Aerial and Tidal Transport of Mosquito Control Pesticides into the Florida Keys National Marine Sanctuary.
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Abstract
This project was undertaken as the initial monitoring program to determine if mosquito adulticides applied along the Florida Keys cause adverse ecological effects in the Florida Keys National Marine Sanctuary (FKNMS). The study monitored the distribution and persistence of two mosquito adulticides, permethrin and dibrom (naled), during three separate routine applications by the Florida Keys Mosquito Control District. The approach was to determine if toxic concentrations of the pesticides entered the FKNMS by aerial drift or tidal transport. The amount of pesticide entering the FKNMS by way of aerial drift was monitored by collection on glass fiber filter pads, set on floats in a grid pattern on either side of the FKNMS. Permethrin was recovered from filter pads on the leeward side for each of the three applications, ranging from 0.5 to 50.1 μg/m² throughout the study. Tidal current transport was monitored by collection of surface and subsurface water samples at each grid site. Tidal transport of naled and dichlorvos (naled degradation product) was apparent in the adjacent waters of the FKNMS. These compounds were detected in subsurface, offshore water at 0.1 to 0.6 μg/l, 14 hr after application. Permethrin was not detected in offshore water samples; however, concentrations ranging from 5.1 to 9.4 μg/l were found in surface water from the canal system adjacent to the application route. Comparison of the observed environmental concentrations with toxicity data (permethrin LC-50, 96 hr for Mysis bahia = 0.02 μg/l) indicated a potential hazard to marine invertebrates in the canals with possible tidal transport to other areas.

Key Words: organophosphate, pyrethroid, naled, dichlorvos, permethrin, mosquito adulticides

Introduction
This project was undertaken to determine if the mosquito adulticides, permethrin and dibrom (naled), applied along the Florida Keys, in Monroe County, Florida, USA, could cause adverse ecological effects in the Florida Keys National Marine Sanctuary (FKNMS). Dibrom-14, active ingredient, naled (1, 2-dibromo-2, 2-dichloroethyl dimethyl ester, phosphoric acid), is a fast acting, nonpersistent organophosphate insecticide. Although hydrolysis occurs fairly rapidly in seawater (half-life @ pH 7 = 16 h, decreasing with increasing pH), the primary degradation product, dichlorvos (2, 2-dichlorovinyl dimethyl phosphate), is also toxic to aquatic invertebrates (Cox 2002). Both naled and dichlorvos are classified as very highly toxic to aquatic invertebrates, with an acute toxicity (96 hr LC50) for Mysis bahia of 8.8 μg/l for naled, and 19 μg/l for dichlorvos (Johnson and Finley, 1980, Peckenpaugh, et al., 1997; Edwards, J., 1999).
Therefore, environmental assessment of the effects of naled must include analyses and toxicological testing for dichlorvos as well.

Permethrin is a broad spectrum, synthetic pyrethroid insecticide, consisting of a mixture of cis and trans isomers. The cis isomer is highly toxic (mouse LC-50, 85 mg/kg), however, the commercial mixture is less toxic (LC-50 of 410 mg/kg) (Kidd and James, 1991). Permethrin is applied with a synergist, piperonyl butoxide (PBO) that inhibits the insect's natural metabolic defense against pyrethroids, greatly enhancing its insecticidal effectiveness (Cox, 1998). Although relatively nontoxic to mammals and birds, permethrin is very highly toxic to aquatic invertebrates and exhibits a high bioconcentration factor in aquatic organisms due to its insolubility in water. Schimmel et al., 1983, reported the 96 hr LC-50 to be 0.02 μg/l for mysid shrimp, 0.2 μg/l for penaeid shrimp and 2.2 μg/l for the silverside minnow, Menidia sp. The half-life of permethrin in water and sediment was found to be <2.5 days, and was dependent upon photolysis in water and on microbial degradation in sediments (Schimmel, et al., 1983).

Previous studies of thermal fog applications showed drift of naled and fenthion into nontargeted (no-spray) terrestrial habitats in the Florida Keys. Pesticide residues collected on cellulose pads exhibited up to 90 μg/m² naled at 15 m inside a Pineland no-spray zone, with naled drift detected up to 750 m into the no-spray zone. At 6 hr after application, naled concentrations on the pads diminished to about 50% that observed at 1.5 h. Fenthion was detected in about the same concentrations 50 m into a wildlife refuge, but none was detected at 500 m (Hennessey et al., 1992). These results show the potential for non-target marine organisms exposure to drift from mosquito adulticide applications.

A study of both permethrin and naled/dichlorvos was conducted in a mangrove community and residential water collection systems on Grand Cayman Island following applications over residential areas as well as coastal mangrove forests (Pierce and Henry, 1988). These results showed persistence of permethrin in mangrove tree leaves and in association with suspended sediment in the water cisterns, but none was detected in saltmarsh water pools. Although no naled or dichlorvos was detected in the mangrove community following application, residues of both were recovered from cistern water up to 16 hr after application, but no pesticide residues were detected in drinking water as it came out of the tap (Pierce and Henry, 1988).

Although naled and permethrin are considered to be relatively nonpersistent in an aquatic environment, even short exposure periods to pesticides can have adverse impacts on susceptible life stages of aquatic invertebrates. The mosquito larvicide, temephos, (an organophosphate) was found to cause 50% mortality in fiddler crab larvae through the first molt, five to seven days after exposure in the field for only 4 h, following ultra low volume (ULV) application in an intertidal saltmarsh (Pierce et al., 2000). This evidence for adverse effects on aquatic invertebrates from short-term exposure to a mosquito pesticide supports the need for establishing the distribution and persistence of pesticides in the FKNMS.

Methods and Materials

The study location was Key Largo, Florida, USA. The geographic location is at latitude 25.18040 and longitude -80.36617. The Florida Keys Mosquito Control District (FKMCD) provided both permanone as well as dibrom applications within the same area of Key Largo, (between mile marker (MM) #89 and MM #91). Dibrom was applied as ultra low volume (ULV) technical product Dibrom-14 (Valent USA Corp.), 0.75 oz/acre using C-47 aircraft. Permethrin was applied as the technical product, Permanone (4% cis + trans permethrin/8%
PBO, Agrevo Corp.) from trucks as ground ULV applied at 5.5 fl oz/acre. Prior to each test application, a grid of nine (9) sampling sites covering an area of approximately 1.5 km along shore to 1 km offshore, was established in both the Florida Bay side as well as the Atlantic side of Key Largo, to provide a control (windward) and a drift (leeward) sampling area, depending on the direction of the wind (Figure 1).

Samples collected at each of the eighteen (18) sampling sites consisted of:

a) Filter pads to determine pesticide deposition as a result of aerial drift, placed on floating platforms above the water surface to avoid water splashing on to the filter surface,

b) Water surface micro layer to determine if the pesticides accumulated from deposition onto the water surface, and

c) Water subsurface (about 20 cm depth) to determine the amount of pesticide in the water column to which marine organisms would be exposed.

A summary of the sample collection locations and times (Table 1) shows that samples were collected just prior to each pesticide application to monitor for residual pesticides, and again at 2 to 4 hr after each application. Persistence in the aqueous environment was monitored by collection of surface and subsurface water samples again at 12 hr intervals post application as deemed necessary for each study according to the procedures of Pierce et al., 1996.

Table 1. Sampling Design for Mosquito Pesticide Transport Study.

<table>
<thead>
<tr>
<th>Collection Timing</th>
<th>Number and location of samples collected for each pesticide</th>
</tr>
</thead>
<tbody>
<tr>
<td>Permanone (PM)</td>
<td>Number</td>
</tr>
<tr>
<td>Naled (AM)</td>
<td>Surface</td>
</tr>
<tr>
<td>Pre Application</td>
<td>3</td>
</tr>
<tr>
<td>3-6 hour</td>
<td>9</td>
</tr>
<tr>
<td>14-16 hour</td>
<td>9</td>
</tr>
<tr>
<td>22-24 hour</td>
<td>9</td>
</tr>
</tbody>
</table>

Water quality parameters were monitored for each site, including temperature, dissolved oxygen, pH, salinity and conductivity. A strict quality control program was followed including field blanks, spiked standard recoveries and surrogate recovery standards added to each sample. Sample custody sheets were maintained and samples processed according to the written standard operating procedure, as approved by the U.S. Environmental Protection Agency. Sample processing was initiated immediately in the field with the addition of dichloromethane solvent to initiate extraction and to reduce degradation of the parent compounds. Samples were brought back to Mote Marine Laboratory (MML) for further processing and gas chromatography-mass spectrometry (GC-MS) analysis.

GC-MS analyses were performed using a Varian Instruments (Sugar Land, TX) Saturn II ion trap Mass Spectrometer coupled with a Varian 3400 gas chromatograph. GC separations were performed on a 30 m glass capillary DB-5 MS column.

Pre-application samples for the first study, June 16, 1998, were collected simultaneously from the Atlantic and Bay sites at 17:00 to 18:40. The permanone was applied by ground ULV starting at 20:00, with the first series of filters, surface and sub-surface water samples collected from 22:00 to 01:00. Dibrom was subsequently applied by air ULV at 07:00 on 6/17/98, with the post application sampling occurring from 09:00 to 11:30. The 24 hr permanone (coincident
with the 12 hr dibrom) sampling occurred from 17:00 to 18:30 on 6/17/98. One final sampling was performed on 6/18/98 from 08:30 to 10:00 on both the Bay and Atlantic sites.

The second study (7/28/98) pre-application samples were collected from 15:00 to 17:00 on both the Bay and Atlantic sides simultaneously, with the permethrin application following at 20:00. The post-application samples were collected from 22:00 to 00:45 on both the Bay and Atlantic sides concurrently. Dibrom was applied the next morning at 07:00. The first pre-application sampling for the Bay side was obtained from 9:45 to 11:15, however, due to boat problems; the Atlantic side was not sampled until 12:38 to 14:00, delaying the post-application samples for that area. The second dibrom sample collection, simultaneous with the third permethrin sampling was obtained from the Bay side from 14:00 to 15:30. No further samples were collected from the Atlantic side.

The third study (9/22/98) pre-application samples were collected from 16:20 to 17:30 in the bay and Atlantic and canal sites simultaneously, with permethrin application initiated at 20:00. Dibrom was not applied during this study due to evacuation of aircraft from the impending Hurricane Georges. The 12 hr post permethrin application sampling was performed from 8:40 to 10:00 on 9/22/98.

Results

First Application - June 16-18, 1998: Pesticide concentrations on the filter pads collected 3 to 6 hr after application recovered total (cis + trans) permethrin, ranging from 4.6 to 15.8 µg/m² on filters in the Florida Bay (down-wind) side of the application area. None was detected on the Atlantic side (Figure 1), which is consistent with the wind direction out of the ESE at 5 kts. Subsequent aerial application of naled the following morning resulted in no detectable naled or dichlorvos on filters from the Atlantic side, and one sample (B6) showing 1.6 µg/m² naled and a different one (B8) showing 0.16 µg/m² dichlorvos on Bayside filters.

No permethrin or naled was detected in the surface water from either side, with dichlorvos indicated at one Atlantic site (A3) at 1.3 µg/l. These results show that although some permethrin was deposited by drift onto the filter papers, none was detected in the surface water microlayer around the filters, indicating rapid dispersion-dissolution into the water, and/or rapid degradation by hydrolysis and photolysis. The low solubility of permethrin would cause it to adsorb to suspended particles, however, permethrin associated with suspended particles in the water column would have been recovered with the solvent extraction method.

Subsurface water samples (collected about 20 cm below the surface) showed no pesticide residues detected on the Bay side (except a trace of dichlorvos at one site) 2 to 4 hr after naled application. Naled was detected in only one sample (0.19 µg/l at A3) in the Atlantic side. Dichlorvos was observed in 50% of the Atlantic samples 2 to 4 hr after application, in concentrations ranging from 0.08 to 0.56 µg/l. Some dichlorvos was still detected in 3 of 9 sites during the 10 to 11 hr sampling period, ranging from 0.05 to 0.33 µg/l, suggesting that the breakdown product of naled (dichlorvos) was transported to these sites in the water column and that dichlorvos, not naled is the contaminant of concern for exposure. The most likely source would be tidal movement out of the canal systems of the residential area adjacent to where the pesticides were applied.

Quality control measures included determination of the lower limits of detection for each pesticide as well as recovery and analysis of standard pesticide mixtures in each matrix (Table 2). The average values for water quality parameters (Table 3) show that the Bay water had lower salinity than the Atlantic with higher dissolved oxygen (DO) and pH.
Table 2. Lower limits of detection and standard recoveries.

<table>
<thead>
<tr>
<th>Lower Limits of Detection</th>
<th>Naled / DDVP</th>
<th>Cis / Trans Permethrin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>0.05 µg/l</td>
<td>0.07 µg/l</td>
</tr>
<tr>
<td>Filter</td>
<td>0.1 µg/m²</td>
<td>0.2 µg/m²</td>
</tr>
<tr>
<td>% Standard Recovery</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Water</td>
<td>56±29</td>
<td>97±25</td>
</tr>
<tr>
<td>Filter</td>
<td>53±30</td>
<td>110±19</td>
</tr>
</tbody>
</table>

Table 3. Water Quality Values during each Sampling event: mean ± std dev.

<p>| | | | | | | |</p>
<table>
<thead>
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</thead>
<tbody>
<tr>
<td></td>
<td>June 16-18</td>
<td>July 28-29</td>
<td>Sept. 22-23</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Temp C°</td>
<td>DO mg/l</td>
<td>Sp Cond.</td>
<td>S%O</td>
<td>pH</td>
<td></td>
</tr>
<tr>
<td>Bay</td>
<td>32.3 ± 0.7</td>
<td>6.3 ± 1.1</td>
<td>44.9 ± 0.3</td>
<td>29.0 ± 0.2</td>
<td>8.2 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>Atlantic</td>
<td>32.0 ± 0.9</td>
<td>5.5 ± 1.6</td>
<td>56.3 ± 0.4</td>
<td>37.4 ± 0.3</td>
<td>7.9 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>Bay</td>
<td>32.5 ± 1.0</td>
<td>6.0 ± 1.1</td>
<td>54.6 ± 0.6</td>
<td>36.2 ± 0.4</td>
<td>8.0 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>Atlantic</td>
<td>31.6 ± 0.4</td>
<td>5.6 ± 1.0</td>
<td>53.0 ± 0.5</td>
<td>34.9 ± 0.3</td>
<td>8.1 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>Bay</td>
<td>31.0 ± 0.5</td>
<td>6.4 ± 1.0</td>
<td>56.6 ± 0.3</td>
<td>37.7 ± 0.3</td>
<td>8.0 ± 0.04</td>
<td></td>
</tr>
<tr>
<td>Atlantic</td>
<td>31.0 ± 0.5</td>
<td>5.6 ± 0.8</td>
<td>51.1 ± 0.4</td>
<td>33.7 ± 0.3</td>
<td>7.8 ± 0.1</td>
<td></td>
</tr>
</tbody>
</table>

Second Application - July 28-29, 1998: Results of the second application were similar to the first, with no permethrin found on filter pads on the Atlantic (up-wind) side and all filter samples in the Bay side exhibiting total (cis + trans) permethrin concentrations ranging from 1.1 to 17.1 µg/m³ at 3-6 hr after application (Figure 2). Dichlorvos was detected at site B4 (0.4 µg/m³). These results are consistent with the wind direction ESE at 12 kts, causing permethrin drift into Florida Bay.

Surface water samples showed no permethrin, dibrom or dichlorvos, indicating no accumulation at the water surface microlayer. Sub-surface water samples exhibited much lower amounts of dichlorvos than were observed from the first application, yet two out of nine sites did indicate trace amounts (0.07 and 0.09 µg/l), one site (B6) containing 0.07 µg/l of permethrin. It is possible that the low dichlorvos concentrations resulted from the delayed sampling time (3 hr late due to boat problems) and that higher wind velocity dispersed the aerially applied pesticides, reducing the amount deposited in any given area of the FKNMS and the canal water. Bay water had a slightly higher temperature than the Atlantic, and higher DO, yet the temperature and pH were about the same for Bay and Atlantic water (Table 3).

Third Application - September 22-23, 1998: The sampling protocol for the third application was changed to include samples of water from the residential canals opening to the Atlantic side of the study area. This sampling episode was scheduled to occur just as hurricane “Georges” was bearing down on the Keys, so it was terminated prematurely. Fortunately, we were able to collect filter and water samples from both the Bay and Atlantic sides and from the canals before and after the evening permethone application. The 12 hr post sampling collection also was obtained the following morning, prior to evacuating from the hurricane. The wind direction during the permethone application was opposite that of the two previous sampling episodes, this time from the north at 5-10 kts. The Bay water had higher S %O, DO and pH than Atlantic water, yet the temperatures were about the same (Table 3).
Pennethrin concentrations found on the filters, including the canal system, are given in Figure 3. These show peramanone collected on the filters in all except one of the Atlantic sites, with cis + trans pennethrin collected on filters in the canal ranging from 13.6 to 51.0 µg/m². Filters on the Bay side exhibited trace amounts of peramanone on 4 out of 10 sample sites.

Surface water was collected only from the canals. These results show peramanone in all of the surface water microlayer samples ranging from 5.1 to 9.4 µg/l at 2-4 hr after application.

Sub-surface water samples also were collected at each site 2-4 hr after application and again at 12 hr after. No pennethrin was detected in any of the subsurface samples, which is consistent with the previous two applications. Unfortunately, we were not able to monitor for naled or dichlorvos in the canal system. Naled was not applied during this study because the airplanes were evacuated due to the pending hurricane.

Discussion

The potential hazard associated with pesticide applications can be assessed by comparing the environmental concentrations with the toxicity to representative organisms that would be exposed. The hazard quotient, the ratio of the environmental concentration divided by the no observable adverse effects level (NOEL), with correction factors for each pesticide for each species considered, provides an estimate for a safe environmental concentration (Gargas et al., 1999). Although a NOEL was not available for representative marine invertebrates for all of the pesticides investigated in this study, a hazard can be assessed by comparison of literature toxicity values for representative invertebrates, with the pesticide concentrations observed in FKNMS water during this study (Table 4).

Table 4. Estimated environmental hazard for acute toxicity data for pennethrin and naled, and for acute and NOEL toxicity data for dichlorvos, to *mysidopsis bahia*, and *Homarus grammarus* (larvae) compared to environmental concentrations from this study.

<table>
<thead>
<tr>
<th>Pesticide</th>
<th>Acute 1 µg/l</th>
<th>NOEL 2 µg/l</th>
<th>EEC 3 µg/l</th>
<th>Hazard Estimate 4</th>
<th>Acute 4</th>
<th>NOEL</th>
</tr>
</thead>
<tbody>
<tr>
<td>pennethrin</td>
<td>0.02</td>
<td>–</td>
<td>6.9</td>
<td>345</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>naled</td>
<td>8.8</td>
<td>–</td>
<td>–</td>
<td>0.19</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>dichlorvos</td>
<td>18.8</td>
<td>0.63</td>
<td>0.56</td>
<td>0.03</td>
<td>0.9</td>
<td></td>
</tr>
</tbody>
</table>

1 96-hr LC-50, µg/l to *mysidopsis bahia*
   Pennethrin (Schimmel et al., 1983; Cox, 1998).
   Naled (Cox, 2002; Peckenpaugh, et al., 1997).
   Dichlorvos (Edwards, 1999)
2 NOEL for lobster (*Homarus grammarus*) larvae (McHenry et al., 1996).
3 Expected Environmental Concentration (observed in field samples).
4 Ratio of EEC to toxicity data.
**Permethrin** - Permethrin (both cis and trans) was found in most of the leeward filter samples, however, concentrations exceeding quantifiable amounts in water were found primarily in the canal surface water samples (range 5.1 to 9.4 \( \mu g/l \)). Permethrin concentrations in canal water greatly exceeded the 96 hr LC-50 values (0.02 \( \mu g/l \)) indicating a potential environmental hazard to aquatic invertebrates in the canals, depending on the duration of exposure. Potential sublethal effects of permethrin on reproduction and development of marine invertebrates also should be considered because the lower limit of detection (0.07 \( \mu g/l \)) was greater than the acute toxicity level (0.02 \( \mu g/l \)), precluding detection of lower toxic permethrin concentrations.

**Naled** - Naled was not detected on filter pads, nor in the surface water samples. The only instance of a detectable amount of naled (0.19 \( \mu g/l \)) was in one sub-surface water sample on the Atlantic side on 6/17/98. This concentration was less than the acute toxicity, indicating no acute toxicity effects. However, no NOEL for naled was found for marine invertebrates, so the sublethal effects could not be determined. The amount of naled (and dichlorvos) deposited into canals during aerial applications should be determined to address the potential hazard.

**Dichlorvos DDVP** - The primary degradation product of naled, dichlorvos, was not found in quantities exceeding the acute toxicity LC-50 of 18.8 \( \mu g/l \) for *mysidopsis bahia* nor the NOEL (0.63 \( \mu g/l \)) for *homarus gammaurus* larvae indicating no toxic impacts likely for those organisms. However, possible higher concentrations in canals raises concern for a potential hazard for sub-lethal effects to sensitive marine invertebrates in the canal systems. Although dichlorvos in the subsurface water could result from degradation of dibrom from aerial deposition, the most likely source would appear to be from tidal flushing of the pesticide residues from the residential canal systems, where the mosquito control pesticides were applied.

**Conclusions:**

These results show permethrin drift into the FKNMS from ground (truck) permanone applications by deposition on glass fiber filters held above the water surface, yet only trace amounts were detected in the surface or subsurface water samples in Florida Bay or in the Atlantic Ocean. Neither naled nor its degradation product, dichlorvos, were detected as drift on the filter pads, however sub lethal concentrations (0.5 to 0.56 \( \mu g/l \)) were found in some subsurface samples several hours after application, indicating aqueous transport from residential canals to the FKNMS. Concentrations of permethrin in canal water samples were of sufficient amounts (5.1 to 9.4 \( \mu g/l \)) to be considered an environmental hazard to sensitive marine invertebrates, depending on the duration and mode of exposure.

Deposition of high amounts of permethrin into the canals suggests the importance of more closely monitoring the amount of pesticides deposited into residential canals and partially enclosed embayments, where susceptible life stages of marine organisms could be exposed to toxic concentrations. Further studies are needed to assess the concentration, distribution and persistence of mosquito control pesticides in the canals and adjacent waters of the FKNMS, along with determining the NOEL for these pesticides on keystone marine organisms, to provide a hazard assessment.

**Acknowledgments**

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References


Figure 1. Concentrations of (cis + trans) permethrin deposited on filter pads in the Florida Bay and Atlantic sides of Key Largo, following ground (truck) application of Permonone, 9/22/98.

Figure 2. Concentrations of (cis + trans) permethrin deposited on filter pads in the Florida Bay and Atlantic sides of Key Largo, following ground (truck) application of Permonone, 6/17/98.

Figure 3. Concentrations of (cis + trans) permethrin deposited on filter pads in the Florida Bay and Atlantic sides of Key Largo, following ground (truck) application of Permonone, 7/28/98.
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