eggplant	20	19	1	95.0	5.0
		All host plants	54	51	3	94.4	5.6
	2008	Cotton	26	1	25	3.8	96.2
		Eggplant	20	4	16	20.0	80.0
		All host plants	46	5	41	10.9	89.1
ZaoZhuang	2005	Cucumber	30	30	0	100.0	0.0
	2006	Cucumber	29	29	0	100.0	0.0
	2008	Pepper	26	0	26	0.0	100.0
		Japanese hop	29	0	29	0.0	100.0
		Cotton	29	0	29	0.0	100.0
		Eggplant	24	0	24	0.0	100.0
		All host plants	108	0	108	0.0	100.0

TABLE 1. (CONTINUED) DATA ON WHITEFLY COLLECTION AND BIOTYPE DETERMINATION BASED ON THE MARKERS.

mtCOI fragment (about 620bp) was first cleaved by the restriction endonucleases VspI (Khasdan et al. 2005) and then the uncut fragment was cleaved by the restriction endonucleases StuI(Ueda 2006). All of the mtCOI that could be cut by VspI should be biotype Q and mtCOI cut by StuIshould be B.

RESULTS AND DISCUSSION

Our results shown in Fig. 1 and Table 1 revealed the following: In 2005, the biotype of *B. tabaci* populations in JiNan, LiaoCheng, ZaoZhuang were determined and biotype Q was only found in LiaoCheng in very low proportion (1.4%). In 2006, biotype Q was found in DeZhou (38.0%), ZiBo (38.9%), ShouGuang (62.6%), LiaoCheng (37.1%) and was absent in JiNan, LinYi and ZaoZhuang. By 2007, biotype Q dominated in most locations, DeZhou (71.8%), ZiBo (92.7%), ShouGuang (96.0%), LiaoCheng (92.4%), and bio-

type Q also was found in JiNan (61.7%) and LinYi (5.6%). In 2008, Q biotype comprised 100.0%, 93.1%, 100.0%, 93.1%, 97.7%, 89.1% and 100.0% of the *B. tabaci* population in DeZhou, ZiBo, Shou-Guang, JiNan, LiaoCheng, LinYi and ZaoZhuang, respectively.

The present results are consistent with previous research on cotton and eggplant (Chu et al. 2010). These results suggest that the changes of *B. tabaci* biotypes occurred not only on the cotton and eggplant but also on the other plants including crops and weeds in Shandong Province during the past several years.

The displacement mechanism of biotypes Qand B remains uncertain, though the increase of biotype Q in many countries may be due to application of insecticides because biotype Q possesses greater resistance to insecticides than biotype B, but ecological and economic factors should be also considered. For example, the host plants that biotype Q preferred also might mediate the competition of B and Q. Multiple introduction of biotype Q from the other provinces or regions through human activities or natural sources should not be neglected.

Overall, our present result showed that the biotypes of *B. tabaci* changed greatly and *B.* tabaci biotype Q has been increasing on all kinds of hosts during the past several years. The displacement of biotypes B by Q has been occurring in the province as a whole. The speed of the displacement of biotypes B and Q was fast and essentially a cryptic invasion (Geller et al. 1997) because the biotypes are morphologically indistinguishable. The displacement mechanism should be further researched and such knowledge might guide the application of the insecticides or adjustment of the crops to effectively control the pest. Differentiation of *B. tabaci* biotypes is important, and molecular markers are important discrimination tools.

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