Sponges have recently been proposed as ideal candidates to act as natural samplers for environmental DNA due to their efficiency in filtering water. However, validation of the usefulness of DNA recovered from sponges to reveal vertebrate biodiversity patterns in Marine Protected Areas is still needed. Additionally, nothing is known about how different sponge species and morphologies influence the capture of environmental DNA and whether biodiversity patterns obtained from sponges are best described by quantitative or qualitative measures. In this study, we amplified and sequenced a vertebrate specific 12S barcode with a set of universal PCR primers (MiFish) for metabarcoding environmental DNA from fishes, to unveil fine-scale patterns of fish communities from natural-sampler DNA retrieved from 64 sponges (16 species) located in eutrophic and well-preserved coral reefs in Nha Trang Bay (central Vietnam). Ninety tropical fish species were identified from the sponges, corresponding to one third of the total local ichthyofauna reported from previous extensive conventional surveys. Significant differentiation in fish communities between eutrophic and well-preserved environments was observed, albeit eutrophication only explained a modest proportion of the variation between fish communities. Differences in efficiency of capturing fish environmental DNA among sponge species or morphologies were not observed. Overall, the majority of detected fish species corresponded to reef-associated small-sized species, as expected in coral reefs environments. Remarkably, pelagic, migratory, and deep-sea fish species were also recovered from sponge tissues, pointing out the ability of sponge natural sampled DNA to detect fishes that were not permanently associated to the biomes where the sponges were sampled. These results highlight the suitability of natural samplers as a cost-effective way to assess vertebrate diversity in hyper-diverse environments.

Keywords: natural samplers, environmental DNA, sponges, fish diversity, coral reefs
INTRODUCTION

Environmental DNA (eDNA) refers to the DNA extracted directly from an environmental sample without the need of isolating the target organisms (Thomsen and Willerslev, 2015; Creer et al., 2016; Cristescu and Hebert, 2018). Environmental DNA metabarcoding is emerging as a fast and efficient method for biodiversity monitoring that provides increased sensitivity and reduced costs compared to conventional surveys (Kelly et al., 2014b; Thomsen and Willerslev, 2015; Creer et al., 2016; Boussarie et al., 2018; Sales et al., 2020). Using eDNA provides different opportunities, such as the detection of cryptic taxa or different life stages, the rapid quantification of species richness and the non-invasiveness of sampling procedures, among others (Deiner et al., 2017). However, challenges remain for its use in different fields of study such as biomonitoring, ecology, conservation and invasion biology (Deiner et al., 2017).

A persistent matter of debate in metabarcoding studies is the quantification of species abundances inferred from eDNA data (Kelly, 2016; Deiner et al., 2017). This quantification issue can be seen in a new light if the actual complexity of eDNA is considered. A natural eDNA sample is always a complex mix, comprised of two inseparable fractions with strikingly dissimilar concentrations. The trace-quantities of extra-organismal DNA released by macrofaunal individuals into their environment (which is what most researchers restrictively understand as “eDNA”) are embedded in an overwhelming background of genomic DNA fragments extracted from the community of living microorganisms (mostly bacteria, but also microeukaryotes) that were present in the environment when the sample was taken. Whether a method is targeting the trace-DNA of macrofauna or the community-DNA of microorganisms is determined by the specificity of the metabarcoding primers used. Methods relying on the detection of trace-DNA can be seriously hindered by non-specific amplification of hyper-abundant bacterial sequences if primers are not specific-enough (Collins et al., 2019). At the other end of the spectrum, primers with high specificity values applied on samples containing too low concentrations of the target sequences usually lead to stochastic amplification results and reproducibility issues (Kelly et al., 2019). The use of multiple PCR replicates has been proposed as a way to overcome this recurrent problem (Sigsgaard et al., 2019), but this solution leads to increased costs and needed sample volumes.

However, the real ability of the proposed methods based on natural sampler DNA (nsDNA) to provide robust ecological information, which could be translated into management decisions, is yet to be tested in a real-life scenario. Although promising, much work remains to be done to validate the use of sponges as natural samplers for marine biodiversity studies.

Environmental management and policy decision making for Marine Protected Areas (MPA) can greatly benefit from genetic monitoring (Kelly et al., 2014b; Shum et al., 2019), which can facilitate the assessment, among other biota, of fish community structure and distribution. Several traditional biodiversity studies have been carried in Nha Trang Bay MPA (Central Vietnam), although few have been published (Vo et al., 2004; Van Nguyen and Kim Phan, 2008). In the present study, we aim to (1) validate the usefulness of nsDNA recovered from sponges to reveal fine-scale fish biodiversity patterns in Nha Trang Bay, (2) assess the efficiency in capturing nsDNA between different sponge species and morphologies, and (3) evaluate whether quantitative or qualitative data represented better the fish communities. With the aim to work toward the validation of sponges as an alternative genetic tool for biodiversity assessments, we also compared our results with visual census data performed in the same area.

MATERIALS AND METHODS

Sponge Samples and Study Site

DNA extracts from sponge tissues were originally collected for research on microbiome structure (Turon et al., 2018, 2019) and were used in the present study as opportunistic samples for the analysis of nsDNA captured by the sponges. These DNA extracts had been previously stored in freezing facilities which also contained other samples from temperate species and which not followed specific routines to avoid cross-contamination by trace-DNA. However, no other tissue or DNA sample from Indo-Pacific origin were ever stored along with these samples. Sponges had been originally collected in April 2015 and intended to reflect the most common sponges found in coral reef areas of Nha Trang Bay (Central Vietnam) at depths between 3 and 9 m (Turon et al., 2019). DNA extractions from small fractions (0.5 cm$^3$) of sponge samples were performed using DNeasy Blood & Tissue Kit (Qiagen). In total, 86 samples belonging to 24 different sponge species distributed across six different locations in Nha Trang Bay were processed (Figure 1). The number of replicates per sponge species ranged between 1 and 12 and sponge morphology were classified as massive, branching, thick encrusting or encrusting (Table 1). Among the sampled locations, Dambay represented an eutrophic area, highly impacted by mariculture activities, and the five other locations represented relatively well-preserved areas within the Nha Trang Marine Protected area (Tkachenko et al., 2016).

Sample Processing and Sequencing

To assess the composition of fish assemblages recoverable from the sponge tissues, total DNA extracted from sponge samples was amplified using the high-specificity MiFish primer set targeting a hypervariable region in the mitochondrial 12S rRNA gene (163–185 bp), which specifically amplifies fish and other vertebrates (Miya et al., 2015). All amplification and library preparation works were performed following strict clean lab routines at the Norwegian College of Fishery Science. PCR amplifications were conducted in 15 µl reactions containing 2 µl of DNA template, 10 µl of AmpliTaq Gold Master mix, 0.16 µl of Bovine Serum Albumin (20 µg/µl), 1 µl of each forward (5'- GTCCGTAAAACTCGTGCCAGC-3') and reverse (5'-CATAGTGGGTATC TAATCCGAGTTG-3') primer (5 µM) and 5.84 µl of H$_2$O. Both primers included 7-base tags in 5’ which uniquely identified amplicons belonging to the same sample, and a variable number (2–4) of leading Ns in order to increase sequence diversity within the sequencer flow-cell. The temperature profile was as follows: 94°C for 10 min;
40 cycles $\times (95^\circ C/30\text{ s}, 60^\circ C/30\text{ s}, 72^\circ C/30\text{ s}); 72^\circ C/5\text{ min}$. Two PCRs were run per sample as technical replicates, and a negative control was run as above but without DNA template added. The success of PCR amplifications was checked by gel electrophoresis in 1% agarose and PCR products were then pooled together into two multiplex sample pools (one per PCR replicate). MinElute PCR purification columns (Qiagen) were used to concentrate the pooled DNA and to remove fragments below 70 bp. Library preparation was performed with the NEXTflex PCR-free library preparation kit (BIOO Scientific) and the exact library concentration was measured in a qPCR using the NEBNext Library Quant Kit (New England BioLabs). Finally, pools were sequenced along with 1% PhiX on an Illumina MiSeq platform using v2 chemistry ($2 \times 150\text{ bp}$).

**Metabarcoding Pipeline**

The OBITools v1.01.22 software suite (Boyer et al., 2016) was used for the initial steps of the bioinformatic analyses. Paired-end reads were aligned using *illumina*·*pairedend* and only sequences with alignment quality score $>40$ were kept. Demultiplexing was done with *ngsfilter*, which also removed primer sequences. Aligned reads with length of 140–190 bp and without ambiguous positions were selected using *obigrep* and then dereplicated with *obiuniq*. Chimeric sequences were removed using the *uchime*-de novo algorithm implemented in vsearch v1.10.1 (Rognes et al., 2016). Clustering of sequences into molecular operational taxonomic units (MOTUs) was performed using SWARM 2.0 algorithm (Mahé et al., 2015) with a $d$ value of 1. Taxonomic assignment of the most abundant (representative) sequence of each MOTU was done with the *ecotag* algorithm (Boyer et al., 2016) against the DUFA_MiFish v2020-05-01 reference database, compiled from sequences of the MiFish fragment available from Genbank and complemented with some in-house sequencing of selected taxa. This database is publicly available from github.com/uit-metabarcoding/DUFA. Further manual refining of the dataset consisted of removing some MOTUs identified as human or terrestrial vertebrates or as fish species clearly not distributed in the area, likely contaminations from other
projects being conducted at the laboratories at Salford University (United Kingdom) and CEAB-CSIC (Spain), where samples had been previously stored.

**Data Analysis**

Statistical analyses were performed in R version 3.1.3 with the vegan package (version 2.5-6; Oksanen et al., 2019) and graphic visualizations were done with ggplot2 package (Wickham, 2016). Species accumulation curve was drawn using the speacaccum function of the vegan package. Reads were first transformed to relative abundances to build a Bray-Curtis dissimilarity matrix (PCR replicates not pooled together), which was used to assess the variance in community composition using Permutational Multivariate Analyses of Variance (PERMANOVA). The samples were categorized as a function of Environment (eutrophic, normal), Location (6 levels), Species (16 levels), and Sponge morphology (4 levels) and the univariate effects of these factors on the community composition were tested using adonis function with 1,000 permutations. Additionally, PERMDISP analysis (betadisper function) was performed to determine if the significance of factors was due to different multivariate mean or to different heterogeneity of the groups. Non-metric multidimensional scaling (nMDS) representation with Bray-Curtis dissimilarities was performed with the metaMDS function with 500 iterations. Shannon diversity and MOTU richness per sample were calculated in vegan (Oksanen et al., 2019) and significant differences between sponge species and sponge morphologies were tested with Kruskal-Wallis test. The remaining analyses were made with presence/absence data (see Results). A heatmap was used to represent species with 0, 1, and 2 detections per sample. Only samples with positive counts were used for graphic representations.

**Fish Metadata**

All the fish MOTUs that were identified at species or genus level were grouped according to the following biological and ecological traits: schooling behavior (solitary, small, large), individual size (small (<15 cm), medium (15–50 cm), large (>50 cm), depth range (surface (0–10 m), shallow (10–60 m), deep (>60 m), habitat (reef-associated, pelagic) and migratory behavior (yes, no). We then calculated the percentage of total detections and total reads that belonged to the different categories of each trait. Moreover, we assessed the mean detections and the mean reads per MOTU according to the traits' categories.

**Visual Census**

We compared our eDNA results with visual census data reported in Van Nguyen and Kim Phan (2008). In that study visual censuses were performed on coral reef fishes in 16 different

### TABLE 1 | Sponge species collected in Nha Trang Bay (Vietnam).

| Species                  | N. of replicates | N. of replicates with detections | Morphology          |
|--------------------------|------------------|----------------------------------|---------------------|
| Aaptos suberitoides      | 12               | 9                                | Massive             |
| Amphimedon paraviridis   | 9                | 3                                | Thick encrusting-massive |
| Amphimedon sulcata       | 10               | 9                                | Thick encrusting-massive |
| Antho (Antho) sp.        | 2                | 2                                | Encrusting          |
| Cladocroce sp.           | 1                | 0                                | Encrusting          |
| Clathria (Thalysas) reinwardti | 11              | 10                               | Branching-massive   |
| Dendroxea sp.            | 1                | 0                                | Encrusting          |
| Didiscus aceratus        | 1                | 0                                | Massive             |
| Dysidea sp.              | 2                | 2                                | Encrusting          |
| Gelliodes fibulata       | 1                | 1                                | Branching           |
| Haliclona (Reniera) sp1  | 1                | 1                                | Encrusting          |
| Haliclona (Reniera) sp4  | 3                | 3                                | Encrusting          |
| Haliclona (Reniera) sp7  | 1                | 1                                | Encrusting          |
| Haliclona (Gellius) toxotes | 3               | 3                                | Encrusting          |
| Hyrtios spinifer         | 1                | 0                                | Encrusting          |
| Monanchora unguiculata   | 9                | 7                                | Massive             |
| Mycale (Arenochalina) sp.| 3                | 3                                | Thick encrusting    |
| Neofibularia sp.         | 9                | 9                                | Thick encrusting    |
| Petrosia sp.             | 1                | 0                                | Encrusting          |
| Phorbas sp.              | 1                | 1                                | Encrusting          |
| Protosuberites proteus   | 1                | 1                                | Thick encrusting    |
| Suberea fusca            | 1                | 0                                | Thick encrusting    |
| Terpios cruciatus        | 1                | 0                                | Encrusting          |
| Thrinachophora cervicomis| 1                | 0                                | Thick encrusting-massive |

Different number of replicates were used to perform PCRs with MiFish primers for each sponge species (second column). Third column indicates the number of replicates with successful detection of fish species. Morphology of each sponge species is indicated in the last column.

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1https://www.R-project.org/
locations in Nha Trang Bay during September-October 2008. Overall, these authors performed 32 transects. Each transect was 100 m long and 5 m wide, thus covering an area of 500 m². Fish present within 5 m above the bottom were recorded. Venn diagrams were used to represent the fish families, genera and species recovered by both sampling methods. Additionally, we searched in Fishbase (Froese and Pauly, 2019)

RESULTS

Fish Richness Recovered From Sponges

Our final dataset consisted of 90 fish MOTUs recovered from 172 PCR samples (86 sponge samples with 2 PCR replicates each). Of these, only 16 sponge species (64 samples), from the original 24 species selected, had positive PCR counts in at least one of the PCR sample replicates after removal of contaminants (Table 1). Indeed, the species accumulation curve did not reach an asymptote, suggesting that more sampling effort is needed to increase the accumulation rate of new MOTUs and accurately capture the whole fish diversity in the area (Figure 2). MOTU richness per sample was rather low, with the majority of samples having only one or two MOTUs. The maximum MOTU richness was 12 in a single sample (Supplementary Figure S1). No significant differences were observed in the number of MOTUs recovered from different sponge species or sponge morphologies (Supplementary Figure S2 and Supplementary Table S1). Within species, no coherent pattern was detected either, as different sponge replicates of a given species presented varying values of MOTU richness (Supplementary Figure S2).

The sequencing produced a highly differential number of reads per sample. In general, the number of reads from samples that only had one positive PCR replicate was lower than the number of reads from samples that had two positive PCR replicates (Supplementary Figure S3). However, the difference in number of reads between the two PCR replicates was high, suggesting an important stochastic component of the PCR amplification of the 12S MiFish primer set (Supplementary Figure S3).

The majority of the MOTUs were identified at the species (76 MOTUs) or genus (11 MOTUs) level, with only three MOTUs that could not be classified at lower taxonomic level than class (Supplementary Table S2). In our dataset, each MOTU belonged to a different fish species, so both terms (MOTU and species) will be used interchangeably hereafter. We identified 40 different families, among which Pomacentridae (11 MOTUs), Labridae (7 MOTUs), Gobiidae (5 MOTUs), and Apogonidae (5 MOTUs) were the ones with most representatives.

Ordination of Fish Communities

The non-metric multidimensional scaling (nMDS) representation based on relative read abundance showed slightly different fish communities between eutrophic and well-preserved environments, although with a large overlap (Figure 3). PERMANOVA analyses of the environment (adonis: $R^2 = 0.02, p < 0.001$) and sponge morphology (adonis: $R^2 = 0.04, p < 0.001$) factors were significant but only explained a small proportion of the total variation in the nsDNA fish communities (Supplementary Table S3). Similarly, sampling location (adonis: $R^2 = 0.08, p < 0.001$) was a significant factor in structuring fish communities, but only explained a low amount of the total variation (Supplementary Table S3 and Supplementary Figure S4). However, PERMDISP test showed that location effect could be explained by differences in dispersion levels across locations. In a similar manner, differences in the nsDNA from fish communities between sponge species (adonis: $R^2 = 0.21, p < 0.001$) were significant, but this effect could be due to significant heterogeneity in dispersion across species (Supplementary Table S3).

Presence/Absence Data

Most of the PCR replicates from the same sample clustered together in Bray-Curtis distance of relative abundance data (Supplementary Figure S5), although there was a high variation in the number of reads recovered from each PCR replicate (Supplementary Figure S3). That is why we thus chose to work with presence/absence data with replicates pooled together, considering “detection” whenever a MOTU was detected in at least one of the PCR replicates from the same sample (Figure 4). Ordination of sponge samples based on Jaccard distances (Supplementary Figure S6) did not show any specific pattern related to sponge species, location or environment (Figure 4). Spratelloides gracilis and Pterocaesio digramma were the fish species most frequently occurring across samples (with more than 15 detections), but the majority of fish species (MOTUs) were detected just once in a single sample (Figure 4).
Nonetheless, the majority of sponge samples had between one and five fish detections when pooling the two PCR replicates per sample.

**Fish Community Ecological Features**

We were able to classify the 87 MOTUs identified at genus or species level according to their schooling behavior, individual size, depth range, habitat, and migratory behavior. We found a differential distribution of the total detections and reads that belonged to different trait categories, considering the whole sampling area (Figure 5). For the schooling behavior, we detected a similar frequency fishes of the three schooling categories: large groups, small groups or solitary individuals (Figure 5A), although a higher percentage of reads was found for fishes that are usually found in large schools, such as *Spratelloides gracilis* (~24%) and *Thunnus obsesus* (~15%). Similarly, individuals of the three size classes were present in our dataset, but small- and medium-sized fishes were more frequently detected, with the highest percentage of the total reads from small fishes (Figure 5B). This was mostly due to the high percentage of reads retrieved from *Spratelloides gracilis* (~24%) and *Plectroglyphidodon dickii* (~17%), two species represented by individuals < 15 cm. Although the collected sponges had a surface distribution (3–9 m), the majority of the fish species retrieved from the sponge samples have a shallow depth preference (10–60 m).

Moreover, there was a non-negligible proportion (~15% detections and reads) of fish species that are associated with a deeper depth range (>60 m) (Figure 5C). Finally, the majority of the MOTUs in our sponge nsDNA survey belonged to non-migratory reef-associated fish species (Figures 5D,E). Among the most common reef species, *Apogon* sp. and *Pterocaesio digramma* had the highest proportion of detections, and *Acanthurus nigrofuscus* and *Plectroglyphidodon dickii* had the highest proportion of reads. Overall, the majority of fish detected in the samples were reef-associated small individuals, with shallow depth range and non-migratory behavior.

There was a significant correlation between the number of detections and the number of reads (log transformed) (Pearson, $R_p = 0.58$, $p < 0.001$, Figure 5F). In general trends, the most frequently detected MOTUs also had the highest number of reads and belonged to fish species with schooling behavior. However, it is important to note that there were many MOTUs with just a single detection that presented a wide range of variability in their number of reads (Figure 5F). The number of total detections and the number of total reads (log transformed) were classified into three abundance categories for each MOTU (Figure 5F): high (> 10), medium (5–10), low (2–5). *Thunnus obsesus* and *Spratelloides gracilis* were the two species found at high abundances considering both, detections and reads (Figure 6). Other highly abundant species were *Apogon* sp. and *Pterocaesio*
digramma in terms of detections (Figure 7F), and Acanthurus nigrofuscus and Plectroglyphidodon dickii in terms of number of reads (Figure 6).

**Mean Detections/Reads per MOTU**

In order to investigate which fish species were more likely to be detected depending on their behavior or physiological characteristics, we calculated the mean detections and reads per MOTU and grouped them according the above explained categories. The most relevant differences were observed for the schooling behavior, for which, species that are usually found in large schools had a much higher detections and reads per individual than solitary individuals (Figure 7A and Supplementary Figure S7A). No differences were observed regarding the individual size for detections (Figure 7B), although small individuals presented a higher number of reads than larger individuals. This latter case is mostly due to the high number of reads belonging to the herring (S. gracilis), which is usually found in large schools. Moreover, shallow, pelagic and migratory fishes had a higher number of reads and detection rates than reef-associated non-migratory species (Figures 7C–E and Supplementary Figures S7D,E).

**Comparisons Between nsDNA Sponge Survey and Visual Census Data**

With the aim of assessing to what extend the nsDNA recovered from sponges captured the fish diversity reported in the same area, we compared our results with the visual census data reported in Van Nguyen and Kim Phan (2008). Although the study area was similar for both studies (Nha Trang Bay), the sampling effort was not comparable, being much higher in the visual survey, where 16 different locations were surveyed and thus covering a much larger area of the bay than our study. The total fish richness recovered was higher for the visual survey than for the nsDNA data with 266 and 90 species, respectively. Among them, 24 species were detected...
FIGURE 5 | Percentage of total detections and total reads associated to specific categories within the studied fish features: (A) Aggregation behavior, (B) Individual size, (C) Depth range, (D) Habitat, and (E) Migration. (F) Relationship between total reads (log transformed) and total detections colored by the aggregation behavior of the represented MOTUs (fish species).

by both methods (Figure 8) and all of them belonged to reef-associated fishes. Thirty different genera were detected by both methods, with 79 genus exclusively detected in visual surveys and 38 genus exclusively detected with nsDNA. At the family level, up to 40 different families were captured by both methods and more than 50% of them were shared. Of the 90 species detected by nsDNA, 62 were reported for Vietnam in the FishBase, and 52 of them had specific reports in Nha Trang Bay. Up to 16 species had other representatives of the same genus reported in Vietnam and only 12 species had no genus representative described in Vietnam according to the FishBase.

DISCUSSION

Sponges are cosmopolitan and the most efficient water filterers in the world (Van Soest et al., 2012). Because of that, they have been recently proposed as ideal candidates to act as natural environmental samplers (Mariani et al., 2019). In this study, we used specific vertebrate primers (Miya et al., 2015) to unveil fine-scale fish patterns from the nsDNA retrieved from the sponge tissues. We were able to identify up to 90 fish species from the sponges collected, although species accumulation curves showed that more replicates are needed to accurately capture the fish diversity of the area. Indeed, a significant proportion
of our samples (34%) did not retrieve any positive PCR counts. This result suggests that fish nsDNA is not always present in sponges, in accordance with previous results from Mariani et al. (2019), in which two out of nine samples contained too few reads/MOTUs to be retained for further analysis. Keep in mind that our sampling was originally intended for microbiome studies, so the samples comprised ca. 0.5 cm$^3$ of sponge material, which may be too small for other purposes. Specific studies should be carried out by repeatedly sampling the same sponge to determine the right sample volume necessary to reliably detect nsDNA and accurately capture the fish diversity in a given locality. From our results, it is clear that fish nsDNA was not abundant or ubiquitous in the sponges analyzed, leading to a relatively high stochasticity in detection levels and in number of reads obtained from the samples. Whether this observed stochasticity is lower or higher than those that would be obtained from comparable filtered seawater, or if it is an artifact due to the high amount of contamination reads detected, is yet to be studied. Moreover, we did not have a control for extraction kit contamination, although this is unlikely to affect our results. Regardless of these constraints, significant differentiation in the fish communities was observed between the eutrophic and the well-preserved environments and between sampling locations, albeit these factors only explained a modest proportion of the observed variation. Sponge species proved also to be a significant factor for detecting fish, albeit a high dispersion between species prevented drawing definite conclusions. The morphological characteristics of the sponge species (encrusting, massive, branching), did not seem to affect the capturing of nsDNA, as no sponge species performed significantly better or worse than others in terms of MOTU richness. Although sponges can have interspecific differences in filtration efficiencies, it has been shown that similar high efficiency is obtained after a few hours of incubation by species with contrasting biological characteristics (Turon et al., 1998).

Reassuringly, consistency was found between PCR replicates of the same sample, indicating that there was no technical artifact related to amplification. Notwithstanding, given that our extracts in general have little DNA of the target group, PCR bias due to the non-linear nature of the amplification reaction can stochastically lead to markedly different number of reads between PCR replicates. In the case of our dataset, with the aim of not adding more sources of variation than the ones already intrinsic to our sampling (environment, location, sponge species), we chose to focus on presence/absence data rather than focusing on differences in the relative abundance of reads. Indeed, presence/absence data has been proven to better differentiate fish communities between sites than read abundances (Port et al., 2016; DiBattista et al., 2017). The majority of sponges only contained between one and five fish MOTUs considering the two PCR replicates per sponge, with the exception of C. (Thalysas) reinwardti, for which up to 12 MOTUs were retrieved. However, other replicates of this species had low MOTU richness, comparable to other sponge species, preventing the consideration of this species as a “better” natural sampler than others. Considering our data, MOTU richness per sponge species is a rather stochastic factor, although a more detailed study focusing in less species and more replicates should better address this issue. No specific pattern for fish communities could be ascribed to sampling location, which were closely located (max ∼2 km apart). Water movements can highly affect the distribution of eDNA in the environment (Goldberg et al., 2016), possibly explaining this lack of specific patterns between fine-scale sampling locations. However, Jeunen et al. (2019) demonstrated...
that eDNA was able to discriminate diverse marine habitats within a small spatial scale subject to tidal and water currents. Indeed, some studies have already shown that eDNA surveys are able to detect fine-scale community composition patterns in heterogeneous aquatic habitats (Port et al., 2016; O’Donnell et al., 2017; Pont et al., 2018; Bakker et al., 2019; Stat et al., 2019;
Nguyen et al., 2020; Oka et al., 2020). In our study, the residency
time and degradation rate of nsDNA in the sponge tissues is an
important factor that can hamper the detection of community
patterns. To date, only one other study has assessed vertebrate
communities using nsDNA from Mediterranean and Antarctic
sponges (Mariani et al., 2019). Thus, further studies are needed
to prove the viability of sponges as natural samplers to detect
community patterns at small geographical scales.

The Most Common Fish Species and
Their Features
With the aim to describe the main features of the fish
sequences retrieved from the sponges of Nha Trang Bay we
considered several ecological and biological traits in our dataset.
A high proportion of the fishes observed were reef-associated
small individuals with shallow depth range and non-migratory
behavior, reflecting the general characteristics of the fishes one
may expect to find in a coral reef environment. Pterocaesio
digramma, Acanthurus nigrofuscus, Plectroglyphidodon dickii,
and Apogon sp. were the most common reef fish species detected
in the sponge tissues. It is relevant to point out that we were
able to detect eDNA from fishes with features not typical of reef-
associated species and thus, not permanently associated to the
environment where the sponges were sampled, such as pelagic,
migratory or deep-sea species. According to our molecular
survey, pelagic fishes might have a more important presence in
coral reefs than previously assessed by visual surveys, such as the
large schools of tuna and herring. Detection of certain species
can be influenced by the efficiency of detecting rare species, and
tiny individuals or larvae (Yamamoto et al., 2017) but also, by
fish movement patterns and residence times (Stat et al., 2019).
In that sense, if residence time of nsDNA in sponges is long
enough, using these biological samplers can provide a picture of the
ichthyofauna integrated over time, rather than a snapshot that
can be acquired by a punctual visual sampling, or by collecting
eDNA directly from water. Clearly, the dynamics of nsDNA in
sponges (and other filter feeders as well) is an interesting field for
further research.

In this analysis we have considered both detections and total
number of reads to assign a certain abundance level to the
observed MOTUs. Although there is a relationship between the
number of reads and the number of detections, it is important to
note the high variability found in the number of reads for single
detected fish species, probably due to PCR bias discussed above.
Overall, among the most abundant fishes for both parameters we
observed MOTUs. Although there is a relationship between the
number of reads and the number of detections, it is important to
note the high variability found in the number of reads for single
detected fish species, probably due to PCR bias discussed above.
In this sense, although we did not have visual data corresponding to
our sampling period (April 2015), we compared the results obtained
with visual censuses performed in the area in September–October
2008 (Van Nguyen and Kim Phan, 2008). In that study, up to 266 species were detected, and fish communities could be
differentiated depending on the substrate were the survey took
place. It is noteworthy that, with a spatially and temporally
restricted sampling of 86 sponges, we could detect one third of
the species found in extensive conventional surveys. We were able
to detect the same number of fish families as in the visual survey
and half of them were coincident. Indeed, the higher density of
Pomacentridae and Labridae species found in Hun Mun area by
visual censuses (Van Nguyen and Kim Phan, 2008) is coherent
with the higher number of representatives detected for these
families in our nsDNA survey. The majority of species detected
by nsDNA from sponges had been described in Vietnam, and the
majority of them had specific detections in Nha Trang according to
FishBase (Froese and Pauly, 2019). The few species for which
no representatives were found in the database can be indicative of
the incompleteness of the molecular reference database or might
represent new reports of species never detected or misidentified
before in the area.

Comparisons With Visual Census Data
A key point when assessing the use of eDNA data for species
surveys is the comparison with traditional visual surveys.
Although we did not have visual data corresponding to our
sampling period (April 2015), we compared the results obtained
with visual censuses performed in the area in September–October
2008 (Van Nguyen and Kim Phan, 2008). In that study, up to 266 species were detected, and fish communities could be
differentiated depending on the substrate were the survey took
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FishBase (Froese and Pauly, 2019). The few species for which
no representatives were found in the database can be indicative of
the incompleteness of the molecular reference database or might
represent new reports of species never detected or misidentified
before in the area.

It has to be considered that results obtained by both sampling
methods should be standardized by their cost-efficiency (time
and expertise needed). Performing 32 transects of 500 m$^2$ each
is much more time consuming than collecting sponges in a single (or a few) dives. Moreover, traditional censuses (visual or captures) are usually invasive, selective, and can be limited by environmental conditions and lack of taxonomic expertise (Thomsen et al., 2012; DiBattista et al., 2017). Some recent studies have pointed out the differences between eDNA metabarcoding methods and visual or capture methods (Tkachenko et al., 2016; Yamamoto et al., 2017; Fujii et al., 2019; Stat et al., 2019). The main advantages of metabarcoding rely on its time efficiency (Yamamoto et al., 2017; Nguyen et al., 2020), its detection capability of eDNA signatures of fishes that cannot be observed during visual surveys (Nguyen et al., 2020) and the dispensable need of taxonomic expertise (DiBattista et al., 2017). In this sense, censuses made by different observers may not be comparable, while results from eDNA are more objective and, particularly, better suited for archiving and future reanalyses (the sequences remain in databases, visual observations are hardly archivable). However, the main drawbacks of the eDNA approach lie on the accuracy of the results obtained, which are biased by the coverage of the reference database (Nguyen et al., 2020), the number of replicates (Yamamoto et al., 2017), the possible PCR inhibition (Fujii et al., 2019), and the differences in the species DNA shedding rates (Port et al., 2016).

CONCLUSION

With the aim of avoiding the problems associated with water filtration and DNA isolation in eDNA research, the use of natural samplers, such as shrimps (Siegenthaler et al., 2019) or sponges (Mariani et al., 2019) has recently been proposed for surveying marine ecosystems. However, their usefulness remains to be tested in different environmental settings and community types. This study contributed to this goal by using nsDNA obtained from tropical sponges under different eutrophication levels. We were able to capture fish diversity from the area with reasonable resolution considering the limited effort required to sample sponge biopsies compared to traditional visual survey techniques. Although we were not able to discriminate fine-scale geographic patterns, for which a more extensive replication would be needed, we observed significant differences in fish communities between an eutrophic area and the five relatively well-preserved areas within the Nha Trang Marine Protected area. Additionally, fish communities were also observed to be structured significantly different among sample locations, suggesting that sponges are effective in monitoring the stability of fish communities and their ecosystems. Our results also show that sponge species and morphology does not impact the efficiency of capturing nsDNA, as also suggested by Mariani et al. (2019).

Overall, our results show that sponges are promising natural samplers: a small number of sponges provided one third the species found in visual censuses of more than 32 transects performed over 2 months. Most species detected correspond to local ichthyofauna and reef fish were the best represented. However, further studies based on properly designed sampling schemes (as compared to our opportunistic study) are required to determine the right sample volumes and replication levels to obtain reliable quantitative information. Investigation on the trace-DNA dynamics and residence time in the sponges is also necessary. For the time being, results are better interpreted in a qualitative way. Comparative studies between sponges and other filter-feeders (such as ascidians or bivalves) can also shed light on the utility of natural samplers for community characterization.

DATA AVAILABILITY STATEMENT

The raw sequence data and the final datasets generated for this study can be found in the Mendele Data repository with the DOI number: 10.17632/3btg5jz9t8.1.

AUTHOR CONTRIBUTIONS

MT performed the sampling and wrote the first manuscript draft. MT, KP, and OW designed the study. KP acquired the funding. MT, CA-P, and OW performed the lab work. OW performed the bioinformatics pipeline. MT, CA-P, and AA performed the analyses. All authors contributed significantly to later versions of the manuscript and approved submission.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2020.605148/full#supplementary-material
plot shows the difference in number of reads between two PCR replicates of the same sample.

Supplementary Figure 4 | Non-metric multidimensional scaling (nMDS) ordination of fish communities obtained from sponge nsDNA based on Bray-Curtis distances. Colors show differences between locations. Each environment is represented by a different shape.

Supplementary Figure 5 | Heatmap based on relative abundance data of fish communities (y-axis) recovered from sponge nsDNA. Replicates of the same sample (x-axis) are indicated by “a” or “b.” Cluster dendrograms of MOTUs and samples are based on Bray-Curtis distances. Scale bar represents the relative abundance of the fish MOTUs in each sample.

Supplementary Figure 6 | Cluster dendrogram based on Jaccard distances of fish communities recovered from sponge nsDNA. Sample replicates pooled together.

Supplementary Figure 7 | Mean reads per MOTU of the studied fish features: (A) Aggregation behavior, (B) Individual size, (C) Depth range, (D) Habitat, and (E) Migration. (F) Names of the fish species recovered from sponge nsDNA sized according to their number of reads.

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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