MinION barcodes: biodiversity discovery and identification by everyone, for everyone

Amrita Srivathsan¹, Leshon Lee¹, Kazutaka Katoh²,³, Emily Hartop⁴,⁵, Sujatha Narayanan Kutty¹,⁶, Johnathan Wong¹, Darren Yeo¹, Rudolf Meier¹

¹ Department of Biological Sciences, National University of Singapore, Singapore
² Research Institute for Microbial Diseases, Osaka University, Japan
³ Artificial Intelligence Research Center, AIST, Tokyo, Japan
⁴ Zoology Department, Stockholms Universitet, Stockholm, Sweden
⁵ Station Linné, Öland, Sweden
⁶ Tropical Marine Science Institute, National University of Singapore, Singapore
Abstract

DNA barcodes are a useful tool for discovering, understanding, and monitoring biodiversity which are critical at a time when biodiversity loss is a major problem for many countries. However, widespread adoption of barcodes requires cost-effective and simple barcoding methods. We here present a workflow that satisfies these conditions. It was developed via “innovation through subtraction” and thus requires minimal lab equipment, can be learned within days, reduces the barcode sequencing cost to <10 cents, and allows fast turnaround from specimen to sequence by using the real-time sequencer MinION. We first describe cost-effective and rapid procedures for obtaining tagged amplicons. We then demonstrate how a portable MinION device can be used for real-time sequencing of tagged amplicons in many settings (field stations, biodiversity labs, citizen science labs, schools). Small projects can use the flow cell dongle (“Flongle”) while large projects can rely on MinION flow cells that can be stopped and re-used after collecting sufficient data for a given project. We also provide amplicon coverage recommendations that are based on several runs of MinION flow cells (R10.3) which suggest that each run can generate >10,000 barcodes. Next, we present a novel software, ONTbarcoder, which overcomes the bioinformatics challenges posed by the sequencing errors of MinION reads. This software is compatible with Windows10, Macintosh, and Linux, has a graphical user interface (GUI), and can generate thousands of barcodes on a standard laptop within hours based on two input files (FASTQ, demultiplexing file). We document that MinION barcodes are virtually identical to Sanger and Illumina barcodes for the same specimens (>99.99%). Lastly, we demonstrate how rapidly MinION data have improved by comparing the performance of sequential flow cell generations. We overall assert that barcoding with MinION is the way forward for government agencies, universities, museums, and schools because it combines low consumable and capital cost with scalability. Biodiversity loss is threatening the planet and the use of MinION barcodes will help with enabling an army of researchers and citizen scientists, which is necessary for effective biodiversity discovery and monitoring.
1. Background

DNA sequences have been used for identification and taxonomic purposes for decades (Hebert, Cywinska et al. 2003, Tautz, Arctander et al. 2003, Meier 2008), but for most of this time been akin to mobile phones in the 1990s: of limited value due to sparse signal coverage and high cost. Obtaining barcodes was problematic due largely to the complicated and expensive procedures on which it relied. Some of these problems have since been addressed by, for example, developing effective DNA extraction protocols and optimizing Sanger sequencing procedures (Ivanova, Dewaard et al. 2006, Ivanova, Borisenko et al. 2009). These improvements enabled the establishment of a centralized barcoding facility in 2006. After 15 years and the investment of >200 million USD, ca. 8 million animal barcodes are available for searches on BOLD Systems, but only ca. 6 million are in the public domain (http://boldsystems.org/index.php/IDS_OpenIdEngine). Combined with barcodes from NCBI GenBank, they are now a valuable resource to the global biodiversity community. However, the cost of barcodes has remained high (http://ccdb.ca/pricing/) and the prevalent approach for sizeable projects is sending specimens from all over the world to one center and then only some back to the country of origin. This interferes with real-time biodiversity monitoring and specimen accessibility. We therefore argue that access to barcodes has to be democratized through decentralization. We here show that this achievable because the application of a technique that is known as “innovation through subtraction” in engineering readily yields simplified and cost-effective solutions for DNA barcoding and the amplification and sequencing of a short mitochondrial COI fragment can be efficiently performed anywhere and by biologists and citizen scientists alike.

A decentralized model for monitoring the world’s biodiversity is necessary given the scale, urgency, and importance of the task at hand. For example, even if there were only 10 million species of metazoan animals on the planet (Stork, McBroom et al. 2015) and a new species is discovered with every 50th specimen that is processed, species discovery with barcodes will require the sequencing of 500 million specimens (Yeo, Srivathsan et al. 2020). Yet,
species discovery is only a small part of the biodiversity challenge in the 21st century. Biodiversity loss is now considered by the World Economic Forum as one of the top three global risks based on likelihood and impact for the next 10 years (World Economic Forum 2020) and Swiss Re estimates that 20% of all countries face ecosystem collapse as biodiversity declines (Swiss Re 2020). Biodiversity loss is no longer just an academic concern; it is now a major threat to human communities and the health of the planet. This also implies that biodiversity discovery and monitoring require completely different scales than in the past. The old approaches thus need rethinking because all countries need real-time distributional and abundance information to develop effective conservation strategies and policies. In addition, they need information on how species interact with each other and the environment (Abrego, Roslin et al. 2021). Many of these biodiversity monitoring and environmental management activities have to focus on terrestrial invertebrates, whose biomass surpasses that of all terrestrial vertebrates combined (Bar-On, Phillips et al. 2018) and who occupy a broad range of ecological guilds. The main obstacles are high numbers of specimens and species and the rapid decline of many of these taxa (Bell, Blumgart et al. 2020, Eisenhauer, Bonn et al. 2019, Hallman, Sorg et al. 2017, Hallman, Ssymank et al. 2021, Stepanian, Entrekin et al. 2020, Wagner, Grames et al. 2021) which means that monitoring should be locally conducted to allow for rapid turnaround. This requires simple and cost-effective procedures that can be implemented anywhere by stakeholders with very different scientific and skill backgrounds.

DNA barcoding was proposed at a time when biodiversity loss was not on the radar of economists. Instead, barcodes were initially intended as an identification tool for biologists (Hebert, Cywinska et al. 2003). Thus, most projects focused on taxa with a large following in biology (e.g., birds, fish, butterflies) (Kwong, Srivathsan et al. 2012). However, this also meant that these projects only covered a small proportion of the terrestrial animal biomass (Bar-On, Phillips et al. 2018) and species-level diversity (Groombridge 1992). Yet, despite targeting taxa with well-understood diversity, the projects struggled with covering >75% of
the described species in these groups (Kwong, Srivathsan et al. 2012). When the pilot barcoding projects ran out of material from identified specimens, they started targeting unidentified specimens; i.e., DNA barcoding morphed into a technique that was used for biodiversity discovery (“dark taxa”: Page 2011, Kwong, Srivathsan et al. 2012). This shift towards biodiversity discovery was gradual and incomplete because the projects used a “hybrid approach” that started with subsampling or sorting specimens to “morphospecies” before barcoding representatives of each morphospecies/sample (e.g., Barrett and Hebert 2005, Hendrich, Pons et al. 2010, Hebert, DeWaard et al. 2013, Ng’endo, Osiemo et al. 2013, Hebert, Ratnasingham et al. 2016, Thormann, Ahrens et al. 2016, Knox, Hogg et al. 2020). This is problematic, as morphospecies sorting is known to be labour-intensive and of unpredictable quality because it is heavily dependent on the taxonomic expertise of the sorters (Krell 2004, Stribling, Pavlik et al. 2008). Thus, such hybrid approaches are of limited value for obtaining reliable quantitative data on biodiversity, but were adopted as a compromise owing to the prohibitive cost of barcoding. The logical alternative is to barcode all specimens and then group them into putative species based on sequence information. Such a “reverse workflow” (Wang, Srivathsan et al. 2018), where every specimen is barcoded as the initial pre-sorting step, yields quantitative data and corroborated species-level units. However, the reverse workflow requires efficient and low-cost barcoding methods that are also suitable for biodiverse countries with limited science funding.

Fortunately, such cost-effective barcoding methods are now becoming available. This is partially due to the replacement of Sanger sequencing with second- and third-generation sequencing technologies that have lowered sequencing costs dramatically (Shokralla, Spall et al. 2012, Shokralla, Porter et al. 2015, Meier, Wong et al. 2016, Hebert, Braukmann et al. 2018, Krehenwinkel, Kennedy et al. 2018, Srivathsan, Baloglu et al. 2018, Wang, Srivathsan et al. 2018, Srivathsan, Hartop et al. 2019, Yeo, Srivathsan et al. 2020). Such changes mean that the reverse workflow is now available for tackling the species-level diversity of those metazoan clades that are so specimen- and species-rich that they have been neglected in
the past (Ponder and Lunney 1999, Srivathsan, Hartop et al. 2019). Many of these clades have high spatial species turnover, requiring many localities in each country to be sampled and massive numbers of specimens to be processed (Yeo, Srivathsan et al. 2020). Such intensive processing is best achieved close to the collecting locality to avoid the unnecessary risks, delays and costs from shipping biodiversity samples across continents. This is now feasible because biodiversity discovery can be readily pursued in decentralized facilities at varied scales. Indeed, accelerated biodiversity discovery is a rare example of a big science initiative that allows for meaningful engagement of students and citizen scientists and can in turn significantly enhance biodiversity education and appreciation (Pomerantz, Peñafiel et al. 2018, Watsa, Erkenswick et al. 2020). This is especially so when stakeholders not only barcode, but also image specimens, determine species abundances, and map distributions of newly discovered species. All of which can be based on specimens collected in their own backyard.

But can such decentralized biodiversity discovery really be effective? Within the last five years, the students and interns in the laboratory of the corresponding author at the National University of Singapore barcoded >330,000 specimens. After analyzing the first >140,000 barcoded specimens for selected taxa representing different ecological guilds, the alpha and beta diversity of Singapore’s arthropod fauna was analyzed based on ~8,000 putative species which revealed that some habitats were unexpectedly species-rich and harboured very unique faunas (e.g., mangroves, freshwater swamp: Yeo, Srivathsan et al. 2020; Baloğlu, Clews et al. 2018). Barcodes even helped with the conservation of charismatic taxa when they were used to identify the larval habitats for more than half of Singapore’s damsel- and dragonfly species (Yeo, Puniamoorthy et al. 2018) and facilitated species interaction research and biodiversity surveys based on eDNA (Lim, Tay et al. 2016, Srivathsan, Nagarajan et al. 2019). Biodiversity appreciation by the public was fostered by featuring newly discovered species and their species interactions on “Biodiversity of Singapore” (BOS >15,000 species: https://singapore.biodiversity.online/) and dozens of new species have

2. Methods for the democratization of DNA barcoding through simplification

Barcoding a metazoan specimen requires the successful completion of three steps: (1) obtaining DNA template, (2) amplifying COI via PCR, and (3) sequencing the COI amplicon. Many biologists learn these techniques in university for a range of different genes – from those that are easy to amplify (short fragments of ribosomal and mitochondrial genes with well-established primers) to those are difficult (long, single-copy nuclear genes with few known primers). Fortunately, amplification of short mitochondrial markers like COI does not require the same level of care as nuclear markers. Learning how to barcode efficiently is hence an exercise of unlearning by applying “innovation through subtraction”. Note that this unlearning is of critical importance for the democratization of biodiversity discovery with DNA barcodes and is particularly vital for boosting biodiversity research where it is most needed: in biodiverse countries with limited science funding.

In this section, we first briefly summarize commonly used procedures for DNA extraction, PCR, and sequencing. For each step we then describe how the procedures can be simplified. In addition to the description, we provide videos for the described procedures which are available from the YouTube channel “Integrative Biodiversity Discovery” (https://www.youtube.com/channel/UC1WowokomhQJRc71FmsUAcg). Note that all techniques have been extensively tested in our lab, primarily on invertebrates preserved in ethanol. Regarding sequencing, we briefly introduce four methods, but our focus is on MinION sequencing because this device is particularly suitable as the default sequencing option for decentralized biodiversity discovery.
Methods for step 1: Obtaining DNA template

Most biologists learn that DNA extraction requires tissue digestion, DNA purification, and DNA elution. This approach is slow and expensive because it frequently involves kits and consumables that are designed for obtaining the kind of high-quality DNA that is needed for amplifying “difficult” genes (e.g., long, single-copy nuclear markers). However COI is a mitochondrial gene and thus naturally enriched. Indeed, the tiny mitochondrial genome (16 kbp) usually contributes 0.5-5% of the DNA in a genomic extraction (Arribas, Andújar et al. 2016, Crampton-Platt, Yu et al. 2016). Furthermore, barcoding requires only the amplification of one short marker (<700 bp) so that not much DNA template is needed. This allows for using the following simplified procedures that are designed for specimens containing DNA template of reasonable quality (e.g., Malaise trap specimens collected within the last 20 years).

Simplified DNA “extraction”: Obtaining template for DNA barcoding need not take more than 20 minutes, does not require DNA purification, and costs essentially nothing. The cheapest, but not necessarily fastest, method is “directPCR”; i.e., deliberately “contaminating” a PCR reaction with the DNA of the target organism by adding the entire specimen or a tissue sample into the PCR reagent mix (Wong, Tay et al. 2014). This method is very fast and effective for small specimens lacking thick cuticle or skin (Wong, Tay et al. 2014) and works particularly well for many abundant aquatic invertebrates such as chironomid midges and larvae. Larger specimens require the use of body parts [leg or antenna: Wong, Tay et al. (2014)]. Such dissections tend to be labour-intensive if large numbers of specimens must be processed, but it is a good method for small numbers of samples or in barcoding experiments that are carried out in poorly equipped labs. Note that the whole body or body part that is used for directPCR can be recovered after amplification, although soft-bodied animals may become transparent.
An alternative to directPCR is buffer-based DNA extraction. This method is also essentially cost-free because it involves alkaline buffers that are inexpensive, usually available in molecular labs (e.g., PBS), or can be prepared easily (HotSHOT) (Truett, Heeger et al. 2000, Thongjued, Chotigeat et al. 2019)). Our preferred method is extraction with HotSHOT, which we have used for barcoding >50,000 arthropods (Yeo, Srivathsan et al. 2020). We use 10-15 μL HotSHOT per specimen. Small specimens are submerged within the well of a microplate while larger specimens are placed head-first into the well. The tissue need not be entirely submerged in HotSHOT. DNA is obtained within 20 minutes in a thermocycler via two heating steps (Truett, Heeger et al. 2000). After neutralization, >20 μl of template is available for amplifying COI and the voucher can be recovered. Note that HotSHOT extraction leaves most of the DNA in the specimen untouched and more high quality DNA can subsequently be extracted from the same specimen. An alternative to obtaining DNA via lab buffers is the use of commercial DNA extraction buffers (Kranzfelder, Ekrem et al. 2016). These buffers have a longer shelf life, and are good alternatives for users who only occasionally barcode moderate numbers of specimens. In the past, we have used QuickExtract (Srivathsan, Hartop et al. 2019) and found that 10 μl is sufficient for obtaining DNA template from most insect specimens. In summary, obtaining DNA templates for barcoding is fast and straightforward and most published barcoding studies greatly overcomplicate this step. It should be noted however, that all DNA extraction methods require the removal of excess ethanol from specimens prior to extraction (e.g., by placing the specimen on tissue paper or replacing ethanol with water prior to specimen processing) and that the DNA extracts obtained with such methods should be stored at -20°C and be used within days.

Methods for step 2: amplifying COI via PCR. Most PCR recipes and reagents are optimized to work for a wide variety of genes and not just for a gene like the COI barcode that is naturally enriched, has a large number of known primers, and is fairly short. Standard PCR recipes can therefore be simplified. However, the use of “modern” sequencing technologies such as Illumina, PacBio, or Oxford Nanopore Technologies introduces one complication:
The amplicons have to be “tagged” (or “indexed”/“barcoded”). This is necessary because pools of amplicons are sequenced simultaneously instead of processing one amplicon at a time (as in Sanger sequencing). Tags are specimen identifiers consisting of short DNA sequences at the 5’ ends of the amplicons. They allow for the assignment of each sequence read obtained during sequencing to a specific specimen in the “demultiplexing” bioinformatics step. Numerous tagging techniques have been described in the literature, but most are too complicated for efficient DNA barcoding.

Simplified techniques for obtaining tagged amplicons

Published protocols tend to have five issues that increase workload and/or inflate cost, while a fifth issue only affects amplicon tagging:

- **Issue 1: expensive polymerases or master mixes.**
  These often utilize high-fidelity polymerases that are designed for amplifying low copy-number nuclear genes based on low-concentration template but rarely make a difference when amplifying COI. Indeed, even home-made polymerases can be used for barcoding. This is important because high import taxes for consumables interfere with biodiversity discovery in many biodiverse countries.

- **Issue 2: indiscriminate use of single-use consumables.**
  Disposable products increase costs and damage the environment. Most biodiversity samples are obtained under “unclean conditions” that create numerous opportunities for cross-specimen contamination long before specimens reach the lab (e.g., thousands of specimens rubbing against each other in sample containers and in the same preservation fluid). Yet numerous studies have shown that the DNA from specimens exposed to such conditions will usually outcompete contaminant DNA that is likely to occur at much lower concentrations. Similarly, the probability that a washed/flushed and autoclaved microplates or pipette tips retain enough viable contaminant DNA to successfully outcompete the template DNA is extremely low. Indeed, we have repeatedly tried and failed to amplify COI using reused plastic consumables and water as template.
That it is safe to reuse some consumables is again good news for biodiversity discovery under severe financial constraints. Note, however, that we do not recommend the re-use of consumables for handling stock chemicals such as primers and sequencing reagents.

- **Issue 3: large PCR volumes (25-50 μl).**

  Pools of tagged amplicons comprise hundreds or thousands of products and there is typically more than enough DNA for preparing a library. Accordingly, even small PCR volumes of 10-15 μl are sufficient, thereby reducing consumable costs for PCR to nearly half when compared to standard volumes of 25-50 μl.

- **Issue 4: using gel electrophoresis for checking amplification success of each PCR product.**

  This time-consuming step is only justified when Sanger sequencing is used or when high-priority specimens are barcoded. It is not necessary when barcoding large numbers of specimens with modern sequencing technologies, because failed amplicons do not add to the sequencing cost. Furthermore, specimens that failed to yield barcodes during the first sequencing run can be re-sequenced or re-amplified and then added to subsequent sequencing runs (Srivathsan, Hartop et al. 2019). We thus only use gel electrophoresis to check a small number of reactions per microplate (N=8-12, including the negative control) in order to make sure that there was no plate-wide failure.

The fifth issue requires more elaboration and concerns how to efficiently tag amplicons. We tag via a single PCR reaction (Meier, Wong et al. 2016) using primers including the tag at the 5' end because it is simpler than the dual-PCR tagging strategy dominating the literature. The latter has numerous disadvantages when applied to one gene: it doubles the cost by requiring two rounds of PCR, is more labour intensive, increases the risk for PCR errors by requiring more cycles, and requires clean-up of every PCR product after the first round of amplification. In contrast, tagging via a single PCR is simple and costs the same as any gene amplification. It is here described for a microplate with 96 templates, but the protocol can be adapted to the use of strip tubes or half-plates. What is needed is a 96-well primer
plate where each well contains a differently tagged reverse primer. This “primer plate” can
yield 96 unique combinations of primers once the 96 reverse primers are combined with the
one forward primer (f-primer x 96 differently tagged r-primers = 96 unique combinations).
This also means that if one purchases 105 differently tagged forward primers, one can
individually tag 10,800 specimens (105 x 96 = 10,800 amplicons). This is the number of
amplicons that we consider appropriate for a MinION flow cell (R10.3; see below).

Assigning tag combinations is also straightforward. For each plate with 96 PCR reactions,
add the same f-primer to a tube with the PCR master mix (Taq DNA polymerase, buffer and
dNTPs) for the plate. Then dispense the “f-primed” master mix into the 96-wells. Afterwards,
use a multichannel pipette to add the DNA template and the tagged r-primers from the r-
primer plate into the PCR plate. All 96 samples in the plate now have a unique combination
of tagged primers because they only share the same tagged forward primer. This makes the
tracking of tag combinations simple because each PCR plate has its own tagged f-primer,
while the r-primer is consistently tied to well position. Each plate has a negative control to
ensure that no widespread contamination has occurred. The tagging information for each
plate is recorded in the demultiplexing file that is later used to demultiplex the reads obtained
during sequencing.

Some users may worry that the purchase of many primers is expensive, but one must keep
in mind that the amount of primer used in a PCR reaction is constant. Therefore, single
PCR-tagging only means a greater upfront investment. Ordering all primers at once,
however, does mean that one must be much more careful about avoiding primer
degeneration and contamination as the stock will last longer. Primer stock should be stored
at -80°C and the number of freeze-thaw cycles should be kept low (<10). This means that
upon receipt of the primer stock, it should be immediately aliquoted into plates/tubes holding
only enough primer for rapid use. For fieldwork, one should only bring enough dissolved
primer for the necessary experiments, or rely on lyophilised reagents.
The choice of tag length is determined by three factors. Longer tags reduce PCR success rates (Srivathsan, Hartop et al. 2019) while they increase the proportion of reads that can be assigned to a specific specimen (demultiplexing rate). Designing tags is not straightforward because they must remain sufficiently distinct (>4bp from each other including insertions/deletions) while avoiding homopolymers. We include a list of 13 bp tags that are suitable in supplementary materials.

Methods for step 3: Amplicon sequencing. The use of the PCR techniques described so far should keep the cost for a tagged barcode amplicon to 0.05-0.10 USD as long as the user buys cost-effective consumables. What comes next is the purification of the amplicons via the removal of unused PCR reagents, the adjustment of DNA concentration, and sequencing. Sequencing can be done with Sanger sequencing, Oxford Nanopore Technologies (ONT) (e.g., MinION: Srivathsan, Hartop et al. 2019), Illumina (Wang, Srivathsan et al. 2018), or PacBio (e.g., Sequel: Hebert, Braukmann et al. 2018). Users select the sequencing option that best suit their needs based on five major criteria: (1) Scaling; i.e., ability to accommodate projects of different sizes, (2) turnaround times, (3) cost, (4) amplicon length and (5) sequencing error rate. Sanger sequencing has fast turnaround times but high sequencing cost per amplicon ($3-4 USD). This is the only method where cost scales linearly with the number of amplicons that need sequencing, while the other sequencing techniques are fundamentally different in that each run has two fixed costs that stay the same regardless of whether only a few or the maximum number of amplicons for the respective flow cells are sequenced. The fixed costs are library preparation (preparing amplicons for sequencing) and flow cell.

The MinION Flongle has the lowest fixed costs (library and flow cell: ca. $120 USD for a routine run) and we show here that it has sufficient capacity for ca. 250 barcodes. The adapter for flongle can be purchased as part of a package that costs $1460 USD, which also
includes 12 flow cells thus increasing the cost of the first 12 runs to ~$170 USD. The turnaround time is fast, so the MinION Flongle is arguably the best sequencing option for small barcoding projects with > 50 barcodes. Full MinION flow cells also have fast turnaround times, but the minimum run cost is ca. 1000 USD, so this option only becomes more cost-effective than Flongle when >1800 amplicons are sequenced. As shown later, one regular MinION flow cell can comfortably sequence 10,000 amplicons. This is a similar volume to what has been described for PacBio (Sequel) (Hebert, Braukmann et al. 2018), but the high instrument cost for PacBio means that sequencing usually has to be outsourced, leading to longer wait times. By far the most cost-effective sequencing method for barcodes is Illumina’s NovaSeq sequencing. The fixed costs for library and lanes are high (3000-4000 USD), but each flow cell yields 800 million reads which can comfortably sequence 800,000 barcodes at a cost of < $0.01 USD per barcode. This high capacity means that the 6 million publicly available barcodes in BOLD Systems could have been sequenced on just <8 NovaSeq flow cells for ~50,000 USD. However, Illumina sequencing can only be used for mini-barcodes of up to 420 bp length (using 250bp PE sequencing using SP flow cell). “Full-length” COI barcode (658 bp) can only be obtained by sequencing two amplicons. Note that while Illumina barcodes are shorter than “full-length” barcodes, there is no evidence that mini-barcodes have a negative impact on species delimitation or identification as long as the mini-barcode is >250 bp in length (Yeo, Srivathsan et al. 2020).

Simplified techniques for sequencing tagged amplicons: Modern sequencing technologies are used to sequence amplicon pools instead of individual amplicons. To obtain such a pool, it is sufficient to combine only 1 μl per PCR product. The pool can be cleaned using several PCR clean-up methods. We generally use SPRI bead-based clean-up, with Ampure (Beckman Coulter) beads but Kapa beads (Roche) or the more cost-effective Sera-Mag beads (GE Healthcare Life Sciences) in PEG (Rohland and Reich 2012) are also viable options (Srivathsan, Hartop et al. 2019). We recommend the use of a 0.5X ratio for Ampure beads for barcodes longer than 300 bp since it removes a larger proportion of primers and
primer dimers. However, this ratio is only suitable if yield is not a concern (e.g., pools consisting of many and/or high concentration amplicons). Increasing the ratio to 0.7-1X will improve yield but render the clean-up less effective. Amplicon pools containing large numbers of amplicons usually require multiple rounds of clean-up, but only a small subset of the entire pool needs to be purified because most library preparation kits require only small amounts of DNA. Note that the success of the clean-up procedures should be verified with gel electrophoresis, which should yield only one strong band of expected length. After the clean-up, the pooled DNA concentration is measured in order to use an appropriate amount of DNA for library preparation. Most laboratories use a Qubit, but less precise techniques are probably also suitable.

Obtaining a cleaned amplicon pool according to the outlined protocol is not time consuming. However, many studies retain “old Sanger sequencing habits”. For example, they use gel electrophoresis for each PCR reaction to test whether an amplicon has been obtained and then clean and measure all amplicons one at a time for normalization (often with very expensive techniques: Ampure beads: (Maestri, Cosetino et al. 2019); TapeStation, BioAnalyzer, Qubit: (Seah, Lim et al. 2020)). This is presumably done to obtain a pool of amplicons where each has equal representation. However, reads are cheap while individual clean-ups and measurements for each PCR product are expensive. Furthermore, weak products that failed to yield a barcode can be re-sequenced (Srivathsan, Hartop et al. 2019).

Another strategy is normalization based on the band strength of a few PCR products per plate as determined by gel electrophoresis. We used this strategy in the current study, but have since determined that pooling without such normalization yields nearly identical success rates based on the same number of reads (>9600 amplicons: 76.8% vs. 77.2%).

3. MinION barcoding with new flow cells (R10.3, Flongle) and high-accuracy basecalling
Oxford Nanopore Technologies (ONT) instruments sequence DNA by passing single-stranded DNA through a nanopore. This creates current fluctuations which can be measured and translated into a DNA sequence via basecalling (Wick 2019). The sequencing devices are small and inexpensive, but the read accuracy is only moderate (85% - 95%) (Wick 2019, Silvestre-Ryan and Holmes 2021). This means that data analysis requires specialized bioinformatics pipelines. The nanopores used for sequencing are arranged on flow cells, with new flow cell chemistries and basecalling softwares regularly released. Recently, three significant changes occurred. Firstly, ONT released a cheap flow cell (Flongle) that only has 126 pores (126 channels) instead of the customary 2048 pores (512 channels) of a full MinION flow cell. We were interested in Flongle because it looked promising for small barcoding projects that needed quick turnaround times. Secondly, ONT released a new flow cell chemistry for full flow cells (R10.3) where the nanopores have a dual instead of a single reader-head. Dual reading has altered the read error profile by giving better resolution to homopolymers and improving consensus accuracy (Chang, Ip et al. 2020, Vereecke, Bokma et al. 2020). Lastly, ONT released high accuracy (HAC) basecalling. We thus obtained amplicons using techniques described in Section 2 and processed them further as described below.

Library preparation. Library preparation was based on 200 ng of DNA for the full MinION flow cells and 100 ng for the Flongle and used ligation-based kits (see Table 1 for details). We generally followed kit instructions, but excluded the FFPE DNA repair mix in the end-repair reaction, as this is mostly needed for formalin-fixed, paraffin-embedded samples. The reaction volumes for the R10.3 flow cell libraries consisted of 45 μl of DNA, 7 μl of Ultra II End-prep reaction buffer (New England Biolabs), 3 μl of Ultra II End Prep enzyme mix (New England Biolabs) and 5 μl of molecular grade water. For the Flongle, only half of the reagents were used to obtain a total volume of 30 μl. We further modified the Ampure ratio to 1x for all steps as DNA barcodes are short whereas the recommended ratio in the manual is for longer DNA fragments. The libraries were loaded and sequenced with a MinION Mk
1B. Data capture involved a MinIT or a Macintosh computer that meets the IT specifications recommended by ONT. The bases were called using Guppy (versions provided in Table 2), under the high-accuracy model in MinIT taking advantage of its GPU.

Sequencing. Six amplicon pools were sequenced (Table 1). For two of the pools, Mixed Diptera (N=511) and Afrotropical (N=4,275) Phoridae, we had comparison barcodes that were obtained with Sanger and Illumina sequencing and the same amplicon pools were previously sequenced with earlier versions of MinION flow cells (Srivathsan, Baloglu et al. 2018, Srivathsan, Hartop et al. 2019) (Table 1). These two pools were used to assess the accuracy of barcodes generated using R10.3 flow cells. Two additional datasets tested the capacity of R10.3 flowcells for mini- and full length barcodes for the same specimens (Palaearctic Phoridae, 658 and 313 bp for ca. 9,930 specimens). The Mixed Diptera Subsample and Chironomidae datasets test the performance of the Flongle. The Mixed Diptera Subsample (N=257) is a subset of the Mixed Diptera (N=511) amplicon pool for which we have Sanger barcodes for comparison. The Chironomidae dataset contains sequences for 313 bp mini-barcodes for 191 specimens of Chironomidae that were newly amplified for this study.
Table 1. Datasets used in the study and the corresponding experimental details.

<table>
<thead>
<tr>
<th>Dataset Name</th>
<th>Number of specimens</th>
<th>Fragment size, primer information</th>
<th>Extraction/PCR setup</th>
<th>PCR cleanup</th>
<th>ONT Library Preparation kit/Flow cell used</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>R10.3 Datasets</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed Diptera (see Srivathsan et al., 2018) - Sanger barcodes available</td>
<td>511 (257 mixed Diptera, 254 Dolichopodidae) 17 negatives</td>
<td>658 bp HCO2198, LCO1490 (Folmer et al., 1994)</td>
<td>Extraction Method: QuickExtract PCR Mix: Total volume: 20 μl 10 x buffer: 2 μl dNTPs (2.5 mM): 1.5 μl Taq polymerase: 0.2 μl BSA (1 mg/ml): 2 μl Primer (5 μM): 2 μl each DNA: 2 μl</td>
<td>Ampure beads (Beckman Coulter)</td>
<td>SQK-LSK110/FLO-MIN111</td>
</tr>
<tr>
<td>Afrotropical Phoridae (see Srivathsan et al., 2019) - Illumina mini-barcodes available</td>
<td>4275 (Phoridae) 45 negatives</td>
<td>658 bp HCO2198, LCO1490 (Folmer et al., 1994)</td>
<td>Extraction Method: QuickExtract PCR Mix: Total volume: 15.16 μl Mastermix (CWBio): 10 μl 25mM MgCl2: 0.16 μl BSA (1 mg/ml): 2 μl Primer (10μM): 1 μl each DNA: 1 μl</td>
<td>Sera-Mag beads (GE Healthcare Life Sciences) in PEG</td>
<td>SQK-LSK109/FLO-MIN111</td>
</tr>
<tr>
<td>Palaearctic Phoridae (658)</td>
<td>9,929 (Phoridae) 105 negatives</td>
<td>658 bp jgHCO2198, jgLCO1490 (Folmer et al., 1994, Geller et al. 2013)</td>
<td>Extraction Method: HotSHOT PCR Mix: Total volume: 16 μl Mastermix (CWBio): 7 μl BSA (1 mg/ml): 1 μl Primer (10μM): 1 μl each DNA: 6 μl</td>
<td>Ampure beads (Beckman Coulter)</td>
<td>SQK-LSK110/FLO-MIN111</td>
</tr>
<tr>
<td>Palaearctic Phoridae (313)</td>
<td>9,932 (Phoridae) 106 negatives</td>
<td>313 bp m1COlIntF, jgHCO2198 (Leray et al. 2013, Geller et al. 2013)</td>
<td>Extraction Method: HotSHOT PCR Mix: Total volume: 14 μl Mastermix (CWBio): 7 μl BSA (1 mg/ml): 1 μl Primer (10μM): 1 μl each DNA: 4 μl</td>
<td>Ampure beads (Beckman Coulter)</td>
<td>SQK-LSK110/FLO-MIN111</td>
</tr>
<tr>
<td>Flongle Datasets</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed Diptera subsample (see Srivathsan et al., 2018) - Sanger barcodes available</td>
<td>257 7 negatives</td>
<td>See “Mixed Diptera” entry for R10.3</td>
<td>See “Mixed Diptera” entry for R10.3</td>
<td>Ampure beads (Beckman Coulter)</td>
<td>SQK-LSK109/Flongle</td>
</tr>
<tr>
<td>Chironomidae</td>
<td>191 (Chironomidae) 1 negative</td>
<td>313 bp m1COlIntF, jgHCO2198 (Leray et al. 2013, Geller et al. 2013)</td>
<td>Extraction Method: HotSHOT PCR Mix: Total volume: 14 μl Mastermix (CWBio): 7 μl BSA (1 mg/ml): 1 μl Primer (10μM): 1 μl each DNA: 4 μl</td>
<td>Ampure beads (Beckman Coulter)</td>
<td>SQK-LSK109/Flongle</td>
</tr>
</tbody>
</table>
Bioinformatics

One of the most significant barriers to widespread barcoding with MinION is the high error rates of ONT reads. In 2018, we developed a bioinformatics pipeline for error correction that was too complex for the average user (Srivathsan, Baloglu et al. 2018, Srivathsan, Hartop et al. 2019). After obtaining data with several R10.3 and new R9.4 flow cells, we initially applied this pipeline (Srivathsan et al. 2019), but we noticed major improvements in terms of MinION read quality and the total number of raw and demultiplexed reads produced by each flow cell. This led to the development of a new user-friendly pipeline after considering alternative, published pipelines which faced one or several of the following problems: they required high read coverage, relied on external sequences, were complex, and/or needed several command line steps, and included external dependencies that limit cross platform compatibility (Menegon, Cantaloni et al. 2017, Maestri, Cosetino et al. 2019, Seah, Lim et al. 2020, Sahlin, Lim et al. 2021). We here present “ONTbarcoder”, which has a graphical user interface (GUI) and is suitable for all major operating systems (Linux, Mac OS, Windows10). Both are requirements for the democratization of barcoding with MinION. In addition, we prepared a simple video tutorial (https://www.youtube.com/channel/UC1WowokomhQJRc71FmsUAcg).

ONTbarcoder. ONTbarcoder (available at: https://github.com/asrivathsan/ONTbarcoder) has three modules. (a) The first is a demultiplexing module which assigns reads to specimen-specific bins. (b) The second is a barcode calling module which reconstructs the barcodes based on the reads in each specimen bin. (c) The third is a barcode comparison module that allows for comparing barcodes obtained via different software and software settings.

a. Demultiplexing. The user provides three pieces of information and two files: (1) primer sequence, (2) expected fragment length, and (3) demultiplexing information (=tag combination for each specimen). The latter is summarized in a demultiplexing file (see supplementary information for format). The only other required file is the FASTQ file
obtained from MinKNOW/Guppy after basecalling. Demultiplexing by ONTbarcoder starts by analyzing the read length distribution in the FASTQ file. Only those reads that meet the read length threshold are demultiplexed (default= 658 bp corresponding to metazoan COI barcode). Technically, the threshold should be the amplicon length plus the length of both tagged primers, but ONT reads have indel errors such that they are occasionally too short and we therefore advise to specify the amplicon length as threshold. Reads that are twice the expected fragment length are split into two parts. Splitting is based on the user given fragment size, primer and tag lengths, and a window size to account for indel errors (default=100 bp).

Once all reads suitable for demultiplexing have been identified, ONTbarcoder finds the primers via sequence alignment of the primer sequence to the reads (using python library edlib). Up to 10 deviations from the primer sequence are allowed because this step is only needed for determining the primer location and orientation within the read. For demultiplexing, the flanking region of the primer sequence is retrieved whereby the number of retrieved bases is equal to the user-specified tag length. The flanking sequences are then matched against the tags from the user-provided tag combinations (demultiplexing file). In order to account for sequencing errors, not only exact matches are accepted, but also matches to “tag variants” that differ by up to 2 bps from the original tag (substitutions/insertions/deletions). Note that accepting tag variants does not lead to demultiplexing error because all tags differ by >4 bp. All reads thus identified as belonging to the same specimen are pooled into the same bin. To increase efficiency, demultiplexing is parallelized and the search space for primers and tags are restricted to user-specified parts of each read.

b. Barcode calling: Barcode calling uses the reads within each specimen-specific bin to reconstruct the barcode sequence. The reads are aligned to each other and a consensus sequence is called. Barcode calling is done in three phases: “Consensus by Length”,


“Consensus by Similarity” and “Consensus by barcode comparison”. The user can opt to
only use some of these methods.

“Consensus by Length” is the main barcode calling mode. Alignment must be efficient in
order to obtain high-quality barcodes at reasonable speed for thousands of amplicons.
ONTbarcoder delivers speed by using an iterative approach that gradually increases the
number of reads (“coverage”) that is used during alignment. However, reconstructing
barcodes based on few reads could lead to errors which are here weeded out by using four
Quality Control (QC) criteria. The first three QC criteria are applied immediately after the
consensus sequence has been called: (1) the barcode must be translatable, (2) it has to
match the user-specified barcode length, and (3) the barcode has to be free of ambiguous
bases (“N”). To increase the chance of finding a barcode that meets all three criteria, we
subsample the reads in each bin by read length (thus the name “Consensus by Length”);
i.e., initially only those reads closest to the expected length of the barcode are used. For
example, if the user specified coverage=25x for a 658 bp barcode, ONTbarcoder would only
use the 25 reads that have the closest match to 658 bp. The fourth QC measure is only
applied to barcodes that have already met the first three QC criteria. A multiple sequence
alignment (MSA) is built for the barcodes obtained from the amplicon pool, and any barcode
that causes the insertion of gaps in the MSA is rejected. Note that if the user suspects that
barcodes of different length are in the amplicon pool, the initial analysis should use the
dominant barcode length. The remaining barcodes can then be recovered by re-analyzing all
data or only the failed read bins (“remaining”, see below) and bins that yielded barcodes that
had to be “fixed”. These bins can be reanalyzed using a different pre-set barcode length.

“Consensus by Similarity”. The barcodes that failed the QC during the “Consensus by
Length” stage are often close to the expected length and have few ambiguous bases, and/or
cause few gaps in the MSA. These “preliminary barcodes” can be improved through
“Consensus by Similarity”. This method eliminates outlier reads from the read alignments.
Such reads often differ considerably from the signal of the consensus barcode and ONTbarcoder identifies them by sorting all reads by similarity to the preliminary barcode. Only the top 100 reads (this default can be changed) that differ by <10% from the preliminary barcode are retained and used for calling the barcodes again using the same techniques described previously (including the same QC criteria). This distance threshold accounts for errors generated by MinION but excludes highly erroneous or contaminating reads. This improvement step converts many preliminary barcodes found during “Consensus by Length” into barcodes that pass all four QC criteria by filling/removing indels or resolving an ambiguous base.

“Consensus by barcode comparison”. The remaining preliminary barcodes that still failed to convert into QC-compliant barcodes tend to be based on read bins with low coverage, but some can yield good barcodes after subjecting them to a further improvement step that fixes the remaining errors. ONTbarcoder identifies these errors by finding the 20 most similar QC-compliant barcodes that have already been reconstructed for the other amplicons. The 21 sequences are aligned and ONTbarcoder finds the errors because they cause insertions and deletions in the MSA. Insertions are deleted, gaps are filled with ambiguous bases (“N”), but mismatches are retained. The number and kinds of “fixes” are recorded and added to the FASTA header of the barcode.

Output. ONTbarcoder extensively documents the barcoding results so that users can check the output and potentially modify the barcode calling parameters. For example, it produces a summary table (Outputtable.csv) and FASTA files that contain the different classes of barcodes. Each barcode header contains information on coverage used for barcode calling, coverage of the specimen bin, length of the barcode, number of ambiguities and number of indels fixed. Five sets of barcodes are provided, here discussed in the order of barcode quality: (1) “QC_compliant”: The barcodes in this set satisfy all four QC criteria without correction and are the highest quality barcodes. (2) “Filtered_barcodes”: this file contains the
barcodes that are translatable, have <1% ambiguities and have up to 5 indels fixed during
the last step of the bioinformatics pipeline. These filtering thresholds were calibrated based
on two datasets for which we have Sanger/Illumina barcodes and the resulting barcodes are
found to be highly accurate. Note that the file with filtered barcodes also includes the
QC_compliant barcodes and that all results discussed in this manuscript are based on
filtered barcodes given that they are of much higher quality than the average barcode in

The remaining files include barcodes of lesser and/or suspect quality. (3)
“Fixed_barcodes_XtoY”: these files contain barcodes that had indel errors fixed and are
grouped by the number of errors fixed. Only the barcodes with 1-5 errors overlap with
Filtered barcodes file, if they have <1% ambiguities. (4) “Allbarcodes”: this file contains all
barcodes in sets (1)-(3). (5) “Remaining”: these are barcodes that fail to either translate or
are not of predicted length. Note that all barcodes should be checked via BLAST against
comprehensive databases in order to detect contamination. There are several online tools
available for this and we recommend the use of GBIF sequence ID tool
(https://www.gbif.org/tools/sequence-id) which gives straightforward output including a
taxonomic summary.

The output folder also includes the FASTA files that were used for alignment and barcode
calling. The raw read bins are in the “demultiplexed” folder, while the resampled bins (by
length, coverage, and similarity) are in their respective subfolders named after the search
step. Note that the raw reads are encoded to contain information on the orientation of the
sequence and thus cannot be directly used in other software without modifications (see
ONTbarcoder manual on Github). Lastly, for each barcode FASTA file (1-5), there are
folders with the files that were used to call the barcodes. This means that the user can, for
example, reanalyze those bins that yielded barcodes with high numbers of ambiguous
bases. Lastly a “runsummary.xlsx” document allows the user to explore the details of the barcodes obtained at every step of the pipeline.

Algorithms. ONTbarcoder uses the following published algorithms. All alignments utilize MAFFTv7 (Katoh and Standley 2013). The MSAs that use MinION reads to form a consensus barcode are constructed in an approach similar to lamassemble (Frith, Mitsuhashi et al. 2020), using parameters optimized for nanopore data by “last-train” (Hamada, Ono et al. 2017) which accounts for strand specific error biases. The MAFFT parameters can be modified in the “parfile” supplied with the software which will help with adjusting the values given the rapidly changing nanopore technology. All remaining MSAs in the pipeline (e.g., of preliminary barcodes) use MAFFT’s default settings. All read and sequence similarities are determined with the edlib python library under the Needle-Wunsch (“NW”) setting, while primer search is using the infix options (“HW”). All consensus sequences are called from within the software. This is initially done based on a minimum frequency of 0.3 for each position. This threshold was empirically determined based on datasets where MinION barcodes can be compared to Sanger/Illumina barcodes. The threshold is applied as follows. All sites where >70% of the reads have a gap are deleted. For the remaining sites, ONTbarcoder accepts those consensus bases that are found in at least >30% of the reads. If no base/multiple bases reach this threshold, an “N” is inserted. To avoid reliance on a single threshold, ONTbarcoder allows the user to change the consensus calling threshold from 0.2 to 0.5 for all barcodes that fail the QC criteria at 0.3 frequency. However, barcodes called at different frequencies are only accepted if they pass the first three QC criteria and are identical. If no such barcode is found, the 0.3 frequency consensus barcode is used for further processing.

c. Barcode comparison. Many users may want to call their barcodes under different settings and then compare barcode sets. The ONTbarcoder GUI simplifies such comparisons. A set of barcodes is dragged into the window and the user can select a barcode set as the reference. The barcode comparisons are conducted using edlib library. The barcodes in the
sets are compared and classified into three categories: “identical” where sequences are a perfect match and lack ambiguities, “compatible” where the sequences only differ by ambiguities, and “incorrect” where the sequences differ by at least one base pair. Several output files are provided. A summary sheet, a FASTA file each for “identical”, “compatible”, and the sequences only found in one dataset. Lastly, there is a folder with FASTA files containing the different barcodes for each incompatible set of sequences. This module can be used for either comparing set(s) of barcodes to reference sequences, or for comparing barcode sets against each other. It furthermore allows for pairwise comparisons and comparisons of multiple sets in an all-vs-all manner. This module was used here to get the final accuracy values presented in Table 3.

4. Performance of flow cells (R10.3, Flongle) and high-accuracy basecalling

The pools used to test the new ONT products contained amplicons for 191 - 9,932 specimens and were run for 15-49 hours (Table 2). The fast5 files were basecalled using Guppy in MinIT under the high accuracy (HAC) model. Basecalling large datasets under HAC is currently still very slow and took 12 days in MinIT for the Palaearctic Phoridae (658 bp) dataset (Table 2) but the reads yielded high demultiplexing rates for three of the four R10.3 MinION datasets (= 30-49%). The exception was the Palaearctic Phoridae (313 bp) dataset (15.5%). Flongle datasets showed overall also lower demultiplexing rates (17-21%).
<table>
<thead>
<tr>
<th>Dataset Name</th>
<th>Flow cell details</th>
<th>Raw reads/reads passing length threshold/reads of suitable length/demultiplexed</th>
<th>Demultiplexing rate/# QC_compliant barcodes/# Filtered barcodes with 1N/# Filtered barcodes with &gt;1N/# Unreliable barcodes</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>MiniON R10.3 Datasets</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed Diptera</td>
<td>R10.3: reused flow cell: 71 pores according to QC, but 500+ active during run</td>
<td>3,864,000/3,425,357/3,560,389/1,544,758</td>
<td>43.39%/495/2/5/8 Total success rate= 502/511 (98.2%)</td>
</tr>
<tr>
<td>(658 bp, N=511)</td>
<td>Runtime: 27.5 hrs</td>
<td>Guppy: 4.2.3+f90bd04</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Afrotropical Phoridae</td>
<td>R10.3: new flow cell: QC: 1,101 pores</td>
<td>6,838,903/5,465,164/5,474,029/2,681,029</td>
<td>48.97%/3,725/121/59/247 Total success rate= 390/4275 (91.3%)</td>
</tr>
<tr>
<td>(658 bp, N=4,275)</td>
<td>Runtime: 49.5 hrs</td>
<td>Guppy: 4.0.11+f1071ce</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Palaearctic Phoridae</td>
<td>R10.3: new flow cell: QC: 1,239 pores</td>
<td>16,595,984/15,658,174/16,100,505/5,012,489</td>
<td>31.13%/8,026/108/231/780 Total success rate= 8,365/9,932 (84.2%)</td>
</tr>
<tr>
<td>(658 bp, N=9,932)</td>
<td>Runtime: 47.5 hrs</td>
<td>Guppy: 4.2.3+f90bd04</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Palaearctic Phoridae</td>
<td>R10.3: new flow cell: QC: 1,297 pores</td>
<td>13,690,869/13,221,764/10,366,455/12,983,260/2,015,135</td>
<td>15.52%/8,705/118/12/899 Total success rate= 8,935/9,929 (90%)</td>
</tr>
<tr>
<td>(313 bp, N=9,929)</td>
<td>Runtime: 37 hrs</td>
<td>Guppy: 4.2.3+f90bd04</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Flongle Datasets</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed Diptera Subsample</td>
<td>Flongle: new QC: 81 pores</td>
<td>294,896/222,189/190,952/33,270</td>
<td>17.42%/185/35/20/9 Total success rate= 240/257 (93.4%)</td>
</tr>
<tr>
<td>(658 bp, N=257)</td>
<td>Runtime: 24 hrs</td>
<td>Guppy: v 4.0.11+f1071ce</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chironomidae</td>
<td>Flongle: new QC: 74 pores</td>
<td>560,062/525,087/504,621/108,574</td>
<td>21.52%/178/12/6 Total success rate= 181/191 (94.8%)</td>
</tr>
<tr>
<td>(313 bp, N=191)</td>
<td>Runtime: 15 hrs</td>
<td>Guppy: 4.2.3+f90bd04</td>
<td></td>
</tr>
</tbody>
</table>

We used ONTbarcoder to analyze the MinION data for all six datasets by analyzing all specimen-specific read bins at different coverages (5-200x in steps of 5x). This means that the barcodes for a bin with 27 reads were called five times at 5x, 10x, 15x, 20x, and 25x coverages while bins with >200x were analyzed 40 times at 5x increments. Instead of using conventional rarefaction via random subsampling reads, we used the first reads provided by
the flow cell because this accurately reflects how the data accumulated during the sequencing run and how many barcodes would have been obtained if the run had been stopped early. This rarefaction approach also allowed for mapping the barcode success rates against either coverage or time.

In order to obtain a “best” estimate for how many barcodes can be obtained, we also carried out one analysis at 200x coverage with the maximum number of “Comparison by Similarity” reads set to 100. This means that ONTbarcoder selected up to 200 reads from the specimen-specific read bin that had the closest match to the length of the target barcode (i.e., 313 or 658 bp), then produced an MSA and consensus barcode using MAFFT. If the resulting consensus barcode did not satisfy all four QC criteria, ONTbarcoder would select up to 100 reads that had at least a 90% match to the preliminary barcode. These reads would then be used to call another barcode with MAFFT. Only if this also failed to produce a QC-compliant barcode, ONTbarcoder would “fix” the preliminary barcode using its 20 closest matches in the dataset. All analyses produced a “filtered” set of barcodes (barcodes with <1% Ns and up to 5 fixes) that were used for assessing the accuracy and quality via comparison with Sanger and Illumina barcodes for Mixed Diptera (MinION R10.3), Afrotropical Phoridae (MinION R10.3), and Mixed Diptera Subsample (Flongle R9.4). For the comparisons of the barcode sets obtained at the various coverages, we used MAFFT and the assess_corrected_barcode.py script in miniBarcoder (Srivathsan et al., 2019).

We investigated barcode accuracy (Figure 1) by directly aligning the MinION barcodes with the corresponding Sanger and Illumina barcodes. We find that MinION barcodes are virtually identical to Sanger and Illumina barcodes (>99.99% identity, Table 3). We then established that the number of ambiguous bases (“N”) is also very low for barcodes obtained with R10.3 (<0.01%). Indeed, more than 90% of all barcodes are entirely free of ambiguous bases. In comparison, Flongle barcodes have a slightly higher proportion of ambiguous bases (<0.06%). They are concentrated in ~20% of all sequences so that 80% of all barcodes
again lack Ns. This means that MinION barcodes more than just match the Consortium for the Barcode of Life (CBOL) criteria for "barcode" designation with regard to length, accuracy, and ambiguity.

Rarefaction at different read coverage levels reveals that 80-90% of high-quality barcodes are obtained within a few hours of sequencing. In addition, the number of barcodes generated by MinION exceeded or was comparable to what could be obtained with Sanger or Illumina sequencing (Figure 1). We then determined the coverage needed for obtaining reliable barcodes. For this purpose, we plotted the number of barcodes obtained against coverage (Figure 2). This revealed that the vast majority of specimen bins yield high-quality barcodes at coverages between 25x and 50x when R10.3 reads are used. Increasing coverage beyond 50x leads to only modest improvements of quality and few additional specimen amplicons yield new barcodes. The coverage needed for obtaining Flongle barcodes is somewhat higher, but the main difference between the R9.4 technology of the Flongle flow cell and R10.3 is that more barcodes retain ambiguous bases even at high coverage for data from R9.4 flow cells. The differences in read quality between R9.4 and R10.3 become even more obvious when the read bins for the “Mixed Diptera Subsample” are analyzed based an identical numbers of R10.3 and R9.4 reads. The barcodes based on Flongle and R10.3 data are compatible, but the R10.3 barcodes are ambiguity-free while some of the corresponding Flongle barcodes retain 1-2 ambiguous bases.

Overall, these results imply that 100x raw read coverage is sufficient for obtaining barcodes with either R10.3 or R9.4 flow cells. Given that most MinION flow cells yield >10 million reads of an appropriate length, this means that one could, in principle, obtain 100,000 barcodes in one flow cell. However, this would require that all amplicons are represented by similar numbers of copies and that all reads could be correctly demultiplexed. In reality, only 30-50% of the reads can be demultiplexed and the number of reads per amplicon fluctuates widely (Figure 3). Very-low coverage bins tend to yield no barcodes or barcodes of lower
quality (errors or Ns). These low-coverage barcodes can be improved by collecting more
data, but this comes at a high cost and increased risk of a small number of contaminant
reads yielding barcodes. For example, we observed that some “negative” PCR controls
yielded low-quality barcodes for 4 of 106 negatives in the Palaearctic Phoridae (313 bp) and
1 of 105 negatives in the Palaearctic Phoridae (658 bp) datasets.

To facilitate the planning of barcode projects, we illustrate the trade-offs between barcode
yield, time, and amount of raw data needed for six amplicon pools (Figure 4: 191-9,932
specimens). These standard curves can be used to roughly estimate the amount of raw
reads needed to achieve a specific goal for a barcoding project of a specific size (e.g.,
obtaining 80% of all barcodes for a project with 1000 amplicons). For each dataset, we
illustrate how many reads were needed to recover a certain proportion of barcodes. The
number of recoverable barcodes was set to the number of all error-free, filtered barcodes
obtained in an analysis of all data. We would argue that this is a realistic estimate of
recoverable barcodes given the saturation plots in Figure 1 that suggest that most barcodes
with significant amounts of data have been called at 200x coverage. Note, however, that
Figure 4 can only provide very rough guidance on how many reads are needed because, for
example, the demultiplexing rates differ between flow cells and different amplicon pools have
very different read abundance distributions (see Figure 3).
Table 3. Quality assessment of barcodes generated by ONTbarcoder at 200X read coverage (Consensus by Length) and 100X coverage (Consensus by Similarity). The accuracy of MinION barcodes is compared with the barcodes obtained for the same specimens using Illumina/Sanger sequencing. Errors are defined as sum of substitution or indel errors. All denominators for calculating percentages are the total number of nucleotides assessed.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>No. of comparison barcodes</th>
<th>No. of barcodes with errors/No. of errors/% identity</th>
<th># of Ns/%Ns</th>
</tr>
</thead>
<tbody>
<tr>
<td>R10.3: Mixed Diptera: Sanger barcodes available</td>
<td>476</td>
<td>2/10/99.997%</td>
<td>19 (0.006%)</td>
</tr>
<tr>
<td>R10.3: Afrotropical Phoridae: Illumina barcodes available*</td>
<td>3316</td>
<td>23/48/99.995%</td>
<td>284 (0.011%)</td>
</tr>
<tr>
<td>Flongle-Mixed Diptera Subsample: Sanger barcodes available</td>
<td>231</td>
<td>5/8/99.994%</td>
<td>91 (0.058%)</td>
</tr>
</tbody>
</table>

*5 barcodes with very high distances from reference were excluded for R10.3: Afrotropical Phoridae dataset as they likely represent lab contamination (see Srivathsan, Hartop et al. (2019).
Figure 1. Rapid recovery of accurate MinION barcodes over time (in hours, x-axis) (filtered barcodes: dark green = barcodes passing all 4 QC criteria, light green = one ambiguous base; lighter green = more than 1N, no barcode = white with pattern, 1 mismatch = orange, >1 mismatch = red). The solid black line represents the number of barcodes available for comparison. White dotted line represents the amount of raw reads collected over time, blue represents number of demultiplexed reads over time (plotted against Z-axis).
Figure 2. Relationship between barcode quality and coverage. Subsetting the data to 5-200X coverage shows that there are very minor gains to barcode quality after 25-50X coverage. (filtered barcodes: dark green = barcodes passing all 4 QC criteria, light green = one ambiguous base; lighter green = more than 1N, no barcode = white with pattern, 1 mismatch = orange, >1 mismatch = red).
Figure 3. Read bin size distribution for six amplicon pools (color-coding as in Figs 1-2). Due to the very generous coverage for the “Mixed Diptera” dataset, we also use grey to show the bin size distribution after dividing the bin read totals by 5.
Figure 4. Relationship between barcoding success and number of raw reads for six amplicon pools (191-9932 specimens; barcoding success rates 84-97%). Percentage of barcodes recovered is relative to the final estimate based on all data.
Discussion

Democratization of Barcoding

Biodiversity research needs new scalable techniques for large-scale species discovery and monitoring. This task is particularly urgent and challenging for invertebrates that collectively make up most of the terrestrial animal biomass. We argued earlier that this is likely to be a task that requires the processing of at least 500 million specimens from all over the world with many tropical countries with limited research funding requiring much of the biodiversity discovery work. Pre-sorting these specimens into putative species-level units with DNA sequences is a promising solution as long as obtaining and analyzing the data are sufficiently straightforward and cost-effective. We believe that the techniques described in this manuscript will help with achieving these goals. Generating DNA barcodes involves three steps. The first is obtaining a DNA template, and we have herein outlined some simplified procedures that render this process essentially free-of-cost, although automation and AI-based solutions will be useful for processing very large numbers of specimens in countries with high manpower cost. The third step is the sequencing of the amplicon. Fortunately, there are now several cost-effective solutions based on 2nd and 3rd generation sequencing technologies so that barcodes can be sequenced for as little as a penny (USD).

We here argue that sequencing with MinION is particularly attractive although the cost is higher (0.10 USD) than with Illumina sequencing. There are several reasons. MinION library preparation can be learned within hours and an automated library preparation instrument is in development that will eventually work for ligation-based libraries. Furthermore, MinION flow cells can accommodate projects of varying scales. Flongle can be used for amplicon pools with a few hundred products, while an R10.3 flow cell can accommodate projects with up to 10,000 specimens. The collection of data on MinION flow cells can be stopped whenever enough have been acquired. Flow cells can then be washed and re-used again although the remaining capacity declines over time because some nanopores will become unavailable. We have re-used flow cells up to four times. Traditionally, the main obstacles to
using MinION have been poor read quality and high cost. Both issues are fading into the past. The quality of MinION reads has improved to such a degree that the laptop-version of our new software “ONTbarcoder” can generate thousands of very high quality barcodes within hours. There is no longer a need to polish reads or rely on external data or algorithms.

The greater ease with which MinION barcodes can be obtained is due to several factors. Firstly, much larger numbers of reads can now be obtained with one MinION flow cell. Secondly, R10.3 reads have a different error profile which allows for reconstructing higher-quality barcodes. Thirdly, high accuracy basecalling has improved raw read quality and thus demultiplexing rates. Lastly, we can now use parameter settings for MAFFT that are designed for MinION reads. These changes mean that even low-coverage bins yield very accurate barcodes; i.e., both barcode quality and quantity are greatly improved.

**Rapid progress in barcode quality and quantity**

We previously tested MinION for barcoding (2018, 2019) and here re-sequenced some of the same amplicon pools. This allowed for a precise assessment of the improvements. In 2018, sequencing the 511 amplicons of the *Mixed Diptera* sample required one flow cell and we obtained 488 barcodes of which only one lacked ambiguous bases. In 2021, we used the remaining ~500 pores of a used R10.3 flow cell (1st use was for 49 hours). After washing, we obtained 502 barcodes and >98% (496) of them were free of ambiguous bases. The results obtained for the 2019 amplicon pools were also better. In 2019, one flow cell (R9.4) allowed us to obtain 3,223 barcodes from a pool of amplicons obtained from 4,275 specimens of *Afrotropical Phoridae*. Resequencing weak amplicons increased the total number of barcodes by approximately 500 to 3,762 (Srivathsan, Hartop et al. 2019). Now, one R10.3 flow cell yielded 3,905 barcodes (+143) for the same amplicon pool, while retaining an accuracy of >99.99% and reducing the ambiguities from 0.45% to 0.01%. If progress continues at this pace, we predict that MinION will be the default barcoding tool for most users. This, too, is because all barcoding steps can now be carried out in one laboratory with a modest set of equipment (see Table 4). With MinION being readily available, there is no
longer the need to outsource sequencing and/or to wait until enough barcode amplicons have been prepared for an Illumina or PacBio flow cell (Ho, Puniamoorthy et al. 2020). This democratizes biodiversity discovery and allows many biologists, government agencies, students, and citizen scientists from around the globe to get involved in these initiatives. Biodiversity discovery with cost-effective barcodes will also facilitate biodiversity discovery in countries with high biodiversity but limited science funding.

Table 4. Equipment required for MinION barcoding

<table>
<thead>
<tr>
<th>Required</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 MinION sequencer (preferably Mk1C for basecalling)</td>
</tr>
<tr>
<td>2 Thermocycler(s)</td>
</tr>
<tr>
<td>3 Gel Electrophoresis setup</td>
</tr>
<tr>
<td>4 Magnetic Separation Rack</td>
</tr>
<tr>
<td>5 Qubit for DNA quantification</td>
</tr>
<tr>
<td>6 Standard equipment: Vortex, Mini-centrifuge, pipettes, freezer, fridge</td>
</tr>
<tr>
<td>7 Standard laptop or PC</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Optional but highly desirable</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Multichannel pipette(s)</td>
</tr>
<tr>
<td>2 Hula Mixer</td>
</tr>
</tbody>
</table>

This raises the question of how much it costs to sequence a barcode with MinION. There is no straightforward answer because the cost depends on user targets. For example, a user who wants to sequence a pool of 5000 barcodes may want a 80% success rate in order to identify the dominant species in a sample. Based on Figure 4, only ca. 1.5 million raw MinION reads would be needed. On average, MinION flow cells yield >10 million reads and cost USD 475-900 depending on how many cells are purchased at the same time. Including
a library cost of ca. USD 100, the overall sequencing cost of a project that requires 1.5
million reads is USD 180-235. This experiment would be expected to yield 4000 barcodes for
the 5000 amplicons (4-6 cents/barcode). Given the low cost of 1 million MinION reads ($50-
90), we predict that most users will opt for sequencing at a greater depth since this will likely
yield several hundred additional barcodes. However, this will then increase the sequencing
cost per barcode, because the first 1.5 million reads already recovered barcodes for all
strong amplicons. Additional reads will predominantly strengthen read coverage for these
amplicons and relatively few reads will be added to the read bins that were too weak to yield
barcodes at low coverage; i.e., there are diminishing returns for additional sequencing.

Overall, we thus predict that most users will, at most, try to multiplex 10,000 amplicons in the
same MinION flow cell so that the sequencing cost per specimen would be 0.06-0.10 USD
depending on the bulk purchase of flow cells. However, we also predict that large-scale
biodiversity projects will switch to sequencing with PromethION, a larger sequencing unit
that can accommodate up to 48 flow cells. This will lower the sequencing cost by more than
60%, as PromethION flow cells have 6 times the number of pores for twice the cost (capacity
per flow cell should be 60,000 barcodes). At the other end of the scale are those users who
occasionally need a few hundred barcodes. They can use Flongle flow cells, which are
comparatively expensive (0.50 USD) because each flow cell costs $70 and requires a library
that is prepared with half the normal reagents (ca. $50). A change of the flow cell chemistry
from that of R9.4 to R10.3 would, however, help with improving the quality of the barcodes
obtained from Flongle. Lastly the initial setup cost for MinION/Flongle, can be as low as
1000 USD, but we recommend purchase of Mk1C unit (currently 4900 USD) for easy access
to a GPU that is required for high accuracy basecalling. Note also, that obtaining flow cells at
low cost often requires collaboration between several labs because it allows for buying flow
cells in bulk.

*ONTbarcoder for large-scale species discovery with MinION*
There are a number of studies that have used MinION for barcoding fungi, animals, and plants (Menegon, Cantaloni et al. 2017, Pomerantz, Peñafiel et al. 2018, Wurzbacher, Larsson et al. 2018, Krehenwinkel, Pomerantz et al. 2019, Maestri, Cosetino et al. 2019, Chang, Ip et al. 2020, Knot, Zouganelis et al. 2020, Seah, Lim et al. 2020, Sahlin, Lim et al. 2021). There is one fundamental difference between these studies and the vision presented here. These studies tended to show that MinION sequencing can be done in the field. Thus only a very small number of specimens were analysed (<150 with the exception of >500 in Chang, Ip et al 2020). The field use is an attractive feature for time-sensitive samples that could degrade before reaching a lab. However, for the time being it is unlikely to help substantially with tackling the challenges related to large-scale biodiversity discovery and monitoring because obtaining few MinION barcodes per flow cell is too expensive for most researchers in biodiverse countries. Additionally, the bioinformatic pipelines that were developed for these small-scale projects were not suitable for large-scale, decentralized barcoding in a large variety of facilities. For example, some of the studies used ONT’s commercial barcoding kit that only allows for multiplexing up to 96 samples in one flow cell (Maestri, Cosetino et al. 2019, Seah, Lim et al. 2020); i.e., each amplicon had very high read coverage which influenced the corresponding bioinformatics pipelines (e.g., ONTrack’s recommendation is 1000x: Maestri, Cosentino et al. 2019). The generation of such high coverage datasets also meant that the pipelines were only tested for such a small number of samples (<60: Menegon, Cantaloni et al. 2017, Maestri, Cosetino et al. 2019, Seah, Lim et al. 2020, Sahlin, Lim et al. 2021) that these tests were unlikely to represent the complexities of large, multiplexed amplicon pools (e.g., nucleotide diversity, uneven coverage).

ONTbarcoder evolved from miniBarcoder, whose barcodes have been assessed for accuracy in four different studies covering >8000 barcodes (Chang, Ip et al. 2020, Chang, Ip et al. 2020, Srivathsan, Baloglu et al. 2018, Srivathsan, Hartop et al. 2019). The new software introduced here addresses two drawbacks of its precursor, miniBarcoder. Firstly,
we dropped the translation-based error correction that tended to increase the number of Ns.

This step used to be essential because indel errors were prevalent in consensus barcodes obtained with older flow cell models. Secondly, ONTbarcoder can be installed by unzipping a file and is easy to maintain on different operating systems. Until now, external dependencies were a major drawback of all MinION bioinformatics pipelines. For example, the one described by Sahlin et al. (2021) involved minibar/qcat and nanofilt, while NGSpeciesID relies on isONclust SPOA, Parasail, and optionally, Medaka (Daily 2016, Krehenwinkel, Pomerantz et al. 2019, Sahlin and Medvedev 2020). These dependencies and complexities meant that Watsa et al. (2020) recommended bioinformatics training before MinION barcoding could be used in schools (e.g., training in UNIX command-line) and additionally required the installation of several software tools onto the teaching computers. Neither is needed for ONTbarcoder, which runs on a regular laptop and has been extensively tested (>4000 direct comparisons to Sanger and Illumina barcodes). In addition, ONTbarcoder is designed in a way that thousands of barcodes can be obtained rapidly without impairing accuracy; i.e., one can run a very fast analysis by using low read coverage. However, at very low coverages, fewer barcodes would be recovered because many would not pass the 4 QC criteria. Speed is also achieved through the parallelization of most steps on UNIX systems (Mac and Linux; parallelization is restricted to demultiplexing in Windows).

ONTbarcoder also allows for updating the parameter file for alignment. This is advisable because MinION continues to evolve quickly. We expect flow cell capacity to increase further and basecalling to improve (see Xu, Mai et al. 2020). For example, a new basecaller (“bonito”) developed by ONT has shown promise by improving raw read accuracy (https://nanoporetech.com/about-us/news/new-research-algorithms-yield-accuracy-gains-nanopore-sequencing). This basecaller is currently suitable for research teams equipped with GPU infrastructure and for advanced users familiar with Linux command lines.

However, our preliminary tests of bonito for barcoding (Flongle: Mixed Diptera Subsample, Chironomidae; R10.3: Palaeartic Phoridae, 313 bp; bonito version=0.3.6) does not yet
significantly affect barcode quality or quantity (unpublished data). However, this may change in the immediate future and readers are advised to watch out for developments. Fortunately, these changes will only further improve MinION barcodes that are already highly accurate and cost-effective.

Biodiversity monitoring

Some readers are likely to argue that large-scale biodiversity discovery and monitoring can be more efficiently carried out via metabarcoding of whole samples consisting of hundreds or thousands of specimens. This would question the need for large-scale, decentralized barcoding of individual specimens. However, large-scale barcoding and metabarcoding will more likely complement each other. For example, large-scale barcoding of individual specimens remains essential for discovering and describing species as it preserves individual voucher specimens associated with the barcode which can be used for further research. Taxonomic research can be guided by examination of putative species units (molecular Operational Taxonomic Units or mOTUs) using species delimitation algorithms (either distance based clustering of sequences: Meier, Shiyang et al. 2006; Puillandre, Brouillet et al. 2020) or tree based methods (Pons, Barraclough et al. 2006; Zhang, Kapli et al. 2013). In this process, it is important to remember that COI lumps recently diverged species and divides species with deep allopatric splits (Hickerson, Meyer et al. 2006), making the ability to relate barcodes to individual specimens critical for barcode cluster validation. High quality barcode databases are important for the analysis of metabarcoding data because they facilitate the identification of numts, heteroplasmy, contaminants and errors. Large-scale barcoding will also be needed in order to benefit from another new technique that may become critical for biodiversity discovery and monitoring; i.e. AI-assisted analysis of images (Valan, Makonyi et al. 2019). Large-scale barcoding generates identified specimens that can be imaged and utilized for training neural networks. With increasing advancements in imaging hardware, computational processing power and machine learning systems, AI-assisted biodiversity monitoring could be the method of choice in the future.
Conclusions

Many biologists would like to have ready access to barcodes without having to run large and complex laboratories or send specimens halfway around the world. Many have been impressed by MinION’s low cost, portability, and ability to deliver real-time sequencing, but they were worried about high cost and complicated bioinformatics pipelines. We here demonstrate that these concerns are no longer justified. MinION barcodes obtained by R10.3 flow cells are virtually identical to barcodes obtained with Sanger and Illumina sequencing. Barcoding with MinION is now also cost-effective and the new “ONTbarcoder” software makes it straightforward for researchers with little bioinformatics background to analyze the data on a standard laptop. This will make biodiversity discovery scalable and accessible to all.

Acknowledgements

We would like to thank John T. Longino and Michael Branstetter for providing valuable comments on the manuscript. For the Palaearctic phorid samples, we would like to thank Dave Karlsson, the Swedish Insect Inventory Project, and the crew at Station Linné that sorted out the phorids. We would also like to thank Wan Ting Lee for help with molecular work, and the numerous staff, students and interns who have contributed to the establishment of the pipeline in the NUS laboratory. We would also like to acknowledge Suphavilai Chayaporn and Niranjan Nagarajan from Genome Institute of Singapore for their help with basecalling using bonito. This work was supported by a Ministry of Education grant on biodiversity discovery (R-154-000-A22-112).

Software and test dataset availability
ONTbarcoder is available at https://github.com/asrivathsan/ONTbarcoder, which also contains the link to download the raw data and demultiplexing files. The manual for the software is included in the repository https://github.com/asrivathsan/ONTbarcoder/blob/main/ONTBarcoder_manual.pdf. The videos tutorials can be found in the YouTube channel Integrative Biodiversity Discovery: https://www.youtube.com/channel/UC1WowokomhQJRc71FmsUAcg.

**Literature cited**


assessments with high phylogenetic resolution across broad taxonomic scale."

Gigascience 8(5): giz006.


Meier, R., W. H. Wong, A. Srivathsan and M. S. Foo (2016). "$1 DNA barcodes for reconstructing complex phenomes and finding rare species in specimen-rich samples."

Cladistics 32(1): 100-110.


Swiss Re. (2020). "Biodiversity and Ecosystem Services A business case for re/insurance." Zurich, Swiss Re Management Ltd.


