Identification of Human-Derived Attractants to *Simulium damnosum* Sensu Stricto in the Madi-Mid North Onchocerciasis Focus of Uganda

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Abstract. Human landing collections (HLCs) have been the standard method for the collection of black flies that serve as vectors for *Onchocerca volvulus*, the causative agent of onchocerciasis or river blindness. However, HLCs are inefficient and may expose collectors to vector-borne pathogens. The Esperanza window trap (EWT) has been shown to be a potential alternative to HLCs for the collection of *Simulium damnosum*, the principal vector of *O. volvulus* in Africa. To improve the performance of the EWT, sweat from individuals highly attractive or less attractive to *S. damnosum sensu stricto* was examined by gas chromatography and mass spectroscopy. Twelve compounds were identified which were solely present or present in increased amounts in the sweat of the highly attractive individuals. Two of these compounds (naphthalene and tert-hexadecyl mercaptan) were found to be attractive to *S. damnosum* sensu stricto in behavioral assays. Traps baited with these compounds outperformed those baited with the current standard bait of worn socks. Using these newly identified compounds as baits will make the EWT more efficient in collecting vector black flies and may enhance the potential utility of the EWT as a local vector control measure.

INTRODUCTION

Onchocerciasis is a parasitic infection caused by the filarial nematode *Onchocerca volvulus*. The parasite is transmitted through the bites of black flies of the genus *Simulium* that develop as larvae in fast running rivers and streams; hence, the disease is commonly called “river blindness.” Currently, 217.5 million people live in areas that are known to be endemic for onchocerciasis, and more than 99% of these cases are concentrated in 28 countries in sub-Saharan Africa.2 Mass drug administration (MDA) with ivermectin has been the main tool in the fight against onchocerciasis in both the Americas and Africa.3 The proof of principle by Diawara et al.4 that ivermectin MDA alone can interrupt the transmission of *O. volvulus* led to a paradigm shift from control to elimination in Africa.5

The major criterion contained in the 2016 WHO guidelines for the verification of onchocerciasis elimination is surveillance of the vector population.6 The standard method used in the collection of the vector (mainly *Simulium damnosum sensu lato*) is human landing collections (HLCs).7 However, HLCs are inefficient and may increase the risk exposure of the collectors to *O. volvulus*. The Esperanza window trap (EWT), developed in Mexico and subsequently evaluated in Burkina Faso and Uganda, has been found to be an effective replacement for HLCs for *S. damnosum*.8–12 In the initial trials of the EWT, sweat-impregnated articles of clothing (trousers and socks) were used in conjunction with CO2 as baits for the EWT.8,9 In an attempt to develop a more consistent and attractive bait, an analysis of compounds that were common to sweat collected from three individuals were identified.13 These compounds, as well as several others that had been shown to be attractive to other species of hematophagous insects, were evaluated for their ability to attract *S. damnosum sensu stricto* in Burkina Faso, using a combination of physiological (electroantennogram) and behavioral (Y-tube) assays.15 Five attractive compounds were identified in this manner. Plastic aroma beads infused with these compounds were found to be as effective, but not more effective than worn socks when used as baits in conjunction with CO2 on the EWT.13 This was perhaps not surprising, as this study focused on compounds that were found in all sweat samples analyzed. However, it has long been known that individuals differ greatly in their attractiveness to other hematophagous insects.14 By identifying differences in the composition of sweat collected from highly attractive versus less attractive individuals, it might be possible to develop a bait formulation that would be more highly attractive than one that incorporated compounds that were common to the sweat of all individuals. Here, we have tested this hypothesis, comparing the composition of sweat collected from individuals who were more attractive to *S. damnosum* s.s. with the sweat of individuals of average or less than average attractiveness. The compounds that were unique to or enhanced in the sweat of the highly attractive individuals were then evaluated for their attractiveness to *S. damnosum* s.s. using Y-tube and field-based assays.

MATERIALS AND METHODS

Study sites. Studies were carried out in Gonycogo and Laminatoo villages in the Koch Goma subcounty of the Nwoya district of Northern Uganda. Details regarding these communities are provided in an earlier publication.15 Both communities are located along the Ayago River. In previous studies, the sibling species of *S. damnosum* s.l. present in these communities was identified as *S. damnosum* s.s.11

Assaying the relative attractiveness of different individuals. A total of 16 individuals (eight each from Gonycogo and Laminatoo) were recruited to serve as vector collectors at the beginning of the study. The vector collectors were assigned to one of eight collection locations situated within a 500-m radius of the major *S. damnosum* s.s. breeding site located near each community. The collectors were trained in standard black fly collection procedures,16 and collections were begun on May 15, 2017. Collections began at sunrise (ca 7 AM) and continued until sunset (ca 6 PM). On the second day of collections, the collectors were rotated to a different...
collection site, and collections were conducted as before. Collections continued approximately twice per week (interrupted occasionally by inclement weather or important community events and holidays) through August 11, 2017, encompassing 24 total collection days. All collectors thus spent 3 days at each collection site, to control for differences in fly density among the collection sites.

The total number of flies collected by each individual in each community was normalized by dividing the total number of flies collected by each individual by the total number of flies collected over the course of the study. This was performed to adjust for the difference in the overall fly density between the two communities.

Collection of sweat samples. Sweat samples were collected from the two most attractive individuals, the two least attractive individuals, and two individuals exhibiting average attractiveness. To collect the sweat samples, each of the individuals enrolled in the study were provided with a new cotton tee shirt, a new pair of cotton underpants, and a new pair of white cotton socks. The shirts, underpants, and socks were laundered three times in plain water before use. Each collector wore the clothing continuously for 72 hours. The worn garments were then collected, placed in ziplock polyethylene bags (one bag per individual, containing the collected shirt, underpants, and socks), and transported to the United States. The clothing samples were stored at −20°C in the sealed bags until analyzed.

Analysis of sweat samples. Sweat samples were analyzed by coupled gas chromatography–mass spectroscopy (GC-MS) as previously described. In brief, a Solid Phase Microextraction - Overcoated (SPME-OC) fiber assembly 65 μm PDMS/DVB (Cat #57439-U, Sigma-Aldrich, St. Louis, MO) was used to detect the compounds in the worn clothes. A fiber was put into the bag containing the dirty clothes for 24 hours at room temperature to absorb volatile compounds. The SPME-OC fibers were analyzed on an Agilent 7200 GC-QToF (Agilent, Inc., Santa Clara, CA). The sweat components were thermally desorbed in the GC inlet at 320°C, and the individual components were resolved on a HP-5ms capillary column (Agilent, Inc., Santa Clara, CA). The sweat components were ionized using methane as the reagent gas and both positive and negative modes (40 eV, compound dependent) to obtain optimum signal-to-noise ratio. The total ion chromatographs were de-convoluted using the s deconvolution algorithm, and the resulting fragmentation patterns were imported into Mass Profiler Professional for statistical analysis. Datasets, prepared for each time point and from both SPME fiber materials, were combined to create an entity list for each subject containing all compounds present in their sweat. Spectra of entities identified as present in the sweat of both highly attractive individuals but absent or in very reduced amounts in the sample from the low and average attractive individuals were compared with the fragmentation patterns contained in the NIST/Wiley mass spectra library for tentative identification (Wiley Interscience, Hoboken, NJ), as previously described.

Y-tube assays. Host-seeking S. damnosum s.s. females were collected by human landing collectors in Gonycoago and Laminatoo. The flies were captured alive into glass tubes which were then wrapped in a moist towel soaked in water and put in a cool box for transportation to the laboratory. Flies were confirmed as S. damnosum s.s. using morphological characters before being used in the Y-tube assays. All flies were used in the Y-tube assays within 24 hours of capture.

Y-tube assays were conducted in a darkened climate-controlled room located at the Molecular Laboratory of Gulu University, essentially as previously described. Air flowing through the chamber was passed through a charcoal filter to remove volatiles in the ambient air. Twenty flies were relocated into the closed release chamber at the trunk of the Y-tube and allowed to acclimate for 10 minutes. During this time, samples were prepared on filter papers which had previously been rinsed in hexane and dried. A total of 20 μL of the sample compound dissolved at various concentrations in hexane was pipetted onto the filter paper, and 20 μL of hexane was pipetted onto a second filter paper, which served as the control. After the acclimation period, the filter papers were loaded into separate end chambers, and the flies were released and allowed to roam freely in the apparatus for 20 minutes. All chambers were then closed, and the number of flies in either arm of the Y-tube apparatus, those remaining in the trunk of the apparatus, and those still in the release chamber, were counted. The compounds were initially tested at 1:100 and 1:1,000 (w/v) dilutions. Those that appeared to exhibit some activity at these dilutions were then tested over a wider range of concentrations. Each concentration of each compound was tested in six independent trials. The arms containing the control and compound samples were interchanged on every trial to eliminate potential position effects. The Y-tube was thoroughly cleaned with ethanol between runs, and control runs were carried out between each experimental trial to ensure that no residual attractant remained in the apparatus. Data were analyzed using a likelihood ratio test (α = 0.05), as previously described.

Field evaluation of compounds as baits. Naphthalene and tert-hexadecyl mercaptan were dissolved in mineral oil at a 1/100 concentration (w/v). This solution was then used to prepare two 10-fold dilutions in mineral oil. Aliquots of 25 g of plastic aroma beads (Bitter Creek Candle Supply [www. candlesupply.com]) were placed into a series of 50 mL conical centrifuge tubes, and the tubes were filled with the solutions of the compounds. The tubes were placed on a tube rotator, and the beads were allowed to absorb the solutions for 2 days. The excess solution was decanted from the beads, and the tubes were sealed and transported to the field sites.

Four EWTs containing a thin black stripe (the optimal design for the collection of S. damnosum s.s. in Uganda) were set up at locations near the larval breeding sites at Gonycoago and Laminatoo. The trap sites were chosen so that they were within 500 m of the breeding site but separated from each other by at least 50 m. All traps were baited with 5-L containers of yeast and sugar dissolved in water to create CO2, as previously described. Baits containing aroma beads impregnated with different concentrations of naphthalene and tert-hexadecyl mercaptan were prepared by placing approximately 20 g of the beads into a woman’s nylon stocking and the top of the stocking tied off. The baits (and worn socks as the control) were attached to the top of the trap by tying them onto the frame with a string.

The traps were allowed to collect flies from sunrise until sunset. The insects collected by the trap were then removed.
and the S. damnosum s.s. identified by morphological keys. On the second day of the trial, baits were rotated among the traps to control for position effects. Each trial was continued for a total of 8 days so that each bait was evaluated twice at each trap location. The yeast solution was replenished daily, and the baits were replaced every other day. The number of flies collected each day by each bait was normalized to the total number of flies collected by all traps at the community during the day, controlling for differences in the fly densities from day to day (resulting from daily changes in weather conditions) and for the differences in fly densities in the two communities.

The proportional data were analyzed for normality using the Shapiro–Wilk test. Where normality assumptions were not violated, the data were analyzed using an Analysis of Variance (ANOVA) with a Tukey post hoc. In cases where normality assumptions were considered violated, the data were analyzed using the Kruskal–Wallis test with a Dwass, Steel, Critchlow-Fligner (DSCF) post hoc. Significance was tested at an $\alpha = 0.05$ level.

Ethical statement. The experiments included in this study were reviewed and cleared by the Institutional Review Committee of Vector Control Division (VCD-REC/071), and final approval was granted by the Uganda National Council for Science and Technology (HS 2154). Approval was also obtained from the University of South Florida Institutional Review Board for Human Subject Research (protocol number CRS_Pro00015108). All individuals involved in the study provided written informed consent to participate.

RESULTS

To identify sweat volatiles specific to highly attractive individuals, it was first necessary to identify individuals who were more or less attractive to S. damnosum s.s. To accomplish this, human landing collectors were recruited in the two study communities of Gonyego and Laminato. Eight collectors were recruited in each community. The collectors were then assigned to eight separate collection sites located near the principal breeding site at each community, as described in Materials and Methods. Fly collections were carried out roughly twice per week, and the collectors were rotated daily among the collection sites. The collections continued for 24 days, so that each collector collected at each site three times, to control for differences in fly density among the different sites. The proportion of all the flies collected in the community by each individual was then calculated. The proportion of flies collected by each individual appeared to follow a normal distribution, ranging from 4.1% of the total community collection to 23.0% of the total community collection (Figure 1). Two individuals collected more than the mean proportion of each community collection plus one SD (16.9%), whereas two individuals collected less than the mean community proportion minus one SD (8.1%). The two most attractive individuals, the two least attractive individuals, and the two individuals with an average level of attractiveness were chosen for further analysis. These six individuals were provided with new cotton shirts, underpants, and socks and asked to wear them for 72 hours without removing them, while going about their normal activities. The clothing samples were then collected and the volatiles analyzed by GC-MS, as described in Materials and Methods. The individual traces were compared with one another to identify compounds that were unique to or present in greater amounts in the samples from the two highly attractive individuals but were absent or present in lesser amounts in the remaining samples (e.g., Figure 2). A total of 12 compounds were identified by this process.
Compounds unique to or in enhanced amounts in sweat from highly attractive individuals

<table>
<thead>
<tr>
<th>Compound</th>
<th>Commercially available</th>
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<tr>
<td>1-Octanol</td>
<td>Yes</td>
</tr>
<tr>
<td>1-Pentadecanol</td>
<td>Yes</td>
</tr>
<tr>
<td>2-Butyl-1-octanol</td>
<td>Yes</td>
</tr>
<tr>
<td>2-Heptanone</td>
<td>Yes</td>
</tr>
<tr>
<td>6,10-Dimethyl-5,9-undecadien-2-one</td>
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</tr>
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<td>Naphthalene</td>
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</tr>
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<td>tert-Hexadecyl mercaptan</td>
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</tr>
<tr>
<td>1-Heptatriocotenol</td>
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</tr>
<tr>
<td>2-Methyl-1-dodecanol</td>
<td>No</td>
</tr>
<tr>
<td>4-Ethyloctane</td>
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</tr>
<tr>
<td>4,8-Dimethyl undecane</td>
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</tr>
<tr>
<td>Pentadecanol</td>
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</table>
reported to occur in humans,22 cattle,23 and dogs,24 and it could be involved in location of hosts for blood meals. Our results also support earlier laboratory observations by Verocai et al.25 that naphthalene was attractive to the North American black fly species *Simulium vittatum* in electroantennogram (EAG) and Y-tube assays. Both *S. damnosum* s.s. and *S. vittatum* are zoophilic species that may attack a variety of hosts. However, naphthalene was previously tested against five species of higher Diptera of veterinary importance and found to be largely repellent.23

The results described here provide useful information necessary to design a more efficient baited trap that could be used to attract large numbers of *S. damnosum* s.l. under field conditions. Onchocerciasis elimination programs in Africa have experienced challenges,26 and alternative strategies to accelerate elimination remain a priority.27 Studies conducted both in Mexico28 and Uganda16 have demonstrated that EWTs, when deployed in areas where people congregate, can significantly reduce biting by the Simulium vectors of *O. volvulus*, providing a supplemental strategy to MDA to accelerate reaching the goal of *O. volvulus* elimination. The EWT is inexpensive and simple to construct from materials that are commonly available in developing countries.16 It is quite possible that aroma bead baits prepared with tert-hexadecyl mercaptan and naphthalene may improve the effectiveness of EWTs, enhancing their potential as a local vector.
control measure. Aroma beads impregnated with these compounds are simple to prepare, and the ingredients are inexpensive. Although we have not yet explored the effective life of the baits, air fresheners prepared in the same way continue to be effective for weeks or months. An aroma bead bait for an EVT will cost $1.87 (for 20 g of beads impregnated with naphthalene) and $2.99 (for 20 g of beads impregnated with tert-hexadecyl mercaptan). Once the beads are prepared, they are stable when kept in a sealed container, are safe to handle, and can be transported at ambient temperatures; these characteristics would make their promotion and marketing much easier. In addition, traps baited with attractive lures can easily find an expanded market, being used for surveillance and evaluation of ivermectin MDA, the major criterion in the recent WHO verification guidelines. This expanded market may increase the availability and decrease the unit cost, both of the baits and the trap platforms.

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