An urban Blitz with a twist: rapid biodiversity assessment using aquatic environmental DNA


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Abstract
As global biodiversity declines, there is an increasing need to create an educated and engaged society. Having people of all ages participate in measuring biodiversity where they live helps to create awareness. Recently, the use of environmental DNA (eDNA) for biodiversity surveys has gained momentum. Here, we explore whether sampling eDNA and sequencing it can be used as a means of rapidly surveying urban biodiversity for educational purposes. We sampled 2 × 1 L of water from each of 15 locations in the city of Trondheim, Norway, including a variety of freshwater, marine, and brackish habitats. DNA was extracted, amplified in triplicate targeting the barcoding fragment of COI gene, and sequenced. The obtained data were analyzed on the novel mBRAVE platform, an online open-access software and computing resource. The water samples were collected in 2 days by two people, and the laboratory analysis was completed in 5 days by one person. Overall, we detected the presence of 506 BINs identified as belonging to 435 taxa, representing at least 265 putative species. On average, only 5.4% of the taxa were shared among six replicates per site. Based on the observed diversity, three distinct clusters were detected and related to the geographic distribution of sites. There were some taxa shared between the habitats, with a substantial presence of terrestrial biota. Here we propose a new form of BioBlitz, where with noninvasive sampling effort combined with swift processing and straightforward online analyses, hundreds of species can be detected. Thus, using eDNA analysis of water is useful for rapid biodiversity surveys and valuable for educational purposes. We show that rapid eDNA surveys, combined with openly available services and software, can be used as an educational tool to raise awareness about the importance of biodiversity.

KEYWORDS
aquatic habitats, cytochrome c oxidase subunit I gene, environmental DNA, environmental education, mBRAVE, metabarcoding
1 INTRODUCTION

Global biodiversity is imperiled with an estimated one million species at risk of extinction (IPBES, 2019). There is an urgent need to build consensus around action and reverse the impending loss of life from our planet. Conducting rapid biodiversity assessments in urban centers such as the City Nature Challenge (http://citynaturechallenge.org/) is increasingly used as means by which to educate all ages and sectors of society about the importance of biodiversity. Many biodiversity observation programs exist to engage with the public (e.g., BioBlitz; Postles & Bartlett, 2018). However, when such surveys occur, the amount of equipment, resources, and time needed from experts to survey diversity across the tree of life quickly becomes intractable for many educators. Thus, educational and outreach programs using rapid biodiversity surveys are often limited in their taxonomic breadth, are infrequent, and are offered mostly by larger organizations (Postles & Bartlett, 2018; Ruch et al., 2010).

New DNA-based technology has the power to solve current limitations of education programs wanting to teach about biodiversity through firsthand experiences. Namely, in the last decade, the scientific community has realized we can survey species by finding trace amounts of their DNA left behind in the environment (i.e., environmental DNA “eDNA”), and this has gained tremendous momentum (Cristescu & Hebert, 2018; Deiner et al., 2017; Thomsen & Sigsgaard, 2019). If biodiversity surveys using DNA from the environment are easy to apply, then they can be used as an educational outreach tool for engaging the public to generate biodiversity knowledge to create awareness. Furthermore, because environmental DNA sequencing services are common and commercially available from many companies around the world, access for educators at all levels is possible. The analysis of the data generated is, however, the real limiting factor, if not provided by such companies. Bioinformatic processing of the sequencing output from metabarcoding requires access to powerful computing servers and a certain degree of experience with command-line bioinformatic software. This obstacle...
can be now overcome by more automated, user-friendly, and with a graphic user interface (GUI), online platforms created to make data processing for metabarcoding accessible to all. Examples of these platforms are mBRAVE or SLIM (https://trtcrd.github.io/SLIM/; Dufresne et al., 2019).

To show how eDNA surveys can potentially be used in an urban setting for educational purposes, we performed a test case in the city of Trondheim, Norway (Figure 1). Trondheim is situated at the Trondheim Fjord in Central Norway at 63° North. It has almost 200,000 inhabitants and is the third largest city in Norway. Although the city center is rather developed, there are numerous parks mixed within residential areas, and outside of the city center itself, there are gardens and small natural forest patches. To the west, the city is bordered by the forest Bymarka, partially a nature reserve, while to the east, the city is bordered by farm fields and the forest Estendstadmarka. Both these natural areas have a rolling landscape, with the highest peak Storheia at 565 m above sea level. They contain the headwaters of multiple streams running toward the city center and the Trondheim Fjord. The river Nidelva originates from the large lake Selbusjøen to the south-east, splits the city, and empties into the fjord north of Trondheim. The river is about 31 km long and is regulated for hydropower purposes, and the water discharge level is heavily influenced by frequent variation in electricity production intensity (“hydropeaking”). However, a minimum discharge of 30–40 m³/s is required, partly to ensure that the viable trout and salmon populations in the river have acceptable living conditions. The river, streams, and ponds in Trondheim are influenced by human activity to a variable degree. In some areas, the aquatic habitats are strongly influenced by land use (e.g., road infrastructure), while other sections are influenced by sewage overflow in the drainage system. In 2016, the city of Trondheim treated seven lakes in Bymarka with rotenone to remove the invasive common roach (Rutilus rutilus (Linnaeus)). Several ponds around Trondheim are known to host great-crested newt (Triturus cristatus (Laurenti)) and common newt (Lissotriton vulgaris (Linnaeus)), but their presence near the city center has declined due to habitat alteration (Tilseth, 2008). The Trondheim Fjord was heavily polluted by heavy metals and organic pollutants but has improved considerably after a large-scale clean-up and dredging in 2015–2016.

With this backdrop, we tested whether eDNA could be rapidly sampled and sequenced from 15 sites and taxa lists determined through the use of DNA-based identifications of observed taxa.
sequences to the BOLD database (Ratnasingham & Hebert, 2007) using the novel Multiplex Barcode Research And Visualization Environment (mBRAVE) platform (http://www.mbrave.net/). To make use of the data and show its potential application for educational purposes, we gathered biodiversity scientists and university students for a workshop at the 8th International Barcode of Life Conference (iBOL2019); most of whom had never sequenced or analyzed environmental DNA data. We spent one intensive day learning the basics of the methods and the online analysis platform mBRAVE, but then returned home and collaborated from all over the world to analyze the data. The diversity of participants and backgrounds is represented by our authors coming from nearly every continent and representative of 14 countries. Together, after our return home, we used our rapid eDNA biodiversity assessment to determine patterns in observed biodiversity. We asked (a) how much total diversity could be detected at the selected locations, and (b) whether patterns of diversity changed due to sampling location or habitat. Additionally, we (c) tested whether we could detect rare amphibians: the great-crested newt (T. cristatus) and the common newt (L. vulgaris), and the invasive common roach (R. rutilus). Lastly, (d) we asked whether eDNA was transported across the landscape and between habitats.

2 | MATERIALS AND METHODS

2.1 | Sampling

Two 1 L water samples were collected from each of 15 sites in Trondheim, Norway (Figure 1) on 13 and 14 2019 May (Table S1). The sites included ponds, small streams, a lake, rivers, and the sea. At each site, water was collected by submerging two sterilized (5% bleach, flushed with the sampling site water) 1 L rectangular polyethylene terephthalate bottles (Nalgene/VWR International) just below the surface, with only bottle and gloved hand touching the water. At the running water sites, water was collected upstream of the collector was standing. Samples were stored in a cooler box. The air temperature was between 0°C and 10°C, so the samples were kept at approximately ambient temperature until filtered.

2.2 | Clean laboratory environment

The samples were processed in a unidirectional controlled facility. Filtration was done in a separate room, where no other environmental or organismal samples were processed, and all surfaces were decontaminated before use (with a 5% bleach solution and 70% ethanol). Each filter was folded with a fresh pair of forceps, which were decontaminated overnight in 5% bleach solution before the second day of filtration. The DNA extraction was done in a laboratory where all surfaces and all necessary equipment were decontaminated with a 5% bleach solution and 70% ethanol, while wearing a full-body suit, face-mask, and double gloves. After a workday, UV-light was set to run overnight. Respective DNA extracts were transferred to a pre-PCR room where PCRs were setup. After amplification, the libraries were prepared in the pre-PCR room and the template was added in the post-PCR room.

2.3 | Sample filtration and DNA extraction

Samples (N = 30) were transferred to the local university laboratory (NTNU University Museum, Norwegian University of Science and Technology) within 2 hr after sampling. All samples were filtered on the same day they were collected and in the order they were sampled using an electrical vacuum pump connected to a manifold (Pall Laboratory) carrying three individually operated filter holder bases. Samples were filtered through sterile and individually packed 0.45 µm mixed cellulose ester filters (ø 47 mm) attached to a 300 ml reservoir (Pall Laboratory). Filters were folded inward three times with decontaminated forceps and placed into 1.5-ml centrifuge tubes with 800 µl of ATL lysis buffer (Qiagen GmbH). To control for contamination during the filtration process on each day, a negative control was made by filtering 500 ml of molecular grade water in the same manner as samples after all field samples were processed (Table S1). Hereafter, these are referred to as sample blanks. Prior to DNA extraction, the lysis buffer was removed from the filter and partitioned into two tubes, each containing ~400 µl of buffer ATL. The DNeasy Blood and Tissue kit (Qiagen) was used for DNA extraction from the lysis according to the manufacturer’s instructions for tissues with the exception that they were shaken horizontally at 250 rpm at 37°C overnight. The purified DNA was eluted with 200 µl of molecular grade water. No negative DNA extraction controls were created, but the sample blanks served as a full process controls for all laboratory procedures. Quantification of the DNA extracts was not performed.

2.4 | PCR amplification and sequencing

The extracted DNA from each of the two samples from the 15 sites and two sample blanks (N = 32) were amplified using a common two-step PCR method, with the highly degenerate BF2-BR2 primers (Elbrecht & Leese, 2017) with attached Illumina adapters 5′-TCGTCGACGCTTACAGATGCTTGATAAGAGACAG-3′ (forward) and 5′-GTCTCTGGGTGTAGCTACAG-3′ (reverse) in the first PCR. The primers target an approximately 420 bp long fragment of the mitochondrial cytochrome c oxidase subunit I gene (COI). Each DNA sample was amplified in three technical PCR replicates and a negative control was created using molecular grade water in place of the sample. Each PCR had a final volume of 25 µl containing 2 µl DNA template (DNA was diluted 1:10 prior to amplification to reduce the chance of inhibition), 17.8 µl molecular biology grade water, 2.5 µl 10X reaction buffer (200 mM Tris HCl, 500 mM KCl, pH 8.4), 1 µl MgCl2 (50 mM), 0.5 µl dNTPs mix (10 mM), 0.5 µl of each primer (10 mM), and 0.2 µl Platinum Taq polymerase (5 U/µl) (Invitrogen). The PCR conditions were, with a heated lid, 94°C
for 5 min, followed by a total of 35 cycles of 94°C for 40 s, 50°C for 1 min, and 72°C for 30 s, and a final extension at 72°C for 2 min. PCR products were visualized on a 1.5% agarose gel to check the amplification success.

In the second PCR, the Illumina tailed amplicons (\(N = 96\)) were dual indexed (File S1), using NextEra XT Index 1 and 2 primers and NextEra XT Index v2 set D (Illumina, FC-131-2004) in a reduced-cycle PCR (15 cycles) according to the manufacturer’s protocol. The amplification reactions contained the same reagent concentrations as above with three modifications: 3 \(\mu\)l of product from the first PCR, 1 \(\mu\)l of each primer, and 0.25 \(\mu\)l of Platinum Taq (Invitrogen). PCRs were not purified between the first and second amplification. Indexed amplicons were pooled by equal volume (5 \(\mu\)l each) into a single library. The library was purified and size-selected (>200 bp fragments retained), using SPRIselect (Beckman Coulter) with a ratio of 0.92. Library was quantified using Qubit™ dsDNA HS Assay Kit (Thermo Fisher), following the manufacturer’s protocol. The library was loaded at a 14 pM concentration with 6% PhiX added. Sequencing was performed on a MiSeq v3 flow cell sequencing paired end 2 \(\times\) 300 bp and index 8 + 8 bp at the Genomics Core Facility, Oslo University Hospital, Oslo, Norway. The run folder was imported into the Illumina Local Run Manager version 2.0.0 and was used to complete the base calling and demultiplex the libraries to generate the fastq files for each sample (\(N = 96\)).

### 2.5 Data upload, filtering criteria, and taxonomic assignment

The resulting 96 fastq files from the Illumina MiSeq run were imported into mBRAVE using the sample batch function. Each demultiplexed library (i.e., fastq file) is now referred to as a “TRD run” for downstream analysis. Every parameter described here was retrieved from the mBRAVE platform and was available to the user as last accessed in July 2019. An extended description of the methods can be accessed in File S1. Thus, we briefly describe the methods used to format, clean, and denoise the raw sequence data before taxonomic identification and biodiversity analyses.

For each library now consisting of a TRD run, the paired end merging of MiSeq reads required a minimum 20 bp overlap between the forward and reverse reads, while allowing up to five nucleotide substitutions. Both front and end part of the resulting sequences were trimmed for 20 bp to remove the primer sequence, followed by trimming of total sequence length down to 500 bp. Low-quality sequences were removed if the average quality value (QV) was less than 20 or sequences were shorter than 200 bp. This filtering step allowed for a max of 2% nucleotides with <20 QV value and max 1% nucleotides with <10 QV value. Sequences fulfilling these criteria were de-replicated and clustered on mBRAVE as operational taxonomic units (OTUs) using a 2.5% similarity threshold. Obtained OTUs were taxonomically assigned using an initial 2% ID distance threshold to the reference sequences. When the obtained sequences could not be assigned to any reference sequence, the threshold levels were relaxed until a hit was found for higher taxonomic ranks: genus (3%), subfamily (4%), family (5%), order (7%), class (10%), and phylum (12%). The publicly available BOLD reference libraries for Insecta, noninsect Arthropods, nonarthropod invertebrates, Chordata, and bacteria, as well as a standard contamination reference database, were used for identification of all OTUs. Whenever a taxonomic assignment from a reference sequence returned more than one species name, taxonomy was assigned to higher taxonomic ranks with no conflicting taxonomy.

After selecting parameter values, mBRAVE automatically applied the same parameters to each TRD run in our dataset. For each run, a summary was generated and checked to ensure a good fit with our filtering criteria. Data visualizations within the mBRAVE environment allowed for the data to initially be reviewed. These visualizations included sequence length distribution, GC composition distribution, run QV score distribution, and BIN (Barcode Index Numbers, Ratnasingham & Hebert, 2013) count versus OTU count. Where low-quality sets were observed, the low-quality runs were removed. The parameters chosen for this study were adjusted after considering the quality comparison in mBRAVE. Data were then combined into “sets” within the mBRAVE environment, where biological and technical PCR replicates for each sample site were combined as a single set that could be further analyzed.

### 2.6 Biodiversity and distribution analyses

To assess general diversity patterns across the city of Trondheim, a summary report for all sets was generated to calculate the number and list of taxa detected in each TRD run. In order to have an overview of which taxa (i.e., Bins) were shared among samples and the negative filter controls, the beta diversity tool with the Jaccard similarity index was used within the mBRAVE environment using the sample sets as units of comparison. A heatmap was generated with either the shared BINS or the index value between the paired sets.

To document the composition of observed diversity in each run, it was considered that freshwater, marine, brackish, and terrestrial taxa could be observed. Thus, sites were designated as belonging to one of three different realms (marine, freshwater, brackish) (Table S1). A brackish site was designated as such due to the tidal activity and temporal admixture of freshwater and seawater. When taxa could be assigned to a species, the known habitat designation was indicated, which included marine, freshwater, brackish, and terrestrial using the WoRMS database (World Register of Marine Species, Horton et al., 2017) or other Internet sources and, in some cases, the species’ original descriptions. When taxa could not be assigned to species, the sequences and records were cross-validated with the following public databases: GBIF (https://www.gbif.org/); Fauna Europaea (https://fauna-eu.org/) and NCBI GenBank (https://www.ncbi.nlm.nih.gov/genbank/).
To assess similarity within sites among the biological and technical replicates, diversity at a local scale was evaluated with the BIN comparison tool in mBRAVE. This generated a Venn diagram with unique and shared BINs. Two approaches were adopted, the first without excluding any BIN, and secondly by accepting only the BINs shared at least by two replicates (biological or technical). We present only the data based on the latter.

Beta diversity analysis was performed in mBRAVE for all TRD runs using the beta diversity grouping runs by sets, composed of all replicates belonging to a particular site or habitat. To assess the impact of singleton/doubleton removal, the beta diversity analyses were run for every BIN (not excluding those found in less than two replicates), and this analysis was run again excluding singletons and doubletons (i.e., selecting in Minimum Sequences per BIN or Species option: 1, 2, or 3, respectively). Heatmaps and similarity measures were generated for the selected samples during the beta diversity analysis. We further explored the effect of including contamination BINs, as indicated by the presence of BINs in sample blanks runs, or the effect of BINs with unclear taxonomic resolution on the diversity observed for all sites.

3 | RESULTS

3.1 | General patterns of diversity and similarity across the city of Trondheim

Overall, we detected the presence of 506 BINs identified as belonging to at least 265 species, representing 85 orders and 17 phyla; 9 BINs could not be assigned to any known taxa (Figure 2, File S2). The full list of OTUs with their associated BINs, taxonomy, and habitat associations is in Table S2. Many OTUs were obtained that could also not be associated with a BIN. These OTUs were not considered further in the analysis. We also note that not all data in the BOLD reference library are public; therefore, some taxonomic names associated with a particular BIN were unavailable when searched through BOLD itself and can only be accessed via mBRAVE. Most of the detected diversity consisted of Arthropoda (312 BINs), with Diptera being the most represented order (141 BINs). The least represented taxa (at most 2 BINs per taxon) were Actinobacteria, Bacteroidetes, Chaetognatha, Gastrotricha, Phoronida, Porifera, and Tardigrada. The most common taxon was Candidatus pelagibacter Rappé, Conn, Vergin & Giovannoni, present in 12 out of 15 sites (Table S2).

According to the Jaccard similarity index, the sites were clustered in three groups, Group A: pond Havstein golf course (TRD_1/2), ponds at Sverresborg Museum (TRD_5/6) and Ringve Botanical Garden (TRD_13/14); Group B: Theisendammen (TRD_3/4), Ilabekken at Bleikvollen (TRD_7/8), Madsjøen at IKEA (TRD_22/23), Stokkbekken upstream (TRD_24/25), Stokkbekken downstream (TRD_26/27) and Leangenbekken (TRD_28/29); and Group C: Pirsenteret (TRD_9/10), Korsvika (TRD_11/12), Nidelva at Trondheim Maritime Museum (downstream; TRD_15/16), Nidelva at Fylkesmannsboligen (midstream; TRD_17/18), Nidelva at Sluppen (upstream; TRD_20/21) and Østmarkneset at Ladekaia (TRD_30/31). Both Groups A and C formed one cluster each in the dendrogram based on the values of the Jaccard similarity index between samples (Figure 3), but four samples of Group B clustered together with Group C, separated from the other two samples of Group B. Group A was the most homogeneous and distinct group with Jaccard similarity indices that ranged from 0.12 to 0.16 among samples within group, and 0.009 to 0.093 among samples between groups, followed by Group C (with ranges of 0.096–0.23 and 0.15–0.23 within and between groups, respectively), while in Group B (ranging 0.036–0.26 and 0.023–0.15), some samples showed greater similarity with samples from Group A than with other samples within Group B. Moreover, the sample blanks were different from the rest of the sites, forming a separate, fourth group. The highest number of BINs (130) was detected in Leangenbekken (TRD_28/29) with the lowest number of BINs (27) detected in the upstream site Sluppen of the Nidelva river (TRD_20/21) (Figure 3).
3.2 | Similarity within sites among biological and technical replicates

The similarity between biological and technical replicates for each site was low (Figure 4). On average, only 5.4% of the BINs were shared among all six replicates for each sampling site (two biological replicates and three technical replicates for each of these). The highest percentage of shared BINs (Pirsenteret/TRD_9/10) was 15%, with two cases (Nidelva at Sluppen/TRD_20/21 and Stokkbekken downstream/TRD_26/27), that did not share BINs among replicates. The number of shared BINs was further impacted by whether BINs were represented by only one or two sequences in the full dataset. However, this mostly affected the sharing of BINs in the blanks with the rest of the sampled sites (File S2).

Among the sample blanks, nine BINs were detected (File S2). All of them were present in other samples (Table S2). The number of BINs observed and the number of reads for any BIN was low. When BINs represented by a single read (singleton) or two reads (doubles) were to be removed, only one BIN remained. Thus, we did not consider this as evidence of extensive laboratory contamination during the processing of samples. Instead, it is more suggestive of sequencing errors in the indexes resulting in a small number of reads being misassigned to a sample when samples were demultiplexed in the bioinformatic processing (known as tag jumping). As this error is random and low, it is unlikely to have a major impact on our results of similarity among samples. Thus, no removal of BINs was done prior to further analysis. In addition to many BINs having few representative sequences, we observed taxonomic incongruences of species names with BINs in the BOLD reference database (presented in detail in File S2 and Table S3).

3.3 | The composition of observed diversity across sites—freshwater, marine, brackish, and terrestrial taxa

On the 11 freshwater sites, 396 BINs were detected, whereas the three marine sites yielded 147 BINs, and the only brackish site had 75 BINs detected. Fifty-six BINs were shared between freshwater and marine sites, 46 between brackish and freshwater, and 40 between brackish and marine sites (Figure 5, Table S4). If we consider the species ecology for each BIN, the brackish site included 44 freshwater taxa, 15 marine taxa, and 3 organisms that can be found in both freshwater and marine environments. It should also be highlighted that although just a few marine taxa were detected in freshwater sites, as many as 53 freshwater BINs were detected in the three marine sites (Table S4). Even though we sampled water, 133 BINs detected were terrestrial taxa, mainly belonging to arthropods and annelids (Table S5). Among them, 125 were detected in freshwater sites, 11 were present in marine locations, and 7 in the brackish site (Table S5). Thus, by subtracting these terrestrial taxa, the effective number of aquatic

FIGURE 3  Heatmap generated in mBRAVE for Jaccard’s similarity index among samples where the darker the blue, the greater the similarity is between sampled sites. Clusters indicated with a dendrogram on top. Numbers correspond to the number of Barcode Index Numbers identified within and among sites. The lines and letters correspond to the group clustering. Site names’ colors correspond to those presented in other figures.
The number of BINs detected differed along the Nidelva river. The highest number was 75 BINs detected downstream, at Sjøfartsmuseet (TRD_15/16), whereas 48 BINs were detected in the middle section of the stream, at Fylkesmannsboligen (TRD_17/18), and 27 upstream, at Sluppen (TRD_20-21). The Jaccard similarity index indicated a higher similarity between the downstream and middle parts of the river, compared to the upper part of Nidelva. Out of all BINs detected upstream, nearly half of them were shared between the middle part of the stream and the downstream Nidelva (13 and 14 BINs, respectively, Figure 6a) and the middle part shared 22 BINs with the downstream site. Among the detected BINs, seven BINs from downstream, seven BINs from Nidelva’s middle part, and three BINs from upstream were terrestrial taxa, and thus, trimming the observed diversity effectively to 68, 41, and 24 aquatic BINs, respectively. From the terrestrial habitat, only human DNA was shared among the three sites. Two more BINs (corresponding to the cow, *Bos taurus* Linnaeus, and the red worm, *Eiseniella tetraedra* (Savigny)) were present in both the middle stream and downstream. The rest of the terrestrial taxa were found only in one of the three parts of the river, and therefore, the terrestrial organisms did not have a profound effect on the similarities and the number of shared BINs reported above.

Regarding the stream Stokkbekken (TRD_24-27), 50 BINs were detected downstream and 80 BINs in the upstream site. The Jaccard similarity value reached 0.25, with 26 BINs shared among the sites (Figure 6b). Among the detected taxa, 16 BINs from downstream and 15 BINs from upstream were terrestrial species, and thus, the observed aquatic diversity was reduced to 34 BINs downstream and 65 BINs upstream. Seven terrestrial taxa were shared between the sites, including DNA from human, four earthworms of the family Lumbricidae, one moth, often present at riverbanks, and one duck species. Therefore, both riverine sites shared 22 aquatic organisms. Theisendammen (TRD_3/4) contained 47 BINs, whereas in the stream Ilabekken (TRD_7/8) 67
BINs were detected. The Jaccard similarity value was 0.15, with 14 detected BINs shared between the sites (Figure 6c). Six BINs from Lake Theisedammen and 16 BINs from the stream Ilabekken were terrestrial taxa; therefore, the observed aquatic diversity was 41 and 51 BINs, respectively. Only one BIN, belonging to order Collembola, was shared between these sites. No common roach was detected.

The diversity in the two ponds at Sverresborg Museum (TRD_5/6) and Ringve Botanical Garden (TRD_13/14) was similar, with 36 and 35 BINs detected. Among those, 10 were shared between the ponds (Figure 6d), including DNA from five arthropods, two annelids, two proteobacteria and human, reaching the Jaccard similarity index value of 0.16. Within those BINs, terrestrial taxa were represented only by human DNA in the pond at Sverresborg and 2 BINs in the pond at Ringve Botanical Garden, including DNA from human and one duck species. Common newt (L. vulgaris) was detected in the pond at Sverresborg Museum.

In the single pond Madsjøen (TRD_22/23), 101 BINs were detected, whereas 130 BINs were present in its underground outflow Leangenbekken (TRD_28/29). The Jaccard similarity value reached 0.25, with 48 BINs shared among sites. Among the detected BINs,
58 BINs from pond Madsjøen and 63 BINs from Leangenbekken were terrestrial taxa, thus, the observed diversity of aquatic BINs was 43 and 67 BINs, respectively. A total of 20 terrestrial taxa were detected from both sites. Great-crested newt was not detected in the pond Madsjøen.

4 | DISCUSSION

In this study, we demonstrate how aquatic eDNA can be sampled, sequenced, and analyzed in an educational setting to rapidly infer both terrestrial and aquatic diversity in the city of Trondheim, Norway. From just 30 liters of water sampled by two people in six hours across the city from different habitats, and sequenced within a short time, 435 taxa spanning 17 phyla could be detected from the total number of sequences obtained. Few detailed analyses of BioBlitz have been performed, but if we consider how many species on average were observed (an estimated 431.2 taxa per event) in the United Kingdom between 2006 and 2013 (Postles & Bartlett, 2018), then our biodiversity observations are equivalent. Additionally, because the data collected here were digitized shortly after collection in the form of DNA sequences and available through mBRAVE, as a team of people from 14 different countries many of whom had never generated or analyzed eDNA information, we could collectively engage in learning about the biodiversity in the city of Trondheim from all over the world. Along similar lines, BioBlitz events implementing DNA barcoding, for example, held in Canada, involving participants from multiple countries (Sobel et al., 2017; Telfer et al., 2015). The number of species documented in these surveys largely exceeds those obtained in BioBlitzes solely based solely on morphological identification, with more than 5,500 BINs representing nearly 1,500 putative species. Even though these numbers are also much higher than those obtained in our study, one has to bear in mind the differences both in time invested and sampling strategy. The sampling of our study was completed within 2 days by two collectors, whereas the BioBlitzes performed in Canada involved sampling stretched over multiple months and involving more than a hundred people. When BioBlitz involving DNA barcoding was performed with sampling completed within 2 weeks and much lower number of collectors, it resulted in less than 50 species retained (Laforest et al., 2013). Moreover, the sampling strategy implemented in our study was based on a noninvasive eDNA metabarcoding sampling method, differing from the methodology used in the referenced studies which relies solely on single-specimen DNA barcoding (Laforest et al., 2013; Sobel et al., 2017; Telfer et al., 2015). Thus, we argue that compared to the rapid biodiversity assessment done using single-specimen DNA barcoding as well as morphological identification, our study supports metabarcoding as time-efficient and reliable method, already validated for rapid biodiversity screening (Yu et al., 2012). The addition of nondestructive eDNA sampling makes it highly attractive for biodiversity assessments. This study also shows that the results obtained can be rapidly analyzed on a user-friendly platform by people with virtually no prior bioinformatic experience, which in overall makes eDNA metabarcoding a potentially great tool for educational outreach. Even though participants in BioBlitz events certainly enjoy observing biodiversity itself, there is educational value and excitement in detecting biodiversity that you cannot immediately observe when visiting a locality, as well as learning about biodiversity of places you could never visit.

As our results suggest, using an eDNA rapid biodiversity assessment in urban settings allows many basic questions about biodiversity to be assessed in and across a landscape. Patterns in overall diversity can be compared allowing educators to use findings to inform people and students about biodiversity. For example, the distribution of detected diversity matches known biodiversity patterns between land, marine, and freshwater. It holds that about 80% of species diversity is on land, 15% is in the oceans, and the remaining 5% is in freshwater (Grosberg et al., 2012). Interestingly, even though we sampled water we still had a high proportion of DNA from terrestrial species including vertebrates such as deer and badgers. Recent studies focused on detection of terrestrial vertebrates from water samples show that it is a powerful way to noninvasively detect mobile and elusive vertebrate species (Harper et al., 2019). However, while vertebrate species detections are exciting, most detections were dominated by terrestrial beetles and annelids. The finding of Diptera as the dominant group of arthropods can be ascribed the high diversity of the midge family Chironomidae where most species have aquatic life stages.

Many improvements to sampling could lead to greater species detection. For example, of the sites sampled, 11 were freshwater, three were marine, and one was classified as brackish, yet the number of detected species reflect the expected patterns of diversity, indicating that had a greater sampling effort (e.g., higher volume of water filtered, more sampling sites) occurred in marine and brackish sites, a greater amount of diversity would be detected as expected from the 15% known diversity in these habitats. Additionally, our sampling was performed at one point in the season (mid-May), with low activity and abundance of some invertebrate groups at this high latitude (63° North). Thus, a greater number of taxa would likely be recorded had our sampling been done at several occasions throughout the seasons, due to differences in the life histories of present taxa. We also acknowledge the observation that we had a very low overlap in diversity among replicates (either biological or technical). Thus, to estimate total diversity even at a site, a greater number of samples from each site or greater sequencing depth could have been performed to increase the likelihood of detections.

In addition to the terrestrial species DNA being sampled from water, there was also evidence of DNA moving across the landscape. Examples include DNA from the lentic opossum shrimp (Mysis relicta Loven) at all three sites in river Nidelva, DNA from the lotic mayfly Baetis rhodani Pictet in the lake Theisendammen as well as lotic caddisfly Hydropsyche siltalai Doehler in the Havstein pond, and about 20 nonbiting midge taxa in the marine sites. Even though the transport of flying insects’ DNA between the sites can be easily explained, the presence of lentic mysid in the Nidelva river is surprising.
Most likely, it can be associated with passive transport via man-made pipes distributing water from the Lake Selbusjøen, where it is known to occur, to the nearby power plant, which then releases the water to upstream Nidelva.

Movement of DNA by the movement of water allows eDNA to measure biodiversity for scales larger than a single sampling site (Deiner et al., 2016). In this study, the sites sampled along the Nidelva river showed higher diversity downstream and a greater number of species were shared the further downstream the site was on the river, suggesting eDNA accumulation. However, one should bear in mind that transport is not necessarily confirmed with these patterns because the species detected have ecologies that would predict their occurrence in both sites along the river and can often be found present at higher densities downstream due to invertebrate drift (Grossman et al., 2010). The terrestrial and freshwater species DNA found in marine sites are more convincing examples of DNA being moved across the landscape. This inherent trait of eDNA moving away from its source is exciting when large areas need documenting and useful for BioBlitz events, but can also be troublesome when specific locations for a species are desired (e.g., as might be the case for an invasive species).

Among the three species of concern (two newts and the common roach), we detected the common newt (*L. vulgaris*) in the pond at Sverresborg cultural museum. The pond itself was restored in 2013–2014 as part of the open air exhibition, and care was taken at Sverresborg cultural museum. The pond itself was restored in 2013–2014 as part of the open air exhibition, and care was taken to preserve the known newt population (https://sverresborg.no/beskrivelse-av-dammen-og-parken). It is nice to see that this was confirmed by analysis of DNA being moved across the landscape. This inherent trait of eDNA moving away from its source is exciting when large areas need documenting and useful for BioBlitz events, but can also be troublesome when specific locations for a species are desired (e.g., as might be the case for an invasive species).

In this survey, we highlight mBRAVE, an open-access computing platform with an interactive interface that performs metabarcoding bioinformatic processing. This platform allows parallel sequence identification for the COI barcoding region by comparison to the Barcode of Life Database for millions of sequences simultaneously. The significance of the mBRAVE platform is that it allows archiving, sharing, and exploration of complex genetic data through visualizations and tools that require no computer science background and can be conducted online. It is noted that further diversity analyses are possible outside the mBRAVE environment by downloading the data and using it elsewhere, but no such external analyses were performed for the purpose of this study.

### 5 | CONCLUSIONS

Our goal was to demonstrate the use of DNA-based detection methods that can be used in educational settings to explore and raise awareness of urban biodiversity. Thus, we propose our study as an example of an entirely new form of a BioBlitz, involving a noninvasive eDNA sampling, next-generation sequencing, and rapid data analyses using a freely accessible, user-friendly online platform. We also suggest improvements for how to implement the field and laboratory protocols, although the latter can be done in collaboration with a local university or company. Moreover, while these studies can be carried out in the absence of direct interaction with taxonomic experts thanks to the digitization of current taxonomic knowledge in the Barcode of Life Database, it is still highly encouraged to collaborate with them. Indeed, even though these DNA-based methods are exciting and have proven efficient, they are still heavily reliant on taxonomic identifications provided by curated reference databases and the accuracy in determining species based on DNA marker divergence. Although much work is still needed to perfect
our abilities in DNA-based detection of life, educational surveys of this type remain useful and exciting explorations for building an engaged public interested in reversing the loss of life from our planet.

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CONFLICT OF INTEREST
None declared.

AUTHOR CONTRIBUTIONS
Sampling, laboratory work and study design: MM, TE. Data analyses: KH, MM, MVC, LS, DM, VI, DAL, SL, FCI, ZC-Z, ISM, AD, LT, KM, DHZA, GK, X-LL, BP, TE, KD. Supervision: KH, KD. Manuscript preparation: all authors edited and commented on the manuscript.

DATA AVAILABILITY STATEMENT
The data are publicly available within the mBRAVE environment (Project Code: MBR-WRMISEQPE), and in the European Nucleotide Archive (ENA) sequence read archive (PRJEB37589).

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Additional supporting information may be found online in the Supporting Information section.