Wet grinding of invertebrate bulk samples – a scalable and cost-efficient protocol for metabarcoding and metagenomics

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Abstract

Most metabarcoding protocols for invertebrate bulk samples start with sample homogenisation, followed by DNA extraction, amplification of a specific marker region, and sequencing. Many of the above-mentioned laboratory steps have been verified thoroughly and best practice strategies exist, yet, no clear recommendation for the basis of almost all metabarcoding studies exists: the homogenisation of samples itself. Two different categories of devices are typically used for homogenisation: bead mills or blenders. Both have upsides and downsides. Bead mills rely on single-use plastics and therefore produce a lot of waste and are expensive. In addition to that, processing times can go up to 30 minutes making them unsuitable for large-scale studies. Blenders can handle larger sample volumes in a shorter time, and be cleaned – yet suffer from an increased risk of cross-contamination. We aimed to develop a fast, robust, cheap, and reliable sample homogenisation protocol that overcomes limitations of both approaches, i.e. does not produce difficult to discard waste and avoid single-use plastics while reducing overall costs. We tested the performance of the new protocol using six size-sorted Malaise trap samples and six unsorted stream macroinvertebrate kick-net samples. We used 14 replicates per sample and included many negative controls at different steps of the protocol to quantify the impacts of i) insufficient homogenisation and ii) cross-contamination. Our results show that 3-min homogenisation is sufficient to recover about 80% of OTUs per sample in each replicate and that a non-hazardous DIY cleaning solution provides an effective and efficient way of cleaning. The improvements of the protocol in terms of speed, ease of handling, an overall reduction of costs as well as the documented reliability and robustness make it an important candidate for sample homogenisation after sampling in particular for large-scale and regulatory metabarcoding but also metagenomics biodiversity assessments and monitoring.

Key Words

bioassessment, biodiversity soup, bulk sample, community metabarcoding, DNA isolation, LTER

Introduction

DNA metabarcoding is an efficient tool to characterize invertebrate species composition in environmental samples. Starting material can be very different and include flying insect samples, soil samples, benthos, or plankton samples (Yu et al. 2012; Hajibabaei et al. 2019b; Meyer et al. 2020). Most metabarcoding protocols start with a complete sample homogenisation, although alternative approaches are also explored (Zizka et al. 2018; Marquina et al. 2019; Nielsen et al. 2019; Martins et al. 2020). For homogenisation, samples can either be already sorted (‘picked’) invertebrate specimens, i.e. without debris such as plant material or sand (Carew et al. 2013; Elbrecht et al. 2017), or they can be complete environmental samples including specimens and debris (Hajibabaei et al. 2019b; Pereira-da-Conceicoa et al. 2019). Given the often high demands in terms of working hours per sample for sorting invertebrate samples with hundreds to thousands of specimens (Haase et al. 2006) there is an increasing demand...
for direct sample homogenisation without picking and sorting (Blackman et al. 2019). After sample homogenisation, DNA is extracted. Different DNA extraction protocols are routinely applied in current metabarcoding studies (e.g. Majaneva et al. 2018; Elbrecht and Steinke 2019; Hajibabaei et al. 2019a). Samples are then amplified, individually labeled, and sequenced, targeting a specific gene region (e.g. commonly cytochrome oxidase subunit I (COI) for macroinvertebrates (Elbrecht et al. 2019; Elbrecht and Leese 2017; Hajibabaei et al. 2019a) before bioinformatics processing and taxonomic assignment using search engines or tools like BLAST on reference databases such as BOLD (Ratnasingham and Hebert 2007) or NCBI GenBank (Johnson et al. 2008). The resulting taxon list is then used for biological interpretations. Given the many advantages such as taxonomic resolution, speed, and the possibility for data validation, the metabarcoding analyses of bulk or environmental DNA samples are used in hundreds of research studies, but are also increasingly considered in biomonitoring programs (Hänfling et al. 2016; Elbrecht et al. 2017; Haase et al. 2018; Hering et al. 2018; Li et al. 2019; Meyer et al. 2020; Pont et al. 2021).

Many of the above-mentioned laboratory steps have been verified thoroughly and best-practice strategies exist. For example, different extraction protocols have been analyzed (Majaneva et al. 2018; Loos and Nijland 2020), DNA polymerases have been studied for biases in metabarcoding studies (Nichols et al. 2018; Sze and Schloss 2019), the choice of markers and primers has been validated to a great extent (Elbrecht and Leese 2015; Elbrecht et al. 2017, 2019; Vamos et al. 2017), and the influence of indexing on taxon recovery has been studied (Schnell et al. 2015; Zizka et al. 2019). For all aspects, good solutions for macroinvertebrate analyses exist. However, only a few studies have so far validated the first laboratory step, i.e. sample homogenisation, despite the fact that this may have a strong effect on all downstream processes.

Sample homogenisation has been done without a drying step (Majaneva et al. 2018; Hajibabaei et al. 2019b; Pereira-da-Conceiccoa et al. 2019) as well as with prior drying of the specimens (Elbrecht and Leese 2015; Elbrecht and Steinke 2019; Zizka et al. 2019). While wet-grinding is probably more time-efficient and the resulting ‘slur’ easier to handle with a pipette, the homogenate itself has to be dried afterward, since ethanol inhibits downstream protocol steps such as DNA extraction and PCR (Schrader et al. 2012). When drying specimens before homogenisation the dry weight can be measured more easily. However, dry tissue powder can be electrostatically charged and therefore hard to handle (Elbrecht and Steinke 2019). More importantly, cross-contamination of the highly volatile components may occur.

For sample homogenisation, different devices are used. They can be divided into two main categories: bead mills and blenders. Bead mills work by accelerating small, hard particles in a closed container like a falcon tube to break down tissue into small fragments. Blenders work with a rapidly rotating blade that slices the tissue. While most bead mills rely on single-use plastics, blenders offer the option to be cleaned. But both methods have downsides to consider: while single-use plastics are ideal in terms of avoiding cross-contamination they produce a lot of waste and are expensive compared to the costs of other parts of the workflow. Prices may vary but go up to 15 € per sample. For sufficient homogenisation, runtime varies from 2 up to 30 mins (Elbrecht et al. 2020; Beermann et al. 2021). Also, sample volumes are low compared to blenders and mostly limited to 100 ml per sample. Bigger grinding chambers are available (e.g. IKA, Staufen im Breisgau, Germany) but increase costs even more.

Blenders on the other hand can handle large sample volumes more easily (e.g. 600 ml in Pereira-da-Conceiccoa et al. 2019) and quickly, but need to be sterilized extensively. In most studies, a combination of bleaching (5–12%) and UV-radiation (up to 30 min) was used to clean the blender (Majaneva et al. 2018; Hajibabaei et al. 2019b; Pereira-da-Conceiccoa et al. 2019). While being the safest way, this is not feasible in large-scale, routine biomonitoring. Furthermore, bleach is also highly corrosive to metals and must be discarded according to strict laboratory guidelines.

Here, we aimed to develop a fast, robust, cheap, and reliable sample homogenisation protocol that overcomes the above-mentioned limitations of both methods, i.e. does not produce difficult to discard waste and avoid single-use plastics. We tested the performance of the new protocol using six sorted Malaise trap and six unsorted stream kick-net samples. We used 14 replicates per sample and included many negative controls at different steps of the protocol to quantify the impacts of insufficient homogenisation and cross-contamination.

Materials and methods

Study design

The design of this study is summarized in Fig. 1.

Sample acquisition

Two different samples types were used in this study: (i) six unsorted stream kick-net samples from a study conducted by Haase and Pilotto (2019), and (ii) six size-sorted (small < 4 mm; large > 4 mm) samples from Malaise traps that had been set up in the Rhine-Mine-Observatory for 2 weeks in July 2020. The Rhine-Main-Observatory is a Long-Term Ecological Research (LTER) site (Haase et al. 2016; Mirlt et al. 2018) east of Frankfurt, Germany (https://deims.org/9f9ba137-342d-4813-a5e58-a60911e3a-bc1). All samples were preserved in 96% technical ethanol.

Tissue homogenisation and cleaning

Samples were homogenized in a common kitchen blender (Mini Blender & Blender Smoothie, Homegeek, China) at 25,000 RPM for 3 min together with the preservation liq-
uid. To reduce heating of the samples, samples were cooled to -20 °C prior to homogenisation. After homogenisation, samples were transferred back to their respective collection container and stored at -20 °C until DNA extraction. The blade and container of the blender were cleaned with ddH₂O until no remainders of the sample were visible. After that, the container and blades were rinsed with ddH₂O and used for the next sample. This cleaning procedure was repeated 1 – 6 times to assess how many rounds of cleaning were necessary for full decontamination. Before each sample, the blender was filled with 100 ml 96% EtOH, which was treated as if it was a sample to produce a blender negative control. While all kick-net samples were treated with ddH₂O, DIY-DS was used for the Malaise trap samples. In between the homogenisation of the 2 size fractions for the Malaise traps we only shortly rinsed the blender container with ddH₂O. Twelve additional negative controls were added to the samples after the homogenisation step to control for any kind of cross-contamination independent of the blending procedure. Homogenisation of the 2 size fractions for the Malaise traps was optically verified using a digital microscope (Keyence VHX-6000, Keyence, Osaka, Japan). Details on the blender container can be provided on request.

**DNA extraction**

Before tissue lysis, the two size fractions of the Malaise trap samples were pooled in a 1:5 ratio (large-small, 5 ml and 25 ml) as suggested by Elbrecht et al. (2020). 500 µl of each sample was used for DNA extraction, which corresponds to roughly 0.1% of the complete 500 ml sample volume. All samples were centrifuged at maximum speed (14,000 × g) for 5 minutes to pellet the tissue. All remaining Ethanol was then evaporated in an Eppendorf Concentrator (Eppendorf, Hamburg, Germany) with a speed of 1400 RPM and at a temperature of 60 °C. Subsequently, 300 µl of TNES buffer was added to each tube as well as a small amount of silicon-carbide beads (0.1 mm diameter, Biospec Products, Bartlesville, USA). The tissue pellet was broken up by 30 s of bead-beating in a FastPrep Bead Beater (MP Biomedicals, Eschwege, Germany). All subsequent processing steps were completed on a Biomek FX liquid handling workstation (Beckman Coulter, Brea, CA, USA). 60 µl of tissue dissolved in TNES buffer was taken out twice of every tube and mixed with 133 µl TNES and 7 µl Proteinase K (10 mg/ml) and digested for 3 h at 55 °C. From this point onwards, the plates containing replicate samples (see Fig. 1 for an overview of the replication) were never opened at the same time. DNA was extracted using a modified version (see Buchner et al. 2021) of the NucleoMag Tissue kit (Macherey Nagel, Düren, Germany). Extraction success was visualised on a 1% agarose gel.
qPCR validation, PCR, and library preparation

To control for possible contamination of the negative controls, all samples were amplified in a quantitative PCR (qPCR) in 20 µl reactions containing 1× perfeCTa FastMix, 300 nM of each primer (fwh2F, fwhR2n (Vamos et al. 2017)) and 1 µl of extracted DNA, filled up with PCR-grade water with the following cycling conditions: 30 s of initial denaturation, 80 cycles of 5 s denaturation at 95 °C, 30 s of annealing at 58 °C and 20 s of extension at 72 °C following a standard melting curve on a StepOne Plus thermocycler (Applied Biosystems, Foster City, CA, USA). The success of the qPCR was verified via melting curve analysis and visualisation on a 1% agarose gel.

The PCR for the metabarcoding library was done in a two PCR step protocol (Zizka et al. 2019). Samples were amplified in a first PCR with the Qiagen Multiplex Plus Kit (Qiagen, Hilden, Germany) with a final concentration of 1× Multiplex Mastermix, 1× Corralload Loading Dye, 100 nM of each primer (fwh2F, fwhR2n (Vamos et al. 2017)), 2.5 µl of DNA filled up to a final volume of 25 µl with PCR-grade water. For amplification, a touchdown protocol was used: 5 min of initial denaturation, 10 cycles of 30 s denaturation at 95 °C, 30 s of annealing at 68–59 °C decreased by 1 degree in each cycle, 30 s of extension at 72 °C followed by 20 cycles of 30 s denaturation at 95 °C, 30 s of annealing at 58 °C, 30 s of extension at 72 °C finished by a final elongation step of 10 min at 68 °C. For subsequent demultiplexing, each of the PCR plates was tagged with a unique combination of inline-tags. The primers also contain a universal binding site for the second step PCR primer to anneal (see Suppl. material 2: Table S1 for detailed information on the primers used in this study).

In the second PCR, samples were amplified with the Qiagen Multiplex Plus Kit with the same final concentrations except that 1 µl of first step PCR product was used as a template. For amplification the following protocol was used: initial denaturation for 5 min at 95 °C, 25 cycles of 30 s denaturation, and 60 s of combined annealing and extension at 72 °C finished with a final elongation of 10 min at 68 °C. In the second PCR, each of the 96 wells was individually tagged so that the combination of inline-tag from the first PCR step and index-read of the second step yields a unique combination. The success of the PCR was visualisation on a 1% agarose gel.

PCR product concentrations were normalised using the SequalPrep Normalisation plate (Invitrogen, Carlsbad, CA, USA). Normalised products were then pooled to the final library in equal parts for all samples. The library was concentrated using the NucleoSpin kit (Macherey Nagel, Düren, Germany) and dual-sided size selected (right ratio: 0.6; left ratio: 0.75) with the NucleoMag size-select kit (Macherey Nagel, Düren, Germany). Library concentration was quantified on a Fragment Analyzer (High Sensitivity NGS Fragment Analysis Kit; Advanced Analytical, Ankeny, USA). The library was then sequenced using the HiSeq X platform with a paired-end (2×151 bp) kit at Macrogen Europe.

Bioinformatics, filtering, and statistical analysis

For analysis of the qPCR results, raw fluorescence values were exported from the instrument and baseline-corrected with the LinRegPCR software (Ruijter et al. 2009). The baseline-corrected data was then used for inspection of amplification- and melting curves.

Raw data of the sequencing run yielded 638,892,616 reads and was delivered demultiplexed by index-reads. Index jump (sensu Schnell et al. 2015) rate for this sequencing run was estimated with a custom python script by counting all index combinations used in the study that were found in the unassigned reads and dividing the number by the sum of all reads. Further demultiplexing by inline-tag was done with the python script “demultiplexer” (v1.0.5; https://github.com/DominikBuchner/demultiplexer). Sequences were subsequently processed with the JAMP-pipeline (v0.67; https://github.com/VascoElbrecht/JAMP). Paired-end reads were merged using Usearch (v11.0.667, Edgar 2010) via the command U_merge(fastq_pctid=75). Primers were trimmed using Cutadapt (v2.5, Martin 2011), and only reads with a length of 205 bp (± 10) were retained for further analysis. Before OTU clustering with a similarity threshold of 97%, reads were dereplicated and singletons were excluded. Only clusters with at least 0.01% abundance in one sample were used in further analysis. Taxonomic assignment was carried out using BOLDigger (v1.2.2, https://github.com/DominikBuchner/BOLDigger; Buchner and Leese 2020). The best hit was determined with the BOLDigger method. This resulted in a raw read table including taxonomic assignment (Suppl. material 3: Table S2).

To control for contamination on the robotic deck, technical replicates of the plates were merged retaining only the mean read number of both replicates if reads were present in both of the replicates. After that, the maximum number of reads for each OTU in all additional negative controls was calculated and subtracted from all reads of the respective OTU to remove noise introduced by the laboratory workflows resulting in a cleaned read table (Suppl. material 4: Table S3). All cleaning steps described above were done with a custom python script (Suppl. material 5: Script 1). All figures were created using the python package “seaborn” (Waskom et al. 2020) with custom python scripts.

To compute the similarity between samples the Jaccard index was used (Jaccard 1912). Rarefaction analysis was computed with a custom python script. Samples were drawn randomly without replacement 1000 times to generate a distribution from which the plots were generated.

Results

Optical verification of homogenisation success

No identifiable parts of the animals were left after 3 min of homogenisation. However, particle size was, overall, coarser for the Malaise trap samples (Suppl. material 6: Fig. S1).
Amplification and melting curve analysis

All invertebrate samples were amplified successfully during qPCR analysis at first try in both technical replicates (Suppl. material 7: Fig. S2). Some of the blender negative controls, as well as the additional negative controls, showed a signal in later cycles that were not consistent across replicates and most likely reflected primer dimers in qPCR. The melting curve analysis showed a clear peak at above 75 °C for all of the invertebrate samples, while negative and mixing controls showed either a double peak or a single peak below 70 °C indicating the formation of primer dimers. Primer dimers were clearly visible in the agarose gel verifying this assumption. The primer used here is known to generate primer dimers (see Vamos 2017, Suppl. material 4: Table S3).

Quantification of tag jumps

Sequencing yielded 110,641,213 reads that could not be assigned to any of the index combinations used in this study. Only 15 of these reads had a combination of the used twin-indices resulting in a very low index jumping rate of 2×10^-8.

OTU richness and potential cross-contamination

Mean OTU richness across all seven replicates ranged from 48.71 (45–50) to 74.00 (67–83) for the kick-net samples. For the blender negative controls that were rinsed with ddH₂O, the mean richness ranged from 0.14 (0–1) to 1.71 (0–6). Regarding the blender negative controls, none of the OTUs was found in all 7 replicates. For the Malaise trap samples mean OTU richness ranged from 293.57 (279–301) to 446.71 (437–456). For the blender negative controls rinsed with DIY-DS, the mean richness ranged from 1.14 (0–4) to 4.00 (0–20). None of the OTUs was found in all 7 replicates either (Fig. 2, top panel).

The mean number of reads per sample was overall higher for the kick-net samples than for the Malaise trap samples (1.28×10⁶–1.72×10⁶ vs. 7.31×10⁵–1.04×10⁶). The mean number of reads in the blender negative controls was overall lower for the DIY-DS treatment than for the ddH₂O treatment (192 vs. 20,718). The mean read numbers were largely influenced by one OTU having a high number of reads for only one of the mixing negative controls (Fig. 2, lower panel).

Replicate consistency

Mean Jaccard similarity between the 7 extraction replicates was overall high for both sample types (kick-net samples: 0.81 vs. Malaise trap samples: 0.84) with the spread being higher for the kick-net samples (0.58 – 0.94) in comparison to the Malaise trap samples (0.78 – 0.9) mainly due to sample number 5 of the kick-net samples (Fig. 3).

Figure 2. Mean OTU richness for kick-net samples (top left panel) and Malaise trap samples (top right panel). Mean sum of all reads across all 7 replicates for one sample for the kick-net samples (bottom left panel) and the Malaise trap samples (bottom right panel). The sample number also indicated the rounds of cleaning after each sample. Error bars indicate the 95% confidence interval ranging from percentile 2.5 to 97.5.

Figure 3. Pairwise comparison of extraction replicates. For each pair of extraction replicates within one sample the Jaccard similarity was computed for the kick-net samples (left panel) and the Malaise trap samples (right panel).

Rarefaction analysis for the extraction replicates of samples (Fig. 1, C) showed that one sample is sufficient in most cases to recover about 80% of the maximum possible OTU richness across all 7 extraction replicates. To recover 90% of the maximum possible OTUs, more replication effort is needed, typically >2. Of the kick-net samples, only sample number 5 is an outlier to that general pattern (Fig. 4).

Discussion

Our study aimed to develop and test an improved invertebrate homogenisation method that is easy to apply, robust,
and reliable, while being cost- and time efficient. Blenders are already used for species homogenisation in metabarcoding studies, however, none of those met the above mentioned criteria. Majaneva et al. (2018) used 5% bleach for 15 minutes and 30 minutes of UV sterilization, Hajibabaei et al. (2019b) used ELIMINase and also UV treatment for 30 minutes, Pereira-da-Conceição et al. (2019) only state that they used 12% bleach without a volume or a time. All methods produce hazardous waste or are high in costs and take at least 30 minutes per sample. In addition to that, all studies did not report OTU/ESV numbers in the negative controls or only used negative controls in PCR, which does not validate the effectiveness of the cleaning approach. In comparison, our newly developed protocol reduces sample processing time significantly without the risk of cross-contamination, which is important for the further implementation of DNA metabarcoding and metagenomic techniques in the increasingly important field of invertebrate bioassessment and monitoring.

**Cross-contamination**

A central concern for bulk sample metabarcoding using blenders rather than single-use plastics is the risk of cross-contamination. The approach we present minimizes this risk effectively due to three points: i) pipetting homogenized samples in ethanol (wet grinding) limits the risk of electrostatic charge and thereby ‘jumping’ specimens. ii) Both tested cleaning procedures, i.e. cleaning with ddH2O and the DIY-DS, proved to be highly effective. While we sporadically saw that some blender negative controls contained low read numbers of single OTUs, this was never the case for all 7 extraction replicates of the given sample. This suggests, that the contamination did not happen in the blender (or only by one of the few left molecules). This was further confirmed by the observation that some of the OTUs found in the blender negative controls were not found in the sample processed before. Furthermore, the DIY-DS reduced this already sporadic and low contamination even further and is thus recommendable. iii) The stringent replication scheme, i.e. performing extraction and downstream analysis twice in physically independent plates that are never open on the benchtop at the same time, further limits the possibility for cross-contamination. This allows to control for low-level cross-contamination by accepting reads or OTUs / ESVs that are only found in both replicates.

**Replicate consistency**

For the two sample types analyzed here, stream benthic macroinvertebrates and insect Malaise trap samples, we observed a high consistency between extraction replicates with typically 80% or higher OTU overlap among replicates (Jaccard similarity). Stream invertebrate kick-net sample 5 was an outlier sample with only 60–70% overlap. The reasons for the lower overlap can be insufficient blending of the sample or independent replicate contamination. Independent contamination seems unlikely, as this was not observed in any of the blender negative controls performed between samples. Microscopic inspection of homogenized tissue did also not indicate systematic differences between sample 5 and all others (Suppl. material 6: Fig. S1). However, this cannot be fully ruled out.

To further improve replicate consistency, i.e. maximize the overlap between replicates, it might be beneficial to first perform lysis on a large fraction of the sample and then perform downstream analysis using two (or more) replicates. Alternatively, and in particular, when the aim is to recover the maximum of species diversity, more extraction replicates should be performed. However, our analysis shows that we already detected 80% of the OTUs found in all 7 extraction replicates with a single replicate.

**Proposition for routine application**

Invertebrate assessment and monitoring using bulk DNA metabarcoding or metagenomics require fast, reliable, and validated protocols that are ideally economically competitive and environmentally friendly (Blackman et al. 2019) and yield comprehensive data on community composition. For this purpose, homogenizing samples
with blenders rather than single-use plastics seems more suitable, yet so far suffered from time-consuming and hazardous chemicals used for cleaning (highly concentrated bleach). Our study proves that the cleaning procedure can be done fast and reliably without aggressive chemicals using DIY-DS. Furthermore, the solution costs only about 1 Cent per cleaning (100 ml) making it economically feasible at the largest scale. Furthermore, while commonly used plastic or glass blenders can break when stones or hard-shelled specimens are homogenized in the blender, we here propose a solution using a stainless steel blending container (Suppl. material 8: Fig. S3).

With this container, wet grinding using ethanol preserved samples can be done fast and reliable even with large volumes of 500 ml. We could process about 30 complete bulk samples per 8 h with one person and one blender. Thus, the approach based on wet-sample grinding is not only technically feasible, scientifically reliable, economically competitive, and environmentally friendly, it offers great speed and is scalable allowing for large-scale DNA-based biomonitoring.

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Data accessibility statement

Demultiplexed raw read data for this publication has been uploaded to Zenodo.org and can be accessed via 10.5281/zenodo.5039930.

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**Supplementary material 1**

**Protocol 1 – DIY-DS**
Authors: Dominik Buchner, Peter Haase, Florian Leese
Data type: text
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Link: https://doi.org/10.3897/mbmg.5.67533.suppl1

**Supplementary material 2**
**Table S1. PCR primers used in this study**
Authors: Dominik Buchner, Peter Haase, Florian Leese
Data type: text
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Link: https://doi.org/10.3897/mbmg.5.67533.suppl2

**Supplementary material 3**
**Table S2. Raw read table**
Authors: Dominik Buchner, Peter Haase, Florian Leese
Data type: text
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**Supplementary material 4**
**Table S3**
Authors: Dominik Buchner, Peter Haase, Florian Leese
Data type: text
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**Supplementary material 5**
**Script 1**
Authors: Dominik Buchner, Peter Haase, Florian Leese
Data type: code
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**Supplementary material 6**
**Figure S1. Pictures were taken with a digital microscope (Keyence VHX-6000, Keyence, Osaka, Japan)**
Authors: Dominik Buchner, Peter Haase, Florian Leese
Data type: image
Explanation note: Top row: Kick-net samples. Bottom row: Malaise trap samples.
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Link: https://doi.org/10.3897/mbmg.5.67533.suppl6

**Supplementary material 7**
**Figure S2. Baseline-corrected amplification curves (left half) and melting-curves (right half) for A & B) kick-net samples and C & D) malaise trap samples**
Authors: Dominik Buchner, Peter Haase, Florian Leese
Data type: statistical data
Explanation note: Different colors indicate different technical replicates of the same sample.
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**Supplementary material 8**
**Figure S3**
Authors: Dominik Buchner, Peter Haase, Florian Leese
Data type: image
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