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Resistance Status of House Flies (Diptera: Muscidae) from Southeastern Nebraska Beef Cattle Feedlots to Selected Insecticides

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This article is the copyright property of the Entomological Society of America and may not be used for any commercial or other private purpose without specific written permission of the Entomological Society of America.
The house fly, *Musca domestica* L., is an important pest of humans and domesticated animals and is also a vector of both human and animal diseases. Greenberg (1971) lists more than 100 pathogen species harbored by house flies, and, of these, 65 are known to be transmitted by the flies. More recently, house flies have been shown to vector enterohemorrhagic colitis caused by *Escherichia coli* 0157:H7 and *Yersinia pseudotuberculosis* (Sasaki et al. 2000 and Zurek et al. 2001). The fly breeds in filth of all kinds, and it is often the primary cause of lawsuits in areas in which urban growth infringes upon agricultural areas, a fact that has greatly increased the pressure on livestock operators to effectively control house fly populations (Thomas and Skoda 1993). Historically, house fly management has been directed at adult populations and relied heavily on chemical control (Campbell 1993). However, the use of insecticides in house fly management programs is becoming more costly, more regulated, and less effective because of resistance problems (Geden et al. 1992). The chronological sequence of pesticide use for control of livestock insects in Nebraska included the chlorinated hydrocarbons such as DDT and methoxychlor in the 1950s and 1960s, the phosphates in the late 1960s and 1970s, including tetrachlorvos, naled, and dichlorvos. Since the late 1980s, pyrethroids have largely replaced the phosphates. Permethrin and fenvalerase were used in ear tags for horn fly control until resistance developed, and permethrin is widely used for house fly and stable fly control at feedlots, dairies, swine, and poultry units. Although the house fly has demonstrated the ability to develop resistance to most insecticides approved for use against it (Bull and Pryor 1991), it is certainly not resistant throughout its geographic range. Susceptible populationsexist (Scott et al. 1989), and regional assessment of susceptibility of local populations of house flies to different insecticides can yield important information for integrated fly control programs that use available insecticides in ways that minimize resistance problems. Pospischil et al. (1996) reported multiresistance in German house fly populations selected from regions with known control problems to pyrethroids, organophosphates, and carbamate insecticides. Scott et al. (2000) found considerable insecticide resistance variation in house fly populations collected from New York caged layer poultry houses. Overall, there was correlation between insecticide use histories and levels of resistance. Kaufman et al. (2001), however, found little variation in resistance levels in fly populations at New York dairy farms. The highest levels of resistance in fly populations collected from both poultry and dairy units was tetrachlorin-
phos, permethrin, and cyfluthrin. Kristensen et al. (2001) also reported considerable variation in insecticide resistance in house fly populations collected from Danish farms with high resistance to pyrethroid organophosphate and carbamate insecticides in some house fly populations.

In assessing pest resistance to pesticides, it is important to use bioassay procedures that are sensitive enough to detect changes in susceptibility status of field populations as they occur. In this study, a direct contact and a residual bioassay were compared with field populations as they occur. In this study, a direct contact and a residual bioassay were compared with field populations as they occur. In this study, a direct contact and a residual bioassay were compared with field populations as they occur. In this study, a direct contact and a residual bioassay were compared with field populations as they occur.

Materials and Methods

Insecticides. All insecticides used were technical grade and diluted in acetone. Permethrin (94.6% [AI]) maximum 55% cis and minimum 45% trans) was provided by FMC Corp. (Philadelphia, PA), stiofos (99% [AI]) was supplied by Fermenta Animal Health (Kansas City, MO), and methoxychlor (95.6% [AI]) was obtained from Kineaid Enterprise (Nitro, WV).

House Flies. The insecticide-susceptible laboratory colony of house flies used in this study originated from the Beltsville W Strain (Livestock Insects Laboratory, Beltsville, MD). This strain was used as a standard reference for insecticide susceptibility because it has been isolated from insecticide exposure since 1969 (Pickens et al. 1972). House flies were collected from two counties in southeastern Nebraska: feedlot A in Saunders County and feedlot B in Lancaster County. Feedlot A had a record of recent insecticide use (adult fly control with permethrin using a mist blower), whereas feedlot B had not been sprayed for 2 yr. We were unable to obtain accurate historical records of insecticide applications from other cattle feeders in the region, although we determined that use of insecticides for fly control was minimal.

Field populations of house flies were used for the resistance bioassay because we felt they represented the situation at the time of collection. Adult house flies were collected by sweeping with an insect net, mostly around feedbunks but also in the general feedlot area. After collection, flies were transferred immediately to cages and transported to the laboratory. Bioassays were performed on the same day as capture, usually within 2 h. Both 3–5-d males and females were used in the bioassays, representing a random sample from field collections. Insecticide-susceptible laboratory flies were tested. After treatment, test insects were maintained at 25°C ± 1°C and photoperiod of 12:12 h (L:D).

Topical Exposure. House flies were separated into groups of 15 and placed in disposable cups before treatment. C cups containing 15 flies were assigned randomly to each of three tests, and within a test, to each dose treatment. Each bioassay consisted of three complete tests using three groups of 15 flies from each population for each treatment level in a test. For each test, serial dilutions were prepared from a stock solution to produce 5–10 dose levels. Dose ranges for each insecticide varied depending on the populations being tested. For permethrin, the doses used to test the susceptible population ranged from 0 to 0.05 µg/fly, and the doses used to test the field populations ranged from 0 to 0.25 µg/fly. For stiofos, doses ranged from 0 to 0.38 µg/fly (susceptible) and from 0 to 193.0 µg/field. For methoxychlor, doses ranged from 0 to 7.77 µg/field (susceptible) and from 0 to 608.0 µg/field (field). Acetone was used as the solvent for all the insecticides. All serial dilutions were prepared the day before being used.

Flies were anesthetized with CO₂ and 1 µl of the appropriate solution was applied to the notum of each fly using an Eppendorf micropipette (0.5–10.0 µl, Brinkmann Instruments, Westbury, NY). Controls were treated with acetone only. After treatment, flies were placed in paper cups (114 ml) that were covered with tulle cloth and secured with rubber bands. A water-saturated dental wick was placed on the bottom of each cup.

Each time a field fly population was tested, a simultaneous test was performed on the reference susceptible laboratory population. Before each bioassay, a sample of ~100 house flies was weighed to determine mean weights for each population on each bioassay date. This allowed us to convert lethal dose values from micrograms per fly to micrograms per milligram of body weight.

Residual Exposure. Serial dilutions were prepared from a stock solution to produce 5–10 insecticide concentrations. Acetone was used as the solvent for all insecticides. Each bioassay consisted of three complete tests using three groups of 20 flies from each population for each treatment level in a test. Concentration ranges for each insecticide varied depending on the population being tested. For permethrin, the concentrations used to test the susceptible population ranged from 0 to 0.12 µg/cm², and the concentrations used to test the field populations ranged from 0 to 6.0 µg/cm². For stiofos, concentrations ranged from 0 to 2.0 µg/cm² (susceptible) and from 0 to 3,000.0 µg/cm² (field). For methoxychlor, concentrations ranged from 0 to 2.0 µg/field (susceptible) and from 0 to 12,000.0 µg/cm² (field). Standard glass Petri dishes were used (interior bottom diameter of 9 cm). For each concentration, 2 ml of the appropriate dilution was applied to the bottoms of three Petri dishes and distributed uniformly by gently rotating the dishes while drying. Acetone alone was applied in a similar manner to each of three control dishes. After treatment, the dishes were allowed to air dry for 3 h and then covered and stored in darkness at room temperature for at least 24 h before use (no longer than 1 wk).

Before each bioassay, test flies were anesthetized using CO₂, separated into groups of 20 and placed into small, screened containers. After all insects had been grouped, flies were anesthetized lightly again and placed in the respective Petri dishes. Test flies were placed on the untreated Petri dish cover to allow them
Results

For both exposure to residues on glass and topical applications, insecticide exposure of house flies for 4 h produced less variable results than when flies were exposed for only 2 h. The 24-h exposure period resulted in high control mortality (>30%) of field-collected flies in many of the bioassays, regardless of technique. Therefore, only results from the 4-h exposure are presented.

Moderate levels of resistance to permethrin were found in house flies from the two feedlots sampled (Table 1). Residual exposure to permethrin on glass allowed for better discrimination between susceptible and resistant individuals which is reflected by the higher resistance ratios (7.3 and 7.0, for feedlots A and B, respectively) as compared with those obtained when the same populations were tested by topical application (4.9 and 3.8, for feedlots A and B, respectively). The insecticide mortality response of house flies from feedlot A was not significantly different from that of house flies from feedlot B.

The field populations tested were extremely resistant to stirofos, whether exposed to insecticide residues on glass or treated topically (Table 1). Insufficient data did not allow the development of significant regression lines because of low mortality (<50%) at the highest concentrations (residual exposure) or doses (topical exposure) tested, which were at least 20,000-fold higher than the LC50 and 2,560-fold higher than the LD50 of the susceptible laboratory colony that was tested simultaneously. Similar to observations with permethrin, the residual exposure provided higher relative resistance ratios.

Resistance to methoxychlor was also very high, both when flies were challenged with residues or by topical application of the insecticide. Dose–response lines could not be generated using topical applications because of low mortality (<50%) at the highest dose tested, 608 µg/ fly (i.e., 55.1 µg/mg body weight for flies from feedlot A). This was 930-fold higher than the LD50 of the simultaneously tested susceptible population. Although quantification of resistance to methoxychlor was not possible using this technique, resistance ratios were at least >930 and >345 for flies

<table>
<thead>
<tr>
<th>Insecticide</th>
<th>Technique</th>
<th>Population</th>
<th>n</th>
<th>Slope ± SE</th>
<th>LC50 (FL95%)</th>
<th>LD50 (FL95%)</th>
<th>RR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Permethrin</td>
<td>Topical exposure</td>
<td>Susceptible</td>
<td>360</td>
<td>4.16 ± 0.39</td>
<td>0.0007 (0.0006–0.0008)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Topical exposure</td>
<td>Feedlot A</td>
<td>356</td>
<td>2.73 ± 0.28</td>
<td>0.0034 (0.0021–0.0051)</td>
<td>4.9*</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Topical exposure</td>
<td>Susceptible</td>
<td>315</td>
<td>6.92 ± 0.82</td>
<td>0.0005 (0.0005–0.0006)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Topical exposure</td>
<td>Feedlot B</td>
<td>267</td>
<td>2.67 ± 0.33</td>
<td>0.0019 (0.0014–0.0025)</td>
<td>3.8*</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Residual exposure</td>
<td>Susceptible</td>
<td>473</td>
<td>4.65 ± 0.39</td>
<td>0.027 (0.022–0.032)</td>
<td>–</td>
<td>–</td>
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<tr>
<td></td>
<td>Residual exposure</td>
<td>Feedlot A</td>
<td>519</td>
<td>3.24 ± 0.30</td>
<td>0.198 (0.172–0.228)</td>
<td>7.3*</td>
<td>–</td>
</tr>
<tr>
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<td>Residual exposure</td>
<td>Susceptible</td>
<td>480</td>
<td>4.77 ± 0.41</td>
<td>0.026 (0.023–0.028)</td>
<td>–</td>
<td>–</td>
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<tr>
<td></td>
<td>Residual exposure</td>
<td>Feedlot B</td>
<td>450</td>
<td>3.37 ± 0.30</td>
<td>0.182 (0.153–0.216)</td>
<td>7.0*</td>
<td>–</td>
</tr>
<tr>
<td>Stirofos</td>
<td>Topical exposure</td>
<td>Susceptible</td>
<td>315</td>
<td>2.78 ± 0.27</td>
<td>0.0025 (0.0016–0.0040)</td>
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<td>–</td>
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<tr>
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<td>Topical exposure</td>
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<td>356</td>
<td>–</td>
<td>&lt;17.63</td>
<td>&gt;7.100</td>
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<td>Topical exposure</td>
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<td>–</td>
<td>&gt;6.40</td>
<td>&gt;2.560</td>
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<td>Residual exposure</td>
<td>Susceptible</td>
<td>420</td>
<td>2.86 ± 0.27</td>
<td>0.151 (0.064–0.270)</td>
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<td>Feedlot A</td>
<td>357</td>
<td>–</td>
<td>&gt;3.000</td>
<td>&gt;20.000</td>
<td>–</td>
</tr>
<tr>
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<td>Residual exposure</td>
<td>Feedlot B</td>
<td>359</td>
<td>–</td>
<td>&gt;3.000</td>
<td>&gt;20.000</td>
<td>–</td>
</tr>
<tr>
<td>Methoxychlor</td>
<td>Topical exposure</td>
<td>Susceptible</td>
<td>360</td>
<td>2.22 ± 0.20</td>
<td>0.0952 (0.0472–0.0743)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
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<td>Topical exposure</td>
<td>Feedlot A</td>
<td>270</td>
<td>–</td>
<td>&gt;55.10</td>
<td>&gt;930.0</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Topical exposure</td>
<td>Feedlot B</td>
<td>235</td>
<td>–</td>
<td>&gt;20.42</td>
<td>&gt;345.0</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Residual exposure</td>
<td>Susceptible</td>
<td>357</td>
<td>1.93 ± 0.18</td>
<td>1.590 (1.010–2.430)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Residual exposure</td>
<td>Feedlot A</td>
<td>410</td>
<td>1.54 ± 0.21</td>
<td>150.9 (79.8–245.9)</td>
<td>94.9*</td>
<td>–</td>
</tr>
<tr>
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<td>Residual exposure</td>
<td>Feedlot B</td>
<td>410</td>
<td>2.00 ± 0.22</td>
<td>207.5 (156.2–279.7)</td>
<td>130.5*</td>
<td>–</td>
</tr>
</tbody>
</table>

Resistance ratio (RR) = lethal concentration (LC) or lethal dose (LD) value of the field population divided by the corresponding value of the reference susceptible population.

* Resistance ratio significantly >1.0 (P > 0.050), as determined by the likelihood test for equality of the lethal concentration or lethal dose values used to calculate the ratio, followed by pairwise comparisons using nonoverlapping fiducial limits (Savin et al. 1977).

Table 1. Comparative toxicity of selected insecticides to house flies from southeastern Nebraska
from feedlots A and B, respectively. The resistance ratios generated for residue assays were 94.5 and 130.5 for feedlots A and B. These data suggest that field populations of house flies were less tolerant to residual contact than they were to direct topical application of Methoxychlor, which is contrary to what was observed for the other two insecticides.

Discussion

Moderate levels of resistance to permethrin were detected in the field-collected house flies from both feedlots using both topical applications or exposure to residues on glass. Farham et al. (1984) have correlated data from topical applications of permethrin and control failures in the field and found that control failures usually occur only when resistance ratios are >15-fold. In our study, the greatest resistance ratio observed was <5-fold, which suggests that permethrin should still be an effective insecticide for control of these house fly populations.

The field populations tested were extremely resistant to both stirofos and methoxychlor. Accurate quantification of resistance was not possible because of low mortality at the highest possible concentrations or doses. Therefore, it is reasonable to infer that the house fly populations tested are minimally affected by stirofos and methoxychlor, either as residues or direct application, and that adult house fly control with either of these insecticides would be largely ineffective. Cross-resistance to other organophosphate or chlorinated hydrocarbons is probable and should be investigated. It is probably not surprising that these house fly populations were resistant to methoxychlor and stirofos. Methoxychlor is a representative of the organochlorine insecticide group, and these were widely used for livestock insect control in the 1960s and 1970s, but is no longer in use, although still registered. Stirofos is an organophosphate insecticide, and these products were popular in the 1980s. Permethrin, a pyrethroid, became a popular livestock insect product in the late 1980s and is still popular today.

In preliminary experiments, exposure of house flies to insecticide residues on filter paper was attempted. However, this technique proved to be inappropriate because little or no mortality was observed in the susceptible laboratory population even at very high concentrations, especially for stirofos and methoxychlor. Apparently, these insecticides became bound to the filter paper thereby preventing exposure of the flies. Significant concentration-response regressions were generated only for permethrin, but the amount of insecticide applied to the filter paper had to be at least 1000-fold higher than residues on glass to elicit a response from the susceptible population. Similar results were observed in another study with stable flies, and a more detailed explanation of the difficulties of this technique is presented elsewhere (Marcon et al. 1997).

Both exposure to residues on glass Petri dishes and topical applications provided suitable results for testing the susceptibility of house flies to permethrin and stirofos. However, a comparison of the resistance ratios across techniques suggested that residual exposure to these two insecticides allows for greater expression of resistance compared with topical exposure. Therefore, it should allow for better discrimination between susceptible and resistant phenotypes (Dennehey et al. 1983). It is important to recognize that residues on glass more closely approximate the mode of exposure of flies to residual insecticides in feedlots. Also, this technique allows for dishes to be prepared in advance and sent to different locations for testing. Development of a diagnostic bioassay using residues on glass should significantly improve the efficiency of this technique in resistance monitoring (Kaufman et al. 2001).

For methoxychlor, the situation seemed to be reversed; that is, topically applied methoxychlor generated higher resistance ratios than did exposure to residues on glass. These results were not unexpected because during the course of the experiments, it was noted that the flies challenged with very high concentrations of methoxychlor on glass became entrapped in the accumulated residues, which most likely contributed to the observed mortality. This problem was not observed for the susceptible laboratory colony because much lower concentrations were needed to generate significant concentration-response curves. This apparent limitation of the technique should only be of concern if quantification of resistance is essential. For detection of resistance using a diagnostic concentration that discriminates between resistant and susceptible individuals, exposure to methoxychlor residues on glass should be a useful technique.

Insecticide susceptibility of these house fly populations greatly contrasts with that of stable flies collected at the same feedlots (Marcon et al. 1997), the latter being highly susceptible to permethrin, stirofos, and methoxychlor. As far as we were able to ascertain, recent insecticide use was minimal at the feedlots sampled. Therefore, the coexistence of highly resistant house fly populations and susceptible stable fly populations in the same feedlots may indicate a greater propensity for dispersal of house flies from areas with higher resistance frequencies.

Boxler and Campbell (1983) found a similar situation with house fly resistance to dichlorvos, which indicated movement of house flies between feedlots. Lysyk and Axtell (1986), in a capture–mark–release–recapture study, found considerable movement of house flies between poultry houses and dairies and into habitats that did not contain house fly breeding or feeding material. Their study indicates that house fly dispersal is considerable, even when habitats seem ideal. Methoxychlor and stirofos resistance may have been retained in the house fly populations as a result of lack of biotic potential disadvantages associated with resistance (Roush and Plapp 1982). However, insecticide resistance may not necessarily be associated with application exposure at the feedlots, but rather the result of migration of resistant house flies into these areas; house flies are more intensely ex-
posed to chemical treatments in domestic settings, and resistant individuals may move to feedlot areas.

These points have important implications for insecticide resistance management because, in practice, pest control efforts are generally the same for both species of flies (Campbell 1993). Further studies are needed to understand the relationship between insecticide resistance, biotic potential, and migration patterns of these two muscid species if we are to design effective control strategies for these pests. Furthermore, monitoring of feedlot fly populations for insecticide resistance should continue not only for the insecticides currently used but also for new insecticides that are developed for cattle pests control. Detecting cross-resistance to new insecticides before they are in common use would be beneficial to both the cattle industry and the insecticide industry. Also, detecting the loss of resistance to currently used insecticides, if it occurs, could be beneficial in developing future pest management strategies.

Acknowledgments

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