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Assessment of the timing and degree of smolt development in southern populations of masu salmon

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Running headline: PARTIAL SMOLTING OF MASU SALMON

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The present study assessed whether non-anadromous masu salmon *Oncorhynchus masou* in Miyazaki, southern Japan, smoltify, and if so at what time of the year. Yearling *O. masou* of Miyazaki and an anadromous population from Hokkaido, northern Japan, were reared in hatcheries in their respective regions and sampled monthly from February to June to examine the spring smoltification period. The Hokkaido population showed a peak of gill Na\(^+\)/K\(^+\)-ATPase (NKA) activity in May, which was accompanied with an increase in mRNA levels of the seawater (SW)-type NKA alpha subunit, *nka α1b*. Increases in gill NKA activity and *nka α1b* levels were not seen in Miyazaki populations. Transferring yearling Miyazaki population to 70% SW (salinity of 23) in mid-April resulted in an increased serum osmolality over four days. These results suggest that they do not smoltify in their second spring. Next, profiles of gill NKA activity and its subunit mRNA levels in under-yearling Miyazaki population in the autumn were examined. Two phenotypes differing in body color during this period were categorized as parr and smolt-like fish. Smolt-like fish had higher gill NKA activity than parr in December while there was no significant difference in gill *nka α1b* levels. Smolt-like fish acclimated to 70% SW better than parr as judged by lower serum
osmolality. However, serum osmolality in smolt-like fish did not return to the basal level seven days after transfer to 70% SW, suggesting that their hypo-osmoregulatory ability was not fully developed to a level comparable to anadromous populations of this species. The present study suggests that, if *O. masou* in Miyazaki go through a smoltification process, it occurs in its first autumn instead of the second spring and is less pronounced compared to anadromous populations.

Key words: masu salmon; smoltification; Miyazaki Prefecture; Na⁺/K⁺-ATPase; osmolality; autumn
Masu salmon *Oncorhynchus masou* (Brevoort 1856) is one of the eight Pacific salmon species distributed in the Asian side of Pacific Ocean. Their distribution is more southerly than other Pacific salmon but spans most of Japan from Hokkaido to Kyushu (Kato, 1991). The southern limit of their distribution in Japan is Miyazaki (32°N) but a land-locked strain is found even further south in Taiwan (Kato, 1991; Kimura, 1989).

Anadromy of *O. masou* declines with latitude (Malyutina *et al.*, 2009; Morita & Nagasawa, 2010) as is the case for other salmonids (Dodson *et al.*, 2013; Morita *et al.*, 2014). In Hokkaido (42°N), the latitudinal middle of their distribution, the majority of females and minority of males are anadromous, going through parr-smolt transformation (smoltification) in the second spring of their life (Kubo, 1980).

Smoltification is a series of pre-adaptive changes in morphology, behavior and physiology by which river-dwelling parr become ocean-type smolt (Wedemeyer *et al.*, 1980; Hoar, 1988; Stefansson *et al.*, 2008; Björnsson *et al.*, 2011; McCormick, 2009, 2013). In one rearing experiment using *O. masou* of Hokkaido, 62% of females and 26% of males became yearling smolts when they were fed twice a ration at 1-2%/body
mass and reared between 5-12°C (Sano and Ozaki, 1969). In Miyazaki, on the other hand, both females and males are believed to be non-anadromous (Kimura, 1989) due to the warm Kuroshio Current running along the Miyazaki coast which reaches 30°C during summer. This lethally high water temperature during the summer is a physical barrier setting them in a unique situation like “seasonal landlocking”.

During the last glacial period, however, *O. masou* in Miyazaki were presumably anadromous since a landlocked strain of this species exists in the high altitude of the far south Taiwan Island (24°N). Increases in water temperature along the Miyazaki coast after the end of the last glacial period should have a negative impact on seaward migration of *O. masou* for thousands of years and reduced selection on anadromous lifestyle over many generations. However, it is not known whether they abandoned the intrinsic rhythm of smoltification or if they smoltify in the spring or at another time of the year.

*O. masou* may undergo smoltification in the autumn. In fact, a subspecies of *O. masou*, amago salmon *O. masou ishikawae* (Jordan and McGregor 1925), smoltify and migrate to the ocean in the autumn to avoid warm seawater influenced by the
Kuroshio Current during the spring and summer (Kato, 1991). They spend a half-year in coastal waters and return to the rivers around May before the seawater temperature rises. Thus, it is possible that *O. masou* in Miyazaki have shifted their timing of smoltification from the spring to the autumn.

Acquisition of hypo-osmoregulatory ability is one of the most important physiological changes during smoltification. The gills are a key organ responsible for extruding sodium and chloride ions, using Na⁺/K⁺-ATPase (NKA; the sodium pump) as a driving force (Evans, 2008; Hiroi & McCormick, 2012; Hwang *et al.*, 2011; Takei *et al.*, 2014). The activity of gill NKA is often used as an indicator of hypo-osmoregulatory ability.

NKA is composed of two essential subunits, α and β, and one regulatory subunit, γ (Blanco & Mercer, 1998; Mobasheri *et al.*, 2000; Geering, 2006, 2008). Multiple isoforms of NKA α and β subunits have been found in vertebrates (Blanco & Mercer, 1998; Geering, 2008), and teleosts possess additional isoforms due to an extra round of whole genome duplication (Rajaran *et al.*, 2001; Serluca *et al.*, 2001; Dalziel *et al.*, 2014). There are five isoforms of NKA α subunits in salmonids, of which an
isoform named $\alpha_{1b}$ is considered as seawater type while $\alpha_{1a}$ is a freshwater type based on their responses to salinity increases (Richards et al., 2003; Bystriansky et al., 2006; Madsen et al., 2009; McCormick et al., 2009). Gill NKA $\alpha_{1b}$ both at mRNA and protein levels increase during smoltification of Atlantic salmon *Salmo salar* (L. 1758) together with gill NKA activity (Nilsen et al., 2003, 2007; McCormick et al., 2013).

Much less is known about profiles of NKA $\beta$ subunit isoforms during smoltification. NKA $\beta$ subunit localizes the catalytic $\alpha$ subunit on the cell membrane and increases the translation efficiency of the $\alpha$ subunit (Blanco & Mercer, 1998; Rajasekaran et al., 2004; Geering, 2008). Four isoforms of NKA $\beta$ subunit ($\beta_{1a}$, $\beta_{1b}$, $\beta_{3a}$ and $\beta_{3b}$) have been identified in rainbow trout *O. mykiss* (Walbaum 1792) and *S. salar* (Gharbi et al., 2004, 2005). A few studies dealt with changes in gill nka $\beta_{1}$ subunit during smoltification of *S. salar* and reported it increased in parallel with nka $\alpha$ subunit and increased NKA abundance during that period (Seidelin et al., 2001; Nilsen et al., 2007). However, there are at present no studies measuring $\beta_1$ isoforms of nka $\beta_{1}$ subunits during smoltification.

Based on their osmoregulatory role during smoltification and seawater
acclimation, comparing profiles of gill NKA activity and its subunits between anadromous and non-anadromous population is a useful tool to reveal the degree of smoltification in *O. masou* in Miyazaki. The aims of the present study were to examine whether these fish undergo smoltification in their second spring or the first fall and, if so, evaluate the degree of smoltification in terms of their hypo-osmoregulatory ability.

**MATERIALS AND METHODS**

**FISH REARING**

*Yearling O. masou in Hokkaido*

*O. masou* of Hokkaido population were obtained from the hatchery of the South Branch of the Salmon and Freshwater Fisheries Institute, Hokkaido Research Organization (42°N; Futami-gun, Hokkaido, Japan). Eggs were collected from returning adults that were released from the hatchery as smolts and returned from the ocean. Alevin were maintained in indoor raceways under dark and fry were moved to outdoor ponds (24.6 x 3.5 m) run through river water in the spring. Fish were maintained in the same outdoor
ponds throughout the sampling period from February to June 2011 (water temperature range: 4 - 14°C) and fed twice or three times a day on a commercial diet (Nippon Formula Feed, Kanagawa, Japan) with standard rations at 0.4-1.9% per body weight.

Yearling *O. masou* in Miyazaki

Yearling *O. masou* of Miyazaki populations were obtained from a local fish farm at Gokase (33°N; Nishiusuki-gun, Miyazaki, Japan) from April to June 2014 (water temperature range: 9-13°C) and Kobayashi Branch, Miyazaki Prefectural Fisheries Research Institute (32°N; Kobayashi, Miyazaki, Japan) from February to April 2015 (water temperature range: 8-15°C). Fish at the local farm were a captive broodstock and were maintained in the river water in outdoor ponds (4 x 20 m) and fed five times a week on a commercial diet (Scientific Feed Laboratory, Tokyo, Japan) at 1.3%/body weight. Fish at Kobayashi Branch were also a captive broodstock and reared in indoor tanks with a standard ration (1.3%/body weight) until use. In February 2015, they were moved to a 500-l circular fibre-reinforced plastic (FRP) tank and reared until April 2015 in cooled water simulating the local river water temperatures (8-15°C).
Under-yearling *O. masou* in Miyazaki

Under-yearling *O. masou* were also reared at the local fish farm in Gokase, Miyazaki as described above. Two phenotypes (parr and smolt-like fish) were sorted based on the silvery color and visibility of parr marks on the body and sampled from September to December 2014.

SEASONAL SAMPLING

Experiments and samplings were carried out in accordance with the guidelines of Hokkaido University Animal Care and Use Committees (#17-0064). Seven to eight fish were sampled from each month, region and phenotype. Fish were anesthetized by 3.3% 2-phenoxyethanol (Kanto Chemical, Tokyo, Japan) and measured for fork length (*L*~*F*~) and body wet mass (*M*~*W*~). Condition factor (*K*) was calculated as follows: $K = \frac{M_W}{L_F^3} \times 100$. Gill arches were excised and a block of gill filaments was immediately frozen on dry ice and stored at -80°C until analyzed for NKA activity. Another block of gill filaments was immersed in RNAlater (Ambion Inc., Austin, TX,
U.S.A.) and stored overnight at 4°C, then frozen and stored in RNAlater at -30°C until laboratory processing for quantification of mRNA.

SEAWATER CHALLENGE TEST

Yearling O. masou in Miyazaki

On 14 April 2015, yearling O. masou of Miyazaki were transferred to 70%, recirculated artificial seawater (70% SW; salinity of 23) (Napqo, Tokyo, Japan) in four 100-l tanks installed with a portable filter system (PowerBox 55, Kotobuki, Nara, Japan) in Kobayashi Branch. Water temperature was maintained at 13.5°C by placing the experimental tanks in a larger tank run through cooled water. Food was withheld during the experimental period. Fish were sampled 0, 1, 2 and 4 days after transfer as described above. Blood was collected using syringe, transferred to a 1.5 ml centrifuge tube and kept overnight at 4°C. After centrifuging at 8,000 g for 10 min at 4°C, serum was collected and stored at -80°C until analysis for osmolality.

Under-yearling O. masou in Miyazaki
On 24 November 2016, parr and smolt-like fish of under-yearling *O. masou* reared at the local fish farm at Gokase were transferred by a truck installed with a freshwater tank with oxygenation to Nobeoka Marine Experimental Station (Nobeoka, Miyazaki, Japan) and placed in 70% artificial, recirculated SW in 500-l tanks with filter system. Water temperature was maintained at 10°C by a water chiller. Fish were sampled 0, 1, 2, 4 and 7 days after transfer as described above. Fish were not fed throughout the experimental period and sampled as described above.

**NA⁺/K⁺-ATPASE ACTIVITY ASSAY**

Gill NKA activity was measured according to Quabius *et al.* (1997) with minor modification (i.e. correction of a wrong concentration of sulfuric acid). Protein concentration was measured by using BCA (bicinchoninic acid) Protein Assay Kit (Thermo Scientific, IL, U.S.A.). The activity was expressed as Pi (µmol) per protein (mg) per time (h).

**RNA EXTRACTION AND CDNA SYNTHESIS**
Total RNA was extracted from the gills using ISOGEN (Nippon gene; Tokyo, Japan) according to the manufacturer’s instruction. One and half micrograms of RNA was reverse-transcribed using SuperScript VILO cDNA Synthesis kit (Invitrogen, Carlsbad, CA, U.S.A.) in a 10-µl reaction according to the manufacturer’s instruction. cDNA was stored at -30°C until use.

REAL-TIME QUANTITATIVE PCR (QPCR)

Sequences of primers for qPCR of nka α1a, nka α1b and ef-1α were the same as described in Nakajima et al. (2014) (Table 1). Primers for qPCR of nka β1a and nka β1b were designed based on their sequences in O. mykiss (Genbank accession #CA374089 and #CB492131; Gharbi et al., 2004) (Table I). Reverse transcribed-PCRs using these primers were performed to prepare assay standards for O. masou. PCR products run on 1.5% agarose gel were excised and purified using QIAEX II Gel Extraction Kit (Qiagen, Valencia, CA, U.S.A.). Copy numbers of the purified amplicon were calculated from the molecular weight of the amplion and concentration. The standard cDNA were serially diluted from 1 x 10^7 to 3 x 10^2 copies.
qPCR was set up using Power SYBR Green PCR Master Mix (Applied Biosystems, Carlsbad, CA, U.S.A.) in a reaction volume of 20 µl with primer concentration of 100 nM. qPCR was run on a 7300 Sequence Detector (Applied Biosystems) using the manufacturer’s recommended cycling conditions: 50°C for 2 min, 95°C for 10 min followed by 40 cycles at 95°C for 15 s and 60°C for 1 min. Measured values were expressed as relative to those of ef-1α. Performance of qPCR was evaluated by confirming a single peak of the dissociation curve in each assay and calculating the amplification efficiencies of the standard curves, which were within the range of 97-100%. Coefficients of determination of the standard curves were also between 0.99-1.00.

SERUM OSMOLALITY MEASUREMENT

Osmolality in serum was measured by using a vapor pressure osmometer (Wescor 5500; Logan, UT, U.S.A.). Ten microliters of serum was used for the measurement.

STATISTICAL ANALYSIS
Results on yearling *O. masou* in Hokkaido and Miyazaki were analyzed by one-way ANOVA using the JMP program (SAS Institute Inc., Cary, NC, U.S.A.) followed by the Fisher’s protected least significant difference (PLSD) test. Results on under-yearling *O. masou* in Miyazaki were first analyzed by two-way ANOVA (phenotype x time). When significant effects were found, differences were further identified by one-way ANOVA followed by the Fisher’s PLSD test. Differences among groups were considered to be significant at *P* < 0.05.

**RESULTS**

Body size (*L*_F and *M*_w) of yearling *O. masou* in Miyazaki was larger than that of Hokkaido salmon throughout the sampling period (from February to June) [*L*_F: two-way ANOVA, *F*<sub>1,70</sub> = 343.11, *P* <0.001; *M*_w: two-way ANOVA, *F*<sub>1,70</sub> = 245.34, *P* <0.001; Fig. 1]. In both groups, average *K* values were the lowest in May followed by significant increases in June [Hokkaido: Fisher’s PLSD ANOVA, *F*<sub>4,30</sub> = 12.10, *P*<0.001; Miyazaki: Fisher’s PLSD ANOVA, *F*<sub>2,18</sub> = 4.42, *P* <0.05; Fig. 1(e), (f)].

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Gill NKA activity in yearling *O. masou* in Hokkaido showed an increase from March to April, peaked in May and decreased to near the basal level in June [Fisher’s PLSD ANOVA, $F_{4,30} = 8.93, P <0.001$, Fig. 2(a)]. *O. masou* in Miyazaki also tended to increase in gill NKA activity from March to May, but its highest values were only about half of those in Hokkaido [Fig. 2(b)].

Gill *nka α1a* mRNA levels were unchanged during spring both in yearling *O. masou* in Hokkaido and Miyazaki and their levels were similar between the two groups [Fig. 3(a), (b)]. On the other hand, yearling *O. masou* in Hokkaido showed a sharp peak of gill *nka α1b* levels in May [Fisher’s PLSD ANOVA, $F_{4,30} = 18.72, P <0.001$; Fig. 3(c)] while that in Miyazaki salmon was remained at the basal level [Fig. 3(d)]. Both gill *nka β1a* and *β1b* levels in fish in Hokkaido also showed peaks in May [*β1a*:Fisher’s PLSD ANOVA, $F_{4,29} = 10.00, P <0.001$; *β1b*: Fisher’s PLSD ANOVA, $F_{4,30} = 12.52, P <0.001$; Fig. 4(a), (b)]. There was a significant increase in gill *nka β1a* levels from February to April in fish in Miyazaki [Fisher’s PLSD ANOVA, $F_{2,18} = 6.03, P <0.01$; Fig. 4(c)] whereas gill *nka β1b* levels remained constant during that period [Fig. 4(d)].

Transferring yearling *O. masou* in Miyazaki from freshwater to 70% SW...
(salinity of 23) in mid-April resulted in an increase in serum osmolality on day 1, after which it gradually decreased until day 4 after transfer but still significantly higher than those of the initial controls in freshwater [Fisher’s PLSD ANOVA, $F_{3,25} = 11.87$, $P <0.001$; Fig. 5(a)]. Gill NKA activity tended to increase, but not significantly, until 4 days after transfer [Fig. 5(b)].

Body size and $K$ were compared between under-yearling parr and smolt-like fish in Miyazaki during fall [Fig. 6]. $L_F$ was not significantly different between the two groups except in November, when smolt-like fish were larger than parr [two-way ANOVA, $F_{1,55} = 8.14$, $P <0.001$; Fig. 6(a)]. $M_N$ was also similar between the two groups except in October, when parr was larger than smolt-like fish [two-way ANOVA, $F_{1,55} = 4.66$, $P <0.01$; Fig. 6(b)]. Overall, both phenotypes decreased $K$ during the autumn and smolt-fish had lower $K$ than parr [phenotype: two-way ANOVA, $F_{1,55} = 16.36$, $P <0.001$; month: two-way ANOVA, $F_{3,55} = 4.66$, $P <0.001$; Fig. 6(c)] .

There was an overall effect of phenotype on gill NKA activity, which was higher in smolt-like fish (two-way ANOVA, $F_{1,56} = 6.48$, $P <0.05$). The activity was similar in parr and smolt-like fish during September and October, but smolt-like fish
exhibited significantly higher NKA activity than parr in November and December

[Fisher’s PLSD ANOVA, $F_{1,56} = 2.81$, $P < 0.05$; Fig. 7].

Overall, gill $nka\ \alpha1a$ mRNA levels were higher in parr [two-way ANOVA, $F_{1,55} = 4.16$, $P < 0.05$; Fig. 8(a)] but $\alpha1b$ mRNA levels were not significantly different between phenotypes and remained relatively constant during September and December [Fig. 8(b)]. Gill $nka\ \beta1a$ showed an increase in October in both phenotypes (two-way ANOVA, $F_{1,55} = 13.66$, $P < 0.001$), but it remained constant thereafter and no significant difference was seen between parr and smolt-like fish [Fig. 8(c)]. Gill $nka\ \beta1b$ in both phenotypes showed a gradual increase from September to December (two-way ANOVA, $F_{3,55} = 6.74$, $P < 0.001$) and did not differ between parr and smolt-like fish [Fig. 8(c)].

When parr and smolt-like fish were transferred from freshwater to 70% SW, there were overall effects of phenotype and time and their interaction on serum osmolality (phenotype: two-way ANOVA, $F_{1,70} = 61.13$, $P < 0.001$; time: two-way ANOVA, $F_{4,70} = 38.47$, $P < 0.001$; interaction: two-way ANOVA, $F_{1,70} = 10.11$, $P < 0.001$). The degree of increase in serum osmolality on day 1 after transfer was lower in smolt-like fish than in parr [Fisher’s PLSD ANOVA, $F_{9,70} = 26.74$, $P < 0.001$; Fig. 9(a)].
Serum osmolality in parr remained relatively high for 7 days. Serum osmolality in smolt-like fish increased over 7 days as compared to the level in initial freshwater controls. Gill NKA activity in smolt-like fish was higher than that in parr throughout the experimental period [two-way ANOVA, $F_{1,70} = 68.30$, $P < 0.001$; Fig. 9(b)].

**DISCUSSION**

*O. masou* in Miyazaki may have been prevented from spring migration by warm seawater temperature, but they still may have opportunities to migrate to the ocean during the autumn and/or winter. Such a migratory strategy may have lead to increased ability of Miyazaki *O. masou* to tolerate seawater in autumn. In order to test this hypothesis, the present study evaluated their smoltification status in the spring and autumn.

Although defining and quantifying smoltification is difficult (Stefansson et al., 2003; Björnsson et al., 2012), one of the characteristic changes during smoltification is an increase in gill NKA activity. Since the activity is generally correlated with
whole-body hypo-osmoregulatory ability, it is often used as an index of smoltification. Moreover, recent findings showed that one of NKA α1 isoforms, α1b, is responsible for branchial NKA function in salmon in seawater and also increases during smoltification. Other parameters commonly used as indices of smoltification are silverying of body color and a reduction of condition factor.

The results of the present study suggest that *O. masou* in Miyazaki do not increase hypo-osmoregulatory ability in their second spring, which contrast with *O. masou* in Hokkaido which show clear evidence of smolt development in spring (Kubo, 1980). Gill NKA activity in Hokkaido population showed a clear peak in May, corresponding to their active migration period. This increase in gill NKA was accompanied with increases in *nka α1b* as reported in a previous study (Nakajima et al., 2014). In addition, *nka β1a* and *β1b* subunits, which were measured separately for the first time, also peaked in May. Parallel increases in *α1b* and *β1* subunits are in good agreement with the findings in *S. salar* (Nilsen et al., 2007). The present study suggests that both NKA β1 subunits play a similar role in enhancing the localization of NKA α1b subunit to the cell membrane and thus promoting the development of
hypo-osmoregulatory ability during smoltification. Although there was a peak in gill NKA activity in Miyazaki population in May, its level was as low as that of parr in Hokkaido and they had no increases in gill \( nka \alpha 1b, \beta 1a \) and \( \beta 1b \) subunits. In addition, their hypo-osmoregulatory ability was not high enough to restore serum osmolality down to the basal level 4 days after transfer to 70% SW (salinity of 23). Although a direct comparison cannot be made, smolts of Hokkaido population were capable of restoring increased serum sodium ion levels within 24 h after full-strength SW (Ban et al., 1987). Based on these findings, they unlikely go through a smoltification process in their second spring.

Exact mechanisms for why \( O. \ masou \) in Miyazaki do not smoltify in the spring are not known, but one reason may be due to a conflict between smolting and maturation. Initiation of maturation in freshwater is known to inhibit smolting (Thorpe, 1986, 1994). High water temperature in Miyazaki accelerates growth, which in turn promotes maturation in freshwater and inhibits smoltification in the second spring. When body size was compared between Miyazaki and Hokkaido populations, the former was much larger. Despite a reduction of condition factor in Miyazaki
populations from March to May, some fish of both sexes had developing gonads (data not shown), suggesting sexual maturation had already begun in this group. In support of this suggestion, Morita & Nagasawa (2010) revealed by a combination of field survey and modeling that warmer water temperature increased freshwater residency of *O. masou* populations in northern Japan through improving early growth conditions.

Next, the possibility that they smoltify in their first autumn was assessed. When comparing parr and smolt-like fish, smolt-like fish had higher gill NKA activity than parr in December. However, the higher gill NKA activity was not accompanied with higher gill *nka α1b* or *β1* subunits. These results suggest that the degree of the acquisition of hypo-osmoregulatory ability in smolt-like fish in Miyazaki was not as high as that of smolts in Hokkaido and that their preparatory changes for marine life were not complete. This assumption was supported by the result of 70% SW transfer experiment. Smolt-like fish had serum osmolality lower than that in parr throughout the experimental period for 7 days. However, serum osmolality in smolt-like fish continued to increase from 2 to 7 days after transfer to a diluted SW (70%).

It is of note that no/little increase in gill *nka α1b* and *β1* subunits in smolt-like
Miyazaki population may be characteristics of autumn smoltification. The patterns observed in the present study were similar to those in *O. masou ishikawae* (Nakajima *et al.*, 2014), where increased NKA activity was not accompanied with *nka α1b* change. Thus, decreasing water temperature or/and photoperiod during the autumn might cause such discordant profiles of NKA activity and *nka α1b*. Our unpublished data showed that gill *nka α1b* levels in smolt-like fish in the autumn increased following seawater transfer (Uchida *et al.*, unpublished data), suggesting that this phenotype does not increase gill *α1b* mRNA levels until they are actually exposed to seawater.

Smolting in the autumn has been reported in Chinook salmon *O. tshawytscha* (Walbaum 1792) (Ewing *et al.*, 1979; Youngson *et al.*, 1983; Healey, 1991; Beckman & Dickhoff, 1998; Schroeder *et al.*, 2016). Autumn smolts in this species had increased gill NKA activity and down migrated the rivers and entered the ocean (Beckman and Dickhoff, 1998; Schroeder *et al.*, 2016). In the case of *S. salar* in southern England, about 25% of juveniles become autumn migrants but they were not sufficiently physiologically adapted to permit permanent or early, entry into the marine environment (Riley *et al.*, 2008). Profiles of smolt-related characters of *O. masou* in Miyazaki may
be comparable to that of *S. salar* in southern England in terms of limited ability to hypo-osmoregulate. Thus, it is not known if smolt-like fish in Miyazaki actually down migrate the river and enter the ocean.

The present study was unable to examine an interaction between genetic and environmental factors since *O. masou* populations with different genetic backgrounds were reared at different environments (i.e. in Hokkaido and Miyazaki). *O. masou* in Miyazaki were reared at higher water temperatures with enough feed throughout the sampling period resulting in larger body size compared to the age-matched Hokkaido population. Thus, the life-history patterns of Miyazaki population described in the present study might be simply environmental responses without genetic difference. It is possible that autumn smoltification might be driven by accelerated growth and consequently spring smoltification might be blocked by the initiation of maturation in freshwater. A common garden experiment using both *O. masou* populations from Hokkaido and Miyazaki should disentangle environmental effect from genetic influence.

The results of the present study are relevant to aquaculture in Miyazaki. There
is a growing interest in sea cage aquaculture for *O. masou* in this region as a local brand. However, a challenge is that fish farmers need to transfer fish to seawater in the winter due to lethally high warm seawater temperatures during the summer. A direct transfer of under-yearling fish in the winter resulted in a high rate of mortality (approximately 70%) (Uchida *et al.*, unpublished data). Thus, unraveling the degree of their smoltification and adopting rearing strategy to stimulate the development of hypo-osmoregulatory ability will be important for the success of sea cage aquaculture in this region. In *O. tshawytscha*, accelerated growth through the summer-fall stimulated the development of smolt characters (Beckman *et al.*, 2003).

In summary, the present study suggests that if *O. masou* in Miyazaki go through the smoltification process, it is the first autumn instead of the second spring. However, it is presumably an incomplete process compared to the robust development of salinity tolerance that occurs in anadromous *O. masou*. The life-history patterns of *O. masou* in Miyazaki provide a unique opportunity to understand how life-history pathways are regulated and evolved.
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References


chinook salmon: relation of size and growth rate to autumn smolting. 


Dalziel, A.C., Bittman, J., Mandic, M., Ou, M. & Schulte, P.M. (2014). Origins and


Nephrology and Hypertension 17, 526-532.


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Figure captions

Fig. 1. Mean fork length ($L_F$; a,b), body mass ($M_w$; c,d) and condition factor ($K$; e,f) of yearling *Oncorhynchus masou* in Hokkaido (a,c,e) and Miyazaki (b,d,f). *O. masou* in Hokkaido were sampled from February to June in 2011 at Kumaishi (circle) and those in Miyazaki were sampled from February to April in 2015 at Gokase (square) and from April to June in 2014 at Kobayashi (triangle). Values are expressed as mean ± S.E. (n=7-8). Symbols sharing the same letters within a group are not significantly different from each other (one-way ANOVA followed by Fisher’s PLSD test, $P < 0.05$). Note that the results of statistical analyses of the data on *O. masou* of Miyazaki from different locations/years are expressed in lower and upper letters.

Fig. 2. Changes in gill NKA activity in yearling *Oncorhynchus masou* in Hokkaido (a) and Miyazaki (b) during the spring. *O. masou* in Hokkaido were sampled from February to June in 2011 at Kumaishi (circle) and those in Miyazaki were sampled from February to April in 2015 at Gokase (square) and from April to June in 2014 at Kobayashi (triangle). Values are expressed as mean ± S.E. (n=7-8). Symbols sharing the same
letters within a group are not significantly different from each other (one-way ANOVA followed by Fisher’s PLSD test, $P < 0.05$). Note that the results of statistical analyses of the data on *O. masou* of Miyazaki from different locations/years are expressed in lower and upper letters.

Fig. 3. Changes in gill $nka\ ala$ (a,b) and $alb$ (c,d) mRNA levels in yearling *Oncorhynchus masou* in Hokkaido (a,c) and Miyazaki (b,d) during the spring. *O. masou* in Hokkaido were sampled from February to June in 2011 at Kumaishi (circle) and those in Miyazaki were sampled from February to April in 2015 at Gokase (square) and from April to June in 2014 at Kobayashi (triangle). Values are expressed as mean ± S.E. (n=7-8). Symbols sharing the same letters within a group are not significantly different from each other (one-way ANOVA followed by Fisher’s PLSD test, $P < 0.05$). Note that the results of statistical analyses of the data on *O. masou* of Miyazaki from different locations/years are expressed in lower and upper letters.

Fig. 4. Changes in gill $nka\ bla$ (a,b) and $blb$ (c,d) mRNA levels in yearling
*Oncorhynchus masou* in Hokkaido (a,c) and Miyazaki (b,d) during the spring. *O. masou* in Hokkaido were sampled from February to June in 2011 at Kumaishi (circle) and those in Miyazaki were sampled from February to April in 2015 at Gokase (square) and from April to June in 2014 at Kobayashi (triangle). Values are expressed as mean ± S.E. (n=7-8). Symbols sharing the same letters are not significantly different from each other (one-way ANOVA followed by Fisher’s PLSD test, *P* < 0.05).

Fig. 5. Changes in serum osmolality (a) and gill NKA activity (b) in yearling *Oncorhynchus masou* in Miyazaki after transfer to 70% seawater (70% SW; salinity of 23). Values are expressed as mean ± S.E. (n=5-8). Symbols sharing the same letters are not significantly different from each other (one-way ANOVA followed by Fisher’s PLSD test, *P* < 0.05).

Fig. 6. Changes in fork length (*L_\text{F}*; a), body mass (*M_w*; b) and condition factor (*K*; c) in under-yearling *Oncorhynchus masou* in Miyazaki during the autumn. Fish were categorized as parr (open circule) or smolt-like fish (closed circule) based on the body
color. Values are expressed as mean ± S.E. (n=8). Overall and interactive effects were indicated by asterisks. Symbols sharing the same letters are not significantly different from each other (one-way ANOVA followed by Fisher’s PLSD test, $P < 0.05$).

Fig. 7. Changes in gill NKA activity in under-yearling *Oncorhynchus masou* in Miyazaki during autumn. Fish were categorized as parr (open circle) or smolt-like (closed circle) based on the body color. Values are expressed as mean ± S.E. (n=8). An interactive effects were indicated by an asterisk. (two-way ANOVA, $P < 0.05$).

Fig. 8. Changes in gill $nka\ \alpha_1a, \ \alpha_1b, \ \beta_1a$ and $\beta_1b$ mRNA levels in under-yearling *Oncorhynchus masou* in Miyazaki during the autumn. Fish were categorized as parr (open circle) or smolt-like (closed circle) based on the body color. Values are expressed as mean ± S.E. (n=8). Symbols sharing the same letters are not significantly different from each other (one-way ANOVA followed by Fisher’s PLSD test, $P < 0.05$).

Fig. 9. Changes in serum osmolality (a) and gill NKA activity (b) in under-yearling
Oncorhynchus masou in Miyazaki after transfer to 70% seawater (70% SW; salinity of 23). Fish were categorized as parr (open circle) or smolt-like (closed circle) based on the body color. Values are expressed as mean ± S.E. (n=5-8). Symbols sharing the same letters are not significantly different from each other (one-way ANOVA followed by Fisher’s PLSD test, P < 0.05).
Inatani et al., Fig. 1
Gill NKA activity (μmol Pi mg⁻¹ h⁻¹)

(a) Hokkaido

(b) Miyazaki

Figure 2

Inatani et al., Fig. 2
Figure 3

Inatani et al., Fig. 3
Inatani et al., Fig. 4

Figure 4

(a) Hokkaido

Relative gill nka β1a mRNA

(b) Miyazaki

Relative gill nka β1b mRNA

(c)

(d)
Inatani et al., Fig. 5

Figure 5

(a) Serum osmolality (mOsmol kg\(^{-1}\)) vs. Days after transfer to 70% SW

(b) Gill NKA activity (µmol Pi mg\(^{-1}\) h\(^{-1}\)) vs. Days after transfer to 70% SW
Figure 6

Inatani et al., Fig. 6
Figure 7

Inatani et al., Fig. 7
Inatani et al., Fig. 8
Figure 9

Inatani et al., Fig. 9

(a) Serum osmolality (mOsmol kg⁻¹)

(b) Gill NKA activity (μmol Pi mg⁻¹ h⁻¹)

Days after transfer to 70% SW