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Author(s)	Awata, Satoshi; Sasaki, Haruka; Goto, Tomohito; Koya, Yasunori; Takeshima, Hirohiko; Yamazaki, Aya; Munehara, Hiroyuki
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## **Marine Biology**

# Host selection and ovipositor length in eight sympatric species of sculpins that deposit their eggs into tunicates or sponges --Manuscript Draft--

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Corresponding Author:	Satoshi Awata, Ph.D. Osaka City University Osaka, JAPAN		
Corresponding Author Secondary Information:			
Corresponding Author's Institution:	Osaka City University		
Corresponding Author's Secondary Institution:			
First Author:	Satoshi Awata, Ph.D.		
First Author Secondary Information:			
Order of Authors:	Satoshi Awata, Ph.D.		
	Haruka Sasaki		
	Tomohito Goto		
	Yasunori Koya, Ph.D.		
	Hirohiko Takeshima, Ph.D.		
	Aya Yamazaki, Ph.D.		
	Hiroyuki Munehara, Ph.D.		
Order of Authors Secondary Information:			
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Abstract:	Interspecific interactions between parasites and hosts can influence the evolution of behavioural and morphological adaptations of both parasites and their hosts. There is, however, little empirical evidence available regarding the evolution of reproductive traits driven by these interactions. In this paper, we investigated host selection and ovipositor length in nine sympatric marine sculpins that oviposit into tunicates or sponges. Field and genetic studies have revealed host use for eight out of nine species of sculpins investigated here: five species of Pseudoblennius, two species of Furcina and one species of Vellitor. For one species studied (V. minutus), no egg masses could be found. Ovipositor length reflects morphology of host species utilised: six sculpin		

### **Original paper**

## Host selection and ovipositor length in eight sympatric species of sculpins that deposit their eggs into tunicates or sponges

Satoshi Awata<sup>1,2</sup> · Haruka Sasaki<sup>1</sup> · Tomohito Goto<sup>1</sup> · Yasunori Koya<sup>3</sup> · Hirohiko Takeshima<sup>4,5</sup> · Aya Yamazaki<sup>6</sup> · Hiroyuki Munehara<sup>6</sup>

<sup>1</sup>Sado Marine Biological Station, Faculty of Science, Niigata University, Sado, Japan

<sup>2</sup>Laboratory of Animal Sociology, Graduate School of Science, Osaka City University, Osaka, Japan

<sup>3</sup>Department of Biology, Faculty of Education, Gifu University, Gifu, Japan

<sup>4</sup>Research Institute for Humanity and Nature, Kyoto, Japan

<sup>5</sup>Department of Marine Biology, School of Marine Science and Technology, Tokai University, Shizuoka, Japan

<sup>6</sup>Usujiri Fisheries Station, Field Science Center for Northern Biosphere, Hokkaido University, Hakodate, Japan

Short title: Host selection and ovipositor length in sculpins

Correspondence

Satoshi Awata

Laboratory of Animal Sociology, Graduate School of Science, Osaka City University, 3-3-138 Sugimoto, Sumiyoshi, Osaka 558-8585, Japan.

Tel.: +81 6 6605 2607

e-mail: sa-awata@sci.osaka-cu.ac.jp

ORCID ID: 0000-0003-3254-7943

#### 1 Abstract

2 Interspecific interactions between parasites and hosts can influence the evolution of 3 behavioural and morphological adaptations of both parasites and their hosts. There is, 4 however, little empirical evidence available regarding the evolution of reproductive traits 5 driven by these interactions. In this paper, we investigated host selection and ovipositor length 6 in nine sympatric marine sculpins that oviposit into tunicates or sponges. Field and genetic 7 studies have revealed host use for eight out of nine species of sculpins investigated here: five 8 species of *Pseudoblennius*, two species of *Furcina* and one species of *Vellitor*. For one species 9 studied (V. minutus), no egg masses could be found. Ovipositor length reflects morphology of 10 host species utilised: six sculpin species had extremely long ovipositors allowing females to 11 attach eggs to the deep atrium of solitary tunicates, whereas the two species that attached their 12 eggs to the small space of atrial syphon of colonial tunicates and the spongocoel of sponges 13 had short ovipositors. Ovipositor length varied between solitary-tunicate spawners and species 14 with longer ovipositors selected larger tunicates. Since the ancestral form is non-parasitic, the 15 ovipositor evolved as an adaptation to utilise sponges and tunicates as hosts. Sculpins found 16 sympatrically may show host specificity to avoid interspecific competition for spawning 17 niches and ovipositors may have evolved depending on the species and size of host 18 invertebrates.

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Keywords: ovipositor; sculpins; tunicates; sponges; marine fish; ostracophils.

#### 20 Introduction

21 Sexual selection, arising from interaction between sexes, has considerable influence on 22 evolutionary adaptations in both sexes, for example, the morphological adaptation of 23 reproductive traits (Andersson 1994; Birkhead and Møller 1998). However, as yet only a few 24 studies have examined the evolution of reproductive traits in parasites through the interactions 25 between parasites and hosts. In parasitic animals that utilise other organisms as oviposition 26 hosts, female selection and oviposition success can strongly affect the survival of offspring 27 and thereby parental fitness (e.g. Smith et al. 2001; Mills and Reynolds 2002, 2003; Kitamura 28 2005; Reichard et al. 2007), whereas hosts often incur costs due to egg deposition (e.g. Peden 29 and Corbett 1973; Poltev and Mukhametov 2009; Reichard et al. 2006). Therefore, such 30 interspecifically antagonistic interactions can influence the evolution of behavioural and 31 morphological adaptations of either or both parasites and their hosts. 32 Some of the parasitic animals that inhabit aquatic environments preferentially deposit

33 their eggs in live invertebrates (known as 'ostracophils': Balon 1975; Leung 2014). For 34 example, bitterlings (Acheilognathinae) deposit their eggs in the gill chambers of freshwater 35 mussels through their atrial siphon (e.g. Smith et al. 2004; Kitamura 2007), the Japanese 36 tubesnout (Aulichthys japonicus) conceals their eggs in tunicates (Akagawa et al. 2008) and 37 snailfishes (Liparidae) deposit eggs within the branchial chambers of crabs (e.g. Hunter 1969; 38 Somerton and Donaldson 1998; Poltev and Mukhametov 2009; Gardner et al. 2016). All of 39 these species display some degree of host specificity, and various degrees of adaptation of 40 reproductive traits, for the deposition of their eggs inside invertebrates (reviewed in Leung 41 2014); for example, the extremely long ovipositors observed in both bitterlings (Smith et al. 42 2004; Kitamura 2007; Kitamura et al. 2012) and snailfishes (Hunter 1969). Similar 43 adaptations of reproductive traits through interactions between parasites and hosts have been

reported for various parasitic wasps including parasitoids that target free-living insects (Le
Ralec et al. 1996), gall-makers (Ghara et al. 2011) and figs (Elias et al. 2018). These
ostracophilous fishes and parasitic wasps have shown that sexual traits can be inter- and
intraspecifically variable, due to interactions between parasites and hosts. However, empirical
evidence regarding these variations in reproductive traits and host selection remain scarce.

49 Sculpins (Teleostei: Cottidae), which belong to the Cottoidea superfamily, are a large 50 group of fish that have 70 genera and 282 species worldwide, and primarily inhabit boreal and 51 cold temperate regions (Nelson et al. 2016). Although evidence is scarce, several species of 52 Cottoidea are known to deposit their eggs inside alive invertebrates (reviewed in Abe and 53 Munehara 2009; Munehara et al. 2011). For example, the little dragon sculpin *Blepsias* 54 *cirrhosus* and the bigmouth sculpin *Hemitripterus bolini* utilise sponges as their spawning 55 host (Munehara 1991; Busby et al. 2012). It has also been reported that four sculpin species in 56 the genus *Pseudoblennius* that inhabit temperate waters seem to be ostracophils (Uchida 1932, 57 1979; Shinomiya and Ikemoto 1987; Okamura and Amaoka 1997; Nishida et al. 2008; Uryu 58 2011). According to these descriptive studies and to underwater photographs taken by divers, 59 these species are all copulating species in which females deposit their eggs exclusively inside 60 tunicates or sponges using their ovipositors (Shiogaki and Dotsu 1974; Shinomiya 1985; Abe 61 and Munehara 2009; Nishida et al. 2008; Uryu 2011). In addition, during a preliminary study 62 of marine sculpins in the nearshore reef of Sado Island, located in the Sea of Japan, it was 63 found that there may be nine species of potentially ostracophilous sculpins (five species of 64 *Pseudoblennius*, two species of *Furcina* and two species of *Vellitor*). They all develop 65 ovipositors and occur sympatrically in shallow water of the island (Awata 2015, 2017). 66 Therefore, these sculpins may be ideal for an examination of the evolution of interspecific 67 variation in reproductive traits that are driven by interactions between parasites and hosts. 68 However, no studies have demonstrated host selection and ovipositor morphology in sculpins,

with the exception of one study that showed host tunicate selection by *Pseudoblennius percoides* (Nishida et al. 2008).

71 In this study, we investigated interspecific variation in host selection and ovipositor 72 length in nine sympatric species of marine sculpins that deposit their eggs into tunicates or 73 sponges on the coast of Sado Island, northern Japan. We hypothesise that species-specific host 74 selection by sculpin females drives the evolution of ovipositor morphology. To confirm our 75 hypothesis, we tested three predictions. Firstly, that the nine ostracophilous sculpin species 76 would utilise different species and sizes of host tunicates or sponges to reduce interspecific 77 competition for hosts (Reichard et al. 2007). This is because these sculpins co-occurred at the 78 study sites. Secondly, that only ostracophilous sculpins, and not species that lay eggs on rock 79 surfaces, would have elongated ovipositors. The morphological and molecular phylogeny of 80 the sculpins suggests that ostracophilous sculpins have evolved from non-ostracophilous 81 sculpins (Yabe 1985; Abe and Munehara 2009; Koya et al. 2011; Knope 2013). Therefore, the 82 development of the ovipositor is likely related to such unique spawning behaviour. Finally, 83 that ovipositor length would reflect the species and size of the hosts utilised.

#### 84 Materials and methods

#### 85 Study sites

86 Fieldwork using self-contained underwater breathing apparatus (SCUBA) was conducted on

- the coast of Sado Island in the Sea of Japan. The island is located about 45 km off the
- 38 Japanese mainland coastal city of Niigata (Fig. 1a). The amplitude of the tides around the
- island is relatively low, ranging from only 0.2 to 0.3 m. Four sites (Fig. 1b: Kitakoura: 38° 16′
- 90 26.90" N, 138° 30' 34.42" E; Mushizaki: 38° 14' 58.66" N, 138° 30' 24.37" E; Tassya: 38° 04'
- 91 28.64" N, 138° 14' 40.81" E and; Kotoura: 37° 48' 13.43" N, 138° 15' 08.26" E) were selected

for fieldwork. All of the host invertebrates and ostracophilous sculpin species were abundant
at the study sites, with the exception of *V. minutus* that was only found at Tassya (Awata
2017). The four study sites were located on rocky shores where seaweed grows abundantly.

#### 95 Fish study species

96 There were nine species of potentially ostracophilous sculpins in the nearshore reef of Sado 97 Island (Nakabo and Kai 2013; Awata 2015, 2017). Microhabitat preferences differ between 98 species (Iwata 1983; Awata 2017), but all species occurred sympatrically in rocky shallow 99 water ranging in depth from 1 to 20 m. They are copulating species and their fertilisation 100 mode is internal gametic association (IGA: Munehara et al. 1989), where fertilisation occurs 101 only when eggs are released into the seawater (Shiogaki and Dotsu 1974; Kimura et al. 1987, 102 1988; Koya et al. 2015). Eggs that are released into seawater are spherical of 1.3–2.0 mm in 103 diameter, demersal and adhesive, forming egg masses of up to 500 eggs (e.g. Kimura et al. 104 1987, 1988; Nishida et al. 2008). There are nine species in three genera, Furcina, 105 Pseudoblennius and Vellitor, which form a closely related and monophyletic group (Yabe 106 1985; Munehara et al. 2011). The reproductive ecology is similar among species, but their 107 body size and shape are notably different: Pseudoblennius sp. ("Kirin-anahaze" in Japanese, 108 see Nakabo and Kai 2013) and Ps. percoides are large sized (130-170 mm in standard length, 109 SL), Ps. cottoides, Ps. marmoratus and Ps. zonostigma are medium sized (80–100 mm SL) 110 and F. ishikawae, F. osimae, V. centropomus and V. minutus are small sized sculpins (40-70 111 mm SL; Okamura and Amaoka 1997; Munehara et al. 2011; Nakabo and Kai 2013; Awata 112 2017).

#### 113 Fish collection and measurement of ovipositor length

114 To examine the ovipositor length of the nine species of ostracophilous sculpins, adult females

115 were captured using hand-nets while SCUBA diving during the spawning season, from late 116 November to late April in 2014, 2015 and 2016, in depth of 2 to 15 m at all four study sites 117 (Fig. 1b). The fish were transported in aerated coolers to the laboratory at the Sado Marine 118 Biological Station (SMBS), Niigata University, Sado, Japan. Species identification of sculpins 119 was based on that of Nakabo and Kai (2013). Fish were anaesthetised with a solution of 120 MS222 or 2-phenoxyethanol buffered with seawater, and their SL (to 1 mm) was measured 121 using a scale. The ovipositor was induced to its maximum extrusion by gently pressing the 122 belly of the fish between ventral fins and anus. The ovipositor was photographed alongside a 123 scale, and ovipositor length (OL) was measured to the nearest 1 mm from the apex to the 124 ventral edge of the ovipositor using image analysis software ImageJ 1.50i. A part of the right 125 pectoral fin was clipped from each individual and preserved in 99 % ethanol for genetic 126 analysis. Of the 124 females analysed, 40 were released after the operation and 84 were used 127 for other studies after sacrification under deep anaesthesia (taxonomy and sperm analyses: 128 Awata 2015, 2017; histology: Koya et al. 2015).

To determine whether only ostracophilous sculpins have evolved elongated
ovipositors, we compared the OL of ostracophilous sculpins and sculpin species that oviposit
on the rock surface. We captured 13 females of four non-ostracophilous sculpin species
(*Alcichthys alcicornis, Bero elegans, Icelinus pietschi* and *Radulinopsis derjavini*) using
SCUBA at Usujiri (41° 56′ 11.85″ N, 140° 56′ 54.66″ E; Fig. 1a), southern Hokkaido, Japan
and at Kotoura, Sado Island, Japan, and measured their SL and OL.

#### 135 Collection of egg masses deposited in host invertebrates

136 Egg masses deposited in the spongocoels of sponges and in the common atrial syphons of

- 137 colonial tunicates were collected in April and May 2012, from March to May 2013 and from
- 138 February to April 2014 at Kotoura and Kitakoura (Fig. 1b). Egg masses were externally

visible, and therefore only whole colonies of sponges and colonial tunicates that contained
egg masses were collected using scrapers. The colonies that contained eggs were distributed
across rock wall habitats at depths of 1–12 m. The sea water temperature was 9–14 °C.

142 Because the egg masses deposited in the atria of solitary tunicates were rarely 143 visible from outside, we haphazardly collected solitary tunicates (n = 1212) in December 144 2011, December 2012 and from December 2013 to May 2014 (approximately once a month) 145 at Kitakoura, Mushizaki and Tassya (Fig. 1b). Of these, 1129 (93 %) tunicates were captured 146 from the wall of Kitakoura Port at a depth of 0.5–4.0 m (Fig. 1c), where the wall was covered 147 with in excess of a hundred thousand solitary tunicates. The remaining samples were collected 148 at 8–18 m depth of off Kitakoura Port, at 8–30 m depth of Mushizaki and at 2–5 m of Tassya. 149 The sea water temperature ranged from 9 to 16 °C (the coldest being in February).

150 All tunicates, sponges and egg masses were transferred to the laboratory in aerated 151 coolers. Sponges and colonial tunicates were cut from the osculum and common atrial siphon, 152 respectively, to the spongocoel and common atrium. The length between the open top and the 153 top of the egg mass was measured using a scale (defined as "egg mass position"). Solitary 154 tunicates were photographed with a scale and their long diameter (LD) and short diameter 155 (SD) were later measured using ImageJ 1.50i software. After being photographed, tunicates 156 were carefully cut in half vertically using a kitchen knife, in order to avoid damaging the egg 157 masses. When egg masses were found inside tunicates, opened tunicates were photographed 158 again with a scale to measure the length from entrance of the atrial siphon to the nearest edge 159 of the egg masses (also defined as "egg mass position"). It is natural to expect that utilisation 160 of larger tunicates will require longer ovipositors, and thus egg mass position will be deeper 161 when larger tunicates are used as hosts. The LD was used as an indicator of the body size of 162 tunicates, given that their LD and SD are strongly correlated (Pearson's r = 0.92, p < 0.0001, 163 *n* = 1212).

Each egg mass removed from the host was counted and transferred to a box-shaped net cage ( $12.5 \times 13.0 \times 8.0$  cm, mesh size: 0.5 mm, s-5330, Sudo & Company, Inc., Nagoya, Japan) that was fixed to the surface of a  $90 \times 45 \times 45$  cm tank at the SMBS. Each egg mass was separately incubated for 1–20 days in the cage at 15 °C. Hatched larvae or late stage eggs (just before hatching) were preserved in 99 % ethanol after being killed by an overdose of anaesthesia.

Note that egg masses of the Japanese tubesnout *Aulichthys japonicas* were also
collected during this study, but the details of this will be published in a later paper.

#### 172 Genetic identification of parent species of egg masses

173 To infer the parental species of the egg masses, 1–3 larvae per egg mass and fin-clip samples 174 of nine potentially ostracophilous sculpins, with the addition of Jordania zonope as an 175 outgroup, were chosen for use in genetic analyses. Fin-clip samples were obtained from 2011-176 2017 (Table S1, Supplementary Information). Whole genomic DNA was extracted from 148 177 adults and from 223 larvae (including late stage eggs) of 120 egg masses, using a Gentra 178 Puregene Tissue Kit (Qiagen, Hilden, Germany). Samples were PCR amplified for the 179 mitochondrial cytochrome b gene (cytb): GLUDG-L: 5'-YGACTTGAARAACCAYCGTTG-180 3' and CB3-H: 5'-GGCAAATAGGAARTATCATTC-3' (Palumbi 1996; Knope 2013). PCR 181 amplifications were performed at a 15  $\mu$ L volume using 7.5  $\mu$ L of 2 × Go Taq Green Master 182 Mix (Promega, WI, USA), 0.6  $\mu$ L (5  $\mu$ M) of each primer, 1.5  $\mu$ L ca. 50 ng/ $\mu$ L of template 183 DNA and 4.8 µL of nuclease free water. PCR thermal cycling was performed using the 184 following protocol for cytb: 2 min of initial denaturation at 95 °C, followed by 35 cycles of 185 94 °C for 30 sec, 50 °C for 30 sec and 72 °C for 1 min. PCR amplifications were successful 186 except for some *Ps. cottoides* and *Ps. zonostigma* and many of the egg masses, for which 187 sequences corresponding to the primers were permutated. Therefore, these samples were PCR

188 amplified using another primer set for cytb: H15915: 5'-CAACGATCTCCGGTTT-3' and 189 L14724: 5'-GTGACTTGAAAAACCA-3' (Schmidt and Gold 1993). The 800 bp sequences 190 amplified by this primer set overlapped with those amplified by the former primer set. PCR products were electrophoresed on 1.5 % agarose gel stained with GelRed<sup>TM</sup> (Biotium, Inc., 191 192 CA, USA). Positive amplicons (evidenced by a clear single band of the correct size) were enzymatically cleaned with illustra<sup>TM</sup> ExoStar<sup>TM</sup> (GE Healthcare UK Limited, Little Chalfont, 193 194 Buckinghamshire, UK). Double stranded PCR products were sequenced directly with the 195 same primers used for the PCR amplifications at Macrogen Japan Corp. (Kyoto, Japan). All 196 sequences were aligned using the ClustalW algorithm implemented in MEGA ver. 7.0.14. 197 (Kumar et al. 2016). 198 The phylogenetic relationships of nine potential parental species (n = 146) and 199 Jordania zonope (n = 2, as the root for the tree) were reconstructed using 800 bp cytb 200 sequences with MEGA ver. 7.0.14, and the Maximum Likelihood method was employed

201 using the Tamura Nei model settings (Tamura and Nei 1993) and 1000 bootstrap replicates.

202 After assessing the usefulness of the sequences for species identification (Fig. S1,

203 Supplementary Information; all these sequences available on GenBank, Table S1), the

204 parental species of egg masses were estimated using the reconstructed phylogeny of the

205 parental species and egg masses. Of 223 eggs analysed, 205 were identified using 800 bp cytb

sequences, and 18 were identified using 628 bp, due to the low quality of the sequencing data.

#### 207 Data analyses

208 Statistical analyses were performed using SPSS ver. 23.0 (IBM Corp., Armonk, NY, USA).

209 Data were analysed using parametric statistics throughout. All reported *p*-values were two

tailed, and results were considered statistically significant at p-values of < 0.05. As we did not

211 obtain all data from all sponges and tunicates, sample size varied between analyses. To

determine whether the relative OL differed between ostracophilous and non-ostracophilous sculpins, we used an analysis of covariance (ANCOVA) with log-transformed OL as the dependent variable and deposition site (sponge, colonial tunicate, solitary tunicate and rock surface) as the independent variable and log-transformed SL as the covariate, including their interaction (full model). If the interaction term (deposition site  $\times \log$  SL) was not statistically significant, then a reduced model was constructed by reducing the interaction term.

#### 218 **Results**

#### 219 Egg masses deposited in sponges and colonial tunicates

Nine egg masses were obtained from sponges (Demospongiae) of nine colonies. Since each colony was small and formed mostly only one spongocoel, multiple egg masses were not found in sponges sampled (Fig. 2a). In contrast, the colony size and shape of colonial tunicates (Didemnidae) were variable, ranging from 20 to 100 mm in maximum width, each of which had 1–38 exhalant syphons and contained 1–5 sculpin egg masses at similar or different developmental stages (Fig. 2b, c). In total, 34 egg masses were obtained from the colonial tunicates of 22 colonies.

#### 227 Egg masses deposited in solitary tunicates

- 228 Of 1212 solitary tunicates collected, seven species were identified. The most commonly
- 229 occurring was *Halocynthia ritteri* (55.9 %, *n* = 678), followed by *Polycarpa cryptocarpa*
- 230 *kroboja* (25.3 %, *n* = 307), *Cnemidocarpa irene* (11.6 %, *n* = 141), *Pyura sacciformis* (3.5 %,
- 231 n = 42), *Microcosmus hartmeyeri* (2.0 %, n = 24), *H. roretzi* (1.6 %, n = 19) and *Herdmania*
- *japonica* (0.1 %, n = 1). Although a considerable number of tunicates were collected, sculpins
- 233 only utilised three species (Fig. 2d–h). The two largest and the smallest species of tunicates

were not used by the sculpins (Fig. 3). The second most abundant tunicate *Po. c. kroboja* was
never used by sculpins, although notably overlapped in size with the most abundant and most
frequently used tunicate *H. ritteri* (Fig. 3).

Of the 89 sculpin egg masses collected, > 90 % were found in *H. ritteri* and the remainder were found in *Py. sacciformis* and *H. roretzi* (Fig. 2f–h; Fig. 3). Although the abundance of *H. ritteri* was more than 16 times greater than that of *Py. sacciformis* at the study sites, the proportion of parasitized tunicates was equal ( $n_{with eggs}/n_{without eggs} = 75/593$  for *H. ritteri*; 6/30 for *Py. sacciformis*: Fisher's exact test, p = 0.29). Note that sculpin eggs were only found in the atria of tunicates, indicating that the ovipositors were inserted through the atrial syphon of the tunicates.

#### 244 **Parental species of egg mass**

245 The reconstructed phylogenetic relationships of the potentially parental species and 120 egg masses showed that sponges were utilised by *Ps. marmoratus* (n = 4 egg masses, Fig. 2a) and 246 247 V. centropomus (n = 4) as spawning hosts (details in Table S2). All 26 egg masses deposited in 248 colonial tunicates were assigned to V. centropomus (Fig. 2b). Solitary tunicates were used by 249 six sculpin species (Table 1; Fig. 2d-h; details in Table S3): egg masses of Ps. cottoides were 250 the most frequent (n = 52 egg masses), followed by those of *Ps. zonostigma* (n = 13), 251 *Pseudoblennius* sp. "Kirin-anahaze" (n = 9), *Ps. percoides* (n = 8), *F. osimae* (n = 3) and *F.* 252 *ishikawae* (*n* =1). Five of six species mainly employed *H. ritteri* as their hosts (Table 1; Fig. 253 2f). Pseudoblennius cottoides and Ps. percoides also utilised Py. sacciformis. Egg masses 254 deposited in *H. roretzi* were identified as belonging to *Pseudoblennius* sp. "Kirin-anahaze" 255 (Fig. 2g). Furcina osimae only utilised Py. sacciformis as the tunicate host (Fig. 2h). No egg 256 masses were genetically assigned to *V. minutus* and so further analyses excluded this species.

#### 257 Egg mass position in different hosts

258 Since egg masses were visible from the outside of sponges and colonial tunicates, but were 259 not for solitary tunicates, the egg mass position would be expected to differ between the hosts. 260 As expected, the egg masses were deposited significantly deeper in solitary tunicates (16.44  $\pm$ 261 6.39 mm, n = 73) than in sponges (2.70 ± 1.69 mm, n = 6) and colonial tunicates (2.02 ± 1.79 262 mm, n = 11) (one-way ANOVA,  $F_{2,87} = 40.46$ , p < 0.0001; Tukey's HSD, p < 0.0001 for both). 263 The egg mass position did not differ between sponges and colonial tunicates (Tukey's HSD, p 264 = 0.97). This indicates that ovipositor length likely differs between sculpin species using 265 solitary tunicates and those using sponges and colonial tunicates.

#### 266 Comparison of ovipositor length between fish using different deposition sites

267 An ANCOVA analysis showed that the interaction between deposition site and log SL did not 268 have a significant effect on log OL (full model,  $F_{3,129} = 2.09$ , p = 0.11), and overall, larger 269 species/individuals had longer ovipositors (reduced model,  $F_{1,132} = 176.77$ , p < 0.0001; Fig. 270 4a). As predicted, the relative OL was significantly related to deposition site (reduced model, 271  $F_{3,132} = 557.23$ , p < 0.0001; Fig. 4a). The relative OL of the ostracophilous sculpins was 272 significantly longer than that of the non-ostracophilous sculpins (Fig. 4b-e). Moreover, the 273 sculpin species that use solitary tunicates as hosts had much longer OLs (ca. 20 % of SL) than 274 that of those using colonial tunicates and sponges. Although the relative OL differed 275 significantly between V. centropomus (ca. 5 % of SL) and Ps. marmoratus (ca. 3 % of SL), the 276 absolute OL did not differ between them (*t*-test, t = 0.99, p = 0.33; Fig. 4a–c).

#### 277 Ovipositor length and the size of solitary tunicates used by sculpins

278 The body size of the six parental species that utilised solitary tunicates as hosts differed

significantly among species (one-way ANOVA,  $F_{5,89} = 166.84$ , p < 0.0001; Fig. 5a). Female

280 *Pseudoblennius* sp. "Kirin-anahaze" were the largest, and two species in the *Furcina* genus 281 were the smallest of all six species. Overall, larger species had longer ovipositors ( $F_{5,89} =$ 282 28.05, p < 0.0001; Fig. 5b), but the OL of *Ps. percoides* was significantly shorter than that of

283 *Ps. cottoides* despite its larger body size (also see Fig. 4a).

284 As expected, egg mass position was deeper when larger tunicates were used as hosts 285 (Pearson's r = 0.49, p < 0.0001, n = 72). The LD of tunicates used as hosts and the egg mass 286 position were significantly different among sculpin species (LD: Fig. 5c; one-way ANOVA, 287  $F_{4.79} = 8.81 \ p < 0.0001$ ; egg mass position: Fig. 5d;  $F_{4.68} = 7.05, \ p < 0.0001$ ). Consistent with 288 the expectation that utilisation of larger tunicates will require longer ovipositors, the LD of 289 tunicates used as hosts and the egg mass position were closely related to the sculpin OL (Fig 290 5); Pseudoblennius sp. "Kirin-anahaze", Ps. cottoides and Ps. zonostigma that have longer 291 ovipositors spawned in a deeper position in larger tunicates, and the genus Furcina with a 292 smaller body size and a shorter ovipositor consequently utilised smaller tunicates.

293 *Pseudoblennius percoides*, which has a larger body size but a shorter ovipositor, selected
294 smaller tunicates as spawning hosts.

295 Pseudoblennius sp. "Kirin-anahaze", Ps. cottoides and Ps. zonostigma used similar-296 sized H. ritteri tunicates as hosts (Fig. 5c), indicating severe competitions for hosts among the 297 three species. However, if host H. ritteri is abundant and sculpins use the most common size 298 of tunicates, competition for hosts would be weakened. Indeed, H. ritteri was used by only 299 11 % of all individuals (Fig. 3d). Additionally, the average and variance of host size were 300 similar between unused H. ritteri (84.20 LD  $\pm$  20.31, n = 595) and that used by 301 *Pseudoblennius* sp. "Kirin-anahaze" (92.84 LD  $\pm$  10.37, n = 7, t-test, t = 1.12, p = 0.26; F test 302 to compare two variances, F = 3.84, p = 0.09), Ps. cottoides (87.16 LD ± 18.38, n = 46, t =303 0.96, p = 0.34; F = 1.22, p = 0.41) or Ps. zonostigma (although slightly larger tunicates were 304 used: 99.12 LD  $\pm$  17.37, n = 11, t = 2.42, p = 0.02; F = 1.37, p = 0.61). These show that these

305 three sculpins used the most common size of *H. ritteri* (Fig. 3d).

#### 306 Discussion

#### 307 Host specificity

308 In this study, we identified oviposition hosts in eight out of the nine ostracophilous sculpins 309 examined. While a number of researchers and divers have observed spawning behaviours of 310 some of these sculpins, such as Ps. marmoratus, Ps. percoides and Ps. zonostigma 311 (Shinomiya 1985; Okamura and Amaoka 1997; Abe and Munehara 2009; Uryu 2011), so far 312 only one study has reported host tunicate selection by *Ps. percoides* (Nishida et al. 2008). 313 Thus, to our knowledge, this is the first study to confirm the host species used by these eight 314 sculpin species, including *Ps. percoides*. There are three patterns of host use by sculpins, such 315 as species that mainly use colonial tunicates, those that only use sponges and those that only 316 use solitary tunicates for oviposition. This result mostly confirms our first prediction, that 317 ostracophilous sculpins would show host specificity. However, as found in ostracophilous 318 bitterlings (Reichard et al. 2007), sculpins also showed variation in their level of specificity. 319 Pseudoblennius marmoratus oviposited only in the spongocoel of sponges, but V. 320 centropomus utilised colonial tunicates and sponges for oviposition. Eggs of six sculpin 321 species were found in solitary tunicates, but three of these fishes used only a single tunicate 322 species and the others used two tunicate species. 323 Egg mass numbers for each sculpin species collected in this study mostly 324 corresponded to the abundance of parental species at the study sites (see Fig. 4, Table 1). 325 Although we captured most tunicates on the wall of Kitakoura Port, the consistency of

326 appearance frequencies between eggs and parental species implies that the six sculpin species

327 do not avoid the artificial habitat for spawning. Since eggs of *V. minutus* were not found and

only one egg mass of *F. ishikawae* was collected, further field sampling including different
sites and seasons are needed to clarify their host use.

330 Sponges are only distantly related to colonial tunicates (e.g. Dunn et al. 2014). Their 331 body structures are therefore notably different. However, they share morphological 332 similarities: both have myriad brachial apertures and shared, common cloacal apertures. These 333 similarities, especially the common structure of the cloacal apertures, may allow female V. 334 centropomus to utilise both colonial tunicates and sponges as hosts. To our knowledge, it is 335 unlikely that any other fishes utilise such distantly related taxa. Pseudoblennius marmoratus 336 only oviposit in sponges, which is consistent with previous underwater observations (Uryu 337 2011). It remains unclear whether *Ps. marmoratus* also utilises colonial tunicates. To confirm 338 this, more information on its spawning behaviour in the field needs to be gathered. These two 339 sculpin species have short and similarly sized ovipositors, suggesting that they are specialised 340 in ovipositing in colonial tunicates and sponges.

341 We found that sculpins used only three out of seven species of solitary tunicates as 342 hosts, even though many species of tunicates co-occurred at the study sites. There are three 343 possibilities to explain sculpin host selection: 1) the size of the tunicates, 2) the internal 344 structure of the tunicates and 3) chemical substances in the atria of the tunicates. First, 345 sculpins may select for tunicates of optimal size. Solitary tunicate spawners have developed 346 long ovipositors, which may allow the placement of their eggs in the atrium of tunicates via 347 the atrial siphon. The elongation of the ovipositor is limited by the cost associated with 348 growing and maintaining a long ovipositor. Sculpin females did not use the large tunicates M. 349 *hartmeyeri* and *Herdmania japonica*, probably due to the excessively long distance from the 350 entrance of atrial siphon to the atrium of the tunicate. In contrast, the small atrium and narrow 351 entrance of the atrial siphon of the smallest tunicate, C. irene, was too small to allow sculpins 352 to use them as hosts.

353 Second, the internal structure of tunicates may be related to host selection by 354 sculpins. Our data showed that the size range of unused tunicate *Po. c. kroboja* greatly 355 overlapped with that of the commonly used tunicate H. ritteri. The unused Po. c. kroboja 356 tunicates are more compressed compared with the H. ritteri tunicates (Awata S. pers. obs.), 357 suggesting that sculpins do not utilise Po. c. kroboja due to the small inner space of their atria. 358 Finally, substances in the atria of tunicates might also affect host use by female sculpins. A 359 number of tunicates produce chemical defences, such as high sulfuric acid and high vanadium 360 content, which may reduce their predation risks (e.g. Stoecker 1980; Paul and Puglisi 2004; 361 Paul et al. 2006). Polycarpa cryptocarpa kroboja has a uniquely strong smell (Awata S. pers. 362 obs.) compared to the other tunicate species dissected. Although the chemical components 363 responsible for this smell were not identified, they may have adverse effects on the fish eggs. 364 Our aquarium experiments have demonstrated that when *Ps. cottoides* females with ripe eggs 365 were allowed to choose *H. ritteri* and *Po. c. kroboja* of the similar size, they spent more time 366 inspecting H. ritteri, and eggs were deposited only in H. ritteri (Awata S, Igarashi N, unpub. 367 data). This indicates that the female sculpins likely discriminate between H. ritteri and Po. c. 368 kroboja prior to spawning, and clearly prefer the former over the latter as oviposition hosts. It 369 remains to be experimentally determined which combinations of these proximate factors such 370 as tunicate size, structure and chemical substances affect host selection by female sculpins.

#### 371 **Ovipositor length**

372 In accordance with our second prediction, only ostracophilous sculpins had elongated

373 ovipositors, while non-ostracophilous sculpins lacked this feature. Since non-ostracophilous

374 sculpins have been suggested to be the ancestral state (Yabe 1985; Koya et al. 2011;

375 Munehara et al. 2011; Knope 2013), we can conclude that the elongated ovipositor has

376 evolved as an adaptation to the ostracophilous reproduction strategy. The ovipositor in species

377 using solitary tunicates as hosts was much longer than that in species using colonial tunicates 378 and sponges. The position of eggs oviposited in colonial tunicates and sponges were as short 379 as they were observed to be from the outside, and perfectly matched the short ovipositor 380 length of V. centropomus and Ps. marmoratus. In contrast, sculpins that oviposited in solitary 381 tunicates had extremely long ovipositors, which in turn matched the distance from the 382 entrance of the atrial siphon to the edge of the egg masses. These results are consistent with 383 the third prediction, that ovipositor length would reflect the species and size of hosts utilised. 384 Both the branchial and atrial siphons of solitary tunicates close immediately by 385 ejecting the water from their body when they detect external stimuli such as vibrations in the 386 water. In contrast, sponges and colonial tunicates shrink a little after the detection of such 387 stimuli (Awata S. pers. obs.). In response to such host behaviour, female sculpins that use 388 solitary tunicates as hosts insert their ovipositors rapidly into the atrium (Movie S1, S2, 389 Supplementary Information). In contrast, species that use sponges as hosts oviposit slowly 390 (Movie S3). Based on these observations, we suggest that spawning behaviour and ovipositor

391 length may have adapted to suit the different hosts.

#### 392 Competition among sculpin species over hosts

393 In this study, six species of sculpins were determined to use solitary tunicates as hosts. 394 Furcina osimae only used the tunicate Py. sacciformis as a spawning host, but the other five 395 species mainly deposited their eggs into H. ritteri. This suggests these five species did not 396 prefer specific host tunicates. However, their host use may be also related to the size and 397 species of the host tunicates. Indeed, two smallest species of genus *Furcina*, which have 398 shorter ovipositors, used smaller tunicates; the largest *Pseudoblennius* sp. "Kirin-anahaze", 399 which has longer ovipositors, spawned in larger tunicates. In contrast, the ovipositor of *Ps*. 400 *percoides* was shorter than that of *Ps. cottoides*, despite its larger body, and matches the size

401 of the host tunicates. Medium-sized sculpins such as *Ps. cottoides* and *Ps. zonostigma* had
402 medium-sized ovipositors but used larger tunicates as hosts. Thus, as expected from our first
403 prediction, sculpins likely choose different sized hosts that fit their ovipositor size, probably
404 to reduce interspecific competition for hosts (Reichard et al. 2007).

405 However, three sculpins (Pseudoblennius sp. "Kirin-anahaze", Ps. cottoides and Ps. 406 zonostigma) showed similar host choice in terms of species and size, indicating that 407 interspecific competition for hosts may be unavoidable (Kitamura 2007; Reichard et al. 2007). 408 Previous studies have shown that host utilisation by female bitterlings may be determined by 409 host availability, as well as by host preference and interspecific competition among bitterlings 410 (Kitamura 2007; Reichard et al. 2007). In this study, the most abundant host, *H. ritteri*, was 411 used by only 11 % of all individuals. Furthermore, these three sculpins used the most common 412 size of tunicates (65–105 mm in LD of host H. ritteri; see Fig. 3d), although Ps. zonostigma 413 oviposited into somewhat larger tunicates. Therefore, their host preferences are strongly 414 related to host availability, and the ovipositor length of those three species may have been 415 shaped by the size of the most abundantly available tunicates. Further experimental studies 416 are needed to identify the "real" host size and species preference by controlling for host 417 availability and interspecific competition among ostracophilous sculpins (Reichard et al. 418 2007). Since most *Pseudoblennius* species are piscivores (Okamura and Amaoka 1997; Uryu 419 2011), it would be necessary to take the risk of predation of smaller species and individuals by 420 larger ones into account when determining interspecific competition at the spawning sites.

#### 421 Costs and benefits for ostracophilous sculpins

422 There are at least three potential benefits for female sculpins that use tunicates and sponges as

423 oviposition hosts. First, the hosts may provide suitable environments for embryonic

424 development (Hunter 1969; Munehara 1991; Spence and Smith 2013). Both tunicates and

425 sponges are filter feeders that draw water and food particles through incurrent siphon or 426 canals and excrete the filtered water through atria and excurrent siphons, or through 427 spongocoels and oscula. Parental care by fishes that exclusively exhibit care for their eggs 428 mainly consists of fanning and mouthing, which are assumed to increase the oxygen supply 429 and to help to remove dead eggs and dust particles (e.g. Keenleyside 1991). Since filtered 430 water constantly flows through the atria of tunicates and the spongocoels of sponges where 431 the eggs are deposited, the hosts may inadvertently serve a similar function to fanning and 432 mouthing for the eggs. Furthermore, sponges may provide another benefit to egg development 433 in sculpins. Many studies have reported that sponges contain compounds with antibacterial 434 and antifungal properties (e.g. reviewed in Mehbub et al. 2014). Considering the absence of 435 dead eggs in sponges, the antibacterial and antifungal activities of sponges may increase the 436 survival rate of sculpin eggs. Although dead, un-decayed eggs of the little dragon sculpin B. 437 *cirrhosus* have been found to be present in sponge tissues (Munehara 1991), such eggs were 438 not found in our study, perhaps due to the difference in embryonic periods (< 1 month in our 439 study species and > 8 months in *B. cirrhosis*; Munehara 1991). Rose bitterling embryos 440 developing on the gills of freshwater mussels are known to gain nutrients and oxygen from 441 the water taken in by the mussels (Spence and Smith 2013). However, this may not be the 442 case in sculpins, because their eggs are placed where few or no nutrients may be available due 443 to the filtered water, and larvae are likely to leave their host soon after hatching since they are 444 capable of active swimming (Kimura et al. 1987, 1988; Awata S. pers. obs.).

Second, eggs are likely protected from predators by being hidden inside the hosts
(Hunter 1969; Munehara 1991). Since eggs are completely hidden in solitary tunicates that
have rigid bodies, predation avoidance may be highly effective. Furthermore, many species of
sponges are chemically defended from predation by marine animals, such as fishes, sea
urchins, sea stars and hermit crabs (e.g. reviewed in Paul and Puglisi 2004; Paul et al. 2006).

Thus, oviposition in sponges may also be effective for predation avoidance due to their
chemical defences. Third, females are not required to provide parental care, which may allow
them to utilise their energy reserves for further reproduction, such as increasing egg numbers
and size.

454 However, it is costly to be ostracophilous. First, the development of elongated 455 ovipositors and unique spawning behaviour are essential. Second, during the spawning 456 season, sculpin females must seek out suitable, limited oviposition hosts, which might 457 increase the probability of predation and consume energy reserves. Despite such costs, the 458 benefits for female sculpins specialising in the use of tunicates and sponges outweigh the 459 costs, which may be the evolutionary forces that produce and maintain the unique spawning 460 behaviours observed in sculpins. This also may be the case for other ostracophilous fishes 461 such as bitterlings, snailfishes and Japanese tubesnouts (Leung 2014).

#### 462 **Costs and benefits for hosts**

463 In contrast to the potential benefits obtained by the sculpins, it is unlikely that sculpins and 464 their eggs have positive impacts on the host tunicates and sponges. Conversely, the eggs 465 deposited by sculpins may adversely affect their hosts. One possible cost for the hosts 466 harbouring sculpin egg masses is the reduction of sea water exchange, as observed in mussels 467 (Smith et al. 2001; Mills and Reynolds 2002, 2003; Kitamura 2005). In many cases, one egg 468 mass was found in a solitary tunicate, and occupied only a small space in the atrium. 469 However, the eggs, especially those oviposited by larger sculpin species such as 470 Pseudoblennius sp. "Kirin-anahaze", sometimes entirely fill up the tunicate atrium due to 471 their large clutch size (see Table S3). In this case, the hosts cannot effectively exchange 472 seawater and consequently might incur some costs such as lowered growth and survival from 473 the overload of fish eggs. Such an overload of fish eggs has been reported for the snailfish-

474 crab system, where the load of the egg masses in the gill cavity of crabs deposited by snail 475 fish caused severe damage to their gills (Love and Shirley 1993; Somerton and Donaldson 476 1998; Poltev and Mukhametov 2009). Spence and Smith (2013) have also found that some 477 host mussels with many bitterling eggs died, probably due to severe competition for oxygen 478 between the host mussels and bitterling embryos. Since sculpin eggs are not oviposited on the 479 gills of tunicates through the incurrent siphon, sculpin eggs unlikely do cause severe damage 480 to the respiratory system of sponges and tunicates. Experimental work is needed to ascertain 481 the cost to the hosts of sculpin eggs.

#### 482 Conclusions

483 In summary, we have shown host selection and variation of ovipositor length in eight out of 484 nine sympatric marine sculpins found off the shore of an island in the Sea of Japan. To date, 485 ostracophilous behaviour has only been described for a small number of fish species, and 486 therefore only limited data on the behavioural and morphological adaptations of parasites and 487 hosts are available, with the exception of the bitterlings. A more recent study, however, has 488 found evidence of ostracophilous behaviour in the tubenose poacher *Pallasina barbata* 489 (Momota and Munehara 2017). Sculpins and tubenose poachers are different fish groups from 490 other ostracophilous fishes (Leung 2014), suggesting that ostracophilous reproductive strategy 491 has likely evolved in parallel across different taxa (Leung 2014). This study has shown the 492 morphological adaptations of sculpin females at the interspecific level. However, ongoing 493 research will need to examine the intraspecific variation in ovipositor size and morphology, 494 depending on the host species used and their size. Studying ostracophils at the population 495 level might be particularly worthwhile to identify the (co-)evolution of behavioural and 496 morphological adaptations of parasites and their hosts.

#### 497 Ethical statement

498 The research reported in this study was carried out in accordance with Animal Care and Use 499 Committees at Niigata University and Osaka City University. All of the procedures described 500 above meet the ABS/ASAB guidelines for the ethical treatment of animals. Research 501 permission was obtained from Fisheries Cooperative Association of Sado, Japan.

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#### 514 **References**

Akagawa I, Hara M, Iwamoto T (2008) Egg concealment in tunicates by females of the
Japanese tubesnout, *Aulichthys japonicus* (Gasterosteiformes), and its subsequent
copulation. Ichthyol Res 55:85–89

Abe T, Munehara H (2009) Adaptation and evolution of reproductive mode in copulating
 cottoid species. In: Jamieson BGM (Ed.) Reproductive Biology and Phylogeny of

520 Fishes (Agnathans and Bony Fishes). Science Publishers, Enfield, pp 221–246 521 Andersson M (1994) Sexual selection. Princeton Univ. Press, Princeton 522 Awata S (2015) Diversity and evolution of reproductive strategies in marine sculpins. 523 Aquabiol 37:614-621 524 Awata S (2017) Taxonomic and ecological notes on marine sculpins on the coast of Sado 525 Island in the Sea of Japan. Nat Hist Sado Isl 5:9–20 526 Balon EK (1975) Reproductive guilds of fishes – proposal and definition. J Fish Res Board 527 Can 32:821-864 528 Birkhead TR, Møller AP (1998) Sperm competition and sexual selection. Academic Press, 529 London 530 Busby MS, Blood DM, Fleischer AJ, Nichol DG (2012) Egg deposition and development of 531 eggs and larvae of bigmouth sculpin (Hemitripterus bolini). Northwest Nat 93:1-16 532 Dunn CW, Giribet G, Edgecombe GD, Hejnol A (2014) Animal phylogeny and its 533 evolutionary implications. Annu Rev Ecol Evol Syst 45:371-395 534 Elias LG, Kjellberg F, Farache FHA, Almeida EAB, Rasplus J-Y, Cruaud A, Peng Y-Q, Yang 535 D-R, Pereira RAS (2018) Ovipositor morphology correlates with life history evolution 536 in agaonid fig wasps. Acta Oecol 90:109–116 537 Gardner JR, Orr JW, Stevenson DE, Spies I, Somerton DA (2016) Reproductive parasitism 538 between distant phyla: molecular identification of snailfish (Liparidae) egg masses in 539 the gill cavities of king crabs (Lithodidae). Copeia 104:645-657 540 Ghara M, Kundanati L, Borges RM (2011) Nature's Swiss Army knives: ovipositor structure 541 mirrors ecology in a multitrophic fig wasp community. PLoS ONE 6:e23642 542 Hunter CJ (1969) Confirmation of symbiotic relationship between liparid fishes (Careproctus 543 spp.) and male king crab (Paralithodes camtschatica). Pac Sci 23:546-547 544 Iwata A (1983) A revision of the cottid fish genus Vellitor. Jap J Ichthyol 30:1–9 545 Keenleyside MHA (1991) Parental care. In: Keenleyside MHA (Ed.) Cichlid fishes: 546 behaviour, ecology and evolution. Chapman and Hall, London, pp 191–208 547 Kimura S, Tsumoto K, Mori K (1987) Development of eggs, larvae and juveniles of the cottid 548 fish, Pseudoblennius cottoides, reared in the laboratory. Jap J Ichthyol 34: 346-350 549 Kimura S, Tsumoto K, Mori K (1988) Development of the cottid fish, Pseudoblennius 550 percoides, reared in the laboratory, with brief descriptions of juvenile P. marmoratus 551 and P. zonostigma. Jap J Ichthyol 35:19-24 552 Kitamura J (2005) Factors affecting seasonal mortality of rosy bitterling (Rhodeus ocellatus 553 kurumeus) embryos on the gills of their host mussel. Popul Ecol 47:41-51 554 Kitamura J (2007) Reproductive ecology and host utilization of four sympatric bitterling 555 (Acheilognathinae, Cyprinidae) in a lowland reach of the Harai River in Mie, Japan. 556 Environ Biol Fish 78:37-55 557 Kitamura J, Nagata N, Nakajima J, Sota T (2012) Divergence of ovipositor length and egg 558 shape in a brood parasitic bitterling fish through the use of different mussel hosts. J Evol 559 Biol 25:566-573

560 Knope ML (2013) Phylogenetics of the marine sculpins (Teleostei: Cottidae) of the North 561 American Pacific Coast. Mol Phylogenet Evol 66:341–349 562 Koya Y, Hayakawa Y, Markevich A, Munehara H (2011) Comparative studies of testicular 563 structure and sperm morphology among copulatory and non-copulatory sculpins 564 (Cottidae: Scorpaeniformes: Teleostei). Ichthyol Res 58:109-125 565 Koya Y, Mitsuhashi N, Awata S, Ito T, Munehara H (2015) Identification of the reproductive 566 mode for internal gamete association in Vellitor centropomus (Cottidae): gonadal 567 histological analysis. Japan J Ichthyol 62:121-131 568 Kumar S, Stecher G, Tamura K (2016) MEGA7: Molecular Evolutionary Genetics Analysis 569 version 7.0 for bigger datasets. Mol Biol Evol 33:1870–1874 570 Le Ralec A, Rabasse JM, Wajnberg E (1996) Comparative morphology of the ovipositor of 571 some parasitic Hymenoptera in relation to characteristics of their hosts. Can Entomol 572 128:413-433 573 Leung TLF (2014) Fish as parasites: an insight into evolutionary convergence in adaptations 574 for parasitism. J Zool 294:1–12 575 Love DC, Shirley TC (1993) Parasitism of the golden king crab, Lithodes aequispinus 576 Benedict, 1895, by a liparid fish. Crustaceana 65: 97-104 577 Mehbub MF, Lei J, Franco C, Zhang W (2014) Marine sponge derived natural products 578 between 2001 and 2010: trends and opportunities for discovery of bioactives. Mar 579 Drugs 12:4539-4577 580 Momota K, Munehara H (2017) Reproductive ecology and morphological changes during the 581 early life stages of Pallasina barbata (Steindachner, 1876). Bull Fish Sci Hokkaido 582 Univ 67:7–12 583 Mills SC, Reynolds JD (2002) Mussel ventilation rates as approximate cue for host selection 584 by bitterling, Rhodeus sericeus. Oecologia 131:473-478 Mills SC, Reynolds JD (2003) The bitterling-mussel interaction as a test case for co-585 586 evolution. J Fish Biol 63:84–104 587 Munehara H (1991) Utilization and ecological benefits of a sponge as a spawning bed by the 588 little dragon sculpin Blepsias cirrhosis. Jap J Ichthyol 38:179-184 589 Munehara H, Goto A, Yabe M (2011) Diversity of Cottoid Fishes -Adaptation and Evolution-. 590 Tokai Univ Press, Kanagawa 591 Munehara H, Takano K, Koya Y (1989) Internal gametic association and external fertilization 592 in the elkhorn sculpin, Alcichthys alcicornis. Copeia 1989:673-678 593 Nakabo T, Kai Y (2013) Cottidae. In: Nakabo T (Ed.) Fishes of Japan with pictorial keys to 594 the species third edition. Tokai University Press, Hadano, pp 1160–1188, 2061–2067 595 Nelson JS, Grande TC, Wilson MVH (2016) Fishes of the World, 5th ed. John Wiley & Sons, 596 Inc, New Jersey 597 Nishida T, Inui R, Onikura N (2008) A note on the spawning bed of Pseudoblennius percoides 598 (Scorpaeniformes, Cottidae) in shallow sea areas around coastal Fukutsu, northern 599 Kyusyu Island, Japan. Biogeography 10:45-51

- 600 Okamura O, Amaoka K Eds. (1997) Seawater Fishes in Japan. Yamatokeikoku-sha, Tokyo
- Paul VJ, Puglisi MP (2004) Chemical mediation of interactions among marine organisms. Nat
   Prod Rep 21:189–209
- Paul VJ, Puglisi MP, Ritson-Williams R (2006) Marine chemical ecology. Nat Prod Rep
  23:153–180
- Palumbi SR (1996) What can molecular genetics contribute to marine biogeography? An
  urchin's tale. J Exp Mar Biol Ecol 203:75–92
- Peden AE, Corbett CA (1973) Commensalism between a liparid fish, *Careproctus* sp., and the
  lithodid box crab, *Lopholithodes foraminatus*. Can J Zool 51:555–556
- Poltev YN, Mukhametov IN (2009) Concerning the problem of carcinophilia of *Careproctus* species (Scorpaeniformes: Liparidae) in the North Kurils. Russ J Mar Biol 35:215–223
- 611 Reichard M, Liu H, Smith C (2007) The co-evolutionary relationship between bitterling fishes
- and freshwater mussels: insights from interspecific comparisons. Evol Ecol Res 9:239–
  259
- Reichard M, Ondrackova M, Przybylski M, Liu H, Smith C (2006) The costs and benefits in
  an unusual symbiosis: experimental evidence that bitterling fish (*Rhodeus sericeus*) are
  parasites of unionid mussels in Europe. J Evol Biol 19:788–796
- 617 Schmidt TR, Gold JR (1993) Complete sequence of the mitochondrial cytochrome *b* gene in
  618 the cherryfin shiner, *Lythrurus roseipinnis* (Teleostei: Cyprinidae). Copeia 1993:880–
  619 883
- 620 Shinomiya A (1985) Studies on the reproductive physiology and ecology in three marine
  621 cottid fish. Dissertation, Hokkaido University
- Shinomiya A, Ikemoto M (1987) Spawning habits of the sculpin *Pseudoblennius percoides* in
  relation to sea squirt. In: Advance abstracts for the 20th annual meeting, The
  Ichthyological Society of Japan, p 26
- 625 Shiogaki M, Dotsu Y (1974) The spawning of the sea sculpin, *Pseudoblennius cottoides*. Bull
  626 Fac Fish Nagasaki Univ 38:71–76
- Smith C, Reichard M, Jurajda P, Przybylski M (2004) The reproductive ecology of the
  European bitterling (*Rhodeus sericeus*). J Zool 262:107–124
- 629 Smith C, Rippon K, Douglas A, Jurajda P (2001) A proximate cue for oviposition site choice
  630 in the bitterling (*Rhodeus sericeus*). Freshw Biol 46:903–911
- 631 Somerton DA, Donaldson W (1998) Parasitism of the golden king crab, *Lithodes aequispinus*,
  632 by two species of snailfish, genus *Careproctus*. Fish Bull 96:871–884
- Spence R, Smith C (2013) Rose bitterling (*Rhodeus ocellatus*) embryos parasitize freshwater
   mussels by competing for nutrients and oxygen. Acta Zool 94:113–118
- 635 Stoecker D (1980) Chemical defenses of ascidians against predators. Ecology 61:1327–1334
- 636 Tamura K, Nei M (1993) Estimation of the number of nucleotide substitutions in the control
- region of mitochondrial-DNA in humans and chimpanzees. Mol Biol Evol 10:512–526
- 638 Uchida K (1932) Fish laying eggs in the body of tunicates. Kagaku 2:56-57
- 639 Uchida K (1979) Chigyo Wo Motomete. Iwanamisyoten, Tokyo

- 640 Uryu T (2011) Marine Fishes of Izu. Kaiyusha, Tokyo
- 641 Yabe M (1985) Comparative osteology and myology of the superfamily Cottoidea (Pisces:
- 642 Scorpaeniformes), and its phylogenetic classification. Mem Fac Fish Hokkaido Univ
- 643 32:1–130

#### 644 Figure legends

657

Fig. 1 Location of sampling sites. (a) Sado Island located 45 km north west of Niigata, Japan.
We collected sculpins that oviposit on rock surface at Usujiri, Hokkaido, Japan. (b) Four
sampling sites on the coast of the island. (c) Kitakoura Port. Monthly collection of tunicates
was conducted along the south breakwater of the port (indicated by an *ellipse*).

649	Fig. 2 Examples of photographs of eggs deposited in sponges and tunicates. (a) Egg masses
650	deposited by females of Pseudoblennius marmoratus in sponges. (b) Eyed eggs of Vellitor
651	centropomus in a colonial tunicate. (c) Colonial tunicate; at least five egg masses were
652	deposited in this colony. Colony width: c.a. 100 mm. (d) Halocynthia ritteri, the solitary
653	tunicates most frequently used by sculpins. (e) Underwater photograph of <i>H. roretzi</i> . (f) Eyed
654	eggs of Ps. cottoides found in H. ritteri. (g) Eyed eggs of Pseudoblennius sp. "Kirin-anahaze"
655	deposited in H. roretzi. (f) Eyed Furcina osimae eggs in Pyura sacciformis. Scale bars
656	indicated in (a), (b), (f), (g) and (h): 10 mm.

658 *Microcosmus hartmeyeri*, (b) *Herdmania japonica*, (c) *Halocynthia roretzi*, (d) *Halocynthia* 659 *ritteri*, (e) *Polycarpa cryptocarpa kroboja*, (f) *Pyura sacciformis* and (g) *Cnemidocarpa irene*. 660 Panels are arranged in decreasing order of the average long diameter (LD) of tunicates. *White*, 661 *grey* and *black bars* indicate tunicates with sculpin egg masses, with Japanese tubesnout egg 662 masses and without any egg masses, respectively.  $n_{sc}$ : number of tunicates with sculpin eggs, 663  $n_{tu}$ : number of tunicates with eggs of Japanese tubesnout (the details of which will be 664 published in a separate paper) and  $n_{ne}$ : number of tunicates with no fish eggs.

Fig. 3 Size frequency distribution of solitary tunicates collected during study periods. (a)

665 Fig. 4 (a) Ovipositor length (OL) increased with standard length (SL) of sculpins and differed 666 among sculpins that utilised four different types of hosts. *Circles, triangles* and *squares* 667 denote the species that used solitary tunicates, colonial tunicates and sponges, and rock 668 surfaces as deposition sites, respectively. Regression lines for different types of hosts are from 669 the reduced model of ANCOVA. Relative OL were statistically significantly different between 670 all combinations of the four types (Tukey's HSD, p < 0.05). (b)–(e) Examples of photographs 671 of ovipositors (indicated by the arrows) in the sculpins of the four types. (b) Pseudoblennius 672 cottoides; (c) Vellitor centropomus; (d) Ps. marmoratus; and (e) Alcichthys alcicornis. Scale 673 bars: 10 mm.

**Fig. 5** Body size and ovipositor length of parasitic sculpins were closely related to the size of solitary tunicates selected by each sculpin species as spawning hosts. (a) Female standard length (SL) and (b) ovipositor length (OL) of six species of sculpins that used solitary tunicates as hosts. (c) Long diameter (LD) and (d) egg mass position of tunicates utilised as hosts. Means ( $\pm$  SD) with different letters are significantly different (Tukey's HSD, *p* < 0.05). Sample sizes are indicated as numerals in bars. *ND*, no data. Bars are arranged in decreasing order of the average SL of sculpins from left to right.

#### 682 Supporting information

Figure S1 Maximum likelihood phylogeny of nine species of potentially ostracophilous
sculpins, with *Jordania zonope* as an outgroup, as reconstructed by MEGA ver. 7.0.14. using
800 bp *cytb* sequences, with the settings of the Tamura Nei model and 1000 bootstrap
replicates. Values at nodes are bootstrap percentages.

Figure S2 Maximum likelihood phylogeny of the potential parental species and 205 eggs
from 113 egg masses as reconstructed by MEGA ver. 7.0.14. using 800 bp *cytb* sequences,
with the settings of the Tamura Nei model and 1000 bootstrap replicates. Values at nodes are
bootstrap percentages.

Figure S3 Maximum likelihood phylogeny of the potential parental species and 18 eggs from
14 egg masses as reconstructed by MEGA ver. 7.0.14., using 628 bp *cytb* sequences, with the
settings of Tamura Nei model and 1000 bootstrap replicates. Values at nodes are bootstrap
percentages.

695 Movie S1 Spawning behaviours of female *Pseudoblennius cottoides* in the aquarium. Six 696 *Halocynthia ritteri* on wire-netting were placed in a  $75 \times 45 \times 45$  cm aquarium. Eggs were 697 deposited in the atrium of a tunicate through the atrial syphon using extremely long 698 ovipositors. Eggs were probably released into the atrium when the female opened her mouth. 699 The female stayed still on the tunicate after spawning. She ejected her ovipositor 1 min 22 sec 700 and left the tunicate 2 min 34 sec after the insert, respectively (not shown). This movie was 701 filmed by H. S. on Feb. 7, 2015. Fish and tunicates were collected on the coast of Sado Island 702 in the Sea of Japan.

Movie S2 Failure of spawning by female *Pseudoblennius cottoides* in the aquarium. The atrial
siphon of the tunicate closed before the sculpin female inserted her ovipositor. Settings were
the same as in Movie S1. This movie was filmed by H. S. on Feb. 1, 2015.

- 706 Movie S3 Spawning behaviours of female *Pseudoblennius marmoratus* in the field. Eggs
- 707 were deposited in the spongocoel of a sponge using short ovipositors. This movie was filmed
- 708 by Tomonobu Uryu on Dec. 28, 2009 at Izu Oceanic Park, Izu Peninsula, Pacific coast of
- Japan. Water temperature was about 16 °C and water depth was about 5 m.