

Research Article

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Effects of Malaise trap spacing on species richness and composition of terrestrial arthropod bulk samples

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Abstract

The Malaise trap is a popular device for assessing diverse terrestrial arthropod communities because it collects large samples with modest effort. A number of factors influence its collection efficiency, placement being one of them. For instance, when designing larger biotic surveys using arrays of Malaise traps we need to know the optimal distance between individual traps that maximises observable species richness and community composition. We examined the influence of spacing between Malaise traps by meta-barcoding samples from two field experiments at a site in Waterloo, Ontario, Canada. For one experiment, we used two trap pairs deployed at weekly increasing distances (3 m increments from 3 to 30 m). The second experiment involved a total of 10 traps set up in a row at 3 m distance intervals for three consecutive weeks.

Results show that community similarity of samples decreases over distance between traps. The amount of species shared between trap pairs drops considerably at about 18 m trap-to-trap distance. This change can be observed across all major taxonomic groups and for two different habitat types (grassland and forest). Large numbers of OTUs found only once within samples cause rather large dissimilarity between distance pairs even at close proximity. This could be caused by a large number of transient species from adjacent habitats which arrive at the trap through passive transport, as well as capture of rare taxa, which end up in different traps by chance.

Key Words

Biodiversity, biomonitoring, experimental design, insects, metabarcoding

"During my extensive travels I have repeatedly found that insects happened to enter my tent, and that they always accumulated at the ceiling-corners in vain efforts to escape at that place without paying any attention to the open tent-door. On one occasion one of the upper tent-corners happened to have a small hole torn in the fabric, and through this hole all the insects pressed their way and escaped. Later on the idea occurred to me, that, if insects could enter a tent and not find their way out, and always persistently tried to reach the ceiling, a trap, made as invisible as possible and put up at a place where insects are wont to patrol back and forth, might catch them much better than any tent and perhaps better than a man with a net..."

Rene Malaise 1937

Introduction

The inclusion of terrestrial invertebrates in biodiversity inventories and surveys has increased substantially over the past years (Dopheide et al. 2019; Drake et al. 2007) but sampling efficiency remains a key consideration when designing larger censuses (Telfer et al. 2015; Timms et al. 2012). Although no single sampling can be used to survey all taxa at a given site, the Malaise trap (Malaise 1937) is often chosen for the assessment of terrestrial arthropod

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communities (Karlsson et al. 2005) because it collects large and diverse samples with fairly little effort. Malaise's invention is a tent-like flight-interception trap made from fine mesh netting with a central screen suspended below a sloping ridge-roof that leads to a collecting bottle at the upper end. Flying insects that hit the screen, subsequently fly or walk along this roof to the bottle which is usually filled with >90% ethanol as preservative. Many traps have different colors in the upper and lower parts, the upper one being white because most insects show positive phototropism. This feature helps directing insects toward the bottle. Traps are usually deployed in such a way that the central mesh intercepts the flying path of insects. There are a number of different designs available although the most commonly used traps are so-called Townes-Style traps (Townes 1972) and derived versions of it (e.g. ez-Malaise traps). The trap is particularly well-suited for inventory because it catches a wide variety of flying insects and some ground active insects that climb up the trap fabric. Malaise trapping is easy, requires modest labour and as such represents one of the best mass-collecting methods available for terrestrial invertebrates (deWaard et al. 2018).

Initially Malaise traps were considered of limited use in conservation evaluation and bio-surveillance because of the huge size of their catch (Drake et al. 2007) which made it difficult to characterize the community using traditional morphology-based methods (Cook et al. 2010). Consequently, larger surveys used total biomass rather than detailed identification of specimens. In fact, one of the recent reports on the dramatic decline of terrestrial arthropod abundance was the result of a long-term study using Malaise traps and catch biomass (Hallmann et al. 2017). The recent advent of DNA barcoding (Hebert et al. 2003) and metabarcoding (Taberlet et al. 2012) opened the door to more comprehensive estimates of species richness and community composition (Braukmann et al. 2019; Yu et al. 2012; Steinke et al. 2021) and Malaise traps are poised to become a ubiquitous tool for biodiversity surveys (Geiger et al. 2016). It seems that large-scale or global high-resolution monitoring networks are within our reach (Hobern and Hebert 2019) but there are still a number of challenges in relation to the small-scale variability of many terrestrial habitats.

There are various factors that influence the efficiency of Malaise traps. Temperature, precipitation, and wind are considered important as largest catches generally occur on hot, dry, and still days (Matthews and Matthews 1971). It has also been noted (Townes 1962) that insects often fly closer to the ground in spring because of the warmer air there, thereby increasing the number of individuals caught during this season. As a Malaise trap samples only those arthropods that happen to fly through a relatively small area, trap orientation and placement become important considerations. Height of surrounding vegetation and location in shade or sun can alter trap performance and efficiency (Matthews and Matthews 1971; Ssymank et al. 2018). Another relevant but not systematically studied variable is the distance between traps in a sampling area. This is a particular important consideration when

designing larger biotic surveys using arrays of traps. For instance, it is not known how many traps at what distance are needed to maximize observable species richness and community composition for a given location.

The main objective of this study was to examine the effects of spacing between traps on species richness and composition of Malaise trap samples. Bulk samples from two field experiments at a site in Waterloo, Ontario, Canada were assessed using metabarcoding to determine if (1) there is a critical distance between traps at which species overlap drops significantly and if (2) structural composition of habitats has an influence on such a distance.

Materials and methods

Site and sampling

Arthropod bulk samples were collected using ez-Malaise traps (Bugdorm, Taiwan). Traps for the first experiments (Fig. 1a) were deployed in both a grassland (Suppl. material 1: Fig. S1) and a forested pond area near Waterloo, Ontario, Canada. Traps for experiment 2 (Fig. 1b, c; Suppl. material 1: Fig. S1) were positioned only in the grassland area. For the first experiment we used two trap pairs that were deployed next to each other (3 m distance between both collecting bottles) respectively. Each week trap spacing for each pair was increased by three meters to a maximum distance of 30 m (Fig. 1a). Samples were collected every week before moving one trap further away (July-September 2019). The second experiment involved a total of 10 traps set up at 3 m distance intervals for three consecutive weeks in September 2019 (Fig. 1b, c; Suppl. material 1: Fig. S1) at the same grassland site that was used for experiment 1. Samples were also collected each week and sample bottles were stored at -20 °C for further analysis. Each time trap heads (collecting area with bottles) were cleaned using bleach and ethanol to minimize cross-contamination between sampling events.

Molecular analysis

All samples were dried at room temperature for three days in a disposable grinding chamber. Each sample was ground to fine powder using an IKA Tube Mill control (IKA, Breisgau, Germany) at 25,000 rpm for 2×3 min. DNA was extracted from approximately 20 mg of ground tissue using the DNeasy Blood & Tissue kit (Qiagen, Venlo, Netherlands) in line with the manufacturer's protocols.

Metabarcoding was carried out using a two-step fusion primer PCR protocol (Elbrecht and Steinke 2019). During the first PCR step, a 421 bp region of the Cytochrome c oxidase subunit I (COI) was amplified using the BF2 + BR2 primer set (Elbrecht and Leese 2017). PCR reactions were carried out in a 25 μ L reaction volume, with 0.5 μ L DNA, 0.2 μ M of each primer, 12.5 μ L PCR Multiplex Plus buffer (Qiagen, Hilden, Germany). The PCR was carried out in a Veriti thermocycler (Thermo Fisher



Figure 1. Sampling design (trap distances over time) for experiment 1 (a) and experiment 2 (b).

Scientific, MA, USA) using the following cycling conditions: initial denaturation at 95 °C for 5 min; 25 cycles of: 30 sec at 95 °C, 30 sec at 50 °C and 50 sec at 72 °C; and a final extension of 5 min at 72 °C. PCR success was checked on a 1% agarose gel. One µL of PCR product was used as template for the second PCR, where Illumina sequencing adapters were added using individually tagged fusion primers (Elbrecht and Steinke 2019). Tagging combinations are available in Suppl. material 3: Table S1. We mainly used the same thermocycler conditions as in the first PCR but the reaction volume was increased to 35 µL by adding water, the cycle number reduced to 20 and extension time increased to 2 minutes per cycle. PCR success was again checked on a 1% Agarose gel. PCR products were purified and normalized using SequalPrep Normalization Plates (Thermo Fisher Scientific, MA, USA, Harris et al. 2010) according to manufacturer protocols. Ten µL of each normalized sample were pooled, and the final library cleaned using left sided size selection with 0.76x SPRIselect (Beckman Coulter, CA, USA). Sequencing was carried out by the Advances Analysis Facility at the University of Guelph using the 600 cycle Illumina MiSeq Reagent Kit v3 and 5% PhiX spike in. The read length of read one was increased to 316 bp, while keeping read 2 to 300 bp. As we only used inline barcodes for sample tagging, both Illumina indexing read steps were skipped.

Data processing

Initial quality control of raw sequence data was done using FastQC v0.11.8. Subsequently, sequence data were processed using the JAMP pipeline v0.69 (github.com/ VascoElbrecht/JAMP) starting with demultiplexing, followed by paired-end merging using Usearch v11.0.667 with fastq pctid=75 (Edgar 2010). Primer sequences were trimmed from each sequence using Cutadapt v1.18 with default settings (Martin 2011), retaining only sequences where primers were successfully trimmed at both ends. Cutadapt was also used to remove sequences shorter than 411 bp and longer than 431 bp. Sequences with poor quality were removed using an expected error value of 1 (Edgar and Flyvbjerg 2015) as implemented in Usearch. Filtered reads of each sample were dereplicated and singletons removed, before pooling all reads for OTU clustering with Usearch cluster otus at a 97% similarity threshold. Previously removed dereplicated and singleton reads were mapped back against generated OTUs using Usearch usearch global, to generate the final OTU table. The maximum read count for each OTU across all 12 negative controls was multiplied by two, and subtracted from corresponding OTU read counts in all samples (Elbrecht and Steinke 2019). Taxonomy was assigned by using OTUs as queries for the BOLD reference database (www.boldsystems.org Ratnasingham and

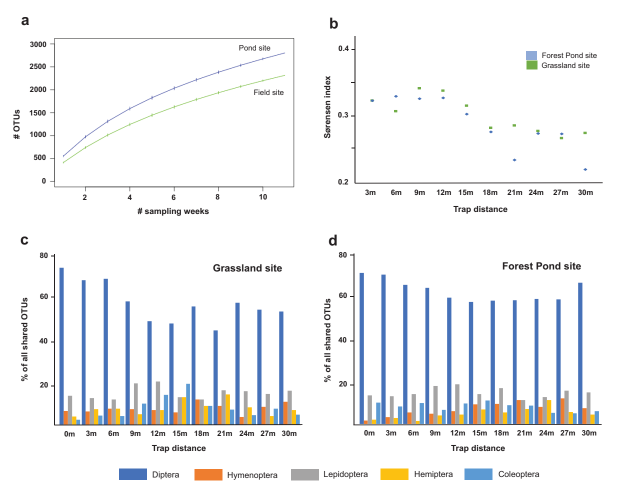


Figure 2. Results for experiment 1. (a) OTU accumulation curves for both sites by sample, (b) Dot plot of Sørensen's similarity coefficient between samples of distance pairs for both sites. Percentage of shared OTUs per trap distance pair for the top five arthropod orders (representing 90% and 96% of all OTUs, respectively) at grassland (c) and forest pond (d) site.

Hebert 2007) utilizing the JAMP Bold_web_hack script with default settings. Only OTUs with a minimum match of 98% were retained for further analysis. For most analysis carried out in R v3.5.1, relative read counts were used, and only reads above 0.01% abundance per sample were considered.

Statistical analysis

OTU tables (Suppl. material 4: Table S2) were used to calculate both the number of OTUs shared as well as the Sørensen similarity coefficient between all trap pairs for both experiments. For experiment 2 we used all possible pairings for a given distance. In order to determine OTU sampling effort for experiment 1 we calculated accumulation curves using the function specaccum and applied the function *specpool* to extrapolate species richness for each week as well as to calculate the Chao 1 (Magurran 2003) species estimate for the total number of OTUs possible with complete sampling. Both functions are part of the vegan package (Oksanen et al. 2018). Pairwise OTU overlap among trap distance pairs was evaluated using the nonparametric multiple comparison function implemented in the R package dunn.test 1.2.4 (Dinno 2016) which is equivalent to the Kruskall-Wallis test.

Results

We were able to extract high quality DNA from all samples, and obtained strong bands for all 70 samples after the second PCR step. Illumina sequencing generated 13,910,614 reads (partial run shared with other projects), with the raw data being available on NCBI SRA with the accession number SRP200574. About 27% of the reads were filtered during data processing, leaving an average of about 137,181 sequences per sample. In total, 10,151,381 post-filtering reads could be used for clustering with Usearch.

Our analysis shows a total of 2,315 OTUs for the grassland site and 2,804 OTUs for the forest pond site in experiment 1 (Fig. 2a). On average, about half of those (49%) were only detected once over the entire experiment. The Chao 1 (Magurran 2003) species estimates calculated with *specpool* for the total number of OTUs possible with complete sampling were 3,847+119 and 4,550+129 respectively. The Sørensen index between samples of distance pairs was generally low (<0.34) for both sites and all distances, however similarity at both sites decreased significantly (Kruskal-Wallis and Dunn's posthoc p < 0.0001) at 18 m distance (Fig. 2b). The average similarity for the trap that remained at the same position was 0.32.

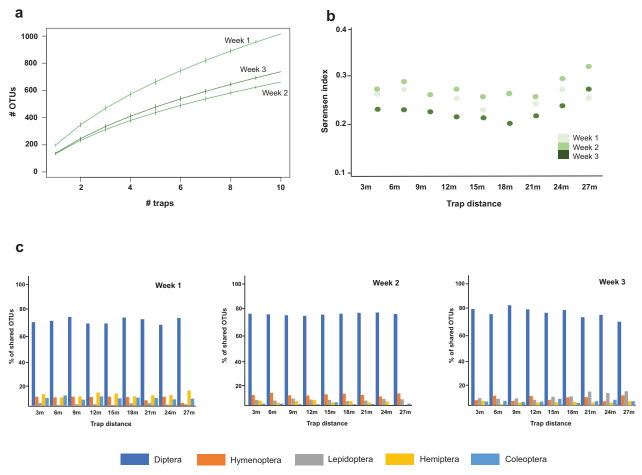


Figure 3. Results for experiment 2. (a) OTU accumulation curves for each sampling week by trap (b) Dot plot of Sørensen's similarity coefficient between samples of all possible distance pairs for all weeks, (c) percentage of shared OTUs by trap distance for the top five arthropod orders (representing 90% of all OTUs) for each week.

Overall, the proportion of OTUs shared between trap pairs and across all distances ranged from 26–27%.

The total OTU count for the grassland site comprised 21 orders with six orders (Coleoptera, Diptera, Hemiptera, Hymenoptera, Lepidoptera, Orthoptera) representing 97% of all OTUs (Suppl. material 5: Table S3). We found 20 orders at the forest pond site with five orders (Coleoptera, Diptera, Hemiptera, Hymenoptera, Lepidoptera) representing 96% of all specimens (Suppl. material 5: Table S3). We observed a distinct drop in the number of OTUs shared between traps at a distance between 15 m and 18 m for both the grassland and the forest pond site (Suppl. material 4: Table S2) but no such changes in the proportion of shared OTUs per order (Fig. 2c, d). Especially the proportion of Diptera varied across the entire experimental run likely as the result of phenological variation.

For experiment 2 we found totals of 1,017 for week 1, 662 for week 2, and 738 OTUs for week 3 (Fig. 3a). The total number of OTUs found over the three weeks was 1,610 (Suppl. material 1: Fig. S1). Chao estimates for the expected amount of total OTUs were 2,007+117, 1,211+80, 1,479+102 for weeks one to three, respectively. Sørensen indices for experiment 2 were generally lower than in experiment 1 with weekly averages ranging from 0.23 (week 3) to 0.27 (week 2) (Fig. 3b). The values remained fairly stable with increasing

distance and in two cases (week 2 and 3) actually increased at the larger distances. In addition, more than half of the OTUs obtained in three weeks of experiment 2 (811) were only detected in a single trap following a common hollow curve species abundance pattern (Suppl. material 2: Fig. S2).

OTUs found during experiment 2 comprised 15–18 orders with the five orders Coleoptera, Diptera, Hemiptera, Hymenoptera, and Lepidoptera representing between 85–95% of all specimens (Suppl. material 5: Table S3). In contrast to experiment 1, we observed a constant decline in the number of species shared between traps with increased distance between them (Suppl. material 4: Table S2) but no variation in relative proportions of orders in shared OTUs.

Discussion

Malaise traps as sampling method for terrestrial arthropod communities represent a rather efficient and economical means for obtaining comprehensive samples with minimal effort (Karlsson et al. 2005). They can be operated continuously in any weather with only occasional attendance and deliver large sample sizes. In conjunction with modern DNA-based methods to assign taxonomy (e.g. metabarcoding) they probably represent the best mass-collecting method available for terrestrial arthropods and are well-suited for large scale biotic surveys using arrays of traps (Yu et al. 2012, deWaard et al. 2018; Steinke et al. 2021). However, so far it was not known how many traps at what distance are needed to maximize observable species richness and community composition for a given location (Noyes 1989). Various strategies have been applied but trap spacing varied considerably (Fraser et al. 2008; Santos et al. 2014). Results from our first experiment (Figs 1a, 2) suggest that deploying traps at about 18 m distance from each other can increase overall species richness and reduce overlap between traps. This is true for all major taxonomic groups collected although the percentage of orders varied independent from distance which is likely a reflection of temporal variation over the 10 weeks of the experiment (Fig. 2c, d). Similarity at the 18 m distance also dropped in comparison to the respective traps that stood in the same place for the entire experiment which indicates that the effect of phenological variation was small. Interestingly, overall habitat type seems to have no effect on the distance observed as both the grassland and the forest pond sites exhibit the same cut-off value. On the other hand, experiment 2 does not show a drop in overlap between adjacent traps (Fig. 3b). For each week we were able to observe a more gradual decline followed by an increase of similarity at large distances in two weeks. The proportion of orders within the group of shared OTUs remained fairly constant and showed a typical distribution for Malaise trap samples (Fig. 3c). This could be the result of the experimental set up we chose. A row of ten traps represents a continuous structure along which some animals can move before being caught. Along the row of traps the amount of observed OTUs varies which is likely the result of variation in microhabitat structure (Suppl. material 1: Fig. S1). The grassland chosen for the experiment was not entirely uniform and characterized by sporadic patches of golden rod (Solidago canadensis) which were in flower over most of the three weeks the experiment lasted. In the weeks prior this site was used for experiment 1 and no such patterns emerged.

The small similarity values observed in both our experiments (Figs 2b, 3b) are influenced by a large proportion of singleton OTUs. We are confident that these are mostly true specimens rather than OTUs derived from sequencing or PCR errors, because we removed OTUs that did not match the BOLD database to at least 98%. Large numbers of OTUs found only once over a sampling period or between traps have been observed several times in other studies using Malaise traps (e.g. Geiger et al. 2016, deWaard et al. 2018; Steinke et al. 2021). This phenomenon has been discussed as an indicator for the presence of transient species (D'Souza and Hebert 2018; Steinke et al. 2021). Transient species have been defined as species that show up only occasionally as a result of dispersal from adjacent habitat (Snell Taylor et al. 2018). Specifically, many smaller species caught are not necessarily living in the sampled habitat but are rather passively transported there e.g. by wind. Additionally, sampling might be stochastic when it comes to rare or low abundant taxa.

Malaise trapping with only a few traps at a single site over a short timescale always provides an incomplete species list. That is no different for our study which suggests that additional trapping efforts by increasing the number of traps or by enlarging the trapping surface (e.g. Gressitt and Gressitt 1962) are needed to approach asymptotic species richness at both experimental sites. The trap results for experiment 1 suggest that it needs a 1.6-fold increase of the full sampling effort for a complete inventory (over the entire 10 weeks of the experiment) based on Chao-1. For experiment 2 sampling efforts would need to be doubled to obtain maximum species richness for the site in any given week. This could perhaps be accomplished by deploying a second row of ten traps at 18 m distance following the findings of experiment 1. The alternative would be to increase the sampling duration (Fraser et al. 2008) or the sampling surface of the traps (Gressitt and Gressitt 1962).

In conclusion, our results suggest the following recommendations for sampling and monitoring terrestrial invertebrate communities with Malaise traps: (a) within a temperate and uniform habitat a number of traps equally spaced at >18 m will likely sample more of the local diversity while at the same time reduce the extend of repetitive sampling, (b) longer trapping duration can help to reach asymptotic species richness and lead to more complete species lists, and (c) future work should include research on the origin and the role of singletons. Are they in fact transient species passively dispersed towards the trap or low abundant resident core species that are not efficiently detected?

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Author contributions

DS developed the concept, DS and VE designed the experiments and carried out the sampling, VE and LM did the laboratory work, DS and VE analysed the data, DS wrote the paper, VE, AW, and TB revised the paper and provided input throughout the study.

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Supplementary material 1

Figure S1

Author: Dirk Steinke, Thomas WA Braukmann, Laura Manerus, Allan Woodhouse, Vasco Elbrecht

Data type: histogram

- Explanation note: Histogram of OTU richness per trap (corresponding trap shown in photograph below) per week of experiment 2.
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Link: https://doi.org/10.3897/mbmg.5.59201.suppl1

Supplementary material 2

Figure S2

- Author: Dirk Steinke, Thomas WA Braukmann, Laura Manerus, Allan Woodhouse, Vasco Elbrecht
- Data type: histogram
- Explanation note: Histogram showing Number of OTU occurrences in traps of experiment 2.
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- Link: https://doi.org/10.3897/mbmg.5.59201.suppl2

Supplementary material 3

Table S1

- Author: Dirk Steinke, Thomas WA Braukmann, Laura Manerus, Allan Woodhouse, Vasco Elbrecht
- Data type: lab protocol information
- Explanation note: Tagging layout.
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- Link: https://doi.org/10.3897/mbmg.5.59201.suppl3

Supplementary material 4

Table S2

- Author: Dirk Steinke, Thomas WA Braukmann, Laura Manerus, Allan Woodhouse, Vasco Elbrecht
- Data type: OTU list
- Explanation note: OTU table (FC, FM, PC, PM Experiment 1; A, B, F Experiment 2; C Controls).
- Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.

Link: https://doi.org/10.3897/mbmg.5.59201.suppl4

Supplementary material 5

Table S3

- Author: Dirk Steinke, Thomas WA Braukmann, Laura Manerus, Allan Woodhouse, Vasco Elbrecht
- Data type: taxonomic
- Explanation note: Taxonomic breakdown by order for both experiments and sites.
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- Link: https://doi.org/10.3897/mbmg.5.59201.suppl5