INTRODUCTION

Salinity is one of the most critical factors governing the invasion of aquatic environments by introduced species and largely determines the survival, abundance, and distribution of migrants (Carrete Vega & Wiens, 2012; Drouin et al., 1985; Romano & Zeng, 2012; Whitehead et al., 2011; Zardi et al., 2006). When faced with novel osmotic conditions, species can respond to salinity stress through phenotypic plasticity in behavioral (Berger & Kharazova, 1997; Ho et al., 2019a; Hoyaux et al., 1976; Michalesc et al., 2010) and physiological traits.

Abstract

Plasticity in salt tolerance can be crucial for successful biological invasions of novel habitats by marine gastropods. The intertidal snail *Batillaria attramentaria*, which is native to East Asia but invaded the western shores of North America from Japan 80 years ago, provides an opportunity to examine how environmental salinity may shape behavioral and morphological traits. In this study, we compared the movement distance of four *B. attramentaria* populations from native (Korea and Japan) and introduced (United States) habitats under various salinity levels (13, 23, 33, and 43 PSU) during 30 days of exposure in the lab. We sequenced a partial mitochondrial CO1 gene to infer phylogenetic relationships among populations and confirmed two divergent mitochondrial lineages constituting our sample sets. Using a statistical model-selection approach, we investigated the effects of geographic distribution and genetic composition on locomotor performance in response to salt stress. Snails exposed to acute low salinity (13 PSU) reduced their locomotion and were unable to perform at their normal level (the moving pace of snails exposed to 33 PSU). We did not detect any meaningful differences in locomotor response to salt stress between the two genetic lineages or between the native snails (Japan vs. Korea populations), but we found significant locomotor differences between the native and introduced groups (Japan or Korea vs. the United States). We suggest that the greater magnitude of tidal salinity fluctuation at the US location may have influenced locomotor responses to salt stress in introduced snails.

KEYWORDS

adaptive divergence, invasive species, locomotion, salinity, snail behavior
Over time, invasive populations can also show various evolutionary changes in response to new habitats (Mooney & Cleland, 2001; Sakai et al., 2001; Suarez & Tsutsui, 2008) including adaptive changes in salinity tolerance (Lee et al., 2003).

The intertidal snail *Batillaria attramentaria* is native to the northwestern Pacific region of Asia along the coastlines of Japan, Korea, and eastern China. In the early 20th century, it spreads via oyster aquaculture (i.e., shipments of *Crassostrea gigas* from Japan) to the bays and estuaries of the northeastern Pacific coast of the United States and Canada (Galtsoff, 1932) and eventually appeared in Monterey Bay, California (Bonnot, 1935; Byers, 1999). Its habitat in Monterey Bay differs strikingly from its native habitat and has much greater temporal salinity fluctuation. Tidal salinity fluctuation can impact perivisceral fluid composition and hemolymph composition (Stickle & Ahokas, 1974, 1975), and osmotic and ionic composition of the body fluid of mollusks and echinoderms (Stickle & Denoux, 1976). Despite these presumably intense challenges, *B. attramentaria* is a common intertidal species in its introduced range and is gradually replacing the native snail *Cerithidea californica* in several marshes in northern California (Byers, 2000a, 2000c), possibly due to its high tolerance for hypoxia (Byers, 2000b). In other places such as Monterey Bay, *B. attramentaria* has recently gone from being very abundant to nearly absent (Wasson et al., 2020). Plasticity or adaptive evolution in response to salinity stress might be a factor in this species’ invasion success and abundance and in that of marine invaders worldwide. However, very little is known about behavioral responses to osmotic stress in marine invertebrates, especially gastropods (Ho et al., 2019a).

*B. attramentaria* is well suited for studying phenotypic changes in invasive species because it (a) exhibits direct development and has limited dispersal capacity (Kojima et al., 2004); (b) quickly forms relatively closed local populations after anthropogenic translocation (Bonnot, 1935; Galtsoff, 1932) or natural disasters (Sato & Chiba, 2016); and (c) has been introduced to areas that differ strongly in salinity conditions from its native region. In addition, this species exhibits a geographic subdivision that apparently corresponds to the main trajectories of the Tsushima and Kuroshio seawater currents which flow around the north and south of the Japanese archipelago, resulting in two divergent mitochondrial lineages termed Tsushima and Kuroshio (Ho et al., 2015; Kojima et al., 2004). Here, we examine population-level variability and plasticity in locomotor behavior in response to salt stress in *B. attramentaria* collected from native and introduced locations. We applied a laboratory culturing and recording method (Ho et al., 2019a) to track horizontal crawling distances of snails during 30 days of exposure to five different salinity levels. To assess the impact of genetic composition on locomotor responses, we also sequenced the mitochondrial CO1 gene for each snail. We present our results in terms of the effects of salinity, geographic distribution, and genetic composition on snail locomotion.

### MATERIALS AND METHODS

#### 2.1 Population sampling

We sampled populations of the mud-tidal snail *Batillaria attramentaria* (G. B. Sowerby I, 1855. Figure 1a) from two sites in Japan and one site in the United States (details in Table S1). We also include here published data which we previously collected from a South Korean population (Ho et al., 2019a). Population locations comprised Hacheon, Cheollabuk-do, South Korea (on June 2016 at 35°32′N, 126°33′E, Ho et al., 2019a); Nemuro city, Hokkaido Prefecture, Japan (June 2017 at 43°15′N, 145°28′E); Matsushima Bay, Miyagi Prefecture, Japan (May 2018 at 38°22′N, 141°4′E); and Monterey Bay, Elkhorn Slough, CA, USA (February 2017 at 36°49′N, 121°45′W) (Figure 1b). For each of the three native populations (one Korean and two Japanese sites), a convenience sample of 100 individuals was collected. These native sites all had high surface salinities of 29–33 PSU at the time of collection. For the introduced (the United States and Canada) population, we collected 100 individuals from Monterey Bay, Elkhorn Slough, CA, USA, in February 2017. We also collected 100 individuals from a South Korean population at Hacheon, Cheollabuk-do, South Korea, in June 2016. All specimens were preserved in 70% ethanol and stored at −20°C until used for analyses. We also sequenced the mitochondrial CO1 gene for each snail. We present our results in terms of the effects of salinity, geographic distribution, and genetic composition on snail locomotion.
States) population, 50 individuals were collected. This site had a low surface salinity of 4 PSU at the time of collection.

Due to river discharge, topography, and tides (e.g., Yoon & Woo, 2013), different estuaries can experience wide fluctuations in daily salinity or very little fluctuation. We characterized salinity fluctuation profiles for each sampling site as low (Nemuro-Japan site; 27–34 PSU), moderate (Hacheon-Korea site; 16–30 PSU), or high (Elkhorn Slough–USA site; 0–30 PSU) based on publicly available data on salinity fluctuation collected over the past several years. These data were obtained from http://www.nemuro.pref.hokkaido.lg.jp for Nemuro (Japan), http://www.khoa.go.kr for Hacheon (Korea), and http://www.mbari.org for the Elkhorn Slough (USA). The Elkhorn Slough has multiple water quality monitoring stations and we used the one closest to our sampling site, which may or may not be typical of the entire area but is most relevant to our sample. We could not find recent records of salinity fluctuation at Matsushima Bay (Japan); however, past data on average monthly salinity levels inside Matsushima Bay indicate that salinity at this site fluctuates from 27 to 34 PSU (Ventilla, 1984) and is similar to the Nemuro site.

In animal locomotion experiments, it can sometimes be advantageous to use individuals of similar size. However, we noticed in the field that the typical body size of the introduced population was larger than the native populations. Since our primary question centers on how different populations respond to salinity stress, and since body size might play a role in both locomotion and salt tolerance, we allowed body size to differ between the native and introduced population samples. This decision somewhat constrains our ability to conclude whether body size is a driving factor behind salinity responses; however, it allows us to use a representative sample of each population rather than using specimens whose size is not typical of their population and thus makes our study more ecologically relevant. Ultimately, the native and introduced samples we collected differed by about 1 cm (average native shell length = 2.1 cm; introduced shell length = 3.1 cm). All collected specimens were maintained in a plastic aquarium with a constant air temperature of 25°C, a water salinity that was the same as their collection site, and a 12-hr L:12-hr D photocycle for 2 days prior to the experiments, in order to reduce the effects of transportation stress and to allow for acclimation to the laboratory.

### 2.2 Salinity stress experiment

For each population consecutively, we conducted a 30-day experiment examining locomotor behavior under different salinity treatments. We randomly divided each population sample into five treatment groups, with 20 individuals per group for the native populations and 10 individuals per group for the US population. These groups were maintained in separate plastic aquaria (40 × 23 × 21 cm³, with an inclined layer of sea sand set up on the bottom and 1 liter of aerated artificial saline water; see supplemental figure S1 in Ho et al., 2019b) at salinities of 43, 33, 23, 13, and 3 PSU for 30 days. Saline water was freshly prepared every 2 days from overnight-aerated distilled water and Instant Ocean Sea Salt (United Pet Group Inc.). Snails in each group were marked with nail polish (Eco Nail color, Innisfree, South Korea) to keep track of their identity. All animals were fed to satiation every 2 days with a commercial brand of fresh, chopped seaweed (Ottogi, South Korea) throughout the 30-day experiment.

### 2.3 Locomotor behavior tracking

We recorded the movement of each snail for 1 hr every 2 days throughout the 30-day experiment following the protocol in Ho et al. (2019a). Briefly, we used a Sony NXCAM camera (AVCHD Progressive MPEG2 SD) to film individual snails in the center of a disposable Petri dish (diameter: 9 cm) filled with artificial seawater which had the same salinity as the snail’s assigned treatment group. We increased the video playback rate using AVS Video Editor v.7.1.2.262 and cropped videos using Avidemux v.2.6.12. We used the Spectral Time-Lapse (STL) toolbox (Madan & Sptch, 2014) implemented in Matlab release R2014a (MathWorks Inc.) to quantify movement distance of the snails.

### 2.4 Shell measurements

We measured shell length of all individuals using images extracted from the recorded videos. We used K-Multimedia Player software (KMP Player, PandoraTV) to extract one video frame (resolution about 1,200 × 1,200 pixels) for each snail and then used tpsDig2 to digitize the most anterior and posterior points of the shell (Rohlf, 2008; Figure S1). The diameter of the petri dish (9 cm) was used as the conversion scale to calculate snail length.

### 2.5 Sequencing and phylogenetic analysis

Because Batillaria attramentaria is very similar in appearance to and has an overlapping distribution with B. zonalis, we sequenced the mitochondrial CO1 gene of all specimens in order to confirm species identity and to identify the haplotype of all individuals. After the salinity experiments, we extracted genomic DNA from fresh foot tissue of all snails using a DNeasy Blood & Tissue kit (Qiagen). We PCR-amplified the mitochondrial CO1 gene using published CO1 primers (Ho et al., 2015) and a Fastmix/Frenchetm PCR kit (IntronBio). PCR products were purified using a Dr. Prep kit (MGmed). Sequencing reactions were performed using a Bigdye Terminator V3.1 Cycle Sequencing kit (Bionics).

We performed a Bayesian phylogenetic analysis based on the CO1 sequences of 5 representative specimens which were identified as Kuroshio and Tsushima haplotypes and were from Korea, Japan, and the United States (GenBank accession no.: MG241503–MG241506 and MT800763), and 53 previously sequenced specimens from the shorelines of Korea (GenBank accession no.
HO et al. (2015) and Japan (GenBank accession no. AB164326–AB164358, Kojima et al., 2004). We used a closely related species, Batillaria multiformis, as an outgroup (GenBank accession no. AB054364, Kojima et al., 2001). We employed MegaX (Kumar et al., 2018) to predict the best substitution model for the CO1 data and found that the Hasegawa-Kishino-Yano+Gamma+Invariable (HKY85+G+I) substitution model was the best fit based on its corrected Akaike information criterion (AICc) value of 1,500.65. We then ran the analyses applying the maximum-likelihood statistical method (Nei & Kumar, 2000) using MegaX with 1,000 bootstrap replications and the neighbor-joining statistical method (Saitou & Nei, 1987; Studier & Keppler, 1988) using Geneious tree builder incorporated in Geneious v.6.1.8 (Kearse et al., 2012) with a random seed of 1,000,000 to construct a phylogenetic tree.

2.6 | Model terms

Our main purpose was to assess the effect of differential salinity exposure (ES) on the locomotion of B. attramentaria (see the description of the 30-day experiment above). In addition to ES, we also included in our analyses four other model terms that might influence snail response to salinity: origin (O), location (LO), population (P), and CO1 lineage (LI). Origin was defined as either native (pooled three populations from Korea and Japan) or introduced (one population from the United States). The location term indicated the countries where the snails were collected (Korea, Japan, or the United States). The population term described the sampling site (Hacheon in Korea, Nemuro city, and Matsushima bay in Japan, and Elkhorn Slough in the United States). Lineage was defined as either Tsushima (comprising the Hacheon and Nemuro populations) or Kuroshio (comprising the Matsushima and Elkhorn Slough populations) based on the individual's position in the CO1 phylogenetic tree.

2.7 | Statistical analyses

For all four populations, snails exposed to 3 PSU did not move at all and died within the first 16 days of the acclimation experiments, apparently due to the extreme osmotic stress represented by such low salinity. Locomotor data for all 3-PSU groups were therefore excluded from all analyses. All other individuals survived the entire duration of the experiment and were included in the analyses.

We applied a linear mixed-effect model (LMM) to assess the impacts of multiple predictor variables (see “Section 2.6” above) on the locomotor response of B. attramentaria to different levels of salinity stress using the package lme4 version 3.1-140 (Pinheiro et al., 2018) implemented in R version 3.0.2 (R Development Core Team, 2011). Since we wished to investigate specifically the impacts of salinity stress, geographic distribution, and genetic composition, we fit a set of competing models with separate single predictor variables (ES, O, LO, P, and LI) and their interactions (ES × O, ES × O + LI, ES × LO, ES × LO + LI, ES × P, ES × P + LI, ES × LI, ES × LI + O, ES × LI + LO, and ES × LI + P). We treated snail identity as a random variable. To avoid overparameterization, we limited the maximum number of predictor variables to three per model. Since body size closely corresponded to the origin factor, which was already in our LMM analysis, we did not include body size as a factor in the LMM analysis. For instance, we knew native snails were smaller than introduced snails. We centered all the predictor variables to mean = 0 and standardized to SD = 0.5 to remove multicollinearity and to directly interpret the results in terms of effect size (allowing us to compare predictors). The response variable was the movement distance of snails, which was measured every 2 days throughout the 30-day acclimation experiments. Prior to the analysis, we square-root-transformed the movement data using the package rcompanion 2.2.1 (Mangiafico, 2017) implemented in R 3.0.2 (R Development Core Team, 2011) to meet the assumption of normal distribution. To determine the best covariance structure for the LMM tests, we tested our response variable against several covariance structures: first order autoregressive (AR1), compound symmetry (CS), and unstructured (UN). We compared their corrected AICc values using the package MuMIn 1.9.13 (Barton, 2009) for R 3.0.2. The AR1 model was the best-supported covariance structure based on the AICc value (3,407.24 vs. 3,710.22 [CS] and 3,708.22 [UN]) and was therefore chosen for the LMM tests as the best available compromise between bias and lack of precision.

We next applied a multimodel inference procedure (Burnham & Anderson, 2003) to the set of competing linear mixed-effect models (LMMs) to select the most parsimonious model that best described snail locomotor response. The models were compared based on AICc values using the aforementioned MuMln package. The model with the lowest AICc value and those satisfying a ΔAICc ≤ 6 cut-off rule (Richards, 2005, 2008) were considered the best fit or most parsimonious models. We then conducted post hoc multiple comparison tests of the best-fit model to examine the effects of each explanatory factor. Additionally, we used MuMln to perform model averaging and estimate the importance of predictor variables by summing the weights of models where the variables appeared.

In addition, we examined differences in shell length, an easily measured trait that might influence locomotion, between native and introduced groups and among native populations using a two-way ANOVA. The purpose of this test was to assess size differences among our population samples, especially with a view to confirm suspected size differences between the native and introduced snails. The first level involved comparisons of snails from four populations, three locations (i.e., countries), and two origins (introduced vs. native). The second level involved comparing individuals belonging to the two mitochondrial CO1 lineages of Kuroshio and Tsushima.

2.8 | Data deposition

Data available from the Dryad Digital Repository: https://datadryad.org/stash/dataset/doi:10.5061/dryad.455m2m and the Mendeley Data repository: http://dx.doi.org/10.17632/ijjmh26c2g.1.
RESULTS

3.1 | Phylogenetic analysis

The CO1-based phylogenetic analysis recovered two distinct clusters, which were called the ‘Tsushima’ and ‘Kuroshio’ lineages after the corresponding ocean currents in the snails’ native range (Figure 2). The Hacheon (Korea) and Nemuro (Japan) individuals were part of the Tsushima lineage, while the Matsushima (Japan) and Elkhorn Slough (USA) individuals belonged to the Kuroshio lineage. These lineage assignments agree with a previous study that identified the region of Japan most likely to be the source of the introduced North American population (Miura et al., 2006).

3.2 | Multimodel inference and model averaging

Based on the criteria of $Z$-scores $> 0$ and $p$-values $< .05$, the average parameter estimates indicated that all the model terms were more or less positively related to changes in snail movement distance, in which the ES term was the most important, followed by O (relative importance $= 98%$), ES $\times$ O (89%), and LI (85%) (Table 1). This result is also supported by the AICc scores (low to high) of the models that included ES, O, and LI (models I, II, and III, Table 2). In particular, the multimodel inference test indicated that the most parsimonious model (based on the lowest AICc score, 3,416.97) was ES $\times$ O + LI (model I). In accordance with the $\Delta$AICc $\leq 6$ cut-off rule, the second and third best-fit models were ES $\times$ O (model II) and ES $\times$ LI + O (model III). These three models, respectively, received 76%, 14%, and 8% of the total weight $w_i$. All the other models, which omitted or included LO and P, received higher AICc scores and lower weights, indicating that these models are not important in describing the locomotion of salt-stressed snails. Therefore, the multimodel analysis supports the conclusion that the geographic distribution-origin and genetic composition substantially influence the movement distance of snails in response to salinity stress.

3.3 | Locomotor performance changes upon variables

The locomotor experiments showed that the intertidal snail *Batillaria attramentaria* from different locations was able to acclimate to a

**FIGURE 2** The CO1 phylogenetic tree that was built based on mitochondrial CO1 gene sequences (286 bp) applying the maximum-likelihood method and HKY85 + G + I substitution model. Vertical black and white bars represent for the two distinct lineages of Tsushima and Kuroshio, respectively.
range of salinity from 13 to 43 PSU. From these experiments, we observed significant locomotion differences between the snail groups exposed to 13 PSU and other treatments of 23, 33, and 43 PSU (LMM_{ModelX}: \( F_{ES} (1, 278) = 99.47, p < .0001, \) Table 3). In particular, snails significantly reduced movement distance when transferred from normal salinity conditions of 33 PSU to the acute low salinity of 13 PSU, but did not significantly change their movement distance when transferred to moderately changed salinities of 23 and 43 PSU (Figure 3a). Post hoc tests using the Tukey post hoc criterion for significance indicated that snails exposed to very low salinity (13 PSU) moved significantly less than the other treatment groups (\( d_{23PSU-13PSU} = 0.43 \pm 0.04 \text{ m}^2/\text{d}, d_{33PSU-13PSU} = 0.48 \pm 0.04 \text{ m}^2/\text{d}, d_{43PSU-13PSU} = 0.45 \pm 0.04 \text{ m}^2/\text{d}, p < .0001, \) Table S2, Figure 3a). The moderately stressed snails (exposed to 23 and 43 PSU) moved slightly less than the control group (exposed to 33 PSU), but this difference was not statistically significant (\( d_{23PSU-23PSU} = 0.05 \pm 0.04 \text{ m}^2/\text{d}, d_{43PSU-43PSU} = 0.03 \pm 0.04 \text{ m}^2/\text{d}, p = .8637, \) and \( d_{43PSU-23PSU} = 0.02 \pm 0.04 \text{ m}^2/\text{d}, \) and \( p = .9559. \))

Notably, when considering all treatment groups, we observed that the origin factor had significant impacts on snail locomotion with \( F_{O} (1, 278) = 31.68 \) and \( p < .0001 \) (Model XII). Subsequent post hoc test of this analysis indicated that the native populations moved significantly more than the introduced population (\( d_{Native-Introduced} = 0.28 \pm 0.05 \text{ m}^2/\text{d}, p < .0001, \) Table S2, Figure 3b). On the other hand, we did not record any significant differences in movement distance between the two CO1-lineages (LMM_{ModelXY}: \( F_{LI} (1, 278) = 0.84, p = .3587, \) Table 3) with \( d_{Yoshima-Kuroshio} = 0.03 \pm 0.04 \text{ m}^2/\text{d}, \) and \( p = .3597. \) Besides this, we also found significant locomotion differences based on location and population (LMM_{ModelXII}: \( F_{LO} (1, 278) = 28.53, p < .0001 \) and LMM_{ModelXIV}: \( F_{P} (1, 278) = 5.28, p = .0223, \) Table 3). Subsequent Tukey post hoc tests revealed that differences in locomotor responses among native snail populations were not statistically significant (\( d_{Korea-Japan} = 0.06 \pm 0.04 \text{ m}^2/\text{d}, d_{Japan-USA} = 0.24 \pm 0.06 \text{ m}^2/\text{d}, d_{Hacheon-Matsushima} = -0.03 \pm 0.05 \text{ m}^2/\text{d}, d_{Hacheon-Elkhorn Slough} = 0.24 \pm 0.06 \text{ m}^2/\text{d}, \) and \( p < .0001, \) Table S2 and Figure 4).

A linear mixed-effect model test of the best model (model I) showed that the interaction of Origin (O) and Salinity Exposure (ES) (LMM_{ModelI}: \( F_{ES \times O} (1, 275) = 11.59, p = .0008, \) was significant, and so was the effect of Lineage (LI) (LMM_{ModelI}: \( F_{LI} (1, 275) = 5.38, p = .0211, \) Table 3). This result corresponds to the significant ES \( \times \) O interaction and LI term in the model outputs (\( Z_{ES \times O} = 3.41, p < .001, \) Table 1). However, when implemented separately, only ES and O significantly influenced the movement distance of the snails independently (LMM_{ModelXII}: \( F_{ES} (1, 278) = 99.47, p < .0001 \) and LMM_{ModelXIV}: \( F_{O} (1, 278) = 31.68, p < .0001, \) Table 3), while in contrast, LI did not influence the movement distance of the snails (LMM_{ModelXIV}: \( F_{LI} (1, 278) = 0.84, p = .36, \) Table 3). Detailed differences in locomotion under the effect of the interaction ES \( \times \) O + LI estimated by Tukey post hoc test can be found in Table S3.

| Model parameter | Multimodel average estimate ± SE | Z | p-Value | Completed model estimate ± SE | Relative importance |
|-----------------|----------------------------------|---|---------|-----------------------------|---------------------|
| Intercept       | 0.99 ± 0.01                      | 70.36 | *** | 0.99 ± 0.01 |                      |
| ES              | 0.14 ± 0.01                      | 11.03 | *** | 0.13 ± 0.01 | 100%                |
| O               | -0.32 ± 0.05                     | 6.63  | *   | -0.31 ± 0.07 | 98%                 |
| LI              | 0.75 ± 0.03                      | 2.24  | *** | 0.01 ± 0.04 | 85%                 |
| ES \( \times \) O | -0.12 ± 0.04                     | 3.41  | *** | -0.11 ± 0.05 | 89%                 |
| ES \( \times \) LI | -0.07 ± 0.03                     | 2.68  | *   | -0.006 ± 0.02 | 8%                  |
| LO              | -0.13 ± 0.02                     | 6.41  | *** | -0.003 ± 0.019 | 2%                   |
| ES \( \times \) LO | -0.06 ± 0.02                     | 3.36  | *** | -0.001 ± 0.009 | 2%                  |
| P               | -0.12 ± 0.03                     | 3.55  | *** | -9.76e-10 ± 1.103e-05 | <1%                 |
| ES \( \times \) P | -0.03 ± 0.01                     | 2.21  | *** | -7.997e-11 ± 1.676e-06 | <1%                 |

Note: Estimates of variables were computed after centering and standardizing the explanatory variables to a mean of 0 and a SD of 0.5. Results are presented in both multimodel inference procedure (Burnham & Anderson, 2003) and completed LMM to account for a greater range of alternative, more parsimonious combinations of explanatory variables with respect to their relative weight of statistical support. The relative importance scores were also measured for each variable based on their frequency within parsimonious models. The variables with higher scores are considered more important than those with lower scores and therefore receive higher support as potential correlates of movement distance. The significant level for the model averaging and relative importance estimations was set at 0.05. *** indicates \( p < .001, \) ** indicates \( 0.001 < p < .01, \) * indicates \( 0.01 < p < .05. \) Abbreviations: ES, exposure salinity; LI, lineage; LMM, linear mixed-effect model; LO, location; O, origin; P, population.
than native ones (Table S4A, B), which is not surprising considering that one of the differences in shell length with F_L from different locations or populations also exhibited significant differences (Figure S2). Simple main effect analyses showed that the average shell length of introduced snails was 31% longer (lIntro = 3.12 cm, lNative = 2.14 cm, Table S4B). These analyses also revealed that snails from different locations or populations also exhibited significant differences in shell length with F_L (2, 277) = 193.7, p < 2e-16 and F_L (3, 276) = 195.1, p < 2e-16 (Table S4A). In particular, the individuals from Korea were smallest, followed by the Japan and the US populations (lKorea, Hacheon) = 1.68 cm, lJapan = 2.38 cm, in which lNemuro = 2.62 cm and lMatsushima = 2.13 cm, and lUSA = 3.12 cm, Table S4B). Furthermore, shell length also significantly varied with lineage (F_L (1, 278) = 19.42; p = 1.5e-05), with lTsushima = 2.15 cm and lKuroshio = 2.62 cm, respectively (Table S4A, B), which is not surprising considering that one of the two lineages includes the introduced (larger) individuals.

### DISCUSSION

Compared to the control groups (exposed to a salinity of 33 PSU, the mean salinity of seawater), we found that all four B. attramentaria populations substantially reduced their movement distance when exposed to low salinity (13 PSU) and did not significantly alter their locomotion in response to moderate increases or decreases in salinity (43 and 23 PSU; Figure 3a). Reduced activity seems to be a general gastropod response to unfavorable environmental conditions (Elfwing & Tedengren, 2002; Hughes et al., 1987; Kitching et al., 1987). Marine invertebrates commonly reduce their activity when experimentally exposed to salt stress (Berger & Kharazova, 1997; De Lange et al., 2006; Felten et al., 2008; Lawrence et al., 2001; Pistart et al., 2007), presumably to conserve energy for ionic-osmotic regulation (discussed in Ho et al., 2019b). Thus, the reduction in movement observed at 13 PSU validates our experimental approach as an appropriate method for measuring stress responses in snails under varying salinity exposures. A lack of differences in locomotion among the 23, 33, and 43 PSU groups suggests that B. attramentaria can successfully acclimate to moderate changes in salinity. The universal failure to thrive among the groups exposed to 3 PSU may indicate that this approximates the lower threshold of this species’ salinity tolerance.

**TABLE 2** Detailed parsimonious linear mixed-effect model of all potential explanatory variables that approximate movement distance of the snail Batillaria attramentaria, ranked by decreasing statistical support

| Model | Intercept | ES | O | LO | P | LI |
|-------|-----------|----|---|----|---|----|
| I     | 0.99 ± 0.01 | 0.14 ± 0.01 | -0.33 ± 0.05 | 0.08 |
| II    | 0.99 ± 0.01 | 0.14 ± 0.01 | -0.28 ± 0.04 |     |
| III   | 0.99 ± 0.01 | 0.14 ± 0.01 | -0.33 ± 0.05 | 0.08 ± 0.03 |
| IV    | 0.99 ± 0.01 | 0.14 ± 0.01 | -0.13 ± 0.02 |     |
| V     | 0.99 ± 0.01 | 0.14 ± 0.01 | -0.13 ± 0.02 | -0.01 ± 0.03 |
| VI    | 0.99 ± 0.01 | 0.14 ± 0.01 | -0.13 ± 0.02 | -0.01 ± 0.03 |
| VII   | 0.99 ± 0.01 | 0.14 ± 0.01 | -0.12 ± 0.03 | 0.18 ± 0.07 |
| VIII  | 0.99 ± 0.02 | 0.14 ± 0.01 | -0.12 ± 0.03 | 0.18 ± 0.07 |
| IX    | 0.99 ± 0.01 | 0.14 ± 0.01 | -0.04 ± 0.01 |     |
| X     | 0.99 ± 0.01 | 0.14 ± 0.01 | -0.03 ± 0.03 |     |
| XI    | 0.99 ± 0.14 | 0.14 ± 0.01 |     |     |
| XII   | 0.99 ± 0.14 | -0.28 ± 0.05 |     |     |
| XIII  | 0.99 ± 0.14 | -0.13 ± 0.02 |     |     |
| XIV   | 0.99 ± 0.14 | -0.04 ± 0.02 |     |     |
| XV    | 0.99 ± 0.14 | -0.03 ± 0.04 |     |     |

Note: AICc is an inverse indicator of model parsimony, considering fit (logLik = log-likelihood) and complexity (df = number of parameters to be estimated in the candidate model). The AICc ≤ 6 cut-off rule was used to define the top-model set (Richards, 2005, 2008). The top-model set with AICc ≤ 6 (AICc difference with the best candidate model) comprises 3 concurrent models of I, II, and III (in bold) with a weight of evidence w_i ranging from about 8 to 76%.

Abbreviations: D, df, denominator degree of freedom; ES, exposure salinity; LI, lineage; LO, location; N, df, numerator degree of freedom; O, origin; P, population.

### 3.4 Variation in shell length with distribution and CO1 lineage

We conducted a two-way ANOVA to examine the effect of geographic distribution and genetic composition on the shell length of all 280 individuals included in the locomotor analyses. We confirmed that introduced B. attramentaria individuals were significantly longer than native ones (F_O (1, 278) = 133.5, p value < 2e-16, Table S4A and Figure S2). Simple main effect analyses showed that the average shell length of introduced snails was 31% longer (lIntro = 2.14 cm, lNative = 3.12 cm, Table S4B). These analyses also revealed that snails from different locations or populations also exhibited significant differences in shell length with F_L (2, 277) = 193.7, p < 2e-16 and F_L (3, 276) = 195.1, p < 2e-16 (Table S4A). In particular, the individuals from Korea were smallest, followed by the Japan and the US populations (lKorea,Hacheon) = 1.68 cm, lJapan = 2.38 cm, in which lNemuro = 2.62 cm and lMatsushima = 2.13 cm, and lUSA = 3.12 cm, Table S4B). Furthermore, shell length also significantly varied with lineage (F_L (1, 278) = 19.42; p = 1.5e-05), with lTsushima = 2.15 cm and lKuroshio = 2.62 cm, respectively (Table S4A, B), which is not surprising considering that one of the two lineages includes the introduced (larger) individuals.
of the LI factor was high (85%) and ranked just after the ES, O, and ES × O factors (Table 1). However, we noted that there was significant difference in movement distance of the Matsushima Bay and the Elkhorn Slough snails, with $d_{\text{Matsushima-Elkhorn Slough}} = 0.33 \pm 0.06 \text{ m}^{1/2}$ and $p < .0001$ (Table S2 and Figure 4). Taken together, our samples from the Matsushima, Japan, population and the Elkhorn Slough, USA, population responded to salt stress quite differently, despite being closely related. Similarly, when the US samples were compared against other countries, the locomotor response also varied: the US snails exposed to 13 PSU exhibited the shortest movement distance, and this location also had lower performance at 33 PSU compared with snails from other countries (Figure 5b). The Elkhorn Slough site, where $B. \text{attramentaria}$ invaded at most 80 years ago, experiences a much wider range of salinity levels due to tidal fluctuation than the Korean and Japanese locations. Differences in responses to salinity stress in the introduced population could be due to local adaptation, phenotypic plasticity, or both. The salinity fluctuation records for Elkhorn Slough suggest that this particular invasive population is frequently exposed to the lower limit of its salinity tolerance and is presumably under strong selective pressure. Exposure to an extended period of 13 PSU would be unusual for the native snails from Korea and Japan but not for the introduced snails at this sampling site in the United States. Although we use the term “introduced” as a general category throughout this paper, it is important to note that we had only one sampling site in the snail’s introduced range, so strictly speaking our findings are limited to this site from the Elkhorn Slough introduced population. The same limitation of interpretation applies to the other categories that lack multiple replicates such as sites with high salinity fluctuation profiles. Additional sampling from other regions of the United States could be helpful in elucidating whether the differences we observed in salt-stress responses are common to the snail’s whole introduced range, or specific to the Elkhorn Slough. Future work could also investigate whether parasitic infection rates, which reportedly differ between native and introduced snails (Torchin et al., 2005), affect locomotor performance.

Our examination of shell length indicated that introduced snails were significantly longer than native ones by about 31% (Table S3B and Figure S2). This is comparable to a previous study reporting a size increase of about 14% in this species (Figure 1, Grosholz & Ruiz, 2003). We also found significant differences in size among native populations (Table S3 and Figure S2). Anatomical and morphological changes in marine gastropods after introduction to a new region are not uncommon (e.g., changes in the excretory system of the littorinid $Cenchristis \text{muricatus}$ (Emson et al., 2002); shell color polymorphisms in White Sea $Littorina \text{saxatilis}$ (Sokolova & Berger, 2000), and increases in body size in $Ilyanassa \text{obsoleta}$ and $Urosalpinx \text{cenereaa}$ (Grosholz & Ruiz, 2003)). Increases in body size in introduced populations might be due to life-history selection, more abundant resources, or absence of key predators or parasites (Mitchell & Power, 2003; Torchin et al., 2001). Alternatively, significant variations in size might be due to age structure; for instance, larger and older snails might be regularly harvested by humans at Hacheon and Matsushima Bay.

| ES × O | ES × LO | ES × P | ES × LI | df | logLik | AICc | ΔAICc | $w_1$ |
|--------|--------|--------|--------|-----|--------|------|-------|------|
| 0.03   |        |        |        | 8   | -1,700.47 | 3,416.97 | 0 | 0.76  |
| -0.12 ± 0.04 |    |        |        | 7   | -1,703.19 | 3,420.40 | 3.43 | 0.14  |
| -0.07 ± 0.03 |    |        |        | 8   | -1,702.66 | 3,421.36 | 4.39 | 0.08  |
| -0.06 ± 0.02 |    |        |        | 7   | -1,705.3  | 3,424.64 | 7.67 | 0.02  |
| -0.05 ± 0.02 |    |        |        | 8   | -1,705.2  | 3,426.44 | 9.47 | 0.01  |
| -0.07 ± 0.03 |    |        |        | 8   | -1,707.34 | 3,430.71 | 13.74 | 0 |
| -0.07 ± 0.03 |    |        |        | 8   | -1,719.17 | 3,454.37 | 37.4 | 0 |
| -0.03 ± 0.01 |    |        |        | 8   | -1,719.96 | 3,455.95 | 38.98 | 0 |
| -0.03 ± 0.01 |    |        |        | 7   | -1,723.81 | 3,461.66 | 44.69 | 0 |
| -0.07 ± 0.03 |    |        |        | 7   | -1,726.13 | 3,466.28 | 49.32 | 0 |
| 5      |        |        |        | 5   | -1,729.77 | 3,469.56 | 52.59 | 0 |
| 5      |        |        |        | 5   | -1,757.55 | 3,525.12 | 108.15 | 0 |
| 5      |        |        |        | 5   | -1,759    | 3,528.01 | 111.05 | 0 |
| 5      |        |        |        | 5   | -1,770.03 | 3,550.08 | 133.11 | 0 |
| 5      |        |        |        | 5   | -1,772.25 | 3,554.51 | 137.54 | 0 |
FIGURE 3  Generic (a) exposure salinity, (b) origin, (c) location, (d) population, (e) lineage–response function of the movement distance of the snail *Batillaria attramentaria*. The predicted variables response derived from the LMM tests, specifying random intercept for subject, with N = 280 individuals split into groups of snails acclimating to different salinities of 13, 23, 33, and 43 PSU (a), from native and introduced ranges (b), coming from different locations (c) and population (d), and having different genetic composition (e). The bottom and top of the box are the 25th and 75th percentile of the movement distance, the straight dash lines show the 50th percentile, and the ends of the whiskers represent the minimum and maximum estimates of the movement distance. Outliers are represented by black dots beyond the whiskers. The solid and dashed black curves represent the statistically significant and insignificant difference between two means, respectively.
Is a larger body size responsible for the reduced locomotor performance that we observed in these invasive snails? We suggest that it is not, but caution the reader that our study was not designed to test this question directly. Body size would be expected to be positively rather than negatively correlated with locomotion speed in marine invertebrates: for example, in the jellyfish *Aurelia aurita* (McHenry & Jed, 2003), sea urchin *Paracentrotus lividus* (Domenici et al., 2003) and sea star *Acanthaster planci* (Mueller et al., 2011), although not in other sea star species including *Linckia laevigata*, *Protopsaster nodosus*, and *Acanthaster planci* (Mueller et al., 2011) and the bat star *Patricia miniata* (Montgomery & Palmer, 2012). Currently, there is no strong evidence for a
relationship between body size and speed in gastropods except for one study of the terrestrial *Cornu aspersum*, which displayed a positive correlation between foot length (but not body mass) and speed (Hemmert & Baltzley, 2016). In contrast, we observed the shortest movement distances in the population with the largest average body size. Furthermore, we did not find any significant difference in movement distance among the native populations or between the two COI lineages, all of which had significant size differences (Table S4). Thus, our results do not support a link between size and locomotion in *B. atramentaria*. However, additional research specifically designed to test whether size contributes to salt tolerance and locomotion is needed.

In conclusion, this paper investigated locomotor responses to salt stress in the intertidal snail *Batillaria atramentaria* from different geographic locations and with different genetic composition. We observed that snails living in native habitats (Korea and Japan) and belonging to different genetic groups (Hacheon/Nemuro vs. Matsushima) did not significantly differ in their responses to salinity stress. However, we found that a population of introduced snails (in the United States) exhibited shorter movement distance than snails from native habitats when exposed to salinity stress. This study demonstrates intraspecific variation in salt tolerance in snails and suggests a correlation between locomotor performance and tidal salinity fluctuations.

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CONFLICTS OF INTEREST

None declared.

AUTHOR CONTRIBUTIONS

Phuong-Thao Ho: Conceptualization (lead); Data curation (lead); Formal analysis (lead); Methodology (lead); Writing-original draft (lead); Writing-review & editing (equal). Hoa Quynh Nguyen: Formal analysis (supporting); Writing-review & editing (equal). Elizabeth M. A. Kern: Writing-original draft (supporting); Writing-review & editing (equal). Yong-Jin Won: Funding acquisition (lead); Writing-review & editing (equal).

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REFERENCES

Barton, K. (2009). *MuMin: Multi-model inference*. Retrieved from http://r-forge.r-project.org/projects/mumin/

Berger, V. J., & Kharazova, A. D. (1997). *Mechanisms of salinity adaptations in marine molluscs*. Paper presented at the Interactions and Adaptation Strategies of Marine Organisms, Petersburg.

Bonnot, P. (1935). A recent introduction of exotic species of mollusc in California waters from Japan. *Nautilus*, 49(1), 1–2.

Burnham, K. P., & Anderson, D. R. (2003). *Model selection and multimodel inference: A practical information-theoretic approach* (2 ed.). Springer Science & Business Media.

Byers, J. E. (1999). The distribution of an introduced mollusc and its role in the long-term demise of a native confamilial species. *Biological Invasions*, 1, 339–352. https://doi.org/10.1023/A:1010038001768

Byers, J. E. (2000a). Competition between two estuarine snails: Implications for invasions of exotic species. *Ecology*, 81(5), 1225–1239.

Byers, J. E. (2000b). Differential susceptibility to hypoxia aids estuarine invasion. *Marine Ecology Progress Series*, 203, 123–132. https://doi.org/10.3354/meps203123

Byers, J. E. (2000c). Effects of body size and resource availability on dispersal in a native and a non-native estuarine snail. *Journal of Experimental Marine Biology and Ecology*, 248(2), 133–150. https://doi.org/10.1016/S0022-0981(00)00163-5

Carrete Vega, G., & Wiens, J. J. (2012). Why are there so few fish in the sea? *Proceedings of the Royal Society B: Biological Sciences*, 279(1737), 2323–2329. https://doi.org/10.1098/rspb.2012.0075

De Lange, H. J., Noordoven, W., Murk, A. J., Lürling, M. F. L. W., & Peeters, E. T. H. M. (2006). Behavioural responses of *Gammarus pulex* (Crustacea, Amphipoda) to low concentrations of pharmaceuticals. *Aquatic Toxicology*, 78(3), 209–216. https://doi.org/10.1016/j.aquatox.2006.03.002

Domenici, P., González-Calderón, D., & Ferrari, R. S. (2003). Locomotor performance in the sea urchin *Paracentrotus lividus*. *Journal of the Marine Biological Association of the United Kingdom*, 83(2), 285–292. https://doi.org/10.1017/S0022258503000794

Drouin, G., Himmelman, J. H., & Béland, P. (1985). Impact of tidal salinity fluctuations on echinoderm and mollusc populations. *Canadian Journal of Zoology*, 63(6), 1377–1387. https://doi.org/10.1139/z85-207

Elfwing, T., & Tedengren, M. (2002). Effects of copper and reduced salinity on grazing activity and macroalgae production: A short-term study on a mollusc grazer, *Trochus maculatus*, and two species of macroalgae in the inner Gulf of Thailand. *Marine Biology*, 140(5), 913–919. https://doi.org/10.1007/s00227-001-0763-8

Emson, R., Morriss, D., Andrews, E., & Young, C. (2002). Life on a hot dry beach: Behavioural, physiological, and ultrastructural adaptations of the littorinid gastropod *Cenchritis* (*Tectarius*) *mucratus*. *Marine Biology*, 140(4), 723–732. https://doi.org/10.1007/S00227-001-0738-9

Felten, V., Charmantier, G., Charmantier-Daures, M., Aujoulat, F., Garric, J., & Geffard, O. (2008). Physiological and behavioural responses of *Gammarus pulex* exposed to acid stress. *Comparative Biochemistry and Physiology Part C: Toxicology & Pharmacology*, 147(2), 189–197. https://doi.org/10.1016/j.cbpc.2007.09.006

Galtsoff, P. S. (1932). *Introduction of Japanese oysters into the United States* (Vol. 12). U.S. Government Printing Office.

Grosholz, E. D., & Ruiz, G. M. (2003). Biological invasions drive size increases in marine and estuarine invertebrates. *Ecology Letters*, 6(8), 700–705. https://doi.org/10.1046/j.1461-0248.2003.00495.x

Helmuth, B. S. T. (1998). Intertidal mussel microclimates: Predicting the body temperature of a sessile invertebrate. *Ecological Monographs*, 68(1), 51–74. https://doi.org/10.1890/0012-9615%281998%29068%3A0051%3AIUMPTB%5B2.C0%5D2

Hemmert, H. M., & Baltzley, M. J. (2016). Intraspecific scaling relations between crawling speed and body size in a gastropod. *The Biological Bulletin*, 230(1), 78–84. https://doi.org/10.1086/BBLV230n1p78

Ho, P. T., Kwan, Y. S., Kim, B., & Won, Y. J. (2015). Postglacial range shift and demographic expansion of the marine intertidal snail *Batillaria*
