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Exploring the vertebrate fauna of the Bird's Head Peninsula (Indonesia, West Papua) through DNA barcodes

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Abstract:

Biodiversity knowledge is widely heterogeneous across the Earth's biomes. Some areas, due to their remoteness and difficult access, present large taxonomic knowledge gaps. Mostly located in the tropics, these areas have frequently experienced a fast development of anthropogenic activities during the last

decades and are therefore of high conservation concerns. The biodiversity hotspots of Southeast Asia exemplify the stakes faced by tropical countries. While the hotspots of Sundaland (Java, Sumatra, Borneo) and Wallacea (Sulawesi, Moluccas) have long attracted the attention of biologists and conservationists alike, extensive parts of the Sahul area, in particular the island of New Guinea, have been much less explored biologically. Here, we describe the results of a DNA-based inventory of aquatic and terrestrial vertebrate communities, which was the objective of a multi-disciplinary expedition to the Bird's Head Peninsula (West Papua, Indonesia) conducted between October 17th and November 20th 2014. This expedition resulted in the assembly of 1,005 vertebrate DNA barcodes. Based on the use of multiple species-delimitation methods (GMYC, PTP, RESL, ABGD), 264 Molecular Operational Taxonomic Units (MOTUs) were delineated, among which 75 were unidentified and an additional 48 were considered cryptic. This study suggests that the diversity of vertebrates of the Bird's Head is severely underestimated and considerations on the evolutionary origin and taxonomic knowledge of these biotas are discussed.

Introduction

The Earth's biosphere has been unevenly explored, with some geographic regions relatively well surveyed and others lacking taxonomic baselines and awaiting large-scale screening of their biotas. The resulting gaps in taxonomic knowledge impede conservation planning and management, a situation particularly evident in species-rich tropical areas experiencing extensive infrastructure development (e.g., roads, dams and agriculture) and deforestation over the past few decades. This is particularly dramatic for highly endemic biotas of tropical insular systems that have become increasingly accessible to human activity and biological invasions and, as a consequence, are mostly endangered (Hoffman et al., 2010; Myers, Mittermeier, Mittermeier, da Fonseca, & Kent, 2000; Schipper et al., 2008). Southeast Asia exemplifies the stakes associated with the conservation of species-rich biota in insular tropical systems. Of the four main biogeographic provinces in this area, three (Sundaland, Wallacea, and the Philippines) have been recognized as biodiversity hotspots because they harbor an exceptional number of endemic species, and intense anthropogenic pressures threaten their existence (Hoffman et al., 2010; Myers et al., 2000; Thiault et al., 2019). Both Sundaland (Java, Sumatra and Borneo) and Wallacea (Sulawesi, Moluccas) have long captured the attention of biologists and served as early model systems in biogeographic studies (Mayr, 1944; Wallace, 1859). In particular, vertebrate assemblages in both regions have been actively documented over the last two decades, resulting in the description of hundreds of new species (Hubert et al., 2015) and the assessment of associated phylogenetic and biogeographic patterns (de Bruyn et al., 2013; den Tex, Thorington, Maldonado, & Leonard, 2010; Esselstyn, Maharadatunkamsi, Achmadi, Siler, & Evans, 2013; Hutama et al.,

2017; Lim et al., 2017; O'Connell et al., 2018; Pinheiro et al., 2017; Rowe et al., 2019; Wood Jr, Heinicke, Jackman, & Bauer, 2012).

In contrast, the northern part of Sahul, i.e. the island of New Guinea, has received comparatively little attention, as a result of limited accessibility. Delimited from Wallacea to the west by the Lyddeker line, its component lineages show multiple affinities with the neighboring Wallacea or Australia, while also displaying high levels of endemism (Crayn, Costion, & Harrington, 2015; Rowe, Reno, Richmond, Adkins, & Steppan, 2008; Unmack, Allen, & Johnson, 2013). The extent of this endemism is presumed to be vastly underestimated, because recent DNA-based species inventories purported large numbers of undescribed taxa (Kadarusman et al., 2012; Riedel, Sagata, Suhardjono, Tänzler, & Balke, 2013). Unfortunately, the rapid acceleration of deforestation in New Guinea during the last decades (Austin, Schwantes, Gu, & Kasibhatla, 2019; Filer, Keenan, Allen, & Mcalpine, 2009; Nelson et al., 2014; Shearman & Bryan, 2011), is putting many species and their habitats at risk, calling for comprehensive inventories of fauna and flora to facilitate conservation planning.

In the last 15 years, DNA barcoding, the use of the mitochondrial Cytochrome Oxidase I gene as an internal species tag (Hebert, Cywinska, Ball, & de Waard, 2003; Hebert, Ratnasingham, & deWaard, 2003), has been providing a major boost to documenting biodiversity. Initially designed to overcome limits of morphology-based species-level identification, it has increasingly been accepted as a tool to capture species boundaries and as a foundation for automated molecular species identification and detection (April, Mayden, Hanner, & Bernatchez, 2011; Blagoev et al., 2015; Delrieu-Trottin et al., 2019; deWaard et al., 2019; Kerr et al., 2007). The utility of DNA barcoding, however, always depends on the taxonomic coverage of the associated DNA barcode reference library which, in turn, requires solid taxonomic knowledge of the biotas under scrutiny (Hubert & Hanner, 2015). Several studies have emphasized the benefits of integrating a standardized DNA-based approach into the inventories of poorly known faunas (Dahruddin et al., 2017; de Araujo et al., 2018; Milá et al., 2012; Monaghan et al., 2009; Riedel et al., 2013; Sholihah et al., 2020; Smith, Fisher, & Hebert, 2005; Smith et al., 2008; Sonet et al., 2018; Tänzler, Sagata, Surbakti, Balke, & Riedel, 2012; Vacher et al., 2020). Newly developed DNA-based species delimitation methods (Fujiwasa & Barraclough, 2013; Kekkonen, Mutanen, Kaila, Nieminen, & Hebert, 2015; Monaghan et al., 2009; Puillandre, Lambert, Brouillet, & Achaz, 2012; Ratnasingham & Hebert, 2013; Jiajie Zhang, Kapli, Pavlidis, & Stamatakis, 2013)

further speeded up the pace of species discovery by dramatically increasing the throughput and lowering analytical costs (Butcher, Smith, Sharkey, & Quicke, 2012; Riedel et al., 2013).

One of the more geologically intricate regions of the island of New Guinea is its northwest portion, known as the Bird's Head Peninsula, particularly the Lengguru karstic massif, in the West Papua province. This massif originated from the subduction of the Australian and Pacific plates that resulted in the development of an accretion prism during the last 10 million years (Myr) (Hall, Cottam, & Wilson, 2011; Lohman et al., 2011). Previous DNA-based inventories of freshwater fishes in this area resulted in the discovery of multiple new taxa (Kadarusman et al., 2012; Nugraha et al., 2015). The co-occurrence of multiple lineages of distinct biogeographic origin, suggests that the Bird's Head Peninsula has been colonized from several regions (Kadarusman et al., 2012; Unmack et al., 2013) and warrants a more thorough faunal inventory, in particular for freshwater fauna, for which freshwater-specific processes of isolation have led to extremely high levels of endemism, much higher than for other continental vertebrates (Leroy et al., 2019).

Between October and November 2014, a large multi-disciplinary expedition to the Bird's Head Peninsula sampled mammals, birds, reptiles, amphibians and fishes across a diverse array of ecosystems, from mangroves to the "cloud" forest habitats, from lakes and rivers to caves of the Lengguru karst system. One of the main aims of the expedition was to conduct a DNA-based inventory of vertebrates in this relatively unchartered part of New Guinea and to extend the taxonomic coverage of the DNA barcode reference library. This survey resulted in 1,005 records for 264 vertebrate Molecular Operational Taxonomic Units (MOTUs) that are presented and discussed in this publication.

Material & methods

Sampling and collection management

The Lengguru expedition, conducted between the 17th of October and the 20th of November 2014, surveyed 35 sites in the Lengguru massif (http://www.lengguru.org/) and an additional set of 20 sites in the Western part of the Bird's Head Peninsula (Fig. 1). Freshwater fishes were sampled using electrofishing gear and cast nets. Bats and birds were trapped using mist nets. Amphibians and reptiles were hand collected or captured with glue traps. Rodents were captured using non-lethal cage traps. Specimens were photographed, individually labeled and their provenance information recorded, including geocoordinates, collection date, and collectors. A muscle tissue or blood sample was taken from each captured specimen and fixed in 95% ethanol. Fish, amphibian

and reptile voucher specimens were fixed in 5% formalin solution and subsequently transferred into a 70% ethanol solution. Mammals were preserved in 80% ethanol and birds were prepared on site as dried study skins by the Indonesian Institute of Sciences (LIPI) personnel. Both tissue and voucher specimens were deposited in the national collections at the Research Center for Biology (RCB) from LIPI.

DNA Sequencing and international repositories

Genomic DNA was extracted from the muscle tissue samples using a Qiagen DNeasy 96 tissue extraction kit following manufacturer's specifications. A 651-bp segment from the 5' region of the cytochrome oxidase I gene (COI) was amplified using the M13 tailed primers cocktails C FishF1t1/C FishR1t1 for fishes, C VF1LFt1/C VR1LRt1 for mammals and reptiles (Ivanova, Zemlak, Hanner, & Hébert, 2007), AmphF2 t1 (TGTAAAACGACGCCAGTTTCAACWAAYCAYAAAGAYATYGG)/AmphR3_t1 (CAGGAAACAGCTATGACTADACTTCWGGRTGDCCRAARAATCA) for amphibians (Prosser, unpublished) and BirdF1 t1/ BirdR2 t1 for birds (Hebert, Stoeckle, Zemlak, & Francis, 2004). PCR amplifications were done on a Veriti 96-well Fast (ABI-AppliedBiosystems) thermocycler with a final volume of 10.0µl containing 5.0µl Buffer 2X, 3.3µl ultrapure water, 1.0µl each primer (10μM), 0.2μl enzyme Phire Hot Start II DNA polymerase (5U) and 0.5μl of DNA template (~50 ng). Amplifications were conducted as follows: initial denaturation at 98°C for 5 min was followed by 30 cycles consisting of denaturation at 98°C for 5s, annealing at 56°C for 20s, and extension at 72°C for 30s, followed by a final extension step at 72°C for 5 min. The PCR products were purified with ExoSap-IT (USB Corporation, Cleveland, OH, USA) and sequenced in both directions. Sequencing reactions were performed at the Centre for Biodiversity Genomics, University of Guelph, Canada, using the "BigDye Terminator v3.1 Cycle Sequencing Ready Reaction" and sequencing was performed on an ABI 3730xl capillary sequencer (Applied Biosystems), following standard protocols described in Hebert et al. (2013). Sequences and collateral information were deposited on BOLD (Ratnasingham & Hebert, 2007) and are available as a public data set DS-LENG (dx.doi.org/10.5883/DS-LENG, Table S1).

Genetic distances and species delimitation

Kimura 2-parameter (K2P) (Kimura, 1980) pairwise genetic distances were calculated using the R package Ape 4.1 (Paradis, Claude, Strimmer, 2004). Maximum intraspecific and

nearest-neighbor genetic distances were calculated using the matrix of pairwise K2P genetic distances and the R package Spider 1.5 (Brown et al., 2012). We checked for the presence of a barcode gap, *i.e.* the lack of overlap between the distributions of the maximum intraspecific and the nearest-neighbor genetic distances, by plotting both distances and examining their relationships on an individual basis, instead of comparing both distributions independently (Blagoev et al., 2015). A Neighbor-joining (NJ) tree was built based on K2P distances and used to visually inspect genetic distances and DNA barcode clusters (Fig. S1).

Several methods have been proposed for delineating species based on DNA sequences (Kapli et al., 2017; Pons et al., 2006; Puillandre et al., 2012; Ratnasingham & Hebert, 2013). Each of these methods has different properties, particularly when dealing with singletons (i.e. delimited lineages represented by a single sequence) or heterogeneous speciation rates among lineages (Luo, Ling, Ho, & Zhu, 2018). A combination of different approaches is increasingly used to overcome potential pitfalls arising from uneven sampling (Kekkonen & Hebert, 2014; Kekkonen et al., 2015; Limmon et al., 2020; Shen et al., 2019; Sholihah et al., 2020). We used six different sequencebased methods of species delimitation to identify Molecular Operational Taxonomic Units (MOTU): (1) Refined Single Linkage (RESL) as implemented in BOLD and used to generate Barcode Index Numbers (BIN) (Ratnasingham & Hebert, 2013), (2) Automatic Barcode Gap Discovery (ABGD) (Puillandre et al., 2012), (3) Poisson Tree Process (PTP) in its single (sPTP) and multiple rates version (mPTP) as implemented in the stand-alone software mptp 0.2.3 (Kapli et al., 2017; Zhang, Kapli, Pavlidis, & Stamatakis, 2013), and (4) General Mixed Yule-Coalescent (GMYC) in its simple (sGMYC) and multiple rate version (mGMYC) as implemented in the R package Splits 1.0-19 (Fujisawa & Barraclough, 2013). A final delimitation scheme was established based on a majority-rule consensus among the six delimitation analyses performed.

Both RESL and ABGD use DNA alignments as input files while a maximum likelihood (ML) tree was used for PTP and an ultrametric tree was used for GMYC. The ML tree was reconstructed using RAxML (Stamatakis, 2014) based on a GTR+I+Γ substitution model. The ultrametric tree was reconstructed using the Bayesian approach implemented in BEAST 2.4.8 (Bouckaert et al., 2014) based on a strict-clock model using a genetic distance of 1.2% per million year (Bermingham, McCafferty, & Martin, 1997). A preliminary analysis indicated that 50 million steps was a sufficient length for the Markov chains to reach ESS>200 for all estimated parameters. Thus, two Markov chains of 50 million steps each were run independently on the entire vertebrate dataset using the Yule pure birth model tree prior and a GTR+I+Γ substitution model. Both runs

were merged using LogCombiner 2.4.8 (Bouckaert et al., 2014) and sGMYC and mGMYC analyses were conducted on 10 chronograms sampled along the merged runs using the complete DNA sequences dataset following Hubert *et al.* (2019). The final delimitation scheme for sGMYC and mGMYC was established based on a majority-rule consensus of all 10 replicates. Further Bayesian chronograms for visual inspection of the topologies and inferred divergence times were built independently for fishes, mammals, birds, amphibians and reptiles using the same parameters. Both runs were combined independently for each group using LogCombiner 2.4.8 and the maximum credibility tree was constructed using TreeAnnotator 2.4.7 (Bouckaert et al., 2014). Sampling coverage at the MOTU level was examined for the five vertebrate groups through accumulation curves generated with BOLD (Smith, Fernandez-Triana, Roughley & Hebert, 2009).

Results

Sequencing yielded a total of 1,005 COI sequences out of 1140 samples. The product lengths for the various primer combinations were as follows: 652bp for C FishF1t1/C FishR1t1, 658 bp for AmphF2 t1/AmphR3 t1, 657bp for C VF1LFt1/C VR1LRt1, and 694bp BirdF1 t1/COIbirdR2 t1. Amplification failures were randomly distributed among species, and at least one individual of each species was successfully sequenced. Average sequence length for all DNA barcodes was 649bp, and no stop codons were detected suggesting that these sequences correspond to functional coding regions. A total of 21 orders representing 61 families and 136 genera were sequenced, including nine orders representing 17 families and 29 genera of fishes, one order with six families and 13 genera of amphibians, five orders comprising 18 families and 37 genera of birds, five orders with 12 families and 27 genera of mammals, and one order representing eight families and 26 genera of reptiles (Table 1). The number of specimens identified to the species level largely varied among classes: 70% of the 343 fish specimens, 70% of the 167 amphibian specimens, 100% of the 155 bird specimens, 85% of the 173 mammal specimens and 53% of the reptile specimens. Intraspecific, interspecific within genus and interspecific within family genetic distances largely vary among classes (Table 2).

The MOTU delimitation analyses yielded varying numbers of MOTUs depending on the algorithm used for all classes (Figs. 2 & 3, Table S1). Numbers of delimited MOTUs were 54 for RESL, 66 for ABGD, 44 for sPTP, 38 for mPTP, 51 for sGMYC and 65 for mGMYC for fishes; 45 for RESL, 46 for ABGD, 43 for sPTP, 33 for mPTP, 44 for sGMYC and 49 for mGMYC for birds; 51 for RESL, 67 for ABGD, 51 for sPTP, 27 for mPTP, 52 for sGMYC and 69 for mGMYC

for mammals; 43 for RESL, 47 for ABGD, 37 for sPTP, 31 for mPTP, 38 for sGMYC and 47 for mGMYC for amphibians; and 63 for RESL, 70 for ABGD, 59 for sPTP, 40 for mPTP, 59 for sGMYC and 67 for mGMYC for reptiles (Table S1). The final consensus consisted of 59 MOTUs for fishes, 46 MOTUs for birds, 53 MOTUs for mammals, 43 MOTUs for amphibians and 63 MOTUs for reptiles. Thus, a total of 264 MOTUs was added to the DNA barcode reference library. Distributions of both maximum intraspecific distance and distance to the nearest neighbor for MOTUs overlapped in a few cases for all classes (Figs. 4a, 4b, 4g, 4h); however, a DNA barcoding gap was observed in most cases with only a few exceptions for fishes (Fig. 4i). The proportion of MOTUs that could be assigned to species varied among classes with 71% for fishes, 100% for birds, 77% for mammals, 65% for amphibians and 48% for reptiles (Table S1). The unidentified MOTUs displayed varying trends of divergence to the nearest neighbor ranging from 0.6% to 22.5% for fishes, 3.5% to 20% for mammals, 1.2% to 23.5% for amphibians and 1.9% to 37% for reptiles (Table 3). Several cases of morphologically unrecognized MOTUs assigned to the same species were detected in all groups (Table 4). As observed for unidentified MOTUs, patterns of genetic distances displayed varying trends, with distances to the nearest neighbor ranging from 1.9% to 10.8% for fishes, 1.9% to 7.5% for amphibians, 1.5% to 2.1% for birds, 2.7% to 8.7% for mammals and 2.7% to 12% for reptiles (Table 4).

MOTU accumulation curves (Fig. 5) indicate that the sampling is nearly representative for fishes and amphibians, with a plateau being almost reached; however, curves are far from reaching a plateau for birds and mammals, suggesting that the number of MOTUs recovered in this study underestimates the true vertebrate diversity in the Lengguru massif. The phylogeographic patterns were mostly congruent across groups in terms of spatial distribution and divergence (Figs. 6 & 7). Multiple cases of closely related lineages that originated during the Pleistocene and occurring in sympatry or at neighboring sites are detected for Fishes (Fig. 6), reptiles (Fig. 7a-7c) and amphibians (Fig. 7d-7f), suggesting a contribution of *in situ* diversification to the diversity build-up in the area. For reptiles and amphibians, however, several cases of deep divergence, tracing back to the Miocene, between unidentified or cryptic MOTUs were detected (Fig. 7a-5f).

Discussion

Ever since the seminal publications on DNA barcoding (Hebert, Cywinska, et al., 2003; Hebert & Gregory, 2005), numerous studies showed how DNA barcoding accelerated the development of molecular diagnostic tools for automated species identification in well-known faunas (April et al.,

2011; Blagoev et al., 2015; Kerr et al., 2007; Knebelsberger, Dunz, Neumann, & Geiger, 2014; Shen et al., 2019). It also significantly helped in speeding up inventories and species discovery (Butcher et al., 2012; Hebert, Penton, Burns, Janzen, & Hallwachs, 2004; Monaghan et al., 2009; Riedel et al., 2013; Smith et al., 2005, 2008; Smith, Wood, Janzen, Hallwachs, & Hebert, 2007; Tänzler et al., 2012). Our study confirms the benefits of integrating DNA barcoding into the taxonomic workflow of a biodiversity inventory in species-rich, yet poorly documented, biotas. This study contributed 1,005 new DNA barcode records to the reference library for the Bird's Head Peninsula, including 264 MOTUs whose delimitation was corroborated by most DNA-based delimitation methods applied (Figs. 2 & 3). The distances to the nearest-neighbor are usually exceeding maximum intra-MOTU distances by an order of magnitude of 12 (Table 2); and a barcode gap is generally observed (Fig. 4). A single case of DNA barcode sharing is observed in mammals, when a specimen of Sus scrofa was nested within S. verrucosus. This was expected, considering the reported introgression among wild Sus species, as well as between domesticated and wild lineages (Scandura, Iacolina, & Apollonio, 2011). Along the same line, species delimitation analyses failed to separate a single species pair, including *Rattus praetor* and *R*. tanezumi (mammals, MOTU199).

Several cases of MOTUs displaying small genetic distances among them were detected among fishes, amphibians and reptiles (Fig. 5) including some newly discovered MOTUs and/or multiple MOTUs within one species entity delineated based on morphology. Several cases of large conflicts between PTP and other algorithms were associated to cases of multiple MOTUs displaying small genetic distances among them. In particular, none of the Melanotaenia goldiei and M. mairasi MOTUs were found by the sPTP and mPTP algorithms, resulting in the lowest estimate of numbers of MOTUs for all methods (Table S1). MOTUs of these two Melanotaenia groups displayed much lower K2P genetic distance to their nearest neighbors than in other fish lineages. Similar discrepancies between PTP and other methods such as GMYC were previously described (Luo et al., 2018; Shen et al., 2019), with PTP being less effective when large number of species and varying divergence levels were involved. Several similar cases were also observed for amphibians (e.g. Asterophrys pullifer MOTUs, Hylophorbus spp.), reptiles (e.g. Cyrtodactylus irianjayensis MOTUs, Emoia spp.) and mammals (e.g. Hipposideros spp.). Along the same line, GMYC is known to produce excessive splitting if based on maximum credibility trees without collapsing sequences into haplotypes. Here, applying GMYC algorithms to sampled trees along the merged Markov chain, using the complete DNA barcode dataset, yielded numbers of MOTUs

that compared favorably to other methods, with mGMYC providing generally a better match to the final consensus. These results confirm the benefits of combining several species delimitation methods and using a consensus instead of a single method when it comes to avoiding artefacts (Blair & Bryson, 2017; Delrieu - Trottin et al., 2020; Kekkonen & Hebert, 2014; Kekkonen et al., 2015; Shen et al., 2019). Aside from these particular cases, methods were largely congruent and resulted in the delimitation of MOTUs with homogeneous maximum intraspecific K2P distance distribution across classes (Fig. 4).

The proportion of MOTUs that were only identified to the genus level is high for fishes, amphibians and reptiles with a total of 75 MOTUs not identified to the species level (Table 3). This trend is markedly different from previous molecular studies of vertebrates in the neighboring biogeographical provinces of Sundaland and Wallacea, where identification to the species level is much more common (Arida, 2017; Bernstein et al., 2020; Conte-Grand et al., 2017; Dahruddin et al., 2017; Connell et al., 2018; Kyle Connell, Hamidy, Kurniawan, Smith, & Fujita, 2018; Sholihah et al., 2020). Along the same line, the number of cryptic MOTUs is substantial with 48 MOTUs delimited within 19 species across the five classes (Table 4), the number of cryptic lineages within species ranging from 2 in most cases to 5 in *Cyrtodactylus irianjayensis* (reptiles). This trend was expected for some genera such as *Melanotaenia* for which DNA-based methods already helped to discover multiple new species in the Western parts of the island of New Guinea (Kadarusman et al., 2012; Nugraha et al., 2015). We found seven new MOTUs of Melanotaenia, which are now awaiting description (Table 3). Multiple cases of high and formally undescribed diversity were also detected here such as for the amphibian genera Hylophorbus (5 MOTUs) and Oreophryne (4 MOTUs), the reptile genera *Emoia* (12 MOTUs) and *Sphenomorphus* (12 MOTUs) and the bat genus *Hipposideros* (3 MOTUs). The result for amphibians was expected as previous similar efforts in tropical species-rich and poorly explored areas, e.g. the Amazon, yielded similar results (Fouquet et al., 2007; Vacher et al., 2020). This study highlights that diversity of the continental vertebrate biotas of the Bird's Head Peninsula is probably largely underestimated. Species accumulation curves are still far from reaching a plateau, especially for birds, mammals and reptiles. This vast diversity is confined to very restricted areas. Our inferences indicate that some of the MOTUs (Melanotaenia golidiei group, Melanotaenia mairasi group, Asterophrys pullifer, Hylophorbus spp., Sphenomorphus spp. and Emoia spp.) may have diversified during the Pleistocene and are distributed in the periphery of the Lengguru massif (Figs. 6 & 7). These particular cases, also pointed out by conflicting PTP and GMYC species delimitation results, suggest a recent *in situ* origin through allopatric speciation on small spatial scales. This trend was expected considering the complex geological history of the Lengguru massif that experienced an intense orogenic activity over the past 5 Million years (Bailly, Pubellier, Ringenbach, de Sigoyer, & Sapin, 2009; Villeneuve et al., 2010). In addition, karsts are highly fragmented landscapes that foster geographic isolation and promote endemism (Clements, Sodhi, Schilthuizen, & Ng, 2006; Polhemus & Allen, 2006). The present study further suggests that the build-up of species diversity in the Lengguru massif likely originated through a combination of immigration and *in situ* diversification over the course of its geological history as previously reported, for example, for the genus *Melanotaenia* (Kadarusman et al., 2012; Unmack et al., 2013). This calls for an increased effort to document further New Guinea's biota and to develop rapidly a critical mass of biological expertise in Indonesia Papua, particularly in times of ongoing deforestation and habitat loss (Austin et al., 2019; Filer et al., 2009; Nelson et al., 2014; Novotny & Molem, 2020; Shearman & Bryan, 2011).

Conclusions

The present study highlights major biodiversity knowledge gaps in the Bird's Head Peninsula, and confirms the utility of standardized DNA-based species delimitation methods in aiding biodiversity inventories. Applied to poorly surveyed faunas, such as those of the island of New Guinea, they facilitate the discovery of previously unknown biodiversity and highlight priorities for further taxonomic study. Here, a total of 123 MOTUs, corresponding 75 unidentified and 48 unrecognized MOTUs based on an initial screening of their morphology, are waiting a reexamination of their morphological characters and potentially a formal description. This number of potential new species is high and still underestimated for several groups such as reptiles, a trend that further points to the need to improve our knowledge of this biodiversity-rich island. Our study clearly shows how much we still do not know about the non-marine vertebrate diversity of the Bird's Head Peninsula. Given that many species are still awaiting discovery and that we are looking at an accelerated loss of forest and other suitable habitat in West Papua, the need for priority conservation is paramount.

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Author contributions

LP, GS, RH, JS, K organized the expedition and designed the survey; AR, PG, AH, AF, EA, M, W, CC, JS, LP, ML, RH, K, SS, EHPM, AA, AY, A, GS, NS, SW, CT, H, BM, HW, MI and S conducted the field sampling and curated the specimens; NH, HD, YF, EA, HA, AB, AF and DS conducted the sequencing and quality control; NH, DS, AB, EDT, YF, GS, AF and EA submitted data records on BOLD and analyzed the data. All authors contributed in drafting and revising the manuscript.

Data accessibility

Sequence data and associated collection information are available on the Barcode of Life Datasystem (BOLD) in the dataset "DS-LENG: DNA barcode reference library of some West Papua vertebrates" (dx.doi.org/10.5883/DS-LENG).

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Figures and Tables captions

- **Figure 1.** Distribution of the 35 collection sites in the Lengguru massif (West Papua, Indonesia) and the 20 collection sites in the western part of the Bird's head peninsula for the 1046 samples analyzed for this study. Map from https://maps-for-free.com/.
- **Figure 2.** Bayesian Chronograms based on a 1.2% of genetic divergence per million years including DNA-based species delimitation derived from sGMYC, mGMYC, sPTP, mPTP, ABGD, RESL and final delimitation schemes based on majority rule consensus among the six methods for fishes (blue), and amphibians and reptiles (green). The black outlined boxes correspond to MOTUs derived from specimens identified to the species level. Regular boxes indicate MOTUs derived from specimens identified to the genus level.
- Figure 3. Bayesian Chronograms based on a 1.2% of genetic divergence per Million years

including DNA-based species delimitation derived from sGMYC, mGMYC, sPTP, mPTP, ABGD, RESL and final delimitation schemes based on majority rule consensus among the six methods for mammals (yellow) and birds (purple). The black outlined boxes correspond to MOTUs derived from specimens identified to the species level. Regular boxes indicates MOTUs derived from specimens identified to the genus level.

Figure 4. Distribution of genetic distances below and above MOTU boundaries for amphibians (a, b, c), birds (d, e, f), fishes (g, h, i), mammals (j, k, l) and reptiles (m, n, o). (a, d, g, j, m) Distribution of maximum intra-MOTU distances (K2P). (b, e, h, k, n) Distribution of nearest neighbor distances (K2P). (c, f, i, l, o) Relationship between maximum intra-MOTU and nearest neighbor distances. Points above the diagonal line indicate species with a barcode gap. MOTUs boundaries correspond to the final delimitation scheme derived from the majority rule consensus among the six delimitation methods.

Figure 5. MOTU accumulation curves for fishes, birds, mammals, amphibians and reptiles. The x-axis varies among classes.

Figure 6. Phylogeographic patterns among selected groups of fishes. MOTUs are represented according to the final delimitation schemes based on majority rule consensus among the six methods. The black outlined boxes correspond to MOTUs derived from specimens identified to the species level. Regular boxes indicate MOTUs derived from specimens identified to the genus level. (a) *Melanotaenia mairasi* group including *M. mairasi* (blue), *M. goldiei* (orange), MOTU008 (green), MOTU007 (yellow), MOTU006 (red), MOTU003 (pink). (b) *Melanotaenia ammeri* group including *M. ammeri* (green), *M. arguni* (orange), *M. veoliae* (red), MOTU002 (blue), MOTU004 (yellow) and MOTU005 (pink).

Figure 7. Phylogeographic patterns among selected groups of reptiles (a, b, c) and amphibians (d, e, f) displaying high diversity of closely related MOTUs occurring on restricted spatial scales. MOTUs are represented according to the final delimitation schemes based on majority rule consensus among the six methods. The black outlined boxes correspond to MOTUs derived from specimens identified to the species level. Regular boxes indicate MOTUs derived from specimens identified to the genus level. (a) *Cyrtodactylus* spp. including *Cyrtodactylus irianjayaensis*

MOTU212 (yellow), MOTU213 (dark green), MOTU214 (orange), MOTU215 (blue), MOTU216 (light green), Cyrtodactylus marmoratus (red) and Cyrtodactylus sermowaiensis MOTU218 (purple) and MOTU219 (pink). (b) Sphenomorphus spp. including MOTU255 (blue), MOTU249 (dark green), MOTU257 (light green), MOTU248 (yellow), MOTU259 (orange), MOTU256 (red) and MOTU258 (pink). (c) Emoia/Sphenormorphus spp. including Emoia jakati (light blue,) MOTU236 (dark blue), MOTU232 (dark green), MOTU235 (yellow), MOTU237 (orange), MOTU233 (brown), MOTU234 (red), MOTU230 (pink), MOTU238 (light purple), MOTU240 (purple) and MOTU239 (dark purple). (d) Cornufer spp. including Cornufer bimaculatus MOTU063 (blue), MOTU065 (dark green), MOTU064 (light green) and MOTU063 (yellow), Cornufer papuensis MOTU061 (orange), MOTU062 (red) and MOTU060 (pink). (e) Asterophrys pullifer including MOTU089 (green), MOTU088 (yellow) and MOTU090 (red). (f) Hylophorbus spp. including MOTU079 (green), MOTU083 (yellow), MOTU080 (orange), MOTU081 (red) and MOTU082 (pink).

Table 1. Summary statistics of the number of DNA barcode records assembled per class, family and genus.

Table 2. Summary statistics of genetic distances including minimum, maximum and average K2P distances within-MOTU, among MOTUs within genus and among MOTUs within family for fishes, birds, mammals, amphibians and reptiles.

Table 3. List of the unidentified MOTUs including their genus assignment, consensus MOTU and BIN numbers, maximum within-MOTU K2P distance and K2P distance to the nearest neighbor.

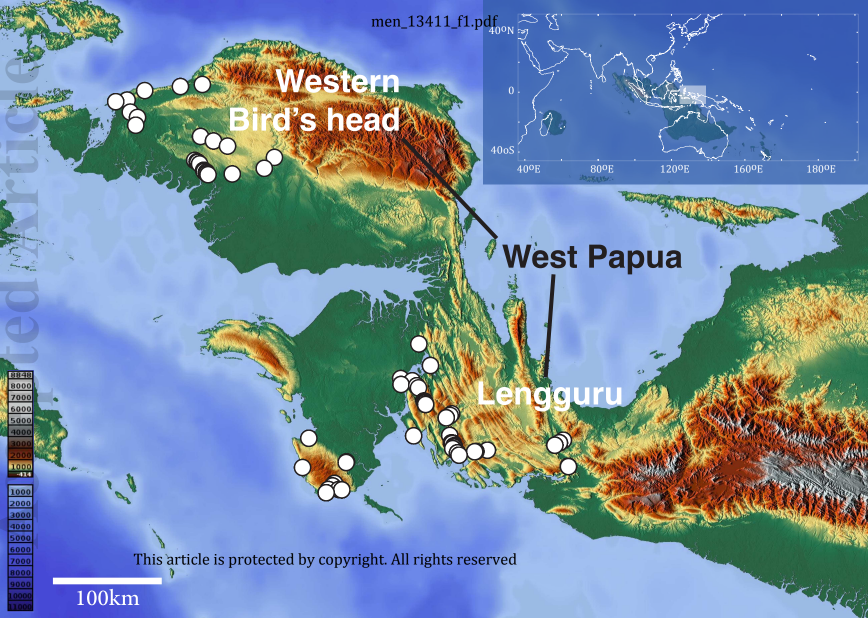
Table 4. List of MOTUs assigned to the same species based on morphological characters, including MOTU and BIN numbers, maximum within-MOTU K2P distance (percent) and K2P distance to the nearest neighbor (percent).

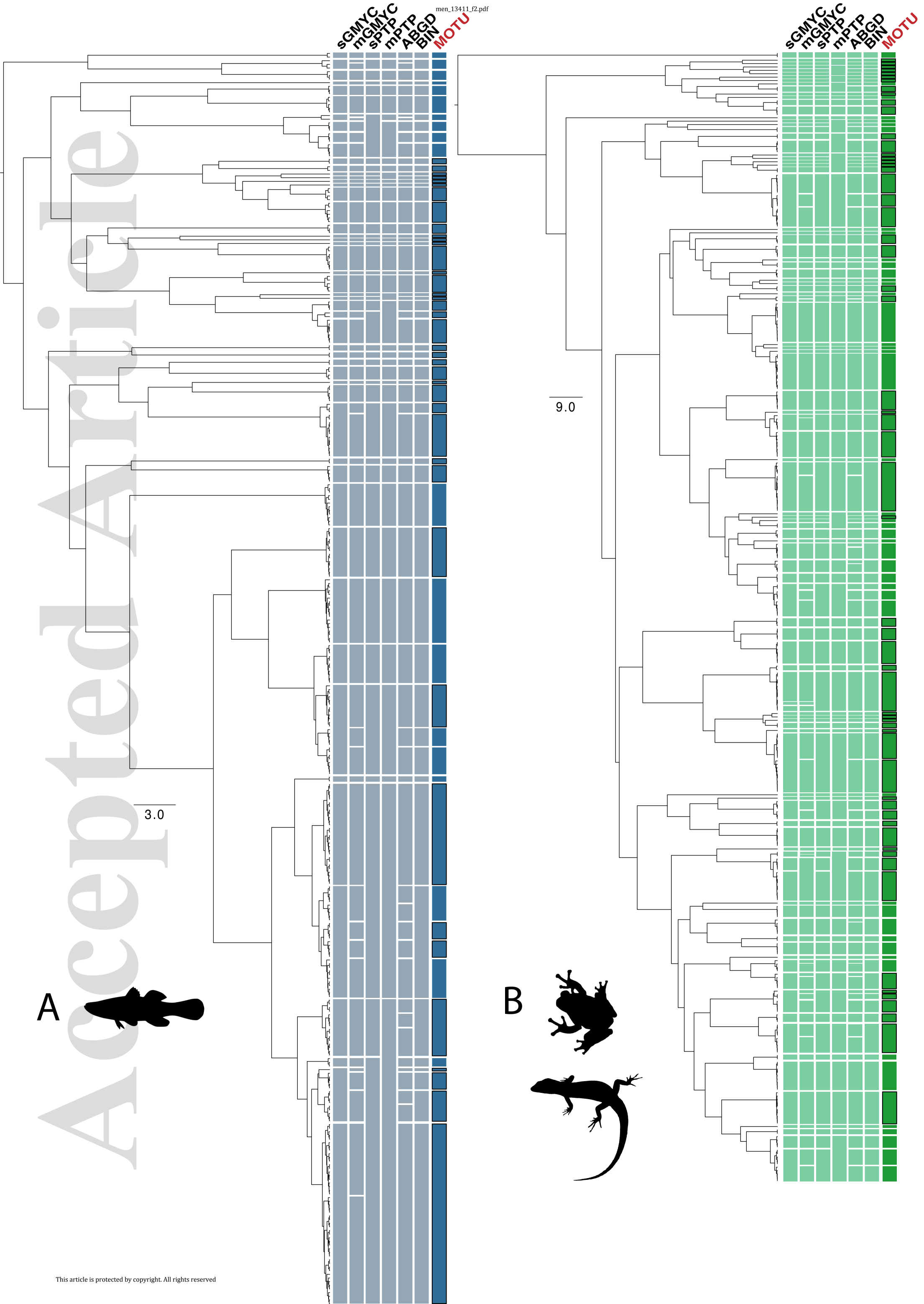
Supplementary material.

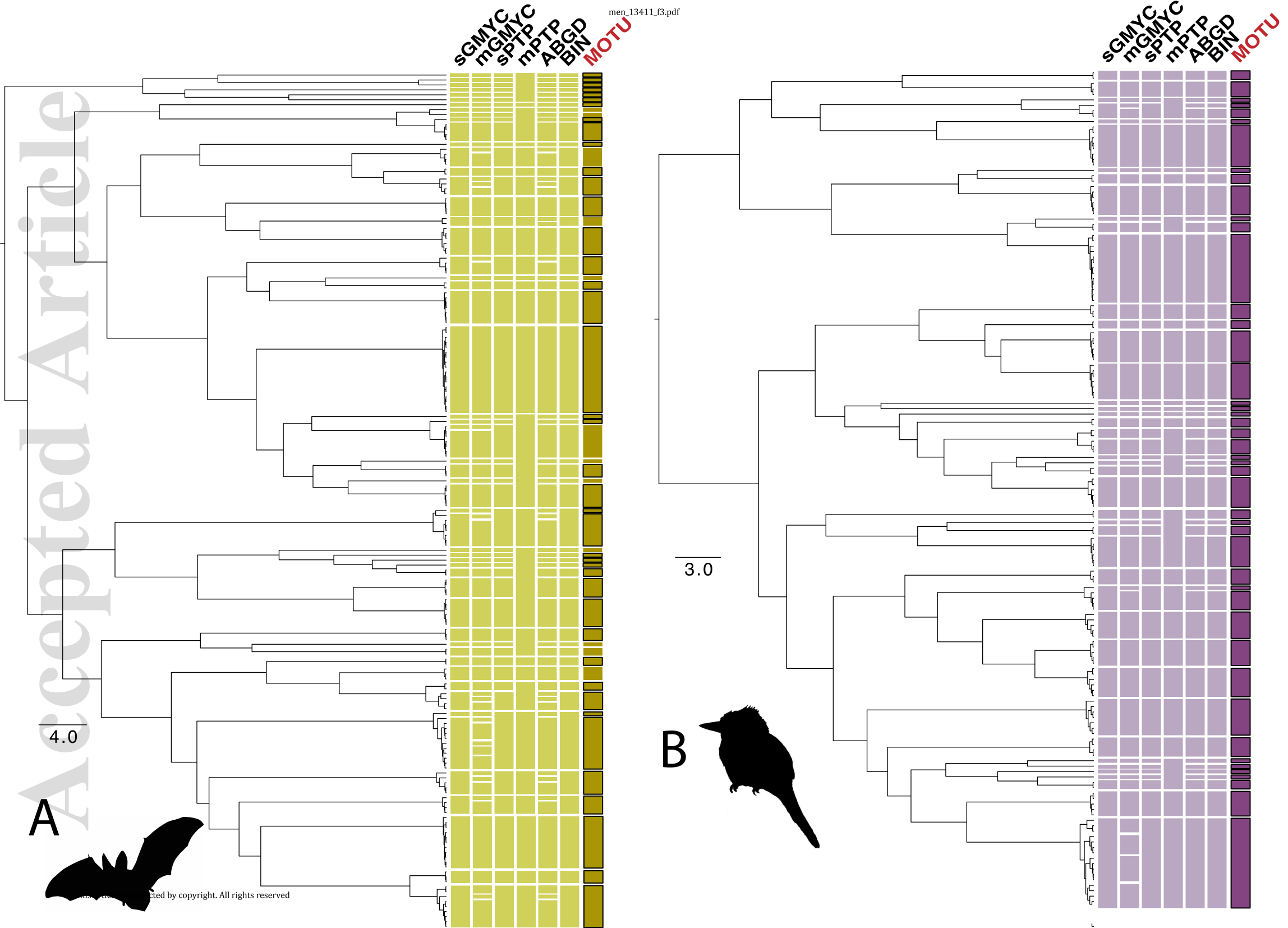
Figure S1. Neighbor-joining tree of the 1005 DNA barcode records.

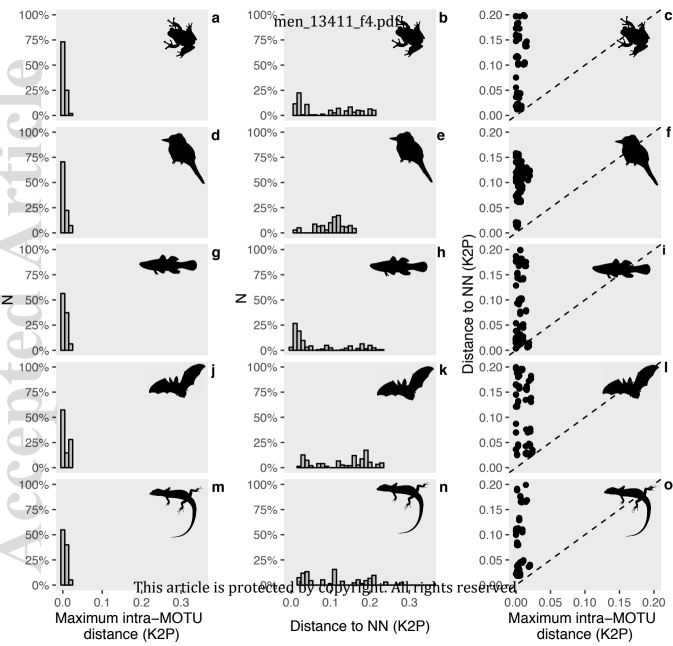
Table S1. List of all specimens barcoded in this study and corresponding results of the MOTU

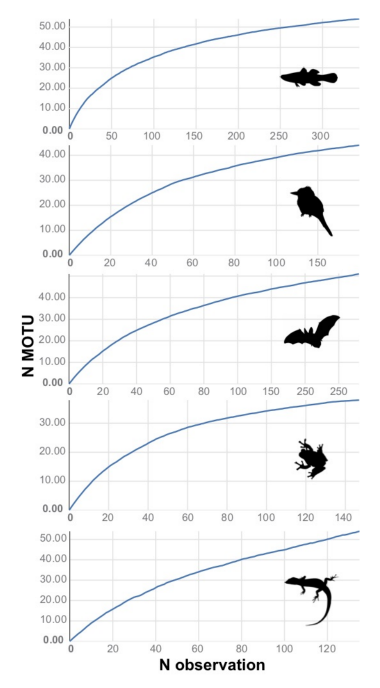
delimitation analyses for sGMYC, mGMYC, sPTP, mPTP, ABGD and RESL.



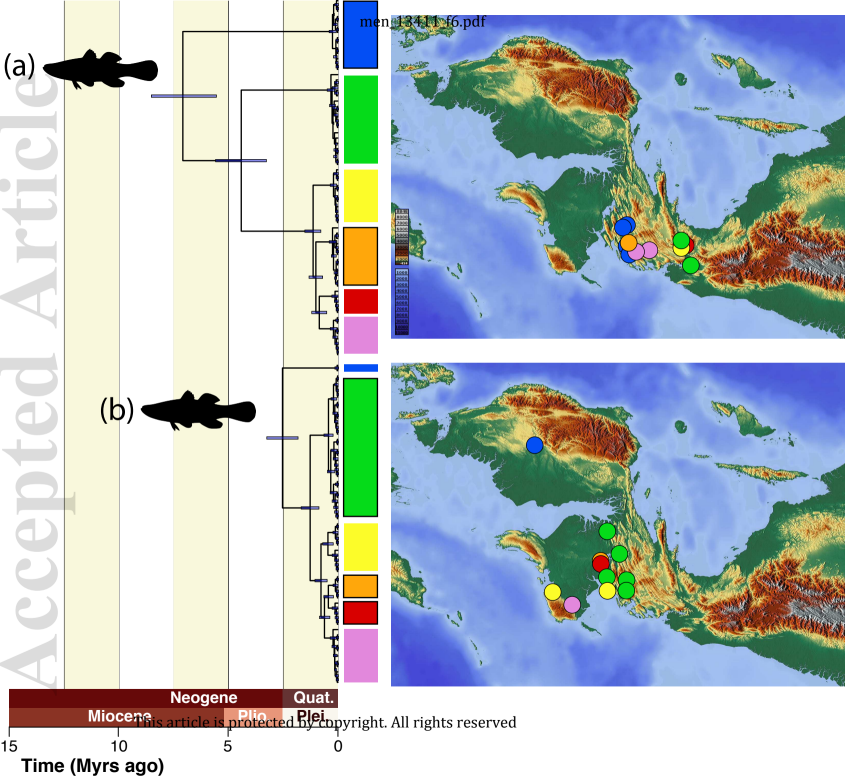


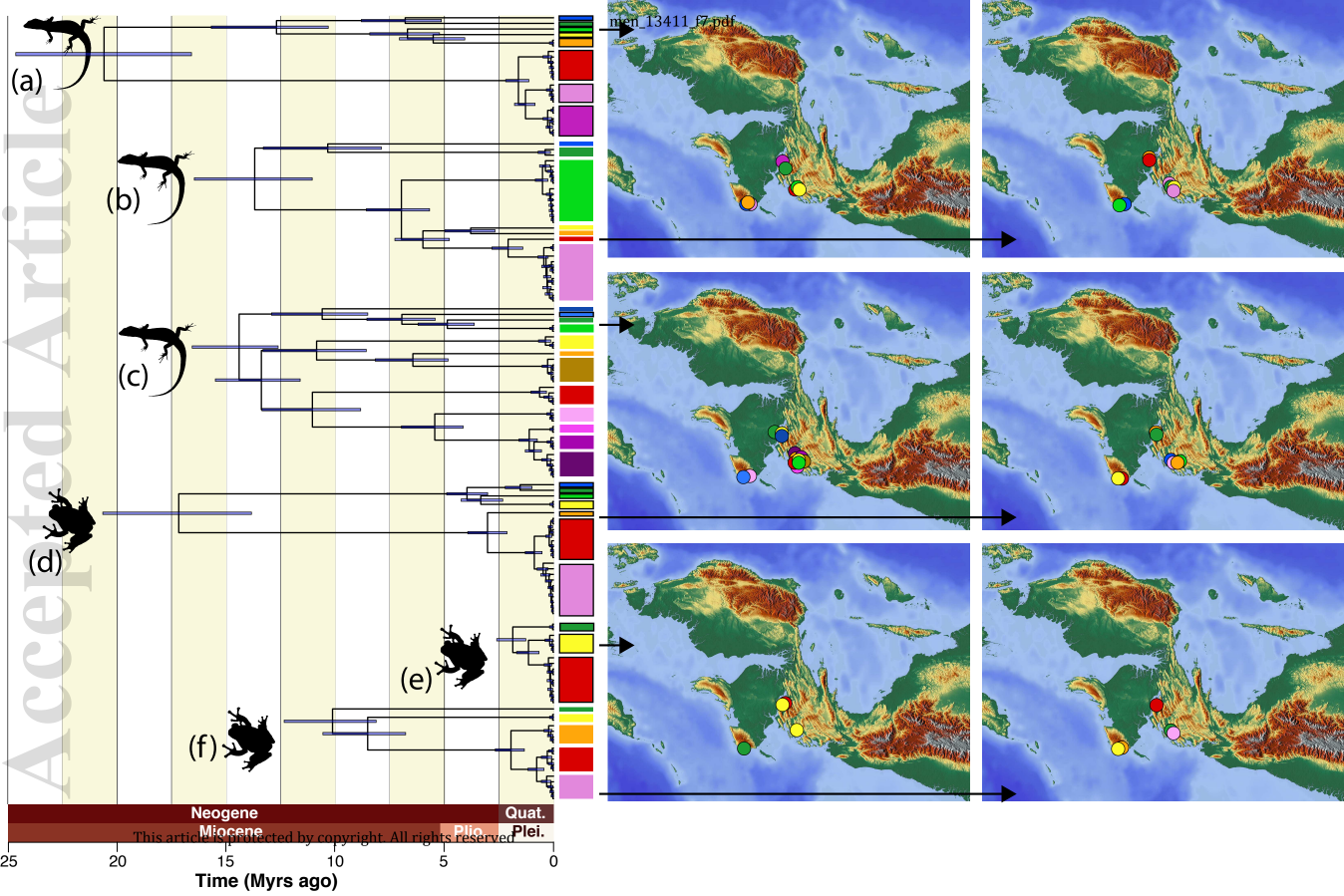






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Systematics	N specimens	
Actinopterygii	343	
Atheriniformes	225	
Atherinidae	12	
Craterocephalus	12	
Melanotaeniidae	213	
Melanotaenia	213	
Clupeiformes	5	
Clupeidae	5	
Nematalosa	5	
Cypriniformes	2	
Cyprinidae	2	
Barbodes	2	
Gobiiformes	51	
Butidae	10	
Butis	8	
Ophiocara	1	
Oxyeleotris	1	
Eleotridae	20	
Eleotris	6	
Giuris	2	
Mogurnda	12	
Oxudercidae	21	
Awaous	2	
Lentipes	1	
Redigobius	3	
Sicyopterus	2	
Stenogobius	2	
Stiphodon	11	
Kurtiformes	1	
Apogonidae	1	
Glossamia	1	
Ovalentaria	2	
Rhyacichthyidae	2	
Rhyacichthys	2	
Perciformes	29	
Kuhliidae	4	
Kuhlia	4	
Leiognathidae	2	
Leiognathus	2	
Lutjanidae	2	
Lutjanus	2	
Terapontidae	6	
Mesopristes	1	
Pingalla	5	
Toxotidae	15	
Toxotes	15	
Siluriformes	20	
Ariidae	12	
Neoarius	12	

Plotosidae	8		
Neosilurus	3		
Porochilus	5		
Syngnathiformes	8		
	8		
Syngnathidae <i>Hippichthys</i>	2		
Microphis	6		
	167		
Amphibia Anura	167		
Ceratobatrachidae	38		
Cornufer	38		
Hylidae	1		
Nyctimystes	1		
Limnodynastidae	1		
Lechriodus	1		
Microhylidae	99		
Asterophrys	16		
Austrochaperina	11		
Callulops	10		
Cophixalus	12		
Hylophorbus	17		
Oreophryne	16		
Sphenophryne	7		
Xenorhina	10		
Pelodryadidae	14		
Litoria	14		
Ranidae	14		
Papurana	14		
Aves	155		
Accipitriformes	4		
Accipitridae	4		
Accipiter	4		
Columbiformes	9		
Columbidae	9		
Chalcophaps	8		
Ptilinopus	1		
Coraciiformes	25		
Alcedinidae	25		
Сеух	16		
Dacelo	2		
Syma	1		
Tanysiptera	6		
Passeriformes	112		
Acanthizidae	18		
Aethomyias	2		
Gerygone	3		
Origma	7		
Sericornis	6		
Artamidae	4		
Melloria	4		
Maluridae	1		
Chenorhamphus	1		

Melanocharitidae	24			
Melanocharis	7			
Toxorhamphus	17			
Meliphagidae	18			
Melilestes	5			
Meliphaga	10			
Myzomela	2			
Xanthotis	1			
Monarchidae	4			
Arses	1			
Symposiachrus	3			
Oriolidae	4			
Pitohui	4			
Pachycephalidae	16			
Colluricincla	6			
Pachycephala	5			
Pseudorectes	5			
Paradisaeidae	2			
Cicinnurus	1			
Diphyllodes	1			
Petroicidae	10			
Heteromyias	1			
Peneothello	2			
Poecilodryas	1			
Tregellasia	6			
Phylloscopidae	2			
Phylloscopus	2			
Pomatostomidae	1			
Garritornis	1			
Psophodidae	3			
Cinclosoma	3			
Rhipiduridae	5			
Rhipidura	5			
Psittaciformes	5			
Psittaculidae	5			
Alisterus	2			
Micropsitta	3			
Mammalia	173			
Artiodactyla	9			
Cervidae	1			
Rusa	1			
Suidae	8			
Sus	8			
Chiroptera	135			
Emballonuridae	13			
Emballonura	12			
Mosia	1			
Hipposideridae	44			
Піррозічениае	44			

Aselliscus	2			
Coelops	1			
Hipposideros	41			
Pteropodidae	55			
Dobsonia	12			
Macroglossus	5			
Nyctimene	9			
Paranyctimene	2			
Pteropus	4			
Rousettus	11			
Syconycteris	12			
Rhinolophidae	7			
Rhinolophus	7			
Vespertilionidae	16			
	10			
Miniopterus	3			
Myotis	3			
Pipistrellus Diprotodontio	6			
Diprotodontia				
Macropodidae	2			
Dorcopsis	2			
Petauridae	2			
Dactylopsila	1			
Petaurus	1			
Phalangeridae	2			
Phalanger	2			
Peramelemorphia	7			
Peramelidae	7			
Echymipera	7			
Rodentia	16			
Muridae	16			
Melomys	1			
Paramelomys	4			
Rattus	10			
Uromys	1			
Reptilia	167			
Squamata	167			
Agamidae	2			
Hypsilurus	2			
Boidae	2			
Candoia	2			
Colubridae	10			
Boiga	1			
Dendrelaphis	1			
Rhabdophis	1			
Stegonotus	5			
Tropidonophis	2			
Elapidae	2			
Aspidomorphus	1			
Micropechis	1			

Total	1005		
Ramphotyphlops	1		
Typhlopidae	1		
Tribolonotus	4		
Tiliqua	3		
Sphenomorphus	39		
Lygisaurus	14		
Eremiascincus	2		
Emoia	47		
Carlia	6		
Scincidae	115		
Simalia	1		
Leiopython	1		
Apodora	1		
Pythonidae	3		
Nactus	1		
Lepidodactylus	1		
Hemidactylus	2		
Gekko	4		
Gehyra	2		
Cyrtodactylus	22		
Gekkonidae	32		

Table 2. Summary statistics of genetic distances including minimum, maximum and average K2P distances within-MOTU, among MOTUs within genus and among MOTUs within family for fishes, birds, mammals, amphibians and reptiles.

Level	Таха	n	N taxa	Comparisons	Min Dist(%)	Mean Dist(%)	Max Dist(%)	SE Dist(%)
Within Species	Fishes	229	27	2215	0.00	0.37	11.14	0.00
	Birds	140	30	426	0.00	0.43	2.48	0.00
	Mammals	140	28	457	0.00	0.64	17.52	0.00
	Amphibians	115	14	580	0.00	1.78	10.87	0.00
	Reptiles	79	14	308	0.00	1.98	17.40	0.01
Within Genus	Fishes	172	5	8541	0.65	9.75	18.02	0.00
	Birds	38	5	81	0.00	8.77	12.87	0.04
	Mammals	82	7	434	0.00	18.66	24.94	0.01
	Amphibians	47	3	191	15.23	21.56	25.99	0.02
	Reptiles	45	4	178	2.81	15.16	24.52	0.05
Within Family	Fishes	57	4	279	11.39	17.79	24.99	0.01
	Birds	131	10	625	6.14	14.24	19.85	0.01
	Mammals	123	6	1254	12.51	20.42	26.95	0.00
	Amphibians	88	2	1339	20.65	25.58	50.00	0.00
	Reptiles	86	5	915	13.67	24.81	36.34	0.00

Table 3. List of the unidentified MOTUs including their genus assignment, consensus MOTU and BIN numbers, maximum within-MOTU K2P distance and K2P distance to the nearest Taxa ▼ MOTU ▼ BIN ▼ Distance max. (%▼ Distance NN (%▼ → † Genus Amphibians Austrochaperina MOTU073 BOLD:ADN9008 0 11.6 Amphibians Austrochaperina MOTU074 BOLD:ADN9009 0.3 11.6 Amphibians *Hylophorbus* MOTU078 BOLD:ADN1146 14.8 0.5 Amphibians Hylophorbus **MOTU080** BOLD:ADN4756 3.5 Amphibians Hylophorbus MOTU081 BOLD:ADO0048 0.2 1.2 MOTU082 BOLD:ADO3938 0.8 Amphibians Hylophorbus 1.2 Amphibians Hylophorbus **MOTU083** BOLD:ADO4150 0 14.8 Amphibians Lechriodus MOTU070 BOLD:ADO3037 22.8 0 Amphibians Oreophryne MOTU071 BOLD:ADN0004 19.7 Amphibians Oreophryne **MOTU083** BOLD:ADO4150 0 17.3 Amphibians Oreophryne MOTU084 BOLD:AED5052 23.5 Amphibians Oreophryne MOTU085 BOLD:AED5053 0.2 17.3 Amphibians Oreophryne MOTU087 BOLD:ADO1683 1.3 19.7 Amphibians Xenorhina MOTU075 BOLD:ADN1930 18.8 Amphibians Xenorhina MOTU094 BOLD:ADO4474 1.5 13.7 MOTU042 BOLD:ADM0638 Fishes Glossamia 21.8 0.4 Fishes Hippichthys MOTU057 BOLD:ADN0840 22.5 0 5.4 Fishes Melanotaenia MOTU002 BOLD:AAY9627 Fishes MOTU003 BOLD:ABY7305 0 1.9 Melanotaenia Fishes MOTU004 BOLD:ABY8664 1.9 0.6 Melanotaenia Fishes MOTU006 BOLD:ACE4002 0 Melanotaenia 1.5 Fishes Melanotaenia MOTU007 BOLD:ADL9884 0 1.5 0.4 Fishes Melanotaenia MOTU008 BOLD:ADM3156 9.1 0.2 Fishes Melanotaenia MOTU009 BOLD:ADM8161 3.6 Fishes **MOTU058** BOLD:ADN5559 1.5 16.7 Microphis 0.4 Fishes **MOTU059** BOLD:ADO3837 Microphis 16.7 Fishes Neoarius MOTU051 BOLD:AAJ9962 1.5 7.7 Fishes Neoarius MOTU052 BOLD:ADL9301 0 3.8 0 Fishes MOTU053 BOLD:ADM1229 2.3 Neoarius 0 Fishes Neoarius MOTU054 BOLD:ADM1230 2.3 Fishes Neosilurus MOTU055 BOLD:ADM9105 0 17.8 Mammals MOTU157 BOLD:ADJ4695 15.7 Coelops Mammals Echymipera MOTU191 BOLD:ADJ2534 3.5 BOLD:ADJ2965 Mammals Echymipera MOTU192 3.5 Mammals MOTU158 BOLD:ADI7709 14.5 Hipposideros Mammals Hipposideros **MOTU159** BOLD:ADI7931 16.5 0 Mammals Hipposideros MOTU160 BOLD:ADJ0463 3.6 Mammals Melomys MOTU195 BOLD:ADI9381 14.7 1.6 Mammals Miniopterus MOTU179 BOLD:ADJ2078 15.4 MOTU170 BOLD:ABV8204 15.9 Mammals Nyctimene 0.5 Mammals Pipistrellus MOTU183 BOLD:ADJ5548 20 Mammals Pipistrellus MOTU184 BOLD:ADJ5623 0 20 Reptilia Boiga MOTU205 BOLD:ADN8334 18.2 MOTU239 BOLD:ADN9150 17.7 Reptilia Emoia

0.2

10.7

Reptilia

Emoia

MOTU240

BOLD:ADO3330

Reptilia	Emoia	MOTU232	BOLD:ADM9606	0	10.9		
Reptilia	Emoia	MOTU233	BOLD:ADM9607	-	10.9		
Reptilia	Emoia	MOTU234	BOLD:ADN0860	0.5	12.9		
Reptilia	Emoia	MOTU235	BOLD:ADN0861	1.6	16.5		
Reptilia	Emoia	MOTU236	BOLD:ADN5383	0.8	16.4		
Reptilia	Emoia	MOTU237	BOLD:ADN9148	-	16		
Reptilia	Emoia	MOTU238	BOLD:ADN9149	-	12.9		
Reptilia	Emoia	MOTU239	BOLD:ADN9150	0	2.2		
Reptilia	Emoia	MOTU240	BOLD:ADO3330	0.6	1.9		
Reptilia	Emoia	MOTU241	BOLD:ADO4309	0.2	1.9		
Reptilia	Eremiascincus	MOTU244	BOLD:ADO2936	-	9.8		
Reptilia	Eremiascincus	MOTU245	BOLD:ADO2937	-	9.8		
Reptilia	Gehyra	MOTU221	BOLD:ADO7212	0.2	24		
Reptilia	Hypsilurus	MOTU202	BOLD:ADN7192	0	37		
Reptilia	Lepidodactylus	MOTU224	BOLD:ADE2841	-	23		
Reptilia	Nactus	MOTU225	BOLD:ADO5284	-	24		
Reptilia	Ramphotyphlops	MOTU264	BOLD:ADN5549	-	31		
Reptilia	Rhabdophis	MOTU207	BOLD:ADN7031	-	10		
Reptilia	Sphenomorphus	MOTU258	BOLD:ADO0519	-	8.7		
Reptilia	Sphenomorphus	MOTU259	BOLD:ADO0520	0.9	16.5		
Reptilia	Sphenomorphus	MOTU260	BOLD:ADO0521	-	21.2		
Reptilia	Sphenomorphus	MOTU252	BOLD:ADN1408	0.2	8.5		
Reptilia	Sphenomorphus	MOTU253	BOLD:ADN1409	0.6	17.5		
Reptilia	Sphenomorphus	MOTU254	BOLD:ADN5384	-	8.5		
Reptilia	Sphenomorphus	MOTU255	BOLD:ADN5577	-	19.7		
Reptilia	Sphenomorphus	MOTU256	BOLD:ADO0517	-	15.8		
Reptilia	Sphenomorphus	MOTU257	BOLD:ADO0518	-	4.3		
Reptilia	Sphenomorphus	MOTU258	BOLD:ADO0519	0.9	10.5		
Reptilia	Sphenomorphus	MOTU259	BOLD:ADO0520	1.1	4.3		
Reptilia	Sphenomorphus	MOTU260	BOLD:ADO0521	-	8.7		

Table 4. List of MOTUs assigned to the same species based on morphological characters, including MOTU and BIN numbers, maximum within-MOTU K2P distance (percent) and K2P distance to the nearest neighbor (percent).

Taxa 🚅	Genus		BIN -	1.	Distance N
	Cornufer papuensis	MOTU060	BOLD:ADN5223	0.8	1.4
-	Cornufer papuensis	MOTU061	BOLD:ADN6050	-	6.5
-	Cornufer papuensis	MOTU062	BOLD:ADN9748	0.3	1.4
Amphibia		MOTU063	BOLD:ADO0083	0	7.5
-	Cornufer bimaculatus	MOTU064	BOLD:ADO0084	-	7.5
	Cornufer bimaculatus	MOTU065	BOLD:ADO0085	-	3.9
-	Cornufer bimaculatus	MOTU066	BOLD:ADO0272	-	3.9
Amphibia	Cornufer punctatus	MOTU067	BOLD:ADN3984	0.6	4.3
-	Cornufer punctatus	MOTU068	BOLD:ADN5222	0.2	4.3
Amphibia	Asterophrys pullifer	MOTU088	BOLD:ADO1683	0.2	2.1
Amphibia	Asterophrys pullifer	MOTU089	BOLD:ADO2921	O	3.7
Amphibia	Asterophrys pullifer	MOTU090	BOLD:ADO2922	0.2	2.1
Amphibia	Sphenophryne cornuta	MOTU091	BOLD:ADN1659	-	2.4
	Sphenophryne cornuta	MOTU092	BOLD:ADO0053	O	1.9
	Sphenophryne cornuta	MOTU093	BOLD:ADO0994	O	1.9
Amphibia	Litoria infrafrenata	MOTU098	BOLD:AAN2556	0.3	2
Amphibia	Litoria infrafrenata	MOTU099	BOLD:ADN6283	0.3	2
Aves	Melilestes megarhynchus	MOTU121	BOLD:AAF2363	O	1.5
Aves	Melilestes megarhynchus	MOTU122	BOLD:AAF2363	0.3	1.5
Fishes	Giuris margaritaceus	MOTU028	BOLD:AAK3399	-	10.8
Fishes	Giuris margaritaceus	MOTU029	BOLD:ADM7171	-	10.8
Fishes	Mogurnda mogurnda	MOTU030	BOLD:AAD3229	O	2.1
Fishes	Mogurnda mogurnda	MOTU031	BOLD:AAD3229	1	2.1
Fishes	Mogurnda mogurnda	MOTU032	BOLD:AAD3229	O	2.5
Fishes	Toxotes oligolepis	MOTU049	BOLD:ADM9057	0.4	1.9
Fishes	Toxotes oligolepis	MOTU050	BOLD:ADM9058	0.2	1.9
Mammals	Sus scrofa	MOTU150	BOLD:AAA3445	2.4	2.7
Mammals	Sus scrofa	MOTU151	BOLD:AAA3445	-	2.7
Mammals	Nyctimene albiventer	MOTU171	BOLD:ABV8022	0.3	4.3
Mammals	Nyctimene albiventer	MOTU172	BOLD:ADI7768	2	4.3
Mammals	Syconycteris australis	MOTU176	BOLD:ABV9982	-	2.7
Mammals	Syconycteris australis	MOTU177	BOLD:ABV9982	1.7	2.7
Mammals	Dorcopsis muelleri	MOTU185	BOLD:ADI7672	-	5.7
Mammals	Dorcopsis muelleri	MOTU186	BOLD:ADI9488	-	5.7
Mammals	Echymipera kalubu	MOTU193	BOLD:ADI7714	0.2	2.5
Mammals	Echymipera kalubu	MOTU194	BOLD:ADJ1796	-	2.5
Mammals	Paramelomys platyops	MOTU197	BOLD:ADI8691	0	8.7
Mammals	Paramelomys platyops	MOTU198	BOLD:ADJ4724	-	8.7
Reptiles	Cyrtodactylus irianjayaensis	MOTU213	BOLD:AED5725	-	10.5
Reptiles	Cyrtodactylus irianjayaensis	MOTU214	BOLD:AED5297	-	12
Reptiles	Cyrtodactylus irianjayaensis	MOTU215	BOLD:ADN4791	0	10.7
Reptiles	Cyrtodactylus irianjayaensis	MOTU216	BOLD:ADN4792	-	12
Reptiles	Cyrtodactylus irianjayaensis	MOTU217	BOLD:ADN9189	_	10.5
Reptiles	Cyrtodactylus sermowaiensis	MOTU219	BOLD:ADO1322	0.5	2.7
Reptiles	Cyrtodactylus sermowaiensis	MOTU220	BOLD:ADO1323	0.2	2.7
Reptiles	Lygisaurus novaeguineae	MOTU246	BOLD:ADN7325	0.3	3.6
Reptiles	Lygisaurus novaeguineae	MOTU247	BOLD:ADN8226	2	3.6
Reptiles	Lygisaurus novaeguineae	MOTU248	BOLD:ADO2052	-	6.1