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## Unveiling hákarl: A study of the microbiota of the traditional Icelandic fermented fish

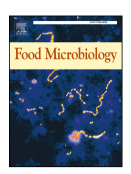
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Andrea Osimani, Ilario Ferrocino, Monica Agnolucci, Luca Cocolin, Manuela Giovannetti, Caterina Cristani, Michela Palla, Vesna Milanovic, Andrea Roncolini, Riccardo Sabbatini, Cristiana Garofalo, Francesca Clementi, Federica Cardinali, Annalisa Petruzzelli, Claudia Gabucci, Franco Tonucci, Lucia Aguilanti



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1	Unveiling hákarl: a study of the microbiota of the traditional Icelandic fermented fish
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#### Abstract

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Hákarl is produced by curing of the Greenland shark (Somniosus microcephalus) flesh, which before fermentation is toxic due to the high content of trimethylamine (TMA) or trimethylamine N-oxide (TMAO). Despite its long history of consumption, little knowledge is available on the microbial consortia involved in the fermentation of this fish. In the present study, a polyphasic approach based on both culturing and DNA-based techniques was adopted to gain insight into the microbial species present in ready-to-eat hákarl. To this aim, samples of ready-to-eat hákarl were subjected to viable counting on different selective growth media. The DNA directly extracted from the samples was further subjected to Polymerase Chain Reaction-Denaturing Gradient Gel Electrophoresis (PCR-DGGE) and 16S ampliconbased sequencing. Moreover, the presence of Shiga toxin-producing Escherichia coli (STEC) and Pseudomonas aeruginosa was assessed via qualitative real-time PCR assays. pH values measured in the analyzed samples ranged from between 8.07±0.06 and 8.76±0.00. Viable counts revealed the presence of total mesophilic aerobes, lactic acid bacteria and Pseudomonadaceae. Regarding bacteria, PCR-DGGE analysis highlighted the dominance of close relatives of Tissierella creatinophila. For amplicon sequencing, the main operational taxonomic units (OTUs) shared among the data set were Tissierella, Pseudomonas, Oceanobacillus, Abyssivirga and Lactococcus. The presence of Pseudomonas in the analyzed samples supports the hypothesis of a possible role of this microorganism on the detoxification of shark meat from TMAO or TMA during fermentation. Several minor OTUs (<1%) were also detected, including Alkalibacterium, Staphylococcus, Proteiniclasticum, Acinetobacter, Erysipelothrix, Anaerobacillus, Ochrobactrum, Listeria and Photobacterium. Analysis of the yeast and filamentous fungi community composition by PCR-DGGE revealed the presence of close relatives of Candida tropicalis, C. glabrata, C. parapsilosis, C. zeylanoides, Saccharomyces cerevisiae, Debaryomyces, Torulaspora, Yamadazyma, Sporobolomyces, Alternaria, Cladosporium tenuissimum, Moristroma quercinum and Phoma/Epicoccum, and some of these species probably play key roles in the development of the sensory qualities of the end product. Finally, qualitative real-time PCR assays revealed the absence of STEC and *Pseudomonas aeruginosa* in all of the analyzed samples.

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Keywords: Tissierella; Pseudomonas; Debaryomyces; 16S amplicon-based sequencing; PCR-DGGE.

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60 Fermentation represents one of the most ancient techniques for food preservation. Traces of this practice can be seen as 61 far back as 6000 B.C. in the Fertile Crescent (Franco et al. 2016). Moreover, the production of fermented foods became 62 very popular among the Egyptian, Greek and Roman civilizations (Huang, 2016). 63 Food fermentation is mainly based on the metabolic activities of microorganisms that are either naturally present in the 64 raw materials or artificially inoculated (Shiferaw Terefe, 2016). The most well-known processes include lactic acid 65 fermentation, fungal fermentation, and alkaline fermentation, where pro-technological microorganisms improve the 66 aroma, flavor, texture, and nutritional characteristics of the raw materials and inhibit spoilage and pathogenic 67 microorganisms (Shiferaw Terefe, 2016). Moreover, species belonging to some microbial groups mostly associated 68 with food fermentation (e.g., lactic acid bacteria) can reduce the health hazards associated with the consumption of food 69 containing some toxic substances (Luz et al., 2018). In this context, microorganisms can be considered human beings' 70 coevolutionary partners responsible for providing a wide variety of fermented foods with enhanced nutritional and 71 sensory characteristics. 72 Most fermented foods available on the market are still produced in accordance with ancient traditions deeply rooted in 73 the territory of origin. The obtained products represent an invaluable source of microbial diversity where complex 74 microbial populations coexist in a dynamic equilibrium. 75 The most popular fermented foods are produced with raw materials from the dairy, meat or vegetable food chains. For 76 such products, an ample scientific literature on both manufacturing technologies and the microbial communities 77 involved during their transformation is available. Regarding fermented foods produced with raw materials from the 78 marine environment, a lack of knowledge on the technological processes and the relevant microbial populations is 79 highlighted (Rajauria et al., 2016). Notwithstanding, fermented marine-based products are currently consumed by 80 several cultural groups worldwide (Rajauria et al., 2016). 81 Brilliant examples of traditional fermented fish products are represented by surströomming and rakfisk, produced in 82 Sweden and Norway, respectively, and hákarl, produced in Iceland (Skåra et al., 2015). The production of such 83 delicacies dates back to the Viking Age, when preservation of foodstuffs with salt was expensive, especially in the 84 remote regions of northern Europe. Therefore, instead of salting, new empiric methods of preservation of caught fish 85 were carried out by local populations, thus leading to the production of edible and safe products. 86 Among the abovementioned fermented fish, hákarl is produced by curing of the Greenland shark (Somniosus 87 microcephalus). As reported by Skåra et al. (2015), the origin of the production technique of hákarl is still not clear,

and it is unknown whether the shark was specifically caught or simply collected from specimens that drifted ashore.

89	The consumption of fresh Greenland shark is considered unsafe, although the toxic substances responsible of poisoning
90	have not been recognized. Different authors have reported cases of poisoning from the flesh of the Greenland shark
91	likely due to a high level of trimethylamine (TMA) (Anthoni et al., 1991; Halsted, 1962; Simidu, 1961).
92	In ancient times, hákarl was produced by cutting the shark into pieces that were left to ferment for weeks or months in
93	gravel pits often close to the sea. The pits were usually covered with stones, seaweed, or turf. These structures were
94	constantly exposed to seawater, which flooded over the fish at high tide (Skåra et al., 2015).
95	In the modern era, the fermentation of shark pieces is carried out in closed containers that allow the resulting leachate to
96	be drained. Such a process can last from 3 to 6 weeks depending on the environmental temperature and season. After
97	fermentation, the shark pieces are further cut and hung to dry in dedicated sheds for weeks or months, depending on the
98	outdoor environmental conditions (Skåra et al., 2015).
99	In both the ancient or the modern processes, the metabolic activities of microorganisms occurring during shark
100	fermentation lead to the conversion of a poisonous raw material into a safe and tasty ready-to-eat food product with a
101	long shelf-life. The hákarl is characterized by a soft texture with a whitish cheese-like appearance, strong ammonia
102	smell and fishy taste (Skåra et al., 2015).
103	Despite the long history of hákarl consumption, a lack of knowledge is available on the microbial consortia involved in
104	the shark fermentation. Indeed, to our knowledge, only one study that dates back to 1984 is available in the scientific
105	literature (Magnússon and Gudbjörnsdottir, 1984).
106	Since many years ago, the cultivation of microorganisms on synthetic growth media was the primary way to study
107	microbial communities in foods. The development of molecular techniques based on the use polymerase chain reaction
108	(PCR) opened new frontiers for the study of microbial ecology in complex matrices (Garofalo et al., 2017). A variety of
109	studies have shown that combinations of different microbiological techniques can provide sound information on the
110	microbial composition of complex food matrices, including those subjected to fermentation. Among the most adopted
111	and sensitive molecular methods, PCR-Denaturing Gradient Gel Electrophoresis (DGGE), real-time PCR and next-
112	generation sequencing provide reliable data for microbiological profiling of foods.
113	Based on these concepts, a polyphasic approach based on both culture and DNA-based techniques was adopted to
114	provide insight into the microbial species present in ready-to-eat hákarl.
115	To this end, samples of ready-to-eat hákarl were subjected to viable counting on different selective growth media. The
116	DNA directly extracted from the samples was further subjected to PCR-DGGE and Illumina sequencing. Moreover, the
117	presence of Shiga toxin-producing E. coli (STEC) and Pseudomonas aeruginosa was assessed via qualitative real-time
118	PCR assays.

2. Materials and methods
2.1. Sampling
Ten samples of ready-to-eat <i>hákarl</i> (Figure 1) codified from H1 to H10 were analyzed, for each sample two 100 g boxes
were purchased (for a total of 20 analyzed boxes). The samples were purchased via the internet from a dealer located in
Iceland. In more detail, the samples were collected through different orders placed from January to May 2018. The
samples were shipped by international express courier in plastic boxes at room temperature in aerobic conditions and
analyzed after 24 hours from shipping. No further information on the samples was provided by the producer.
2.2. pH measurements
pH values of the hákarl samples were determined with a pH meter equipped with an HI2031 solid electrode (Hanna
Instruments, Padova, Italy). For each sample, the measurements were performed in duplicate.
2.3. Microbial viable counts
Twenty-five grams of each hákarl sample were homogenized for 5 min at 260 rpm in 225 mL of sterile peptone water
(bacteriological peptone 1 g L <sup>-1</sup> , Oxoid, Basingstoke, UK) using a Stomacher 400 Circulator apparatus (VWR
International PBI, Milan, Italy). The obtained suspensions were diluted 10-fold and subjected to microbial counts of
total mesophilic aerobes, lactic acid bacteria, Pseudomonadaceae, Enterobacteriaceae and eumycetes. Briefly, total
mesophilic aerobes were counted as reported by Osimani et al. (2011); presumptive mesophilic lactobacilli and
lactococci were enumerated in De Man, Rogosa and Sharpe (MRS) agar medium incubated at 30 °C for 48 h and M17
agar medium incubated at 22 °C for 48 h, respectively as previously described (Aquilanti et al., 2013). The enumeration
of Pseudomonadaceae was carried out using Pseudomonas Agar Base (PAB) with cetrimide-fucidin-cephalosporin
(CFC) selective supplement (VWR International, Milan, Italy), incubated at 30 °C for 24–48 h (Garofalo et al., 2017),
whereas Enterobacteriaceae were counted in Violet Red Bile Glucose Agar (VRBGA) incubated at 37 °C for 24 h
(Garofalo et al., 2017). Finally, the enumeration of eumycetes was carried out on Wallerstein Laboratory Nutrient
(WLN) agar medium supplemented with chloramphenicol (0.1 g/L) to inhibit the growth of bacteria and incubated at 25

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°C for 72 h (Taccari et al., 2016).

A miniVIDAS apparatus (Biomerieux, Marcy l'Etoile, France) was used to assess the presence of *Listeria* monocytogenes through the enzyme-linked fluorescent assay (ELFA) method in accordance with the AFNOR BIO 12/11-03/04 validated protocol (Aquilanti et al., 2007).

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2.4. DNA extraction from hákarl samples

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Aliquots (1.5 mL) of each homogenate (dilution 10<sup>-1</sup>) prepared as described above were centrifuged for 5 min at 16000 g, and the supernatants were discarded. The cell pellets were then used for the extraction of total microbial DNA using an E.Z.N.A. soil DNA kit (Omega bio-tek, Norcross, GA, USA) following the manufacturer's instructions. A Nanodrop ND 1000 (Thermo Fisher Scientific, Wilmington, DE, USA) was used to measure the quantity and purity of the extracted DNAs, which were then standardized to a concentration of 25 ng  $\mu$ L<sup>-1</sup> for further analysis. DNA extracts obtained from the *hákarl* from each of the two boxes representing one sample (H1-H10) were then pooled and subjected to *PCR-DGGE* analyses and 16S rRNA gene amplicon target sequencing (Milanović et al., 2018).

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2.5. PCR-DGGE analysis of bacteria

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166 The extracted DNA was first amplified by PCR in a My Cycler Thermal Cycler (BioRad Laboratories, Hercules, CA, 167 USA) using the universal prokaryotic primers 27F and 1495R described by Weisburg et al. (1991) for the amplification 168 of 16S rRNA gene. In detail, 2 µL (approximately 50 ng) of DNA from each sample was amplified in a 25 µL reaction 169 volume composed of 0.5 U of Taq DNA polymerase (Sibenzyme, Novosibirsk, Russia), 1X reaction buffer, 0.2 mM 170 dNTPs and 0.2 μM of each primer using the cycling program described by Osimani et al. (2015). The PCR products 171 were checked by routine electrophoresis on 1.5% agarose (w/v) gels and then purified using the Illustra GFX PCR DNA 172 and Gel Band Purification Kit (GE Healthcare Life Sciences, Buckinghamshire, UK) according to manufacturer's 173 instructions. 2 µL of the purified PCR products was used as a template for the amplification of the V3 region of the 16S 174 rRNA gene with the 338F-518R primer pair (Alessandria et al., 2010). The forward primer, 338F, was attached with the 175 GC clamp necessary for the following DGGE analysis as described by Ampe et al. (1999). The PCR conditions were 176 those described by Osimani et al. (2015), except for the Taq polymerase (Sibenzyme) used in the present study. 5 µL of 177 PCR amplicons was loaded on a 1.5% agarose (w/v) gel with a 100 bp molecular weight marker (HyperLadder<sup>TM</sup> 100 178 bp) to check for the expected PCR product size of 180 bp prior to the PCR-DGGE analysis. Subsequently, 20 μL of the 179 PCR products was loaded on a 30-60% urea-formamide (w/v) gradient DGGE gel (100% corresponds to 7 M urea and 180 40% (w/v) formamide), and the gel was run at a constant voltage of 130 V for 4 h at 60 °C in 1× TAE buffer (0.04 mol

L<sup>-1</sup> Tris–acetate, 0.001 mol L<sup>-1</sup> EDTA) in a DGGE Bio-Rad D-code<sup>TM</sup> apparatus (Bio-Rad Laboratories). After the DGGE run, the gel was stained with SYBR Green I Stain 1X (Lonza, Walkersville, MD, USA) in 1X TAE for 30 min, visualized under UV light and photographed with a Complete Photo XT101 system (Explera). All of the single bands visible by eye after UV light exposure were excised with gel cutting pipette tips, introduced into 50  $\mu$ L of molecular biology grade water and left overnight at 4 °C to allow the elution of the DNA. 5  $\mu$ L of the eluted DNA was amplified by PCR as described above but using the 338F and 518R primers without the GC clamp. The amplicons were checked by electrophoresis and sent to Genewiz (Takeley, UK) for purification and sequencing. The resulting sequences in FASTA format were compared with those previously deposited in the GenBank database (http://www.ncbi.nlm.nih.gov) using Basic Local Alignment Search Tool (BLAST) (Altschul et al., 1990), and only the sequences showing ≥ 97% similarity was unambiguously assigned into species or genus levels.

2.6. PCR-DGGE analysis of the yeast and filamentous fungal communities

DNA extracted as previously reported was used for the analysis of the yeast and filamentous fungal communities. An approximately 250 bp long fragment of the D1/D2 region of the 26S rRNA gene was amplified using NL1 (5'-GCC ATA TCA ATA AGC GGA GGA AAA G-3') and LS2 (5'-ATT CCC AAA CAA CTC GAC TC-3') primers (Cocolin G-3') was added to the NL1 forward primer. The amplification reactions and conditions were carried out as described in Palla et al. (2017). The presence of amplicons was confirmed by electrophoresis in 1.5% (w/v) agarose gels stained with 20,000X REALSAFE Nucleic Acid Staining Solution (Durviz, s.l., Valencia, Spain). All gels were visualized using UV light and captured as TIFF format files using the UVI 1D v. 16.11a program for the FIRE READER V4 gel documentation system (Uvitec Cambridge, Eppendorf, Milan, Italy). The amplicons were analyzed using the DCode<sup>TM</sup> Universal Mutation Detection System (Bio-Rad, Milan, Italy). Twenty µL of the PCR products in 20 µL of a 2x buffer consisting of 70% glycerol, 0.05% xylene cyanol and 0.05% bromophenol blue were loaded on an 8% polyacrylamide-bisacrilamide (37.5:1) gel with a urea-formamide denaturing gradient ranging from 20% to 80%. The gels were run at 80 V and 60 °C for 16 hours and stained for 30 min in 500 mL of 1x TAE buffer containing 50 µL of Sybr® Gold Nucleic Acid Gel Stain (Life Technologies, Milan, Italy). The profiles were visualized as previously described. The bands of interest in the DGGE profiles were cut out from the gels for sequencing. DNA was extracted by eluting for 3 days in 50 μL 10 mM TE at 4 °C. One μL of the supernatant diluted 1:100 was used to reamplify the D1/D2 regions of the DNA according to the PCR protocol described above using an NL1 primer without the GC clamp. The amplification products were then purified with the UltraClean PCR

212	CleanUp Kit (MO-BIO Laboratories, CABRU Sas, Arcore, Italy) according to the protocol of the manufacturer,
213	quantified and 5' sequenced at the Eurofins Genomics MWG Operon (Ebersberg, Germany). Sequences were analyzed
214	using BLAST on the NCBI website (http://blast.ncbi.nlm.nih.gov/Blast.cgi). The related sequences were collected and
215	aligned using MUSCLE (Edgar, 2004a, b), and phylogenetic trees were constructed using the maximum likelihood
216	method based on the Kimura 2-parameter model (Kimura, 1980) using Mega 6 software
217	(http://www.megasoftware.net/) with 1000 bootstrap replicates (Tamura et al., 2013). The sequences were submitted to
218	the European Nucleotide Archive under the accession numbers from LS990841 to LS990863.

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2.7. 16S rRNA gene amplicon target sequencing

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- DNA directly extracted from hákarl samples was quantified using a QUBIT dsDNA Assay kit (Life Technologies,
- 223 Milan, Italy) and standardized at 20 ng μL<sup>-1</sup> and used a template in the PCR amplifying the V3-V4 region of the 16S
- rRNA gene using the primers and protocols described by Klindworth et al. (2013).
- The PCR amplicons were cleaned using the Agencourt AMPure kit (Beckman Coulter, Milan, Italy), and the resulting
- products were tagged using the Nextera XT Index Kit (Illumina Inc. San Diego. CA) according to the manufacturer's
- instructions. After the second clean-up, the amplicons were quantified using a QUBIT dsDNA Assay kit and equimolar
- amounts of the amplicons from different samples were pooled. The pooled samples were analyzed with an Experion
- workstation (Biorad, Milan, Italy) for quality analysis prior to sequencing. The sample pool was denatured with 0.2 N
- NaOH, diluted to 12 pM, and combined with 20% (vol/vol) denatured 12 pM PhiX prepared according to Illumina
- 231 guidelines. The sequencing was performed with a MiSeq Illumina instrument (Illumina) with V3 chemistry to generate
- 232 250 bp paired-end reads according to the manufacturer's instructions.

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234 2.7.1. Bioinformatics analysis

- After sequencing, the paired-end reads were first joined using FLASH software (Magoc and Salzberg, 2011) with
- default parameters. Joint reads were quality filtered (at Phred < Q20) using QIIME 1.9.0 software (Caporaso et al.,
- 238 2010) and the pipeline recently described (Ferrocino et al., 2017). Briefly, USEARCH software version 8.1 (Edgar et
- al., 2011) was used for chimera filtering and Operational Taxonomic Units (OTUs) were clustered at a 99% similarity
- threshold using UCLUST algorithms (Edgar, 2010). Centroid sequences of each cluster were mapped against the
- Greengenes 16S rRNA gene database version 2013 for taxonomic assignment. To avoid biases due to different
- sequencing depths, OTU tables were rarefied at 11010 sequences. The OTU table displays the higher taxonomy

243	resolution that was reached, and the two biological replicates from each sampling point were averaged. The tables were
244	then imported in the Gephi software (Bastian et al., 2009), and an OTU network was built.
245	All of the sequencing data were deposited in the Sequence Read Archive of the National Center for Biotechnology
246	Information (SRA accession number: SRP153795).
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248	2.7.2. Statistical analysis
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250	Statistics and plotting were carried out in the R environment (www.r-project.org). Alpha diversity indices were
251	calculated using the diversity function of the vegan package (Dixon, 2003). Weighted and unweight UniFrac distance
252	matrices and OTUs table were used to find differences between the samples using Anosim and Adonis statistical tests
253	through the function vegan in the R environment. Pairwise Wilcoxon tests were used, as appropriate, to determine
254	significant differences in alpha diversity or OTU abundance. The principal component analysis was plotted using the
255	function dudi.pca through the made4 R package.
256	
257	2.8. Real-time PCR analyses for the detection of foodborne pathogens
258	
259	Real-time PCR analyses were performed on a RotorGene Q thermal cycler (Qiagen, Hiden, Germany) exploiting
260	TaqMan chemistry. All target probes employed were dual-labeled with 5'-FAM and a 3'-nonfluorescent quencher (as
261	specified below). The oligonucleotides were purchased from ThermoFisher Scientific (Milan, Italy) and from LCG
262	Biosearch Technologies (Petaluma, CA, USA). The reaction mixtures were all prepared at a final 25 µl reaction
263	volume. Molecular-grade H <sub>2</sub> O was included in each analytical session as a negative control, as well as DNA from
264	reference strains as positive controls. Fluorescence was measured in the green channel for the target genes, and in the
265	yellow channel for the Internal Amplification Control.
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267	2.8.1. Detection of Shiga-toxin E. coli (STEC)
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269	STEC detection was performed according to the standard ISO/TS 13136:2012 specifications, as also previously
270	reported (Petruzzelli et al. 2013). Briefly, this method initially targets the Shiga-toxin genes stx1 and stx2, followed by
271	the eae adhesion factor and serogroup-specific genes (O157, O145, O103, O111, O26 and O104:H4). Amplification of
272	2 μl of template DNA was performed using the QuantiFast Pathogen PCR+IC kit (Qiagen) in combination with the

273	previously reported primer set and 5'-FAM-3'-MGBNFQ dual-labeled probes (Osimani et al. 2018). An Internal control
274	DNA and Internal Control Assay to be added to the reaction mix were provided with the kit.
275	DNA from STEC strains provided by the EU Reference Laboratory for STEC - Istituto Superiore di Sanità (Rome,
276	Italy) were included as positive controls.
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278	2.8.2. Detection of Pseudomonas aeruginosa
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280	P. aeruginosa was detected using the QuantiFast Pathogen PCR+IC kit together with a primer-probe set identifying the
281	presence of the ecfX gene, which encodes an extracytoplasmic sigma factor. The probe was dual-labeled with 5'-FAM
282	and 3'-BHQ1, and the assay mixture was prepared as described by Amagliani et al. (2013). Amplification of 2 $\mu$ l of
283	template DNA was performed following the thermal protocol indicated by the same authors. DNA from a previously
284	prepared boiled extract of <i>P. aeruginosa</i> ATCC 27853 was included as a positive amplification control in this assay.
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286	3. Results
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288	3.1. pH measurements and microbial viable counts
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290	The pH values measured in the analyzed samples ranged from between 8.07±0.06 and 8.76±0.00 (Table 1). The results
291	of the viable counts are reported in Table 1. In more detail, the mean values of the total mesophilic aerobes ranged from
292	between 1.00±0.00 and 5.71±0.09 log cfu g <sup>-1</sup> . Low counts of LAB on MRS at 30 °C were recorded with mean values
293	from between $< 1$ and $1.60\pm0.43$ log cfu g <sup>-1</sup> . Regarding the LAB counted on M17 at 22 °C, the mean values ranged
294	from between $1.95\pm0.07$ and $4.51\pm0.04$ log cfu g <sup>-1</sup> . Pseudomonadaceae counts showed mean values from between $<1$
295	and $1.59\pm0.16$ log cfu g <sup>-1</sup> . For both the Enterobacteriaceae and Eumycetes counts, mean values <1 log cfu g <sup>-1</sup> were
296	recorded. Finally, no Listeria monocytogenes was detected.
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298	3.2. PCR-DGGE analyses
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300	3.2.1. Bacteria
301	Regarding the bacteria, the results of PCR-DGGE analysis of the hákarl samples are reported in Table 2, while
302	Supplementary Figure 1 shows the DGGE profiles obtained from the analysis of the microbial DNA directly extracted
303	from the samples.

In more detail, the dominance of close relatives to Tissierella creatinophila was clear in all of the pooled samples with
sequence identities from between 85 and 98%. Moreover, close relatives to Anaerosalibacter species were detected in
the pooled samples H1, H2 and H10. Finally, close relatives to Murdochiella massiliensis, Sporanaerobacter acetigenes
and <i>Pontibacillus marinus</i> were also found in samples H5, H6 and H10, respectively.

- 3.2.2. Yeast and filamentous fungal communities
- A DNA fragment of approximately 250 bp containing the partial D1/D2 domain of the 26S rRNA gene was successfully amplified from all of the samples, except H1. DGGE analysis of the PCR products showed distinctive patterns characterized by intense and clearly defined fragments (Supplementary Figure 2). With the goal of identifying the microbial species present in hákarl, the main DGGE bands were excised, sequenced and matched to species by using BLAST and phylogenetic trees analyses (Table 3 and Figure 2). The sequences matched the yeast species Candida tropicalis, C. glabrata, C. parapsilosis, C. zeylanoides, Saccharomyces cerevisiae and the yeast genera Debaryomyces, Torulaspora, Yamadazyma, Sporobolomyces. Figure 3 shows the relative percentages of the most abundant fungal genera detected in the different samples. For each sample, the percentage was calculated by dividing the number of fragments referring to a genus by their total number. Each hákarl sample showed a different yeast composition, with Debaryomyces occurring in all the samples, although at different levels. In three hákarl samples (H2, H6 and H7), sequences related to filamentous fungal species were also found. In particular, Cladosporium tenuissimum occurred in the H2 sample, while Moristroma quercinum was found in the H7 sample. The H6 sample was characterized by the presence of Alternaria, and genera belonging to the family Didymellaceae, such as Phoma and Epicoccum.

3.3. 16S rRNA gene amplicon target sequencing

The total number of paired sequences obtained from the 16S rRNA gene sequencing reached 3,208,571 raw reads. After merging, a total of 997.224 reads passed the filters applied through QIIME, with an average value of 49.861,2  $\pm$ 17.399,72 reads/sample, and a mean sequence length of 456 bp. The rarefaction analysis and Good's coverage, expressed as a median percentage (95%), also indicated satisfactory coverage for all samples (Supplementary Table 1). Alpha-diversity indicated a higher level of complexity and the highest number of OTUs when only taking into the account the sample H3 (P < 0.05). Adonis and analysis of similarity (ANOSIM) statistical tests based on weighted and unweighted UniFrac distance matrices showed significant differences among the samples (P < 0.001; R= 0.980). Differences between the samples were further demonstrated by principal-component analysis (PCA) based on the

335	relative abundance of the main OTUs (Fig. 4). The PCA clearly showed a separation between samples (ANOSIM
336	statistical test $P < 0.01$ ). As shown in Fig. 7, the main OTUs shared among the data set were <i>Tissierella</i> (78.6% of the
337	relative abundance) (Table 4), Pseudomonas (8.4%), Abyssivirga (4.0%), Oceanobacillus (6.7%) and Lactococcus
338	(0.2%), and based on the size of the edges, it was possible to see that <i>Tissierella</i> can be considered a core OTU of the
339	hákarl (Fig. 5 and Table 4). Several minor OTUs (<1%) were also detected, including Listeria, Staphylococcus,
340	Photobacterium and Acinetobacter (Table 4).
341	
342	3.4. Real-time PCR analyses
343	
344	Fluorescence signals resulting from real-time PCR assays were analyzed by manually positioning the cycle threshold at
345	the take-off point of the positive control's amplification curve relative to the gene under investigation.
346	All DNA samples analyzed yielded negative results for both pathogens of interest. Every reaction mixture, regardless of
347	the type of master mix or internal amplification control used, yielded positive signals in the yellow channel, thus
348	ensuring the absence of inhibition and excluding false negative results.
349	As for STEC strain, the first detection step (aimed at revealing the two Shiga toxin-encoding genes stx1 and stx2) were
350	performed in singleplex, since the specific probes were labeled with the same fluorophore. All of the DNA samples
351	tested negative for both sequences; therefore, no further analysis of eae or serogroups was necessary according to the
352	ISO/TS 13136:2012. The reference strain used in this experiment was EURL-VTEC D07 E. coli O26 (stx1+, stx2+,
353	eae+).
354	
355	4. Discussion
356	
357	Among the fermented fish products of northern European countries, hákarl represents a masterful example of a delicacy
358	and niche product that, in former times, nourished the Icelandic populations (Skåra et al., 2015). It is noteworthy that
359	only a few producers are currently carrying out the production of hákarl in accordance with ancient traditions that
360	maintain the use of Greenland shark flesh.
361	Greenland shark is a slow growing, coldwater shark that can reach 21 feet in length. As reported by MacNeil et al.
362	(2012), the physiology of the Greenland shark is not generally well studied. It is noteworthy that high levels of
363	trimethylamine N-oxide (TMAO) have been detected in its flesh by different authors (Anthoni et al., 1991; Bedford et
364	al., 1998; Goldstein et al., 1967; Seibel and Walsh, 2002). The role of such a compound is not completely understood;
365	nevertheless, it is thought that the high concentrations found in polar fish suggest that this osmolyte may contribute to

366	the enhancement of osmotic concentrations, thus lowering the freezing point of the bodily fluids (MacNeil et al., 2012).
367	Moreover, both TMAO and its reduced form TMA represent low-density molecules that can increase the buoyancy of
368	the shark. It is also thought that, due to the high urea concentrations present in elasmobranchs like the Greenland shark,
369	TMAO might act as a counteracting solute that protects proteins from destabilization (MacNeil et al., 2012; Seibel and
370	Walsh, 2002).
371	The attention of the food industry and consumers towards locally produced traditional food is constantly increasing.
372	Although both the ancient and modern processing steps used to obtain hákarl from Greenland shark are mostly
373	recognized and standardized, less is known about the chemical and microbiological traits of such a food product. Based
374	on the physiology of the Greenland shark, it is thought that the microbiota occurring in hákarl might be strongly
375	influenced by the peculiarities of the flesh used as raw material. A 30-year-old study attempted to identify the bacterial
376	species in hákarl (Magnússon and Gudbjörnsdottir, 1984) with no mention of the possible occurrence of eumycetes.
377	Hence, to our knowledge, the microbiology of such a fermented food product has not yet been unveiled. The present
378	study aimed to reveal the microbial species occurring in ready-to-eat hákarl samples using a combination of traditional
379	microbiological culture-dependent (viable counts) and -independent methods (namely, PCR-DGGE, amplicon-based
380	sequencing and qualitative real-time PCR).
381	The samples under study were first subjected to pH measurements. The detected values were in agreement with those
382	reported by Skåra et al. (2015) for dried ready-to-eat hákarl. High pH values can be explained by microbial metabolic
383	activities that led to the conversion of urea, which is naturally present in Greenland shark flesh, into ammonia (Skåra et
384	al., 2015). Interestingly, Jang et al. (2017) reported similar pH values (8.4-8.9) in alkaline-fermented skate, which
385	represents a typical fermented seafood in South Korea. It is known that skates retain urea and trimethylamine N-oxide
386	in their muscles; hence, similarly to hákarl, the fermentation of skate, which is carried out at low temperature, leads to
387	the production of a distinctive odor due to the ammonia and trimethylamine produced during fermentation (Reynisson
388	et al., 2012).
389	In the present study, the viable counts of total mesophilic aerobes, presumptive mesophilic lactobacilli and lactococci,
390	Pseudomonadaceae, Enterbacteriaceae and eumycetes were assessed.
391	Regarding the total mesophilic aerobes, the counts were generally lower than the value of 8 log cfu g <sup>-1</sup> reported by
392	Skåra et al. (2015) in <i>hákarl</i> at the end of ripening.
393	As far as the counts of lactic acid bacteria are concerned, little is known about the magnitude of their presence in
394	hákarl, though this microbial group is known to play a primary role in the production of many other traditional
395	fermented fish, such as jeotgal from Korea, shidal from India, rakfisk from Norway, and numerous fermented fish
396	products from China (Françoise, 2010; Majumdar et al., 2016; Skåra et al., 2015; Thapa et al., 2016; Xu et al., 2018;

397	Zang et al., 2018a). In addition to this, the occurrence of lactic acid bacteria in the marine environment has been
398	reported (Gómez-Sala et al., 2016). Of note, the differences in the counts of presumptive lactococci in comparison with
399	those of presumptive mesophilic lactobacilli in $h\acute{a}karl$ . This difference agrees well with the results from metagenomic
400	sequencing that highlighted the sole presence of lactococci among lactic acid bacteria. Such a difference might
401	tentatively be ascribed to both the higher adaptation of lactococci to shark flesh fermentation conditions and their
402	acknowledged higher competitiveness in protein-rich substrates explained by their higher proteolytic and peptidase
403	activities in respect with lactobacilli (Requena et al., 1993; Quigley et al., 2013; Terzic-Vidojevic et al., 2014),
404	especially under alkaline conditions (Addi and Guessas, 2016).
405	The presence of Pseudomonadaceae has already been described in alkaline-fermented skate, where <i>Pseudomonas</i> was
406	reported to be among the dominant genera (Jang et al., 2017; Reynisson et al., 2012).
407	Regarding Enterobacteriaceae, it is noteworthy that their minimum pH for growth is 3.8, with an upper limit of
408	approximately 9.0; hence, even if they were present, the high pH of the samples did not allow Enterobacteriaceae to
409	grow in the fermented flesh. Among Enterobacteriaceae, STEC strains represent a major health treat for humans due to
410	their ability to produce food-borne infections (EFSA and ECDC, 2016). Although E. coli is not naturally harbored by
411	fish, STEC strains have recently been isolated by Cardozo et al. (2018) from the fish Nile tilapia. Of note is the absence
412	of STEC strains in all of the analyzed hákarl samples, as shown by real-time PCR assays.
413	In the present study, viable counts < 1 log cfu g <sup>-1</sup> were reported for eumycetes.
414	The use of PCR-DGGE and amplicon sequencing allowed major and minor microbial species to be detected in the
415	analyzed <i>hákarl</i> samples.
416	In more detail, a massive presence of <i>Tissierella</i> was detected by both PCR-DGGE and amplicon sequencing in all the
417	analyzed samples. Tissierella belongs to the Tissierellia classis nov., which includes the genera Anaerococcus,
418	Anaerosphaera, Finegoldia, Gallicola, Helcococcus, Murdochiella, Parvimonas, Peptoniphilus, Soehngenia,
419	Sporanaerobacter, and Tepidimicrobium. Moreover, the misclassified species Bacteroides coagulans and Clostridium
420	ultunense are also included.
421	Members of the class <i>Tissierellia</i> are Gram-positive or Gram-variable anaerobic obligate cocci and rods. As reported by
422	Chen et al. (2018), the genus <i>Tissierella</i> comprises protein-degrading anaerobes that can produce volatile fatty acids
423	such as acetic, butyric, and propionic acids. Moreover, it is acknowledged that Tissierella creatinophila grows on
424	creatinine as sole source of carbon and energy (Harms et al., 1998a). Tissierella species can degrade creatinine via
425	creatine, sarcosine, and glycine into monomethylamine, ammonia, carbon dioxide, and acetyl phosphate (Harms et al.,
426	1998b). Interestingly, active Tissierella cells have already been reported among the major genera involved in the
427	production of alkaline-fermented skate (Jang et al., 2017), thus confirming the possible adaptation of this microbial

428	genus to saline and alkaline conditions, as suggested by Jang et al. (2017). It is noteworthy that species belonging to the
429	Tissierella genus have also been described as psychrotolerant microorganisms (Prevost et al., 2013), thus likely
430	explaining the presence of members of such genus in the analyzed hákarl samples.
431	Regarding the presence of Oceanobacillus, which was detected only through amplicon sequencing, it is worth noting
432	that this bacterial genus comprises Gram-positive, spore forming rods that are obligate aerobes, extremely halotolerant
433	and facultatively alkaliphilic (Lu et al., 2001). Moreover, Oceanobacillus species can grow at temperatures between 15
434	and 42 °C and in pH range from 6.5 to 10, with an optimum pH between 7.0 and 9.5 (Lu et al., 2001). Kumar et al.
435	(2012) demonstrated that halotolerant bacteria, including Oceanobacillus, can produce enzymes that are salt stable and
436	active under extreme conditions. Interestingly, Yukimura et al. (2009) reported the isolation of Oceanobacillus strains
437	from a glacial moraine in Qaanaaq, Greenland, and this finding likely explains the presence of these spore forming
438	bacteria in the analyzed <i>hákarl</i> samples.
439	Amplicon sequencing supported the detection of Abyssivirga. This bacterial genus belongs to the Lachnospiraceae
440	family and comprises strictly anaerobic, mesophilic and syntrophic organisms. To the our best knowledge, the sole
441	species that belongs to this genus is represented by Abyssivirga alkaniphila, which is an alkane-degrading, anaerobic
442	bacterium recently isolated from a deep-sea hydrothermal vent system (Catania et al., 2018; Schouw et al., 2016). This
443	observation explains the presence of this bacterium in the marine environment and, hence, in the hákarl samples. This
444	species can grow between 14-42 °C and within a pH range between 7.0 and 8.2. As reported by Schouw et al. (2016),
445	who first described this species, A. alkaniphila can ferment carbohydrates, peptides and aliphatic hydrocarbons.
446	As for the presence of <i>Pseudomonas</i> , which was detected through amplicon sequencing, it is noteworthy that this
447	bacterial genus has already been reported among the most abundant isolates in fermented skate (Jang et al., 2017). The
448	genus Pseudomonas encompasses many alkaliphiles; among these, Yumoto al. (2001) reported the isolation of a novel
449	facultatively psychrothrophic alkaliphilic species of <i>Pseudomonas</i> , being <i>Pseudomonas alcaliphila</i> sp. nov. This species
450	of marine origin can grow at 4 °C and at pH 10 in the presence of NaCl; these features likely explain the presence of the
451	genus Pseudomonas in the analyzed hákarl samples. Moreover, it is widely acknowledged that the Pseudomonas
452	species responsible for fresh fish spoilage can also be present in fish-based foods (Stanborough et al., 2018). In this
453	regard, Liffourrena et al. (2010) reported that some species of <i>Pseudomonas</i> possess TMA degradation pathways. More
454	specifically, in Pseudomonas aminovorans TMAO is oxidized by a trimethylamine monooxygenase (TMA
455	monooxygenase), whereas TMA can be directly dehydrogenated to formaldehyde and dimethylamine (DMA) by a
456	trimethylamine dehydrogenase (TMA dehydrogenase). Similarly, in <i>Pseudomonas putida</i> the same enzymes, namely,
457	TMA monooxygenase and TMA dehydrogenase, oxidize TMA under aerobic conditions (Liffourrena et al., 2010). On
458	the one hand, these metabolic pathways represent a selective advantage for <i>Pseudomonas</i> , and on the other hand, they

459	lead to the removal of TMA (Liffourrena et al., 2010). Although the amounts of TMAO and TMA were not assessed in
460	the present study, it can be hypothesized that the metabolic activity of Pseudomonas species could presumably lead to
461	the detoxification of TMAO or TMA in the shark meat or during fermentation.
462	Of note, the pathogenic species <i>P. aeruginosa</i> was absent in all of the samples, as shown by qualitative real-time PCR
463	assays.
464	In the analyzed hákarl samples, the presence of Lactococcus was also detected through amplicon sequencing. The
465	presence of lactococci in the marine environment has already been demonstrated (Gómez-Sala et al., 2016); moreover,
466	Alonso et al. (2018) recently reported the isolation of <i>Lactococcus</i> strains from the gut of marine fishes. As reported by
467	Guan et al. (2011), lactic acid bacteria were among the dominant genera isolated from saeu-jeot, a Korean salted
468	fermented food produced made with shrimp (Acetes japonicas). Although their role has not yet been fully clarified,
469	lactococci were also detected in fermented skate and in narezushi, which is produced through the fermentation of salted
470	fish with rice (Jang et al., 2017). As reported by Françoise (2010), lactic acid bacteria are generally less competitive in
471	fish flesh than psychrotrophic Gram-negative bacteria; nevertheless, Ji et al. (2017) highlighted the essential role of
472	lactic acid bacteria (including Lactococcus) in flavor definition during fish fermentation. It is also noteworthy that lactic
473	acid bacteria can exert a potential biopreservative activity in seafood products (Ghanbari et al., 2013).
474	Regarding Alkalibacterium, the presence of this marine lactic acid bacteria has already been reported in marine
475	environments and organisms (Jang et al., 2017) as well as in marine-based foods such as <i>jeotgal</i> (Guan et al., 2011).
476	Interestingly, species belonging to the Alkalibacterium genus were found in fermented Spanish-style green table-olives
477	and blue-veined raw milk cheese, thus confirming the high adaptation of this genus to halophilic and alkaline conditions
478	(Lucena-Padrós et al., 2015; Yunita and Dodd, 2018).
479	Photobacterium, detected through amplicon sequencing, encompasses species that are naturally present in the marine
480	environment with both symbiotic or pathogenic relationships with marine organisms (Labella et al., 2018). This
481	bacterial genus was also detected by Reynisson et al. (2012) in fermented skate, where it was thought to play a key role
482	in the fish flesh fermentation.
483	Although found in a limited number of samples, Proteiniclasticum and Anaerobacillus were also detected. The former
484	genus includes anaerobic bacteria (e.g., <i>Proteiniclasticum ruminis</i> ) with proteolytic activity (Zhang et al., 2010),
485	whereas the latter genus includes species that can grow in alkaline environments (e.g., Anaerobacillus alkalilacustre)
486	(Zavarzina et al., 2009).
487	The low occurrence of Staphylococcus suggests a possible contamination during the processing of hákarl, even though
488	Ji et al. (2017) suggested a possible role of this genus in the bacterial amino acid metabolism occurring during the
489	fermentation of the fish Siniperca chuatsi, together with Acinetobacter. Of note is that although found as a minority

490 species in the hákarl samples analyzed in the present study, Magnússon and Gudbjörnsdottir (1984) reported 491 Acinetobacter as one of the fermentation-driving microorganisms during the production, along with Lactobacillus. 492 Regarding Ochrobactrum, it is noteworthy that members belonging to this halophilic genus have recently been isolated 493 by Jamal and Pugazhendi (2018) in Red Sea saline water and sediments. 494 Erysipelothrix was sporadically detected in the analyzed hákarl samples through metagenomic sequencing. 495 Interestingly, species of this genus have already been detected in soils collected from the Ross Sea region of Antarctica, 496 which is characterized by extreme low temperatures and high water salinity (Aislabie et al., 2008). 497 Listeria was found as minority OTU by amplicon sequencing; notwithstanding, no viable cells belonging to the 498 pathogenic species L. monocytogenes were found in any of the analyzed hákarl samples in accordance with the results 499 reported by Jang et al. (2017) in fermented skate. It is noteworthy that in seafood from cold environments L. 500 monocytogenes represents a foodborne pathogen of increasing public health and food safety concern (Elbashir et al., 501 2018; Jami et al., 2014). 502 Although the bacterial biota in hákarl has already tentatively been studied by Magnússon and Gudbjörnsdottir back in 503 1984, to our knowledge, the present study represents the first attempt to gain insight into the fungal biota present in this 504 fermented fish product. 505 The assessment of the diversity of yeast communities present in the hákarl samples revealed the occurrence of 4 yeast 506 genera: Debaryomyces, Candida, Saccharomyces, Torulospora, which are commonly found in several traditional 507 fermented beverages and food products, including fermented fish (Tamang et al., 2016), along with two other genera, 508 Yamadazyma and Sporobolomyces. The occurrence of sequences affiliated to Debaryomyces in all of the hákarl 509 samples suggests that this genus may be the main organism responsible for the late stages of hákarl fermentation. This 510 yeast, retrieved from the skin and inside the intestines of fresh fish by Andlid et al. (1995) and Gatesoupe (2007), was 511 also reported to be able to grow at extremely high salt concentrations and low water activity (a<sub>w</sub>) (Asefa et al., 2009; 512 Viljoen and Greyling, 1995), characteristics of the ripened hákarl. 513 Our findings are consistent with previous reports showing the occurrence of this genus in salted and traditionally 514 fermented fish from Thailand and Ghana (Paludan-Muller et al., 2002; Sanni et al., 2002). The yeasts of the genus 515 Debaryomyces might positively contribute to the development of the sensory qualities of fermented fish, as they are 516 known to occur in cheeses and dry-cured meat products. More specifically, in cheeses with high salt content, D. 517 hansenii was found to predominate, being responsible for the acceleration of lipolysis and proteolysis (Andrade et al., 518 2009). 519 Several sequences affiliated with different species from the genus Candida were retrieved in almost all the hákarl

samples. In particular, C. tropicalis, C. glabrata, C. parapsilosis, and C. zeylanoides were identified. Our results agree

521	with a previous report showing the dominance of the genus Candida in ripened Suan yu, a Chinese traditional fermented
522	fish (Zang et al., 2018b) and a study demonstrating the presence of C. zeylanoides and C. tropicalis in a salted and
523	fermented traditional fish called "adjuevan", which is produced in Ivory Coast (Clementine et al., 2012).
524	Candida was reported to produce more flavoring substances than other yeasts by metabolizing branched-chain amino
525	acids (BCAAs) through the Ehrlich pathway (O'Toole, 1997). In particular, C. tropicalis was also isolated from
526	Burukutu, a Nigerian traditional fermented beer, and it was shown to produce protease, phytase, lipase and esterase
527	enzymes (Ogunremi et al., 2015). These last two enzymes improve the aromatic profile of fermented foods by
528	increasing their free fatty acid content, which are precursors to the formation of different aromatic compounds (Arroyo-
529	Lopez et al., 2012).
530	Three sequences affiliated with Saccharomyces were retrieved from the nine hákarl samples, confirming the presence
531	of such yeasts in fermented fish (Clementine et al., 2012; Zang et al. 2018a).
532	Close relatives of Yamadazyma and Sporobolomyces were also detected in the H9 and H10 hákarl samples,
533	respectively. These two yeast genera were not previously recovered from fermented fish products, although
534	Sporobolomyces is a marine yeast commonly found in deep-sea waters (Kutty and Philip, 2008).
535	As for the occurrence of filamentous fungi, sequences affiliated with the genera Alternaria, Cladosporium,
536	Phoma/Epicoccum, and Moristroma were retrieved from the analyzed hákarl samples. Among such fungal genera,
537	Cladosporium was previously found during the fermentation of the Chinese traditional fermented fish Suan yu (Zang et
538	al. 2018). The presence of filamentous fungi, such as Aspergillus, Penicillium and Mucor, was also reported from some
539	Japanese fermented fish products, i.e., Katsuobushi and Narezushi, where they produced enzymes such as amylase,
540	protease, and lipase, which are important for the improvement of the nutritional and functional traits of fermented goods
541	(Fukuda et al., 2014).
542	
543	Conclusions
544	
545	Overall, the combination of culture-dependent and -independent methods allowed major and minor microbial species
546	harbored by the ready-to-eat hákarl samples to be detected. The culture-dependent approach provided insight into the
547	viable microbial species, whereas the culture-independent methods were pivotal in preventing the possible

minor components of the bacterial biota, the use of PCR-DGGE led to the identification of both bacterial and fungal

underestimation of bacterial diversity caused by culturing biases or the presence of microbial cells in the "viable but

non culturable" (VBNC) state. It is noteworthy that although amplicon sequencing was essential for detecting major and

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551	populations at the species level, thus contributing to the development of a first overview of the microbiota occurring in
552	this poorly studied food product.
553	Based on our results, hákarl revealed a complex and heterogeneous biodiversity. The bacterial community was
554	characterized by species well adapted to alkaline and saline environments; the dominant genus was Tissierella, followed
555	by Oceanobacillus, Pseudomonas and Abyssivirga. Moreover, based on the presence of Pseudomonas in the analyzed
556	samples, a role of this bacterial genus in the detoxification of TMAO or TMA in shark meat during fermentation may be
557	hypothesized. The fungal community was mainly represented by Debaryomyces, Candida and, to a lesser extent,
558	Saccharomyces species, which through interactions with the bacterial community might play key roles in the late stages
559	of hákarl fermentation, especially contributing to the development of the sensory qualities of the end product. Further
560	studies are needed to establish the roles and the viabilities of the detected microbial species occurring during shark
561	fermentation, as well as their interactions and relationships with the physical-chemical and rheological parameters of
562	hákarl. Moreover, the occurrence of spore-forming bacteria should also be evaluated since the presence of these
563	microorganisms has already been reported by different authors in fresh or fermented fish products (Metcalf et al., 2011,
564	Reynisson et al.,2012). It is noteworthy that, among spore forming bacteria, Clostridium botulinum type E spores and
565	toxins (produced even at low temperatures) can commonly be found in seafood (Elbashir et al., 2018; Iwamoto et al.,
566	2010), thus representing a serious health threat for the consumers.
567	
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569	
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572	
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Planalto de Bolona: an artisanal cheese produced in uncommon environmental conditions in the Cape Verde

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806	FIGURE LEGENDS
807	
808	Fig. 1 Ready-to-eat hákarl
809	
810	Fig. 2. Affiliation of the sequences retrieved from DGGE gel fragments (marked in Supplementary Figure 2) with the
811	existing sequences of the partial D1/D2 region of the large sub-unit rRNA gene.
812	
813	Phylogenetic analysis was inferred by using the Maximum Likelihood method based on the kimura 2-parameter model.
814	Bootstrap (1000 replicates) values below 70 are not shown. Evolutionary analyses were conducted in MEGA6. The
815	sequences from the database are indicated by their accession numbers. The DNA sequences retrieved in this work are
816	indicated by their corresponding band number and their accession number.
817	
818	Fig. 3. Relative abundance (%) of yeast and filamentous fungal genera detected in each hákarl sample.
819	
820	For each sample the percentage was calculated as follows: the number of fragments referring to a genus divided by their
821	total number.
822	
823	Fig. 4 PCA based on the OTU abundance of the hákarl grouped as a function of the samples.
824	
825	The first component (horizontal) accounts for the 40.66% of the variance and the second component (vertical) accounts
826	for the 22.53 %
827	
828	Fig. 5 OTU network summarizing the relationships between taxa and samples.
829	
830	Only OTUs occurring at 0.2% in at least 2 samples are shown. The abundances of OTUs in the two biological replicate
831	were averaged. The sizes of the OTUs were made proportional to weighted degree (i.e., for OTUs, this measures the
832	total occurrence of an OTU in the whole data set) using a power spline. OTUs and samples are connected with a line
833	(i.e., edge) to a sample node, and its thickness is made proportional to the abundance of an OTU in the connected
834	sample.

**Table 1** Results of pH and viable counting (log cfu per gram) of bacteria and eumycetes in ready-to-eat *hákarl* samples

Sample	pН	Total mesophilic	Presumtive mesophilic	Presumptive mesophilic	Pseuodomonadacea	e Enterobacteriaceae	Eumycetes	
		aerobes	lactobacilli	lactococci				
H1	8.07±0.06	3.61±0.02	<1	4.51±0.04	<1	<1	<1	
H2	$8.23\pm0.01$	$5.71\pm0.09$	$1.60\pm0.43$	$4.39\pm0.01$	$1.54\pm0.34$	<1	<1	
H3	$8.20\pm0.01$	$1.24\pm0.34$	<1	$4.27 \pm 0.05$	<1	<1	<1	
H4	$8.09\pm0.01$	$1.30\pm0.00$	<1	$3.84\pm0.08$	1.59±0.16	<1	<1	
H5	8.37±0.01	$2.40\pm0.02$	<1	$3.20\pm0.17$	$1.30\pm0.00$	<1	<1	
H6	$8.53\pm0.04$	$2.01\pm0.15$	<1	$3.22\pm0.31$	1.15±0.21	<1	<1	
H7	$8.41\pm0.00$	$1.15\pm0.21$	<1	$1.95 \pm 0.07$	$1.00\pm0.00$	<1	<1	
H8	$8.41\pm0.01$	$1.00\pm0.00$	<1	$2.39\pm0.06$	<1	<1	<1	
H9	$8.46 \pm 0.01$	$2.24\pm0.09$	<1	$3.30\pm0.08$	1.24±0.34	<1	<1	
H10	$8.76\pm0.00$	$2.01\pm0.15$	<1	$4.04\pm0.15$	<1	<1	<1	

Values are expressed as means  $\pm$  standard deviation

Table 2. Sequencing results of the bands excised from the DGGE gel obtained from the amplified fragments of bacterial DNA extracted directly from the pooled hákarl samples

Sample	e Band <sup>a</sup> Identification		% Identity <sup>b</sup>	Most closely related GeneBank sequence				
H1	1	Tissierella creatinophila	98%	NR_037028				
	2	Anaerosalibacter massiliensis	97%	NR_144694				
	3	Anaerosalibacter sp.	93%	LT598565				
	4	Tissierella creatinophila	97%	NR_117377				
	5	Tissierella creatinophila	95%	NR_117377				
H2	6	Anaerosalibacter massiliensis	97%	NR_144694				
	7	Tissierella creatinophila	97%	NR_117377				
Н3	8	Tissierella creatinophila	85%	NR_117377				
H4	9	Tissierella creatinophila	98%	NR_117377				
H5	10	Murdochiella massiliensis	97%	NR_148568				
	11	Tissierella creatinophila	98%	NR_117377				
Н6	12	Tissierella creatinophila	96%	NR_117377				
	13	Sporanaerobacter acetigenes	97%	NR_117381				
	14	Tissierella creatinophila	95%	NR_117377				
H7	15	Tissierella creatinophila	97%	NR_117377				
	16	Tissierella creatinophila	94%	NR_117377				
H8	17	Tissierella creatinophila	97%	NR_117377				
H9	18	Tissierella creatinophila	97%	NR_117377				
	19	Tissierella creatinophila	90%	NR_117377				
H10	20	Tissierella creatinophila	97%	NR_117377				
	21	Pontibacillus marinus	97%	LT992038				
	22	Anaerosalibacter massiliensis	97%	NR_144694				
	23	Anaerosalibacter massiliensis	93%	NR_144694				

 <sup>&</sup>lt;sup>a</sup> Bands are numbered as indicated in Supplementary Figure 1
 <sup>b</sup> Percentage of identical nucleotides in the sequence obtained from the DGGE bands and the sequence of the closest relative found in the GenBank database.

Table 3 Identification of yeasts and filamentous fungi occurring in the hákarl by sequencing the fragments obtained from DGGE profiles.

DGGE	Identification	Identity (%) <sup>b</sup>	Most closely related		
fragments <sup>a</sup>	Identification	identity (%)	GeneBank sequence		
H2-1	Candida tropicalis CBS:2320	99%	KY106851.1		
H2-2	Torulaspora delbrueckii CBS 1146	100%	NG 058413.1		
	Torulaspora pretoriensis CBS 11124		KY109883.1		
	Torulaspora quercuum CBS 11403		NG 058413.1		
H2-3	Cladosporium tenuissimum QCC:M024/17	100%	KY781762.1		
H3-4	Candida parapsilosis CBS:2915	100%	Y102317.1		
H3-5	Debaryomyces hansenii CBS:11096	100%	KY107560.1		
	Debaryomyces prosopidis JCM 9913		NG_055701.1		
	Debaryomyces subglobosus JCM 1989		NG_055699.1		
	Debaryomyces fabryi CBS:4373		KY107483.1		
H3-6	Saccharomyces cerevisiae CBS:7961	100%	KY109313.1		
H4-7	Debaryomyces hansenii CBS:11096	100%	KY107560.1		
	Debaryomyces prosopidis JCM 9913		NG_055701.1		
	Debaryomyces subglobosus JCM 1989		NG_055699.1		
	Debaryomyces fabryi CBS:4373		KY107483.1		
H6-8	Candida zeylanoides CBS:947	100%	KY106918.1		
H6-9	Alternaria sp strain QCC/M011/17	99%	KY744118.1		
H6-10	Debaryomyces hansenii CBS:11096	100%	KY107560.1		
	Debaryomyces prosopidis JCM 9913		NG_055701.1		
	Debaryomyces subglobosus JCM 1989		NG_055699.1		
	Debaryomyces fabryi CBS:4373		KY107483.1		
H6-11	Alternaria sp. isolate 1A1	100%	MF379649.1		
H6-12	Epicoccum sp CN018	100%	KX954392.1		
	Phoma herbarum JN0408		MG004796.1		
H7-13	Moristroma quercinum BN1678	98%	AY254051.1		
H7-14	Candida zeylanoides CBS:947	100%	KY106918.1		
H7-16	Debaryomyces hansenii CBS:11096	100%	KY107560.1		
	Debaryomyces prosopidis JCM 9913		NG_055701.1		
	Debaryomyces subglobosus JCM 1989		NG_055699.1		
	Debaryomyces fabryi CBS:4373		KY107483.1		
H7-17	Candida glabrata CBS:859	99%	KY106478.1		
H8-18	Candida parapsilosis CBS:2193	99%	KY102320.1		
H9-19	Yamadazyma mexicana CBS 7066	97%	NG_058439.1		
H9-20	Debaryomyces hansenii CBS:11096	100%	KY107560.1		
	Debaryomyces prosopidis JCM 9913		NG_055701.1		
	Debaryomyces subglobosus JCM 1989		NG_055699.1		
	Debaryomyces fabryi CBS:4373		KY107483.1		
H9-21	Saccharomyces cerevisiae CBS:7961	100%	KY109313.1		
H10-22	Debaryomyces hansenii CBS:11096	100%	KY107560.1		
	Debaryomyces prosopidis JCM 9913		NG_055701.1		
	Debaryomyces subglobosus JCM 1989		NG_055699.1		
	Debaryomyces fabryi CBS:4373		KY107483.1		
H10-23	Sporobolomyces salmoneus CBS:488	100%	KY109767.1		
	Sporobolomyces roseus OL10		KF273854.1		
H10-24	Saccharomyces cerevisiae CBS:7961	100%	KY109313.1		

 <sup>&</sup>lt;sup>a</sup> Bands are numbered as indicated in Supplementary Figure 2
 <sup>b</sup> Percentage of identical nucleotides in the sequence obtained from the DGGE bands and the sequence of the closest relative found in the GenBank database.

**Table 4** Incidence of the major taxonomic groups detected by 16S amplicon target sequencing. Only OTUs with an incidence above 0.2% in at least 2 biological replicate were averaged.

OTU	H1	H2	Н3	H4	Н5	Н6	H7	Н8	Н9	H10
Abyssivirga	0.19	2.88	2.92	5.73	3.56	5.52	2.93	6.88	4.81	4.41
Acinetobacter	0.00	0.04	0.90	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Alkalibacterium	0.06	0.15	0.10	0.24	0.13	0.19	0.18	0.18	0.21	0.23
Anaerobacillus	0.00	0.91	0.01	0.00	0.40	0.06	0.00	0.02	0.00	0.00
Erysipelothrix	0.05	0.15	0.23	0.20	0.13	0.13	0.05	0.11	0.13	0.02
Lactococcus	0.46	0.14	0.05	0.26	0.07	0.11	0.09	0.06	0.29	0.67
Listeria	0.00	0.29	0.00	0.00	0.16	0.05	0.00	0.02	0.01	0.00
Oceanobacillus	43.69	1.84	2.17	3.76	0.71	3.32	5.08	1.52	4.05	1.15
Ochrobactrum	0.00	0.57	0.00	0.00	0.27	0.06	0.00	0.01	0.01	0.00
Photobacterium	0.01	0.67	0.07	0.10	0.14	1.00	0.35	0.10	0.20	0.20
Proteiniclasticum	0.00	0.14	0.43	0.35	0.24	0.03	0.00	0.08	0.13	0.00
Pseudomonas	42.99	0.29	0.31	0.25	0.65	8.24	13.04	4.65	2.57	10.96
Staphylococcus	0.00	0.17	0.00	0.00	2.71	0.00	0.00	0.00	0.00	0.00
Tissierella	12.37	87.77	92.56	88.66	90.42	80.75	77.82	86.07	87.09	82.11

OTU Operational Taxonomic Units

Fig. 1



Fig. 2

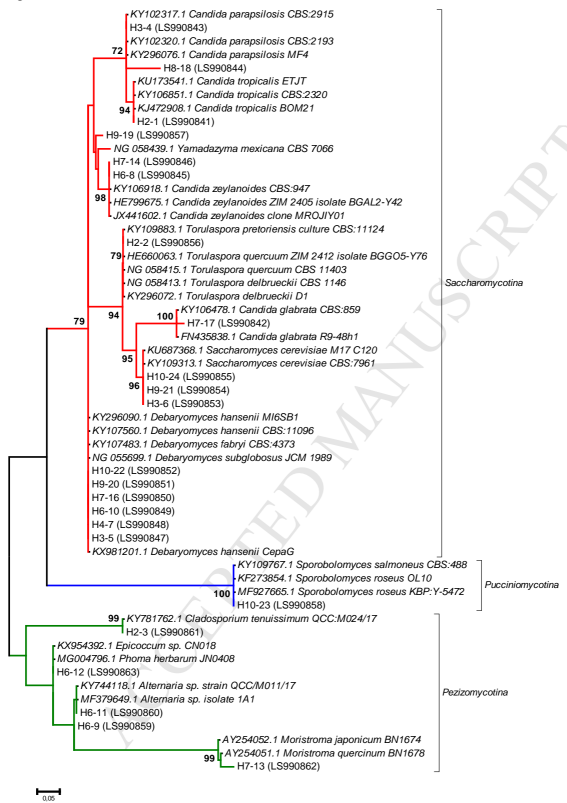


Fig. 3

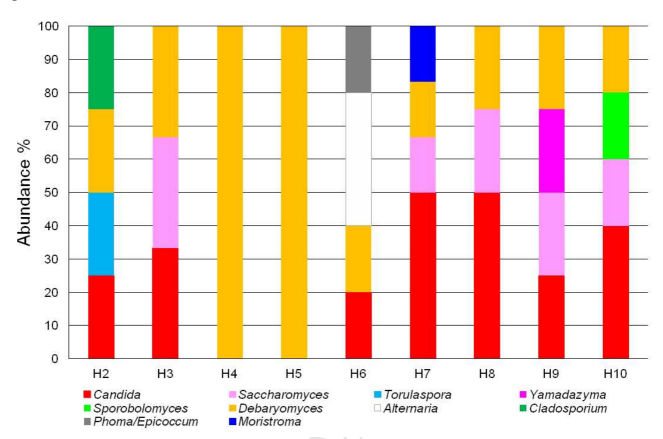


Fig. 4

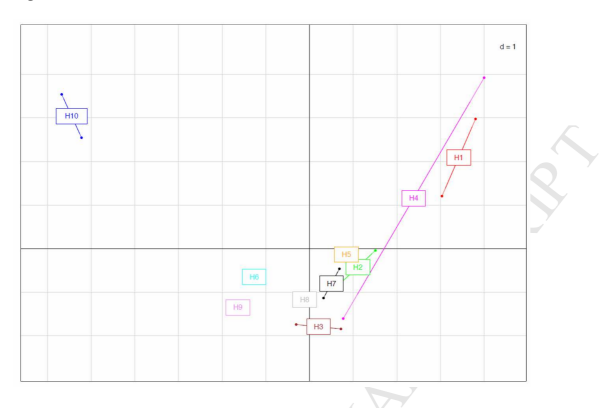
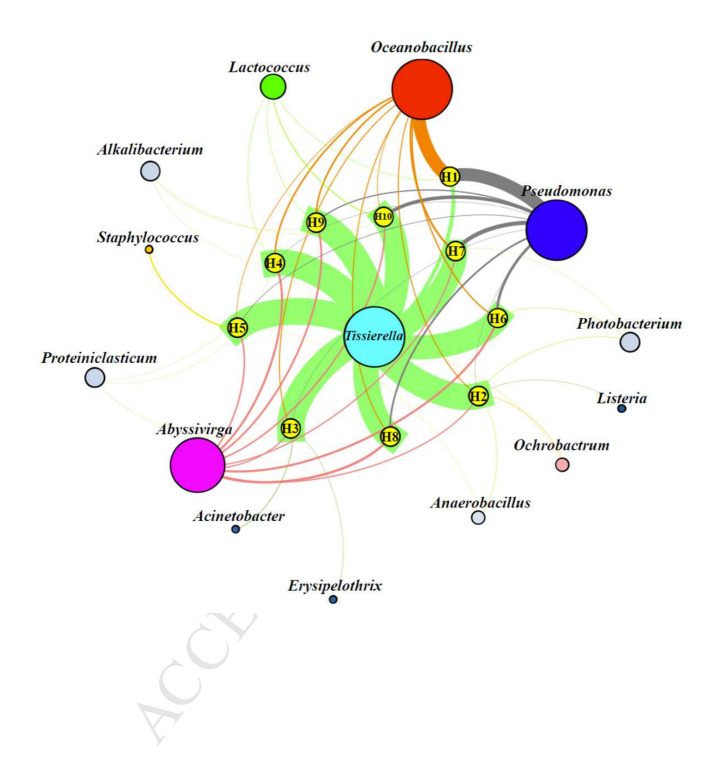


Fig. 5



## Highlights

The microbiota of hákarl was studied using a polyphasic approach

Both metagenomic sequencing and PCR-DGGE highlighted the dominance of *Tissierella* sp.

The contribution of Pseudomonas in the detoxification of shark meat was hypothesized

The fungal community was mainly represented by Debaryomyces and Candida species

Real-time PCR showed the absence STEC and Pseudomonas aeruginosa in all the samples