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# Sorting the wheat from the chaff: a review of BINs associated with groupers of Vietnam and the implications for species identification from DNA barcoding

Jean-Dominique Durand<sup>1</sup>, Manh Hung Pham<sup>2</sup>, Thanh Thi Viet Tran<sup>3</sup>, Duc Huy Hoang<sup>2</sup>, Quang Van Vo<sup>4</sup>

The Epinephelidae is a marine fish family that has been the focus of worldwide taxonomical attention due to its economic value. Recent molecular-based phylogenies have improved understanding of species diversity within this family. Nonetheless, species misidentification or hybridization has led to a barcode index number (BIN) being associated with several species, which hampers a clear species identification within the BOLD system. In this study, grouper diversity in Vietnam was investigated to check the species inventory, provide a DNA barcode library for Vietnam, and reevaluate BINs associated with some species that are present in Vietnam. To this end, 157 specimens were barcoded corresponding to 30 species and 31 BINs. Nine species were new records, bringing to 49 the number of species inventoried for Vietnam. In BOLD, these species are associated with 75 BINs, 31 species being represented by more than one BIN. A careful review of these BINs, considering the present results from Vietnam, the species composition and frequency of each BIN in BOLD, and reference species sequences from phylogenetic studies, revealed those misidentified or hybrid specimens that artificially increased the number of BINs per species. After appropriate revision, 15 species still remained associated with more than one BIN. A phylogeographic analysis of these species demonstrated that most of the BINs derived from evolutionary lineages with geographic distributions that match well known biogeographic units in the Indo-Pacific. Beyond the species identification, these multiple BINs of each species can be used to track the biogeographic origin of each of the associated specimens.

Keywords Cytochrome oxidase I · BOLD · Misidentification · Biogeography · Epinephelidae

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### Introduction

DNA barcoding in fishes is now widely used for taxonomical investigations (Hubert and Hanner 2015), biodiversity evaluations (Galdino Brandao et al. 2016; Shen et al. 2016; Durand et al. 2017), and food control (Barcaccia et al. 2015; Brito et al. 2015). One of the main results of the broad scan of fish diversity across geographic areas, using cytochrome oxydase I (COI) polymorphisms, has been the revelation of numerous species complexes or cryptic species, leading to a profound change to our vision of biodiversity (Hubert et al. 2012, 2017; Jaafar et al. 2012; Winterbottom et al. 2014; Durand and Borsa 2015; Williams and Viviani 2016; Habib et al. 2018; Delrieu-Trottin et al. 2019). This important result justifies more barcoding investigations because gaps in taxonomical coverage limit accuracy of the species identification based on a DNA barcode. However, if the BOLD database is used as reference library and is continuously enriched with new barcodes, then there is also an increase in mis-labelled sequences due to misidentification, which is a serious concern (Vilgalys 2003; Ward et al. 2009). By comparing sequences of a marine fish family from a reference library to those available in Genbank, an important source of barcodes for BOLD, Durand et al. (2017) demonstrated that misidentified species of a fish family were more frequent in barcoding studies than other types of studies like molecular phylogeny and phylogeography. This may be due to the difficulties of developing reliable systematic expertise for a wide range of fish taxa, for studies that do not target specific families.

As a consequence, while barcoding studies are still necessary to fill taxonomical knowledge gaps and solve some taxonomical issues, it is also necessary to review the DNA barcodes in BOLD, to better define species boundaries and track species misidentification. To reach this objective, new barcoding projects could be (1) more taxonomically restricted so as to limit potential species identification problems, (2) increase the number of specimens and sampling localities over the species distribution range so as to better estimate levels of intraspecific diversity, and (3) consider all DNA barcodes stored in the BOLD library, to track both mislabelled sequences and putative undescribed or cryptic species. While individual review of DNA barcodes in the BOLD library would be particularly tedious, the process could be facilitated by the barcode index number (BIN) provided by BOLD system that corresponds to an operational taxonomical unit (OTU). This OTU consists of barcodes that are identical or phylogenetically closely related and separated from others in the library by a barcode gap. OTUs are delimited in BOLD, using the Refined Single Linkage (RESL) algorithm (Ratnasingham and Hebert 2013). The BIN was created to circumvent taxonomical issues linked to identification problems, undescribed species or, more generally, to reveal cryptic diversity (Ratnasingham and Hebert 2013).

The grouper, Epinephelidae, is a marine fish family present in all temperate and tropical marine coastal waters. They comprise economically important species that are the target of fisheries and aquaculture. For this reason, the family has been the focus of a series of molecular phylogenetic and barcoding studies (Craig and Hastings 2007; Craig et al. 2011; Chu et al. 2011; Sachithanandam et al. 2012; Alcantara and Yambot 2016; Ma et al. 2016, 2018; Basheer et al. 2017). Among these studies, molecular phylogenetic or phylogeographic studies highlighted systematics issues such as the paraphyly of Cephalopholis (Bloch & Schneider, 1801) and Epinephelus (Bloch, 1793) genera, and the presence of cryptic species (Craig and Hastings 2007; Ma et al. 2016; Borsa et al. 2016; Ma and Craig 2018). In contrast, although all region-based barcoding studies provided new COI barcodes for groupers collected in countries such as India (Sachithanandam et al. 2012; Basheer et al. 2017), Malaysia (Chu et al. 2011), or the Philippines (Alcantara and Yambot 2016), none considered the barcodes in BOLD in order to review intraspecific diversity over a broader geographic scale than their barcoded species or the BIN diversity per species, which represent an alternative way to both track mis-labelling sequences or polyphyletic species.

Within global grouper diversity, 34 to 39 species may occur in Vietnamese waters (Heemstra and Randall 1993, 1999; Craig et al. 2011). Among these species, some are uncommon or have a limited distribution range. Due, however, to the difficulty in identifying some species by morphometric or phenotypic characters (Heemstra and Randall 1993; Guo et al. 2014; Alcantara and Yambot 2016) the species inventory should be considered with caution. Furthermore, new grouper species are still being described, such as Epinephelus kupangensis Tucker et al., 2016, Epinephelus craigi Frable et al., 2018 and Epinephelus fuscomarginatus Johnson & Worthington Wilmer, 2019. Other species have debateable taxonomic status, such as Epinephelus moara (Temminck & Schlegel, 1843) that is considered as valid by several authors (Guo et al. 2009; Liu et al. 2013; Guo et al. 2014) but a junior synonym of Epinephelus bruneus (Bloch, 1793) by others (Heemstra and Randall 1993; Craig et al. 2011; Fricke et al. 2019). For all these reasons, phylogenetic and barcoding studies are needed to clarify the species diversity present in Vietnam where barcoding analyses are rare (Thai et al. 2007; Tran et al. 2016; Ha et al. 2018; Huang et al. 2018) and missing for the Epinephelidae family.

In this context, this study aimed to create a reference barcoding library for Vietnam and test whether this approach can be used to update the species checklist and flag cryptic or undescribed species. Our ambition with this region-based barcoding study was also to review all BINs in BOLD for species present in Vietnam and determine significance of the presence of several BINs associated with a species. The last goal requires combining all available data to determine the congruency between species name associated with a BIN and some species' sequence considered as a reference (see hereafter), and investigate the phylogeographic structure of species associated with more than one BIN. Finally, this study represents a good opportunity to review the BOLD library and tracked misidentified specimens that can limit accuracy of species identification.

## **Material and methods**

#### Taxonomy and species checklist for Vietnam

The taxonomy and classification of groupers is debated (Ma and Craig 2018) because successive molecular phylogenetic studies have highlighted a number of inconsistencies, such as the non-monophyly of the Serranidae family and of some genera within the Epinephilinae subfamily (Maggio et al. 2005; Ding et al. 2006; Craig and Hastings 2007; Craig et al. 2011; Zhuang et al. 2013; Schoelinck et al. 2014; Ma et al. 2016). To resolve this issue, some authors have proposed an updated taxonomy (Smith and Craig 2007; Ma and Craig 2018). Within this, the subfamily "Epinephelinae" (sensu Johnson 1983) was elevated to family status and the tribes Diploprionini, Epinephelini, Grammistini, and Liopropomini to subfamilies within Epinephelidae (sensu Smith and Craig 2007). Furthermore, the species classification of the Epinephelinae (sensu Smith and Craig 2007) was revised according to their molecular phylogeny (Ma et al. 2016).

In the current study, the taxonomy proposed by Smith and Craig (2007) has been adopted but we have conserved the "traditional" genus naming of species of the Epinephelinae, since this has not yet been adopted by Eschemeyer's Catalog of Fishes (Fricke et al. 2019) and thus not recognized by BOLD or GenBank. However, in Table 1, the second column has the revised taxonomic nomenclature of Ma and Craig (2018), involving the genus replacement of *Aethaloperca* by *Cephalopholis*, *Anyperodon*, and *Cromileptes* by *Epinephelus*; *Triso* by *Hyporthodus*; and *Epinephelus* by *Mycteroperca* for the following species: *E. epistictus*, *E. heniochus*, *E. morrhua*, and *E. poecilonotus*.

The species checklist of Epinephelidae present in Vietnam (Table 1) has been established on the basis of two references, Heemstra and Randall (1993) and Craig et al. (2011). One additional species have been added to this list, *Epinephelus craigi* because it was recently described in the South China Sea, probably including Vietnamese waters (Nha Trang) according to Frable et al. (2019).

### Sampling

Grouper specimens were collected between 2015 and 2017 from major fish landings and markets at 18 locations belonging to 11 provinces in Vietnam (Fig. 1). We strived to obtain as many species as possible, to provide the broadest overview of diversity in Vietnam. When possible, several specimens from the same species were collected at different locations, to provide some information about intraspecific diversity and distribution ranges in Vietnam. A total of 157 groupers were collected, plus one specimen of Diploprion bifasciatum belonging to the Diploprioniae subfamily, used as outgroup (Table S1). An initial species identification was performed on site using the FAO identification sheet (Heemstra and Randall 1993) and Craig et al. (2011). Each specimen was photographed and a small section of the pectoral fin stored in ethanol 70° in a 1.5 ml Eppendorf, which were then stored at -20 °C in the laboratory.

#### **DNA extraction and barcoding**

Genomic DNA was isolated from fins using the G-Spin Total DNA extraction mini kit (iNtRON biotechnology, Korea) following the manufacturer's recommendations. Approximately 655 base pairs (bp) of the cytochrome oxydase I (COI) were amplified using the reverse primer FishR1 and the two forward primers in cocktail: FishF1 and FishF2 (Ward et al. 2005). The 40  $\mu$ l PCR reaction mixes included 20  $\mu$ l of MyTaq PCR Mastermix (Bioline), 16  $\mu$ l of ultrapure water, 0.8  $\mu$ l of BSA (Euromedex), 0.6  $\mu$ l of each primers (3  $\mu$ M), and 2  $\mu$ l of DNA template. Amplifications were performed in a PTC-100 BIORAD Thermal cycler. The thermal regime consisted of an initial step of 2 min at 92 °C followed by 35 cycles of 45 s at 92 °C, 45 s at 52 °C, and 1 min at 72 °C, followed in turn by 5 min at 72 °C and then held at 4 °C. PCR products were visualized on 1–2% agarose gels and the most intense products were selected for sequencing by Macrogen (https://dna.macrogen.com/).

Electropherograms were visually checked using 4Peaks by A. Griekspoor and Tom Groothuis, nucleobytes.com, and sequences aligned using MUSCLE implemented in MEGA7 (Kumar et al. 2016). DNA barcodes and specimens' data are available in the project file DS-SERV: Epinephelidae of Vietnam on BOLD systems site dx.doi.org/10.5883/DS-SERV). Each DNA barcode was assigned to a BIN by the BOLD system and deposited in GenBank under accession number MN708800 to MN708957 (Table S1). Sequence divergences (Table S2) were calculated using the Kimura two parameters (K2P) distance model (Kimura 1980). Neighborjoining (NJ) trees of K2P distances were drawn to highlight pattern of divergence between species (Saitou and Nei 1987). A bootstrapping test was performed in MEGA3 (Kumar et al. 2004) with 500 replications, to test the relevance of the tree's nodes.

#### Species identification and species boundaries

The accuracy of the OTU delineation provided by BOLD using the RESL algorithm was first evaluated by comparing BINs to OTUs identified by the Automatic Barcode Gap Delineation (ABGD) algorithm (Puillandre et al. 2012). These two methods used DNA alignments as inputs, consisting of ABGD, of all DNA barcodes (with a core segment of a 502 bp) constituting BINs associated with species name listed in Table 1, plus all Vietnamese barcodes generated in this study (Table S3). The final fasta file contained 1648 sequences and was uploaded to the website that hosted the ABGD algorithm to run the analyses (https://bioinfo.mnhn. fr/abi/public/abgd/abgdweb.html).

The first phylogenetic analyse was performed on a dataset composed exclusively of DNA barcodes obtained in this study, i.e., the groupers sampled in Vietnam. The tree was rooted with *Diploprion bifasciatum*. The species name recovered using the morphology and the BIN provided by BOLD was mapped onto this tree, to flag polyphyletic species and to reveal intraspecific and interspecific diversity within the Epinephelidae.

Craig et al. (2011) each species, the a	and this study. In bol associated BINs in BC	d, species name that DLD, in Ma et al. (20	represent new rec [16] and in this s	ords for tudy, are	Vietnar Vietnar e listed.	a. For with the species name are ment in the (see Table S3 for the rationale)	tioned in brackets,	probable species mi	isidentific	ations are in italics
Classification "Catalog of Fishes"	Classification according Ma and Craig (2018)		Heemstra and Randall 1993 (FAO)	Craig et al. 2011	This study	BIN's BOLD	Ma et al. (2016)	BIN present in Vietnam (this study)	Multi- BIN species	Phylogeographic pattern
Aethaloperca	Cephalopholis roadd	Forsskål, 1775	X	×		BOLD:AAD9665 [16/15], BOI D:ACC0322 [3/3]	BOLD:AAD9665	NA	έΥ	z
nogua Anyperodon leucogrammic- us	Epinephelus leucogrammicus	Valenciennes in Cuvier & Valenciennes, 1828	X	X		BOLD:AAC8328 [7/6], BOLD:AAC8327 [5/5]	BOLD:AAC8328	NA	Y	Y
Cephalopholis argus	I	Bloch & Schneider, 1801	Х	Х		BOLD:AAC4474 [26/24], BOLD:AAC0217 [21/1] , BOLD:AAC0217 [21/1] ,	BOLD:AAC4474	NA	Z	Z
Cephalopholis boenak	I	Bloch, 1790	×	x	×	BOLD:ACT7478 [32/31], BOLD:ACT7478 [32/31], BOLD:AAB3684 [7/7], BOLD:ACT7479 [1/7],	BOLD:ACT7478	BOLD:ACT7478	Y	Y
Cephalopholis cyanostigma	I	Valenciennes in Cuvier & Valenciennes,	×	×	×	BOLD:AAC0217 [21/15]	BOLD:AAC0217	BOLD:AAC0217	z	Z
Cephalopholis 5	I	Shaw & Nodder,	Х	Х	Х	BOLD:AAF0621 [29/27]	BOLD:AAF0621	BOLD:AAF0621	z	Z
jormosa Cephalopholis	Ι	1812 Lacepède, 1802		Х		BOLD:AAD9032 [7/7]	BOLD:AAD9032	NA	Z	I
teoparaus Cephalopholis	I	Bleeker, 1852	Х	X		BOLD:AAC0217 [21/4],	BOLD:AAC0217	NA	Z	Í
microprion Cephalopholis	I	Forsskål, 1775	X	Х	Х	BOLD:AAF0629 [4/4] BOLD:AAC0216 [28/28],	BOLD:AAC0216	BOLD:AAC0216	z	N
mınıata Cephalopholis sonnerati	I	Valenciennes in Cuvier &	х	х	х	BOLD:AAC021/ [21/1] BOLD:AAD4737 [26/25], BOLD:AAB5431 [15/14].	BOLD:AAD4737	BOLD:AAD4737	Υ	Υ
		Valenciennes, 1828				BOLD:ACC0647 [5/5], BOLD:AAB5432 [6/2]				
Cephalopholis urodeta	I	Forster in Bloch & Schneider 1801	X	X	X	BOLD:AAC4202 [234/93]	BOLD:AAC4202	BOLD:AAC4202	z	Z
Cromileptes altivelis	Epinephelus altivelis	Valenciennes, 1001 1828	Х	Х		BOLD:AAC1876 [16/16]	BOLD:AAC1876	NA	z	I
Epinephelus akaara	-	Temminck & Schlegel 1842	Х	Х		BOLD:AAF1053 [83/81], BOI D:AAC5362 [50/11	BOLD:AAF1053	NA	z	I
Epinephelus amblycephalus	I	Blecker, 1857	×	X	X	BOLD:AAB1345 [67/48] , BOLD:AAB1346 [57/48] , BOLD:AAB1346 [3/3], BOLD:AAB1346 [3/3],	BOLD:AAB1345	BOLD:AAB1346	Z	Z
Epinephelus areolatus	I	Forsskål, 1775	×	Х	Х	BOLD:AA9821 [37]36], BOLD:AA9822 [37]36], BOLD:AA9822 [37]33], BOLD:AA9822 [37]33],	BOLD:AAA9821	BOLD:AAA9821, BOLD:AA- A0877	Y	Y
Epinephelus awoara	I	Temminck & Schlegel, 1842	Х	Х	х	BOLD:AAF0829 [8/7], BOLD:AAB8391 [113/2]	BOLD:AAF0829	BOLD:AAF0829	z	

Table 1 (contin	nued)									
Classification "Catalog of Fishes"	Classification according Ma and Craig (2018)		Heemstra and Randall 1993 (FAO)	Craig et al. 2011	This study	BIN's BOLD	Ma et al. (2016)	BIN present in Vietnam (this study)	Multi- BIN species	Phylogeographic pattern
Epinephelus	1	Vaillant, 1878	X	x	x	BOLD:AAC5362 [50/48]	BOLD:AAC5362	BOLD:AAC5362	z	N
Dieekeri Epinephelus hruneus	I	Bloch, 1793			Х	BOLD:AAJ0588 [6/6] , BOLD:ACS0266 [3/3]	BOLD:AAJ0588	BOLD:ACS0266	Z	Z
Epinephelus chlorostigma	I	Cuvier & Valenciennes, 1828	Х	×	X	BOLD:AC5363 [19/14] , BOLD:ABA7404 [5/5] BOLD:ABA7404 [5/5]	BOLD:AAA9822	BOLD:AAC3591	Υ	I
Epinephelus coioides	I	Hamilton, 1822	Х	Х	×	BOLD:ADB339[[1137/6], BOLD:ADB339[[1137/6], BOLD:ADB3399[35/1], BOLD:AAC7807 [19/1]	BOLD:AAB8391	BOLD:AAB8391	Z	N
Epinephelus corallicola	I	Valenciennes in Cuvier & Valenciennes, 1828		×	×	BOLD:AAD8682 [9/9]	BOLD:AAD8682	BOLD:AAD8682	Z	I
Epinephelus	I	Frable, Tucker &				The name "craigi" is not in BOLD but the	NA	NA	I	I
craugt Epinephelus	I	w alker, 2019 Richardson, 1846	Х	х		sequence match BOLD:AAD1767 [16/3]	BOLD:AAD1767	NA	Z	I
cyanopodus Epinephelus	Mycteroperca	Temminck &	Х	х		BOLD:ADB3169 [13/13],	BOLD:AAF1066	NA	Υ	Υ
epistictus Enirophology	epistictus	Schlegel, 1842	>	>	>	BOLD:AAF1066 [4/4] BOLD:ACC0501 [15/11]	NIA		Z	Z
erythrurus erythrurus	I	valencientes III Cuvier & Valenciennes, 1828	<	<	<		<b>W</b>	DULDAUCUU	Z	2
Epinephelus fasciatomacul- osus	1	Peters, 1866	Х	×		BOLD:AAC5392 [11/2], BOLD:AAF0829 [8/1] , BOLD:AAF1053 [8/2]	BOLD:AAC5392	NA	Z	I
Epinephelus fasciatus	I	Forskål, 1775	×	×	×	BOLD: AABI 332 [30/29] BOLD: AABI 332 [30/29] BOLD: AABI 333 [23/18], BOLD: AABI 333 [4/3], BOLD: AABI 335 [1/1]	BOLD:AAB1333	BOLD:AAB1332	Y	Y
Epinephelus	I	Forsskål, 1775			х	BOLD:AAD8872 [23/18],	BOLD:AAD8872	BOLD:AAD8872	Z	Y
<b>Juscogunatus</b> Epinephelus	Mycteroperca	Fowler, 1904	Х	X		BOLD:AAD3855 [5/5]	NA	NA	I	I
heniochus Eninanhalus	heniochus	Forster in Bloch &			×	BOLD: A A D3458 [11/10]	BOI D. A AD3458	ROI D-A AD3458	Z	~
Lpureprieus hexagonatus		Schneider, 1801			<			OCTONIC ALLON	-	T
Epinephelus	I	Bloch, 1790	Х	X		BOLD:AAD8914 [6/6], BOLD:AAD8872	BOLD:AAD8914	NA	Z	I
Epinephelus	I	Temminck &	Х	Х		BOLD:AAD8911 [9/9],	BOLD:ADM3791	NA	Y	Y
latifasciatus Epinephelus	I	Schlegel, 1842 Bleeker, 1855		x		BOLD:ADM3791 [5/5] BOLD:AAE1882 [11/9],	BOLD:AAE1882	NA	Z	
macrospilos Epinephelus malabaricus	I	Bloch & Schneider, 1801		×	×	BOLD:AAB8386 [42/4] BOLD:AAB8389 [35/27], BOLD:AAB8391 [113/1]	BOLD:AAB8389	BOLD:AAB8389	Z	Y

Table 1 (contin	ued)									
Classification "Catalog of Fishes"	Classification according Ma and Craig (2018)		Heemstra and Randall 1993 (FAO)	Craig et al. 2011	This study	BIN's BOLD	Ma et al. (2016)	BIN present in Vietnam (this study)	Multi- BIN species	Phylogeographic pattern
Epinephelus merra	1	Bloch, 1793	x	×		BOLD:AAB8386 [42/38], Rol D:AAB8387 [13/4]	BOLD:AAB8386	NA	Y	Y
Epinephelus morrhua	Mycteroperca morrhua	Valenciennes in Cuvier & Valenciennes, 1833	×	×	×	BOLD:ACE6821 [15/14], BOLD:ADL0994 [13/4], BOLD:AAC6284 [22], BOLD:AAC6284 [22],	BOLD:AAC6284	BOLD:ACE6821	¥	Z
Epinephelus poecilonotus	Mycteroperca poecilonotus	Temminck and Schlegel, 1842	×	Х	>	BOLD:AD3854 [30/28], BOLD:AAD3855 [5/1] BOLD:AAD3855 [5/1]	BOLD:AAD3854		z >	I >
Epunepnetus polyphekadion	1	Bleeker, 1049			<	BOLD:ABX5597 [29/1], BOLD:ABX5597 [9/9]	BULD:AAD00/2	BULD:AAD	I	I
Epinephelus quoyanus	1	Valenciennes in Cuvier & Valenciennes, 1830	X	×	×	BOLD:AAC2809 [18/18], BOLD:AAC5362 [50/1]	BOLD:AAC2809	BOLD:AAC2809	z	Y
Epinephelus rivulatus	I	Cuvier and Valenciennes, 1830	X	X	×	BOLD:ACZ9919 [22/17], BOLD:AAF0433 [3/1]	BOLD:ACZ9919	BOLD:AAF0433	Y	Y
Epinephelus sexfasciatus	I	Valenciennes in Cuvier & Valenciennes, 1828	×	×	×	BOLD:AAC5392 [11/9], BOLD:ACM3827 [13/13]	BOLD:AAC5392	BOLD:ACM3827	Z	Z
Epinephelus stictus	Ι	Randall and Allen, 1987	X	x		BOLD:AAB1345 [67/18]	NA	NA	I	I
Epinephelus trimaculatus	I	Cuvier & Valenciennes, 1828		Х		BOLD:AAJ0586 [6/4]	BOLD:AAJ0586	NA	Z	1
Plectropomus leonardus	I	Lacepède, 1802	Х	×	x	BOLD:AAC9348 [86/54]	BOLD:AAC9348	BOLD:AAC9348	z	Y
Plectropomus maculatus	I	Bloch, 1790			Х	BOLD:ACK7859 [16/16], BOLD:AAC9348 [86/9] , BOLD:ADL8508 [4/4]	BOLD:ACK7859	BOLD:ACK7859	¥	Y
Plectropomus oligacanthus	I	Bleeker, 1855			х	BOLD:AAL4434 [12/12]	BOLD:AAL4434	BOLD:AAL4434	z	I
Triso dermonterus	Hyporthodus dermonterus	Temminck and Schlegel 1842			х	BOLD:AAD9769 [6/6]	BOLD:AAD9769	BOLD:AAD9769	z	I
Variola		de Baissac, 1953			Х	BOLD:AAC9675 [22/20]	BOLD:AAC9675	BOLD:AAC9675	z	Υ
unnunginuu Variola louti	I	Forsskål, 1775			X	BOLD:AAC5719 [30/25], BOLD:AAC5720 [3/3]	BOLD:AAC5720	BOLD:AAC5719	Υ	Y

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**Fig. 1** Sampling locations in Vietnam for the DNA barcoding of the Epinephelidae subfamily. 1 Tien Yen, Quang Ninh; 2 Co To Island, Quang Ninh; 3 Ha Long bay, Quang Ninh; 4 Do Son, Hai Phong; 5 Xuan Thuy, Nam Dinh; 6 Hau Loc, Thanh Hoa; 7 Nui Thanh, Quang Nam; 8 Quy Nhon, Binh Dinh; 9 Cu Mong, Phu Yen; 10 Thuy Hoa, Phu Yen; 11 Nha Trang, Khanh Hoa; 12 Binh Ba Island, Khanh Hoa; 13 Ninh Thuan, Ninh Thuan; 14 Loc An, Ba Ria Vung Tau; 15 Con Dao, Ba Ria Vung Tau; 16 Nam Du, Kien Giang; 17 Ha Tien, Kien Giang; 18 Phu Quoc, Kien Giang

The second phylogenetic analysis was performed on a large dataset that comprised:

 One sequence representative of each species and lineage (BIN) observed in this study.

- All sequences published by Ma et al. (2016) that represent 87% of the known species diversity in the Epinephelidae and that may be considered as the reference.
- The COI sequences of Epinephelus moara resurrected by Liu et al. (2013), Epinephelus craigi described by Frable et al. (2019), and Epinephelus fuscomarginatus + E. magniscuttis described and redescribed respectively by Johnson and Worthington Wilmer (2019).
- One COI barcode of each BIN associated with a species name in BOLD, i.e., those present in the updated species checklist for Vietnam. BINs have been identified through the public data portal of BOLD (consultation 29/11/ 2018).

The aim of this second phylogenetic tree was to check concordance between species identification based on morphological criteria (present data) and DNA barcode (BOLD), to reveal misidentification or introgression in BINs associated with grouper species present in Vietnam and, finally, to highlight species that are polyphyletic due to the presence of several BINs. Species misidentification or introgression is responsible for more than one species name within a BIN. However, it is possible to determine specimens in a BIN that were misidentified or a hybrid, by considering:

- the number and origin (laboratory and geography) of specimens of different species within a BIN: the more barcodes were similarly identified by independent laboratories from various locations, the more the species identification may be considered reliable,
- the phenotype of the specimen when a picture is available in BOLD,
- the phylogeny, by considering the phylogenetic position of reference species sequence and BINs.

Following this rationale, we reassigned BINs to a probable species.

A phylogeographic analysis was used to further investigate processes responsible for intraspecific diversity in the groupers. A special attention was paid to polyphyletic species, those that present several BINs that form a clade in the phylogenetic tree, in order to determine the evolutionary significance of these BINs. All public COI barcodes belonging to BINs associated with a species name were uploaded and added to "Vietnamese" grouper data (Table S4). Haplotypic networks were estimated using the TCS method of Clement et al. (2002) implemented in PopART (French et al. 2014; http://popart. otago.ac.nz), and the geography and BIN were mapped



◄ Fig. 2 Phylogenetic relationships among cytochrome oxydase I barcodes and BINs recovered among Vietnamese groupers. The pattern of divergence between species is represented by a neighbor-joining (NJ) tree based on K2P distances. The relevance of tree nodes are highlighted by bootstrap values (only values > 70% are indicated). The tree is rooted using *Diploprion bifasciatum* 

onto these TCS networks to determine a phylogeographic structure.

### Results

# Known diversity of groupers in Vietnam and their associated BINs

By compiling knowledge from two major grouper species checklists for the West Pacific or worldwide (Heemstra and Randall 1993; Craig et al. 2011) and reviewing all recent grouper species description or redescription (Liu et al. 2013; Tucker et al. 2016; Frable et al. 2019), 40 species were recorded for Vietnam (Table 1). These species belong to either six or four different genera, according Eschmeyer's Catalog of Fishes (Fricke et al. 2019) or Ma and Craig (2018), respectively (Table 1). With the exception of *Epinephelus craigi* described recently (Frable et al. 2019) and still not referenced in BOLD, all these species names were associated with one or several BINs (up to 4, Table 1). Only 1/3 of the species were associated with only a single BIN.

#### Species diversity and BINs recovered in Vietnam

A total of 158 DNA barcodes (with a core segment of a 603 bp) were uploaded to the DS-SERV project (dx.doi.org/ 10.5883/DS-SERV). These DNA barcodes correspond to 31 species matching 32 BINs (Fig. 2), only one species *Epinephelus areolatus* presented haplotypes that belonged to two BINs (Tables 1 and 2). With the exception of *Epinephelus areolatus* that presented 3.5% of divergence between the two BINs, the intraspecies diversity (similar to the intra BIN) was extremely low and ranged from 0% for several species sampled more than one time to 0.4% for *Cephalopholis boenak* (Table S2). Interspecies divergence ranged from 3.8% between *Epinephelus malabaricus* and *Epinephelus coioides* to 24.3% between *Epinephelus malabaricus* and *Diploprion bifasciatum* (Table S2)

Among the species identified, nine were new records for Vietnam: *Epinephelus bruneus*, *Epinephelus fuscoguttatus*, *Epinephelus hexagonatus*, *Epinephelus polyphekadion*, *Plectropomus maculatus*, *Plectropomus oligacanthus*, *Triso dermopterus*, *Variola albimarginata*, and *Variola louti*. The grouper checklist for Vietnam therefore now includes 49 species. These species were associated with 76 BINs. The ABGD algorithm identified 77 OTUs that broadly matched the RESL delineation (i.e., BINs) (Table S3). Actually, if we exclude three cases, there are no differences between OTUs delineated by the ABGD or RESL algorithms. The OTU6 included sequences belonging to two BINs BOLD:AAD4737 and BOLD:AAB5431 while one sequence of this last BIN is highlighted as a distinct OTU: OTU7 (Table S3). All sequences belonging to OTU6, OTU7, BOLD:AAD4737, and BOLD:AAB5431 come from specimens identified as Cephalopholis sonnerati. The last discrepancy concerned a unique sequence that belonged to the BIN BOLD:AAC5719 but was assigned by ABGD to a specific OTU (OTU72). Because the RESL algorithm appeared more conservative by comparison to ABGD, which was able to create a specific OTU for only one sequence, we hereafter only considered BINs for further analyses.

Multi-BIN species may correspond to polyphyletic species, but specimens that were misidentified or hybridized would also artificially increase the number of BIN per species. For these latter cases, the careful reexamination revised the species associated with them (Table 2). For 23 BINs, it was possible to sort hybrids and misidentified specimens, and thus limit these BINs to one species name (see Table S5 for rationale). For one BIN (BOLD:ADE3544) originally assigned to *E. chlorostigma*, it was not possible to check the species identification because no picture was provided for any BIN members while its phylogenetic position did not show any close relationships with either *E. chlorostigma* lineages or the species barcode reference.

In the phylogenetic tree (Fig. 3) we also noticed some issues for a number of barcodes published by Ma et al. (2016). The COI sequence corresponding to E. chlorostigma in Ma et al. (2016) is grouped with sequences identified as E. areolatus in BOLD and the present study (Fig. 3 (5)). The Ma et al. (2016) COI sequence of E. bruneus matched the sequence of the resurrected species E. moara published by Liu et al. (2013) but not the E. bruneus sampled in Vietnam that belonged to the BIN BOLD:ACS0266 that was also associated with E. bruneus. (Fig. 3 (3)). Their sequence labelled as E. sexfasciatus was not different to their sequence labelled as E. fasciatomaculosus while, in this study, we identified a specimen as E. sexfasciatus that presented a different sequence that belonged to the BIN BOLD:ACM3827 that only comprised E. sexfasciatus members (Fig. 3 (6)). Their sequence of E. amblycephalus corresponded to the DNA barcode later described as E. craigi (Frable et al. 2019) (Fig. 3 (6)). Their COI sequence corresponding to E. irroratus was phylogenetically extremely close to E. fasciatus that comprised 4 BINs. Similarly, no genetic divergence was observed between their sequence of Cephalopholis nigripinnis and C. urodeta (Fig. 3 (7)), or between C. cyanostigma and C. microprion (Fig. 3 (8)). Considering this revision, several conditions can be highlighted:

Table 2	List of BINs present in the	Table 1 and	associated species	s name after revision	(see rationale in	Table S5)	or in BOLD
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BIN	According to this review	In BOLD			
	Species name	Species 1	Species 2	Species 3	Species 4
BOLD:AAB1332	Epinephelus fasciatus	Epinephelus fasciatus			
BOLD:AAB1333	Epinephelus fasciatus	Epinephelus fasciatus	Epinephelus irroratus		
BOLD:AAB1334	Epinephelus fasciatus	Epinephelus fasciatus			
BOLD:AAB1335	Epinephelus fasciatus	Epinephelus fasciatus			
BOLD:AAB3684	Cephalopholis boenak	Cephalopholis boenak			
BOLD:ACT7478	Cephalopholis boenak	Cephalopholis boenak	Cephalopholis igarashiensis		
BOLD:ACT7479	Cephalopholis boenak	Cephalopholis boenak	igurusmensus		
BOLD:AAB5431	Cephalopholis sonnerati	Cephalopholis sonnerati			
BOLD:AAD4737	Cephalopholis sonnerati	Cephalopholis sonnerati	Cephalopholis argus		
BOLD:ACC0647	Cephalopholis sonnerati	Cephalopholis sonnerati			
BOLD:AAA9821	Epinephelus areolatus	Epinephelus areolatus			
BOLD:AAA9822	Epinephelus areolatus	Epinephelus areolatus			
BOLD:ACC0344	Epinephelus areolatus	Epinephelus areolatus			
BOLD:AAC6284	Epinephelus morrhua	Epinephelus morrhua			
BOLD:ACE6821	Epinephelus morrhua	Epinephelus morrhua			
BOLD:ADC2027	Epinephelus morrhua	Epinephelus morrhua			
BOLD:AAD9665	Aethaloperca rogaa	Aethaloperca rogaa			
BOLD:ACC0322	Aethaloperca rogaa	Aethaloperca rogaa			
BOLD:AAC8327	Anyperodon leucogrammicus	Anyperodon			
BOLD:AAC8328	Anyperodon leucogrammicus	Anyperodon leucogrammicus			
BOLD:AAC3591	Epinephelus chlorostigma	Epinephelus spilotoceps			
BOLD:ABA7404	Epinephelus chlorostigma	Epinephelus chlorostigma			
BOLD:AAF1066	Epinephelus epistictus	Epinephelus epistictus			
BOLD:ADB3169	Epinephelus epistictus	Epinephelus epistictus			
BOLD:AAD8911	Epinephelus latifasciatus	Epinephelus latifasciatus			
BOLD:ADM3791	Epinephelus latifasciatus	Epinephelus latifasciatus			
BOLD:AAB8386	Epinephelus merra	Epinephelus merra	Epinephelus macrospilos		
BOLD:AAB8387	Epinephelus merra	Epinephelus merra	Epinephelus faveatus		
BOLD:AAD8873	Epinephelus polyphekadion	Epinephelus polyphekadion	Epinephelus fuscoguttatus		
BOLD:ABX5597	Epinephelus polyphekadion	Epinephelus polyphekadion			
BOLD:AAF0433	Epinephelus rivulatus	Epinephelus rivulatus			
BOLD:ACZ9919	Epinephelus rivulatus	Epinephelus rivulatus			
BOLD:ACK7859	Plectropomus maculatus	Plectropomus maculatus			
BOLD:ADL8508	Plectropomus maculatus	Plectropomus maculatus			
BOLD:AAC5719	Variola louti	Variola louti			
BOLD:AAC5720	Variola louti	Variola louti			
BOLD:AAB1346	Epinephelus amblycephalus	Epinephelus amblycephalus			
BOLD:AAJ0588	Epinephelus moara	Epinephelus bruneus			

#### Table 2 (continued)

BIN	According to this review	In BOLD			
	Species name	Species 1	Species 2	Species 3	Species 4
BOLD:ACS0266	Epinephelus bruneus	Epinephelus bruneus			
BOLD:AAC4474	Cephalopholis argus	Cephalopholis argus			
BOLD:AAB5432	Cephalopholis aurantia	Cephalopholis aurantia	Cephalopholis sonnerati	Cephalopholis spiloparaea	
BOLD:AAC0217	Cephalopholis cyanostigma	Cephalopholis cyanostigma	Cephalopholis microprion	Cephalopholis miniata	Cephalopholis argus
BOLD:AAF0621	Cephalopholis formosa	Cephalopholis formosa			
BOLD:AAD9032	Cephalopholis leopardus	Cephalopholis leopardus			
BOLD:AAF0629	Cephalopholis microprion	Cephalopholis microprion			
BOLD:AAC0216	Cephalopholis miniata	Cephalopholis miniata			
BOLD:AAC1876	Cromileptes altivelis	Cromileptes altivelis			
BOLD:AAF1053	Epinephelus akaara	Epinephelus akaara	Epinephelus fasciatomaculosus		
BOLD:AAF0829	Epinephelus awoara	Epinephelus awoara	Epinephelus fasciatomaculosus		
BOLD:AAC5362	Epinephelus bleekeri	Epinephelus bleekeri	Epinephelus quoyanus	Epinephelus akaara	
BOLD:ADL0994	Epinephelus coeruleopunctatus	Epinephelus coeruleopunctatus	Epinephelus morrhua		
BOLD:AAB8391	Epinephelus coioides	Epinephelus coioides	Epinephelus tauvina	Epinephelus awoara	Epinephelus malabaricus
BOLD:AAD8682	Epinephelus corallicola	Epinephelus corallicola			
BOLD:AAB1345	Epinephelus craigi	Epinephelus amblycephalus	Epinephelus stictus		
BOLD:ACC0501	Epinephelus erythrurus	Epinephelus erythrurus	Epinephelus coeruleopunctatus		
BOLD:AAC5392	Epinephelus fasciatomaculosus	Epinephelus sexfasciatus	Epinephelus fasciatomaculosus		
BOLD:AAD8872	Epinephelus fuscoguttatus	Epinephelus fuscoguttatus	Epinephelus lanceolatus		
BOLD:AAD3855	Epinephelus heniochus	Epinephelus heniochus	Epinephelus poecilonotus		
BOLD:AAD3458	Epinephelus hexagonatus	Epinephelus hexagonatus			
BOLD:AAD8914	Epinephelus lanceolatus	Epinephelus lanceolatus			
BOLD:AAE1882	Epinephelus macrospilos	Epinephelus macrospilos			
BOLD:AAB8389	Epinephelus malabaricus	Epinephelus malabaricus	Epinephelus coioides		
BOLD:AAC2802	Epinephelus ongus	Epinephelus ongus	Epinephelus coioides		
BOLD:AAD3854	Epinephelus poecilonotus	Epinephelus poecilonotus			
BOLD:AAC5363	Epinephelus polylepis	Epinephelus chlorostigma	Epinephelus polylepis		
BOLD:AAC2809	Epinephelus quoyanus	Epinephelus quoyanus			
BOLD:ACM3827	Epinephelus sexfasciatus	Epinephelus sexfasciatus			
BOLD:ADA9689	Epinephelus stictus	Epinephelus amblycephalus			
BOLD:ADE3544	Epinephelus sp. 1	Epinephelus chlorostigma			
BOLD:AAJ0586	Epinephelus trimaculatus	Epinephelus trimaculatus	Epinephelus fario		
BOLD:AAC9348	Plectropomus leopardus	Plectropomus leopardus	Plectropomus areolatus	Plectropomus maculatus	Plectropomus pessuliferus

Table 2 (continued)

BIN	According to this review	In BOLD			
	Species name	Species 1	Species 2	Species 3	Species 4
BOLD:AAL4434	Plectropomus oligacanthus	Plectropomus oligacanthus			
BOLD:AAD9769	Triso dermopterus	Triso dermopterus			
BOLD:AAC9675	Variola albimarginata	Variola albimarginata			
BOLD:AAC4202	Cephalopholis urodeta/C. nigripinnis	Cephalopholis urodeta	Cephalopholis nigripinnis		
BOLD:AAD1767	Epinephelus cyanopodus/E. flavocaeruleus	Epinephelus flavocaeruleus	Epinephelus cyanopodus		

- One BIN–several species: Only two BINs corresponded to this situation where there was no apparent misidentification: BOLD:AAD1767 composed of members either identified as *E. cyanopodus* or *E. flavocaeruleus*, and BOLD:AAC4202 composed of members that can either corresponded to *Cephalopholis urodeta* or *C. nigripinnis*.
- One BIN-one species: This concerned 37 species listed in Table 2. Chromileptes altivelis, Cephalopholis argus, C. aurantia, C. cyanostigma, C. formosa, C. leopardus, C. microprion, C. miniata, Epinephelus akaraa, E. amblycephalus, E. awoara, E. bleekeri, E. bruneus, E. coioides, E. corallicola, E. craigii, E. heniochus, E. coeruleopunctatus, E. erythrurus, E. hexagagonatus, E. fasciatomaculosus, E. fuscoguttatus, E. lanceolatus, E. malabaricus, E. macrospilos, E. moara, E. ongus, E. poecilonotus, E. polylepis, E. quoyanus, E. sexfasciatus, E. stictus, E. trimaculatus, Plectropomus leopardus, P. oligacanthus, Triso dermopterus, Variola albimarginata.
- One species-several BINs: This concerned 15 species: Aethaloperca rogaa, Anyperodon leucogrammicus, Cephalopholis boenak, C. sonnerati, Epinephelus areolatus, E. chlorostigma, E. epistictus, E. fasciatus, E. latifasciatus, E. merra, E. morruha, E. polyphekadion, E. rivulatus, Plectropomus maculatus, Variola louti.

#### Phylogeographic structure of polyphyletic species

The TCS networks built for species that remained polyphyletic (several BINs for a unique species name) after BIN/species reassignment showed various phylogeographic patterns.

 A geographic overlap of BINs/clades associated with a species name. This was only observed for *Epinephelus morrhua*. However, it is noticeable that, among the three clades corresponding to 3 BINs, two are limited to one geographic location (East Australia for BOLD:ADC2027 and the South China Sea for BOLD:AAC6284) while the third was widespread and present in the whole Indo-Pacific Ocean (Fig. 41).

- A strict allopatric/parapatric distribution for at least one BIN associated with a species. This was observed for: Aethaloperca rogaa, Epinephelus areolatus and Plectropomus maculatus with the BINs BOLD: ACC0322, BOLD: ACC0344, and BOLD: ADL8508 respectively, restricted to the Andaman Islands (Fig. 4a, f, o); Cephalopholis sonnerati with BINs BOLD:AAB5431 and BOLD:ACC0647 restricted to SW Indian Ocean and the Andaman Islands, respectively (Fig. 4d); Epinephelus epistictus with the BIN BOLD: ADB3169 and the BIN BOLD: AAF1066 restricted to the Indian Ocean and the South China Sea. respectively (Fig. 4h); E. fasciatus and E. rivulatus with their BINs (BOLD:AAB1334 and BOLD:ACZ9919, respectively) restricted to the West Indian Ocean and the Red Sea (Fig. 4i, k); E. polyphekadion for which the distribution of the two BINs showed a clear geographic barrier located in the central part of the Indian Ocean (Fig. 4m), and Variola louti with the BIN BOLD:AAC5720 restricted to the Central Pacific Ocean: (Fig. 4p).
- Partial parapatry corresponded to species that presented several BINs at one location that, elsewhere, were in parapatry. This was the case for: *Anyperodon leucogrammicus* with two BINs present in Taiwan whereas BOLD:AAC8328 presented a distribution usually

Fig. 3 Phylogenetic relationships among all BINs recovered in Vietnam (white dot), BINs associated with species inventoried in Vietnam (black square), and some species COI sequence references from Ma et al. (2016, gray triangle), Frable et al. (2019, red triangle), Liu et al. (2013, red triangle), and Johnson and Worthington Wilmer (2019, red triangle). The pattern of divergence among species is represented by a neighborjoining (NJ) tree based on K2P distances. The relevance of tree nodes are highlighted by bootstrap values (only value > 70% are indicated). \*Flags polyphyletic species (more than one BIN but phylogenetically close); °flags paraphyletic species





Fig. 3 (continued)

restricted to the Indian Ocean (Fig. 4b); *Cephalopholis boenak* with two BINs present in Western Australia whereas BOLD:AAB3684 was restricted to Australian shores (Fig. 4c); *Epinephelus areolatus* with two BINs present in Vietnam while BIN BOLD:AAA9822 was mainly observed in the Indian Ocean (Fig. 4f), and *E. merra* (Fig. 4k) with two BINs observed in Indonesia while their distribution range were allopatric outside this location and presented a clear separation between the Indian and the Pacific Oceans.

Lastly, for some multi BIN species a phylogeographic structure was also observed within a BIN. This was the case for: *Plectropomus maculatus* (Fig. 4o) with the BIN BOLD:ACK7859 that showed two sets of haplotypes either distributed along the Australian shore or in the South East Asia, and *Variola louti* (Fig. 4p) with the BIN BOLD:AAC5719 that presented three subclades restricted to the Red Sea, the Andaman Islands and the Indian Ocean.

## Discussion

#### Grouper diversity in Vietnam

To date, the species checklist for Vietnam consisted of 40 groupers and, while this study identified 30 species, up to nine are new records. As a result, up to 50% of known Vietnamese grouper diversity was not recovered in this study, despite sampling on a large spatial and temporal scale. Several species listed are, however, uncommon and can present a patchy geographical distribution or occurrence in Vietnam that is limited to some provinces. This is the case for Anyperodon leucogrammicus, Cephalopholis leopardus, C. microprion, Epinephelus akaara, E. craigi, E. cyanopodus, etc. Furthermore, the absence of some species may be a consequence of the intense fishing pressure on many groupers. In 1996, 43.5% of groupers were considered as threatened and two, Cromileptes altivelis and Epinephelus akaara as endangered, in the IUCN red list (Moriss et al. 2000). These two endangered species were not observed in the present study despite historical records in Vietnam. Consequently, our results suggest that more grouper species may be locally endangered but the absence, in Vietnam, of temporal series of fish biomass landing hampers our ability to monitor population dynamics and evaluate species status (Stocks et al. 2019). Life history traits of groupers, such as large body size, slow growth, long life-span, late reproduction, plus behaviors such as territoriality and spawning aggregations, make their survival uncertain and the chances of local population collapse high (Moriss et al. 2000; Sadovy de Mitcheson et al. 2012). Beyond biological reasons that render groupers vulnerable to over-exploitation, pollution, and habitat destruction are also a major threat. Urbanization of coastal areas in Vietnam has degraded and even destroyed many aquatic habitats. Mangrove forests, known to be important nursery areas for marine fishes including groupers, declined dramatically from 4000 km<sup>2</sup> in 1950 to 1570 km<sup>2</sup> in 1994 (Son and Thuoc 2003), due to conversion for agriculture or pond aquaculture, and unsustainable use for timber (Hong 1993). Similarly, coral reefs that concentrate a high proportion of grouper species have decreased to 15–20% of their original surface due to pollution, siltation, and blast fishing (Lutaenko 2011). For all these reasons, it is likely that many groupers species in Vietnam have become rare, so explaining their absence in the present sampling.

Curiously, up to nine species are new records for Vietnam. This contrasts with the species rarefaction described above. These new records may also, however, be explained by exploitation. Vietnamese fisheries are primarily small-scale and inshore, but, due to the decline of many high-value inshore resources, many fishermen now focus on deeper waters (Thuoc 2001). As a consequence, some species that are not present inshore and thus were not targeted historically by the small scale fisheries, have become more visible on fish markets (Sadovy de Mitcheson et al. 2012). Among the nine new records up to six (E. bruneus, P. maculatus, P. oligacanthus, T. dermoptrus, V. albimarginatus and V. louti) present a vertical distribution extending into deep water below above 100 m. None are, however, strictly restricted to deep waters, which limits this interpretation such that it cannot by itself explain why these species were not previously recorded. Nevertheless, in recent Vietnamese investigations, six of these species were observed (Le et al. 2011; Tran et al. 2015; Vo et al. 2016; Vo 2018) while E. fuscoguttatus is known to be reared in aquaculture at Quang Ninh based on larvae fished in the wild (Le 2004). Last, there are still two species: E. polyphekadion and V. albimarginata that remain new records for Vietnam whatever species checklist is considered and regardless of any aquaculture practices in Vietnam. The distribution range of some groupers are known to be very patchy, such as E. polyphekadion, such that they may be limited to small areas in Vietnam and have therefore not been recorded before. Concerning V. albimarginata, it is more difficult to explain why it was not recorded before because the species seems abundant in Vietnam (pers obs) and is difficult to misidentify with other groupers.

# BIN diversity in Indo-Pacific groupers present in Vietnam

Specimens mislabelled in BOLD may have two origins: taxonomic confusion or hybridization. Grouper taxonomy is traditionally based on color patterns, morphological distinctiveness, and osteological characters (Heemstra and Randall 1993) but, for some species, these criteria overlap from a species to another leading to confusion (Alcantara and Yambot 2016; Qu et al. 2018). Furthermore, recent taxonomical investigations describing new species confound with other species in previous DNA barcoding studies. This is the case for *E. craigi* recently described by Frabe et al. (2018) and previously confounded or still misidentified with *E. amblycephalus* and *E. stictus*. Similarly, despite the redescription of *E. moara* by Liu et al. (2013), this species is still ignored in BOLD and its BIN:BOLD:AAJ0588 is associated with *Epinephelus bruneus*.

In addition, hybridization within groupers can produce specimens with atypical body coloration, influencing our ability to recognize overall species. Several studies already identified grouper hybrids in the wild (Bostrom et al. 2002; Frisch and van Herwerden 2006; Randall and Justine 2008; Payet et al. 2016; Qu et al. 2018). Genetic characterization of a hybrid requires the comparison of its phylogenetic position estimated by a mitochondrial marker such as the COI (maternally inherited) against a nuclear marker (bi-parentally inherited), as proposed by Payet et al. (2016) or Qu et al. (2018). However, hybridization does not necessarily lead to conflicting phylogenies since an absence of interspecies differentiation by genetic markers can also be interpreted as a consequence of hybridization and introgression. This was already pointed out for two couples of species that share an unique BIN:BOLD:AAC4202 for Cephalopholis urodeta and C. nigripinnis and BOLD:AAD1767 for E. cyanopodus and E. flavocaeruleus. Despite clear differences of color pattern, these species share a unique BIN and are indistinguishable with nuclear markers (Payet et al. 2016; Qu et al. 2018). According to Payet et al. (2016), such absence of interspecies differentiation maybe due either to hybridization and introgression, taxonomical characters that are phylogenetically irrelevant, or incomplete lineage sorting. While Qu et al. (2018) suggested that E. cyanopodus and E. flavocaeruleus may be considered as a unique species, Payet et al. (2016) considered Cephalopholis urodeta and C. nigripinnis as real species that able to interbreed because these authors depicted various intermediate phenotypes between the two species where they occur in sympatry.

The question that arises for species that present two to four specific BINs is of their evolutionary significance. Among 15 species presenting this situation, only three, *E. chlorostigma*, *E. fasciatus*, and *E. epistictus* were associated with BINs that does not form monophyletic clade. Identification of *E. chlorostigma* seems confused in light of the phylogenetic tree based on sequences of various origin (Fig. 3). While the COI barcode of the specimen of Ma et al. (2013) corresponded to *E. areolatus* (for this study and for BOLD), among the BINs associated with *E. chlorostigma* in BOLD, none corresponded to the Vietnamese specimens that have been

Fig. 4 Haplotypic networks recovered in multi BIN grouper species for which BINs form a clade in Fig. 3. Dot color in the network represents the geographic origin of the haplotypes, while its size represents the abundance in BOLD. **a** *Aethaloperca rogaa*; **b** *Anyperodon leucogrammicus*; **c** *Cephalopholis boenak*; **d** *Cephalopholis sonnerati*; **e** *Epinephelus amblycephalus*, *Epinephelus stictus*, *Epinephelus craigii*; **f** *Epinephelus areolatus*; **g** *Epinephelus bruneus*, *Epinephelus moara*; **h** *Epinephelus epistictus*; **i** *Epinephelus fasciatus*; **j** *Epinephelus latisfasciatus*; **k** *Epinephelus merra*; **l** *Epinephelus morrhua*; **m** *Epinephelus polyphekadion*; **n** *Epinephelus rivulatus*; **o** *Plectropomus maculatus*; **p** *Variola louti* 

unambiguously identified as E. chlorostigma. By contrast, the BIN of the Vietnamese E. chlorostigma (BOLD:AAC3591) was associated in BOLD to seven specimens identified by the same institute as E. spilotoceps and collected in the South China Sea close to the Vietnamese sampling location. These specimens have been misidentified considering that (1) available pictures do not present the four blackish brown saddle-blotches at the base of the dorsal fin that are characteristic of E. spilotoceps (FAO), (2) the specimen identified by Ma et al. (2016) as E. spilotoceps belongs to a different phylogenetic lineage (Fig. 3), and (3) there is a BIN (BOLD:AAC3590) consisting of 17 members (including the specimen of Ma et al. 2016) that was unambiguously identified by BOLD as E. spilotoceps. Finally, there are two unrelated BINs (BOLD:AAC3591 and BOLD:ABA7404) associated to the name E. chlorostigma which suggest that there is a cryptic species in this taxon. Considering the type locality of E. chlorostigma (Seychelles), and BIN's geographic distributions which are based on only a few specimens, more work is needed to determine the BIN that corresponds to E. chlorostigma.

Concerning BINs recovered in specimens identified as E. fasciatus, they form a monophyletic clade that is paraphyletic with E. retouti and E. irroratus (Fig. 3). E. irroratus is endemic to the Marguesas Island and shares the BIN BOLD: AAB1333 with E. fasciatus, which is mainly observed in the SW and Central Pacific (Fig. 4i). By contrast, the Indo-Pacific E. retouti was associated with a BIN (BOLD:AAX8350) that did not correspond to any BINs identified in E. fasciatus but was closely phylogenetically related to the E. fasciatus BIN (BOLD: AAB1334). This last BIN (BOLD:AAB1334) diverged deeply from all other E. fasciatus BINs (Fig. 3) and presented a strong allopatric distribution limited to the Red Sea and the West Indian Ocean (Fig. 4i). All these results stress the presence of a species complex in E. fasciatus and, because the type locality of this species is the Red Sea, this suggests that the valid E. fasciatus should be limited to the Red Sea and the West Indian Ocean (= the BIN BOLD:AAB1334). All other BINs associated with the name E. fasciatus may correspond to another species for which phylogenetic relationships with E. irroratus is not clear. More taxonomic and phylogenetic work is necessary to solve





Fig. 4 (continued)



Fig. 4 (continued)

this question, given that genetic data concerning *E. retouti* is limited to only 7 COI barcodes.

Last, the two BINs (BOLD:AAF1066 and BOLD:ADB3169) associated with specimens identified as *E. epistictus* form a well-supported clade that also included the recent described species *E. fuscomarginatus* and its sister species *E. magniscuttis* (Fig. 3). This result confirmed findings of Johnson and Worthington Wilmer (2019) who demonstrated that *E. epistictus* is a species complex composed of at least 4 species. Because the type locality is Nagasaki, Japan, it is probable that the BIN BOLD:AAF1066 observed in specimens collected in the NW Pacific (Taiwan and China) corresponds to *E. epistictus*, but see Johnson and Worthington Wilmer (2019) for a more detailed and complete analysis.

For all other polyphyletic species or multi-BIN species, the present study demonstrated that their BINs form well supported clades in the majority of cases (Fig. 3). This result may be interpreted in three ways: BINs of a species correspond to (1) undescribed or cryptic sister species; (2) intraspecific biogeographic lineages, or (3) artifacts due to bad nucleotide scoring in the sequencing file (ab1 file) generated by the sequencing apparatus. This last situation cannot be excluded when a BIN is observed at only one location and thus consists of few members analyzed by only one institute. Up to seven BINs correspond to this situation and, curiously, 4 of them where described only in the Andaman Islands, such as BOLD:ACC0322 (Aethaloperca rogaa), BOLD:ACC0647 (Cephalopholis sonnerati), BOLD:ACC0344 (Epinephelus areolatus), and BOLD:ADL8508 (Plectropomus maculatus). Given that it is rare in groupers that a BIN is restricted to only one location and unless there is a hotspot of diversity in the Andaman Islands, such results remain doubtful and call for further investigation.

Now, when polyphyletic species are composed of BINs with members from different geographic origin that do not match any geographic pattern, it is probable that these BINs flag sister species. This is the case for BINs highly associated with the name *E. amblycephalus* and that correspond to three sister species: *E. amblycephalus*, *E. stictus*, and the recently described species *E. craigi* (Figs. 3 and 4e). This is also true for the two BINs associated with the species name *E. bruneus*, since one of these corresponds to the COI sequence of the species *E. moara* that is still not recognized as valid in Eschmeyer's Catalog of Fishes (Fricke et al. 2019), despite arguments provided by Liu et al. (2013).

Lastly, in all remaining multi-BIN species, a phylogeographic structure has been revealed, with BIN geographic distributions that match biogeographic units. This suggests that the presence of more than one BIN in a species mirrors a complex evolutionary history. Several studies have already stressed phylogeographic structures in groupers such as *Cephalopholis argus*, *Epinephelus areolatus*, *E. polyphekadion*, *E. merra*, *Plectropomus areolatus*, and P. leopardus (Gaither et al. 2011; DiBattista et al. 2013; Borsa et al. 2016; Ma et al. 2018). If we exclude Plectropomus areolatus that was not reviewed in this study due its absence in Vietnam, three of five of these species are multi BIN species. Most of these studies interpret the genetic structure of these species as a result of paleogeographic changes that directly impact the phylogeographic structure of groupers (Gaither et al. 2011; DiBattista et al. 2013; Borsa et al. 2016). A number of important biogeographic barriers in the Indo-Pacific have shaped the present-day genetic structure, such as the mid Indian Barrier or the Indo/Pacific Ocean divide. But Ma et al. (2018) also noticed that reproductive behavior is an important factor that can modulate the strength of a biogeographic barrier by either limiting or increasing population connectivity. According to Ma et al. (2018) discrete reproduction in small groups of Plectropomus leopardus individuals may have favored a stronger genetic structure of Pleistocene origin in the West Indo-Pacific than Plectropomus areolatus and Ephinephelus polyphekadion, which breed in large numbers on a few spawning grounds. Secondary contact and genetic connectivity may be stronger for species presenting such behavior, explaining why some genetic breaks in the West Indian Pacific are not observed.

To conclude, the association of several BINs to a species cannot be interpreted straightforwardly. If, in a number of cases, misidentification and hybridization artificially increase the apparent genetic diversity of a species, the presence of several BINs can also flag putative cryptic species or a strong phylogeographic structure. More barcoding studies focusing on fish families are needed to clarify the association between a species and BINs and, thereby, improve the validity and reliability of the species identification engine provided by BOLD. In any case, beyond species identification, once clarified for multi-BIN species, it would certainly be possible to use the BINs not only to identify a specimen but also to track its biogeographic origin.

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#### **Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no conflict of interest.

**Ethical approval** All applicable international, national, and/or institutional guidelines for the care and use of animals were followed by the authors.

**Sampling and field studies** All necessary permits for sampling and observational field studies have been obtained by the authors from the competent authorities and are mentioned in the acknowledgements, if applicable.

**Data availability** All data generated or analyzed during this study are included in this published article [and its supplementary information files]. DNA barcodes generated during the current study are available in the project file SERV: Epinephelidae of Vietnam on BOLD systems website (http://boldsystems.org).

Author contribution JDD conceived and designed research. JDD, MHP, TTVY, DHH, and QVV conducted experiments. JDD analyzed data and wrote the manuscript. All authors read and approved the manuscript.

## References

- Alcantara SG, Yambot AV (2016) DNA barcoding of commercially important grouper species (Perciformes, Serranidae) in the Philippines. Mitochondrial DNA Part A 27:3837–3845. https://doi.org/10.3109/ 19401736.2014.958672
- Barcaccia G, Lucchin M, Cassandro M (2015) DNA barcoding as a molecular tool to track down mislabeling and food piracy. Diversity 8:2. https://doi.org/10.3390/d8010002
- Basheer VS, Vineesh N, Bineesh KK, Kumar RG, Mohitha C, Venu S, Kathirvelpandian A, Gopalakrishnan A, Jena JK (2017) Mitochondrial signatures for identification of grouper species from Indian waters. Mitochondrial DNA Part A 28:451–457. https://doi. org/10.3109/19401736.2015.1137899
- Bleeker P (1849) Bijdrage tot de kennis der Percoïden van den Malaijo-Molukschen Archipel, met beschrijving van 22 nieuwe soorten. Verhandelingen van het Bataviaasch Genootschap van Kunsten en Wetenschappen. v. 22 [13]: 1-64.
- Bleeker P (1852) Nieuwe bijdrage tot de kennis der ichthyologische fauna van Amboina. Natuurkundig Tijdschrift voor Nederlandsch Indië 3: 545–568
- Bleeker P (1855) Derde bijdrage tot de kennis der ichthyologische fauna van Batjan. Natuurkundig Tijdschrift voor Nederlandsch Indië 9: 491–504
- Bleeker P (1857) Achtste bijdrage tot de kennis der vischfauna van Amboina. Acta Societatis Regiae Scientiarum Indo-Neêrlandicae 102 pp.
- Bloch ME (1790) Naturgeschichte der ausländischen Fische. Siebenter Theil J Morino & Co, Berlin, 128 pp
- Bloch ME (1793) Naturgeschichte der ausländischen Fische. Siebenter Theil J Morino & Co, Berlin, 144 pp
- Bloch ME, Schneider JG (1801) Systema Ichthyologiae Iconibus cx Ilustratum. Post obitum auctoris opus inchoatum absolvit, correxit, interpolavit Jo. Gottlob Schneider, Saxo. Berolini. Sumtibus Auctoris Impressum et Bibliopolio Sanderiano Commissum. 584 pp.
- Borsa P, Durand J-D, Chen W-J, Hubert N, Muths D, Mou-Tham G, Kulbicki M (2016) Comparative phylogeography of the western Indian Ocean reef fauna. Acta Oecol 72:72–86. https://doi.org/10. 1016/j.actao.2015.10.009
- Bostrom MA, Collette BB, Luckhurst BE, Reece KS, Graces JE (2002) Hybridization between two serranids, the coney (*Cephalopholis*

*fulva*) and the creole-fish (*Paranthias furcifer*), at Bermuda. Fish Bull 100:651–661

- Brito MA, Schneider H, Sampaio I, Santos S (2015) DNA barcoding reveals high substitution rate and mislabeling in croaker fillets (Sciaenidae) marketed in Brazil: the case of "pescada branca" (Cynoscion leiarchus and Plagioscion squamosissimus). Food Res Int 70:40e46. https://doi.org/10.1016/j.foodres.2015.01.031
- Chu C, Rizman-Idid M, Chong VC (2011) The relationship of groupers (subfamily: Epinephelinae) from Peninsular Malaysia based on COI. Conference: 8th International Scientific Symposium, Ocean Climate and Marine Ecosystems in the Western Pacific, Busan, Korea
- Clement M, Snell Q, Walker P, Posada D, Crandall K (2002) TCS: Estimating gene genealogies. Parallel and Distributed Processing Symposium, International Proceedings 2:184
- Craig MT, Hastings PA (2007) A molecular phylogeny of the groupers of the subfamily Epinephelinae (Serranidae) with a revised classification of the Epinephelini. Ichthyol Res 54:1–17. https://doi.org/10. 1007/s10228-006-0367-x
- Craig MT, Sadovy de Mitcheson YJ, Heemstra PC (2011) Groupers of the world. NISC, Grahamstown
- Cuvier G, Valenciennes A (1828) Histoire naturelle des poissons. Tome second. Livre troisième. Des poissons de la famille des perches, ou des percoïdes. FG Levrault, Paris. 490 pp. [Valenciennes author of pp. 238-249, 262-386, Cuvier the remainder.]
- Cuvier G, Valenciennes A (1830) Histoire naturelle des poissons. Tome Sixième. Livre sixième. Partie I. Des Sparoïdes; Partie II. Des Ménides. FG Levrault, Paris. 559 pp. [Valenciennes is author of pp. 1-425, 493-559; Cuvier 426-491.]
- Cuvier G, Valenciennes A (1833) Histoire naturelle des poissons. Tome neuvième. Suite du livre neuvième. Des Scombéroïdes. FG Levrault, Paris. 512 pp. [Cuvier authored pp. 1-198, 330-359, 372-427; Valenciennes the balance.]
- de Baissac JB (1953) Contribution à l'étude des poissons de l'Ile Maurice. V. Proc Roy Soc Mauritius 1: 185-240.
- Delrieu-Trottin E, Williams JT, Pitassy D, Driskell A, Hubert N, Viviani J, Cribb TH, Espiau B, Galzin R, Kulbicki M, de Loma TL, Meyer C, Mourier J, Mou-Tham G, Parravicini V, Plantard P, Sasal P, Siu G, Tolou N, Veuille M, Weigt L, Planes S (2019) A DNA barcode reference library of French Polynesian shore fishes. Sci Data 6:114
- DiBattista JD, Berumen ML, Gaither MR, Rocha LA, Eble JA, Choat JH, Craig MT, Skillings DJ, Bowen BW (2013) After continents divide: comparative phylogeography of reef fishes from the Red Sea and Indian Ocean. J Biogeogr 40:1170–1181. https://doi.org/10.1111/ jbi.12068
- Ding SX, Zhuang X, Guo F, Wang J, Su Y, Zhang Q, Li Q (2006) Molecular phylogenetic relationships of China seas groupers based on cytochrome b gene fragment sequences. Sci China Ser C 49:235– 242. https://doi.org/10.1007/s11427-006-0235-y
- Durand JD, Borsa P (2015) Mitochondrial phylogeny of grey mullets (Acanthopterygii: Mugilidae) suggests high proportion of cryptic. C R Biol 338:266–277
- Durand J-D, Hubert N, Shen K-N, Borsa P (2017) DNA barcoding grey mullets. Rev Fish Biol Fisheries 27:233–243. https://doi.org/10. 1007/s11160-016-9457-7
- Forsskål PS (1775) Descriptiones animalium avium, amphibiorum, piscium, insectorum, vermium; quae in itinere orientali observavit [...]. Post mortem auctoris edidit Carsten Niebuhr, Hauniae, 164 pp
- Fowler HW (1904) A collection of fishes from Sumatra. J Acad Nat Sci Phila. Second series. 12: 495-560.
- Frable BW, Tucker SJ, Walker HJ Jr (2019) A new species of grouper, *Epinephelus craigi* (Perciformes: Epinephelidae), from the South China Sea. Ichthyol Res 66:215–224. https://doi.org/10.1007/ s10228-018-0669-9
- French N, Yu S, Biggs P, Holland B, Fearnhead P, Binney B, Fox A, Grove-White D, Leigh JW, Miller W, Muellner P, Carter P (2014)

Evolution of *Campylobacter* species in New Zealand. In: Sheppard SK, Méric G (eds) *Campylobacter* ecology and evolution. Caister Academic Press, Norfolk, pp 221–240

- Fricke R, Eschmeyer WN, Van der Laan R (2019) Catalog of Fishes: genera, species, references. http://researcharchive.calacademy.org/ research/ichthyology/catalog/fishcatmain.asp. Accessed 29 January 2019
- Frisch AJ, van Herwerden L (2006) Field and experimental studies of hybridisation between coral trouts, *Plectropomus leopardus* and *Plectropomus maculatus* (Serranidae), on the Great Barrier Reef, Australia. J Fish Biol 68:1013–1025. https://doi.org/10.1111/j. 0022-1112.2006.00977.x
- Gaither MR, Bowen BW, Bordenave T-R, Rocha LA, Newman SJ, Gomez JA, van Herwerden L, Craig MT (2011) Phylogeography of the reef fish *Cephalopholis argus* (Epinephelidae) indicates Pleistocene isolation across the Indo-Pacific barrier with contemporary overlap in the Coral Triangle. BMC Evol Biol 11:189. https:// doi.org/10.1186/1471-2148-11-189
- Galdino Brandao JHS, Araújo Bitencourt J, Borges Santos F, Almeida Watanabe L, Schneider H, Sampaio I, Mello Affonso PRA (2016) DNA barcoding of coastal ichthyofauna from Bahia, northeastern Brazil, South Atlantic: high efficiency for systematics and identification of cryptic diversity. Biochem Syst Ecol 65:214–224. https:// doi.org/10.1016/j.bse.2016.02.012
- Guo M, Su Y, Zhang Z, Ding S, Wang J (2009) Differentiation of *Epinephelus moara* from *E. bruneus* by improved nest-tetraprimer-specific PCR. Prog Nat Sci 19:1221–1226. https://doi.org/ 10.1016/j.pnsc.2009.04.003
- Guo M, Wang S, Su Y, Zhou Y, Liu M, Wang J (2014) Molecular cytogenetic analyses of *Epinephelus bruneus* and *Epinephelus moara* (Perciformes, Epinephelidae). PeerJ 2:e412. https://doi.org/ 10.7717/peerj.412
- Ha TTT, Nguyen TH, Nguyen PNP, Guiguen Y (2018) Species identification using DNA barcoding on processed Panga catfish products in Viet Nam revealed important mislabeling. Turk J Fish Aquat Sc 18: 457–462. https://doi.org/10.4194/1303-2712-v18\_3\_11
- Habib KA, Neogi AK, Oh J, Lee Y-H, Kim C-G (2018) Chelonodontops bengalensis (Tetraodontiformes: Tetraodontidae): a new species of puffer fish from the Northern Bay of Bengal based on morphology and DNA barcode. Ocean Sci J 54:79–86. https://doi.org/10.1007/ s12601-018-0054-7
- Hamilton F (1822) An account of the fishes found in the river Ganges and its branches. Edinburgh & London, 405 pp
- Heemstra PC, Randall JE (1993) Groupers of the world. FAO Fisheries synopsis no. 125, vol 16. FAO, Rome
- Heemstra PC, Randall JE (1999) Serranidae groupers and sea basses (also, soapfishes, anthiines, etc.). In: Carpenter KE, Niem VH (eds) FAO species identification guide for fishery purposes. The living marine resources of the Western Central Pacific, vol 4. Bony fishes part 2 (Mugilidae to Carangidae). FAO, Rome, pp 2069–2790
- Hong PN (1993) Aspects of mangrove ecosystem and status of shrimp culture in Ngoc Hien District, Minh Hai Province pp 49
- Huang WC, Nguyen VQ, Liao TY (2018) First record of the snowflakepatched moray *Gymnothorax niphostigmus* Chen, Shao, & Chen, 1996 (Anguilliformes; Muraenidae) in Vietnam and its validity confirmed by DNA barcoding. J Appl Ichthyol 34:687–690. https://doi. org/10.1111/jai.13684
- Hubert N, Hanner R (2015) DNA barcoding, species delineation and taxonomy: a historical perspective. DNA Barcodes 3:44–58. https://doi.org/10.1515/dna-2015-0006
- Hubert N, Meyer CP, Bruggemann HJ, Guérin F, Komeno RJL, Espiau B, Causse R, Williams JT, Planes S (2012) Cryptic diversity in Indo-Pacific coral-reef fishes revealed by DNA-barcoding provides new support to the centre-of-overlap hypothesis. PLoS One 7(3):e28987. https://doi.org/10.1371/journal.pone.0028987

- Hubert N, Dettai A, Pruvost P, Cruaud C, Kulbicki M, Myers RF, Borsa P (2017) Geography and life history traits account for the accumulation of cryptic diversity among Indo-West Pacific coral reef fishes. Mar Ecol Prog Ser 583:179–193. https://doi.org/10.3354/ meps12316
- Jaafar TNAM, Taylor MI, Mohd Nor SA, De Bruyn M, Carvalho GR (2012) DNA barcoding reveals cryptic diversity within commercially exploited Indo-Malay carangidae (Teleosteii: Perciformes). PLoS ONE 7:e49623. https://doi.org/10.1371/journal.pone.0049623
- Johnson GD (1983) *Niphon spinosus*: a primitive epinepheline serranid, with comments on the monophyly and intrarelationships of the Serranidae. Copeia 1983:777–787
- Johnson JW, Worthington Wilmer J (2019) Epinephelus fuscomarginatus (Perciformes: Epinephelidae), a new species of grouper from off the Great Barrier Reef, Australia. Zootaxa 4674: 329–348. https://doi.org/10.11646/zootaxa.4674.3.2
- Kimura M (1980) A simple method for estimating evolutionary rate of base substitutions through comparative studies of nucleotide sequences. J Mol Evol 16:111–120. https://doi.org/10.1007/ BF01731581
- Kumar S, Tamura K, Nei M (2004) MEGA3: integrated software for molecular evolutionary genetics analysis and sequence alignment. Brief Bioinform 5:150–163. https://doi.org/10.1093/bib/5.2.150
- Kumar S, Stecher G, Tamura K (2016) MEGA7: molecular evolutionary genetics analysis version 7.0 for bigger datasets. Mol Biol Evol 33: 1870–1874. https://doi.org/10.1093/molbev/msw054
- Lacepède BGE (1802) Histoire naturelle des poissons. Tome Quatrième, Plassan Paris, 728 pp
- Le AT (2004) Grouper farming in Vietnam: status and technical problems. Vietnamese J Fish Sci Technol, Special issue, 174–179
- Le TTT, Vo QV, Nguyen VUP (2011) Check list of grouper (family Serranidae) in Vietnam waters. Proceedings of the National Conference on Marine Science and Technology 5<sup>th</sup>. Nat Sci Technol 4:145–153
- Liu M, Li J-L, Ding S-X, Liu Z-Q (2013) *Epinephelus moara*: a valid species of the family Epinephelidae (Pisces: Perciformes). J Fish Biol 82:1684–1699. https://doi.org/10.1111/jfb.12112
- Lutaenko KA (2011) Proceedings of the workshop coastal marine biodiversity and bioresources of Vietnam and adjacent areas to the South China Sea, Nha Trang, Vietnam. Dalnauka, Vladivostok-Nha Trang, p 123
- Ma KY, Craig MT (2018) An inconvenient monophyly: an update on the taxonomy of the groupers (Epinephelidae). Copeia 106:443–456. https://doi.org/10.1643/CI-18-055
- Ma KY, Craig MT, Choat JH, van Herwerden L (2016) The historical biogeography of groupers: clade diversification patterns and processes. Mol Phylogenet Evol 100:21–30. https://doi.org/10.1016/j. ympev.2016.02.012
- Ma KY, van Herwerden L, Newman SJ, Berumen ML, Choat JH, Chu KH, Sadovy de Mitcheson Y (2018) Contrasting population genetic structure in three aggregating groupers (Percoidei: Epinephelidae) in the Indo-West Pacific: the importance of reproductive mode. BMC Evol Biol 18:180. https://doi.org/10.1186/s12862-018-1284-0
- Maggio T, Andaloro F, Hemida F, Arculeo M (2005) A molecular analysis of some Eastern Atlantic grouper from the *Epinephelus* and *Mycteroperca* genus. J Exp Mar Biol Ecol 32:83–92. https://doi. org/10.1016/j.jembe.2005.01.004
- Moriss A, Roberts CM, Hawkins JP (2000) The threatened status of groupers (Epinephelinae). Biodivers Conserv 9:919–942. https:// doi.org/10.1023/A:1008996002822
- Payet SD, Hobbs J-P A, DiBattista JD, Newman SJ, Sinclair-Taylor T, Berumen ML, McIlwain JL (2016) Hybridisation among groupers (genus *Cephalopholis*) at the eastern Indian Ocean suture zone: taxonomic and evolutionary implications. Coral Reefs 35:1157–1169. https://doi.org/10.1007/s00338-016-1482-4

- Peters WCH (1865) Über einige Bloch'sche Arten der Fisch-Gattung Serranus. Monatsberichte der Königlichen Preussischen Akademie der Wissenschaften zu Berlin 1865:97–111
- Puillandre N, Lambert A, Brouillet S, Achaz G (2012) ABGD, automatic barcode gap discovery for primary species delimitation. Mol Ecol 21:1864–1877
- Qu M, Tang W, Liu Q, Wang D, Ding S (2018) Genetic diversity within grouper species and a method for interspecific hybrid identification using DNA barcoding and RYR3 marker. Mol Phylogenet Evol 121:46–51. https://doi.org/10.1016/j.ympev.2017.12.031
- Randall JE, Allen GR (1987) Four new serranid fishes of the genus Epinephelus (Perciformes: Epinephelinae) from Western Australia. Rec West Aust Mus 13:387–411
- Randall JE, Justine JL (2008) *Cephalopholis aurantia* x *C. spiloparaea* a hybrid Serranid fish from New Caledonia. Raffles B Zool 56:157–159
- Ratnasingham S, Hebert PDN (2013) A DNA-based registry for all animal species: the barcode index number (BIN) system. PLoS One 8: e66213. https://doi.org/10.1371/journal.pone.0066213
- Richardson J (1846) Report on the ichthyology of the seas of China and Japan. Report of the British Association for the Advancement of Science 15th meeting [1845]: 187-320.
- Sachithanandam V, Mohan PM, Muruganandam N, Chaaithanya IK, Dhivya P, Baskaran R (2012) DNA barcoding, phylogenetic study of *Epinephelus* spp. from Andaman coastal region, India. Indian J Mar Sci 41:203–211
- Sadovy de Mitcheson Y, Craig MT, Bertoncini AA, Carpenter KE, Cheung WWL, Choat JH, Cornish AS, Fennessy ST, Ferreira BP, Heemstra PC, Liu M, Myers RF, Pollard DA, Rhodes KL, Rocha LA, Russell BC, Samoilys MA, Sanciangco J (2012) Fishing groupers towards extinction: a global assessment of threats and extinction risks in a billion dollar fishery. Fish Fish. https://doi.org/10. 1111/j.1467-2979.2011.00455.x
- Schoelinck C, Hinsinger DD, Dettai A, Cruaud C, Justine J-L (2014) A phylogenetic re-analysis of groupers with applications for ciguatera fish poisoning. PLoS One 9:e98198. https://doi.org/10.1371/ journal.pone.0098198
- Shaw G, Nodder FP (1812) The Naturalist's Miscellany, or coloured figures of natural objects; drawn and described from nature. J Cooper, London. 24 vols. unnumbered pages.
- Shen Y, Guan L, Wang D, Gan X (2016) DNA barcoding and evaluation of genetic diversity in Cyprinidae fish in the midstream of the Yangtze River. Ecol Evol 6:2702–2713. https://doi.org/10.1002/ ece3.2060
- Smith WL, Craig MT (2007) Casting the percomorph net widely: the importance of broad taxonomic sampling in the search for the placement of serranid and percid fishes. Copeia 2007:35–55. https://doi.org/10.1643/0045-8511(2007)7[35:CTPNWT]2.0.CO;2
- Son DM, Thuoc P (2003) Management of coastal fisheries in Vietnam. In Silvestre G, Garces L, Stobutzki I, Ahmed M, Valmonte-Santos RA, Luna C, Lachica-Aliño L, Munro P, Christensen V, Pauly D (eds) Assessment, management and future directions for coastal fisheries in Asian countries. World Fish Center Conference Proceedings 67, pp 957–986
- Stocks AP, Foster SJ, Bat NK, Vincent ACJ (2019) Local fishers' knowledge of target and incidental seahorse catch in southern Vietnam. Hum Ecol 47:397–408. https://doi.org/10.1007/s10745-019-0073-8
- Temminck CJ, Schlegel H (1843) Pisces. Siebold PF de (ed.): Fauna Japonica, sive descriptio animalium, quae in itinere per Japoniam

... suscepto annis 1823-1830 collegit, notis, observationibus et adumbrationibus illustravit Ph. Fr. de Siebold. Lugduni Batavorum [Leiden] (A. Arnz et soc.). Part 1: 1-20.

- Thai BD, Si VN, Phan PD, Austin CM (2007) Phylogenetic evaluation of subfamily classification of the Cyprinidae focusing on Vietnamese species. Aquat Living Resour 20:143–153. https://doi.org/10.1051/ alr:2007025
- Thuoc P (2001) Scientific basis for sustainable management of coastal fisheries resources in Vietnam. In Proceedings of marine fisheries research, Volume II. Research Institute for Marine Fisheries, Ministry of Fisheries, pp 279–300
- Tran CT, Vo VQ, Le TTT, Nguyen PUV, Tran THH (2015) Species composition of the grouper specimens (Serranidae) in museum of oceanographic. The Proceedings of the Vietnam Conference of Ecology and Biological resources 6<sup>th</sup>, pp 327–333 (in Vietnamese)
- Tran VTT, Pham LK, Durand J-D (2016) Diversity and distribution of cryptic species within the *Mugil cephalus* species complex in Vietnam. Mitochondrial DNA Part A 28:493–501. https://doi.org/ 10.3109/24701394.2016.1143467
- Tucker SJ, Kurniasih EM, Craig MT (2016) A new species of grouper (*Epinephelus*; Epinephelidae) from the Indo-Pacific. Copeia 104: 658–662. https://doi.org/10.1643/CI-16-398
- Vaillant LL (1878) in Vaillant LL, Bocourt F (1874-1915) Études sur les poissons. In: Mission scientifique au Mexique et dans l'Amérique centrale. Recherches zoologiques. Quatrième partie. Paris. Part 4, 265 pp.
- Vilgalys R (2003) Taxonomic misidentification in public DNA databases. New Phytol 160:4–5. https://doi.org/10.1046/j.1469-8137. 2003.00894.x
- Vo VQ (2018) Diversity of grouper fish (Serranidae) in coastal zone waters from Da Nang to Binh Thuan. J Mar Sci Technol 4A:101–113 (in Vietnamese)
- Vo VQ, Le TTT, Nguyen TTV, Tran THH, Nguyen PUV, Tran CT (2016) Diversity and catching status of grouper (Serranidae) in coastal waters off Da Nang and Quang Nam. J Mar Sci Technol 4: 405–417 (in Vietnamese)
- Ward RD, Zemlak TS, Innes BH, Last PR, Hebert PDN (2005) DNA barcoding Australia's fish species. Philos Trans R Soc Lond B 360: 1847–1857. https://doi.org/10.1098/rstb.2005.1716
- Ward RD, Hanner R, Hebert PDN (2009) The campaign to DNA barcode all fishes, FISH-BOL. J Fish Biol 74:329–356
- Williams JT, Viviani J (2016) Pseudogramma polyacantha complex (Serranidae, tribe Grammistini): DNA barcoding results lead to the discovery of three cryptic species, including two new species from French Polynesia. Zootaxa 4111:246–260. https://doi.org/10.11646/ zootaxa.4111.3.3
- Winterbottom R, Hanner R, Burridge M, Zur MA (2014) Cornucopia of cryptic species - a DNA barcode analysis of the gobiid genus Trimma (Percomorpha, Gobiiformes). Zookeys 381:79–111
- Zhuang X, Qu M, Zhang X, Ding S (2013) A comprehensive description and evolutionary analysis of 22 grouper (Perciformes, Epinephelidae) mitochondrial genomes with emphasis on two novel genome organizations. PLoS ONE 8:e73561. https://doi.org/10. 1371/journal.pone.0073561