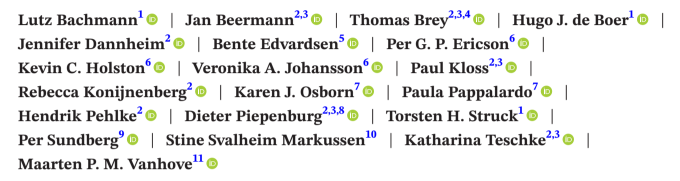
ORIGINAL ARTICLE



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The role of systematics for understanding ecosystem functions: Proceedings of the Zoologica Scripta Symposium, Oslo, Norway, 25 August 2022



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Abstract

On 25 August 2022, the Zoologica Scripta - An International Journal of Systematic Zoology and the Norwegian Academy of Sciences and Letters arranged a symposium entitled 'The role of systematics for understanding ecosystem functions' in the Academy's premises in Oslo, Norway. The symposium aimed at offering a forum for exploring and discussing trends and future developments in the field of systematics. Eleven international experts contributed expertise on various issues related to global challenges, such as biodiversity assessments, databases,

Authors are listed in alphabetical order, and all authors contributed equally to the paper.

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cutting-edge analysis tools, and the consequences of the taxonomic impediment. Here, we compiled a multi-author proceedings paper of the symposium contributions that are arranged in chapters and presents the content and the key conclusions of the majority of the presentations.

KEYWORDS

animal systematics, biodiversity, climate change, databases, ecosystems, taxonomy

1 | MOTIVATION AND INTRODUCTION

Lutz Bachmann, Per G. P. Ericson, Per Sundberg.

Many biologists are concerned about the taxonomic impediment recognized already in 1992 by the Convention on Biological Diversity signed in Rio de Janeiro, Brazil. The decline of expertise in taxonomy and systematics; even more so in the context of ongoing biodiversity loss and climate change. At the same time, we experience an upgrading of the portfolio of methods to highly advanced and complex 'big data' analyses, either project generated, retrieved from databases or a combination of both. A frequently heard concern is that this trend may further decrease expertise in taxonomy and systematics and increase the knowledge gap. The trends towards more complex bioinformatics analyses as well as addressing and understanding systematics in an ecological context are also reflected in the manuscripts submitted to the Zoologica Scripta - An International Journal of Systematic Zoology.

The taxonomic impediment is not only a scientific issue but also of importance for societal development. Roughly 10 years ago, the Intergovernmental Science-Policy Platform for Biodiversity and Ecosystem Services (IPBES) was established, aiming, among others, at advancing knowledge and data as well as at building capacities that strengthen the science-policy interface for biodiversity and ecosystem services for the conservation and sustainable use of biodiversity. In addition, national initiatives such as, for example the Norwegian Biodiversity Information Centre (artsdatabanken.no),

invest substantially in monitoring biodiversity and compiling public databases.

The Zoologica Scripta and its publisher, the Norwegian Academy of Sciences and Letters (DNVA), organized a symposium entitled 'The role of systematics for understanding ecosystem functions' in order to offer a forum for exploring and discussing trends and future developments in the field of systematics. International experts contributed a series of talks that focused on various issues related to among others global challenges, biodiversity assessments, databases and cutting-edge analysis tools. The detailed program can be found at https://dnva.no/detskjer/2022/06/role-systematics-understanding-ecosy stem-functions. Roughly, 70 participants attended the symposium that was held at DNVA's premises in Oslo, Norway, on 25 August 2022 and contributed to the lively discussions.

We editors of the Zoologica Scripta think that the contributions to the symposium are of great interest to many scientists in the field that could not attend the meeting. We, therefore, suggested to the presenters to contribute to a proceedings paper. This multi-author paper covers the major topics and conclusions of the symposium and beyond. It is arranged in several chapters based on the individual talks presented at the meeting. However, the chapter titles may differ from the original titles of the symposium talks.

The symposium closed with an evening dinner to which Matthias Glaubrecht contributed an inspiring talk. The essence of his evening lecture entitled 'On the end of evolution – Humankind and the annihilation of species'

will be provided as a separate contribution to this issue of the Zoologica Scripta (Glaubrecht, 2023).

Finally, we wish to express our gratitude to all contributors to and participants of the symposium that turned the arrangement into an exciting and stimulating event.

1.1 | Acknowledgements

We highly appreciate the generous sponsoring of the symposium by the Norwegian Academy of Sciences and Letters (dnva.no) represented by general secretary Gunn Elisabeth Birkelund and director Øyvind Sørensen. We are particularly thankful for the extraordinary support of project leaders Aslaug K. L. Fuchs and Eirik Lislerud in planning and arranging the symposium.

2 | TAXONOMY IS CHANGING BUT WHERE IS IT HEADING?

Hugo J. de Boer, Stine Svalheim Markussen.

2.1 | Taxonomy and species discovery

In order to understand the role of taxonomy, we should never be afraid to ask ourselves why taxonomy and systematics matter. Numerous studies highlight the gap between expected and described biodiversity and how this differs from one organism group to another (Colwell & Coddington, 1994; Costello, 2015; Laurance & Edwards, 2011). To many scholars of taxonomy, there is no doubt that discovering and describing all diversity on Earth is a matter of general importance (Bebber et al., 2010; Mace, 2004; Padial et al., 2010), and accurate species recognition underpins our knowledge of global biodiversity (Chapman, 2009; Pitman & Jørgensen, 2002; Stuart et al., 2010). The discipline of taxonomy has nevertheless long been neglected and underfunded (Cotterill & Foissner, 2010; Pearson et al., 2011; Wägele et al., 2011; Wheeler et al., 2004; Wilson, 2004). Furthermore, across taxonomic groups, new species continue to be described at rates that suggest a backlog in both species discovery and description (Antonelli et al., 2020; Barreca et al., 2020). Not all species groups yield as many new species as others. Some groups are well known, others are poorly known. Some are easy to study, whereas others are very challenging (Jörger & Schrödl, 2013). Whether we look at terrestrial or marine ecosystems - new species continue to be discovered everywhere (Cheek et al., 2020; Glaw et al., 2012; Rouse et al., 2004).

Also, species are not discovered at equal rates across the world. In Europe and North America, few species are discovered, whereas in Asia and South America many. In Africa, very few new species are discovered due to a lack of research (Antonelli et al., 2020). While Norway and the other Nordic countries have relatively low species diversity compared to most countries and are seen as well investigated when it comes to species richness, the Norwegian Taxonomy Initiative and related projects have shown that there still is a significant number of species to be discovered (Elven & Søli, 2021). Since its beginning in 2009, the Norwegian Taxonomy Initiative has funded 120 projects aimed to investigate poorly known species groups, and these have discovered more than 4000 species new to Norway. Elven and Søli (2021) estimate that only about 65% of the total number of species present in Norway have been found to date. Based on species discoveries, we can also predict the potential of future discoveries. An analysis of terrestrial vertebrate taxa discovery identifies those groups were the highest potential for new species discovery remains (Moura & Jetz, 2021). Taxonomy is not only important for the description of new species per se but is also a great way of communicating the value of biodiversity to an audience that is increasingly disconnected from nature. The tiny chameleon Brookesia nana (Glaw et al., 2021), which may be the world's smallest reptile species, is an example of a species description that has generated a lot of interest in biodiversity. Another example is the discovery of a new Giant Waterlily hidden in plain sight at Kew (Smith et al., 2022). Victoria boliviana Magdalena and L.T.Sm. had been growing at London's Kew Gardens for 177 years and in the National Herbarium of Bolivia for 34 years, but botanists had not realized that it was distinct from its closely related *V. amazonica* (Poepp.) J.C. Sowerby. The latter is also a good example of how not all undiscovered species are hiding in pristine rainforests, but that some are already collected in natural history museums and botanic gardens.

2.2 | Species loss – a race against the clock

However, species discovery is a race against the clock, with species loss far outpacing species discovery (Humphreys et al., 2019). Around 1 million animal and plant species are now threatened with extinction, many within decades, and more than ever before in human history (IPBES, 2019). The decline in wildlife and the number of mammal species lost are indicators of the attrition of biodiversity across the board. Thirty-two million hectares of primary or recovering forest were lost between 2010 and 2015 (IPBES, 2019). This is an area nearly the size of

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Norway, and Norway is not a small country. A less widely communicated phenomenon is that in some of the most economically developed countries, we are losing species and rediscovering them too (Humphreys et al., 2019). This highlights that species loss can only be accurately quantified if species diversity is known. Assessing species loss in the rest of the world can only be measured by extrapolation and approximation. Taxonomic impediment is the worldwide shortage of this important taxonomic information, the gaps in our taxonomic knowledge and the shortage of trained taxonomists and curators to fill this need. The taxonomic impediment is also often invoked to stress the unequal distribution of taxonomic knowledge. The study by Lücking (2020) highlights this dismayingly. Nearly, all the 200 most prolific fungal taxonomists, responsible for nearly half of all approximately 360,000 species names described, lived or live in the global north.

2.3 The value of biodiversity

Communicating the value of biodiversity can be done in many ways but putting it in economic terms makes it easier to balance with other economic activities. The 2021 World Bank report, 'The Economic Case for Nature: A global Earth-economy model to assess development policy pathways' (Johnson et al., 2021), estimates that protecting nature could avert global economic losses of \$2.7 trillion per year by 2030. This type of doom value is alarming but does not help us understand in what ways biodiversity contributes to ecosystem services, in terms of livelihoods and sustainability. The Intergovernmental Science-Policy Platform on Biodiversity and Ecosystem Services (IPBES) uses 18 'Nature's Contributions to People' to analyse and value ecosystem services (Chaplin-Kramer et al., 2019; Díaz et al., 2018; Hill et al., 2021; Pascual et al., 2017). A clear framework helps stakeholders communicate the value of nature to human livelihoods and sustainability. Similarly, the UN Sustainable Development Goals (SDGs) provide a framework to communicate integrated sustainability objectives. This visualization by the Stockholm Resilience Centre shows the hierarchical organization of the SDGs. Many SDGs are related to the economy and society but all rely upon a healthy biosphere. Understanding biodiversity underpins the SDGs for Life below Water and Life on Land and its heavy dependence on biodiversity knowledge.

Is species discovery a priority? 2.4

However, can we equate biodiversity knowledge with systematics and taxonomy? How much do we need to know

about biodiversity if we can already assess and quantify its value? If we know the species we need, do we need to describe the rest? Do we need to stick a label on each creature in our ecosystem? The apparent gap between taxonomic knowledge and biodiversity has been highlighted by many scholars (Giangrande, 2003; Kim & Byrne, 2006; McNeely, 2002). We have arguably entered the sixth mass extinction, with species loss being accelerated by humaninduced effects like never before (Ceballos et al., 2015). We are currently in a situation where for many species, we are unaware that they are at risk of extinction because we know them so poorly (Boehm & Cronk, 2021; Reaka-Kudla, 2001; Scheffers et al., 2012; Tedesco et al., 2014). Although new species are continuously being described, at the same time, others are going extinct. Plant extinctions are shown to occur up to 500 times faster today than in pre-industrial times (Humphreys et al., 2019). Unfortunately, many more species are going extinct without us knowing about it or even having discovered them (Boehm & Cronk, 2021; Scheffers et al., 2012; Tedesco et al., 2014). If species discovery is a priority, we are in a hurry to ramp things up if we want to describe a fraction of the species being lost.

Taxonomy today - new ways in 2.5 which taxonomy contributes

The CBD Global Taxonomy Initiative Forum for the post-2020 Global Biodiversity Framework (Abrahamse et al., 2021) launched a call for action stressing that 'Innovative and emerging technologies provide unprecedented opportunities for generating and sharing knowledge about the biosphere, when combined with essential taxonomic knowledge, techniques and skills'. Taxonomic knowledge is essential to generating and sharing knowledge about the biosphere, but in order to accelerate this work, we need to harness the opportunities provided by emerging technologies. New methods empower taxonomists like never before. DNA evidence can provide a lead to reassess previously overlooked diversity (Hebert & Gregory, 2005); and molecular species delimitation can aid in separating species that are hard to recognize using morphology alone (Luo et al., 2018). The advent of DNA sequencing technologies has also helped to uncover a previously unknown world of cryptic species (Struck et al., 2018a, 2018b). Studies looking at environmental DNA (eDNA) have opened Pandora's box of taxonomic needs with many studies being able to identify only a minority of operational taxonomic units (OTUs) (Arroyo et al., 2016; Carvalho et al., 2019; Gleason et al., 2022; Sinniger et al., 2016). In order to increase the resolution and use of biodiversity assessments through eDNA, we

need to generate DNA barcodes from reference specimens, which in turn are based on morphologically identified specimens deposited in natural history collections, which in turn require taxonomists and species descriptions. The more we want to employ molecular biodiversity assessments for biodiversity monitoring the greater the need for taxonomic expertise. Within the Norwegian setting, these interlinked dependencies support each other. The Norwegian Environment Agency has a strong focus on the development of sampling, data and analysis standards for eDNA biodiversity monitoring. This helps the private and institute sector use validated methods for reproducible studies. The Taxonomy Initiative program of the Norwegian Biodiversity Information Centre supports species discovery and mapping of poorly known biodiversity and liaises with the distributed Norwegian Barcode of Life (NorBOL) infrastructure to ensure that all specimens are barcoded and sequences deposited in public data repositories. The Taxonomy Initiative projects stimulate the transmission of taxonomic knowledge, generation of new taxonomic knowledge and help train a new generation of taxonomists within the academic and institute sector.

2.6 | Taxonomy tomorrow – what does the future need?

Whereas a taxonomist in the past might have been able to focus mostly on the collection, identification and description of species, a taxonomist today can benefit from a more diverse skillset to succeed within academia. The Norwegian Biodiversity Information Centre funded Scandinavian Research School in Biosystematics - ForBio, coordinated by the four Norwegian university museums, provides training in core taxonomy subjects as well as other relevant methods, including genomics, phylogenetics, phylogenomics, biogeography, grant writing, publication and presenting. It is hard to know what skillset will be helpful to have in the future but having expert taxonomic skills that can contribute to integrative and collaborative projects is certainly essential. Taxonomic skills are in increasing demand in collaborative research and being able to understand and highlight where these skills can yield synergies is important in promoting the field. Recent publications drawing on taxonomic expertise suggest that a future taxonomist should be a taxonomic expert first and foremost, but also able to accelerate species description by complementing traditional morphology with novel data such as molecular evidence and deep-learning approaches (Coleman & Radulovici, 2020; Padial et al., 2010; Valan et al., 2021; Yang et al., 2022). As a field, we should increase rates of species diversity monitoring to track ecosystem health, decline and recovery and advocate for a more equal sharing of taxonomic knowledge through Findible, Accessible, Interoperable and Reusable data.

3 | THE POTENTIAL AND POWER OF BIODIVERSITY DATABASES

Veronika A. Johansson, Kevin C. Holston.

3.1 | Biodiversity databases – purpose and procedures

As the scale and scope of biological data grow and become available for use and reuse, examining its persistence in various aggregated forms provides insight into the range of intended purposes, or target user groups, for biodiversity databases. The category 'biological data' encompasses any information derived from living organisms and their products (Dhillon, 2019; Gauthier et al., 2019), but the scope of biodiversity databases is often limited to a single data class. These can include species traits, nucleotide and protein sequences, or observation records with proprietary and additional descriptive information often associated with these individual records. In practice, these data classes usually represent the transformation and refinement of raw data into data sets that are fit for use. Methods in biodiversity informatics generate these raw data via measurements from biosensors or human observations; equivalent methods generate and refine laboratory data into research data sets, for example DNA sequence or genomic data.

Application of similar analysis methods to legacy, as well as novel data sets, reinforces standardization of practices in data set compilation, which characterize scientific works subject to peer review. In this way, community-driven practices can support the growth and maintenance of specialized databases relevant to data from particular research communities (e.g. DNA sequence and genomic data). The disadvantages of centralization often outweigh, however, the benefits of managing encyclopedic digital resources on biological or ecological units of interest, such as living cells or species. There are, however, strong, parallel interests in maintaining data warehouses to leverage the vast amounts of available biological data towards, for example developing ecosystem and whole cell models (Karr et al., 2012; Gauthier et al., 2019; Teschke et al., 2022).

In the domain of biological data categorized as species observations, associated vouchers and public repositories, the Global Biodiversity Information Facility (GBIF; www.gbif.org) plays a central role in guiding data discovery and use (Heberling et al., 2021). GBIF functions as a warehouse of these classes of biological data (Lane &

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Edwards, 2007), a central registry and aggregator whose application of Darwin Core (DwC) data standards supports efficient transformation into accessioned data sets for research, from taxonomic revisions to ecosystem studies at various geospatial scales. The associated international, open-access biodiversity data network and infrastructure provide open access to biological data as an integrated resource, whose contributions necessitate the broad application of best practices throughout the community.

Maintaining data aggregations requires technical support for unique, persistent and actionable identifiers for data and data sets. Although data providers aim to present reliable and traceable data, this challenge is addressed by tools such as the Integrated Publishing Toolkit (IPT), which guides the interpretation of potential GBIF data sets as DwC archives while supporting the publication of accessioned data sets with rich metadata (Edwards, 2000; Robertson et al., 2014). GBIF is a distributed facility, comprising a network of participant nodes and its corresponding network of systems and data managers. Strategic changes in data pipelines are intended to expand this network through the coordinated deployment of new features across the biodiversity data publishing community (Martin et al., 2020).

3.2 Biodiversity databases - potential

Coherence and persistence of these databases represent the intention of scientists to anchor reproducibility to their analyses and establish comprehensive knowledge bases relevant to core research areas within their disciplines. Standards for data storage and exchange are meant, in this context, to increase interoperability among datasets and reduce barriers to contributions and effective use of data by stakeholders. Additionally, guidelines for applying fundamental characterizations to data sets, like taxonomic determinations and geospatial descriptors, become increasingly important elements of systems operations.

The most promising methods underscore the importance of facilitating even higher levels of interoperability across domains, such as establishing persistent identifiers (Güntsch et al., 2017), reinforcing the fundamental role of biodiversity databases as discipline-specific resources. A long history of publishing scientific research is increasingly complemented by data publication, exemplified in the FAIR (Findability, Accessibility, Interoperability and Reuse) and CARE principles (Wilkinson et al., 2016; https://www. go-fair.org/fair-principles/; https://www.gida-global.org/ care). In this way, modern biodiversity databases are managed with a greater focus on the democratic responsibility of research communities to provide access to data while opening possibilities for scientific investigations spanning multiple domains. Improving deficiencies in metadata can be a major step towards establishing FAIR data practices by increasing data discovery across disciplines and between data silos, such as individual peer-reviewed journals (Contaxis et al., 2022).

This transformational perspective characterizes GBIF data, where cross-disciplinary use spans all biological sciences, well beyond its original focus on species occurrences essential to ecology (Heberling et al., 2021). Considering the unprecedented attention of GBIF in its formative years on mobilizing biodiversity informatics data from public collections, this discipline-specific data have been aggregated in a format that emphasizes transformation and subsequent data repurposing.

Guidelines for generating GBIF metadata to promote data discovery have expanded to include harmonization identifying the publishing institutions, specimen repositories or collections and individual data set contacts. This is a function of the continuous work within this community that reaches beyond the technical infrastructure into external development having a significant impact on the data available to users. In this way, GBIF exemplifies how the potential of biodiversity databases still depends on discipline-specific influences for their establishment but is positively affected by community efforts to incorporate incentives to improve interoperability.

Biodiversity databases – power

Biodiversity data are rapidly increasing in magnitude and scope (König et al., 2019), which allows it to be leveraged to answer increasingly difficult questions in ecology and evolution while elevating its status within disciplines as critical specialist resources. Management strategies for these discipline-specific biological databases are shifting towards enhanced interoperability of systems supporting effective data pipelines. Archives of biological data are also shifting towards repositories under various levels of the federation, with best practices promoting data persistence in technical frameworks supporting accessibility and reuse.

Biodiversity databases are heterogeneous with respect to data origin and underlying motivations for compiling the data. They are, however, coherent by design with the research communities defining circumscriptions for included data, generating biodiversity data and guiding its long-term management as its primary stakeholders. Successful community management of discipline-specific data can result in greater levels of access through, for example a micro services-centric perspective (Sheffield et al., 2022), particularly when status as transparent, reliable resources has been established.

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Due to its focus on data interoperability, GBIF is well positioned to lead the community in a shift from domain-specific databases to data hubs for biodiversity informatics. The concept of the Digital Extended Specimen, or DES, describes the emerging paradigm in which specimen data from natural history collections are digitally linked to ecological, environmental and refined biological data from other domains (Wilkinson et al., 2016; Macklin et al., 2022). Outside the context of GBIF, however, motivation to normalize biodiversity data and support DES development is low when the emphasis is on sharing results from analysis or data aggregations rather than sharing the underlying data.

FAIR principles lead instead towards technical implementations handling community-supported data accession procedures as well as persistent identifiers for data and data types (Stocks et al., 2016; McMurry et al., 2017). GBIF publication protocols promote data standardization so it is discoverable and interpretable in a comparative context, allowing users, for example, to apply search criteria across individual datasets. Capturing variation among datasets as structured metadata, such as environmental sampling methods, greatly improves rigour in critical post-discovery tasks, like vetting data.

Heberling et al. (2021) presented comprehensive analyses demonstrating how biological data published by GBIF enables science in research areas beyond biology. Topics of the study emphasized the diversity of data used within disciplines, with interdisciplinary research successfully mediated by data standardization and integration. The authors also recognize the importance of interoperability as a guiding principle in future efforts by GBIF in support of access and reuse of biodiversity informatics data. Removing data from silos of origin improves the efficacy of attempts to identify biases and gaps in global data sets and achieve levels of taxonomic and geographical data coverage in analyses that would otherwise be impossible (König et al., 2019). Decreasing levels of heterogeneity among datasets improve the ability to find, access and combine data, securing its reuse in further scientific analysis and its long-term availability (Teschke et al., 2022).

3.4 Biodiversity databases – conclusions

Biodiversity databases constitute an important component of studies in ecology, evolution and ecosystem dynamics – securing access to data is considered a necessary precursor to, as well as an expected result from research programs in these fields. As a target data repository in this domain, GBIF is a central registry and data aggregator whose application of community standards allows biodiversity data to be published as fit-for-use data sets. Assessing the impact

of GBIF data has shifted from descriptive metrics, such as record totals and taxonomic scope, to measuring interoperability with other user domains and systems of biological data. The increasing magnitude of data published by GBIF will continue to make impressive gains in the upcoming years, considering the growth of citizen science platforms that report species observations. Leveraging the power of available biological data will be a major challenge for GBIF, an endeavour that will improve its role as a data publisher and data hub, focused on interoperability to improve the potential for novel data use.

3.5 | Acknowledgements

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4 | UNDERSTANDING BIODIVERSITY CHANGE - THE POTENTIAL OF COMBINING GENOMIC DATA AND TAXONOMIC CHALLENGES LIKE CRYPTIC SPECIES

Torsten H. Struck.

In the last two decades, biology is experiencing a paradigm shift driven by technological advances. Like physics before, biology is changing into big data science on different fronts. Large amounts of data are generated by remote sensing technology, high-throughput imaging technologies, large databases for environmental parameters and so forth and are accessible to big data analysis in biology (Devictor & Bensaude-Vincent, 2016; Leonelli, 2019). With high-performance computing and tools like deep learning and artificial intelligence, we have the analytical power at hand to analyse such large-scale data in reasonable times and sensible ways (e.g. Li et al., 2019). One such example is sequencing technologies, which have changed tremendously allowing the generation of sequence data of several times the human genome in just a few days or hours (e.g., Lang et al., 2020). Moreover, some of the sequencing and other molecular biological technologies such as MinION have become so small that one can take them out into the field directly (Pomerantz et al., 2022). This allows molecular biomonitoring on the spot instead of transporting the samples back to the lab for this kind of analysis. Pomerantz et al. (2022) provided a protocol and a list of necessary equipment, which allowed species identification in situ based on a metabarcoding approach. This included DNA isolation, amplicon sequencing of barcodes using MinION and downstream data analysis without reliance on Internet connectivity and could be conducted in less than 10h. This changes the way fieldwork can be done in biology and allows a much faster reaction time to, for example adjust a study design. For example, if a rare species, which is not often detected, or a new invasive species, which has not been recorded from that region before, is detected this way in a bulk collection of species (e.g. a Malaise trap) in a very remote region, one can react to this directly on site and adjust the sampling design to take more similar samples. These can also be analysed directly on the spot or taken to the laboratory back home depending on available time. In former times, the bulk samples would have been taken back to the lab, sorted and the species identified there. If one would like to follow up on how abundant that species actually is one might have to go back to the remote region the next year or even only a couple of years later depending on possibilities. A consequence of these new possibilities might be that scientific excursions are planned differently in the near future including time allocations also for spot data generation and flexible slots for tailored extended sampling. Srivathsan et al. (2019) showed the potential of the combination of metabarcoding with high-throughput sequencing. In their study, they caught 7059 specimens in one Malaise trap placed for 8 weeks in Kibale National Park, Uganda. With an accuracy of 99.99% and 0.46% of undetermined nucleotides in the barcodes, they could assign specimens to a total of 650 phorid species, which exceeded the number of phorid species currently described for the entire Afrotropical region. Ninety per cent of the detected species belong to the neglected, megadiverse genus Megaselia (Srivathsan et al., 2019). Hence, the approach has tremendous potential. However, metabarcoding itself has some disadvantages such as primer fit and dropout, and other amplification biases, which can be overcome by metagenomics approaches (Obiol et al., 2020). Using metagenomics would also simplify procedures further to some degree as no amplification step is necessary. On the other hand, computational demands might increase depending on the quality of the generated data. Another requirement for metagenomics approaches is the availability of a sufficient and representative amount of reference genomes for all possibly collected groups.

Fortunately, also in this respect, the new sequencing technologies brought substantial progress. Nowadays, it is possible to sequence high-quality genomes for different plants, fungi, protists and animals including non-model species and species with very large and complex genomes, with very small body sizes, with high ploidy levels or with low-quality preservation status (Cerca et al., 2022; Kingan et al., 2019; Kwiatkowski et al., 2021; Martín-Durán

et al., 2020; Meyer et al., 2021; Schloissnig et al., 2021; Schneider et al., 2021; Varney et al., 2022). This has led to the proposal of a biological moonshot mission, the sequencing of a reference genome for each known eukaryotic species on Earth (Lewin et al., 2018). Since its proposal, this endeavour has gained enormous momentum and support with already several large consortia being established and often funded (e.g. see Table 1). The advantage of genomic sequences for our understanding of ecosystem functioning, biodiversity assessment and conservation biology does not only rest in the advantages of metagenomics but much more importantly in a more detailed understanding of the evolutionary, population genetic and demographic history of the investigated species (e.g. Brandies et al., 2019). For example, studies of the Atlantic cod revealed that besides geographic divergences, there are also clear genetic differences between migratory and non-migratory cod populations on the US and the European Atlantic coast (Berg et al., 2017). These differences are associated with major inversions in four linkage groups, possibly chromosomes. These inversions comprise hundreds of genes, which facilitate the coevolution of complex behavioural traits. The knowledge of the inversions allows for an efficient and reliable ecotype identification and hence monitoring of ecological diversity within and among harvested cod populations. Another example from conversation biology is the Iberian lynx (Abascal et al., 2016). Genomic studies showed a series of severe population bottlenecks within the Iberian lynx populations. These bottlenecks were associated with drastically reduced rates of weak-to-strong substitutions, low genome-wide genetic diversity with long stretches of regions of homozygosity, multiple signatures of genetic erosion and a high frequency of potentially deleterious variants. Moreover, the genomic data allowed the development of genome-wide SNP markers for monitoring (Kleinman-Ruiz et al., 2017). These markers resulted in improvements in the identification of individuals, assignment of parentage and estimation of relatedness, ancestry and admixture. Hence, more powerful, efficient and flexible tools for the genetic management and non-invasive monitoring of Iberian lynx populations could be used. Finally, genome-scale data can also allow better detection of species boundaries. For example, based on only the barcoding gene COI, it was assumed that the earthworm Lumbricus rubellus comprised four cryptic species, but RADseq data revealed the presence of only one species (Giska et al., 2015). Hence, this species is characterized by deeply divergent mitochondrial lineages within populations, which result in genetic differences comparable to interspecific distances between other species. The deep population structure is possibly due to the admixture of lineages from geographically separated glacial refugia,

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TABLE 1 Examples of genome project in the animal kingdom or geographically.

Project name	Main focus	Webpage
Vertebrate Genome Project (VGP)	Vertebrates	https://vertebrategenomesproject.org/
10,000 Bird Genomes (B10K)	Birds	https://b10k.genomics.cn/
Fish 10,000 Genomes (Fish 10 K)	Fish	https://english.cas.cn/newsroom/research_ news/life/201910/t20191008_219736.shtml
Squalomix	Sharks and rays	https://github.com/Squalomix/info/
5000 Insect Genomes (I5K)	Insects	http://i5k.github.io/
Global Ant Genomics Alliance (GAGA)	Ants	https://db.cngb.org/antbase/project
Beenome100	Bees	https://www.beenome100.org/
Global Invertebrate Genomics Alliance (GIGA)	Invertebrates without insects	http://www.gigacos.org/
InvertOmics	Lophotrochozoa	https://www.frontiersinevolutionaryzoology. com/research-1
Soil Invertebrate Genome Initiative (SIGI)	Soil invertebrates	https://tbg.senckenberg.de/sigi/
Aquatic Symbiosis Genomics Project	Freshwater and marine species	https://www.sanger.ac.uk/collaboration/aqua ic-symbiosis-genomics-project/
Deep-Ocean Genomes Program	Deep Sea species	https://www.oceandecade.org/actions/deep- ocean-genomes-program/
Earth Biogenome Project (EBP)	Global	https://www.earthbiogenome.org/
Africa BioGenome Project (AfricaBP)	Africa	https://africanbiogenome.org/
Australian amphibian and reptile genomics (AusARG)	Australia – Reptiles and Amphibians	https://researchprofiles.canberra.edu.au/en/ projects/ausarg-australian-amphibian-and- reptile-genomics-initiative-colla
Oz Mammals Genomics initiative	Australia – Mammals	https://ozmammalsgenomics.com/
Canada 150 Sequencing Initiative (CanSeq150)	Canada	http://www.cgen.ca/canseq150
California Conservation Genomics Project (CCGP)	California (USA)	https://www.ccgproject.org/
Illinois EBP Pilot	Illinois (USA)	https://blogs.illinois.edu/view/7447/79070152
European Reference Genome Atlas (ERGA)	Europe	https://www.erga-biodiversity.eu/
Biodiversity Genomics Europe (BGE)	Europe	https://biodiversitygenomics.eu/
Darwin Tree of Life (DToL)	UK and Ireland	https://www.darwintreeoflife.org/
EBP-Norway (EBP-Nor)	Norway	https://www.ebpnor.org/english/
Endemixit	Italy – Endemic species	https://endemixit.com/the-project/
Catalan Initiative for the Earth BioGenome Project (CBP)	Catalonia (Spania)	https://www.biogenoma.cat/en/home/

which did not separate reproductively, large effective population sizes with low migration rates, rapid population expansion after a single transient bottleneck on a large ancestral population or a combination of these.

Genomic data might be very powerful in revealing the difference between cryptic species and deep population structure, but cryptic species in themselves are also of importance in our understanding of ecosystem functioning. With the advent of molecular species identification, it has become apparent that many species, which were only morphologically delineated, are likely to be genetically different species (Bickford et al., 2007). Several papers have shown how erroneously recognized species

can affect analyses of biodiversity estimates and with that ecosystem functioning (e.g. Bálint et al., 2011; Fišer et al., 2018; Pante et al., 2015; Poulin & Pérez-Ponce de León, 2017). This might lead to wrong conclusions concerning the threat level of species or habitats. For example, for African giraffes, it has been shown that it was not one, but four species increasing the endangered species status for some (Fennessy et al., 2016). However, ongoing taxonomic research in recent years has also shown that the recognition of cryptic species can be challenging and potentially misleading. This has recently been related to the fact that the definition of cryptic species was ambiguous as it was based on taxonomic history

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rather than biological properties (Fišer et al., 2018; Struck et al., 2018a, 2018b). Amended definitions of cryptic species separated the species delineation process from the assignment if a species is a cryptic species or not (Fišer et al., 2018; Struck et al., 2018b). Only species for which it could be shown that they are phenotypically more similar than one could expect should be considered cryptic species. This allows a more accurate definition and detection of cryptic species and hence insights into evolutionary processes (Struck & Cerca, 2019, 2022; Struck et al., 2018b). With respect to ecosystem function, climate change and the biodiversity crisis, one evolutionary process is of particular interest. It is the process of morphological stasis. Several cryptic species could be shown not to change at all or much and hence remain unchanged for millions of years (Fišer et al., 2018; Struck & Cerca, 2022; Struck et al., 2018b). On the other hand, this also means that these species experienced quite substantial environmental changes already and hence can be used as study systems to understand how species can withstand strong environmental changes seemingly without evolutionary adaptation. For example, the annelid Stygocapitella species complex, while it is globally distributed, consists of at least 12 species, which exhibit only four morphotypes (Cerca, Meyer, Purschke, & Struck, 2020; Cerca, Meyer, Stateczny, et al., 2020). Some of these morphotypes evolved tens of millions of years ago up to 140 million years ago (Cerca, Meyer, Purschke, & Struck, 2020; Cerca, Meyer, Stateczny, et al., 2020). Hence, while the dinosaurs went extinct the morphotype of S. pacifica, S. furcata and S. australis remained unchanged and identical even though S. pacifica and S. furcata occur in the Northern hemisphere and S. australis in the Southern one. For the Northern Atlantic species S. subterranea, S. westheidei and S. josemariobrancoi, it could be shown that recent gene flow does not play a role in the maintenance of their identical morphologies (Cerca et al., 2021). However, shared ancestral polymorphism and standing genetic variation could potentially play a role. Hence, cryptic species in general could be ideal systems to understand stasis in evolution and the responsiveness of species to ecosystem changes (Fišer et al., 2018; Struck & Cerca, 2019, 2022).

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MODEL PARASITES OF MODEL HOSTS? EVOLUTION AND ECOLOGY OF CICHLID-INFECTING MONOGENEAN FLATWORMS

Maarten P. M. Vanhove.

Parasites are the most diverse, abundant and threatened metazoan lifeforms. Despite their critical role in ecosystem functioning, most species are scarcely documented or remain undiscovered (Carlson et al., 2020). Given the assumption all species harbour parasites, and the estimate that most species are parasites, their evolutionary influence on biodiversity is considerable (Windsor, 1998, 2021). Apart from this species richness and evolutionary and ecological importance, other features render parasites excellent target organisms for research into a range of fundamental and applied biodiversity-related topics. Host organisms present their parasites with a living and hence changing niche, which leads to ample opportunities and mechanisms for parasite speciation. However, their potential as speciation models is far from optimally harnessed (Huyse et al., 2005). On a more applied note, they may serve, for example as markers for migration and invasion of their hosts (Gagne et al., 2022).

Their often minute size and 'hidden' lifestyle pose logistical challenges to collecting, identifying and sequencing parasites. For example, genomic work on helminths is hampered by difficulties to acquire them fresh and in sufficient quantities and by a lack of experimental protocols. This delayed advances in helminth genomics compared with the progress made in free-living model organisms (McVeigh, 2020). Sequencing effort is biased towards helminths of medical, zoonotic, veterinary or conservation concern (Poulin et al., 2019). On a practical level, most parasite taxa can indeed hardly be considered established models.

Conversely, cichlid fishes are well-known, wellstudied model organisms in many fields of (evolutionary) biology, including the study of behaviour (Jordan et al., 2021) and (ecological) diversification (Burress, 2015; Salzburger, 2018). To stay within the realm of genomics, it is, therefore, unsurprising, as Parsons et al. (2021) put it, '[c]ichlid biology has a continuing history of benefitting from the latest technical advances in sequencing technology'. The latter authors even underline the potential of cichlids in clinical disease research.

Despite this (disease) model status, cichlids' parasites and other symbionts are scarcely studied and often in a quite fragmented way. A group of cichlid parasites, the monogenean flatworms infecting West African tilapias, have however been proposed as models in the study of parasite biodiversity by Pariselle et al. (2003). This proposal rested on the species richness of both the fish and worm taxon, the close phylogenetic relatedness of a set of ecologically comparable and often sympatrically occurring hosts, the one-host lifecycle and quite narrow host-specificity of monogeneans, the substantial variation in monogenean species richness among host species and the idea that under natural conditions monogeneans exert little selection pressure on their hosts. Now, two decades later, what is the state-of-the-art in the parasitological approach of cichlids as a candidate model for host–parasite interactions? Moreover, how is cichlid parasitology relevant to illustrate the role of systematics in understanding ecosystem functioning?

Case studies from our team's work on cichlid parasitology mainly focus on monogeneans parasitising African tilapias (of high policy relevance in view of their economic importance and their invasiveness), and cichlids from the African Great Lakes, 'natural experiments' well known in evolutionary biology but hardly ever studied in the context of an explicit link between host evolution and parasite diversity. While these lakes are well known for their species flocks in various taxa, resulting from radiation events, cichlid-infecting monogeneans belonging to Cichlidogyrus Paperna, 1960 provided the first known case of parasite radiation in any of the African Great Lakes (Vanhove et al., 2015). What is more: these lineages of Cichlidogyrus, sequenced by Vanhove et al. (2015) from the gills of tropheine cichlids in Lake Tanganyika, appeared more species-rich than their hosts. Indeed, as had already been shown for other dactylogyrid monogeneans infecting cichlids and non-cichlid freshwater fishes, the co-phylogenetic analysis indicated that within-host speciation often led to several species of Cichlidogyrus infecting a given host species. Much less common was the proposal of co-speciation as an important diversification mechanism in these dactylogyrids. It soon became clear, however, that not all cichlid lineages in Lake Tanganyika harbour such species-rich monogenean assemblages as the littoral tropheine cichlids, whose parasites belonging to Cichlidogyrus exhibit a quite narrow host specificity. Indeed, Kmentová et al. (2016, 2021) showed that the members of Bathybatini, cichlids from the (deep) open waters of Lake Tanganyika, share a single species of Cichlidogyrus, Cichlidogyrus casuarinus Pariselle et al., 2015. They proposed that the lower density of non-littoral hosts explains the reduced host specificity of this parasite. This exemplifies how host ecology influences parasite diversification processes. Moreover, the broad host range of C. casuarinus was already suggested together with the species' morphology-based description (Pariselle et al., 2015), illustrating how morphological systematics may underpin hypothesis building in ecology.

Knowing that there is no single pattern in parasite speciation in the context of host radiation throughout the Tanganyika cichlids, the question arises of how radiation phenomena in cichlids influence host–parasite combinations at an African scale. Applying various forms of

network analysis on all available infection data for members of Cichlidogyrus (including the nested Scutogyrus Pariselle & Euzet, 1995), Cruz-Laufer, Artois, et al. (2022) point to the role of the evolutionary history of the host in determining the host repertoire of these cichlid parasites. Cichlid radiations in a more stable environment seem conducive to more specialization in host-parasite metacommunities. The authors also show the host environment to be an important predictor of these cichlid-monogenean interactions, with host-switching more likely between ecologically similar hosts. This role of ecological opportunity has implications for aquaculture-environment interactions: it can contribute to the invasive potential of parasites that are co-introduced with the countless tilapia populations that have been anthropogenically translocated. As tilapia-infecting monogeneans may be (on their way to become) the most widespread tropical fish parasites worldwide (Shinn et al., 2023), the importance of a better understanding of these possibly co-invasive parasites can hardly be overstated.

Unfortunately, parasitological aspects of the consequences of tilapia introductions are poorly investigated (Deines et al., 2016). The fact that in Africa and other tropical regions (fish), parasite diversity is disproportionately understudied (e.g. Poulin et al., 2019, 2020) exacerbates this research gap. For want of baseline data, it is difficult to establish whether tilapia parasites in a region that underwent tilapia introductions are there as a result of co-introduction (potentially followed by transmission to native fishes) or are naturally present in the area in the first place. A potential solution is the use of historical museum collections to establish a pre-introduction baseline of parasite communities. This approach allowed Jorissen et al. (2020) to propose certain changes in the monogenean gill parasite fauna of cichlids in Central Africa to be a result of tilapia introductions. Their study showcases the promise that biodiversity infrastructure and in particular host collections hold for advancing parasite ecology (Wood & Vanhove, 2023) and disease research in general (Thompson et al., 2021) - a notion that garnered a lot of scientific and media attention in the context of COVID-19. As collection-based parasitology largely hinges on morphological parasite identification, it proves the importance of systematics in tackling timely ecological questions (Wood et al., 2023).

Even though the work by Jorissen et al. (2020) demonstrated that morphology-based taxonomy allows species-level identification of most of these gill-infecting worms retrieved from cichlids, looking at invasion biology through a parasitological lens gives rise to questions that require genetic methods. For example, even when parasite species shared between introduced tilapia and native fishes are native to the area, there may still be 'hidden transmission' or 'cryptic invasion' of conspecific parasite strains that

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cannot be discerned morphologically. Conclusively determining whether native and introduced parasite strains differ, and whether any of these strains have been transmitted between native and introduced hosts, requires a genetic approach. Indeed, a barcoding approach using a fragment of the COI gene allowed Jorissen et al. (2022) and Geraerts, Huyse, et al. (2022b) to identify such transmissions of alien strains (of native monogenean species) to indigenous host species (and in the latter study, also spill-back of native parasite strains to non-native hosts).

The resolution of such genetic approaches can also be increased to the extent that parasite genetics complement knowledge on their hosts, in this case on tilapia introduction history. Indeed, the magnifying glass hypothesis predicts that parasites because their generation time is shorter than their host's, shed additional light on the evolution and biogeography of their hosts (Nieberding & Olivieri, 2007). Geraerts, Huyse, et al. (2022a) demonstrated this using the highest-resolution markers currently available for these cichlid-infecting monogeneans: mitochondrial genome sequences. In particular, for Cichlidogyrus thurstonae Ergens, 1981, the highest intraspecific diversity was found in the same part of Africa based on the results of COI barcoding of this parasite (Jorissen et al., 2022) and RAD-sequencing data of its Nile tilapia host (Geraerts, Vangestel, et al., 2022). Moreover, the intraspecific genomic differentiation was, between rivers in the Congo Basin and between African countries, several times higher for this monogenean than for Nile tilapia, confirming the magnifying glass potential (Geraerts, 2022). Importantly, of four tilapia-infecting dactylogyrid species whose mitogenomes were sequenced, C. thurstonae was the only candidate to test the magnifying glass hypothesis, as it was the only one with a clear geographical genetic structure (Geraerts, Huyse, et al., 2022a). Its host specificity probably precluded other cichlid species to act as hosts and cloud the picture. It is remarkable that patterns of genetic diversity markedly differ between closely related monogenean species parasitizing the same hosts. This should again be an argument for the importance of thorough systematic studies before embarking on ecological hypothesis testing. Indeed, as co-infections are common, sequencing without prior worm identification would render it much less straightforward to discern patterns and processes in parasite populations.

As an example of recent work in which systematic, genetic and ecological understanding of this cichlidmonogenean model fed into each other, Cruz-Laufer, Pariselle, et al. (2022) provided an updated multimarker phylogeny of the lineage of Cichlidogyrus and Scutogyrus. Using this reconstruction in combination with machine learning, they assessed the diagnostic value, and use, of morphological characters and host repertoire data typically

used for phenotyping and classifying these monogeneans. It is hoped such work provides inspiration on how to improve the identification and systematics of understudied animal taxa and shows why parasite taxonomy should not be allowed to become a 'lost art'. Indeed, since the discovery of new species and new host-parasite combinations in the cichlid-Cichlidogyrus system still seems to be in its exponential stage (Cruz-Laufer et al., 2021), taxonomic and faunistic research on these animals will remain crucial in the foreseeable future. In view of the economic and ecological importance of cichlids, especially in the Global South, (capacity building in) morphological diagnostics of their parasites is a crucial step in acquiring baseline data to assess the health of aquatic environments and how it is anthropogenically impacted (Kapepula Kasembele et al., 2023).

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EXPLORING MARINE **INVERTEBRATE BIODIVERSITY:** CRITTERBASE - A ROADMAP TO THE SOLUTION OF THE FAIR **BIODIVERSITY DATA CHALLENGE**

Dieter Piepenburg, Thomas Brey, Jennifer Dannheim, Katharina Teschke, Jan Beermann, Rebecca Konijnenberg, Hendrik Pehlke, Paul Kloss.

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In times of rapidly increasing climate change (Doblas-Reyes et al., 2021; Gutiérrez et al., 2021) and multiple anthropogenic impacts on marine ecosystems and biodiversity (Meredith et al., 2019; Constable et al., 2022), sound scientific investigations to enhance understanding, sustainable-use management and protection of biotas are of major importance (Wassmann et al., 2011). Research on marine organisms, populations and communities, and their interactions with each other and the environment, is fundamental in this context, but available data are still diverse and scattered. Data on marine biota exist in many formats and sources, such as published literature, data repositories and unpublished materials (Costello et al., 2018). Because of this heterogeneity, information can be difficult to find, access and combine, severely impeding its reuse for further scientific analysis and its long-term availability for future generations. Therefore, scientists, decision-makers and the public require versatile tools to compile, synthesize and manage biodiversity data in a transparent, efficient and comprehensible way with high-level quality assurance (Teschke et al., 2022).

To address this challenge, we developed, implemented and utilize CRITTERBASE (https://critterbase.awi.de; Teschke et al., 2022), a publicly accessible data warehouse and interactive portal that complies with the FAIR data principles (Findability, Accessibility, Interoperability and Reusability). The purpose of CRITTERBASE is to complement (not substitute) existing long-term data storage repositories, such as the Ocean Biodiversity Information System (OBIS) or the Data Publisher of Earth and Environmental Science (PANGAEA), through providing a versatile platform facilitating management and analysis of verified biodiversity data across multiple spatial and temporal scales and

in various contexts (research, governance, management and public information). To this end, CRITTERBASE features an operational modular data management environment including powerful and easy-to-use data ingest, retrieval and exploration options (Collector App, Data Space, and Analyst App, respectively; Figure 1) for handling sample-based organism-related data from marine environments on a global scale.

The Data Space contains a data model that covers a variety of biotic data types (e.g. presence/absence, abundance, biomass), sub-sample processing options and sampling methods and can be expanded easily. The data ingestion process via the Collector App includes a thorough quality check, for example through the validation of taxonomic names against the current classification provided by the World Register of Marine Species (WoRMS) (Figure 1). Through its web interface hosted by the Computing Centre of the Alfred Wegener Institute (AWI), Helmholtz Centre for Polar and Marine Research Bremerhaven, CRITTERBASE is open to the public. Direct machine-to-machine communication with the web service of CRITTERBASE is also possible through a REST interface to allow for software-based data queries.

The Analyst App, currently under development, already features a key data extraction tool, aiding users with filtering options (by region, time frame, gear and data set) to identify and download exactly the data they are looking for. These filter options will be expanded to include, for example the search for specific taxa. Moreover, it will also provide basic data exploration applications by means of which, for example the distribution of data values, the presence of outliers and relationships between different data variables can be explored to provide data users with greater insight into the raw data.

COLLECTOR APP DATA SPACE ANALYST APP

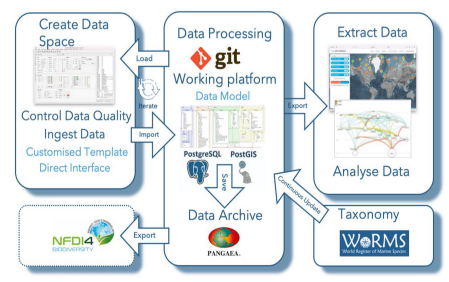


FIGURE 1 Modular structure of the CRITTERBASE data warehouse.

A simple use case demonstrates an example of basic data exploration options by means of CRITTERBASE in combination with R (v4.2.0; R Core Team, 2022; RStudio Team, 2022). To examine habitat preferences of six North Sea polychaete species for certain sediment grain sizes (Armonies, 2021), R packages DBI (v1.1.3; R Special Interest Group on Databases (R-SIG-DB) et al., 2022) and RPostgres (v1.4.4; Wickham et al., 2022) are used to query data from four CRITTERBASE data sets with a total of 143 grab samples, containing species abundance data and information about median sediment grain size at sampling stations. By means of R package ggplot2 (v3.4.0; Wickham, 2016), a scatterplot of the species's abundances (individuals m⁻²) versus sampled sediment grain sizes is produced (Figure 2), visualizing their preference for fine sand environments. In addition, the R package ggOceanMaps (v.1.3.4, Vihtakari, 2022) is used to produce maps showing the geographic location and number of the stations, where the species were sampled (Figure 3). The use case exemplifies the ease and simplicity with which basic data exploration can be conducted by tying R to CRITTERBASE, to visualize with only a few R commands distributions and relationships of the queried

data and identify a potentially interesting area for further investigation.

The use case also highlights the role of user choices in data query, analysis and interpretation (Teschke et al., 2022). For instance, not all CRITTERBASE data sets contain the same type of biotic information. While some of the four data sets queried feature standardized abundance data (individuals m⁻²), others provide only the total number of individuals alongside information about the sample area (m²). The user must choose how to address these differences and run any necessary standardizations. In this use case, we also did not discriminate among life stages, meaning the number of samples given in Figure 3 differs slightly from the number of actual data points in Figures 2 and 3, as some stations contained two or more entries of the same species but for a different life stage. Moreover, some stations were also sampled repeatedly. Users must carefully evaluate whether further data refinement is needed. Lastly, Figure 3 underscores that users should also consider whether they can interpret a lack of data to present true absences. In this use case, the number of grab samples differed between species, as not all species were present at each station (Figure 3). A detailed

Density of various Polychaete species versus median sediment grain size

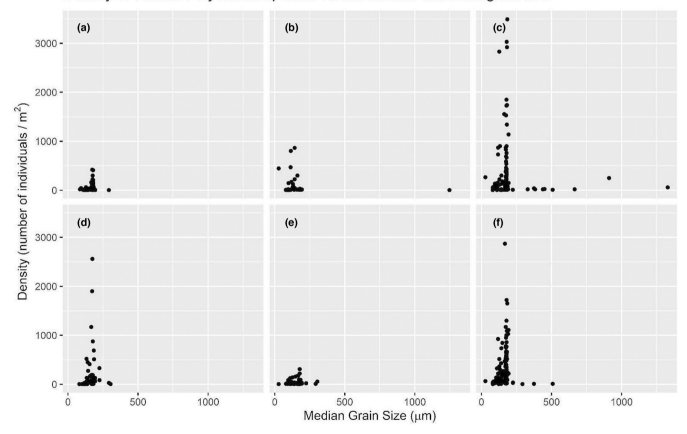


FIGURE 2 Scatterplot showing abundances (individuals per m⁻²) of six polychaete species in the southern North Sea in relation to the median grain size of surficial bottom sediments (μm). (a) Chaetozone setosa, (b) Lagis koreni, (c) Lanice conchilega, (d) Magelona johnstoni, (e) Scoloplos armiger, (f) Spiophanes bombyx.

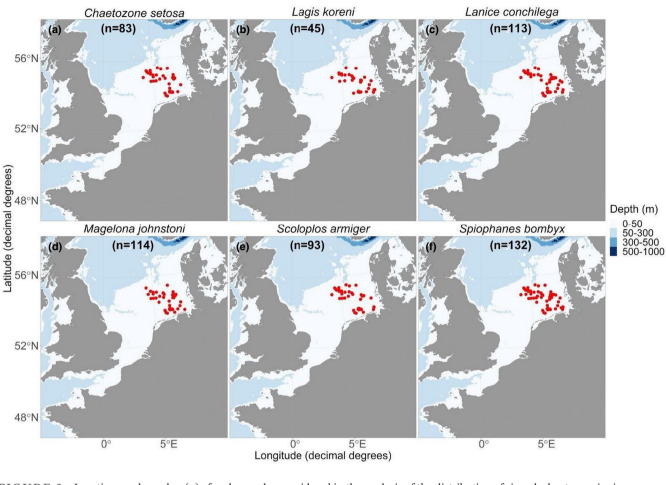


FIGURE 3 Locations and number (*n*) of grab samples considered in the analysis of the distribution of six polychaete species in relation to sediment type in the southern North Sea (Figure 2). Produced using R package ggOceanMaps, the bathymetric data contained therein (Amante & Eakins, 2009; GEBCO, 2021; EMODnet Bathymetry Consortium, 2020) and the continent boundary vectors (Natural Earth, 2022).

examination of metadata provided in CRITTERBASE can help to decide whether the lack of data of a species at certain stations could be interpreted as true absences.

At the moment, CRITTERBASE contains mostly benthic biodiversity data (a total of 414,666 records of 4550 taxa identified in approx. 30,000 samples taken at 19,900 stations by means of grabs, trawls and underwater imaging in the Arctic, North Sea and Antarctic). However, its comprehensive and flexible data model allows for handling geo-referenced field data on any type of marine biota (benthos, plankton, nekton, etc.) and any type of sampling technique and gear. Its connection to the Research Data Commons (RDC), a cloud-based data infrastructure that is being developed under NFDI4Biodiversity, a consortium under the umbrella of the German National Research Data Infrastructure (NFDI), will help make CRITTERBASE the core of a national information system for biodiversity data on marine organisms. In the future, CRITTERBASE will also feature interfaces for data exchange with long-term repositories such as OBIS and PANGAEA. We envision

CRITTERBASE to become a valuable and continuously expanding tool for a wide range of usages, such as studies of spatiotemporal biodiversity patterns, impacts and risks of climate change or evidence-based design of marine protection measures.

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7 | CRITICAL TAXONOMIC CONTRIBUTIONS FOR METABARCODING STUDIES

Paula Pappalardo, Karen J. Osborn

Assigning taxon names is essential for biodiversity research and conservation (Mace, 2004). A detailed morphological assessment is a traditional approach to identifying an organism to the lowest possible taxonomic level. The morphological study of an individual organism has some advantages, such as the potential to identify life stages, but it requires high taxonomic expertise and is time-consuming. Barcoding, a cost-effective and fast method for species identification, was developed to help identify species and cryptic diversity. Barcoding for identification uses a DNA sequence for a specific genetic marker from the sample in question and compares it to known sequences in a DNA reference database, hoping to match the unknown sequence to a known one. A natural extension, made possible by recent sequencing technology, is metabarcoding, where multiple organisms from a mixed sample (e.g. insect trap, plankton net) can be identified simultaneously in this same manner. When the sample is environmental (e.g. water, sediment), metabarcoding techniques are known as eDNA. Metabarcoding can facilitate rapid, large-scale biodiversity sampling of diverse environments, for example monitoring for invasive species. But the list of organisms found using metabarcoding is affected by multiple factors, and the reliability of the taxonomic identifications depends heavily on the completeness and accuracy of the reference database.

Taxonomists have two key roles in improving taxonomic assignment for metabarcoding: (1) identifying understudied organisms and contributing high-quality barcode sequences for those organisms to DNA reference databases and (2) helping to curate publicly available DNA reference databases and report taxonomic issues. Ideally, a sequence included in a DNA reference database will have an associated voucher so that it can be reexamined if there is a taxonomic conflict. BOLD (Ratnasingham & Hebert, 2007), one of the first barcode reference databases, was built with this in mind and it even requires images to complete a specimen record. Other large sequence repositories such as GenBank do not require an associated voucher to publish a genetic sequence but do have an identifier available to provide voucher numbers when submitting sequences. In addition, for some institutions (e.g. Smithsonian National Museum of Natural History) GenBank can link the voucher number with the institution's online catalogue for additional information such as collection and handling metadata. In cases, where specimens are very small and have been fully utilized for DNA extraction, storage of images and DNA extractions help with future efforts to re-examine the identification if needed. Large barcoding efforts using museum specimens (when feasible) have proven useful not only to generate DNA barcodes associated with vouchers but also to make available the associated metadata (Hebert et al., 2013).

Regardless of how inclusive DNA reference databases are, there are large gaps in representation documented for many taxonomic groups, even bioindicator species that are used to assess aquatic ecosystem quality (Weigand et al., 2019). The gaps in representation are usually larger for invertebrates than vertebrates and are pervasive even for the most commonly used genetic markers. We used the MIDORI2 reference database (Machida et al., 2017; Leray et al., 2022) to showcase how even for COI, the genetic marker with the largest number of available sequences, there is highly variable species representation (Figure 4). Species representation can be as low as 0.3 (for Nematoda using the maximum richness estimates) or as high as 100% (phylum Cycliophora with just two known species). This means that taxonomists can have a big impact if they contribute reference sequences, particularly those of traditionally understudied taxa.

In our experience, a small group of researchers can generate a large number of novel barcodes in a short time (Pappalardo et al., 2021). The Smithsonian StreamCode project, which assessed zooplankton biodiversity in the Gulf Stream during a single field season (2 weeks, 14 taxonomists, 12 staff and students) contributed 2150 sequences (Pappalardo et al., 2021). At the time of submission, 109 were novel sequences to GenBank at the species level (43 for 18S and 66 for COI), 99 were novel at the generic level (33 for 18S and 66 for COI) and 44 were novel at the family level (11 for 18S and 33 for COI) (Pappalardo et al., 2021). These sequences were compiled to create a local reference database that outperformed other databases in the taxonomic assignment of metabarcoding samples of zooplankton to phylum (Pappalardo et al., 2021). Similarly, other initiatives involving taxonomic experts and citizen scientists to combine intense biodiversity surveys of a specific area (in a short amount of time) with DNA barcoding and specimens' vouchers (e.g. https://www.abol.ac.at/en/ abol-bioblitz-2019/) are an effective way to populate reference databases.

There are multiple DNA reference databases available – they differ in the genetic marker, which taxonomic groups are included, the source of the sequences, the taxonomic backbone, curation level, submission standards and the

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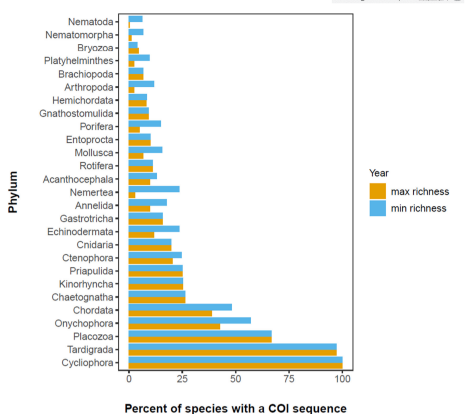


FIGURE 4 Representation gaps for COI at the phylum level. Percent of species that have a COI sequence in the reference database MIDORI2 based on minimum (light blue) or maximum (orange) estimates of species richness (when a range of species richness estimates was available). Data on species richness were extracted from table 2 in Machida et al. (2017). We revised the two phyla reporting 1 species and updated them to 2 for Cycliophora (based on the species reported in the World Register of Marine Species, https://www.marinespecies.org/aphia.php?p=taxdetails&id=22586 on 2022-10-31) and 3 for Placozoa (Osigus et al., 2019). Counts on unique species per phylum for the COI genetic marker were calculated using data from MIDORI2 version GB249 downloaded on 13 May 2022 from the MIDORI server (http://www.reference-midori.info/download.php#).

number of sequences (among other variables). Even for a specific marker and taxonomic group, there may be multiple reference databases available. And unfortunately, the choice of reference database does matter. For example, the different family-level composition was recovered when four different reference databases were used to analyse bacterial and archaeal community composition (Robeson et al., 2021). Similarly, in our work, we have seen that the per cent of correct species-level assignments using COI for a test set of zooplankton can vary from 69% to 100% when using a 99% identity cut-off for BLASTn among different reference databases (Pappalardo et al., in prep). In general, an increase in the number of sequences included in the reference database increases the number of correct assignments. For example, we observed that the MIDORI2 'unique' database (Leray et al., 2022), which includes all the unique haplotypes for each species, performed better that the 'longest' version, which only contains the single longest sequence for each species (Pappalardo et al., in prep).

Additionally, the likelihood of a sequence being assigned correctly is also influenced by the presence of errors in the reference database. There are several studies reporting errors in sequences published in GenBank (Nilsson et al., 2006; Leray et al., 2019; van den Burg et al., 2020). Inspecting some of the papers that report errors (Leray et al., 2019; van den Burg et al., 2020), we noticed that in most of the cases, the errors reported are related to taxonomy. These may be cases where similar or identical sequences have different names because of synonyms or taxonomic revisions (e.g. new genus placement). The errors related to taxonomy can be easily fixed if reported because the NCBI taxonomic team (Schoch et al., 2020) can revise and fix them without the need to contact the sequence author. Taxonomic issues can be flagged by writing to the NCBI address info@ncbi.nlm.nih.gov. Consider writing 'taxonomy question' in the subject line to help the issue reach the taxonomic team faster. The taxonomic issues for which we or our collaborators submitted an issue were solved within a week.

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If another type of problem is found with a sequence (e.g. suspected contamination), it can be reported to gb-admin@ncbi.nlm.nih.gov for them to contact the sequence author. Currently, only the submitter is able to modify the identity of a submission. When alerted of a possible error, GenBank can inspect the issue, contact the sequence author and if deemed appropriate, flag the sequence with the tag 'UNVERIFIED'. Sequences with the UNVERIFIED tag do not appear in BLAST searches (https://www.ncbi.nlm.nih.gov/genbank/unverified/). Depending on the response of the submitting author, these errors can also be corrected.

Another issue is that sequence authors can help with keeping their taxon names updated when the taxonomic status changes. Even after publishing a new name, Schoch et al. (2020) reported that most submitters do not update their taxon names. Updates can be reported to gb-admin@ncbi.nlm.nih.gov and in our experience, revision and changes occur quickly.

To fulfil the promises of metabarcoding, we must all work together to improve DNA reference databases. Reference databases can be improved by keeping taxon names updated in GenBank, reporting errors when we find them, and publishing sequences associated with vouchers. Incorporating molecular information with species descriptions could be another way to help expand reference databases (e.g. Osborn et al., 2011; Stoev et al., 2013). Understanding the strength and limitations of the reference database is also very important, and a sensitivity analysis comparing results from two different databases is an effective way to show the robustness of the results. In terms of the choice of genetic marker, not one genetic marker is perfect for all taxonomic groups, and multiple genetic markers can increase the chances of correct species detection. Our general advice for metabarcoding projects is to have an interdisciplinary team, including expertise in taxonomy, bioinformatics, ecology and evolution. We think that the scientific community should encourage training opportunities for the new generation of taxonomists that will use both morphological assessment and molecular approaches for species identification.

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8 | METABARCODING AND SYSTEMATICS OF PROTISTS IN THE OCEAN TO UNDERSTAND ECOSYSTEM FUNCTIONING

Bente Edvardsen

8.1 | Protist diversity

The massive loss of biodiversity due to human activities and climate change is accelerating. We need to know the organisms, who they are, their distribution in time and space and their ecological role in order to be able to conserve biodiversity. We also need to understand the driving forces and processes that shape biodiversity and the community structure to be able to predict the consequences of human activities and mitigate losses. By combining different methods in an integrative approach, a more detailed picture of the community composition, structure, distribution and function can be obtained.

Protists are all eukaryotes that are not animals, plants or fungi and are thus a polyphyletic group. They are mostly tiny, single-celled organisms and are much less studied than animals, plants and fungi, despite that they dominate eukaryotic diversity and play key functional roles in all ecosystems. They span a wide range of sizes, forms and lifestyles and their nutrition can be phototrophic as the algae, heterotrophic consumers as the protozoans, and mixotrophic which is everything in-between, and some are symbionts or parasites living inside other organisms. Phytoplankton, consisting of different algal lineages and cyanobacteria, are primary producers and the basis for most marine food webs and fisheries. They have therefore been studied by light microscopy and experiments for more than a century to reveal production, community composition and seasonal dynamics and relate this to abiotic factors (e.g. Hjort & Gran, 1900, Braarud et al., 1958). Microscopical phytoplankton long-time series data form the basis for analysis to understand changes over time (e.g. Lundsør et al., 2022). Many scientific questions in plankton ecology are the same today as 100 years ago, and we still ask: who are there? Where, when and how do they occur and what are the driving factors? How much is produced, what are they doing and what are their ecological roles? New technologies have steadily emerged and changed the methodologies and expanded the possibilities of what we can discover. DNA-based methods such

as metabarcoding have during the last decades become a common method in microbial diversity studies as it provides high taxonomic resolution and may also detect and identify tiny, fragile and rare taxa with few morphological traits (Lopes dos Santos et al., 2022). Here I will elucidate how different methods can be combined in biodiversity studies of protists in the ocean and discuss challenges and possibilities.

8.2 | Metabarcoding of protists

In metabarcoding, the organisms present in an environmental sample (e.g. sea water, sediment, soil) can be identified by extracting DNA from the sample, amplifying a marker gene of all the targeted organisms, sequencing the obtained DNA fragments by high-throughput sequencing and comparing the sequences to a reference library with sequences of known and identified organisms (see also Burki et al., 2021). The most wide marker gene region used in protist studies is the V4 or V9 regions of the 18S rRNA gene, which is present in all eukaryotes, it has both conserved and variable regions to enable classification from kingdom to species level, and it is the DNA region with most reference sequences available in gene databases, with a curated DNA reference sequence library for a taxonomic assignation, the PR2 (Guillou et al., 2012; Lopes dos Santos et al., 2022). High-throughput sequencing (HTS) with the Illumina Miseq pair-end technology gives the needed sequence length (up to ca 500 base pairs, bp) and quality to separate most taxa to genus and many to species level. Other HTS technologies are emerging that enable long-read sequencing, from 1500 to 5000 bp, and thus higher taxonomic resolution, such as PacBio and Nanopore, that are already used as a complement to Illumina (Burki et al., 2021). Metabarcoding has during the last 15 years revealed an unprecedented diversity and provided massive data on how protist diversity is distributed in space and time and combined with metadata (environmental information) it has contributed to a better understanding of driving forces structuring protist communities (e.g. de Vargas et al., 2015; Massana et al., 2015).

8.3 | Challenges with metabarcoding

Metabarcoding also has some major challenges and limitations. The laboratory step includes a polymerase chain reaction (PCR), where the marker gene is not amplified equally well for all taxa. No primer pair for the amplification of the V4 18S rRNA gene match 100% to the DNA of all eukaryotic taxa (Vaulot et al., 2022). This step may also introduce false diversity by combining sequences

from different organisms (chimeras, Egge et al., 2013). Different bioinformatic treatments may also give different results (Egge et al., 2013). Another challenge is that the sequences with Illumina technology are rather short (about 500 bp) and some species have identical DNA sequences in the marker region (e.g. some diatom species in the V4 region, Luddington et al., 2012). Also, the number of copies of the marker gene in each cell varies, not only by cell volume (a large species usually has more copies than a small species within the same taxonomic group) but also between different taxonomic groups. So, the quantification ability is disputed and at best it gives the relative abundance of a taxon in each sample. Another challenge is the lack of sufficient reference sequences for taxonomic classification.

8.4 | An integrative approach

The ability for quantification by metabarcoding has been tested in several studies, for example by comparing with the traditional microscopical cell counts. To be able to link available quantitative protist time-series and historical data based on light microscopy to future metabarcoding data, we need to know how they compare. Santi et al. (2021) compared the protist composition and relative abundance from marine waters by metabarcoding and microscopical cell counts. They found that some taxonomic groups were represented rather equally with the two methods, such as diatoms, whereas others did not (dinoflagellates and ciliates) and that the differences became larger at low taxonomic levels. They concluded that metabarcoding provided a better estimate of the taxonomic richness of a community, while microscopy provided more accurate quantitative data of abundance and biomass. The study showed that the two methods are complementary, and by using both methods, a more detailed information on taxonomic composition, richness and abundance can be obtained (Santi et al., 2021).

The lack of sufficient DNA reference sequences is presently a major limitation for species identification by metabarcoding. For most protist groups, only a fraction of the described species has been isolated into the culture and been sequenced in some marker gene region. For the algal division Haptophyta, 320 species have been formally described and given a name, but only 100 of these have been sequenced in the 18S rRNA gene (Edvardsen et al., 2016). Environmental sequencing has detected more than 900 different genotypes that may represent species (Edvardsen et al., 2016), and thus the majority of these cannot be identified at the species level by molecular methods. Some haptophyte taxa known only from environmental sequences may even represent novel classes. In a metabarcoding

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study of the haptophyte diversity in Oslofjorden, Norway, genotypes were detected that fell into clades representing three putatively novel classes, one novel order and one genus (Egge et al., 2015).

In order to infer the phylogeny and improve the taxonomy and reference databases of haptophytes, we and collaborators have isolated 30 new haptophyte strains into the culture, sequenced and determined their phylogeny and obtained new haptophyte reference sequences (e.g. Edvardsen et al., 2000, 2011). We have also described several novel haptophyte species (e.g. Eikrem & Edvardsen, 1999; Eikrem, 1996; Seoane et al., 2009). We steadily discover protist species new to science, and there is a strong need to formally describe more protists.

Metabarcoding studies in the Arctic have shown a large unknown protist diversity and a need for more reference sequences (e.g. Egge et al., 2021). In the project TaxMArc, we have isolated more than 200 strains from Arctic waters and ice either by single cell picking under the microscope or by a dilution series. The strains have been sequenced in rRNA genes, identified under the microscope and are now maintained in and available from the Norwegian Culture Collection of Algae (NORCCA, norcca.scrol.net). With cultures, we can obtain both the morphology and long sequences of many genes, which enables the discovery and description of novel species and phylogenetic placement. Šupraha et al. (2022) discovered that many species in the Arctic are morphologically similar to temperate species but are genetically different and may represent novel species or varieties. A search against the 18S rRNA gene database metaPR2 (Vaulot et al., 2022), presently containing 41 metabarcoding data sets from 4000 samples from marine waters, can reveal the geographical distribution where a genotype has been found before in the data sets and under which environmental conditions. By using the metaPR2 database, Šupraha et al. (2022) discovered that several diatom genotypes isolated from Arctic Svalbard seem restricted to the Arctic, whereas other diatom genotypes had either an Arctic temperate or cosmopolitan distribution.

8.5 | Linking taxonomic diversity to functional diversity

In order to link a genotype to an ecological function, we need to know different traits of the organisms such as their size, shape, motility, growth rate, preferred environment for growth, pigments, trophic mode, geographical distribution, etc. With a trait table containing this information on all species and genotypes in a sample, we can obtain information on the ecological functionalities from metabarcoding data (see Martini et al., 2021). Ramond et al. (2019) have compiled and tested a trait table for protists from a few marine localities. However, much work remains to obtain a more complete trait table for marine protist species. With algal cultures available, growth responses to various growth conditions should be screened. Experiments for physiological and ecological traits such as organism interactions, (host-parasite and food-grazing) are needed to be able to better understand how the ecosystems function. Increasing genomic data from protists will also add to a better understanding of physiological and ecological functioning.

Conclusion 8.6

A majority of marine protists have not yet been characterized genetically and morphologically and lack reference DNA sequences. There is a need to isolate cultures, characterize and describe more protist species and provide more reference DNA sequences and expand and curate reference libraries. To be able to link taxonomic diversity to functional diversity, we need more information on different traits (e.g. morphology, ecology, physiology, behaviour) and more comprehensive trait tables. Further, to be able to use metabarcoding in monitoring and link available time series and historical data based on light microscopy to new molecular data, we need to know how they compare. By combining several methods in an integrated approach, a more detailed analysis of the community composition, structure, distribution, ecological function and ecosystem functioning can be obtained.

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