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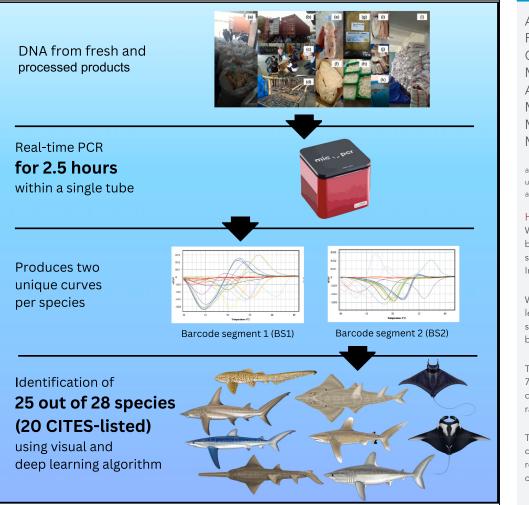
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Article

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Highlights

We used a portable qPCRbased tool to identify shark and ray species in Indonesia

We built a deep machine learning model to assign species from two barcodes

The model achieved 79.41% accuracy for classifying 28 shark and ray species

This *in-situ* diagnostic tool could greatly help in regulating trade operations

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Universal closed-tube barcoding for monitoring the shark and ray trade in megadiverse conservation hotspots

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SUMMARY

Trade restrictions for endangered elasmobranch species exist to disincentivise their exploitation and curb their declines. However, trade monitoring is challenging due to product variety and the complexity of import/export routes. We investigate the use of a portable, universal, DNA-based tool which would greatly facilitate *in-situ* monitoring. We collected shark and ray samples across the Island of Java, Indonesia, and selected 28 commonly encountered species (including 22 CITES-listed species) to test a recently developed real-time PCR single-assay originally developed for screening bony fish. In the absence of a bespoke elasmobranch identification online platform in the original FASTFISH-ID model, we employed a deep learning algorithm to recognize species based on DNA melt-curve signatures. By combining visual and machine-learning assignment methods, we distinguished 25/28 species, 20 of which were CITES-listed. With further refinement, this method can improve monitoring of the elasmobranch trade worldwide, without a lab or species-specific assays.

INTRODUCTION

Biodiversity is depleting more rapidly than at any time in human history. Within the last 50 years, animal species have declined by an average of almost 70% due to continued and increasing anthropogenic stressors.^{1,2} Shark and ray populations (hereafter referred to as "elasmobranchs") have one of the highest extinction risks across the animal kingdom due to fishing pressure, whether targeted or as by-catch,^{3–5} compounded by their conservative life histories.⁶ Although some elasmobranch fisheries can be sustainably managed,⁷ the market demand for shark and ray products typically leads to overexploitation.^{3,8}

The rapid global decline of elasmobranch populations requires collaborative management and conservation measures to ensure the long-term benefits of these populations to the wider ecosystem, including, where sustainable, for human resource use. Binding international trade consortia, such as CITES (Convention on International Trade in Endangered Species of Wild Fauna and Flora), regulate and provide the framework to restrict the international trade of species of priority conservation concern by creating species listing (CITES appendix I and II). Indeed, there has been an increasing number of elasmobranch listings in CITES Appendix II over the last decade with 38 of the 47 species regulated by CITES added at the 16th (2013), 17th (2016), and 18th (2019) Conference of the Parties conventions (Booth et al., 2020). The number of Appendix II listings then more than tripled at the 19th Conference of the Parties (CoP19) in 2022 where parties agreed to add all remaining (54) species of requiem sharks (Carcharhinidae spp.), 6 species of hammerhead sharks, and 37 species of guitarfishes to Appendix II. Seven species of Brazilian freshwater stingrays were also adopted for Appendix II listing. The scale and pace of these listings (now 151 species) present an important implementation challenge for countries with large and diverse landings of sharks and rays, such as Indonesia.

As a result of substantial bycatch, Indonesian fisheries hold the world's largest volume of elasmobranch landings.^{9,10} This exploitation contributes to the high vulnerability rate of elasmobranch populations in Indonesian waters,¹¹ including the populations in its coral reef ecosystems.⁴ This is particularly concerning as Indonesia harbors almost a quarter of the world's elasmobranch diversity.^{12,13} Several measures have been established by the Indonesian authorities to reduce the decline of elasmobranch populations,



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Figure 1. Various types of derivative shark and ray products are being traded

Condition of inspection and some derivative products from sharks and rays i.e. large volumes of mixed cartilages waiting for inspection (A); two containers full of dried shark and ray skin (B); inspectors checking a mixed bag of small fins and finding some hammerhead species' fins (C); caudal fins being dried (D); shark teeth (E); processed ray skin (F); shredded fins "hissit" in brine ready for exporting to Japan (G); blue shark cartilages soaked for processing (H); dried meat from small sharks (I); dried meat from a large shark (J); live bowmouth guitarfish for the aquarium market; and dried fins of silky and hammerhead sharks waiting for quota to export (L).

such as: increasing the number of protected species, extensive outreach programmes, improvement of data collection and stock assessment, expansion of marine protected areas, as well as the establishment of port state measures to combat illegal fishing.^{14–17}

The issue around elasmobranch fisheries is rendered even more challenging by the myriad of shark and ray product derivations, which add another layer of complexity.^{18–20} Due to their similarity in appearance and the lack of distinctive features in most derivative products, elasmobranch species can be deliberately or accidentally mislabeled by those involved in the trade (Figure 1). The general lack of transparency in the trade of living resources is an ongoing concern for fisheries and conservation management²¹ and can have a negative impact on stock management and damages the reputation of entire sectors and countries.^{21,22} Furthermore, the continuous increase of elasmobranch species listed in the CITES Appendices requires constant improvements of national and transnational capabilities in monitoring the supply chain.²³

The rapid development of DNA-based diagnostic tools offers an ever-expanding option for wildlife identification, which have greatly assisted elasmobranch biology and forensics. Established DNA barcoding²⁴ and mini-barcoding²⁵ approaches can robustly identify species in fresh and processed samples. However, these traditional DNA-barcoding methods require longer processing time and high costs for their sequencing processes. More recently, advances in real-time PCR have eliminated the sequencing stage, thereby allowing species identification to be conducted in the field. This approach uses target-specific primers and fluorescent dyes to detect the presence of the targeted nucleic acid template during PCR amplification and has been successfully applied to detect several CITES-listed shark species in a single run tube²⁶ and Multiplex LAMP.²⁷ However, given their reliance on species-specific primers and probes, these methods are better suited to screening large numbers of specimens from one or few target organisms rather than from a wide variety of species. Thus, the need remains for a fast and easy way to identify any sample, by-passing the need to design species-specific assays.

This issue is particularly glaring when inspectors are dealing with multiple types of products from different species across many locations and with a limited time frame to investigate species compositions.²⁸ This year, the magnitude of the challenge has more than tripled, with the number of CITES-listed species going from 47 to 151.^{29,30} Since CITES regulations still allows species listed on Appendix II to be traded by





considering the sustainability of exploitation through a non-detrimental findings (NDF) framework, trade monitoring is more crucial than ever before.

In an attempt to circumvent the limits of species-specific methods, a universal single-tube assay marketed as FASTFISH-ID was recently developed for use in the seafood industry.³¹ This method uses LATE (Linear-After-The-Exponent) PCR to amplify one strand of the full 650bp COI barcoding region,³² and uses a set of fluorescent probes to target two distinct mini-barcode regions selected for their high intra-specific variability which will then produce unique species-specific fluorescent signatures.³¹ The fluorescent signatures.

However, this approach and its libraries were originally designed and validated for bony fishes³¹ and no elasmobranch fluorescence fingerprints are publicly available in the FASTFISH-ID cloud. We therefore chose to test (i) whether the existing FASTFISH-ID diagnostics could produce a diverse range of fluorescent signatures unique and specific to each of the 28 elasmobranch species frequently found in Indonesian trade; and (ii) whether a deep machine learning method could quantitatively assign signatures to the correct species, irrespective of the visual appearance of the fluorescence. Deep learning algorithms are highly flexible and well suited for undertaking these tasks,^{33,34} and have recently been applied in marine science, including fish size estimation,³⁵ bycatch detection and shark identification from photos and videos.^{36–38}

Therefore, this study aimed to examine the power of FASHFISH-ID technology to distinguish between elasmobranch species from trade commodities collected in the whole trade chain (from fishing ports, local buyers, processing plants, and export hubs) across Java Island, Indonesia. We also explore the application of machine learning to tackle the absence of reference fluorescence fingerprints on these species. This portable methodology that can be performed in the field allows for accelerated inspection times for local authorities and could be a vital tool in tacking the illicit trade in endangered sharks and rays.

RESULTS

Fluorescent signature of species

After filtering and removing 33 inconsistent runs (datasets with poor probe-barcode hybridization or inconsistent fluorescent signature), 357 pairs of fluorescent signatures from 28 species were generated, including 14 sharks and 14 rays, with 22 of those species (12 sharks, 10 rays) being CITES-listed species. Within 2.5 h, all types of samples—from fresh to processed samples sourced from different body parts—were amplified and produced one or two fluorescent signatures (referred to as BS1 and BS2 for barcode segment one and barcode segment two) (Tables S1 and S2). These two barcode segments refer to the two mini-barcode regions within the amplified COI target sequence that emitted fluorescent to be read by the real-time PCR machine.

Many species were distinguishable using a combination of both barcode segments and had unique signatures, such as *Alopias pelagicus* (pelagic thresher), *Alopias superciliosus* (bigeye thresher), and *Isurus paucus* (longfin mako shark). However, some species displayed probe-barcode hybridization difficulties (see Methods), with more shark species (7) than ray species (3) being affected, namely *Carcharhinus falciformis* (silky shark), *Carcharhinus longimanus* (oceanic whitetip shark), *Isurus oxyrinchus* (shortfin mako shark), *Lamna nasus* (porbeagle shark), *Carcharhinus brevipinna* (spinner shark), *Galeocerdo cuvier* (tiger shark), *Prionace glauca* (blue shark), *Rhynchobatus laevis* (smoothnose wedgefish), *Glaucostegus typus* (giant shovelnose ray), and *Pristis pristis* (Largetooth sawfish). Nevertheless, some of the species displaying poor probe-barcode hybridization remained distinguishable using the alternative barcode segment (Table 1 and Figures S1–S4).

Based on visual evaluations, the generated melt curves showed different fluorescent signatures for closely related species, such as thresher sharks (*Alopias* spp.) and hammerheads (*Sphyrna* spp.; Figure 2). Across the two species of thresher sharks, FASTFISH-ID produced visually distinguishable curves in BS1 at the initial stages of the hybridization process and produced a similar drop at ~74–79°C, while the signatures in BS2 were clearly distinct in the initial stages (about 42–47°C). Some species, on the other hand, have virtually identical BS1 signatures but are distinguishable using BS2, such as in the case of zebra shark (*Stegostoma fasciatum*) and spot-tail shark (*Carcharhinus sorrah*) (Figure 3). However, there are problematic species pairs that have highly similar signatures with both segments and therefore appear visually indistinguishable. This is the case between the tiger shark and giant shovelnose ray, between the silky and blue



	CITES status			Amplification condition		Distinguishable	
No.		Scientific name	English name	Barcode segment 1 (BS1)	Barcode segment 2 (BS2)	Visual	Deep learning
	Yes	Alopias pelagicus	Pelagic thresher	Yes	Yes	Yes	Yes
2		Alopias superciliosus	Bigeye thresher	Yes	Yes	Yes	Yes
		Carcharhinus falciformis	Silky shark	Yes	No	No	Yes
		Carcharhinus Iongimanus	Oceanic whitetip shark	No	Yes	Yes	No
		lsurus oxyrinchus	Shortfin mako shark	No	Yes	Yes	Yes*
		lsurus paucus	Longfin mako shark	Yes	Yes	Yes	Yes*
		Lamna nasus	Porbeagle shark	No	Yes	Yes	Yes
		Sphyrna lewini	Scalloped hammerhead	Yes	Yes	Yes	Yes
		Sphyrna mokarran	Great hammerhead	Yes	Yes	Yes	Yes
0		Carcharhinus brevipinna	Spinner shark	Yes	No	Yes	Yes
1		Carcharhinus sorrah	Spot-tail shark	Yes	Yes	Yes	No
2		Prionace glauca	Blue shark	Yes	No	No	Yes*
3		Anoxypristis cuspidata	Knifetooth sawfish	Yes	Yes	Yes	Yes
4		Glaucostegus typus	Giant shovelnose ray	No	No	No	No
5		Mobula birostris	Giant oceanic manta ray	Yes	Yes	No	Yes
6		Mobula mobular	Giant devil ray	Yes	Yes	No	Yes
7		Mobula tarapacana	Sicklefin devil ray	Yes	Yes	Yes	Yes
8		Pristis pristis	Largetooth sawfish	No	Yes	Yes	Yes
9		Rhina ancylostoma	Bowmouth guitarfish	Yes	Yes	Yes	Yes
0		Rhynchobatus australiae	Whitespotted guitarfish	Yes	Yes	Yes	Yes
1		Rhynchobatus laevis	Smoothnose wedgefish	No	Yes	Yes	Yes*
2		Rhynchobatus springeri	Broadnose wedgefish	Yes	Yes	Yes	Yes*
3	No	Galeocerdo cuvier	Tiger shark	No	No	No	No
4		Stegostoma fasciatum	Zebra shark	Yes	Yes	Yes	No
5		Gymnura poecilura	Longtail butterfly ray	Yes	Yes	Yes	Yes
5		Himantura imbricata	Bengal whipray	Yes	Yes	Yes	Yes
7		Neotrygon orientalis	Oriental bluespotted maskray	Yes	Yes	Yes	Yes
8		Telatrygon zugei	Pale-edged stingray	Yes	Yes	Yes	Yes
otal d	istinguishable s	species				22	23

Amplification condition denotes whether the species amplified at either or both segments (BS1 and BS2) and whether the species was distinguishable from all other species by its fluorescent signature(s) and deep learning.

Note: species with Asterix "*" mark have probability of mis-assignment by the deep learning model.

sharks, and between the giant oceanic manta and giant devil ray (two *Mobula* species), which have nearly identical signatures in both barcode segments (Figure 4). Overall, six out of 28 species were deemed visually indistinguishable, four of which are CITES-listed. We also found seven species that amplified inconsistently; shortfin mako shark (*I. oxyrinchus*), oceanic whitetip shark (*C. longimanus*), porbeagle shark (*L. nasus*), tiger shark (*G. cuvier*), largetooth sawfish (*Pristis pristis*), giant shovelnose ray (*G. typus*), and smoothnose wedgefish (*R. laevis*). It was observed that the rightmost trough in the BS1 fluorescent signature labeled "TM" corresponds to ThermaMark, an internal marker for correction of artifactual temperature variation (Figure S5). However, in BS2, some segments were amplified and unique for each of these species.

Half of the samples were highly processed products, but they still amplified well. In some of these, there were differences in the intensity of the signatures, as reflected in signature variation from BS2 of great



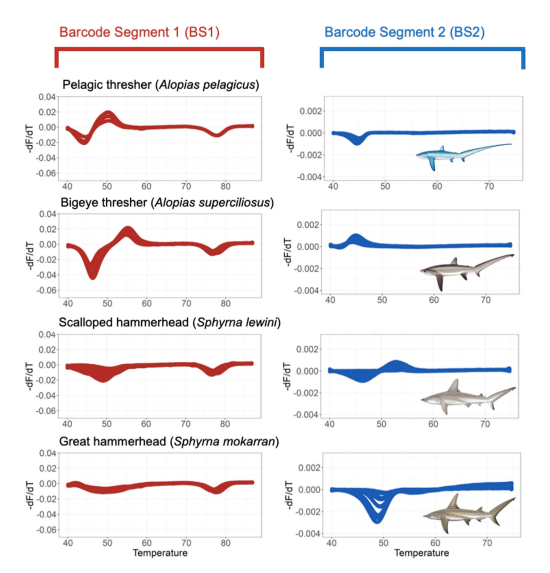


Figure 2. Some species that have visually distinguishable signatures in both barcode segments i.e., pelagic thresher, bigeye thresher, scalloped hammerhead, and great hammerhead

hammerhead, zebra shark, and bowmouth guitarfish (Figures 2, 3 and S4), which may in part be ascribed to the actual state of degradation of the original DNA template.

Machine learning for species assignment

We transposed data for the training sets and then used fluorescence values at 8,152 temperature intervals (>4,000 per each barcode segment) as variables and identified variable importance as a key feature for species assignment. We ranked variable states according to their relative importance, scaled importance and percentage of variance explained, for each barcode segment (see Table S3). We generated 301 potential deep learning models, aiming for high accuracy and minimizing error. The best deep-learning model was chosen as the one with the highest accuracy (98.20%; Table S4). When the model was applied to melt curve data from the independent specimens, accuracy dropped to 79.41%, with 54 out of 68 specimens correctly assigned (Figure 5). Misassignments were consistent with the species that also proved problematic during visual assessments, i.e., the spinner and blue shark. The model also misidentified spot-tail shark as zebra shark despite it visually having a unique signature in BS2 (Figure 3). During the testing, some samples from hammerhead sharks (*Sphyrna* spp.), smoothnose wedgefish (*R. laevis*), and broadnose wedgefish (*Rhynchobatus springeri*) were assigned to the wrong species, even though each of these species had their own unique fingerprint (Figure S1–S4).





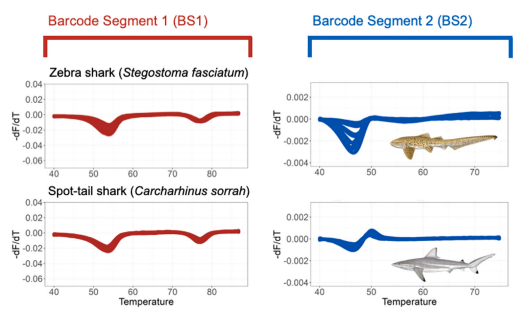


Figure 3. Some species that have similar signature in one barcode segment but visually unique in other segment i.e., zebra and spot-tail shark

DISCUSSION

Within a couple of hours and without the need to adjust the existing FASTFISH-ID assay from teleost fish to elasmobranchs, this real-time PCR method offered a portable monitoring tool that reliably enabled the identification of 25 elasmobranch species (20 of which are CITES-listed). The device used to conduct the runs, the MIC, is a convenient portable real-time PCR thermocycler weighing no more than 2 kg and allowing for the simultaneous inspection of 48 specimens per run.³¹ More importantly, the use of probes targeting mini barcodes with high inter-specific variation offers a universality that other qPCR-based assays do not currently provide, and the automatic amplification of the full COI barcode as part of the same reaction offers downstream opportunities for further in-depth screening, if necessary.

While existing genetic-based monitoring tools continue to be useful in many situations,^{24–27} FASTFISH-ID seems poised to significantly expand the horizons of DNA-based control: alongside its speed, portability, and universality, the method exhibits single nucleotide resolution³⁹ which can minimize the risk of similar fluorescent signatures, particularly when more species are added to a reference library.³¹ This is a particularly compelling argument for its implementation, as CITES lists are likely to continue to expand in the future. Additionally, the amplification of the whole COI universal barcode segment embeds a forensic dimension⁴⁰ that is not necessarily afforded by other portable tools.

A difficulty typically encountered in genetic-based trade monitoring is the handling of processed products, and this is particularly true for elasmobranchs which tend to be heavily processed in a variety of ways.^{41,42} Despite the issues of fragmented DNA due to the effect of various processing techniques,⁴³ FASTFISH-ID shows notable robustness and reliability, with 83.6% of processed samples yielding reliable melt curve profiles (51 of 61 processed samples). Since FASTFISH-ID uses real-time PCR and relies on fluorescent signatures, some species display variation in signature amplitude (the variation in peak heights and valley depths) especially when the DNA was degraded, as observed with processed products and displayed by the signature of both hammerhead species on BS2 (Figure 2). This deviation may be problematic for species assignment, especially when the assignment depends on a deep learning algorithm. The high probability of the features being similar to those of other species caused misassignments. Other issue that may have occurred is variation in the fluorescence signature from the same species. This could be due to single nucleotide polymorphisms (SNPs) within species or possibly to contamination in the case of the BS2 signature of the pale-edged stingray (*Telatrygon zugei*; Figure S4).



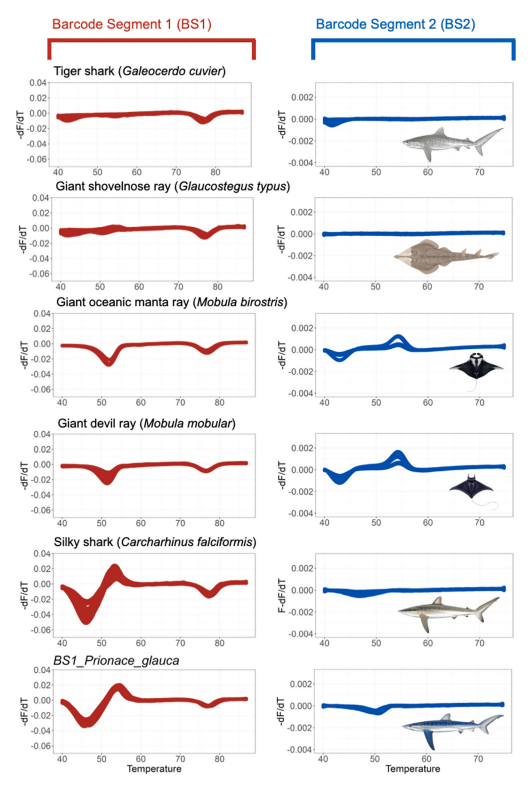


Figure 4. Problematic species that visually have highly similar signatures at both barcode segments i.e., tiger shark and giant shovelnose ray; giant oceanic manta ray and giant devil ray; silky shark and blue shark



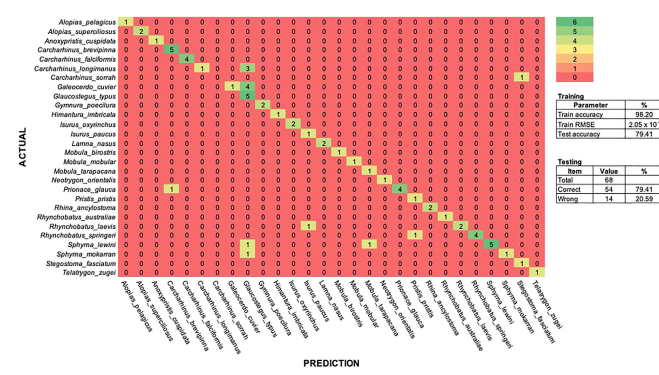


Figure 5. Confusion matrix of 28 shark and ray species assignment

Visual assessment could distinguish 22 species out of 28 with more than half of these (N = 17) being CITESlisted. Even in this preliminary phase, the method could therefore readily be applied by inspectorswithout the application of computational tools-and reliably reveal cases of illegal activities. Three pairs of species had spectral features that are difficult to distinguish, e.g., these ambiguities were present between tiger shark and giant shovelnose ray, between two species of Mobula rays (giant oceanic manta ray and giant devil ray), and between silky and blue shark (Table 1 - Visual). Thus, it must be acknowledged that the barcode segments have the same sequence of nucleotides and produced similar signatures for those species. The technology was originally designed for bony fish,³¹ and the database is currently being expanded to various important species that are globally traded as seafood. Yet, the much lower diversity of elasmobranchs (~1/30th that of teleosts) will make any effort to produce spectral reference databases a far less onerous task than that currently encountered with bony fishes. While it has been known that the COI gene is more slowly evolving in chondrichthyans than teleosts,^{44–46} this is seldom a major issue in most DNA barcoding applications,⁴⁷⁻⁴⁹ so an optimized iteration of the FASTFISH-ID method is poised to be transformational for elasmobranch conservation and management. A gualitative investigation on the full length of COI sequences (Sanger sequencing results) based on visual and simple comparison (https://www. bioinformatics.org/sms2/ident_sim.html) revealed that for those problematic three pairs of species mentioned above for that particular segment, there is a sufficient level of divergence in their sequence (70–98%) to allow discrimination via this sensitive method⁵⁰ using a modified set of probes.

In the absence of an online reference database of elasmobranch fluorescent signatures, machine learning was developed for this study. One of the machine learning applications is pattern recognition.^{38,51} Deep learning (also known as deep structured learning) is broadly applied in machine learning applications, especially pattern recognition.^{38,51} and has advantages in its flexibility to develop learning styles i.e., supervised, semi-supervised, or unsupervised.^{33,34} Deep learning models have been chosen and deployed with independent testing datasets to measure their accuracy. We found that the accuracy of our test model was 79.41%, which is lower than the training accuracy (98.20%; Table S5), and yet the model could identify similar species that could not be distinguished visually. In fact, the model enabled us to differentiate the two *Mobula* species that have similar signatures in both barcode segments. Machine learning could also recognize silky shark, a problematic species for the authorities as the species belongs to the Carcharhinidae, a diverse family that has plenty of look-alike species. In particular, the silky shark spectral profiles



appeared visually indistinguishable from blue shark. However, the new CITES listing agreed during CoP19 added all requiem sharks into Appendix II (including blue shark along with the other 53 species shark from Carcharhinidae family) will make implementing action manageable since requiem sharks make up a large proportion of the products found in the global shark fin trade hubs in China.⁵² Although international trade in all requiem sharks will now be regulated, a non-detriment finding (NDF; CITES's mechanism that allows certain species listed in Appendix II to be traded with strict quotas) which is specific to each species will still require the capability of identification at the species level.

Five out of 28 species could not be assigned accurately using the model, i.e., between spot-tail and zebra shark as well as mis-assignments among oceanic whitetip shark, tiger shark, and giant shovelnose ray (Table 1 – Deep Learning). Curiously, there were also misassignments for species that had quite unique fluorescent signatures. We argue that these misassignments could be due to variation in amplitude, where some species actually have similar signatures but different amplitudes, ⁵³ the cause of which is undetermined, but could be due to degraded DNA. For instance, the signature in BS2 of zebra shark has high-amplitude variations that may challenge the model to assign the species (Figure 3). Increasing training datasets may be required as this should improve the robustness of the model, ³³ while future re-tailoring of the barcode regions to elasmobranch variation may also remove some of the within-species noise. Despite the assignment problems, when we combine visual and deep-learning assignments, we could distinguish 25 out of 28 species, 20 of which are listed in CITES Appendix II.

Limitations of the study

The probe hybridization problems (which occurred when the barcode segments have a high degree of mismatches with the designed probes) encountered in seven species prevented the machine learning tool from adequately assigning fluorescent signatures to a given species. Since BS1 failed to hybridize for most of these species, the species assignment in these cases was solely reliant on BS2, which, in many cases also exhibited poor hybridization. To address this issue, it seems that going forward the designing of new probes tailored to elasmobranch sequence variation will be a necessary solution to increase the versatility and reliability of FASTFISH-ID, this process may also consider the evaluation of other gene regions, such as NADH2,^{45,54,55} which has proven to be highly diagnostic in elasmobranchs. An increased set of elasmobranch species may also inflate misassignments due to the higher degree of similarity among species in both visual-based and machine learning-based systems. There are also limitations in using fully supervised deep-learning approaches in the selection of important features from highly variable training sets (e.g., signatures from the two barcode segments).⁵⁶ The addition of more species to the database will require more training images. However, with such improvements, this method will help authorities (i.e., fish inspectors, customs, and guarantine officers) by providing a single, agile testing option, at any point in the supply chain, to disentangle the complexity of the shark and ray product trade, and ultimately reduce the consequential risk of extinction for these endangered and iconic taxa.

STAR*METHODS

Detailed methods are provided in the online version of this paper and include the following:

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- METHOD DETAILS
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 - FASTFISH-IDTM closed-tube barcoding protocol
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 - O Machine learning for species assignment

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.isci.2023.107065.

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AUTHOR CONTRIBUTIONS

Conceptualization (A.P.P., A.D.M., and S.M.), Funding acquisition (S.M. and J.M.M.), Methodology (A.P.P., M.C., A.D.M., and S.M.), Resources (A.D.M. and S.M.), Investigation (A.P.P. and M.C.), Formal Analysis and Visualization (A.P.P.), Project Administration (A.P.P., J.M.M., F.A., A.D.M., and S.M.), Supervision (A.D.M. and S.M.), Writing—original draft (A.P.P., A.D.M., and S.M.), Writing—review & editing (M.C., J.M.M., F.A., and E.M.).

DECLARATION OF INTERESTS

The authors declare that they have no competing interests.

INCLUSION AND DIVERSITY

We support inclusive, diverse, and equitable conduct of research.

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STAR*METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER	
Biological samples			
198 tissue samples (specimens) belonging to 28 species	This paper	https://www.ncbi.nlm.nih.gov/sra/? term=PRJNA850687	
Chemicals			
Mu-DNA extraction reagents	57	https://www.protocols.io/view/mu-dna-a- modular-universal-dna-extraction-method-a- 6qpvryj2gmkn/v2	
Critical commercial assays			
FASTFISH-ID [™] Probe Mix	Ecologenix, LLC. Natick, MA - USA	https://www.fastspecies-id.com/	
Deposited data			
Training and testing datasets	This paper	https://doi.org/10.5281/zenodo.7997300	
Software and algorithms			
H2O	H2O.ai	https://h2o.ai/platform/ai-cloud/make/h2o/	
pandas	The pandas development team	Library https://pandas.pydata.org	
Deep learning algorithm for species recognition	This paper	https://github.com/andhikaprima/FastSharkID	
Oligonucleotides			
M13F primer	Macrogen™	Ecologenix, LLC. Natick, MA - USA	
Fish02 primer sets	Macrogen™	58	
Leray-XT primer sets	Macrogen™	59	

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Andhika Prasetyo (a.p.prasetyo@edu.salford.ac.uk) and/or A. D. McDevitt (allan. mcdevitt@atu.ie).

Materials availability

This study did not generate new unique reagents. FASTFISH-ID reagents were manufactured by Ecologenix, LLC. Natick, MA - USA.

Data and code availability

- All original code is deposited at the Github repository and are publicly available of the date of publication database: https://github.com/andhikaprima/FastSharkID.
- The datasets associated with the study are provided in a dedicated Zenodo repository: https://doi.org/10.5281/zenodo.7997300.
- Any additional information required to reanalyse the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Tissue sample of shark and ray specimens were collected in several sites nested in six locations across cities on Java Island, the most populous island in Indonesia (Figure S6, namely Jakarta, Indramayu, Tegal, Cilacap, Surabaya and Banyuwangi. Collected specimens were gathered without prior knowledge of their



exact harvest location and were available for collection at a variety of sites, such as fishing ports (FP), traditional markets (TM), processing plants (PP), export hubs (EH) and an inspector station (AU).

Sample collection was granted by research permit no.251/BRSDM/II/2020 issued by Agency for Marine and Fisheries Research and Human Resources AMFRAD, the Ministry of Marine Affairs and Fisheries (MMAF), Republic of Indonesia. Research ethics no. STR1819-45 issued by the Science and Technology Research Ethics Panel, University of Salford. Export permits no. 00135/SAJI/LN/PRL/IX/2021 (CITES-listed specimens) and 127/LPSPL.2/PRL.430/X/2021 (non-CITES-listed specimens) were granted under the authority of the Ministry of Marine Affairs and Fisheries (MMAF), Republic of Indonesia. Sample were imported into the UK under import permit no. 609191/01-42 from the Animal and Plant Health Agency (APHA), United Kingdom.

METHOD DETAILS

Sample collection and DNA extraction

579 specimens were opportunistically collected at the above-mentioned sites and processing factories throughout January and February 2020. The tissue, which could either be fresh, frozen, partially or heavily processed, was then stored in 2.0mL screw-cap microcentrifuge tubes, submerged in 90% ethanol and stored at 4°C. DNA was extracted from samples following the Mu-DNA protocol for tissue samples⁵⁷ with an overnight incubation at 55°C on the thermomixer with a medium mixing frequency and a final elution volume of 100 μ L. All surfaces were sterilised with 50% bleach and then washed with 70% ethanol, in-between and after extracting each sample, to reduce cross-contamination risks (Figures S7A–S7B).

Of these, we excluded specimens of unclear taxonomy, and all species represented by less than 3 individuals. We refined the collection to 130 tissue samples (specimens) belonging to 28 species; for each species, we used three replicates per specimen as training sets (390 runs) (Table S1). We also had another 68 tissue samples without replication and used them as testing datasets (Table S2). As sampling was conducted opportunistically, we did not have an equal number of samples per species. Some species had a limited number of specimens, so we took out some training sets to be used as testing datasets. Datasets were then filtered, and ambiguous qPCR runs (i.e. poor probe-barcode hybridisation or inconsistent fluorescent signature) were removed. A poor probe-barcode hybridisation was checked using a reference point created by ThermaMarkTM (TM) in the signature produced from BS1. If only ThermaMarkTM (TM) amplified in the BS1 fluorescent signature, those runs would have failed to hybridize. Inconsistent fluorescent signatures within a replication or species were re-run a second time. If the re-runs kept failing, those runs were removed. In the end, we used 357 (number of replications varied by specimens) and 68 runs for training and testing datasets, respectively.

FASTFISH-ID[™] closed-tube barcoding protocol

PCR reaction and amplification conditions

In the first instance, the FASTFISH-IDTM method requires the amplification of the full cytochrome c oxidase I (COI) gene (~650 bp) and in the second instance, it targets the two mini-barcodes (~80 bp) using a set of probes. PCR master mixes were prepared in low-adhesion Eppendorf tubes.³¹ The major components of this method are ThermaStopTM, ThermaMarkTM and FASTFISH-IDTM Probe Mix (Ecologenix, LLC.). ThermaStopTM is a novel hot-start reagent that prevents non-specific amplification prior to the start of the reaction, while ThermaMarkTM (hereafter referred as TM) is a temperature-dependent marker for correction of melt-curve analysis (Ecologenix, LLC.). The FASTFISH-IDTM probe mix consisted of two sets of positive/negative probe pairs labelled in two different colours that hybridize along the length of two mini-barcode regions within the amplified COI target sequence, hereafter referred to as Barcoding Segment 1 (BS1) and Barcoding Segment 2 (BS2). A M13 primer was used as a priming site that facilitates the sequencing process for eventual species validation through Sanger sequencing.

FASTFISH-ID[™] uses asymmetric PCR to produce more single stranded amplicons which allow the probes to hybridize more easily.³² After amplification, mismatch tolerant positive/negative probe pairs bind to their single-stranded DNA targets. Each positive-probe is formed of a target binding sequence that is 20–35 nucleotides long and has a higher fluorescent signal when it is bound to its target sequence but a low background fluorescence when it is not. Negative-probes are only quenchers that reduce the fluorescent signal when they are bound next to their paired positive-probe. Positive/negative probe pairs can bind



to both perfectly matching strands and target sequence variants with one or more nucleotide polymorphisms. This means that they can tolerate mismatches, which is one of the most important features of this technology as a single set of reagents can be used to identify a large number of species.³¹ Target sequences that are similar but different, even if only by one nucleotide, almost always have different fluorescent signatures. Positive/negative probe sets therefore have the potential to discriminate among thousands of fish species and their variants.³¹

PCR amplification was performed on a Magnetic Induction Cycler (MIC) which is a real-time PCR thermocycler designed by Bio Molecular SystemsTM (Upper Coomera, Queensland, Australia). Thermocycling conditions were 94°C for 2 mins, 5 cycles of 94°C for 5 secs, 55°C for 20 secs, 72°C for 45 secs, then 65 cycles of 94°C for 5 secs, 70°C for 45 secs (in total: 2 hrs, 20 mins and 44 secs). Following a total of 70 amplification cycles, the reaction leads to a 10- to 20-fold excess of single-stranded DNA which is critical for probe/target hybridization in a single closed tube.^{32,60} At the completion of PCR, the temperature was decreased down to 40°C for 10 mins to enable the fluorescent probes in the FASTFISH-IDTM probe mix to hybridize to the excess single-stranded DNA. This step was followed by a melting curve analysis where the temperature was gradually increased from 40°C to 87°C at 0.1°C /secs with sequential fluorescent acquisition first in the MIC PCR Cycler's Orange Channel (suitable for detection of CalRed 610- labelled probes; max excitation: 590 nm; max emission 610 nm) and then detection in the Red Channel (suitable for detection of Quasar 670-labelled probes; max excitation: 647 nm; max emission 670 nm). The first derivative of the melt curve was then used as the fluorescent signature. Species assignment was revealed by comparing a distinct mix of Cal-Red 610 and Quasar 670 fluorescent signatures (Figures S7C–S7F). Those multiple combinations allow FASTFISH-IDTM to identify a large number of species with the same reagents.^{31,39,50}

DNA barcoding and species validation

The same single strand DNA products used to generate a fluorescent signature can also be sequenced by DNA barcoding for further investigation. The sequencing protocol uses the M13 tail sequence in the FASTFISH-IDTM FISH COI HBCts excess primer (5' CACGACGTTGTAAAACGAC 3', a modified version of the M13F primer) as a sequencing primer to generate the sequence of the excess primer strand. By design, the excess primer-strand sequence can be queried directly in the NCBI nucleotide database⁶¹ or the Barcode of Life Database⁶² for species identification. In addition, we also used Fish F2 (5' TCGACTAATCATAAAGATATCGGCAC 3') and Fish R2 (5' ACTTCAGGGTGACCGAAGAATCAGAA 3') primer sets⁵⁸ for several initial specimens for comparison with HBCts excess primer (M13). Sequencing was outsourced to Macrogen EuropeTM. Samples were prepared according to the service provider protocols (https://www.macrogen-europe.com/services/sanger-sequencing). We also added species and/or specimens after identification using a highly degenerated primer set using a high throughput barcoding (HTB) method (A.P. Prasetyo et al., *unpublished data*); Leray-XT primer sets (313 bp). This set included the primers jgHCO2198 (5' TAIACYTCIGGRTGICCRAARAAYCA 3') and mICOlintF-XT (5' GGWA CWRGWTGRACWITITAYCCYCC 3').⁵⁹

QUANTIFICATION AND STATISTICAL ANALYSIS

Machine learning for species assignment

Since the two probing barcode segments and the algorithm were developed for teleost fishes, they are not expected to maximise differentiation among the melt curves of elasmobranch species. Furthermore, the existing cloud-based reference library does not contain any elasmobranch signatures. We therefore developed our own species identification system by using machine learning using the H2O platform (Figures S7G and S7H). H2O is an open source, fast and scalable machine learning and predictive analytics platform that allows building machine learning models on big data, and improving reproducibility.⁶³ The deep learning algorithm was deployed to address the problem of species assignment by considering its capability to arrange multiple nonlinear transformations to model high-level abstractions in data. H2O's Deep Learning is based on a multi-layer feedforward artificial neural network (FANN) that is trained with a stochastic gradient descent using a backpropagation environment.⁶³ Deep learning is also advantaged by extracting the optimal input representation from raw data without user intervention.⁶⁴

The fluorescent signature datasets (BS1 and BS2) were extracted, with the species identity serving as the "response", and the transposed PCR profile temperature values being used as the predictor "variables" (each barcode fragment is recorded at about 4,000 temperature values), and fluorescent values serving as the "feature". In deep learning, "response" refers to the individual value that served as the output





(species name in our case); while "variable" refers to properties of the "response" and is evaluated through the "feature".

The performance of deep learning algorithms depends heavily on the extracted features, so it's important to choose the right group of features that best represent the input data.⁶⁵ Data filtering was conducted to exclude poor probe-barcode hybridisation or inconsistent fluorescent signature datasets and provided the best representative of the data input. Two datasets (BS1 and BS2) were then merged by specimen ID with species name used as an input to the model. Our model was divided using a 70–30 ratio of training data to validation data (i.e. 246 and 111 runs respectively) and then tested with 68 independent datasets. Default parameters of H2O's Deep Learning were optimized, with a process called "grid-search", this process tried to adjust several parameters to find the optimal "stopping criteria" (list of parameters provided on Table S6). We setup a "stopping criteria" to limit the computational load in searching for the best deep learning algorithm, which was based on random discreteness, the number of generated models, and model runtime (Table S7). The best model was chosen based on model accuracy and Root Mean Square Error (RMSE) optimization. A confusion matrix is used to visualize model accuracy.

As for other algorithms, larger databases are required to improve predictive abilities by optimizing distributed representation, activation function non-linearity, and flexible architecture depth in terms of hidden layers and nodes.⁶⁶ The main challenges in applying deep learning is overfitting due to a dominant influence on the generalization ability of a deep neural network model.⁶⁷ However, regularization methods such as Ivakhnenko's unit pruning⁶⁸ or sparsity (I₁-regularization) or weight decay (I₂-regularization) can be applied during training to combat overfitting.⁶⁹ The sparsity and weight decay were used in this study.