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Correct Species Identification and Its Implications for Conservation Using Haploniscidae (Crustacea, Isopoda) in Icelandic Waters as a Proxy

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Correct identification of species is required to assess and understand the biodiversity of an ecosystem. In the deep sea, however, this is only possible to a limited extent, as a large part of the fauna is undescribed and the identification keys for most taxa are inadequate or missing. With the progressive impact of climate change and anthropogenic activities on deep-sea ecosystems, it is imperative to define reliable methods for robust species identification. In this study, different techniques for the identification of deep-sea species are tested, including a combination of morphological, molecular (DNA barcoding, and proteomic fingerprinting), biogeographical and ecological modeling approaches. These are applied to a family of isopods, the Haploniscidae, from deep waters around Iceland. The construction of interactive identification keys based on the DELTA format (DEscription Language for TAXonomy) were a major pillar of this study, the evaluation of which was underpinned by the application of the supplementary methods. Overall, interactive keys have been very reliable in identifying species within the Haploniscidae. Especially in a deep-sea context, these types of keys could become established because they are easy to adapt and flexible enough to accommodate newly described species. Remarkably, in this study, the interactive key enabled identification of a supposedly new species within the Haploniscidae that was later verified using both molecular genetic – and proteomic methods. However, these keys are limited given that they are based on purely morphological characteristics, including where species with strong ontogenetic or sexual dimorphism occur as both genders are not always described. In this case, integrative taxonomy is the method of choice and the combination presented here has been shown to be very promising for correct identification of deep-sea isopods.

Keywords: Peracarida, deep-sea, taxonomy, interactive keys, COI barcoding, MALDI-TOF MS, depth distribution, geographical distribution

INTRODUCTION

Species determination is the first step in conducting almost any kind of biological research. Without identifying known and scientifically describing newly discovered species, drawing conclusions about species diversity, their distribution and their ecology is impossible. Species delimitation describes the process by which species boundaries are identified and defined involving the identification of (known) species as well as of species new to science (e.g., Wiens et al., 2007). The question of how one species can be differentiated from another is irrevocably linked to the question of “what is called a species?”

Just like in other habitats in the deep sea, defined as depth >200 m (Gage and Tyler, 1991), species are often identified based on their morphological appearance. However, deep-sea sampling is known to be challenging; long hauls over several hours and subsequent sampling and sorting processes can damage specimens and thus conceal important morphological characters. In addition, morphological identifications are limited by the high rate of new species discoveries (Brandt et al., 2007), combined with a high diversity and typically low densities of deep-sea communities. The latter in turn imposes restrictions on the evaluation of intra- and interspecific variations (e.g., Brandt et al., 2007; Lim et al., 2012). Since the deep-sea environment also has a high proportion of morphologically similar but genetically different species, so-called cryptic species (Raupach and Wägele, 2006; Vrijenhoek, 2009; Brasier et al., 2016), the morpho-species approach tends to underestimate true biodiversity and, conversely, to overestimate the range of species and their population size. In addition, intraspecific morphological variability, including ontogenetic variation, polymorphism, or sexual dimorphism, poses a challenge to species assignment based on morphological characters alone (Raupach and Wägele, 2006; Riehl et al., 2012; Zaharias et al., 2020; Paulus et al., 2021).

In the last two decades, major leaps have been made in the development of taxonomic approaches and methodologies, including DNA-based taxonomy and (meta-) barcoding, “-omic” techniques, imaging tools, and integrative approaches linking different types of taxonomic information (Dayrat, 2005; Boistel et al., 2011; Raupach et al., 2016; Paulus et al., 2021). Many of these methods, despite their apparent advantage in facilitating and accelerating species delimitation, have been slow to find their way into deep-sea taxonomy. However, with increasing human impacts on deep-sea ecosystems, new methods and ideas for identifying and delimiting species are urgently needed (Brix et al., 2020).

In this study, we use a relatively well known deep-sea benthic group, isopod crustaceans (Brix and Svavarsson, 2010; Brix et al., 2018), from a relatively well known region, waters around Iceland (Brökeland and Svavarsson, 2017), to apply a range of (morphological and molecular) methods and procedures for the identification of deep-sea species. Situated right at the border between the Northern North Atlantic and the Nordic seas and separated by the Greenland-Iceland-Faroe (GIF) ridge, Icelandic waters provide a very heterogeneous marine environment linked to complex water masses and diverse submarine topography that strongly influence faunal distributions. Additionally, the

fauna is threatened by climate change, including warming, acidification, freshening and productivity changes (Hanna et al., 2006; Arnason, 2007; D’alba et al., 2010; Pecl et al., 2017) which in turn could lead to changes in species diversity and geographic ranges (e.g., D’alba et al., 2010; Pecl et al., 2017). To properly assess these future impacts, understanding where species occur and how they are distributed is imperative, and thus they must be properly identified.

Isopod crustaceans are commonly found in deep-sea benthic communities, and they are also widespread in Icelandic waters (Svavarsson et al., 1993; Brix and Svavarsson, 2010; Brökeland and Svavarsson, 2017; Schnurr et al., 2018). The isopod family Haploniscidae Hansen, 1916 is considered a typical deep-sea family, although members are also found at shelf depth (Brökeland and Svavarsson, 2017; Johannsen et al., 2020). Due to their ubiquity and widespread occurrence in Icelandic waters, Haploniscidae, and the genus *Haploniscus* Richardson, 1908 in particular, are well suited as proxies for assessing species distributions around Iceland.

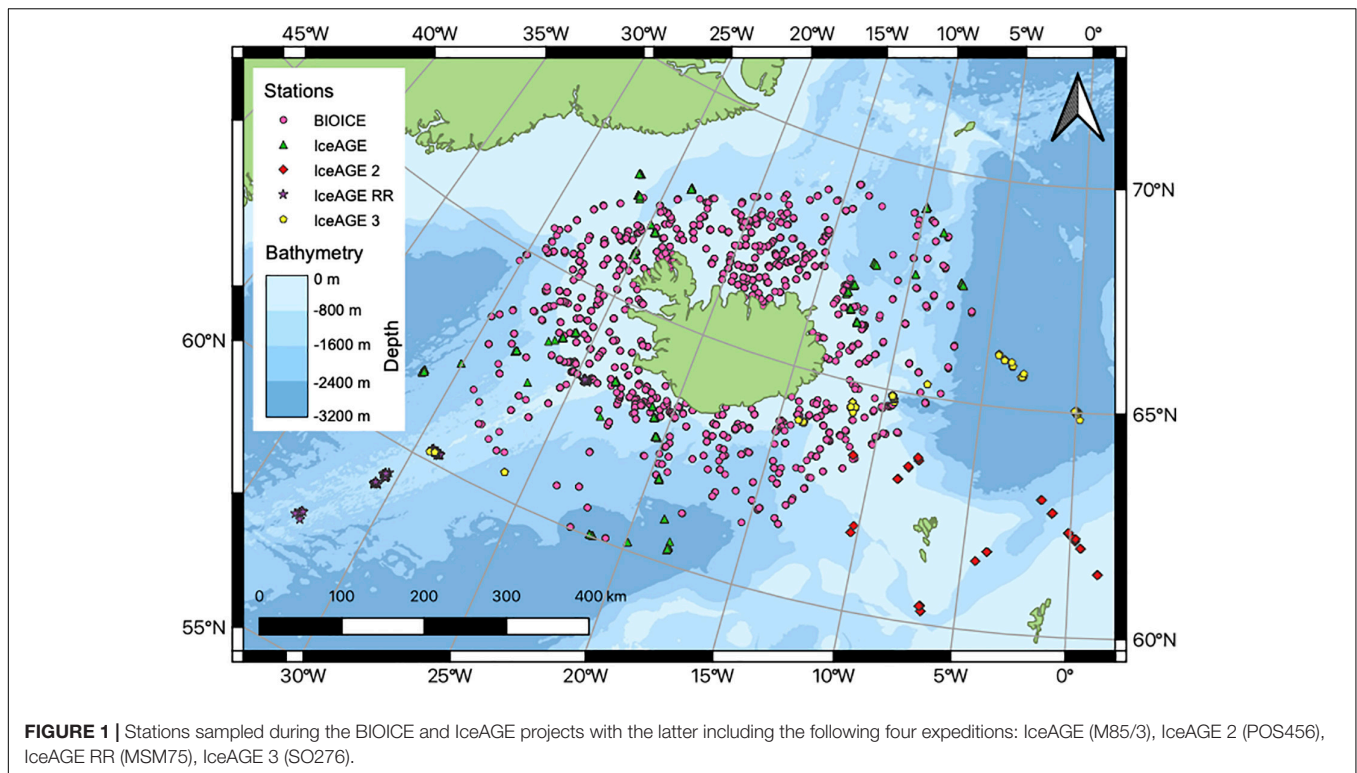
The basis for the current study is provided by sampling carried out in the course of the BIOICE (Benthic Invertebrates of Icelandic Waters) and the IceAGE (Icelandic marine Animals:

Genetics and Ecology) projects leading to a very comprehensive data set that is also suitable for genetic and “omic” studies (Brix et al., 2014; Riehl et al., 2014; Meißner et al., 2018). Morphology-based approaches (interactive identification keys) were used to separate haploniscid genera occurring around Iceland (*Antennuloniscus* Menzies, 1962, *Chauliodoniscus* Lincoln, 1985b and *Haploniscus*) and to identify species within the genus *Haploniscus*. In addition, bathymetric and geographic information for species in the family Haploniscidae around Iceland are compiled as part of an integrative taxonomic framework. These data are complemented by genetic [using Cytochrome oxidase I (COI) as barcoding marker] and proteomic fingerprinting tools to foster species identification. The assumption was that many of the Icelandic haploniscid species are described and can therefore be reliably recognized as such with the interactive key as not many cryptic species are to be expected. That is, we sought to test whether the species identified by the key are “valid,” and what were the limiting factors for correct species identification? This combined approach has the power to overcome problems of taxonomic standardization with data being made publicly available (e.g., via BOLD, OBIS) and enhancing species identification, for instance by fellow taxonomists, para-taxonomists and students, through online-identification keys (Balke et al., 2013).

MATERIALS AND METHODS

Study Area, Sampling and Data Collection

Specimens of the isopod family Haploniscidae were collected around Iceland during BIOICE (Benthic Invertebrates of Icelandic waters) with yearly expeditions between 1992 and 2004 and IceAGE with four expeditions: IceAGE (2011), IceAGE2 (2013), IceAGE_RR (2018) and IceAGE3 in 2020 (**Figure 1**).



For the molecular analyses, only specimens were taken that were collected during the IceAGE, IceAGE2 and IceAGE_RR cruises respectively using an epibenthic sledge (EBS) or a van Veen grab (Brix and Devey, 2019). These samples were fixed in cooled high-grade ethanol to ensure genetic analysis. The material was first examined morphologically and given a unique database number at the DZMB (German Centre for Marine Biodiversity Research, Hamburg). For the species distribution models as well as bathymetry and geological distribution analyses, specimens from BIOICE and IceAGE (IceAGE, IceAGE2, IceAGE_RR, IceAGE3) were utilized.

During the BIOICE project, specimens were collected using a detritus sledge (Sneli sled; Sneli, 1998), a RP sledge (Rothlisberg and Percy sledge; Rothlisberg and Percy, 1976), a triangle dredge and an Agassiz trawl. The RP samples were elutriated through a 0.5 mm sieve, and the remaining sediment and animals were processed through a series of sieves (4, 2, 1, and 0.5 mm). The detritus sledge samples were then processed through two vertically stacked (1 and 0.5 mm) sieves. The Agassiz and triangle dredge samples were hand-picked on board. After the sorting process the BIOICE samples were stored in 5% buffered formalin (Brökeland and Svavarsson, 2017). Samples that were collected during the four different IceAGE expeditions and included in the analyses come from the EBS (EBS, Brenke, 2005) and triangle dredge.

Interactive Identification Keys

Interactive keys were developed for identifying genera of the family Haplonicidae as well as for identifying species of the genus *Haplonicus* distributed around Iceland by examining the

relevant literature of species and genera descriptions to derive distinguishing morphological characters (Lincoln, 1985a,b; Sars, 1896-1899; Richardson, 1908; Hansen, 1916; Menzies, 1962; Hessler, 1970; Chardy, 1974; Brökeland and Wägele, 2004; Brökeland, 2005; Brökeland and Svavarsson, 2017). The genus *Haplonicus* was present with nine species: *H. aduncus* Lincoln, 1985a, *H. ampliatus* Lincoln, 1985a, *H. angustus* Lincoln, 1985a, *H. astraphes* Brökeland and Svavarsson, 2017, *H. bicuspis* (Sars, 1877), *H. borealis* Lincoln, 1985a, *H. foresti* Chardy, 1974, *H. hamatus* Lincoln, 1985a and *H. spinifer* Hansen, 1916. The genera *Antennuloniscus* and *Chauliodoniscus* were only represented by one species each with *Antennuloniscus simplex* Lincoln, 1985a and *Chauliodoniscus armadilloides* (Hansen, 1916).

Length-width ratios of body segments and appendages were measured according to Hessler (1970); total width refers to the widest part of the habitus, while total length was measured from the anterior beginning of the head (excluding the rostrum if present) to the exterior medial end of the pleotelson. Length and width measurements for the appendage ratios always describe the greatest length of the articles. Arabic numerals were used to refer to body segments (Wolff, 1962; Hessler, 1970).

For distinguishing characters as well as digital habitus drawings were created. To illustrate these, literature drawings were modified, first by scanning the image and then tracing them manually using a Wacom Intuos M Drawing Pad, while using the vector-based software Adobe Illustrator CC (Lincoln, 1985a,b; Sars, 1896-1899; Hessler, 1970; Coleman, 2003; Brökeland and Wägele, 2004; Brökeland, 2005; Coleman, 2009; Brökeland and Svavarsson, 2017; Adobe Inc, 2019).

To create the interactive keys, the software package DELTA (DEscription Language for TAXonomy) (Dallwitz, 1980; Dallwitz et al., 2006; Coleman et al., 2010) was utilized, running in a virtual machine (Oracle, 2021) with Microsoft Windows XP as operating system. Characters, character states, species-specific links to worms.org and obis.org as well as taxa were added and put into the DELTA editor together with previously created character and taxon illustrations. Afterward this was exported into natural language descriptions and a multi-access key (INTKEY) using the actions set's directive files tonatr and toint. Finally, the interactive keys were published online.¹

Molecular Analysis

Cytocrome Oxidase I Barcoding

The focus of the molecular analysis was on the genus *Haploniscus*. Five individuals of each species were studied, where available. *Haploniscus bicuspis* has been previously shown to represent a species complex (see Paulus et al., 2021) and thus representative sequences of all three cryptic species identified by Paulus et al. (2021) were included.

DNA extraction was performed using the Marine Animal Tissue Genomic DNA Extraction Kit (Neo Biotech) or the Genomic DNA from tissue kit with NucleoSpin technology (Macherey-Nagel) following the manufacturer's recommendations. The gut was removed prior to DNA extraction and only the midsection of the specimens was utilized. A 70 μ L elution buffer was added to elute the DNA. For PCR amplification of COI, 1 μ L of the DNA extract was utilized together with PuReTaq Ready-To-Go PCR Beads (GE Healthcare) and 1 μ L of either dgLCO (GGT CAA CAA ATC ATA AAG AYA TYG G; Meyer, 2003)/dgHCO (TAA ACT TCA GGG TGA CCA AAR AAY CA; Meyer, 2003) or LCOJJ(CHACWAAYCATAAAGATATYGG; Astrin and Stüben, 2008)/HCOJJ (AWACTTCVGGRTGVCCAAARAATCA; Astrin and Stüben, 2008) primers as well as 22 μ L nuclease-free water. The PCR program comprised of an initial denaturation step at 95°C for 5 min, followed by 38 cycles of 95°C for 45 s, 45°C for 50 s and 72°C for 1 min as well as a final elongation at 72°C for 5 min. Successful amplification was assessed via gel electrophoresis (1% TAE gels). Excess primers were removed with ExoSAP and the final PCR products were bidirectionally sequenced by Macrogen using the PCR primers. Forward and reverse sequences were assembled and quality checked in GENEIOUS Prime version 2019.2.3. Sequences were aligned with MUSCLE (v 3.8.425, Edgar, 2004) including *Chauliodoniscus* sp. (GenBank accession: JF283447) as an outgroup.

To identify genetic lineages that may correspond to species, two different computational approaches were employed: general mixed Yule coalescent (GMYC; Pons et al., 2006) and assembling species by automatic partitioning (ASAP; Puillandre et al., 2021). ASAP partitions species by ranked genetic distances. Pairwise uncorrected *p*-distances were pre-computed with MEGA-X 10.0.5 (Kumar et al., 2018). The online version of ASAP was run using standard settings, except for increasing the maximum

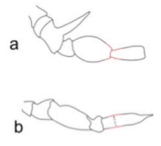
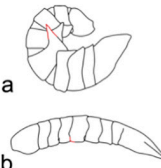
Character	Drawing	Character states	
Articles 5 and 6 of antenna 2		Not fused (a) <i>Haploniscus</i> <i>Chauliodoniscus</i>	Fused (diffuse suture still visible) (b) <i>Antennuloniscus</i>
Anterior angles of pereonite 2-4 not prolonged		At least one pereonite prolonged (a) <i>Chauliodoniscus</i>	Not prolonged (b) <i>Haploniscus</i> <i>Antennuloniscus</i>

FIGURE 2 | Summary of morphological characters and character states used to create the interactive identification key for genera within Haploniscidae.

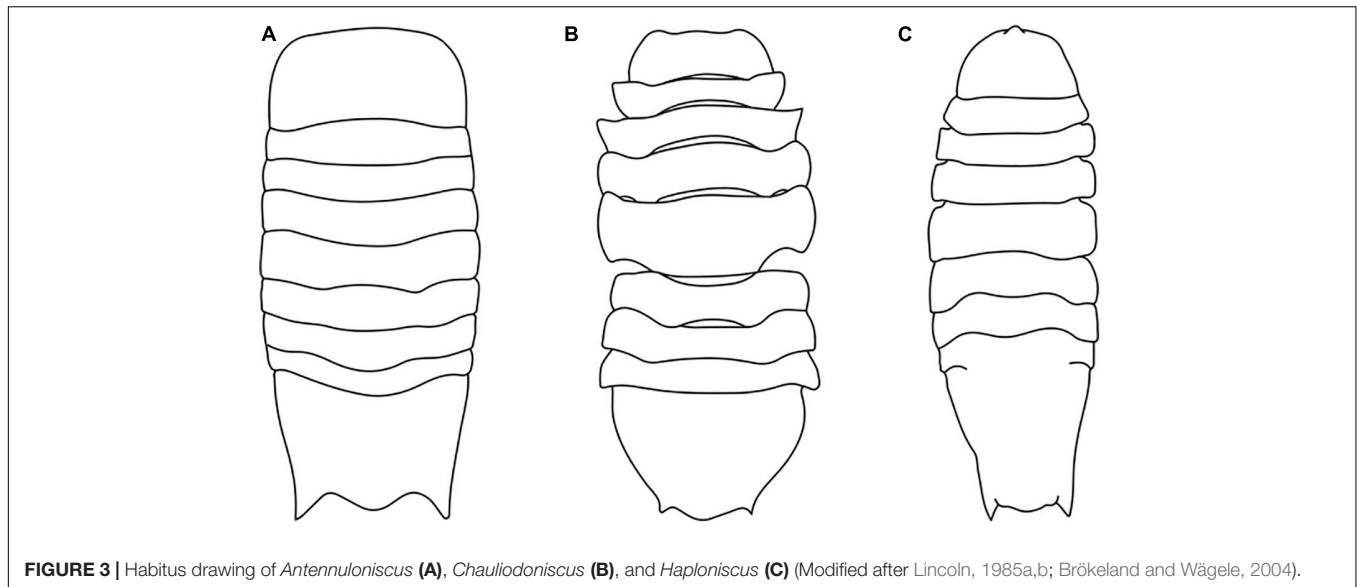
considered distance to 15%. GMYC delimits species based on branching patterns in an ultrametric tree. The ultrametric tree was computed with BEAST2 2.63 (Bouckaert et al., 2019), employing a Yule prior and enforcing the monophyly of the ingroup. Each haplotype was included only once. BEAST2 was run for 10⁶ generations, sampling every 1000th tree. Convergence was assessed with Tracer and the final tree annotated with TreeAnnotator (BEAST2 package), removing the first 25% of retained trees as burn-in. GMYC was run in R (R Core Team, 2020) using the once the single and once the multiple threshold method.

Phylogenetic analysis was performed with MrBayes (v 3.2.7a, Ronquist et al., 2012) using the best-fitting model (GTR + G + I) with four runs and six chains for 10⁷ generations. Every 5000th tree was retained and the first 25% were removed as burn-in. The best-fitting model was determined with MEGA-X following the AIC, and the analysis was performed on the CIPRES Science Gateway (Miller et al., 2010). The resulting tree was visualized with FigTree (Version 1.4.4).

Matrix-Assisted Laser Desorption/Ionization Time-of-Flight Mass Spectrometry

From the same specimens used for COI barcoding, a single pereopod with the attached muscles was used for matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF MS) measurements. The tissue was incubated in 5 μ l of a matrix solution containing α -Cyano-4-hydroxycinnamic acid (HCCA) as a saturated solution in 50% acetonitrile, 47.5% molecular grade water and 2.5% trifluoroacetic acid. After 5 min of incubation, 1.5 μ l of the extract solution was applied to one spot for crystallization on a target plate. Measurements were carried out on a Microflex LT/SH System (Bruker Daltonics), employing the flexControl 3.4 (Bruker Daltonics) software. Masses were measured in a range from 2,000 to 20,000 Dalton (Da). During measurements, peaks were evaluated using a centroid peak detection algorithm, a signal-to-noise threshold of 2, and a minimum intensity threshold of 600, with a peak resolution higher than 400 in a range from 2,000 to 10,000 Da.

¹ <https://www.researchgate.net/project/Use-of-computers-for-taxonomy/update/5d78f8333843b0b98263acf3>



For fuzzy control the Proteins/Oligonucleotide method was employed with a maximal resolution ten times above the threshold. Analyses included mass spectra from *H. bicuspis* (I-III = 6, IV = 5, V = 5), *H. foresti* A ($n = 3$), *H. foresti* B ($n = 1$), *H. angustus* ($n = 3$), *H. hamatus* ($n = 4$) and *H. n. sp.* A ($n = 4$).

Data were processed in R (R Core Team, 2020) using R packages MALDIquant and MALDIquantForeign (Gibb and Strimmer, 2012; Gibb, 2015), trimming mass spectra to an identical range from 2,000 to 20,000 Da. Data were smoothed with the Savitzky-Golay method (Savitzky and Golay, 1964) and the baseline was removed using the SNIP baseline estimation method (Ryan et al., 1988) using 15 iterations. After normalization was done using the TIC method, a noise reduction using a signal to noise ratio (SNR) of 3 was applied. For peak picking, a half window size of 14 was used, applying the MAD method implemented in MALDIquant. Repeated peak binning was carried out with a tolerance of 0.002 in a strict approach. Missing values were interpolated from the corresponding mass spectra and all signals below a SNR of 1.75 were assumed to be below the detection limit and set to zero. The resulting data matrix was Hellinger transformed (Legendre and Gallagher, 2001).

To test group differentiation for classification approaches, a Random Forest (RF) (Breiman, 2001) analysis was carried out using R package Random Forest (Liaw and Wiener, 2002) ($n_{tree} = 2000$, $m_{try} = 35$). Only species with at least three specimens were used for RF models. Significant deviation from random of the observed model errors was calculated with the function `MVSF.test` from package `RFtools`² (Rossel and Martínez Arbizu, 2018). Classification success was tested by creating RF models whereby one sample was omitted in each case and this was then classified using the model. Classification was tested using the *post hoc* test described by Rossel and Martínez Arbizu (2018) from the R package `RFtools` using a 1% alpha value for

false positive recognition. Data were visualized in R using a Barnes-Hut implementation of t-Distributed Stochastic Neighbor Embedding (Krijthe and Van der Maaten, 2015) (perplexity = 5, max.iter = 4,000).

Biogeography

Geographical and Depth Distribution of Haploniscidae Species Distributed Around Iceland

To analyze the geographical species distribution, one map for each haploniscid species distributed around Iceland was created as well as one projection encompassing all species together. Data were visualized using QGIS 3.4.7-Madeira (QGIS Development Team, 2020) with the WGS 84 (ESPG: 32631) coordinate reference system. Individuals of *Halploniscus bicuspis* were assigned to the three species found within the species complex in Paulus et al. (2021) based on their distribution. To analyze the species' depth distribution, a box and whisker plots was created with Excel 16.53 (Microsoft Corporation, 2021). The whiskers boxplot is based on the same data as utilized in the geographical distribution maps although each station finding was only included once per species.

Species Distribution Models

Species distribution models (SDM) were calculated using the random forest approach (Breiman, 2001). Random Forest is a non-parametric and non-linear modeling approach based on decision trees (Hastie et al., 2009), which has been successfully applied to the modeling of species distributions around Iceland in both classification (Meißner et al., 2014; Paulus et al., 2021) and regression problems (Ostmann and Arbizu, 2018).

Predictive variables include particulate organic carbon (POC) flux, bottom water temperature, salinity, oxygen, and depth. The georeferenced predictive layers with a pixel resolution of 5-arc minute were retrieved from Global

²<https://github.com/pmartinezarbizu/RFtools>

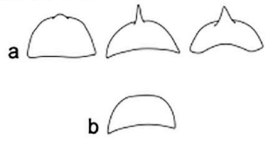
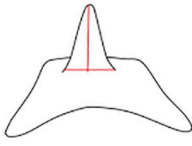
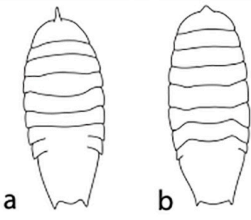
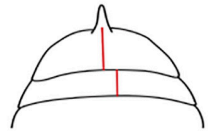
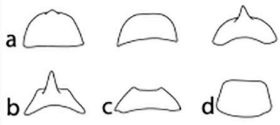
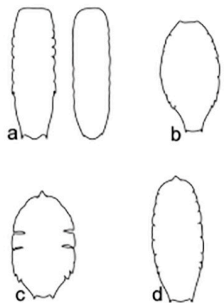
Character	Drawing	Character state			
		Present (a)	Absent (b)		
Rostrum presence		<i>H. ampliatus</i> <i>H. aduncus</i> <i>H. bicuspis</i> <i>H. borealis</i> <i>H. hamatus</i>	<i>H. angustus</i> <i>H. foresti</i> <i>H. spinifer</i> <i>H. astraphes</i>		
Rostrum length-width ratio		Longer than wide <i>H. aduncus</i> <i>H. hamatus</i>	About as wide as long or wider than long <i>H. ampliatus</i> <i>H. bicuspis</i> <i>H. borealis</i>		
Pleotelson fusion		Fused with pereonite 6 and 7 (a) <i>H. foresti</i> <i>H. aduncus</i>	Fused with pereonite 7 (b) <i>H. ampliatus</i> <i>H. angustus</i> <i>H. astraphes</i> <i>H. bicuspis</i> <i>H. borealis</i> <i>H. hamatus</i> <i>H. spinifer</i>		
Head length compared to first segment length		Length of first segment fits over 1.5 times into head length <i>H. aduncus</i> <i>H. angustus</i> <i>H. astraphes</i> <i>H. bicuspis</i>	Length of first segment fits under 1.5 times into head length <i>H. ampliatus</i> <i>H. borealis</i> <i>H. foresti</i> <i>H. hamatus</i> <i>H. spinifer</i>		
Head shape		Half circular (a) <i>H. aduncus</i> <i>H. ampliatus</i> <i>H. angustus</i> <i>H. bicuspis</i> <i>H. borealis</i> <i>H. spinifer</i>	Trapezoid (b) <i>H. hamatus</i>	Slightly concave (c) <i>H. foresti</i>	Rectangular (d) <i>H. astraphes</i>
Body shape		Straight (a) <i>H. angustus</i> <i>H. astraphes</i>	Fish-like (b) <i>H. foresti</i>	Plate-shaped (c) <i>H. ampliatus</i>	Slightly convex (d) <i>H. aduncus</i> <i>H. bicuspis</i> <i>H. borealis</i> <i>H. hamatus</i> <i>H. spinifer</i>

FIGURE 4 | Summary of characters and character states used to create the interactive identification key for species within *Haploniscus* occurring around Iceland (Modified after Lincoln, 1985a; Brökeland and Wägele, 2004; Brökeland and Svavarsson, 2017).

Marine Environment Dataset (Basher et al., 2014). The training dataset consists of 1,511 sampling events with information on presence or absence of the isopod species under consideration. A separate model was calculated for each of the 14 taxa. Haploniscidae was found in 189 samples, while

1,322 samples had no Haploniscidae. Because of the great unbalanced distribution of classes presence and absences, for training the models, the sample size argument was adjusted, so that the number of absence samples in each of the 5,000 random trees was set to half of the number of

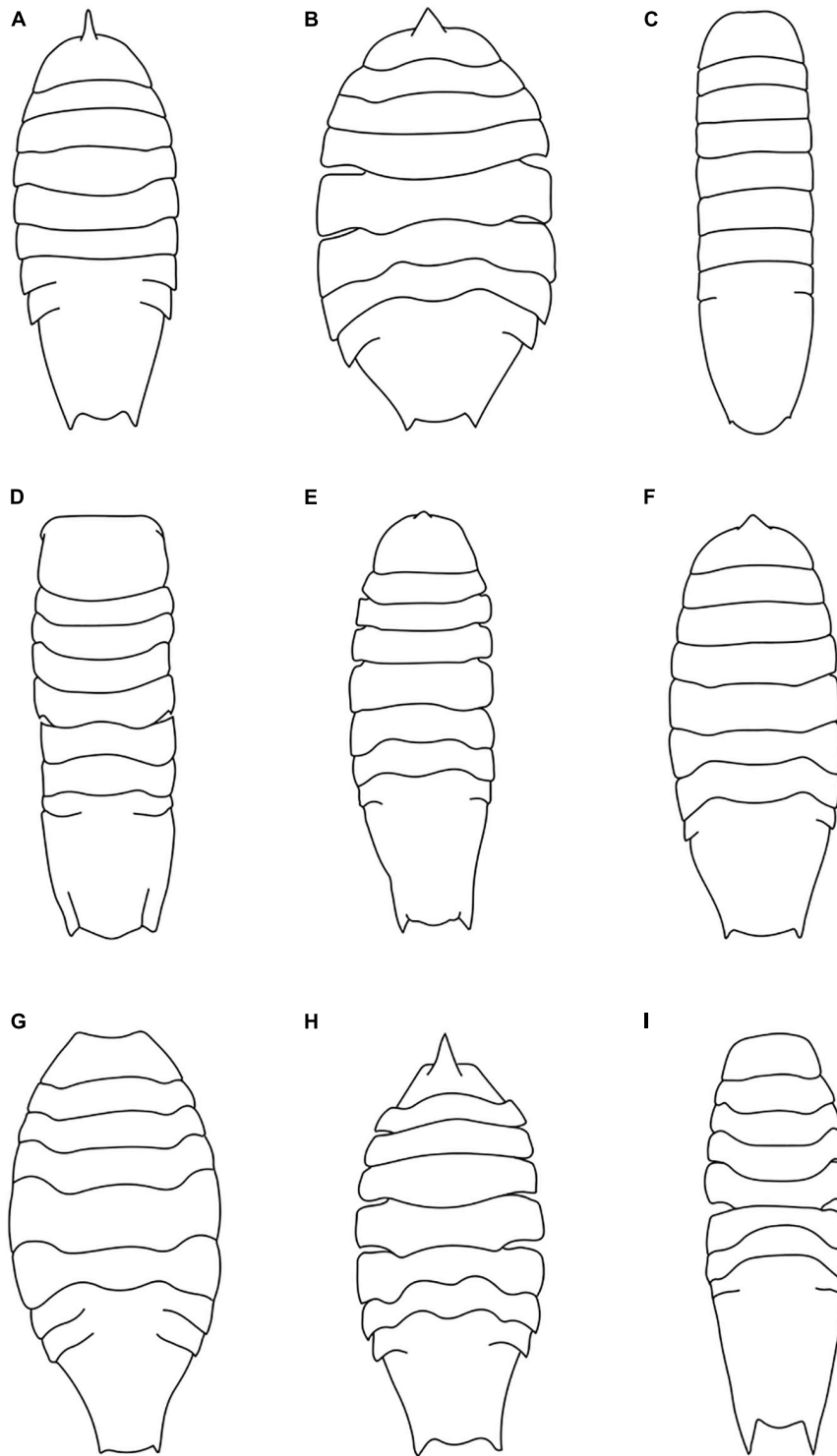


FIGURE 5 | Habitus drawings of Haploniscus aduncus **(A)** *H. ampliatus* **(B)** *H. angustus* **(C)** *H. astraphes* **(D)** *H. bicuspis* **(E)** *H. borealis* **(F)** *H. foresti* **(G)** *H. hamatus* **(H)** *H. spinifer* **(I)** (Modified after Brökeland and Wägele, 2004; Brökeland and Svavarsson, 2017).

presences. The probability of occurrence of the haploniscid species was predicted on 88,785 locations around Iceland. Computations were carried out in the statistical environment

of R using the package randomForest (Liaw and Wiener, 2002). The spatial distribution was visualized in QGIS (QGIS Development Team, 2020).

RESULTS

Interactive Determination Keys

Generic Level – Haploniscidae

To distinguish between the three haploniscid genera that are distributed around Iceland, two determination characters were utilized in the Interactive key (Figure 2) the articles five and six of the second antenna show a visible suture only in the genus *Antennuloniscus*. The remaining two genera are distinguishable from each other as at least one of the anterior angles of the pereonites two to four, which are prolonged in *Chauliodoniscus* but not in *Haploniscus*. For the interactive key, each character as well as habitus are illustrated (Figures 2, 3). The key was published online: <https://doi.org/10.5281/zenodo.5682763>.

Species Level – *Haploniscus*

To distinguish between the nine described *Haploniscus* species that are distributed around Iceland, six identification characters were utilized. Each distinguishing character as well as habitus are illustrated in the interactive key (Figures 4, 5). The species can be categorized into four different body shapes: (1) a slightly convex body shape (*H. aduncus*, *H. bicuspis*, *H. borealis*, *H. hamatus* and *H. spinifer*); (2) a rather straight body shape (*H. angustus* and *H. astraphes* B); (3) a plate-shaped body (*H. ampliatus*); and lastly (4) a fish-like body shape (*H. foresti*).

Furthermore, four head shapes are found in the species: (1) half circular (*H. aduncus*, *H. ampliatus*, *H. bicuspis* and *H. borealis*) (2) trapezoid (*H. hamatus*) (3) slightly concave (*H. foresti*) and (4) rectangular (*H. astraphes*). Furthermore, the absence (*H. angustus*, *H. foresti*, *H. spinifer* and *H. astraphes*) or presence (*H. ampliatus*, *H. aduncus*, *H. bicuspis*, *H. borealis*, *H. hamatus*) of the rostrum was used as another distinguishing character. In addition, of the species with a rostrum, its length- and width ratio differs between species. In some species (*H. aduncus* and *H. hamatus*), the rostrum is longer than wide, while in others (*H. ampliatus*, *H. bicuspis*, *H. borealis*), the rostrum is at least as wide as long or even wider than long. The species can also be categorized into two different types of pleotelson fusion: (1) the pleotelson is fused with sixth and seventh pereonite (*H. aduncus*, *H. foresti*); or (2) the pleotelson is only fused with the seventh pereonite (*H. ampliatus*, *H. angustus*, *H. astraphes*, *H. aduncus*, *H. borealis*, *H. bicuspis*, *H. hamatus*, *H. spinifer*). Finally, comparing the ratio of the head length (without rostrum if present) relative to the first segment length can be used to differentiate between the nine different *Haploniscus* species. The length of the first segment either fits over 1.5 times (*H. aduncus*, *H. angustus*, *H. astraphes*, *H. bicuspis*) or under 1.5 times into the head length (*H. ampliatus*, *H. borealis*, *H. foresti*, *H. hamatus* and *H. spinifer*). The key was published online: <https://doi.org/10.5281/zenodo.5701346>.

During the testing of the interactive key for *Haploniscus*, a new species has been discovered, which was not identifiable with the produced key. In addition, we consulted species descriptions of all other described species in the genus, which confirmed that it is new to science. This species is most similar to *Haploniscus aduncus*, *Haploniscus astraphes* and *Haploniscus hamatus*. It differs from the other *Haploniscus* species around Iceland in

possessing the unique character combination of: The pleotelson is only fused with pereonite 7, no rostrum is present, and the species has a different head shape. In the following, we provisionally name this undescribed species “*Haploniscus* sp. A” and include it into all molecular, proteomic and biogeographical analyses, while it does not occur in the interactive key.

Molecular Analyses

Cytochrome Oxidase I

The COI alignment included sequences of 37 *Haploniscus* specimens with a length of 652 bp. No indels or stop codons were present. ASAP (Threshold 11%, p -value $1.22e^{-02}$) and the single threshold analysis of GMYC suggested the presence of seven putative species: *H. angustus*, *H. hamatus*, *H. bicuspis* (all three cryptic species identified by Paulus et al. (2021) grouped together), *H. sp. A* and three putative species identified as *H. foresti* (*H. foresti* A–C) (Figure 6). These putative species are all separated by > 17% interspecific p -distances (Table 1). Two of the putative *H. foresti* species are potential sister species (*H. foresti* A and B, ~17% genetic distance), while *H. foresti* C is more divergent from the other two with ~24% genetic distance. The GMYC multiple threshold resulted in ten putative species by additionally delimiting the three cryptic species of *H. bicuspis* (following Paulus et al., 2021) and by further splitting *H. sp. A* into two putative species (Figure 6). While the three cryptic species of *H. bicuspis* are separated by ~4–6% genetic distance from each other (Table 1), the two putative species are separated by only 2.4%. The 2nd and 3rd highest scoring ASAP partitions resulted in either six (grouping *H. foresti* A and B together; threshold 18.2%, p -value $2.97e^{-01}$) or 12 (splitting *H. angustus* and *H. sp. A* into two and *H. bicuspis* into four putative species; threshold 1.7%, p -value $3.67e^{-01}$) putative species.

Haploniscus foresti A and C occurred sympatrically at station 983 (South of Iceland), *H. foresti* B was recorded from station 106 (Reykjanes Ridge). The relatively large intraspecific distances of up to 2.3% within *H. angustus* were observed among specimens collected at the same station (Station 1148). The two most divergent putative species within *H. sp. A* did not occur sympatrically, but one occurred along the Reykjanes Ridge (Stations 106 and 137) and the other from the Irminger Sea (Station 1054; Close to the Reykjanes Ridge) to eastern Iceland (Station 1219).

Matrix-Assisted Laser Desorption/Ionization Time-of-Flight Mass Spectrometry

In total, mass spectra from 31 specimens of eight species were assessed (Figure 7A). The PCA of the raw data (Figure 7B) depicts clear differences between the different species, except for the highly similar *H. bicuspis* complex. Significant differences were found between all groups using a pairwise Wilcoxon test. Constraining the data toward the respective species in a RF model (OOB error = 0.03), results in a clear distinction of the different species, even for the specimens belonging to the *H. bicuspis* complex of which only a single specimen was misclassified within the RF model (Figure 7C). Creating RF models, leaving out one specimen for each species respectively and subsequently using this model to classify the left-out specimen resulted in 100%

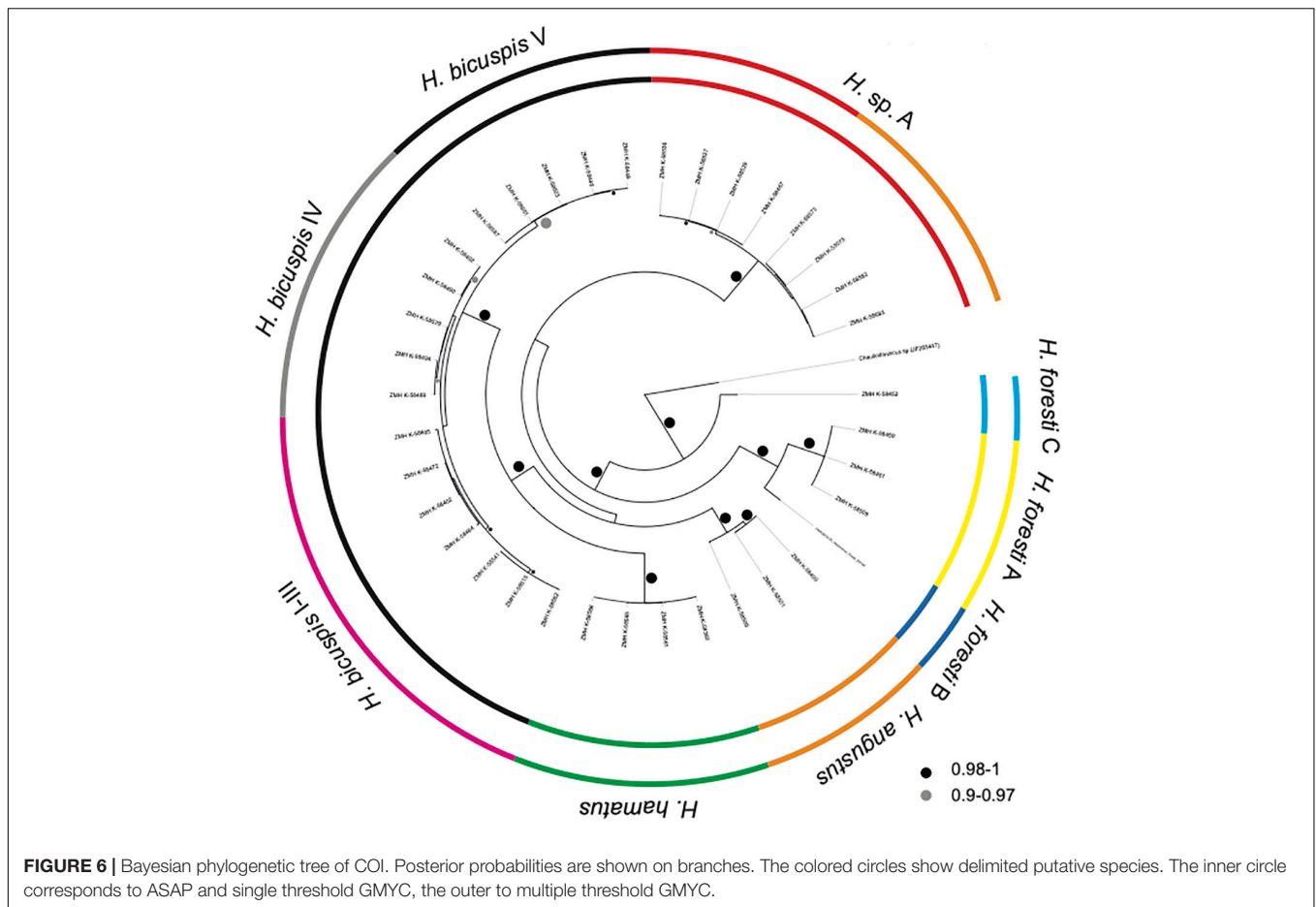


FIGURE 6 | Bayesian phylogenetic tree of COI. Posterior probabilities are shown on branches. The colored circles show delimited putative species. The inner circle corresponds to ASAP and single threshold GMYC, the outer to multiple threshold GMYC.

TABLE 1 | Uncorrected COI *p*-distances among putative Haploniscus species.

	<i>H. foresti</i> A (EP72)	<i>H. foresti</i> B (EP199)	<i>H. foresti</i> C (EP74)	<i>H. angustus</i>	<i>H. hamatus</i>	<i>H. sp. A</i>	<i>H. bicuspis</i> I-III	<i>H. bicuspis</i> IV	<i>H. bicuspis</i> V
<i>H. foresti</i> A (EP72)	0.0–0.3								
<i>H. foresti</i> B (EP199)	17.3–17.6	–							
<i>H. foresti</i> C (EP74)	24.4–24.7	23.8	–						
<i>H. angustus</i>	23.2–23.5	21.9–22.4	20.1–20.4	0.0–2.3					
<i>H. hamatus</i>	23.2–23.5	23.3–23.5	23.6	21.8–22.4	0.0–0.2				
<i>H. sp. A</i>	25.2–26.2	23.3–23.9	20.4–21.3	19.9–20.4	24.1–25.8	0.0–2.4			
<i>H. bicuspis</i> I-III	25.6–26.4	23.9–25.9	23.6–23.9	20.6–21.0	20.4–20.9	22.4–23.7	0.0–3.5		
<i>H. bicuspis</i> IV	22.9–24.8	22.2–23.8	23.1–23.5	20.2–21.0	20.2–20.7	20.9–22.7	4.1–5.2	0.0–1.5	
<i>H. bicuspis</i> V	25.5–25.8	24.4–24.7	23.5–23.8	19.2–20.6	20.4	22.0–22.9	5.5–6.4	4.8–5.4	0.0–0.5

Intraspecific distances are along the diagonal, interspecific distances below the diagonal.

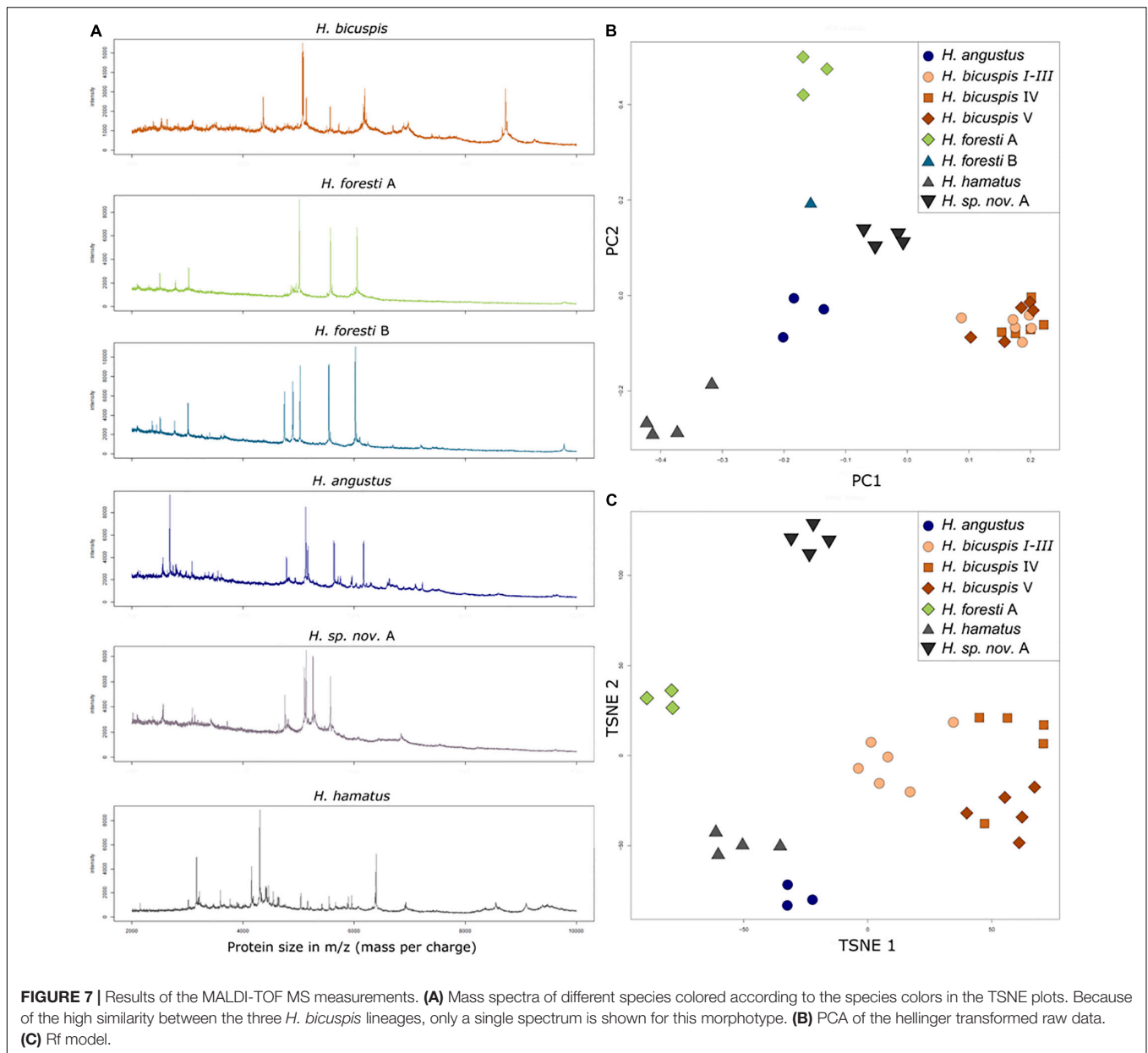
identification success supported by the *post hoc* test in all cases. None of the specimens were misclassified and no classification was recognized as a false positive by the *post hoc* test.

Biogeography Geographical Distribution

Distribution maps for thirteen haploniscid species identified from the BIOICE and IceAGE samples were created in QGIS (Figure 8). Except for *Haploniscus borealis* Lincoln, 1985a, our

data included all species previously recorded from Icelandic waters as well as *Haploniscus* sp. A. Where possible records of *Haploniscus bicuspis* were assigned to the respective cryptic species (Paulus et al., 2021), whose distribution was shown separately (Figure 9). The distribution of the three potentially cryptic species suggested for *H. foresti* by the genetic data was not analyzed separately as we cannot assign the various distribution records accordingly.

Only one species each of the genera *Antennuloniscus* and *Chauliodoniscus* were present, both species occurring only to



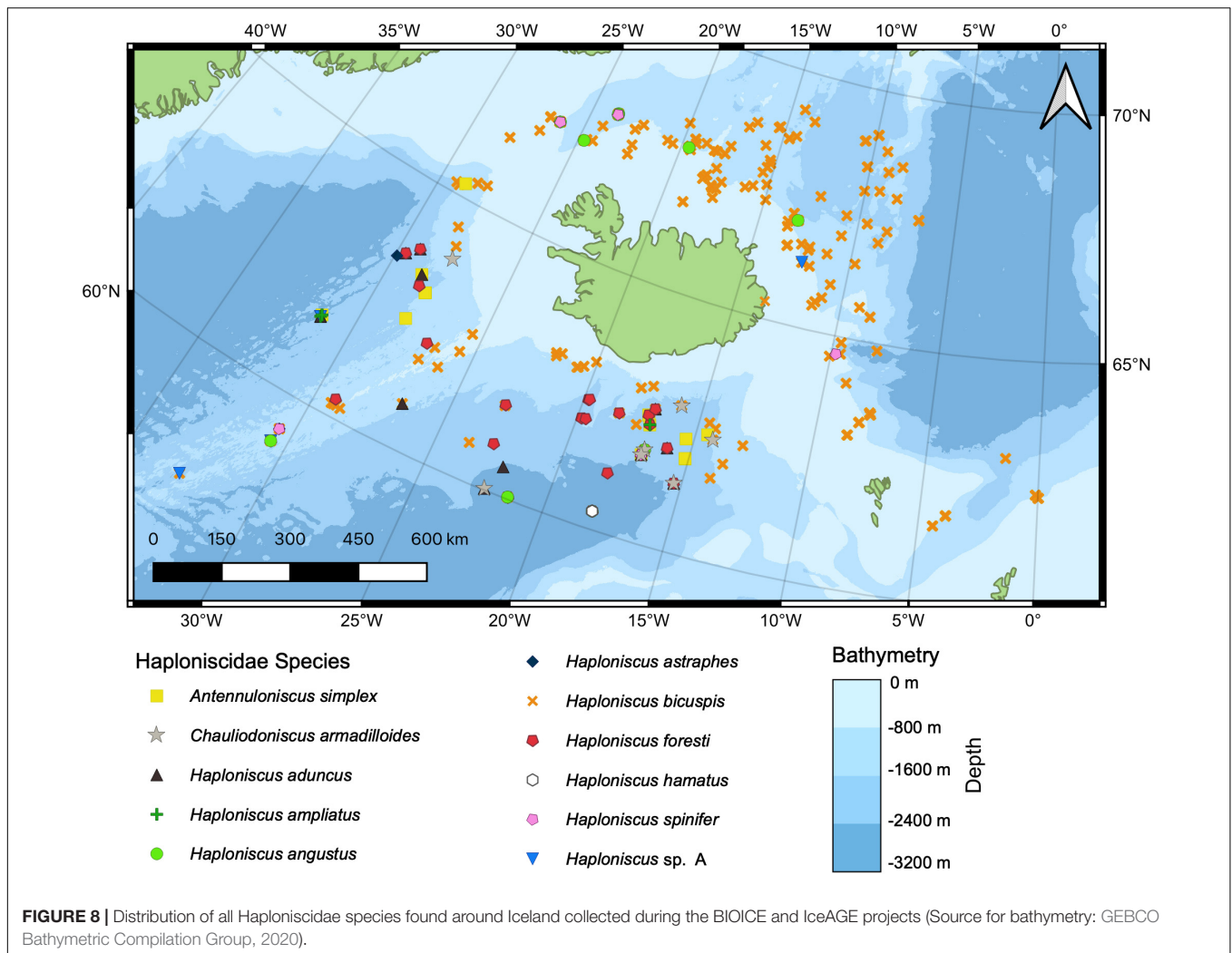
the south of the GIF ridge. Within the genus *Haploniscus*, the distribution of seven species was restricted to either the north or south of the GIF ridge. *H. aduncus*, *H. ampliatus*, *H. bicuspis* IV (compare Paulus et al., 2021), *H. bicuspis* V (compare Paulus et al., 2021), *H. foresti* and *H. hamatus* occurred solely south of the ridge, while *H. bicuspis* I–III (compare Paulus et al., 2021) was restricted to the north of the ridge.

Four species showed a near-circum-Iceland distribution. *Haploniscus angustus* Lincoln, 1985a was present at four stations in the south as well as at five stations north of the GIF ridge and *Haploniscus spinifer* occurred at two stations south as well as at three stations north of the ridge. *Haploniscus astraphes* and *H. sp. A* were both found at three stations south and only at one station north of the ridge.

Bathymetric Distribution

Haploniscid species are primarily distributed at bathyal depths (Figure 10), with only one species, *Haploniscus bicuspis* I–III (compare Paulus et al., 2021), recorded at shelf depth (minimum depth: 64 m). Most species occur across a relatively large depth range (>1000 m), with *Haploniscus bicuspis* I–III exhibiting the widest depth range (64–2613 m [median: 1108 m]). In contrast, *H. hamatus* was bathymetrically the most restricted species, having been found between 2568 and 2670 m (median: 2619 m). Notably, species within the *H. bicuspis* species complex (compare Paulus et al., 2021) all show fairly broad, overlapping depth distributions (Figure 10).

Species found only in the south of the GIF ridge and those found north of the ridge showed some differences in



bathymetric distribution; most southern species (*A. simplex*, *C. armadilloides*, *H. aduncus*, *H. ampliatus*, *H. bicuspis* V, *H. foresti* and *H. hamatus*) were only found in a depth deeper than 800 m, except *H. bicuspis* IV which is present between 778 and 2850 m (median: 1412 m). In contrast, *H. angustus*, *H. astraphes*, *H. bicuspis* V, *H. spinifer* and *H. sp. A*, whose distribution is either limited to the north or that occur on both sides of the ridge, can also be found above 800 m.

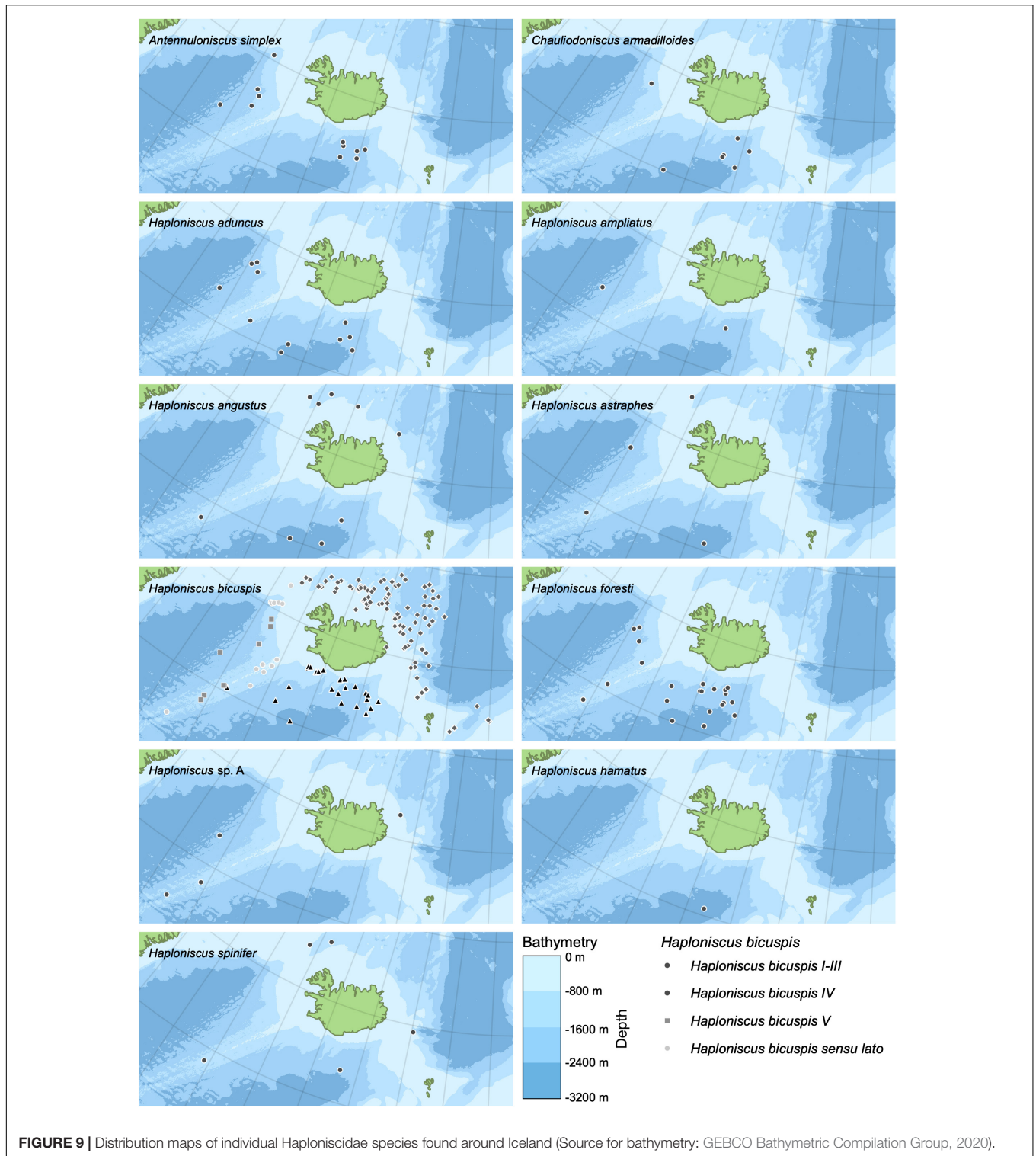
Species Distribution Models

The prediction of the probability of occurrence of the species in geographic space is shown in **Figure 11**. Probabilities below 0.5 are considered “absence” and are not shown. The models show that four of the species, viz. *A. simplex*, *H. aduncus*, *H. foresti* (here the three potential cryptic species were not treated separately) and *H. bicuspis* IV, are sympatric having their main distribution in deep waters in the Icelandic basin, but they slightly differ in the probability of occurrence in the Irminger Sea. Only *H. bicuspis* I-III shows a clear preference for areas influenced by the colder deep-waters in the Iceland and Norwegian Seas.

Model accuracy is shown in **Table 2**. The prediction error is larger for the class “n” (absent) and is relatively low for class “y” (presence). This results in a relatively high combined error for the model, which is however, not compromising accuracy for presence. In other words, the models are failing to predict where the species is truly absent.

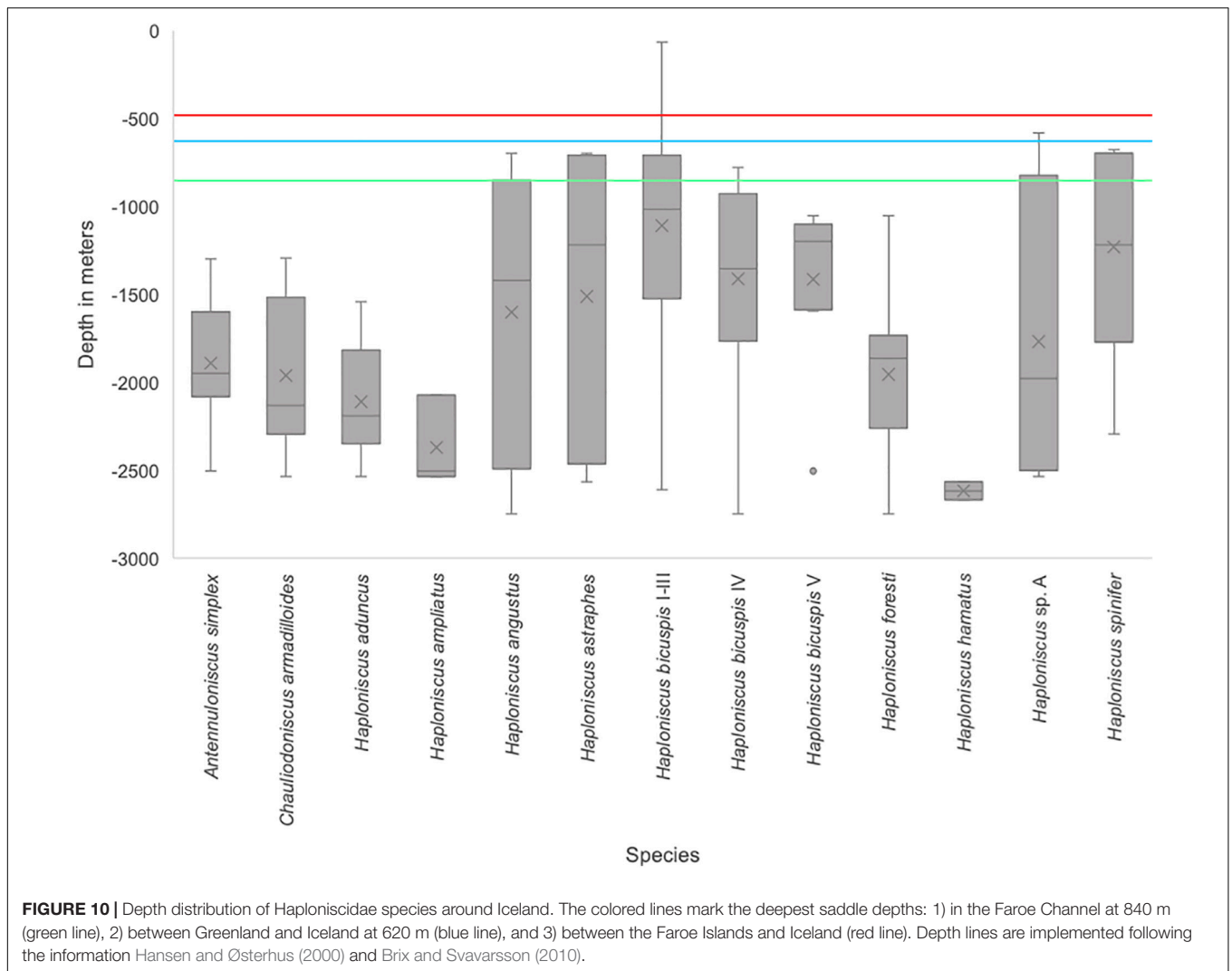
DISCUSSION

Progressive destruction and/or transformation of deep-sea ecosystems and associated loss of biological diversity has raised concerns about the consequences for overall ecosystem functioning (Danovaro et al., 2008). Concomitantly a call for sustainable use of the services provided by the deep sea (Armstrong et al., 2012; Thurber et al., 2014; Glover et al., 2018). Deep-sea ecosystems and fauna found around Iceland could be significantly impaired primarily by climate-related changes (Arnason, 2007; Astthorsson et al., 2007). Mitigating and managing the impacts of such changes on the marine environment, however, requires broadening of taxonomic



knowledge in order to identify patterns and drivers of diversity, forecast potential alterations of the deep-sea environment, and ultimately inform marine spatial planning (Howell et al., 2020). Correct identification of the species is central to achieving this, but often poses a challenge when dealing with deep-sea samples.

Interactive keys have proven to be a valuable tool to aid species identification of Icelandic haploniscids. Originating in the 1970s, interactive keys using Descriptive Language for Taxonomy (DELTA) methodologies as an instrument for identifying species are not new (Dallwitz, 1974), but their use in deep-sea taxonomy

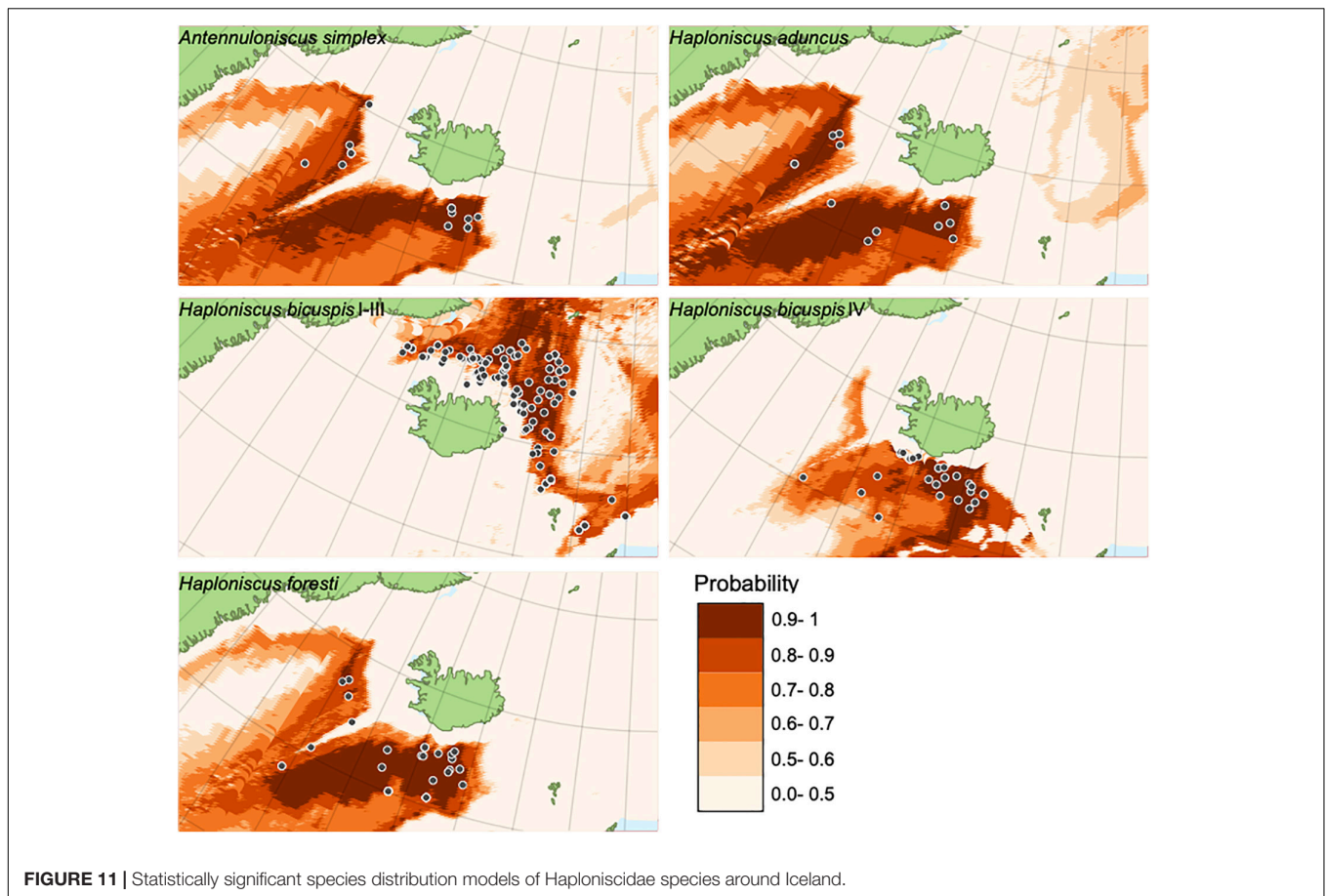


has thus far been rudimentary (Saucède et al., 2021). Overall, there is a paucity of any type (paper or digital) of identification key and guide in general³ and for deep-sea taxa in particular (Miljutin et al., 2010; Gollner et al., 2014; Bergmeier et al., 2017); instead, identifications are mostly based on descriptive works that are distributed throughout the literature (cf. Glover et al., 2021). This makes the identification process on the one hand arduous and lengthy, and on the other hand exclusive, since non-experts are not familiar with or have no access to often historical monographs. In addition, these historical monographs are often inadequately illustrated, further hindering easy species identification. Yet, given the increasing human pressures on the deep sea, the need to generate important biodiversity information is currently enormous. At the same time, the number of professional taxonomists performing this vital task is declining. As an antidote, non-experts could increasingly be called upon to identify species (Langenkämper et al., 2019; Saucède et al., 2021). To this end, it is also critical that increased efforts are

made to develop keys, particularly those that are easy to use and access, such as interactive keys deposited in open repositories. Accompanying this, taxonomic efforts to describe species from the deep sea should be intensified by trained taxonomists, since only what we know can be preserved (Glover et al., 2018).

Both traditional and interactive keys, including those created in this paper, are based on formal species descriptions (e.g., Drinkwater, 2009). The applicability of these keys therefore strongly depends on the level of known and described biodiversity in the area for which they were created. Around Iceland, the deep-sea isopod fauna is considered to be relatively well known (Brix et al., 2018) and the assumption was that most species should therefore be easily identifiable with the generated key. When creating keys for taxa in other deep-sea areas, the situation is different. In these areas, typically a high percentage of undescribed crustacean species is expected (e.g., Poore, 2008; Poore et al., 2015). Here, created keys would be less applicable, as these undescribed species would not be contained in the key and thus naturally not be identifiable. However, this would also be the case with conventional dichotomous keys. In fact, where a major

³<https://www.cbd.int/gti/problem.shtml>



advantage of interactive keys over conventional keys comes into effect as these keys have been proven to be highly customizable. This is particularly important for deep-sea crustaceans, where much of the fauna is unknown (Appeltans et al., 2012). Indeed, interactive keys can keep pace with the dynamics of continuous new species discoveries (and descriptions) as well as a changing taxonomy (through revisions), as these can be expanded and added to rather swiftly (Coleman et al., 2010). However, this presupposes that the keys are publicly accessible (e.g., on a website provided for this purpose), easy to find (e.g., advertised on commonly used taxonomic websites) and that the necessary capacities are available to maintain them over the long term (Coleman and Radulovici, 2020).

Lately there have been numerous initiatives to bring taxonomy to the web and thus make it accessible to a wider public. Among these efforts, EDIT (European Distributed Institute of Taxonomy⁴), CATE (Creating a taxonomic e-science, Godfray et al., 2007) and Scratchpad⁵ deserve special mention. The advantage of these services is that they are backed by institutes that guarantee that the infrastructure remains in place for the long term and can be brought up to date. Web-based services that are provided by individuals often lack the manpower

to consistently renew their content and make it sustainable (Coleman and Radulovici, 2020). An example of this is crustaceanet⁶ of the Australian Museum, Sydney, which has provided interactive keys for numerous crustacean taxa, yet its owner is now retired and the project is currently not being continued (Coleman and Radulovici, 2020). For deep-sea taxa, the World Register of Deep-Sea Species (WoRDSS) collects (traditional and interactive) keys and relevant literature that help in species identification, which now contains records from more than 600 publications – yet with a notably very low representation of sources for isopods (Glover et al., 2021). There are also individual researchers behind WoRDSS, but it is linked to the World Register of Marine Species (WoRMS, WoRMS Editorial Board, 2021), providing a catalog of all marine species names. As it is supported by the mass of deep-sea taxonomists, it is likely to be carried forward. Through this type of web-based taxonomic projects, a variety of other meta information could be added to complement species identification, including biogeographical, ecological, and collection data, as well as images and sketches (Farr, 2006). The latter would facilitate the exchange of data between researchers on undescribed species, as this is the case in many deep-sea regions, and

⁴<https://cybertaxonomy.eu/>

⁵<http://scratchpads.org/about/concept>

⁶<https://australian.museum/learn/collections/natural-science/marine-invertebrates/crustacea/>

TABLE 2 | Model accuracy.

Species	n	y	n error	n	y	y error	Model error
1 <i>Antennulonicus simplex</i>	1,296	214	0.142	0	14	0	14.042
2 <i>Haploniscus aduncus</i>	1,265	248	0.164	1	10	0.091	16.339
3 <i>Haploniscus bicuspis</i> I-III	1,081	326	0.232	9	108	0.077	21.982
4 <i>Haploniscus bicuspis</i> IV	1,209	289	0.193	0	26	0	18.963
5 <i>Haploniscus foresti</i>	1,267	232	0.155	2	23	0.080	15.354

thus facilitating taxonomic intercalibration (International Seabed Authority [ISA], 2020; Lins et al., 2021). In practice, one could imagine integrating undescribed species based on preliminary identifications into interactive keys (cf. Yamasaki et al., 2020), whereby the identifications are certainly not robust, but would help immensely in solving problems of species identification in highly diverse deep-sea areas. Since many interactive keys not only offer a web-based user interface, but also an offline version for download (such as Intkey), they are flexible enough to be used in the field or on a research vessel, regardless of internet access.

Interactive keys have a number of further assets compared to conventional dichotomous keys, which can also be advantageous in a deep-sea context. For example, the identification process does not require a strict order of the identification features. Instead, the user can freely choose the sequence of the determination characters and avoid those which are either unclear to the identifier or that are simply not present because specimens are damaged (Dallwitz, 2018). This is especially important when recruiting non-experts for species identification as they tend to have difficulties in recognizing described identification features. In addition, the use of interactive keys can be more efficient compared to dichotomous keys, as programs, such as DELTA, can autonomously specify the most suitable identification features (Dallwitz, 2018), and thus the user is able to choose the identification character that will identify individuals most quickly. In some cases, this can be tied to just one diagnostic character, for example in *Haploniscus hamatus* the unique, trapezoid head shape.

In general, a sensible selection of the identification features influences the applicability of interactive and binary keys equally. It is desirable to utilize conspicuous determination features that are retained even after rough sampling of the animals; fragile crustacean specimens in particular can lose appendages when retrieving samples from greater depth and subsequent sample processing. Therefore, determination features that can break off easily, such as antennae or legs, are often unsuitable, since damaged or missing determination features can lead to the identification in traditional keys being aborted. On the other hand, some flashy characters harbor the risk that crucial but less conspicuous features for species identification could be overlooked. Within Desmosomatidae, *Eugerdella serrata* Brix, 2006, for example, a pronounced jagged rostrum is present, with this feature enabling identification from several oceans (Brix, 2006; Lörz et al., 2013; Kaiser et al., 2021). But it is now clear that it represents a complex of species, containing at least two (Brix et al., 2018), but probably even more species. The Haploniscidae family is morphologically very conservative, and often only subtle

differences are decisive in distinguishing between species and genera (Park, 2000; Brökeland, 2005). Here, in particular, the overall body form and the shape of the rostrum were emphasized as valuable diagnostic features for intraspecific delimitation and identification (Brökeland and Raupach, 2008). Further characters, such as setation patterns of pereopods or uropods have been also studied to deduce intraspecific differences, but these are often not useful as segregating characters in identification keys, as they are not visible at first glance and setae are often broken off (Brökeland, 2005). We have therefore tried to strike a balance between the practicality of the keys in the sense of easily recognizable features and those that are robust and constant within a species. Therefore, for haploniscid taxa in this study, preferably habitus characteristics and the shape of the rostrum were used to distinguish genera and species within Haploniscidae.

Interactive keys reach their limits in similar places as conventional keys but can partially counteract them. For example, many traditional keys only focus on female characters, which is largely due to the fact that female isopods are more prevalent in the deep sea, and male descriptions are often lacking (Riehl et al., 2012; Kaiser, 2015). In addition, there is a strong sexual dimorphism encountered in many isopod families (incl. Haploniscidae), with males and females differing greatly in certain features (Riehl et al., 2012); in Haploniscidae this is, for example, expressed in the form that in males the second antenna is more robust and setose, and pleotelson processes are more distinct (Park, 2000; Brökeland, 2010; Paulus et al., 2021). As a result, many of the original species' descriptions only describe the female, since the male, if any, was difficult to assign and, therefore many identification keys are ultimately better suited to identify female individuals.

The identification keys produced here were based on gender-neutral characters, where a description for both sexes was available. With the help of this, however, it was not possible to identify males and females in the same way if they exhibit strong sexual dimorphism or even different male stages (such as within *Haploniscus bicuspis*, cf. Paulus et al., 2021). So, it becomes clear that using interactive keys, despite its flexible nature, would not solve the problem of identifying highly sexually dimorphic species. The same applies to ontogenetic dimorphism and the identification and differentiation of potentially cryptic species or severely damaged specimens. Many interactive keys for Crustacea, including the ones presented in this paper, are only able to identify adult individuals (e.g., Dallwitz, 2021). This is because of differing determination features depending on the developmental stage of the animals (e.g., Hessler, 1970; Wilson et al., 2011). The result could be misidentification or termination of the identification process. While there have been few studies on selected species on how the various ontogenetic stages differ (e.g., Hessler, 1970; Brökeland, 2010; Riehl and Kühn, 2020), and these characteristics could easily be fitted into an interactive key, ultimately there is often not enough material from deep-sea samples to investigate this in greater detail.

Molecular approaches, such as DNA barcoding, are a great tool to identify individuals which cannot be easily determined with morphological characters, for example in juveniles, species

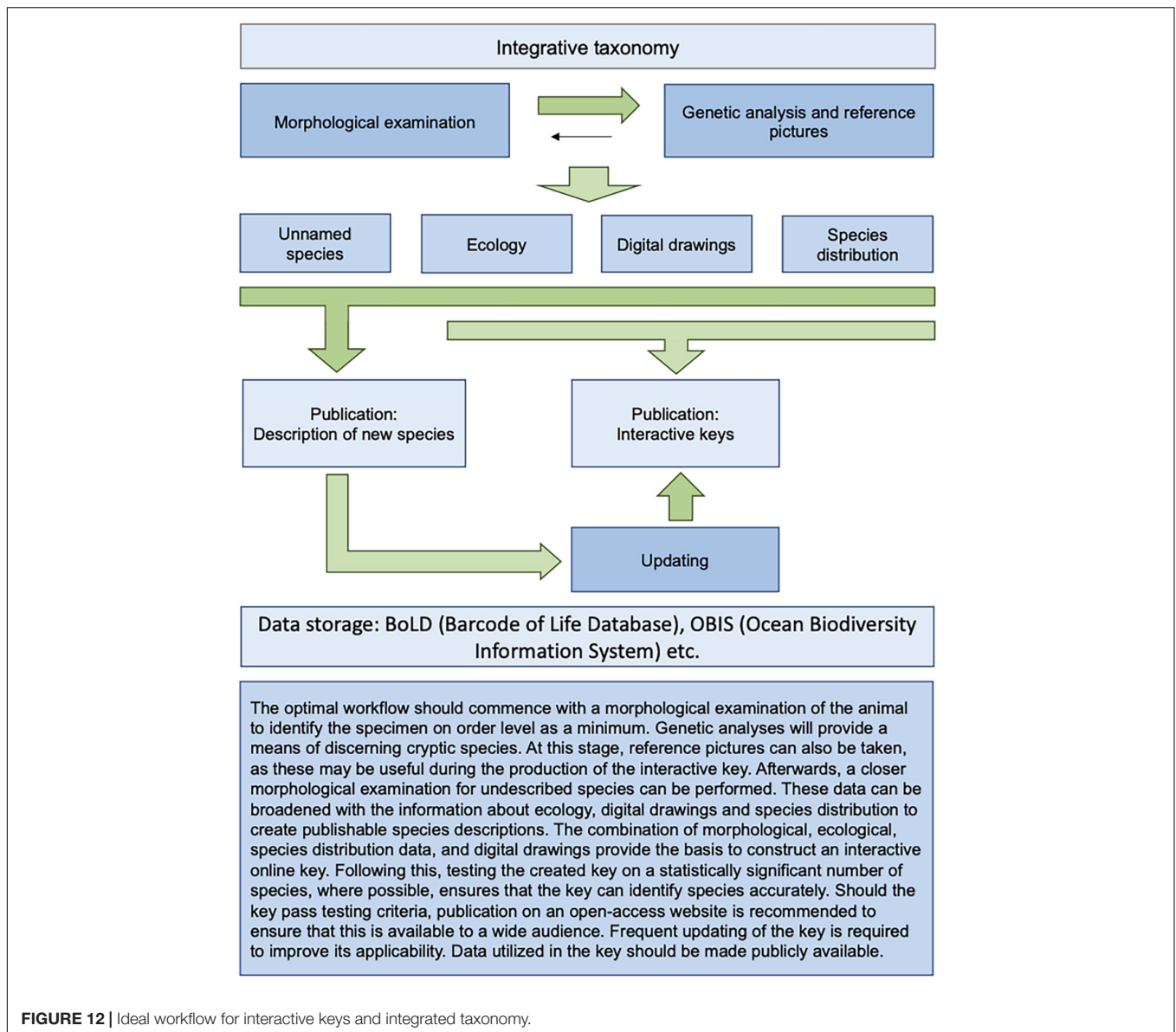


FIGURE 12 | Ideal workflow for interactive keys and integrated taxonomy.

with strong sexual dimorphism or damaged individuals. The identification of known species requires that DNA sequences of the respective species are available. DNA barcoding can further assist in the delimitation of hitherto unknown species by revealing genetically highly divergent lineages, which might correspond to (cryptic) species new to science.

Our molecular-genetic analysis of the haploniscids consistently separated all of the morphologically identified species from each other, and further uncovered a number of cryptic species, notably within *H. foresti* and *H. bicuspis* (see also Paulus et al., 2021), as well as to delimit the newly discovered species within *Haploniscus* (sp. A).

Similar to COI barcoding, MALDI-TOF MS can help in the identification of those animals which are difficult to determine with interactive keys alone. In our data proteomic fingerprinting further supported the distinctiveness of *H. bicuspis* (see also

Paulus et al., 2021). Nearly all of the genetically suggested species, including the cryptic species of *H. foresti*, differed distinctly in their proteomic spectrum as well. However, differences between the putative species of *H. bicuspis* were very small and referred to a recent speciation process of this complex (Paulus et al., 2021). In a classification test, all specimens were classified correctly, showing the use of mass spectra for specimen identification. These results are in concordance with previous studies showing the high success of proteome fingerprinting in metazoan specimen identification on taxa such as fish (Mazzeo and Siciliano, 2016; Rossel et al., 2021), insects (Dieme et al., 2014; Raharimalala et al., 2017) or other crustaceans (Bode et al., 2017; Rossel et al., 2019; Renz et al., 2021). Overall, MALDI-TOF MS seems to be a promising fast and low-cost tool for the identification of deep-sea isopods. However, reference spectra need to be available to facilitate good identification

success especially for highly similar groups such as the different *H. bicuspis* lineages. It can be worthwhile to create internal databases with mass spectra for species from a defined area, especially for ongoing projects such as IceAGE.

In addition to the morphological and molecular approaches, analyzing geographical and depth distribution may provide further indications for correct species identification. This is because the distribution of benthic species around Iceland is strongly influenced by water mass properties, depth and topography, and here in particular by the GIF ridge. The latter is a major obstacle for marine invertebrates due to topographic constraints and confluence of different water masses (e.g., Svavarsson et al., 1993; Weisshappel and Svavarsson, 1998; Jöst et al., 2017; Lörz et al., 2021). This probably applies in particular to brooding taxa such as isopods, which do not have a swimming larval stage and are therefore likely to have limited dispersal capacity (Pearse et al., 2009; Brix et al., 2020). Although we recognize that this could be a circular argument, since the true geographic distribution of many deep-sea species is unknown, we have derived the assumption from the foregoing that the isopod species would be confined to a certain depth or geographic area. Hence, we could, for example, deduce a possible identification error from an unusually large geographic or depth distribution, which will need to be further assessed using an integrative taxonomy. Overall, distribution patterns helped to predict species entities to some extent, including the example of *H. bicuspis* I–III occurring north of the GIF ridge from the remaining putative species occurring only south of the ridge. However, there were many overlapping distributions both geographically and bathymetrically (Figures 8–10), which therefore did not aid species identification. Furthermore, distribution patterns of some species raise the question of whether they are not actually two species; for example, *Haplونiscus angustus* and *H. spinifer* both have records north and south of the GIF ridge. Unfortunately, no species records of any of these two species from the north and south of the ridge are included in our molecular data to refute or confirm this hypothesis. In view of their depth distribution, there would be at least the possibility for both species to cross the GIF ridge at its deepest point (i.e., the Faroe channel at 840 m, Figure 10).

Finally, it was investigated whether species can be better differentiated from one another and thus identified using species distribution models. The underlying assumption here was that each species occupies its own niche space, which is defined by a certain set of environmental variables. The models presented here suffer from the fact that the number of sampling events with presence of the species is remarkably low compared to the number of sampling events where the species were not found. Yet, not finding a species in a sample does not mean that the species is not living in this location, it just means that the species was eventually not found at this sampling event. Because of the “pseudoabsence” nature of our class “n”, it is not dramatic that the prediction accuracy in the training dataset for class “n” is relatively high (14–23% error). In contrast the prediction accuracy for presence (class “y”) was low 0–9%, indicating that the models provide a good estimate of the potential distribution of the species. Using the distribution of *Haplونiscus bicuspis*

species as an example where an adequate number of data points were available, it was, however, possible to demonstrate the potential of this method; it could be shown that the predicted occurrences diverge in at least two species or groups of species (*H. bicuspis* sp. I–III restricted to the north of the GIF and *H. bicuspis* IV to the south). The addition of more environmental layers and biogeographic data will likely contribute to a better prediction of species distributions.

In summary, biogeographic and ecological species demarcation and identification is only helpful in areas in which biodiversity and distribution have been well studied and sampled. But even in one of the better-known areas like Iceland, our results have shown that knowledge gaps still exist, as demonstrated by a number of cryptic and supposedly new species. This can only be remedied through ongoing morphological examination, together with molecular methods and additional sampling. Certainly, the creation of interactive keys when combined with molecular, bathymetric and environmental datasets can increase their effectiveness as a tool for robust species identification (Figure 12).

CONCLUSION

In our analysis of Haplونiscidae around Iceland, the use of interactive keys was proven to be a powerful tool to identify described, but also to recognize potentially new species. In particular, the easier involvement of non-experts in the identification process is highlighted, especially when additional information, such as images or biogeographical data, is included. Since interactive keys are based solely on morphology, they are subject to some restrictions due to “phenomena” such as cryptic species, sexual and ontogenetic dimorphism, or when species are severely damaged. Therefore, genetic or -omic techniques such as COI barcoding, metabarcoding, and proteomics amongst others, are essential to complement the morphological assessment.

We deliberately examined specimens from a comparatively well-known group from a comparatively well-known area, but nonetheless discovered some supposedly new isopod species. Our intention was not to evaluate various morphological and genetic methods for species identification, but to test how robustly species can be identified. Nevertheless, we found molecular tools very helpful to support the morphological identification of species, in particular proteomics as a novel tool for identifying isopod species. Despite some limitations (e.g., issues of intercalibration of mass spectra between different data sets), the latter could become a valuable alternative to the more expensive and time-consuming DNA barcoding approach.

In the longer term, all newly discovered species in this study – be it part of a species complex (*H. bicuspis*) or entirely new (*H. sp. A*) – should be included in the key. Ultimately, interactive keys for other Icelandic crustacean taxa are to be developed and made publicly available in order to build a framework of taxonomic information that will help professionals and non-experts to identify the marine fauna of Iceland and thus to gain a more complete picture of the local biodiversity. The interactive key

presented in this paper is the first for isopods, and the first step toward completing an atlas of the Icelandic marine isopod fauna. At present, even dichotomous keys on paper do not exist although the BIOICE inventory of the Icelandic fauna is used as baseline knowledge for species distributions. As mentioned in the beginning, we rely on original species descriptions and their comparison for most taxa. These missing keys highlight the need for taxonomic expertise in times of the “taxonomic impediment” (Coleman, 2015). While part of this is due to the lack of skilled taxonomists, there is an argument that taxonomists could improve accessibility of species descriptions through online databases and syntheses of taxonomic information to create identification keys. Through summarizing published species description and allowing easy adaptation, interactive online keys can help achieve this goal.

This study shows that interactive keys are an important, but hitherto underutilized tool, for obtaining an understanding of species within a given ecosystem. Within the context of the UN’s Ocean Decade (2021 to 2030), interactive keys may provide the solution toward the objective of re-establishing taxonomic knowledge in the young generation of scientists, which is imperative for assessing ecosystem function and future change within the Anthropocene.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: http://www.boldsystems.org/index.php/Public_SearchTerms?query=DS-HAPLB, accession number(s) can be found in the additional information for this paper: <https://doi.org/10.5061/dryad.r2280gbc>; <https://zenodo.org/record/5710070#.YZqyzC2l2u4>; <https://zenodo.org/record/5701346#.YZqy873P3eo>; <https://zenodo.org/record/5682763#.YZqzAi2l2u4>.

AUTHOR CONTRIBUTIONS

JS was one of the organizers of the BIOICE expeditions and sampling. SB organized the IceAGE expeditions and sampling. JS and SB conducted the species identification. A-NL supervised

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the theses of KK. EP sequenced the COI data. MS performed the genetic distance and phylogenetic analyses. SR, JP, EP, and SB created the MALDI-TOF MS data and discussed with PM. PM created the species distribution models and KK visualized them. KK, CC, A-NL, and SB constructed the interactive keys. KK performed geological mapping and species bathymetry distribution analyses. KK, SK, MS, SR, JP, PM, CC, and SB drafted the first version of the manuscript. All authors contributed to the final version of the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fmars.2021.795196/full#supplementary-material>

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