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1 MinION sequencing of seafood in Singapore reveals creatively labelled flatfishes, confused

2 roe, pig DNA in squid balls, and phantom crustaceans

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- 10 Abstract

Food mislabelling is a growing world-wide problem that is increasingly addressed 11 12 through the authentication of ingredients via techniques like mass spectrometry or DNAsequencing. However, traditional DNA sequencing methods are slow, expensive, and 13 14 require well-equipped laboratories. We here test whether these problems can be 15 overcome through the use of Nanopore sequencing. We sequenced 92 single and 13 mixed-species samples bought in supermarkets and restaurants in Singapore which has a 16 large and diverse seafood trade. We successfully obtained DNA barcodes for 94% and 17 18 100% of the single- and mixed-species products after correcting the numerous sequencing errors of MinION reads with a correction pipeline optimized for DNA 19 barcodes. We find comparatively low levels of clear-cut mislabelling for single-species 20 21 samples (7.6 %) while the rates are higher for mixed-species samples (38.5 %). These low 22 rates are somewhat deceptive, however, because of the widespread use of vague 23 common species names that do not allow for a precise assessment of the expected 24 ingredients. With regard to the clearly mislabelled single-species products, higher-value 25 products (e.g., prawn roe, wild-caught Atlantic salmon, halibut) are replaced with lowervalue ingredients (e.g., fish roe, Pacific salmon, arrowtooth flounder) while more serious 26 27 problems are observed for mixed-species samples. Cuttlefish and prawn balls repeatedly contained pig DNA and 100% of all mixed samples labelled as containing crustaceans 28 ('crab', 'prawn', 'lobster') only yielded fish barcodes. We conclude that there is a need 29 for more regular testing of seafood samples and suggest that due to speed and low-cost, 30 MinION would be a good instrument for this purpose. We also emphasize the need for 31 32 developing clearer labelling guidelines.

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35 1. Introduction

36 In today's globalised economy, seafood readily moves across borders. Fish caught in the

- 37 Arctic and the Antarctic is served in restaurants on the equator, while scallops, oysters, and
- 38 sea cucumbers harvested from the shores of North America satisfy the ever-increasing

39 demand of consumers in East Asia. This increased demand has also led to the expansion of

- 40 seafood farming worldwide. However, increased demand has also created incentives for
- 41 seafood fraud via mislabelling. Such fraud is particularly common for fillets and heavily
- 42 processed seafood products because they are not readily identifiable by eye (Boughattas, Le
- 43 Fur, & Karoui, 2019; Carvalho, Palhares, Drummond, & Gadanho, 2017; Di Pinto et al., 2013;
- 44 Giusti, Armani, & Sotelo, 2017; Veneza et al., 2018).

45 In recent years, seafood fraud and mislabelling have attracted much attention and the scope

- 46 of the problem has become more apparent. This is partly because new technologies have
- 47 made it easier to detect fraud. Most fraud appears driven by the desire to maximize profit
- 48 because profit margins can be significantly increased by substituting expensive and
- 49 desirable food species with less desirable and cheaper ones. For example, tilapia
- 50 (Oreochromis spp.) or pangasius (Pangasianodon hypophthalmus) are occasionally sold as
- 51 more expensive fish such as snapper or cod (Hu, Huang, Hanner, Levin, & Lu, 2018; Kappel &
- 52 Schröder, 2015; Khaksar et al., 2015; Nagalakshmi, Annam, Venkateshwarlu, Pathakota, &
- 53 Lakra, 2016; Pardo et al., 2018). Similarly, farmed Atlantic salmon (*Salmo salar*) is sold as
- 54 wild-caught Pacific salmon (*Onchorhynchus* spp.) (Cline, 2012), and farmed rainbow trout
- 55 (*Oncorhynchus mykiss*) as wild-caught brown trout (*Salmo trutta*) (Muñoz-Colmenero,
- 56 Juanes, Dopico, Martinez, & Garcia-Vazquez, 2017).

57 But seafood mislabelling is sometimes more than "just" consumer fraud. It can also affect

- 58 food safety when toxic or unpalatable species such as pufferfish or escolar enter the market
- 59 by relabelling them as palatable species (Huang et al., 2014; Lowenstein, Amato, &
- 60 Kolokotronis, 2009; Xiong et al., 2018). In addition, mislabelling frequently interferes with
- 61 the conservation of species and populations when they are sold although they are protected
- by law (Almerón-Souza et al., 2018; Marko et al., 2004; Marko, Nance, & Guynn, 2011;
- 63 Wainwright et al., 2018). Finally, an additional and underappreciated problem is that
- 64 mislabelled food may contain ingredients that violate religious rules or ethical preferences,

65 given that the consumption of some ingredients are disallowed or discouraged by specific

- 66 religions.
- 67 The number of studies examining seafood fraud have increased greatly in recent years
- 68 (Cawthorn, Baillie, & Mariani, 2018; Harris, Rosado, & Xavier, 2016; Pardo et al., 2018;
- 69 Shehata, Bourque, Steinke, Chen, & Hanner, 2019). Several methods have been developed
- that are able to identify the ingredients of commercially sold seafood. This includes
- 71 chromatographic, spectroscopic, proteomic and genetic methods. Protein-based methods
- are particularly well-established for the identification of commonly traded fish species. They
- 73 were the first molecular method for identifying ingredients of seafood products to species
- and they remain very popular in the form of mass spectroscopy (MS) which has the
- 75 advantages of being fast and comparatively low-cost (Black et al., 2017; Mazzeo & Siciliano,
- 76 2016; Stahl & Schröder, 2017; Wulff, Nielsen, Deelder, Jessen, & Palmblad, 2013). However,
- identification requires comprehensive databases of MS profiles for the traded seafood,
- 78 which are difficult to develop for rare species, heavily processed samples, and samples
- 79 consisting of mixtures of multiple species.

For these reasons, genetic methods have recently received more attention. They have high 80 accuracy and specificity (Haynes, Jimenez, Pardo, & Helyar, 2019) and benefit from the large 81 number of seafood species that have been characterized with DNA barcodes. Genetic 82 testing of seafood ingredients generally relies on the standard DNA barcode for animals; i.e., 83 an approximately 650bp long segment of the mitochondrial cytochrome oxidase I (COI) 84 gene. Reference sequences for this barcode are available for a large number of 85 commercially traded species. This has the advantage that most sequences obtained from 86 87 seafood products can be assigned to species or species-groups. In addition, mixed- and 88 heavily processed samples can still be characterized because they still contain trace amounts of DNA. However, DNA barcodes are only slowly becoming popular for food 89 authentication because of the comparatively high cost of sequencing when they are 90 91 obtained with Sanger sequencing (e.g., cost per barcode at the Canadian Centre for DNA barcoding is USD 17: http://ccdb.ca/pricing/). Furthermore, Sanger sequencing does not 92 93 allow for sequencing products that contain signals from multiple species. Fortunately, these problems can be overcome by using new sequencing methods that are often collectively 94 95 referred to as Next-Generation sequencing (NGS) or High Throughput Sequencing 96 technologies (HTS). DNA barcodes obtained on platforms such as Illumina, Ion Torrent, and 97 PacBio have been used for food authentication (Carvalho et al., 2017; Giusti et al., 2017; Xing et al., 2019), but they have several disadvantages. The equipment and maintenance 98 99 cost for Illumina and PacBio instruments are so high that these sequencers are mostly found in sequencing centres that have fairly long turnaround times for submitted samples. In 100 addition, due to the high cost of flowcells, the cost per DNA barcode is high unless 101 102 thousands of products are sequenced at a time (Ho, Foo, Yeo, & Meier, 2019; Kutty et al., 2018; Srivathsan et al., 2018; Wang, Srivathsan, Foo, Yamane, & Meier, 2018; Yeo, 103

104 Puniamoorthy, Ngiam, & Meier, 2018).

Fortunately, these issues can now be addressed with Oxford Nanopore sequencing which is 105 implemented on small and portable MinION[™] sequencers. This technology could potentially 106 have three key advantages for food authentication. Firstly, the sequencer and the flowcells 107 are sufficiently inexpensive to make them suitable for routine testing in many laboratories 108 and regulatory agencies. In addition, the cost per sample is quickly dropping because recent 109 110 advances in bioinformatic pipeline now allow for obtaining up to 3500 barcodes on a single standard flowcell (Srivathsan et al., 2018). Furthermore, even less expensive flowcells with 111 lower capacity have become available that will be suitable for processing a few hundred 112 samples. Secondly, obtaining barcodes with MinION requires minimal lab equipment and 113 the data can even be obtained under difficult field conditions ranging from hot, humid 114 115 tropical rainforest (Pomerantz et al., 2018; Schilthuizen et al., 2019) to freezing Antarctic habitats (Johnson, Zaikova, Goerlitz, Bai, & Tighe, 2017). This is why MinION is not only 116 suitable for rapid species discovery (Schilthuizen et al., 2019; Srivathsan et al., 2018, 2019) 117 but also for identifying species under challenging circumstances (Blanco et al., 2019; Parker, 118 119 Helmstetter, Devey, Wilkinson, & Papadopulos, 2017; Pomerantz et al., 2018). Lastly, MinION devices generate data within minutes of loading a flowcell and allow for data 120 collection in real-time. Given all these advantages, one may ask why MinION sequencers are 121 not the default for food authentication with DNA sequences. Presumably, the main reason is 122

the high sequencing error rate of 10-15% (Wick, Judd, & Holt, 2019), but fortunately these
errors can now be effectively corrected using a range of new bioinformatics pipelines that
are optimized for obtaining animal barcodes with nanopore sequencers (Maestri et al.,
2019; Srivathsan et al., 2018).

Currently, MinION sequencing has apparently only been used in one study addressing 127 seafood authentication (Voorhuijzen-Harink et al., 2019). It compared the accuracy of 128 MinION results with those of other high-throughput sequencing techniques and found them 129 130 to be similar. However, the study did not examine seafood products sold commercially and 131 only examined two artificially mixed samples. The study also predated recently improved bioinformatics pipelines for obtaining DNA barcodes with MinION (Srivathsan et al., 2019). 132 These limitations are here overcome by studying >100 samples of seafood sold in Singapore. 133 The data are analysed using these newly developed techniques and we analyse both single-134 135 and mixed-species samples using two different primer pairs. Our study furthermore contributes to the still very limited amount of information available on the prevalence of 136 137 seafood fraud in Southeast Asia (Labrador, Agmata, Palermo, Follante, & Pante, 2019; Sarmiento, Pereda, Ventolero, & Santos, 2018; Sultana et al., 2018; Too, Adibah, Danial 138 139 Hariz, & Siti Azizah, 2016; Tran, Nguyen, Nguyen, & Guiguen, 2018). Note that Singapore is a 140 good area for developing seafood authentication methods because it is a very large seafood market. The city state imported 129,439 tonnes of seafood in 2017 (70% being fish and 30% 141 being other seafood), while only producing 6,498 tonnes (91% fish and 9% other 142 seafood)(Agri-Food and Veterinary Authority, 2018). Average per capita consumption is an 143 144 estimated 21 kg (71% fish and 29% other seafood), which is slightly above the world average of 20.5 kg (FAO, 2018). Overall, Singapore residents obtain nearly 30% of their animal 145 protein from seafood, yet seafood products purchased in Singapore have only been included 146 in two authentication studies. The first established the identity of commercially sold 147 'snappers' in six English-speaking countries (Cawthorn et al., 2018) while the second 148 149 examined the species identity of commercially available elasmobranchs. The latter revealed that in Singapore, shark meat (Carcharhinidae) was being sold as Indian threadfin 150 (Leptomelanosoma indicum) (Wainwright et al., 2018). 151

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153 2. Materials and Methods

154 2.1 Sample collection

We obtained 105 samples of fresh and frozen seafood from 6 supermarkets (Table 2, Table 3, Table 5) and 2 seafood restaurants (Table 4) in Singapore. All samples were purchased in the first week of May 2018, and each location was visited only once. The products were divided into two categories, single-species products (e.g., frozen fillets) and mixed-species products (e.g., fish or squid balls). All samples did not undergo any cooking or processing after purchase. We tested 92 single-species products (21 from restaurants and 71 from supermarkets) and 13 mixed-species products (all from supermarkets).

162 2.2 DNA extraction and PCR

DNA extraction was conducted using an automated extraction system (Bioer Automatic 163 Nucleic Acid Purification system) using MagaBio plus Tissue Genomic DNA purification kit 164 using the manufacturer's protocols. Afterwards, we amplified two barcodes that differed in 165 length. In order to obtain full length DNA barcodes, we used a COI-3 primer cocktail 166 (C FishF1t1–C FishR1t1, (Ivanova, Zemlak, Hanner, & Hebert, 2007)), while a shorter mini-167 barcode (313 bp) was obtained using m1COlintF: 5'-GGWACWGGWTGAACWGTWTAYCCYCC-168 3' (Leray et al., 2013) and a modified jgHCO2198: 5'-TANACYTCNGGRTGNCCRAARAAYCA-3' 169 (Geller, Meyer, Parker, & Hawk, 2013). In order to multiplex a large number of samples in a 170 171 single MinION run, we adopted a tagged amplicon strategy (Meier et al. 2016) where each primer was tagged with a 13-bp unique sequence at the 5' end of the primer. Eleven 172 forward and ten reverse tagged primers allowed for the amplification of 110 products using 173 a dual-indexing strategy. For this study we used the tags developed by Srivathsan et al. 174 (2019, F: HL001-HL011, R: HL001-HL010) and the PCR conditions for all amplifications were 175 as follows: 8 µl Mastermix (CWBio), 7.84 µl molecular grade H₂O, 0.16 µl of 25mM MgCl₂, 1 176 μ l of 1 mg/ml BSA, 1 μ l of each primer, and 1 μ l of sample DNA. The PCR conditions were 5 177 min initial denaturation at 94°C followed by 35 cycles of denaturation at 94°C (1 min), 47°C 178 179 (2 min), 72°C (5 min), followed by final extension of 72°C (5 min). PCR products were pooled 180 in equal volumes for library preparation and MinION sequencing. The libraries were prepared using SQK-LSK108 kit as per instructions using 1 µg of starting DNA. The only 181 modification to the protocol recommendation by the manufacturer was the use of 1X 182 Ampure beads for clean-up instead of the customarily recommended 0.4X. Sequencing was 183 carried out using MinION R9.4.1 over 24 hours. 184

185 2.3 Bioinformatics

The nanopore reads were base-called in real-time using MinKNOW. The resulting fastq file 186 187 was converted to a fasta file and the data were processed using *miniBarcoder* (Srivathsan et 188 al., 2018, 2019). In short, the reads were split into two sets based on lengths (1) 300-600 bp and (2) >600 bp. The first read set was demultiplexed to obtain sequences corresponding to 189 the COI minibarcode while the second read set included the reads pertaining to the full-190 length barcode. For this set, we first demultiplexed the reads using one pair of primers 191 (FishF2 t1 and FishR2 t1) that were then removed from the read set. Next we used the 192 193 second pair of primers (VF2 t1- FR1d t1) for demultiplexing the remaining reads in the second set. The average coverage for two combinations was >1000 X (median 770X) with all 194 195 specimens having >10X coverage. Hence, we did not proceed to recover additional reads by demultiplexing the remaining primer combinations. 196

A bioinformatics pipeline for single-species barcodes from sets of reads developed by
Srivathsan et al. (2018, 2019) was used here. Briefly, it first obtains a "MAFFT barcode" by
aligning the reads using MAFFT (Katoh & Standley, 2013) and obtaining a majority rule
consensus with subsequent removal of gaps. These MAFFT barcodes are further corrected
using RACON (Vaser, Sovic, Nagarajan, & Sikic, 2017) to generate a second set of consensus
barcodes. The MAFFT and RACON barcodes are then corrected for indel errors based on
amino-acid translations. Lastly these barcode sets are consolidated to obtain final barcodes.

- 204 For mixed species products, we modified the bioinformatics procedures. For each sample,
- the demultiplexed reads were matched by BLAST to GenBank (e-value threshold of 1e-5).
- 206 The BLAST matches were then parsed using *readsidentifier* (Srivathsan, Sha, Vogler, &
- 207 Meier, 2015) to summarize the taxonomy using the Lowest Common Ancestor approach and
- retaining only the best scoring matches. Read sets were grouped by genus, and the
- abovementioned pipeline was used to obtain a consensus barcode for each genus specific
- read set. This approach was also applied to read sets for samples for which we failed to get
- clean barcodes using the single-species approach. This is because bacterial signals can be co-
- amplified with a seafood product, and a clean barcode sequence can only be obtained after
- 213 the removal of the bacterial reads.
- All barcode sequences were matched by BLAST to NCBI NT database and the 50 best
- 215 matches were retrieved. These were aligned with the barcode datasets using MAFFT and
- 216 queried with SpeciesIdentifier (Meier, Shiyang, Vaidya, Ng, & Hedin, 2006) to find the best
- 217 matching sequence.

218 3. Results and Discussion

219 3.1. Amplification success

The use of two different sets of primers amplifying the full-length and a mini-barcode of 220 221 313bp length allowed us to obtain sequences for 87/92 (94.5%) of the single-species and 13/13 (100%) of the mixed-species products. These barcodes were derived from 158,329 222 223 short and 91,901 long nanopore reads that were successfully demultiplexed into read sets representing the different amplicons. This overall high success rate is due to combining the 224 225 data for both amplicons. We obtained mini-barcodes for 72 and full-length barcodes for 70of the 92 single-species samples, but only 55 samples (60%) have data for both. We thus 226 227 strongly recommend the use of different primer sets in order to increase the overall success 228 rates. The usage of two different PCR reactions furthermore helps with overcoming 229 potential primer biases and allows for cross-validation. For example, one sample (FM095) was expected to contain frozen scallop but a prawn DNA barcode was obtained when using 230 the full-length primer cocktail. In contrast, the mini-barcode reads revealed the expected 231 232 scallop signal. Once this sample is excluded from the analysis, our total success rate for single-species products is 93.4%, since no other samples failed this cross validation. Note 233 that for mixed products, the success rates were higher than for single-species products. This 234 applies to both sets of primers (12 of 13 samples had at least one sequence successfully 235 236 barcoded) and was surprising because we had expected that such samples would be more 237 difficult to sequence. By matching the barcodes to publicly available reference sequences, 238 we classified seven single-species samples (7.6%: Table 2) and five mixed species samples (Table 5) as being clearly mislabelled. However, we submit that an additional seven mixed-239 species samples are borderline mislabelled and the labelling could be considered fraudulent 240 if stricter rules were applied to the equivalence of scientific and common names. 241

242 3.2. Identification of seafood samples

243 Several of the clear-cut cases of mislabelling involved flatfish for which about 40% of all 244 single-species samples were affected (3 out of 7). This includes two cases of halibut

(Hippoglossus sp.) being substituted by arrowtooth flounder (Atheresthes stomias) and one 245 sample of sole (Solea sp.) being substituted by Indian halibut (Psettodes erumei). Similar to 246 cases reported elsewhere in the literature, salmon were also targeted with two samples of 247 chum salmon (Onchorhynchus keta) being sold as wild-caught Atlantic salmon (Salmo salar). 248 We also found that one sample of capelin roe (*Mallotus villosus*) was sold as prawn roe. 249 Arguably, the most serious case of mislabelling for a multi-religious society like Singapore 250 involved pig DNA in cuttlefish and prawn balls. We initially suspected lab contamination, but 251 the same seafood brand repeatedly yielded pig DNA in five samples which were bought at 252 253 different times and places. Pig DNA was also consistently amplified by both primer sets and 254 were not found in any of the other seafood samples. This ingredient in a seafood product is a serious problem given that many consumers avoid pork for religious, ethical, or health 255 256 reasons (e.g., allergies). Fortunately, the samples were not labelled as halal or kosher, but such cases do highlight the need for regular testing of heavily processed, multi-species 257 258 seafood samples. Note that a similar case of pig DNA in seafood balls had also recently been reported from the Philippines (marketed as fish, squid, or shrimp balls). These seafood balls 259 also included chicken meat (Sarmiento et al., 2018). 260

261 In most mislabelling cases, the substituted product was less valuable than the species 262 indicated on the label. For example, halibut is a more highly valued fish compared to arrowtooth flounder, which tends to develop a soft and mushy texture when cooked 263 (Greene & Babbitt, 1990). Arrowtooth flounder is found throughout the Eastern Pacific, 264 from the Bering Sea to the coast of Baja California. Historically, it was not targeted by 265 266 commercial fisheries because it was considered unpalatable, but new technology and population declines of other species have led to the exploitation of arrowtooth flounder 267 populations (Grandin & Forrest, 2017). However, this does not change the fact that 268 arrowtooth flounder can at best be considered a 'low-value' or even 'nuisance' species 269 270 (Kasperski, 2016). Yet, it is starting to regularly show up in mislabelling studies, with recent 271 cases reported from Brazil (Carvalho, Palhares, Drummond, & Frigo, 2015) and China (Xiong 272 et al., 2016). We submit that explicit regulation is needed that requires that arrowtooth flounder be labelled as such. In addition, fast detection techniques targeting this species 273 274 should be developed.

275 Not surprisingly, other cases of mislabelling involved salmon. "Wild-caught" Atlantic salmon 276 (Salmo salar) was found to be chum salmon (Onchorhynchus keta). The latter species usually 277 commands a lower price than wild-caught king or coho salmon (O. tshawytscha; O. kisutch) (Alaska Department of Fish and Game, 2018). This is presumably due to the fact that the 278 279 commercial fishery for wild Atlantic salmon has now virtually collapsed due to significant population declines. Worldwide, the mislabelling of salmon usually involves farmed S. salar 280 labelled as wild caught Onchorhynchus sp. or less valuable species of Onchorhynchus being 281 substituted by more valuable ones (Cline, 2012; Muñoz-Colmenero et al., 2017; Warner et 282 al., 2015). It appears that Singapore's case of O. keta being labelled as "wild-caught" S. salar 283 284 is a new addition to the numerous mislabelling problems in Salmo and Onchorhynchus.

285 Many mixed-species products were labelled as 'crab', 'prawn', or 'lobster' sticks or balls. 286 Only fish were listed as ingredients in 6 out of 8 mixed-species samples while two more 287 explicitly listed shrimp meat or prawn powder in addition to fish in their ingredients. 288 However, we were unable to find any crustacean DNA in all eight samples. Fish DNA was abundant and we suspect that overall, many of these products do not include any or only 289 minuscule amounts of crustacean tissues. One additional sample, which was simply labelled 290 'crab legs' without any ingredient list and was treated as a single-species product, proved to 291 only contain fish DNA as well. One way or another, we submit that the average consumer 292 would consider extremely low proportions of crustacean protein to be unacceptable given 293 294 that the label highlights the crustacean component ('crab', 'prawn', 'lobster'). This is in 295 contrast to cuttlefish balls which usually contained cephalopods, usually from the cuttlefish genus Sepia. We suggest that this 'creative labelling' misleads consumers because the main 296 product label indicates crustacean content and the fine print needs to be examined in order 297 298 to determine that the product does not actually contain crustaceans. Note that the lack of 299 crustacean signal is not due to primer biases because we used a mini-barcode primer mix 300 that that is known to amplify a wide range of marine invertebrates; i.e., we would have expected to find crustacean DNA if it had been there. 301

302 3.3. Implications and suggestions

303 Overall, our study suggests that the level of clear-cut mislabelling of seafood products in Singapore is not particularly high when compared to results from other Southeast Asian 304 countries. Studies from Malaysia, Vietnam and the Philippines found levels of mislabelling to 305 306 be around 60% (Sarmiento et al., 2018; Sultana et al., 2018; Tran et al., 2018) with the only outlier study being by Too et al., (2016) who only detected seafood fraud in 16% of the 307 tested seafood products in Malaysia. Unfortunately, establishing a baseline for overall levels 308 of seafood mislabelling in the region is difficult because the studies are not directly 309 comparable due to differences in methodology and sampling criteria. Hence, the next step 310 311 for understanding and reducing the problem would be developing standardised sampling 312 and analysis criteria. Sampling criteria could be the sales volume of a product (e.g., highdemand species like salmon, grouper, or cod)(Anjali et al., 2019; Cline, 2012; Muñoz-313 Colmenero et al., 2017; Xiong et al., 2016) or conservation concerns (Logan, Alter, Haupt, 314 Tomalty, & Palumbi, 2008; Marín et al., 2018; Wainwright et al., 2018). Such standardised 315 sampling would allow for a direct comparison across studies and regions. They would also 316 allow for studying seafood mislabelling rates over time. 317

We would argue that the main problem with Singapore's seafood products is 'creative 318 labelling', especially for heavily processed products. This is likely due to the lack of clear 319 regulations defining which species should be included in products carrying a particular 320 321 common names. The Sale of Food Act (Cap. 283, RG 1) only states that labels need to 322 provide a name or description which is "sufficient to indicate the true nature of the food", as well as defining 'fish' as any aquatic organism commonly consumed by humans, excluding 323 324 mammals, but explicitly including crustaceans and molluscs. This rules out egregious cases 325 of mislabelling such as the use of pork in seafood products, but it allows for creative labelling. Arguably, this state of affairs is no longer in line with the expectation of today's 326 consumers who expect labels to be precise. This suggests that there may be a need for a 327 regulatory update that could follow the example set by the European Union. The EU 328

mandates that both the commercial and scientific name should be listed and that the 329 330 commercial name be taken from approved lists published by EU member countries (Regulation (EU) No 1379/2013). The implementation of these rules resulted in a drop in the 331 incidence of mislabelling of commercially sold seafood in EU supermarkets (from ca. 20% to 332 ca. 8%: (Mariani et al., 2015), while countries with less strict laws continue to have 333 mislabelling rates of about 20-30% (Carvalho et al., 2015; Hu et al., 2018; Nagalakshmi et al., 334 2016). Levels of seafood mislabelling may also drop in Singapore's supermarkets if such 335 legislation were to be enacted. Note, however, that the seafood mislabelling rates in 336 337 Europe's restaurants did not benefit from the new regulations (Christiansen, Fournier, Hellemans, & Volckaert, 2018; Horreo, Fitze, Jiménez-Valverde, Noriega, & Pelaez, 2019; 338 Pardo et al., 2018), but this may not be a major concern in Singapore where all seafood 339

340 samples obtained from restaurants were correctly labelled (N=21).

341 4. Conclusions

Our results suggest that MinION is ready for DNA-based monitoring for seafood. MinION 342 reads can be used to identify key ingredients in single- and multi-species products even if 343 they were heavily processed. We surmise that methods based on MS are likely to be the 344 345 best choice for the routine identification of single-species samples of common species, but we would argue that DNA sequencing is the most suitable tool for mixed-species samples or 346 samples of rare species lacking MS profiles. Developing better techniques for mixed-sample 347 products is particularly important because some contain ingredients that should be 348 highlighted on the labels while others appear to lack ingredients that are listed. Testing such 349 samples can now be accomplished rapidly with MinION at a reasonable cost. The barcodes 350 in our study still cost ca. USD 10 per sample, but this was an artefact of only sequencing 105 351 samples on one flowcell. The correct capacity is closer to 1000 samples (Srivathsan et al., 352 353 2019) even if two sets of primers are used. Fortunately, sequencing at smaller scales can 354 also be cost-effective because flowcells can be used multiple times. Each re-use lowers the capacity which allows for having flowcells that are suitable for experiments of different 355 sizes. In addition, small-scale projects can be carried out on new, lower-capacity flowcells 356 ("Flongle"). Overall, we would thus predict that the consumable cost of MinION barcodes 357 will be <USD1 per sample. Of course, implementing a fully developed monitoring scheme 358 would require more than just a good sequencing method. It will require well-designed 359 sampling methods, the development of explicit labelling guidelines, user-friendly 360 bioinformatics software, and experimentally determined detection levels for ingredients in 361 362 mixed-species samples.

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Table 1: DNA barcoding data obtained with MinION

620

	COI full length barcode	COI minibarcode
Number of reads demultiplexed	158,329	91,901
Coverage (Average/median/range)	1508/773/13-14,211	875/403/12-6071
Number of single-species sample barcodes	70	68
Number of mixed-species samples with	14	15
barcodes		

- Table 2. Mislabelled single-species samples of seafood products obtained from supermarkets in Singapore. Ambiguities in identification are
- 623 separated by slashes.

	Seafood sold/marketed as: (expected species)	ID of barcode (best match %)
FM027	Premium Halibut fillet (<i>Hippoglossus</i> sp.)	313 bp: Atherethes stomias (100%)
		658 bp : Atherethes stomias (100%)
FM041	Wild caught Atlantic Salmon boneless fillet (Salmo	313 bp: Oncorhynchus keta (100%)
	salar)	658 bp: Unsuccessful
FM049	Sole fillet (Soleidae sp.)	313 bp: <i>Psettodes erumei</i> (100%)
		658 bp: Unsuccessful
FM072	Premium Halibut fillet (<i>Hippoglossus</i> sp.)	313 bp: Atherethes stomias (100%)
		658 bp : Atherethes stomias /A. evermanni (100%)
FM073	Wild-caught Atlantic Salmon boneless fillet (<i>Salmo salar</i>)	313/658 bp: Oncorhynchus keta (100%)
FM074	Crab leg (Brachyura sp.)*	313 bp:
		Nemipterus mesoprion/N. randalli (100%)
		658 bp:
		Epinephelus diacanthus (99.85%); Nemipterus mesoprion/N. randalli (100%);
		Panna microdon (99.85%); Priacanthud hamrur/P. prolixus (100%); Scolopsis taenioptera (100%)

FM077	Prawn Roe (Penaeidae sp.)	313/658 bp: <i>Mallotus villosus</i> (100%)

⁶²⁴ * This product was expected to be single-species but analysis revealed multiple species (separated by semicolons)

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Table 3: Correctly labelled single-species samples of seafood products obtained from supermarkets in Singapore. Ambiguities in identification
 are separated by slashes.

	Seafood sold/marketed as: (expected species)	ID of barcode (best match %)
FM004	White Fish (Oreochromis sp.)	313 bp: Oreochromis niloticus (100%)
		658 bp: Oreochromis aureus/mossambicus/niloticus (100%)
FM005	Red Spot Emperor (Lethrinus lentjan)	313 bp: Lethrinus lentjan (100%)
		658 bp: Unsuccessful
FM006	Barred Spanish Mackerel (Scomberomorus	313 bp: <i>Scomberomorus commerson</i> (100%)
	commerson)	658 bp: Unsuccessful
FM008	Salmon Fish tail (Salmo salar / Onchorhynchus sp.)	313/658 bp: Salmo salar (99.68%/99.85%)
FM009	Arrowtooth Flounder (Atheresthes stomias)	313/658 bp: Atheresthes evermanni/Atheresthes stomias (100%)
FM011	Red Snapper (<i>Lutjanus</i> sp.)	313 bp: Lutjanus madras/ophoyusenii/xanthopinnis (100%)
		658 bp: Lutjanus madras/ophoyusenii (99.85%)
FM012	Scallop (Pectenidae sp.)	313/658 bp: Unsuccessful
FM013	White Snapper (Lutjanidae sp.)	313/658 bp: <i>Pristipomoides multidens</i> (100%/ 99.85%)
FM014	Barramundi (Lates calcarifer)	313/658 bp: Lates calcifer (100%)
FM015	Oyster Meat (Ostreidae)	313/658 bp: Unsuccessful
FM017	Grouper (Epinephelus sp. / Plectropomus sp.)	313 bp: Epinephelus sexfasciatus (100%)
		658 bp: Unsuccessful

FM018	Crab Meat (Brachyura sp.)	313 bp: Monomia gladiator (99.36%)
		658 bp: Unsuccessful
FM019	Toman (Channa micropeltes)	313/658 bp: Channa micropeltes (99.36%/100%)
FM020	Barramundi (<i>Lates calcarifer</i>)	313 bp: Unsuccessful
		658 bp : Lates calcifer (100%)
FM021	Norwegian Saba Mackerel (Scombridae)	313/658 bp: <i>Scomber scombrus</i> (100%)
FM022	Blue Swimmer Crab (Portunus pelagicus)	313 bp: <i>Portunus pelagicus/reticulatus</i> (100%)
		658 bp: Unsuccessful
FM023	Wild caught Alaska Cod (<i>Gadus</i> sp.)	313 bp: Gadus macrocephalus/ogac/morhua (100%)
		658 bp : Gadus macrocephalus/ogac (100%)
FM024	Grouper (Epinephelus sp. / Plectropomus sp.)	313/658 bp: Epinephelus areolatus (100%)
FM025	Alaskan pollock fillet (Gadus chalcogrammus)	313/658 bp: Gadus chalcogrammus/Theragra finnmarchica
		(99.68%/99.85%)
FM026	Crab leg meat (Brachyura sp.)	313/658 bp:
		Monomia gladiator (98.74%/99.26 %)
FM028	Threadfin head-bone (Polynemidae sp.)	313/658 bp: Leptomelanosoma indicum (100%)
FM029	Wild pacific sole (Soleidae sp.)	313/658 bp: Leptoidopsetta polyxstra (100%)
FM030	Batang fillet slice (Scomberomorus sp.)	313/658 bp: Scomberomorus commerson (99.68%/99.85%)
FM031	Toman (Channa micropeltes)	313/658 bp: Channa micropeltes (100%)
FM032	Cod morue (<i>Gadus morhua</i>)	313 bp: Gadus macrocephalus/ogac/morhua (99.68%)
		658 bp : Gadus macrocephalus/ogac (100%)
FM033	Haddock (Melanogrammus aeglefinus)	313/658 bp: <i>Melanogrammus aeglefinus</i> (100%)
FM034	Tilapia fillet (Oreochromis sp.)	313 bp: Unsuccessful
		658 bp: Oreochromis aureus/mossambicus/niloticus (100%)
FM035	Toman fillet slice (Channa micropeltes)	313 bp: Unsuccessful
		658 bp : Channa micropeltes (100%)

Silver fish	313 bp: Protosalanx hyalocranius (100%)
	658 bp : <i>Protosalanx chinensis/hyalocranius</i> (100%)
Tuna in salt (<i>Thunnus</i> sp.)	313 bp: Thunnus atlanticus/albacres /tonggol (100%)
	658 bp: Unsuccessful
Toman slice (Channa micropeltes)	313 bp: Unsuccessful
	658 bp: Channa micropeltes (99.54%)
Wild cod morue (Gadus morhua)	313 bp: Gadus macrocephalus/ogac/morhua (100%)
	658 bp: Gadus macrocephalus/ogac (99.85%)
Bay scallop meat (Argepecten irradias)	313 bp: Argopecten irradias (100%)
	658 bp: Unsuccessful
Frozen red snapper fillet (<i>Lutjanus</i> sp.)	313/658 bp: Lutjanus vitta (99.82%)
Frozen grouper fillet (<i>Epinephelus</i> sp. / <i>Plectropomus</i>	313 bp: Unsuccessful
sp.)	658 bp : Epinephelus areolatus (97.58%)
Grouper fillet (Epinephelus sp. / Plectropomus sp.)	313/658 bp: Epinephelus areolatus (100%)
Red snapper fillet (<i>Lutjanus</i> sp.)	313 bp: Lutjanus malabaricus (99.35%)
	658 bp: Unsuccessful
Threadfin fillet (Polynemidae sp.)	313/658 bp: Leptomelanosoma indicum (100%)
White Fish fillet (<i>Oreochromis</i> sp.)	313 bp: Oreochromis niloticus (100%)
	658 bp : Oreochromis aureus/mossambicus/niloticus (100%)
Batang steak (Scomberomorus sp.)	313/658 bp: Scomberomorus commerson (100%)
Threadfin meat (Polynemidae sp.)	313/658 bp: Leptomelanosoma indicum (100%)
Salmon fish meat (seafood from Norway) (Salmo salar)	313/658 bp: Salmo salar (100%/99.85%)
Pike conger eel (<i>Muraenesox</i> sp.)	313 bp: Unsuccessful
	658 bp: Muraenesox bagio (86.81%)
Sea bass meat (<i>Lates calcarifer</i>)	313 bp: Unsuccessful
	658 bp : Lates calcifer (99.85%)
Tasmanian salmon fillet (Salmo salar)	313 bp: Salmo salar (100%)
	Tuna in salt (Thunnus sp.)Toman slice (Channa micropeltes)Wild cod morue (Gadus morhua)Bay scallop meat (Argepecten irradias)Frozen red snapper fillet (Lutjanus sp.)Frozen grouper fillet (Epinephelus sp. / Plectropomus sp.)Grouper fillet (Epinephelus sp. / Plectropomus sp.)Red snapper fillet (Lutjanus sp.)Threadfin fillet (Polynemidae sp.)White Fish fillet (Oreochromis sp.)Batang steak (Scomberomorus sp.)Threadfin meat (Polynemidae sp.)Salmon fish meat (seafood from Norway) (Salmo salar)Pike conger eel (Muraenesox sp.)Sea bass meat (Lates calcarifer)

		658 bp: Unsuccessful
FM071	Batang mid cut (Scomberomorus sp.)	313/658 bp: Scomberomorus commerson (100%)
FM075	Seasoned jellyfish	313/658 bp: Unsuccessful
FM076	Flying Fish roe (Exocoetidae sp.)	313 bp: Cheilopogon pitcairnensis (98.36%) 658 bp : Cheilopogon heterurus (98.05%)
FM080	Frozen tuna portion (<i>Thunnus</i> sp.)	313 bp: Unsuccessful658 bp: Thunnus tonggol (99.85%)
FM081	Deep sea flounder skin on fillet (Pleuronectoidei sp.)	313/658 bp: Atheresthes stomias (100%/99.68%)
FM082	Frozen snakehead fillet (<i>Channa striata / Channa micropeltes</i>)	313 bp: Unsuccessful 658 bp : Channa micropeltes (100%)
FM088	Norwegian Salmon (Salmo salar / Onchorhynchus sp.)	313/658 bp: Salmo salar (100%)
FM089	Toman fillet slice (Channa micropeltes)	313/658 bp: Channa micropeltes (99.68%/99.83%)
FM096	Prawn meat (Penaeidae sp.)	313 bp: <i>Metapenaeopsis barbata</i> (98.72%) 658 bp : Unsuccessful
FM097	Red snapper fillet with skin on (Lutjanus sp.)	313/658 bp: Lutjanus vitta (96.8%/99.69%)
FM098	Grouper fillet with skin on (<i>Epinephelus</i> sp. / <i>Plectropomus</i> sp.)	313 bp: <i>Epinephelus sexfasciatus</i> (100%) 658 bp: Unsuccessful
FM099	Frozen prepared Squid (Cephalopoda sp.)	313/658 bp: Unsuccessful
FM100	Red grouper slices (<i>Plectropomus s</i> p.)	313 bp: <i>Plectropomus leopardus</i> (100%) 658 bp : Unsuccessful
FM101	Toman slices (Channa micropeltes)	313 bp: Unsuccessful 658 bp : Channa micropeltes (99.85%)
FM102	Frozen grouper fillet (<i>Epinephelus</i> sp. / <i>Plectropomus</i> sp.)	313 bp: <i>Epinephelus sexfasciatus</i> (100%) 658 bp: Unsuccessful
FM103	Alaska cod (<i>Gadus</i> sp.)	313 bp: Gadus macrocephalus/ogac/morhua (100%) 658 bp : Gadus macrocephalus/ogac (100%)

FM104	Wild cod morue (Gadus morhua)	313 bp: Unsuccessful
		658 bp: Gadus macrocephalus/ogac (99.85%)
FM105	Frozen bay scallop meat (Argopecten irradias)	313 bp: Argopecten irradias (100%)
		658 bp: Unsuccessful
		Ambiguous
FM095	Frozen scallop (Pectinidae sp.)	313 bp: 3/5 reads match to <i>Pecten</i> sp- thus this is likely to be correct
		658 bp: Litopenaeus vannamei/Penaeus vannamei (100%)

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Table 4: Single-species samples of seafood products obtained from restaurants in Singapore. Ambiguities in identification are separated by

631 slashes.

	Seafood sold/marketed as: (expected species)	ID of barcode (best match %)
FM058	Eel Sushi Roll (<i>Anguilla</i> sp.)	313 bp: Unsuccessful
		658 bp : Anguilla anguilla (100%)
FM059	Salmon Roll (Salmo salar / Onchorhynchus sp.)	313/658 bp: Onchorhynchus keta (100%/99.84%)
FM060	Scallop (Pectenidae sp.)	313/658 bp: Unsuccessful
FM061	Red Sea Bream/Amberjack (Pagrus sp. / Pagellus sp. /	313 bp: Pagrus major/auratus (100%)
	Seriola sp.)	658 bp: Unsuccessful
FM062	Yellow Fin Tuna (Thunnus albacares)	313 bp: Unsuccessful
		658 bp : Thunnus albacares (99.85%)
FM063	Red Sea Bream/Amberjack (<i>Pagrus</i> sp. / <i>Pagellus</i> sp. / <i>Seriola</i> sp.)	313/658 bp: Seriola dumerili (100%/99.85%)
FM064	Northern Spot Prawn (<i>Pandalus platyceros</i>)	313 bp: Unsuccessful
		658 bp : Pandalus borealis (100%)
FM065	Salmon (Salmo salar / Onchorhynchus sp.)	313/658 bp: Salmo salar (100%/99.69%)
FM066	Eel Sushi (Anguilla sp.)	313/658 bp: Anguilla japonica/anguilla (100%/99.85%)
FM067	Codroe with Kelp (Gadus morhua)	313 bp: Gadus macrocephalus/ogac/morhua (100%)

		658 bp: Gadus macrocephalus/ogac (99.85%)
FM068	Yellowfin tuna (Thunnus albacares)	313 bp: Thunnus atlanticus/albacres /tonggol (100%)
		658 bp : Thunnus albacares (100%)
FM069	Golden Cuttlefish (Sepia esculenta/ Sepia elliptica)	313 bp: Sepia prashadi/pharaonis/ramani (99.68%)
		658 bp : Sepia pharaonis (99.84%)
FM084	Salmon (Salmo salar / Onchorhynchus sp.)	313/658 bp: Salmo salar (100%/99.69%)
FM085	Swordfish (Xiphias gladius)	313/658 bp: <i>Xiphias gladius</i> (100%)
FM086	Bluefin tuna/Skipjack Tuna (Thunnus sp. / Katsuwonus	313 bp: Thunnus obesus/atlanticus/albacares (100%)
	pelamis)	658 bp : Thunnus obesus (99.85%)
FM087	Bluefin tuna/Skipjack Tuna (Thunnus sp. / Katsuwonus	313/658 bp: Katsuwonus pelamis (100%/99.85%)
	pelamis)	
FM090	Tara kirimi (cod fish fillet) (Gadus sp.)	313/658 bp: Gadus macrocephalus/ogac (100%)
FM091	Kajiki kirimi (swordfish fillet) (Xiphias gladius)	313/658 bp: <i>Xiphias gladius</i> (100%)
FM092	Gindara kirimi (black cod fillet) (Anoplopoma fimbria)	313/658 bp: Anoplopoma fimbria (100%)
FM093	Buri kirimi (Yellowtail fillet) (Seriola quinqueradiata)	313 bp: Unsuccessful
		658 bp : Seriola lalandi/quinqueradiata (100% - 99.66%)
FM094	Seasoned cod roe (Gadus morhua)	313/658 bp: Gadus chalcogrammus/Theragra finnmarchica
		(100%/100% - 99.23%)

Table 5: Mixed-species samples obtained from supermarkets in Singapore. Ambiguities in identification are separated by slashes.

Seafoo	d sold/marketed as: (expected species/ingredients listed)	ID of barcodes (best match %)		
Pig DNA detected				
FM001	Prawn Ball (Penaeidae sp. unknown fish, Sepiida)	313 bp:Sepia pharaonis/prashadi/ramani (100%)Sus scrofa (99.68%)Trichiurus gangeticus/lepturus (100%)658 bp:Lutjanus lutjanus (99.85%)Nemipterus mesoprion/randalli (100%)Priacanthus macracanthus (100%)Sepia pharaonis (99.85%)Sus scrofa (100%)		
FM002	Cuttlefish Ball (unknown fish, Sepiida)	313 bp: Sus scrofa (100%) Sepia prashadi/pharaonis/ramani (98.74%) Trichiurus gangeticus/lepturus (99.68%) 658 bp: Nemipterus mesoprion/randalli (99.71%) Sus scrofa (100%) Sepia pharaonis/ramani (99.87%)		
FM016	Cuttlefish Ball (Unknown fish, Sepiida)	313 bp: Nemipterus mesoprion/randalli (100%) Sepiella japonica (99.68%) 658 bp: Nemipterus mesoprion/randalli (100%) Sepia pharaonis/ramani (99.85%) Sus scrofa (99.85%)		
FM078	Cuttlefish Balls (Unknown fish, Sepiida)	313 bp:		

		Sepia prashadi/pharaonis/ramani (100%) Sus scrofa (100%) Sepiella japonica (100%) 658 bp: Sepia pharaonis/ramani (99.87%) Nemipterus mesoprion/randalli (100%) Sus scrofa (99.71%)
FM079	Prawn Balls (Unknown fish, Sepiida, Penaeidae)	313 bp: Nemipterus mesoprion/randalli (100%) Sepiella inermis (100%) 658 bp: Nemipterus mesoprion/randalli (100%) Sepia pharaonis (100%) Sus scrofa (99.71%) Priacanthus tayenus (99.79%)
	Key ingredi	ent not present
FM003	Crab Stick (Brachyura, sp. unknown fish)	313 bp: Nemipterus mesoprion/randalli (99.68%) Scolopsis taenioptera (99.68%) 658 bp: Dussumieria acuta (100%) Nemipterus mesoprion/randalli (100%) Pentapodus setosus (100%) Priacanthus macracanthus (99.87%) Upeneus margarthae (99.75)
FM007	Flavoured Crab Ball (Brachyura sp., Surimi, Threadfin Bream (<i>Nemipterus virgatus</i>)	313 bp: Unsuccessful 658 bp: <i>Atule mate</i> (100%)

FM010	Breaded Fish Fingers (Threadfin Bream (<i>Nemipterus</i>	Decapterus maruadsi (100%)Jaydia striata/trunctata (100%)Parastromateus niger (99.85%)Liza parsia/melinoptera (100%)Psettodes erumei (100%)Scolopsis taenioptera (100%)Scomberomorus commerson (100%)Sphyraena flavicauda (99.54%)Lutjanus lutjanus (100%)Saurida macrolepis/tumbil (99.75%)Siganus fuscescens/canaliculatus (100%)Trichiurus lepturus (100%)Upeneus sulphureus (99.85%)313 bp:
FINIOLO	japonicus)	SIS bp: Sardinella jussieu (99.33%) Selar crumenophthalmus (100%) 658 bp: Priacanthus hamrur/prolixis/tayenus (100%)
FM039	Lobster Ball (Decapod sp., Unknown fish)	313 bp: Nemipterus furcosus (99.04%) Priacanthus macracanthus (99.68%) Melanogrammus aeglefinus (100%) 658 bp: Scolopsis taenioptera (100%) Nemipterus marginatus (99.74%) Priacanthus macracanthus (99.87%)
FM040	Crab Stick (Brachyura sp., Unknown fish)	313 bp: Nemipterus mesoprion/randalli (100%)

		Upeneus sulphureus (98.71%)
		658 bp:
		Nemipterus mesoprion/randalli (100%)/
		Priacanthus hamrur/prolixus (99.87%)
		Scolopsis taenioptera (100%)
		Upeneus sulphureus (98.90%)
FM047	Crab Leg (Brachyura sp., Threadfin bream (Nemipterus sp.,	313 bp:
	Alaska Pollock (Gadus chalcogrammus)	Gadus chalcogrammus/ Theragra finnmarchica (100%)
		Nemipterus mesoprion/randalli (100%)
		658 bp:
		Gadus chalcogrammus/ Theragra finnmarchica (100%)
		Nemipterus mesoprion/randalli (100%)
FM083	Crab leg (Unknown fish, Brachyura sp.)	313 bp:
		Priacanthus hamrur/prolixus (100%)
		658 bp:
		Unsuccessful
		Clean
FM042	Cuttlefish Ball (Unknown fish, Sepiida)	313 bp:
		Scolopsis taenioptera (100%)
		Sepia pharaonis (99.68%)
		Trichiurus gangeticus/lepturus (100%)
		658 bp:
		Lepturacanthus salava (100%)
		Priacanthus macracanthus (100%)
		Upeneus vittatus/supravittatus (100%)