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# 1 Detection and characterization of lipids in eleven species of fish by

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# non-targeted liquid chromatography/mass spectrometry

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26 **1. Abstract** 

27 Fish is an important nutrition source because its lipids, which are rich in  $\omega$ -3 fatty acids, are beneficial for human health. However, studies focusing on their detection, composition, and 28 nutritional value are limited. In this study, we applied a non-targeted lipidomic approach 29 based on ultra-high performance liquid chromatography coupled with linear-ion trap-30 Orbitrap mass spectrometry (UHPLC/LTQ-Orbitrap-MS) to comprehensively profile, 31 compare, and detect unknown lipids in eleven types of dietary fish. A total of 287 molecular 32 species from five major lipid classes were characterized by MS/MS analysis. Multivariate 33 34 principal component analysis revealed the distinct lipid composition in shishamo smelt and Japanese sardine compared to other fish types. The assessment of nutritional indices 35 based on the levels of free fatty acid suggested that among the eleven fish types, shishamo 36 37 smelt is highly beneficial for health. Further, lipids such as N-acyl 38 lysophosphatidylethanolamine were detected and characterized for the first time in fish 39 fillets. Hierarchical cluster correlations indicated the predominance of glycerophospholipids 40 (GPs) and sphingolipids in sardine, whereas fatty acyls and triacylglycerols (TAGs) were 41 predominant in shishamo smelt. The high levels of polyunsaturated fatty acid-enriched GPs 42 and TAGs in dietary fish endow it with great potential as a health-promoting food for human 43 consumption. This study offers a comprehensive analysis of lipids and their compositions in 44 fish fillets, demonstrating their potential use in the nutritional assessment of functional foods.

Keywords: Fish fillet, lipidomics, nutritional indices, N-acyl lyso-phosphatidylethanolamine,
liquid chromatography, mass spectrometry, correlation analysis

#### 47 **1. Introduction**

48 Fish are an important component of human nutrition and contain many valuable nutrients, such as bioactive lipids, essential amino acids, minerals, vitamins, and proteins(Neff et al., 49 2014). The benefit of this meat matrix is highly related to the quality of their lipid content. 50 Lipids such as long-chain fatty acids account for one of the key constituents of fish fillets 51 with potential health benefits (Xu et al., 2020). Fatty acids are classified into saturated fatty 52 acid (SFA), monounsaturated fatty acid (MUFA), and polyunsaturated fatty acid (PUFA), 53 depending on the number of double bonds. Fish and its products are a known source of 54 55 long-chain  $\omega$ -3 PUFAs such as docosahexaenoic acid (DHA) and eicosapentaenoic acid (EPA) (Xu et al., 2020). Extensive studies have demonstrated that the intake of fish 56 products is associated with the reduction of major chronic diseases, inflammation, and 57 58 cardiovascular events (Virtanen et al., 2008). The fatty acid content and types mainly vary 59 according to species, seasons, and geographic conditions (Zhang et al., 2020).

60 Diets rich in SFAs increase the blood cholesterol levels and cardiovascular disease risks (Garonzi et al., 2021). Conversely,  $\omega$ -3 PUFAs offers protection against cardiovascular 61 disease (Yagi et al., 2017), diabetes (Farrell et al., 2021), and neurodegeneration to reduce 62 inflammation (Borsini et al., 2021). However, men with high blood levels of  $\omega$ -3 PUFAs are 63 64 reported to have an increased risk of prostate cancer (Brasky et al., 2013). Hence, a 65 balanced intake of SFA and PUFA in the diet is preferred to reduce the disease risk (Kang et al., 2005). In addition to the PUFA:SFA ratio, several nutritional indices have been 66 67 developed to provide further knowledge on fish lipid quality and its association with diseases (Fernandes et al., 2014). Moreover, several functional lipids other than fatty acids, 68 such as phospholipids (PLs), glycerolipids (GLs), and sphingolipids, are yet to be 69 70 characterized well in dietary fishes. Notably, the abundant nutritional and healthy 71 characteristics of fish are primarily attributed to the abundance of PLs and GLs rich in high-72 level of  $\omega$ -3 PUFAs (Schuchardt & Hahn, 2013).

73 Although several analytical methods focus on the targeted determination of lipids in various 74 fish types by gas or liquid chromatography/mass spectrometry(LC/MS), they are limited to 75 the analysis of a specific class of lipids such as fatty acids (de Souza et al., 2020; 76 Devadason et al., 2016). Recently, untargeted mass spectrometric approaches have been 77 widely used in comprehensive food and fish lipidome profiling (Siddabasave et al., 2021; 78 Zang et al., 2018; Zhou et al., 2018). However, studies focusing on the comprehensive lipid 79 fingerprinting of dietary fish are limited. In our previous study, we profiled fatty acids and 80 annotated furan-fatty acids in seafood by nontargeted analysis (Siddabasave et al., 2021). 81 In this study, we present a comprehensive lipidome profiling of dietary fish using a highly 82 sensitive ultra-high performance liquid chromatography coupled with linear-ion trap-Orbitrap 83 mass spectrometry (UHPLC/LTQ-Orbitrap-MS) technique. Comparative correlations of 84 multiple lipids among the characterized eleven common dietary fish were studied, and their 85 nutritional significance from the dietary aspect was discussed. Further, previously unknown lipids such as N-acyl lysophosphatidylethanolamine were characterized in fish fillets using 86 87 the un-targeted technique.

#### 88 2. Materials and method

#### 89 **2.1. Materials**

Solvents such as isopropanol, chloroform, and methanol of LC/MS grade were purchased from Wako Pure Chemical Industries, Ltd., (Osaka, Japan). Ammonium acetate (eluent additive for LC/MS, 1 M) was obtained from Sigma-Aldrich (St. Louis, USA). Zirconium ceramic oxide bulk beads (1.4 mm, catalog no. 15-340-159, Fisherbrand) for fillet homogenization were purchased from Thermo Fisher Scientific (Tokyo, Japan). The EquiSPLASH LIPIDOMIX quantitative standard for mass spectrometry and oleic acid-d9 internal standards were purchased from Avanti Polar Lipids, Inc (Alabaster, USA).

## 97 2.2. Fish samples

98 Commercially available refrigerated fish such as Pacific cod (Gadus macrocephalus, Madara in Japanese, Hokkaido, Japan), saffron cod (*Eleginus gracilis*, Komai, Hokkaido, 99 100 Japan), Lindberg skate (Raja pulchra, Makasube, Hokkaido, Japan), blackin flounder (Microstomus achne, Nametagarei, Hokkaido, Japan), Japanese sardine (Sardinops 101 melanostictus, Maiwashi, Hokkaido, Japan), Japanese pond smelt (Hypomesus 102 103 nipponensis, Wakasai, Hokkaido, Japan), red seabream (Pagrus major, Madai, Ehime, Japan), flatfish (Paralichthys olivaceus, Hirame, Hokkaido, Japan), shishamo smelt 104 105 (Spirinchus lanceolatus, Shisyamo, Hokkaido, Japan), skipjack tuna (Katsuwonus pelamis, 106 Katuo, Shizuoka, Japan), and Okhotsk Atka mackerel (Pleurogrammus azonus, Mahokke, 107 Hokkaido, Japan) were purchased from different supermarkets located in Sapporo, Hokkaido, Japan and were stored at -80 °C until the analysis. Triplicate analysis was 108 109 performed for each fish species.

#### 110 **2.3. Lipid extraction**

111 Total lipid extraction of the fish fillet was performed by the method described by Folch with 112 minor modifications (Folch et al., 1957; Siddabasave et al., 2021). A freeze-dried fish fillet was weighed and homogenized in 10 volumes of methanol in a homogenizer (Bead Mill 4, 113 Fisherbrand, Tokyo, Japan) for 30 s (x 2 cycles). Subsequently, 100 µL of the homogenate 114 (10 mg) was transferred to a 1.5 mL Eppendorf tube, and 100 µL of the pre-mixed internal 115 116 standard in methanol was added (10 ng of each of the following: phosphatidylcholine (PC) (15:0-18:1(d7)), phosphatidylethanolamine (PE) (15:0-18:1(d7)), phosphatidylglycerol (PG) 117 (15:0-18:1(d7)), phosphatidylserine (PS) (15:0-18:1(d7)), phosphatidylinositol (PI) (15:0-118 18:1(d7)), lysophosphatidylethanolamine (LPE) (18:1(d7)), lysophosphatidylcholine (LPC) 119 (18:1(d7)), sphingomyelin (SM) (d18:1/18:0(d9)), ceramide (Cer) (d18:1/15:0 (d7)), 120 triacylglycerol (TAG) (15:0-18:1(d7)-15:0), diacylglycerol (DAG) (15:0-18:1(d7)), cholesterol 121 ester (18:1(d7)), and monoacylglycerol (MAG) (18:1(d7)) and 100 ng of oleic acid (d9)) and 122 123 vortexed at 3500 rpm for 30 s. Then, 400 µL of chloroform was added and vortexed for

5 min; subsequently, 100  $\mu$ L of milli-Q was added and the solution was further vortexed for 30 s. The lipid extracts were centrifuged at 15,000 rpm for 10 min. After centrifugation, the chloroform layer was transferred to a new Eppendorf tube, and the aqueous layer was reextracted with an additional 400  $\mu$ L of chloroform. The combined chloroform layer containing lipids was collected and evaporated under a vacuum in a centrifuge evaporator, which was then dissolved in 100  $\mu$ L of methanol, centrifuged, and transferred to an LC/MS vial.

#### 131 2.4. LC/MS analysis

The UHPLC system (Shimadzu Corporation, Kyoto, Japan) was equipped with an Atlantis 132 T3 C18 column (2.1 x 150 mm, 3 µm, Waters, Milford, USA). The LC/MS conditions were 133 134 slightly modified compared to those reported in our previous studies (Gowda et al., 2021; Siddabasave et al., 2021). The column temperature was maintained at 40 °C; the flow rate 135 and injection volume were 0.2 mL/min and 10 µL, respectively. Mobile phases used were 136 10 mM ag.  $CH_3COON_4$  (A), isopropanol (B), and methanol (C). The gradient elution is as 137 138 follows: 0-1 min, 30% B and 35% C; 1-9 min, 75% B and 15% C; 9-21 min, 82.5% B and 15% C; 21-25 min, 95% B and 5% C; 25-26 min, 30% B and 35% C. The column was pre-139 equilibrated for 4 min; the total run time was 30 min. Mass spectrometry analysis was 140 performed on a LTQ Orbitrap mass spectrometer (Thermo-Fisher Scientific Inc., San Jose, 141 142 USA) in the positive and negative ionization modes. The electron spray ionization (ESI) conditions were set as follows: capillary temperature 330 °C; sheath gas flow 50 units; and 143 auxiliary gas 20 and 30 units for the positive and negative modes, respectively. For the 144 145 negative mode, the source and capillary voltage was set to 3 kV and 10 V, respectively. For 146 the positive mode, source and capillary voltage was set to 4 kV and 25 V, respectively. The 147  $MS^1$  scan range was set at m/z 160–1900 and 100–1750 for negative and positive modes, respectively in a Fourier transform mode with a resolving power of 60,000 collision energy 148

of 35 V to obtain MS spectra for high-resolution masses. Low-resolution MS/MS spectra
were obtained at a collision energy of 40 V in the ion-trap mode.

#### 151 **2.5. Data processing for identification of lipids**

The raw data obtained from the MS was processed for alignment, peak extraction, 152 153 identification, and peak area integration using MS-DIAL software (ver 4.2). The following parameters were set during data processing in MS-DIAL: peak height cut-off 1000 154 155 amplitude, mass slice width 0.1 Da, smoothing level and minimum peak width 3 and 5 156 scans, sigma window 0.5, and signal intensity 5 folds (> blank) (Siddabasave et al., 2021). The retention time and MS<sup>1</sup> tolerance were 0.5 min and 0.015 Da, respectively. The 157 identification of lipid molecular species was confirmed by their MS/MS spectra. Extracted 158 159 ion chromatograms and MS spectra illustrated herein were obtained using Xcalibur 2.2 (Thermo Fisher Scientific, Waltham, USA). The relative quantification of lipid metabolites 160 was performed according to the guidelines of Lipidomics Standards Initiative level 2 and 161 level 3. Relative quantification was achieved considering that the lipid molecule was 162 163 quantified by the labeled internal standard of similar lipid subclasses or a typical lipid class. The relative amount was calculated by considering peak area ratios of the analyte to the 164 internal standard and multiplying it using the added internal standard. 165

#### 166 **2.6. Calculation of lipid health quality indexes**

The nutritional assessment of eleven fish types was conducted based on their values of index of atherogenicity (IA), health promotion index (HPI), hypo/hypercholesterolemic (HH) ratio, and fish lipid quality (FLQ). These parameters were calculated based on the equations suggested in a previous study (J. Chen & Liu, 2020). IA indicates the relationship between the total SFAs and the total unsaturated fatty acids (UFAs); it assesses the atherogenic potential of FA (Ulbricht & Southgate, 1991). The SFAs, which include C12:0, C14:0, and C16:0, are considered pro-atherogenic whereas UFAs are considered to be anti-atherogenic properties. The IA value is ranges from 0.21 to 1.41 for fish (S. Chen et al.,

175 2004) and was calculated using the following equation:

176 IA =

 $IA = [C12:0+(4 \times C14:0) + C16:0]/\Sigma UFA.$ 

The HPI is the reciprocal of IA and is an index for evaluating the nutritional value of dietary fat (S. Chen et al., 2004). This value determines the effect of fatty acid composition on cardiovascular diseases (CVD). It is mainly used in research on dairy products and a value ranges from 0.16-0.68 (S. Chen et al., 2004). The HPI was calculated using the following equation:

182 HPI =  $\Sigma UFA/[C12:0+(4 \times C14:0)+C16:0]$ .

Further, the HH ratio was used to assess the effect of FA composition on cholesterol and 183 cardiovascular risk. lt 184 predict the characterizes the relationship between hypocholesterolemic fatty acid (C18: 1 and PUFA) and hypercholesterolemia FA. It is 185 186 assumed that, HH ratio could accurately reflect the effect of the FA composition on CVD. The HH ratio is ranges from 1.54 to 4.83 for fish (S. Chen et al., 2004). and was calculated 187 using the following equation: 188

189  $HH = (C18:1+\Sigma PUFA/(C12:0+C14:0+C16:0))$ 

In the HH equation, the term "cis-C18:1" was revised to "C18:1" because we did not 190 characterize cis or trans isomers in the present method. FLQ calculates the sum of EPA 191 and DHA as a percentage of total fatty acids. This parameter is used to assess the quality 192 of the fish lipids and is more suitable for marine products because of their high proportions 193 194 of EPA (FA 22:6) and DHA (FA 20:5). FLQ is widely used for marine products and it calculated using the ratio of sum of EPA and DHA as percentage of total FAs. The FLQ is 195 ranges from 13.01 to 36.37 for fish (S. Chen et al., 2004) and was calculated using the 196 following equation: 197

198 FLQ =100×(C22:6 + C20:5)/ΣFA.

#### 199 **2.7 Statistical analysis**

The data were subjected to one-way analysis of variance (ANOVA), principal component analysis (PCA), and cluster correlation analysis using MetaboAnalyst (ver 5.0). The data were plotted in GraphPad Prism 8 software as the mean  $\pm$  standard error (analysis of triplicate of each sample). Student's *t*-test (p < 0.05) or one-way ANOVA (\* p < 0.05, \*\* p <0.01, \*\*\* p < 0.001, ns: not significant) were used to study statistically significant differences between the groups.

#### 206 **3. Results and Discussion**

#### **3.1. Lipid annotation and principal component analysis of eleven types of fish**

208 The lipidomic profiles of fish fillets were analyzed by UHPLC/LTQ-Orbitrap-MS in both 209 positive and negative modes. A total of 287 lipid molecular species were annotated based 210 on their exact masses and MS/MS behavior by MS-DIAL software. After the identification and relative quantification using the internal standard method, the multivariant analysis such 211 as one-way ANOVA (p < 0.05) with Tukey's post-hoc analysis was applied. The results are 212 shown in Fig 1a; 282 lipid molecular species (red) were statistically significant, and 5 were 213 214 insignificant (green). The PCA demonstrated the differences based on the liquid composition changes of the groups. The PCA analysis of the 287 annotated lipids from 215 eleven types of fish was performed, and the results are illustrated in **Fig 1b**. A previous 216 study demonstrated comparative lipid profiles of various commercial fishes from Japan. 217 218 However, their analysis was limited to only the fatty acid class (Devadason et al., 2016). The score plots showed distinct grouping and clear separation between the Japanese 219 sardine and the shishamo smelt, suggesting their distinct lipid composition compared to 220 221 other analyzed fish types. Although these two species were previously studied for their lipid 222 content, a comprehensive class-specific determination was not conducted (Shirai et al., 2002). The two components, PC1 and PC2, accounted for 90.3% of the total model 223 variance, which was described by 56.9% of PC1. The loading plot signifies the distribution 224 225 of the important variables, the two first principal components, and the groupings among the

samples. A loading plot of components that affects the differences is visualized in the score plot, as shown in **Fig 1b.** The larger positive or negative loading scores indicate that a variable strongly affects the components. Lipids such as triacylglycerols (TAG (16:0/18:1/18:1)), cholesterol esters (CE 20:5, CE 22:6), and sphingomyelin (SM (d18:1/24:0)) have large positive loading scores (for PC1) and contributed mainly for the group separation. These results indicate the differential lipids responsible for group separation in the two-dimensional PCA plot.

#### 233 **3.2 Distributions of lipid classes in eleven different types of fish**

234 The percentage distributions based on the total amount of each lipid class for all the fish 235 types are represented by pie charts, and the results are illustrated in Fig 2a. The lipid 236 molecular species were grouped based on their main categories, such as glycerophospholipids (GP), sphingolipids (SP), glycerolipids (GL), fatty acyls (FA), and 237 238 sterols (ST), according to the LIPID MAPS guidelines. Although these lipids are beneficial to 239 human health, their percentage distribution studies in fish are limited. Profiling of phospholipids and eicosanoids in Atlantic salmon was attempted using shotgun lipidomics; 240 however, other lipid classes were not examined (Yeo & Parrish, 2021). Recently, interest in 241 GPs has dramatically increased owing to their biochemical features in the lipid fraction 242 243 (Pongsetkul et al., 2017). In our results, the Lindberg skate (83.61%), skipjack tuna (74.80%), and Okhotsk atka mackerel (70.68%) demonstrated the highest abundance of 244 GPs. In contrast, shishamo smelt (86.39%), Japanese sardine (64.27%), and red seabream 245 246 (50.64%) had higher levels of GLs. The abundance of different molecular species of GPs has been reported in fermented fish (Zang et al., 2018). However, information on the lipid 247 248 molecular species in commercially available dietary fish is still limited. Hence, in our 249 analysis, we annotated the lipids from multiple classes at molecular species level using 250 state-of-the-art mass spectrometry. Blackin flounder (34.40%) and flatfish (24.48%) 251 demonstrated the highest abundance of sphingolipids, whereas saffron cod (49.23%) had 252 the highest percentage of fatty acyls. In a previous study, the amount of fatty acids in

saffron cod was present in the milligram-scale with a higher amount of neutral lipids than
polar lipids (Copeman et al., 2020). Remarkably, we also determined sterols (mainly CEs as
major lipids in Pacific cod (68.89%). Although sterols, including various cholesterols esters,
were reported in the red fish muscle (Voronin et al., 2021) and cod liver (Hammann et al.,
2015), they were not observed in the Pacific cod fillets. Thus, the present study is the first to
demonstrate the abundance of sterols in fillets.

#### **3.3 Fish fatty acid compositions and aspects of nutritional indices**

Untargeted analysis of fish samples detected 36 fatty acid molecular species from  $C_{12}$  to 260  $C_{28}$  carbon chain. These species were further classified as saturated fatty acid (SFA), 261 262 mono-unsaturated fatty acid (MUFA), and poly-unsaturated fatty acid (PUFA), based on their number of double bonds. The relative amounts of total SFAs, MUFAs, and PUFAs in 263 different types of fish studied are illustrated in Fig 2b. The highest amount of SFAs 264 265 (10.50±0.21 mg/100g) and MUFAs (11.59±0.32 mg) was observed in shishamo smelt 266 followed by saffron cod (8.22±0.36 mg, 3.23±0.06 mg), whereas Japanese sardine had the lowest amount of MUFAs (0.15±0.01 mg). Furthermore, the highest amount of bioactive 267 PUFAs was observed in shishamo smelt (12.7±0.30 mg) and the lowest in skipjack tuna 268  $(0.16\pm0.01 \text{ mg})$ . A previous study had established that fish-derived  $\omega$ -3 PUFAs, particularly 269 270 DHA and EPA, had favorable effects on cardiovascular, neurological, inflammatory, and 271 metabolic disorders (Parolini, 2019). Considering this aspect, shishamo smelt could be a 272 promising source as a dietary supplement of sufficient amount of PUFAs compared to other 273 groups. Subsequently, the evaluation of PUFA/SFA (P:S) ratio was conducted, and the 274 results are shown in Fig 2c.

The P:S ratio is considered as the most common index to assess the dietary food nutritional value. It is also used to evaluate the impact of diet on the cardiovascular health; a value in the range 1–1.5 in salmon trout is favorable for reducing the risk of cardiac diseases (Kang et al., 2005). Our results indicate that shishamo smelt has the highest P:S ratio (1.21)

followed by saffron cod (0.79), whereas skipjack tuna had the lowest value (0.04). 279 Therefore, shishamo smelt could be considered a valuable source in diet due to its 280 appropriate levels of P:S ratio. Recently, the P:S ratio of various food products was 281 282 extensively reviewed, and the ranges were as follows: sea weeds: 0.42-2.12, meat: 0.11-283 2.04, fish: 0.11–1.62, and shellfish: 0.20–2.10 (J. Chen & Liu, 2020). Our previous study 284 also demonstrated the highest P:S value of 1.52 in salmon trout (Siddabasave et al., 2021). 285 Hence, the P:S value of dietary fish evaluated in this study could be useful to assess their 286 potential nutritional value. The relative concentrations of each identified fatty acid are 287 represented by a heatmap visualization, as shown in Fig 3a, classifying each group and clustered with each other. The intense red color indicates the high concentration of fatty 288 acids in that group. Conversely, the intense blue color implies the low concentration of fatty 289 290 acids. These results suggest that fatty acids, including FA 20:5 and FA 22:6, are 291 considerably high in shishamo smelt and Saffron cod.

292 Fish with different fatty acid profiles may contribute differently toward human health. These 293 benefits can be evaluated using the nutritional indices, such as the IA, HPI, HH ratio, and 294 FLQ; their respective equations are described in Section 2.6. The nutritional qualities of the 295 lipid profiles from this study and related indices are shown in Fig 3b. The IA value was 296 higher in skipjack tuna (4.86±0.39) followed by Lindberg skate (3.09±0.15) and lower in shishamo smelt (0.30±0.01) and saffron cod (0.48±0.02). A previous study reported that 297 consuming foods with lower IA values could decrease the total cholesterol and low-density 298 lipoprotein-cholesterol in human blood plasma (Yurchenko et al., 2018). Additionally, the IA 299 value of fish was reported to range from 0.21 to 1.41 (J. Chen & Liu, 2020). Among all the 300 fish analyzed, the highest HPI value was in shishamo smelt (3.30±0.06) and saffron cod 301 (2.09± 0.11), and the lowest value was in skipjack tuna (0.21±0.02) and Lindberg skate 302 303 (0.32±0.02). Previously, the HPI was mainly used to evaluate the nutritional value of milk and cheese; the value ranges from 0.16 to 0.68 (J. Chen & Liu, 2020). 304

305 Additionally, the HH ratio was the highest in shishamo smelt (4.18  $\pm$  0.09) and saffron cod 306  $(2.22 \pm 0.13)$  and the lowest in skipjack tuna  $(0.21 \pm 0.02)$ . Compared with the P:S ratio, the 307 HH ratio can precisely reflect the effect of the fatty acid composition on cardiovascular 308 diseases. The HH ratio indicates the effect of fatty acids on cholesterol metabolism, and the 309 higher values are desirable for human health. A recent review reported that the HH levels in 310 fish were in between 0.87 and 4.83 (J. Chen & Liu, 2020). Finally, the quality of lipids is 311 assessed by FLQ indices. The highest value was obtained for saffron cod (23.51±0.65), 312 Pacific cod (22.45 $\pm$ 0.34), and shishamo smelt (21.95 $\pm$ 0.79). Skipjack tuna (1.29  $\pm$  0.07) 313 demonstrated the lowest FLQ indices among all fish types. Fish lipid quality is the percentage of EPA and DHA in total fatty acids. These fatty acids reduce the risk of CVD 314 (Breslow, 2006) and dementia (Johnson & Schaefer, 2006). Previous studies reported the 315 316 quality of fish lipids by FLQ indices, with the value in the 13.01–36.37 range (J. Chen & Liu, 317 2020). Additionally, the FLQ value assessed the seasonal changes in the fatty acid profile of the fillet of red bream (Senso et al., 2007). Because this index was high in saffron cod, 318 this fish has a great scope for further development as a nutritional supplement. 319

## 320 **3.4. Identification of N-acyl-lysophosphatidylethanolamine (LNAPE) in fish fillets**

Untargeted LC/MS analysis of total lipid extracts from eleven different types of fish detected 321 322 the previously discovered lipids, particularly N-acyl-lysophospahtidylethanolamine (LNAPE), 323 which were the partially hydrolyzed products of N-acyl-phospahtidylethanolamine (NAPE). 324 Although these lipids belong to the family of phosphatidylethanolamines (PEs), their amine 325 moieties are esterified with fatty acids. NAPEs were previously detected in the brain of bony 326 fish; however, they were not detected in cartilaginous skeletons (Schmid et al., 1990). 327 Nevertheless, the study lacks a complete characterization of the lipids. The occurrence of 328 NAPEs in mammals and other organisms, such as yeast, insects, reptiles and worms, were also reported (Wellner et al., 2013). However, to the best of our knowledge, no studies have 329 330 detected LNAPEs in fish fillets. NAPEs are the precursors of bioactive N-acylethanolamines (NAE), which are endogenous signaling molecules. The incorporation of NAPE-producing 331

bacteria into the gut retards high-fat diet-induced obesity in wild-type mice (Chen et al.,
2014) and improves cardiometabolic disease indices in low-density lipoprotein null mice
(May-Zhang et al., 2019). Furthermore, multiple studies demonstrated the anti-inflammatory
effects of NAPEs (Dalle Carbonare et al., 2008; Solorzano et al., 2009). Despite their great
benefits, the characterization of NAPEs in food is still limited.

In this study, we detected seven molecular species of LNAPE with FA 20:5 as the 337 predominating N-acylated fatty acid. The extracted ion chromatograms of LNAPE are 338 shown in Fig 4a; their relative levels and other lysophospholipids in each fish type are 339 340 visualized by a heatmap, as illustrated in Fig 4b. The identification of these lipids was 341 confirmed by their high-resolution masses obtained from their MS spectra and fragmentation behavior observed in low-resolution MS/MS spectra. The MS and MS/MS 342 343 spectra of all the seven major LNAPE species are shown in Fig 5. These lipids ionized in 344 negative mode to provide [M-H]<sup>-</sup> ion as the precursor ion. The compound elutes with a 345 retention time of 13.92 min and as a m/z value of 688.4929 [C<sub>37</sub>H<sub>72</sub>NO<sub>8</sub>P, calculated m/z: 346 688.4923, error: -0.05 ppm], and ionizes to generate m/z 452.3 by the neutral loss of FA 16:1 and H<sub>2</sub>O. Additionally, peaks at *m/z* 255.3 (FA16:0) and 253.3 (FA 16:1) in the MS/MS 347 348 spectra suggest that the detected compound is LNAPE-(N-(FA16:0)/16:1). The lipid eluted 349 at 14.25 min has m/z 740.5242 [C<sub>41</sub>H<sub>76</sub>NO<sub>8</sub>P, calculated m/z: 740.5236, error: 0.28 ppm] and ionizes to generate m/z 478.5 by the neutral loss of FA 18:2 and H<sub>2</sub>O. Moreover, 350 predominant peaks at m/z 279 (FA 18:2) and FA m/z 281 (FA 18:1) in the MS/MS spectra 351 indicate that the detected compound is LNAPE (N-(FA18:1)/18:2). The lipid eluted at 13.91 352 min has *m/z* 736.4936 [C<sub>41</sub>H<sub>72</sub>NO<sub>8</sub>P, calculated *m/z*: 736.49231, error: 1.23 ppm] and 353 ionizes to provide m/z 452.3 by the neutral loss of FA 20:5 and water. Further, peaks at m/z354 255 (FA 16:0 and m/z 301 (FA 20:5) in the MS/MS spectra imply that the detected 355 356 compound is LNAPE-(N-(FA16:0)/20:5).

The compound eluted at 13.64 min has m/z 748.4928 [C<sub>42</sub>H<sub>72</sub>NO<sub>8</sub>P, calculated m/z: 748.4923, error: 2.28 ppm] and demonstrates an MS behavior similar to that mentioned

359 earlier; it ionizes to afford m/z 464.4 (neutral loss of FA 20:5 and water), 267 (FA 17:1), and 301 (FA 20:5) in the MS/MS spectra, suggesting that the detected compound is LNAPE (N-360 361 (FA17:1)/20:5). Similarly, the lipid eluted at 13.96 min has m/z 762.5084 [C<sub>43</sub>H<sub>74</sub>NO<sub>8</sub>P, 362 calculated m/z: 762.5079, error: 1.39 ppm] and ionizes to afford m/z 478 by the neutral loss 363 of FA 20:5 and water. Peaks at m/z 281(FA 18:1) and 301(FA 20:5) in the MS/MS spectra suggest that the detected compound is LNAPE (N-(FA18:1)/20:5). The lipid eluted at 13.27 364 min has *m/z* 758.4775 [C<sub>43</sub>H<sub>70</sub>NO<sub>8</sub>P, calculated *m/z*: 758.4766, error: 1.18 ppm] and ionizes 365 366 to provide m/z 474 (neutral loss of FA 20:5 and water); the peaks at m/z 277 (FA 18:3) and 367 m/z 301(FA 20:5) in the MS/MS spectra suggest that it is also an isomeric mixture and mainly comprises LNAPE (N-(FA18:3)/20:5). Finally, the lipid eluted at 13.90 min has m/z368 810.5081 [C<sub>47</sub>H<sub>74</sub>NO<sub>8</sub>P, calculated *m/z*: 810.5078, error: 2.65 ppm], and it ionizes to give 369 m/z 520 (loss of FA 20:5 and water) and m/z 500 (loss of FA 22:6 and water). Moreover, 370 371 peaks of FA 22:6 (*m/z* 327), FA 20:5 (*m/z* 301), FA 20:4 (*m/z* 303), FA 22:6 (*m/z* 283), and FA 20:5 (m/z 301) in the MS/MS spectra indicate that it is an isomeric mixture of LNAPE (N-372 (FA 20:4)/22:6) and LNAPE (N-(FA 22:5)/20:5). 373

374 The concentrations (mean  $\pm$  SE, in mg/100g) of total LNAPE were as follows: red seabream (0.25±0.01), flatfish (0.33±0.05), Okhotsk atka mackerel (0.44±0.02 mg), Japanese sardine 375 376 (0.75±0.02 mg), blackin flounder (0.75±0.02 mg), Lindberg skate (0.25±0.01 mg), skipjack tuna  $(0.19\pm0.03 \text{ mg})$ , saffron cod  $(0.15\pm0.00 \text{ mg})$ , shishamo smelt  $(0.33\pm0.02 \text{ mg})$ , Pacific 377 cod (0.19±0.01 mg), and Japanese pond smelt (0.83±0.03 mg). LNAPE (N-(FA18:1)/18:2) 378 is abundant in red seabream, LNAPE (N-(FA20:4)/22:6) and LNAPE (N-(FA16:0)/16:1) 379 were abundant in blackin flounder. LNAPE (N-(FA16:0)/20:5), LNAPE (N-(FA18:1)/20:5), 380 and LNAPE (N-(FA17:1)/20:5) were abundant in Japanese pond smelt, whereas LNAPE (N-381 (FA18:3)/20:5) and LPE (N-(FA17:1/20:5) were abundant in Japanese sardine. The 382 383 abundance of PUFAs as LNAPEs acyl chains suggest that these lipids are also a potential 384 source of dietary PUFAs. To the best of our knowledge, the present study is the first to

characterize and demonstrate the relative levels of LNAPEs in multiple fish types. Further
 investigations are necessary to determine their biological effects.

#### 387 **3.5.** Hierarchical cluster analysis of complex lipids profiled in eleven types of fish

388 Hierarchical clustering heatmaps visualizes the concentrations of complex lipids detected in 389 fish fillets. The visualization of lipids such as phosphatidylcholines (PC), phosphatidylserine 390 (PS), phosphatidylinositol (PI), phosphatidylglycerols (PG), and cardiolipins (CL) in eleven types of fish is represented by a heatmap and is shown in Fig 6a, 6b. A tight clustering 391 392 among the sample groups was observed. The concentration of PC, PS, and PI was relatively higher in Japanese pond smelt and Lindberg skate compared to other fish types. 393 394 These phospholipids are highly enriched with PUFAs such as FA 22:6, FA 20:5, and FA 20:4. Hence, these membrane lipids could be the potential sources of PUFAs in the diet. 395 Moreover, fish-derived  $\omega$ -3 PUFAs were reported to significantly affect intestinal microbiota 396 397 and regulate immune function (Parolini, 2019).

The energy metabolism associated lipids, such as PGs and CLs, were relatively high in 398 399 shishamo smelt. Further, lipids such as Cer, SM, CE, MAG, DAG, TAG, and NAE in eleven types of fish are visualized by a heatmap, and the results are illustrated in Fig 6c,6d. The 400 concentration of sphingolipids (Cer and SM) and sterols (CE) was higher in Japanese pond 401 402 smelt whereas that of MAG, DAGs, and TAGs was higher in shishamo smelt. Remarkably, 403 TAGs were also enriched with many PUFAs as acyl chains. The NAE lipids were relatively 404 high in shishamo smelt and saffron cod. NAEs are a functionally diverse family of signaling lipids that have been implicated as metabolic signals regulating the nutritional status and 405 lifespan (Lucanic et al., 2011). These lipids may have important implications for biomedical 406 407 research on aging and obesity (De Petrocellis & Di Marzo, 2011). A previous study demonstrated the application of the untargeted technique for comprehensive lipid profiling 408 in round scad (He et al., 2020) and sea urchin (Zhou et al., 2018); several studies focused 409 on profiling specific lipids such as phospholipids (Yeo & Parrish, 2021; Zang et al., 2018) or 410

fatty acids (Devadason et al., 2016; Zhang et al., 2020) in various other fish types. However,
no evident studies have demonstrated a comprehensive profile of lipids in the fish types
explored in the present study.

#### 414 **4. Conclusion**

415 In summary, various lipid species and their relative content in the eleven types of fish were analyzed using UHPLC/LTQ-Orbitrap MS. GP was the most abundant lipid class in many 416 groups. The PUFAs, P:S ratio, and HPI was significantly higher in shishamo smelt, 417 suggesting it as a potential health-beneficial dietary fish type. The previously 418 uncharacterized and biologically beneficial lipids such as LNAPEs were detected and 419 420 characterized for the first time in fish fillets. In addition, the comprehensive profile of multiple lipid classes and their distribution among each fish type was revealed by the cluster 421 correlation analysis. Shishamo smelt was enriched with TAGs, and Japanese sardines had 422 423 a relatively large amount of PC, PI, SM, and CEs. Therefore, lipids in dietary fishes can 424 account for their nutritional and health-beneficial functions. Because exploring the sources of these lipids is of significant interest, we aim to apply the untargeted technique to 425 determine the unidentified lipids in other food types. 426

#### 427 **CRediT authorship contribution statement**

Siddabasave Gowda B. Gowda: Conceptualization, Methodology, Supervision, Writing original draft. Yusuke Minami: Formal analysis, Visualization, Writing - original draft.
Divyavani Gowda: Data curation, visualization, Writing - review & editing. Hitoshi Chiba:
Supervision, Writing - review & editing. Shu-Ping Hui: Supervision, Resources, Writing review & editing.

## 433 **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Lipids	Red seabream	Flatfish	Okhotsk atka mackerel	Japanese sardine	Blackin flounder	Lindberg skate	Skipjack tuna	Saffron cod	Shishamo smelt	Pacific cod	Japanese pond smelt
LPC	4.17±0.28	00.0±00.0	0.43±0.10	2.29±0.05	2.25±0.18	0.04±0.01	1.35±0.11	0.40±0.05	0.56±0.03	3.20±0.37	1.31±0.22
LPE	0.23±0.01	0.08±0.00	0.57±0.00	0.59±0.03	1.33±0.04	0.15±0.01	0.19±0.02	0.44±0.01	0.51±0.02	0.17±0.01	0.53±0.01
A-LPE	0.03±0.00	0.04±0.00	0.02±0.00	0.01±0.00	0.39±0.01	0.04±0.00	0.09±0.01	0.29±0.02	0.28±0.01	0.06±0.00	0.22±0.01
LNAPE	0.25±0.01	0.33±0.03	0.44±0.02	0.75±0.02	0.75±0.02	0.25±0.00	0.19±0.01	0.15±0.00	0.34±0.02	0.39±0.02	0.83±0.03
LPI	L	ı	0.01±0.00	0.01±0.00	0.05±0.01	0.01±0.00	0.01±0.00	0.03±0.00	0.04±0.00	0.03±0.01	0.02±0.01
PC	42.3±2.39	25.05±0.57	20.44±2.29	25.46±1.84	21.36±1.52	53.24±1.95	16.93±2.33	4.06±0.33	23.47±0.97	4.18±0.22	197.01±10.49
AA-PC	10.12±0.13	8.53±0.20	8.52±1.26	8.68±0.55	5.98±0.47	17.74±3.7	11.59±2.02	1.53±0.07	6.69±0.61	2.46±0.12	10.84±2.23
PE	9.95±0.58	6.12±0.19	12.9±2.58	17.78±0.90	8.54±0.80	20.86±2.95	1.62±0.20	0.77±0.01	4.98±0.17	1.52±0.08	11.60±0.20
AA-PE	3.98±0.21	3.00±0.14	2.28±0.47	3.01±0.16	3.80±0.42	13.16±1.99	3.05±0.36	0.49±0.01	1.94±0.06	0.69±0.04	4.22±0.03
PS	1.81±0.07	1.85±0.07	1.44±0.04	1.95±0.05	1.69±0.05	2.23±0.05	1.33±0.14	0.88±0.01	1.72±0.04	0.27±0.01	1.51±0.04
Ā	3.27±0.73	3.07±0.07	1.83±0.04	5.43±0.10	3.17±0.06	3.03±0.07	2.32±0.21	2.13±0.04	2.46±0.07	0.55±0.02	11.17±0.15
PG	0.04±0.00	0.02±0.00	0.01±0.01	0.14±0.00	0.08±0.00	0.04±0.00	0.07±0.01	0.08±0.00	0.32±0.01	0.01±0.00	0.20±0.01
СГ	0.62±0.03	0.11±0.00	0.06±0.00	0.33±0.01	0.11±0.01	0.34±0.02	0.21±0.02	0.08±0.00	0.17±0.01	0.11±0.01	0.35±0.01
Cer	1.16±0.50	1.03±0.09	0.75±0.11	1.39±0.06	1.94±0.04	1.27±0.16	0.29±0.04	0.30±0.01	3.02±0.15	0.61±0.05	10.21±0.94
SM	14.72±2.06	16.56±1.06	5.28±1.54	14.88±1.92	21.31±0.57	12.00±1.56	6.05±0.88	1.53±0.25	9.79±0.36	0.75±0.07	299.00±38.36
CE	1.25±0.39	3.67±0.18	1.54±0.03	2.52±0.46	31.56±2.23	10.09±1.16	1.99±0.11	6.54±0.61	37.43±7.8	64.27±4.60	210.94±31.76
MAG	0.11±0.00	0.08±0.01	0.14±0.01	0.16±0.01	0.02±0.00	0.03±0.00	0.12±0.03	0.15±0.01	1.29±0.04	0.06±0.00	0.32±0.01
NAE	0.17±0.01	0.52±0.03	1.28±0.06	0.45±0.02	1.46±0.02	0.20±0.01	0.16±0.01	1.92±0.10	2.16±0.06	1.78±0.07	1.14±0.00
TAG	98.16±9.61	0.16±0.00	2.79±0.07	182.33±11.09	0.15±0.01	0.16±0.01	0.11±0.03	0.34±0.06	845.60±0.06	0.47±0.04	329.68±13.55
Total lipic (mg/100g	I ) 192.34±17.03	3 70.33±2.63	60.82±8.65	268.16±17.28	105.96±6.45	134.89±13.64	47.68±6.55	22.11±1.58	942.77±133.26	81.56±5.74	1124.43±98.67

**Table 1:** Amount (mg/100 g of fillet) of lipid characterized in eleven types of dietary fish. (A-

601 LPE: Alkyl lysophosphatidylethanolamine, AA-PE: Alkyl acyl phosphatidylethanolamine)

602 Figure legends:

Fig 1 Multivariant analysis of lipid metabolites annotated in fish fillet samples. a. One-way
ANOVA analysis of 287 identified lipid molecular species (Tukey's HSD with p<0.05). b.</li>
Principal component analysis score plot and loadings plot of lipid metabolites from eleven
fish types.

607 Fig 2 Distribution of lipid classes and fatty acid compositions. a. Percentage distribution of five major lipid classes in eleven fish types (GP: Glycerophospholipids, SP: Sphingolipids, 608 GL: Glycerolipids, FA: Fatty acyls, St: Sterols). b. Fatty acid composition a based-on 609 unsaturation in eleven fish types (p<0.05, are considered as statistically significant. The 610 groups which are significant are shown here: SFA: a<sub>1</sub>-[c<sub>1</sub>,e<sub>1</sub>,h<sub>1</sub>,i<sub>1</sub>,j<sub>1</sub>,k<sub>1</sub>] i.e. group a1 is 611 significantly different with the groups shown in square bracket, b<sub>1</sub>-[c<sub>1</sub>,e<sub>1</sub>,h<sub>1</sub>,i<sub>1</sub>,j<sub>1</sub>,k<sub>1</sub>], c<sub>1</sub>-612  $[d_1, f_1, g_1, h_1, i_1], d_1 - [e_1, h_1, i_1, j_1, k_1], e_1[f_1, g_1, h_1, i_1], f_1 - [h_1, i_1, j_1, k_1], g_1 - [h_1, i_1, j_1, k_1], h_1 - [i_1, j_1, k_1], i - [j, k].$ 613 614 **MUFA:**  $a_2$ -[ $e_2$ , $h_2$ , $i_2$ , $j_2$ , $k_2$ ],  $b_2$ -[ $e_2$ , $h_2$ , $i_2$ , $j_2$ , $k_2$ ],  $c_2$ -[ $d_2$ , $f_2$ , $g_2$ , $h_2$ , $l_2$ , $j_2$ , $k_2$ ],  $d_2$ -[ $e_2$ , $h_2$ , $i_2$ , $j_2$ , $k_2$ ],  $e_2$ [ $f_2$ , $g_2$ , $h_2$ , $l_2$ , $j_2$ ], 615  $f_2$ - $[h_2, i_2, j_2, k_2]$ ,  $g_2$ - $[h_2, i_2, j_2, k_2]$ ,  $h_2$ - $[i_2, j_2, k_2]$ ,  $i_2$ - $[j_2, k_2]$ ,  $j_2$ - $k_2$ . **PUFA:**  $a_3$ - $[c_3, e_3, h_3, i_3, j_3, k_3]$ ,  $b_3$ - $[c_{3}, e_{3}, h_{3}, i_{3}, j_{3}, k_{3}], \quad c_{3}-[d_{3}, e_{3}, f_{3}, g_{3}, h_{3}, l_{3}, j_{3}], \quad d_{3}-[e_{3}, h_{3}, i_{3}, j_{3}, k_{3}], \quad e_{3}[f_{3}, g_{3}, h_{3}, l_{3}, k_{3}], \quad f_{3}-[h_{3}, i_{3}, j_{3}, k_{3}], \quad g_{3}-(h_{3}, h_{3}, h_{3},$ 616 [h<sub>3</sub>,i<sub>3</sub>,j<sub>3</sub>,k<sub>3</sub>], h<sub>3</sub>-[i<sub>3</sub>,j<sub>3</sub>,k<sub>3</sub>], i<sub>3</sub>-[j<sub>3</sub>,k<sub>3</sub>], j<sub>3</sub>-k<sub>3</sub>]. **c.** PUFA to MUFA ratio (P:S) in dietary fish [a<sub>0</sub>-617 618  $[c_0, e_0, g_0, h_0, i_0, j_0, k_0],$  $b_0-[c_0,e_0,f_0,g_0,h_0,i_0,j_0], \quad c_0-[d_0,e_0,f_0,g_0,h_0,l_0,j_0,k_0],$  $d_0$ -[ $e_0$ , $h_0$ , $i_0$ , $j_0$ , $k_0$ ],  $e_0$ -619 [f<sub>0</sub>,g<sub>0</sub>,h<sub>0</sub>,I<sub>0</sub>,k<sub>0</sub>], f<sub>0</sub>-[h<sub>0</sub>,i<sub>0</sub>,j<sub>0</sub>,k<sub>0</sub>], g<sub>0</sub>-[h<sub>0</sub>,i<sub>0</sub>,j<sub>0</sub>,k<sub>0</sub>], h<sub>0</sub>-[i<sub>0</sub>,j<sub>0</sub>,k<sub>0</sub>], i<sub>0</sub>-[j<sub>0</sub>,k<sub>0</sub>], j<sub>0</sub>-k<sub>0</sub>]. (Blackin founder

Fig 3 Heatmap visualization of free fatty acids and evaluation of nutritional indices of fish. **a.** Hierarchical cluster correlation analysis of free fatty acids, **b.** Index of atherogenicity [IA], **c.** Health promotion index [HPI], **d.** Hypo/Hyper-cholesterolemic ratio [HH], and e. Fish lipid quality [FLQ]. (Pacific cod (PC), saffron cod (SC), Lindberg skate (LS), blackin flounder (BF), Japanese sardine (JS), Japanese pond smelt (JP), red seabream (RB), flatfish (FF), shishamo smelt (SS), skipjack tuna (ST), and Okhotsk Atka mackerel (OM).

Fig 4 a. Extracted ion chromatograms of LNAPEs b. Hierarchical cluster correlation analysisof lyso-phospholipids detected in fish fillet samples.

Fig 5 The MS and MS/MS spectra of LNAPEs detected in various fish types [\*isomericmixture].

Fig 6 Hierarchical cluster correlation analysis of glycerophospholipids. a. PC and PS. b. PI,
PG, and CL c. sphingolipids, N-acyl ethanol amines, cholesterol esters, mono-and diacylglycerols. d. Triacylglycerols characterized in eleven fish types. (clustering method: ward,
distance measure: Euclidean)

# Figures

Fig 1











Fig 4



Fig 5







# **Graphical Abstract**

