DIFFERENCES IN THE DETOXIFICATION METABOLISM BETWEEN TWO CLONAL LINEAGES OF THE APHID Myzus persicae (SULZER) (HEMIPTERA: APHIDIDAE) REARED ON TOBACCO (Nicotiana tabacum L.)

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ABSTRACT

Myzus persicae (Sulzer) is a highly polyphagous aphid species, with a subspecies (M. persicae nicotianae) well adapted to tobacco (Nicotiana tabacum L.). We evaluated the effect of this host plant on the aphid performance and detoxification enzymes, in order to test the participation of xenobiotic metabolism on the ability of this aphid to overcome the tobacco chemical defences. Two genotypes, one corresponding to the only M. persicae nicotianae genotype reported in Chile on tobacco, and one genotype belonging to M. persicae sensu stricto were reared on tobacco and pepper (Capsicum annuum L.), respectively. M. persicae nicotianae showed a significantly higher intrinsic rate of increase (rm) on pepper than on tobacco, and M. persicae s.s. performed similarly, but with no reproduction at all on tobacco. In order to evaluate the effect of tobacco on detoxification enzymes, esterases, glutathione S-transferases (GST) and cytochrome P-450 monoxygenases (MO) were determined in both selected aphid genotypes after 12, 24, 36, 48 and 72 h of rearing on tobacco and pepper. M. persicae nicotianae exhibited the higher total esterase activities when reared on tobacco than on pepper after 48 h of rearing, while the activities of GST and MO did not show any significant difference between host-plants and duration of treatment. For M. persicae s.s., no significant differences were observed among host-plants for the studied enzymes. These results suggest a participation of the esterases, on the ability of this M. persicae nicotianae to overcome the tobacco defences.

Key words: Myzus persicae nicotianae, Myzus persicae sensu stricto, detoxification enzymes, esterases, glutathione S-transferases, monoxygenases, intrinsic rate of increase.

INTRODUCTION

Aphids are ubiquitous herbivorous insects that frequently constitute agricultural pests. Among the most relevant aphid pests the peach-potato aphid, Myzus persicae (Sulzer), is a polyphagous insect able to utilise a wide range of weeds and crops belonging to more than 40 plant families (Blackman and Eastop, 2000). The tobacco aphid, Myzus persicae nicotianae, is a subspecies particularly well adapted to tobacco (Nicotiana tabacum L.), unlike M. persicae sensu stricto (Blackman, 1987; Eastop and Blackman, 2005; Blackman et al., 2007). Interclonal variation in the performance of M. persicae on different host-plants has been reported, but M. persicae s.s. frequently shows a poor performance on tobacco in relation with M. persicae nicotianae (Nikolakakis et al., 2003; Olivares-Donoso et al., 2007). Interestingly, the tobacco aphid is also found on other host-plants, including peach Prunus persica, field crops of potato (Solanum tuberosum L.), pepper (Capsicum annuum L.) and cabbage (Brassica oleracea L.), as well as weeds (Semtner et al., 1998; Blackman et al., 2007; Margaritopoulos et al., 2007). Since wild and cultivated tobacco species show a number of anti-herbivore defences, the use of tobacco by M. persicae nicotianae should involve a fitness trade-offs on this host-plant, which could be based on the anti-herbivore defences produced by tobacco (Heidel and Baldwin, 2004), among which sugar esters (Neal et al., 1994; Xia and Johnson, 1997; Xia et al., 1997; Johnson et al., 2002; Puterka et al., 2003) and alkaloids (Thurston et al., 1966) produced by glandular trichomes are the major component of resistance against aphids. Those cuticular allelochemicals have been suggested as targets for tobacco breeding programs against M. persicae nicotianae on tobacco (Johnson et al., 2002; Goundoudaki et al., 2003), and could be related with the poor performance reported for M. persicae s.s. comparative with M. persicae nicotianae on this crop (Nikolakakis et al., 2003).
In Chile, *M. persicae nicotianae* was not observed or reported until 1998 (Fuentes-Contreras et al., 2004). A previous work has shown the presence of a single predominant genotype on tobacco, which has been found to be widespread not only in Chile (Fuentes-Contreras et al., 2004), but also observed in high frequencies in USA, Brazil and Argentine populations (Zepeda-Paulo et al., 2010).

The success of *M. persicae* colonizing different host-plants has been related to the presence of aphid enzymatic mechanisms of detoxification, which are responsible for the metabolization of host-plant allelochemicals (Francis et al., 2005; 2006). Therefore, the performance of *M. persicae nicotianae* on tobacco should have a genetic/biochemical base, which could be related to the ability of this subspecies to colonize a well defended host plant (*i.e.* tobacco with glandular trichomes and cuticular sucrose esters).

We hypothesise that the *M. persicae nicotianae* enzymatic detoxification system plays a role in the ability of this aphid to overcome the tobacco defences. Hence, the aim of this work was to compare the activity of three well-known enzymes that detoxify xenobiotics in insects: esterases (EST), glutathione S-transferases (GST) and cytochrome P-450 monooxygenases (MO) in two aphid clonal lineages reared on contrasting hosts (chemically defended and non-defended plants). The study was conducted comparing the performance and detoxification activity in two highly common field genotypes of *M. persicae*: the Chilean single predominant genotype belonging to *M. persicae nicotianae* and one *M. persicae s.s.* genotype chosen based on its higher genetic distance shown with respect to *M. persicae nicotianae* genotype.

**MATERIALS AND METHODS**

**Aphids and plants**

A total of 59 individual aphids were sampled in Central Chile. Thirty-six individuals were collected from *N. tabacum*, 12 individuals were collected from weeds including *Brassica rapa* L. (Brassicaceae), *Datura stramonium* L. (Solanaceae), *Polygonum persicaria* L. (Polygonaceae), and *Chenopodium album* L. (Chenopodiaceae), and eleven individuals were collected from crops including *Beta vulgaris* var. *saccharifera* L. (Chenopodiaceae), *Prunus persica* (L.) Batsch. (Rosaceae), and *Prunus domestica* L. (Rosaceae).

Once in the laboratory, *M. persicae nicotianae* aphids collected from tobacco were individually reared on *N. tabacum* var. burley var. BY64 (Profigen do Brazil, Brazil), while *M. persicae s.s.* aphids collected from weeds and non-tobacco crops were individually reared on pepper (*C. annuum* var. *grossum*) cv. Resistant (PetoSeed, Chile). Those host-plants were chosen because aphids exhibit their optimal growth (Olivares-Donoso et al., 2007).

Aphids and plants were kept in climate rooms under controlled conditions (20 ± 0.1 °C, 16:8 photoperiod), which ensure the parthenogenetic reproduction of aphids. Hence, each individual produced a genetically identical colony or clonal lineage. Samples from each clonal lineage were used to confirm species following taxonomic keys. Before experiments, aphids were left to reproduce on their optimal host-plants for at least three generations, to prevent any maternal effect or grand-maternal effects, a consequence of the telescopic reproduction in aphids.

**Microsatellite genotyping**

Once clonal lineages were established, they were genotyped using 10 microsatellite loci previously described (Sloane et al., 2001; Wilson et al., 2002). The DNA isolation was carried out following the “salting-out” method (Sunnucks and Hales, 1996) using one individual from each of the 59 laboratory clonal lineages. Microsatellite loci were amplified by the polymerase chain reaction (PCR) and the products separated through an electrophoresis in polyacrylamide gels stained with silver nitrate. The size of each fragment (microsatellite alleles) was determined by size comparison with the sequence of the pGEM -3ZF(+) vector (Promega, Madison, Wisconsin, USA) loaded in the same gel.

**Evaluation of aphid intrinsic rate of increase**

Synchronized second instar aphids from the selected *M. persicae nicotianae* and *M. persicae s.s.* genotypes were obtained from clip-cages placed on leaves of their respective host-plants (tobacco or pepper). Nymphs were gently removed with a paint brush and carefully placed on leaves of either tobacco or pepper. All evaluations were performed on plants maintained in a growth chamber with controlled temperature and photoperiod (20 ± 0.1 °C, 16:8 h). The pre-reproductive duration (T) and fecundity (M) of the two *M. persicae* genotypes on both tobacco and pepper, according to the method described by Wyatt and White (1977). Each experiment was replicated (n = 12) and differences analyzed with Kruskal-Wallis test (Sokal and Rohlf, 1995).

**Host-plant transfer experimental design**

Synchronized wingless adult aphids from the chosen *M. persicae nicotianae* and *M. persicae s.s.* genotypes (clonal lineages) were reared on tobacco and pepper, respectively, and then transferred to: (i) their same optimal host-plant (control), and to (ii) a different plant (treatments from tobacco to pepper and from pepper to tobacco). All
evaluations were performed using plants maintained in a growth chamber with controlled temperature and photoperiod (20 ± 0.1 ºC, 16:8 h). Twelve, 24, 36, 48 and 72 h after the aphids were transferred to a new host-plant; they were immediately used for enzymatic determinations. The enzymatic determinations for each time period mentioned above were conducted in three blocks of 10 replicates each (30 each time).

Detoxification enzymes

Total EST activity was determined by photometry (Devonshire et al., 1992), while GST and MO activity were determined by fluorometry (De Sousa et al., 1995; Nauen and Stumpf, 2002). Total proteins were quantified following the Bradford (1976) method using bovine serum albumin as standard.

Esterases. The total activity of EST was determined using α-naphyl-acetate (α-NA) as substrate (Devonshire et al., 1992). One single aphid from each treatment was homogenized in 100 µL buffer PBS/TWEEN 20. Then, 25 µL of each homogenized were transferred to a transparent 96-wells microplate, adding 25 µL of buffer PBS/TRITON X-100. Next, 150 µL of α-NA (in buffer PBS/TRITON X-100) were added and incubated for 5 min at room temperature. The reaction was stopped with 25 µL of dying solution (Fast blue RR salt) and left in darkness for 20 min at room temperature. The absorbance was read at 620 nm using a Victor 3 Wallac microplate reader (Perkin Elmer Life and Analytical Sciences, USA), and the results were expressed as nmoles α-naphtol min⁻¹ mg⁻¹ protein.

Glutathione S-transferase. The activity of GST was determined using monochlorobimane as substrate (Nauen and Stumpf, 2002). Five aphids of the same clonal lineage from each treatment were homogenized in 300 µL of buffer PBS 1X and centrifuged at 10 000 RPM during 5 min at 4 ºC. Then, 30 µL of the supernatant (enzymatic extract) were transferred to a black 96-wells microplate, adding 30 µL of PBS 1X and 150 µL of substrate (reduced Glutathione in HEPES buffer). Next, 20 µL of monochlorobimane (in HEPES buffer) were added to each well, incubating at room temperature for 20 min. The fluorescence was measured at 390 nm for excitation and 465 nm for emission using a microplate reader. Because the product bimane-glutathione adduct (GS-mCB) is not commercially available, the results were expressed as fluorescence units of GS-mCB per insect (Nauen and Stumpf, 2002).

Cytochrome P-450 monooxygenases. The activity of MO was determined using the 7-ethoxycoumarine-O-deethylase activity (De Sousa et al., 1995). Three aphids of the same genotype from each condition were gently crushed in 150 µL of buffer HEPES in a black 96-wells microplate, adding 100 µL of substrate (0.4 mM ethoxycoumarin in 50 mM buffer phosphate, pH 7.2). For the controls, the same volume of buffer glycine/ethanol (50% v/v) was added to avoid the reaction. Then, the reaction was incubated for 4 h at 37 ºC and stopped by the addition of 100 µL of buffer glycine/ethanol (50% v/v). Fluorescence was read at 390 nm for excitation and 460 nm for emission, using a microplate reader. The results were expressed in amount of fluorescent product 7-hydroxycoumarin, ng min⁻¹ (De Sousa et al., 1995).

Data analysis

Enzymatic activities of EST, GST and MO were analyzed separately for each aphid belonging to the chosen M. persicae nicotianae and M. persicae s.s. clonal lineages. Independent variables were: i) transfer between host-plant species (tobacco to tobacco as control and tobacco to pepper for M. persicae nicotianae) (pepper to pepper as control and pepper to tobacco for M. persicae s.s.), and ii) hours since host-plant species transfer (12, 24, 36, 48 and 72 h), using an ANOVA for repeated measures on time (Sokal and Rohlf, 1995).

RESULTS

The genotyping showed that most microsatellite alleles were in the size range previously described for M. persicae (Terradot et al., 1999; Sloane et al., 2001; Wilson et al., 2002; Fuentes-Contreras et al., 2004; Blackman et al., 2007; Margaritopoulos et al., 2007; Zepeda-Paulo et al., 2010). After combining the genotypes from all the amplified loci, 15 multilocus genotypes were identified among the 59 clonal lineages. Fourteen genotypes were found on weeds and non-tobacco crops, while a single genotype was detected on tobacco, as previously reported. Based on the proportion of shared alleles, the most genetically distant M. persicae s.s. genotype in relation to the single M. persicae nicotianae genotype was selected for further comparisons.

Aphid intrinsic rate of increase (rₘ) was significantly lower for M. persicae s.s. reared on tobacco, where no reproduction was observed in T and 2T. Also, both M. persicae nicotianae (rₘ = 0.259 ± 0.011) and M. persicae s.s. (rₘ = 0.282 ± 0.005) reared on pepper showed a significantly higher rₘ than M. persicae nicotianae on tobacco (rₘ = 0.181 ± 0.018) (H₂, ₃₄ = 14.95, P < 0.001) (Table 1).

The enzymatic activity of EST showed no significant differences between host-plant species for both M. persicae nicotianae (F₁, ₅₈ = 3.4, P = 0.07) and M. persicae s.s. (F₁, ₅₈ = 0.7, P = 0.39). However, there were
significant differences between hours since host-plant transfer for both *M. persicae nicotianae* ($F_{4, 232} = 4.6, P = 0.001$) and *M. persicae s.s.* ($F_{4, 232} = 2.6, P = 0.04$) aphids. Similarly, there was a significant interaction between host-plant transfer and hours since host-plant transfer effects for *M. persicae nicotianae* ($F_{4, 232} = 5.2, P < 0.001$), but not for *M. persicae s.s.* ($F_{4, 232} = 2.0, P = 0.09$). Multiple comparisons revealed that such statistical interaction was based on a significant increase of EST activity of *M. persicae nicotianae* genotype at 48 h since transfer from tobacco to tobacco in relation to all remaining hours (LSD test, $P < 0.05$), which was not observed when transferred from tobacco to pepper at any hours since host-plant transfer (Figure 1).

Both aphid clonal lineages on pepper showed a significantly higher $r_m$ than *M. persicae nicotianae* on tobacco ($H_{2, 34} = 14.95, P < 0.001$).

### Table 1. Intrinsic rate of increase ($r_m$) of *Myzus persicae* s.s. and *M. persicae nicotianae* reared on tobacco and pepper (n = 12).

<table>
<thead>
<tr>
<th>Aphids</th>
<th>On pepper</th>
<th>On tobacco</th>
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<tr>
<td><em>M. persicae nicotianae</em></td>
<td>0.259 ± 0.011</td>
<td>0.181 ± 0.018</td>
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<tr>
<td><em>M. persicae s.s.</em></td>
<td>0.282 ± 0.005</td>
<td>NR</td>
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NR: no reproduction was observed.

**Table 1. Intrinsic rate of increase ($r_m$) of *Myzus persicae* s.s. and *M. persicae nicotianae* reared on tobacco and pepper (n = 12).**

![Graphs showing enzyme activities of *M. persicae nicotianae* and *M. persicae s.s.*](image)

**Figure 1. Enzyme activities of *Myzus persicae nicotianae* and *M. persicae s.s.* (black on tobacco; grey on pepper).**

a and b: Specific esterases (EST) activities (nmoles $\alpha$-naphtol min$^{-1}$ mg$^{-1}$ protein) ($n = 30$ enzymatic determinations for each time); c and d: specific glutathione S transferase (GST) activities (fluorescence units of GS-mCB per insect) ($n = 30$ enzymatic determinations for each time); e and f: specific cytochrome P-450 monooxygenases (MO) activities (ng of fluorescent product 7-hydroxycoumarin min$^{-1}$) ($n = 30$ enzymatic determinations each time).

*Significant differences ($F_{4, 232} = 5.2, P < 0.001$) (Error bars: standard error).
The enzymatic activity of GST showed no significant differences between host-plant species for both *M. persicae nicotianae* (F1, 58 = 2.7, P = 0.11) and *M. persicae s.s.* (F1, 58 = 1.02, P = 0.32). However, there were significant differences between host-plant species for *M. persicae nicotianae* (F4, 232 = 24.1, P < 0.001) and *M. persicae s.s.* (F4, 232 = 20.4, P < 0.001) aphid genotypes. No significant interactions between host-plant transfer and hours since host-plant transfer effects for both *M. persicae nicotianae* (F4, 232 = 1.3, P = 0.28) and *M. persicae s.s.* genotype (F4, 232 = 0.96, P = 0.43) were found. Multiple comparisons showed a significant decrease of GST activity at 72 h since host-plant transfer in relation to all remaining hours (LSD test, P < 0.05) for both *Myzus* subspecies and host-plant transfer treatments (Figure 1).

Cytochrome P-450 monoxygenases activity showed no significant differences between host-plant species for both *M. persicae nicotianae* (F1, 58 = 2.0, P = 0.16) and *M. persicae s.s.* (F1, 58 = 0.18, P = 0.67). However, there were significant differences between hours since host-plant transfer for both *M. persicae nicotianae* (F4, 232 = 74.0, P < 0.001) and *M. persicae s.s.* (F4, 232 = 41.6, P < 0.001) aphid. No significant interactions between host-plant and time treatment for both *M. persicae nicotianae* (F4, 232 = 1.9, P = 0.11) and *M. persicae s.s.* (F4, 232 = 0.34, P = 0.85) were found. Multiple comparisons showed a significant decrease of MO activity at 72 h since host-plant transfer (Least Significant Difference test, P < 0.05) for both subspecies and host-plant transfer treatments (Figure 1).

**DISCUSSION**

Evidences of the involvement of detoxification enzymes in the ability of *M. persicae s.s.* to face secondary metabolites (such as glucosinolates and isothiocyanates) from Brassicaceae plants have been reported (Francis et al., 2005; 2006). Those studies have shown that aphids fed for 2-wk on *Brassica* plants have evidenced an induction of GST activities, which has been discussed in terms of host-plant transfer and hours since host-plant transfer effects for both *M. persicae nicotianae* (F4, 232 = 24.1, P < 0.001) and *M. persicae s.s.* (F4, 232 = 20.4, P < 0.001) aphid genotypes. No significant interactions between host-plant transfer and hours since host-plant transfer effects for both *M. persicae nicotianae* (F4, 232 = 1.3, P = 0.28) and *M. persicae s.s.* genotype (F4, 232 = 0.96, P = 0.43) were found. Multiple comparisons showed a significant decrease of GST activity at 72 h since host-plant transfer in relation to all remaining hours (LSD test, P < 0.05) for both *Myzus* subspecies and host-plant transfer treatments (Figure 1).

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In our study we did not found any significant differences in GST or MO activity attributable to the change of host-plant, but a significant increase in the total activity of EST was found after 48 h of transferring *M. persicae nicotianae* aphids from tobacco to tobacco. It is important to note here, that our results were obtained after a 3-d rearing period on tobacco and pepper, while the involvement of GST in response to plant allelochemicals was reported after 2-wk of rearing on *Brassica* plant and 1-wk of rearing on artificial diets (Francis et al., 2005; 2006).

This higher total activity of EST observed after 48 h on tobacco, could be related with the settlement phase of the aphids on the tobacco leaf surface. After transferring aphids, they get in contact with tobacco trichomes, which release sugar esters known to produce deleterious effects on *M. persicae nicotianae* performance (Xia and Johnson, 1997; Xia et al., 1997; Johnson et al., 2002; Puterka et al., 2003; Goundoudaki et al., 2003). At present, there are no studies about the mechanisms of toxicity of sucrose and glucose esters from tobacco trichomes on insects, although it has been proposed that they could disrupt the integrity of cellular membranes, and uncouple oxidative phosphorylation, similar to the action of insecticidal soaps (Neal et al., 1994; Xia et al., 1997; Puterka et al., 2003). Similarly, it is unclear how these leaf surface cuticular components, bound to water soluble sugars could cross the aphid cuticle and interact with enzymes inside the aphid. However, it is possible that microbial activity in the phyloplane could naturally break-down the sugars after several hours, releasing fatty acids with lipophilic acyl chains and free hydroxyls that could affect insect cuticle (Puterka et al., 2003). The increase of total EST activity after 48 h on tobacco observed in our experiments could be related with a time lag necessary for the sugar esters to be modified and induce the increase of enzyme activity inside the aphid body.

Detoxification of plant allelochemicals in insects is frequently related with insecticide detoxification, often sharing the same enzymatic mechanisms (Li et al., 2007). One of the main mechanisms of insecticide resistance in *M. persicae* is the amplification of one of two closely related genes, which express carboxylesterase enzymes (E4 and FE4). These carboxylesterases hydrolyse or sequester insecticidal esters (Devonshire et al., 1992; 1997; Xia et al., 1994; Puterka et al., 2003). The increase of total EST activity after 48 h on tobacco observed in our experiments could be related with a time lag necessary for the sugar esters to be modified and induce the increase of enzyme activity inside the aphid body.

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M. persicae s.s. and M. persicae nicotianae show the same mechanism of insecticide resistance based on E4 and FE4 activity (Field et al., 1994). We have found that M. persicae nicotianae could be able to increase the total EST activity following its transfer to tobacco plants. However, since epigenetic control of E4 and FE4 expression, based on chromosomal translocation or gene methylation processes, involve heritable and more permanent changes in the aphid genome, it is unlikely that they could be involved in the transient increase of total EST activity that we observed within a scale of hours. In addition, processes, involve heritable and more permanent changes on chromosomal translocation or gene methylation.

Following its transfer to tobacco plants. However, since epigenetic control of E4 and FE4 expression, based on chromosomal translocation or gene methylation processes, involve heritable and more permanent changes in the aphid genome, it is unlikely that they could be involved in the transient increase of total EST activity that we observed within a scale of hours. In addition, the M. persicae nicotianae genotype used in this study is the same previously reported to be predominant in Chile (Fuentes-Contreras et al., 2004), which was characterized as a chromosome translocated R1 genotype with a rather low to mid level of total EST activity. Similarly, the M. persicae s.s. genotype selected for this study showed the same mechanism of detoxification enzymes, pheromone expression, based on the detoxification enzymes, which are involved in the aphid response, which is a point to consider during host specialization to tobacco.

**ACKNOWLEDGEMENTS**

This work is part of the activities of the Centre for Research on Plant-Insect Interactions (iCROP), and was funded by FONDICYT 3020051 and 1050644 grants. Partial support was also obtained from CONICYT-PBCT Anillos ACT-38 grant. We also thank Luis Castañeda for his help with statistical analysis and criticisms.

**CONCLUSIONS**

The findings here reported suggest the presence of fitness costs associated with the use of tobacco as a host-plant, and the involvement of esterases as an aphid response, which was observed following the host plant transfer to tobacco (a defended plant). Detoxification enzymes seem to be one of the main traits allowing M. persicae nicotianae to overcome tobacco defences, which is a point to consider during host specialization to tobacco.

**RESUMEN**

Diferencias en el metabolismo de detoxificación entre dos linajes clonales del áfido Myzus persicae (Sulzer) (Hemiptera: Aphididae) criados sobre tabaco (Nicotiana tabacum L.). Myzus persicae (Sulzer) es un áfido polífago que incluye a Myzus persicae nicotianae, una subespecie altamente adaptada sobre tabaco (Nicotiana tabacum L.). Evaluamos el efecto del tabaco sobre el desempeño biológico y sobre determinadas enzimas de detoxificación en áfidos, para estudiar su participación en la capacidad de Myzus persicae nicotianae de superar las defensas químicas del tabaco. Dos genotipos de Myzus persicae, uno correspondiente al único genotipo perteneciente a M. persicae nicotianae reportado en Chile sobre tabaco y el otro correspondiente a M. persicae sensu stricto, fueron criados sobre tabaco y pimentón (Capsicum annuum L.), respectivamente. Myzus persicae nicotianae mostró una tasa intrínseca de crecimiento (rm) significativamente mayor sobre pimentón que sobre tabaco, mientras que M. persicae s.s. no se reprodujo sobre tabaco. Para evaluar los efectos del tabaco sobre el metabolismo de detoxificación, se determinó la actividad de las enzimas esterasas, glutación S-transferasas (GST) y citocromo p450 monooxigenasas (MO), transcurridas 12, 24, 36, 48 y 72 h desde que ambos genotipos fueron transferidos desde su planta hospedera óptima a una nueva planta de tabaco y pimentón. En M. persicae nicotianae, se observaron diferencias significativas en las esterasas cuando se alimenta sobre tabaco, transcurridas 48 h desde el inicio del experimento. Para M. persicae s.s. no se observaron diferencias significativas en las actividades de las enzimas esterasas. Estos resultados sugieren la participación de las esterasas en la capacidad de Myzus persicae nicotianae para enfrentar las defensas químicas del tabaco.

**Palabras clave:** Myzus persicae nicotianae, Myzus persicae s.s., enzimas de detoxificación, esterasas, glutación S transferasa, monooxigenasas, tasa intrínseca de crecimiento.
LITERATURE CITED


