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Individual behavioral type captured by a Bayesian model comparison of cap making by sponge crabs

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ABSTRACT

'Animal personality' is considered to be developed through complex interactions of an individual with its surrounding environment. How can we quantify the 'personality' of an individual? Quantifying intra- and inter-individual variability of behavior, or individual behavioral type, appears to be a prerequisite in the study of animal personality. We propose a statistical method from a predictive point of view to measure the appropriateness of our assumption of 'individual' behavior in repeatedly measured behavioral data from several individuals. For a model case, we studied the sponge crab Lauridromia dehaani known to make and carry a 'cap' from a natural sponge for camouflage. Because a cap is most likely to be rebuilt and replaced repeatedly, we hypothesized that each individual crab would grow a unique behavioral type and it would be observed under an experimentally controlled environmental condition. To test the hypothesis, we conducted behavioral experiments and employed a new Bayesian model-based comparison method to examine whether crabs have individual behavioral types in the cap making behavior. Crabs were given behavioral choices by using artificial sponges of three different sizes. We modeled the choice of sponges, size of the trimmed part of a cap, size of the cavity of a cap, and the latency to produce a cap, as random variables in 26 models, including hierarchical models specifying the behavioral types. In addition, we calculated the marginal-level widely applicable information criterion (mWAIC) values for hierarchical models to evaluate and compared them with the nonhierarchical models from the predictive point of view. As a result, the crabs of less than about 9 cm in size were found to make caps from the sponges. The body size explained the behavioral variables namely, choice, trimmed cap characteristics, and cavity size, but not latency. Furthermore, we captured the behavioral type as a probabilistic distribution structure of the behavioral data by comparing WAIC. Our statistical approach is not limited to behavioral data but is also applicable to physiological or morphological data when examining whether some group structure exists behind fluctuating empirical data.

Subjects Animal Behavior, Marine Biology, Data Science **Keywords** Camouflage behavior, WAIC, Bayesian approach, Animal personality, Repeated measurement

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INTRODUCTION

An individual is an important hierarchical structure in biology. We aim to capture intraand inter-individual variations in behavior as a probabilistic distribution structure, because it is a prerequisite for the study of 'animal personality' (Sih, Bell & Johnson, 2004; Niemelä & Dingemanse, 2018). Because the term 'individual difference' sometimes means only interindividual difference, we use 'individual behavioral type' to refer to the two variations. Behavioral ecologists and evolutionary biologists have been interested in the behavioral type because behavior can be a selective trait. At the evolutionary time scale, the distribution structure is very likely to be related to the evolvability of behavior (Kirschner & Gerhart, 1998). At the behavioral time scale, the behavioral type can be caused through complex and dynamic interactions of individual properties such as behavioral plasticity based on physiological processes, with surrounding dynamic environments. A typical interaction can be observed in body extending behaviors such as tool making and usage (e.g., Hunt, 1996; Wang, Brodbeck & Iida, 2014; Matsui & Izawa, 2017; Sonoda et al., 2012). A body extending behavior, which is basically a behavior involving the attachment of non-living things to a body, seems to require at least some information processing to infer a current body size in order to achieve an adaptive extension through complex interactions. Uncertainty in the inference, and accumulation of experiences accompanying the realization of body extensions, might result in the emergence of some behavioral type. Here we examine the hypothesis that individual behavioral types would emerge in the body extending behavior. As an example of the body extending behavior, the sponge crabs behavior of cap making and carrying is experimentally examined and statistically modeled in this study.

To capture the structure, we need repeated measurements and specific statistical modeling considering a hierarchical structure (Niemelä & Dingemanse, 2018). Hierarchical models such as a generalized linear mixed effect model (GLMM) are widely considered appropriate for the repeated data (Zuur et al., 2009; Niemelä & Dingemanse, 2018; Reinhart, 2015). However, empirical data has poorly examined the appropriateness of a hierarchical model relative to a non-hierarchical one such as the generalized linear model (GLM). One well-known statistical measure used in GLM from the predictive point of view is the Akaike Information Criterion (AIC) (Akaike, 1974; Sakamoto, Ishiguro & Kitagawa, 1986). To calculate AIC, the maximum log-likelihood needs to be calculated, but in general, prediction by the maximum likelihood (ML) method is inappropriate for hierarchical models (Watanabe, 2005). This is because a model with hierarchical structures is a statistically non-regular model and the assumptionsset in the ML estimation are considered inappropriate (Watanabe, 2005; Watanabe, 2010b; Watanabe, 2018). Then, how much is the degree of inappropriateness? Alternatively, a Bayesian procedure to construct a predictive distribution is known to perform better than the ML method in the hierarchical models in terms of the predictive point of view (Watanabe, 2018). The Bayesian framework can give a unified measure of the inappropriateness.

Although the basic Bayesian framework and its mathematical foundation of measuring the predictability of an arbitrary pair of a statistical model and a prior distribution, has been rigorously established (*Watanabe, 2010b*; *Watanabe, 2010a*; *Watanabe, 2018*),

there are few applications of the framework to behavioral data containing repeated measurements (*Wakita, Kagaya & Aonuma, 2020*). Specifically, the performance of a predictive distribution can be inferred using the Widely-Applicable Information Criterion (WAIC); it is a measure of the generalization error defined as the extent to which a specified predictive distribution is approximated with respect to an unknown true distribution that generates data (*Akaike, 1980*; *Watanabe, 2018*). Furthermore, there are almost no appropriate applications of WAIC to hierarchical models for repeatedly measured data. To construct a predictive distribution using a hierarchical model, we are usually interested in a new observation from a new cluster other than from the clusters that provided the initial observations. Therefore, we need to marginalize the parameters assigned to each cluster when training a model to calculate WAIC in that situation (*Watanabe, 2018*; *Millar, 2018*). However, this point does not seem to be recognized well not only in biological communities but also generally in other real-world data analyses.

Therefore, we propose and adopt a Bayesian model comparison framework using WAIC to study a specific individual behavioral type in the body extending behavior of the crab. In previous research, one field study dealt with the preference of dromiid crabs for materials and examined the association between a cap size with a body size (*McLay*, 1983). Additionally, it is reported that *Cryptodromia hilgendorfi* use caps made from many species of sponges, but they particularly prefer the sponge *Suberites carnosus*, and the crabs make sponge caps twice as large as the carapace area. In previous experimental research, the preference for material size and their suitability for the body size and cap size are scarcely investigated. It is reported that Dromia personata mainly uses sponges and ascidians (Bedini, Canali & Bedini, 2003), although they could also make caps with paper (Dembowska, 1926). Dembowska (1926) reported that a non-breaking space is used qualitatively and that the cap size made by Dromia personata (reported as D. vulgaris) using paper is as large as the size of the caps originally carried by the crabs. Because these studies once sampled a body size and a camouflage size for an individual, it is unclear whether there is an individual behavioral type. In addition, it is unknown whether a behavioral type that is conditional on the body size exists in the cap making behavior. Thus, although the crabs in the family Dromiidae have been known to make a cap (Guinot & Wicksten, 2015), the behavior of the Lauridromia dehaani has not been examined so far.

Accordingly, the lack of experimental data on the cap making behavior of the crab *Dromia personata* and the limitations of the statistical approach, we set four goals to study the individual behavioral type in the body extending behavior: (1) to perform behavioral experiments by sampling behavioral data repeatedly, (2) to formulate an individual behavior type in statistical models, and simultaneously to construct other models assuming no such behavior type, (3) to measure the predictive performance of those models by WAIC, including hierarchical models that assume a particular individual behavior type and compare the findings with those of non-hierarchical models assuming the existence of no such behavior type, and (4) to infer a relationship between the behavioral data and the body size by conditioning the behavioral variables by the body size.

MATERIALS & METHODS

Animal collection

From December 2015 to April 2017, 40 individuals (21 males, 19 females) of Lauridromia dehaani (Brachyura: Dromiidae) (Fig. 1A) were captured using a gill net at the Sakai fishing port, Minabe town, Wakayama, Japan (33°44'N, 135°20'E). We conducted behavioral experiments of cap making on 38 individuals (20 males, 18 females) and video recorded the behaviors of 2 individuals (4.30 cm and 7.19 cm of the carapace widths for each) in a tank filled with filtered natural seawater (about 3.4 % of the salinity) at Shirahama Aquarium, Seto Marine Biological Laboratory, Kyoto University (33°41'N, 135°20'E), from December 2015 to June 2017. For the behavioral experiments, we successfully sampled 8 individuals repeatedly (3 or 5 times for each). Thus, we only sampled one observation from one of the other 30 individuals. Note that although the sample sizes of the behavioral acts for each individual are different our method is still applicable. Before the experiments, all individuals were retained in the tanks (19.5–23.8 °C, light on: 8–17, light off: 17–8) of the aquarium for more than two days for acclimation. We measured their carapace width (cm) (Fig. 1B) as a proxy for the body size, and divided them into three levels depending on whether they lacked any of the fourth and fifth pereiopods: (O) none of the fourth and fifth pereiopods were absent, (1) one of them was absent, (2) both fourth and fifth of each side were absent.

Experimental setup and procedure

We prepared three sizes of white melamine foam that is found commonly worldwide (most notably manufactured by BASF of Germany) and often used in general households (in Japan, it is called Gekiochi-kun, LEC, Inc.) (S: 20 mm \times 30 mm \times 40 mm; M: 30 mm \times 60 mm \times 85 mm; L: 30 mm \times 140 mm \times 150 mm). We used this sponge because it is easy to sink.

First, to confirm that the cap making behavior of the crab *Lauridromia dehaani* is similar to the behavior in the reports (*Dembowska*, 1926; *McLay*, 1983), we video-recorded the behavior from two crabs. The two crabs were used only for video recording in the aquarium (310 mm \times 180 mm \times 240 mm, W \times L \times H). We started the recording from 9 am to 10 am in the morning, and stopped for 2 h after the crabs stopped cap making. We used a red light transmitted through a polyvinyl chloride board and excluded another light source by enclosing the aquarium. We obtained 5 recordings for each crab.

Secondly, we performed experiments on choice of cap size, trimming, and excavating behaviors. S size sponge was smaller than all crabs, whereas the L size was larger than all crabs. Each sponge was put pseudo-randomly on either side and at the back center of the cage (700 mm \times 470 mm \times 190 mm, W \times L \times H, Fig. 1C), which floated in the tank. Then, crabs were introduced to the front center of a cage floating in the tank, thereby the distance between each sponge and the crab was equal. We started a trial from 9 am to 10 am and checked whether the crab carried any sponge once a day. We counted the days when the crab carried a sponge. The latency, measured as the number of days to produce a cap, is modeled as a random variable. Note that the crab was assumed to make a cap at night, because it is considered nocturnal (*McLay*, 1983). If it did, we collected the



Figure 1 Experimental animals and their setup. (A) *Lauridromia dehaani*; p—propodus of fifth pereiopod; d—dactylus of fifth pereiopod; c—chela (1st pereiopod); 2p—second pereiopod; 3p—third pereiopod; 4p—fourth pereiopod; 5p—fifth pereiopod. (B) Carapace width we measured. (C) Experimental cage floating in an aquarium tank with three different sizes of sponges. The drawings are by Keita Harada. Full-size DOI: 10.7717/peerj.9036/fig-1

sponge; otherwise, the crab and the three sponges remained in the cage. When the crab did not carry any sponge for five days, we stopped the trial. We destroyed all sponges that the crabs processed and measured their whole area (cm^2) , and area of the concave part (cm^2) excavated by a crab from the pictures taken 46 cm above the sponges. The trimmed area and concave area are modeled as random variables. Although in the beginning we only performed one trial for one individual $(N_{animal} = 30)$, we obtained five trials for one individual after February 2017 $(N_{animal} = 8)$ to examine the behavior type. Our hypothesis that a behavioral type would be formed was conceived after the day, but we did not change the experimental condition.

Statistical modeling

To quantify the behavioral type in the experiment, we constructed 26 statistical models (Table 1) for the four different aspects: (1) choice of sponge size (6 models), (2) amount of sponge trimmed by cutting (8 models), (3) size of cavity (6 models), and (4) latency to produce a cap (6 models). In each case, we built the statistical models specifying individual behavioral types as hierarchical structures with parameters and performed MCMC samplings from the posterior distribution. Also, we conditioned the variables with the carapace width, levels of leg absence, and gender. We specified the models in the probabilistic programming language Stan (*Stan Development Team, 2018*). We used

non-informative uniform priors for some parameters unless otherwise explicitly described. The performed samplings from the posterior distributions using No-U-Turn Sampler (NUTS) implemented as a Hamiltonian Monte Carlo (HMC) sampler in Stan. Sampling convergence was visually diagnosed by trace plots and quantitatively via the Gelman-Rubin convergence statistic, R_{hat} (*Gelman & Rubin, 1992*). All sampled draws were judged to be converged when $R_{hat} < 1.10$, were used to construct predictive distributions with WAIC on each model. All computations were performed in the R statistical environment (*R Core Team, 2018*), and the Stan codes for each model were compiled and executed through the R package *rstan* (*Stan Development Team, 2018*).

We compared the predictive performances of the models using WAIC (*Watanabe*, 2018; *Watanabe*, 2010b). It should be emphasized that the WAIC of a hierarchical model can be defined in several ways depending on how a predictive distribution is defined. In our case, as we would like to construct a new distribution regarding a new act of a new individual, we have to marginalize the intermediate parameters assigned to each individual in the statistical model (*Watanabe*, 2018). This is because we are interested in the prediction of a new behavioral act when we get a new individual and a new behavioral act instead of the prediction of a new behavioral act from the individuals sampled in this study. By performing this procedure, we can equally compare a hierarchical model with a non-hierarchical model, because the focus of the prediction in a non-hierarchical model is on a new behavioral act of a new individual.

Here we briefly describe the basic procedure based on *Watanabe* (2018). Let $X^n = (X_1, ..., X_n)$ be an sample from the unknown true distribution and p(x|w) a statistical model with w assigned to each individual. Furthermore, w is assumed to be taken from $\varphi(w|w_0)$ and w_0 from φ (w_0) to form a hierarchical structure. In learning step, $w_1, ..., w_{N_{animal}}$ is prepared. In prediction step, our statistical model is built like $p_{model}(x|w_0)$ by marginalizing w out:

$$p_{model}(x|w_0) = \int p(x|w)\varphi(w|w_0)dw.$$
(1)

WAIC is a measure for the degree of accuracy of an approximation of a predictive distribution to the true distribution generating data. For our hierarchical model, the predictive distribution is defined as $\mathbb{E}_{w_0}[p_{model}(x|w_0)]$. Then, the marginal-level WAIC for a hierarchical model is defined as:

$$WAIC_{h} = -\frac{1}{N} \sum_{i=1}^{N} \log \mathbb{E}_{w_{0}}[p_{model}(X_{i}|w_{0})] + \frac{1}{N} \sum_{i=1}^{N} \mathbb{V}_{w_{0}}[\log p_{model}(X_{i}|w_{0})]$$
(2)

where $\mathbb{E}_{w_0}[]$ and $\mathbb{V}_{w_0}[]$ are the average and variance operators, respectively, of the posterior distribution of w_0 . w_0 is estimated so that practically the MCMC sample is used, thus numerical integration is required. In this study, the computation is implemented in the 'function' block in the Stan codes using the Simpson's rule for the numerical integration and the *log_sum_exp* function provided in Stan (see Supplemental Information 3).

Response variable	model	Hierarchical structure	Conditioning variables	Link function	Distribution	WAIC (nat)	dWAIC (nat)	Plot
Choice	1_1	intercept_L	CW_L Leg_L CW_NO Leg_NO	softmax	categorical	-2.13	0.00	Fig. 3A
Choice	1_2	intercept_L	CW_L CW_NO	softmax	categorical	-1.87	0.26	_
Choice	1_3	intercept_L	_	softmax	categorical	-0.88	1.25	_
Choice	1_4	intercept_L	Leg_L Leg_NO	softmax	categorical	-0.78	1.35	_
Choice	1_5	-	CW_L CW_NO	softmax	categorical	0.85	2.99	Fig. 3C
Choice	1_6	-	CW_L Leg_L CW_NO Leg_NO	softmax	categorical	0.87	3.01	_
Trimmed size	2_1	intercept_1 intercept_2	CW Choice	logit log	ZIP	-2.08	0.00	Fig. 4A
Trimmed size	2_2	intercept_2	Choice	logit log	ZIP	0.81	2.89	_
Trimmed size	2_3	intercept_2	CW Choice	logit log	ZIP	0.86	2.95	_
Trimmed size	2_4	intercept_2	_	logit log	ZIP	1.23	3.32	_
Trimmed size	2_5	intercept_2	CW	logit log	ZIP	1.37	3.46	_
Trimmed size	2_6	-	CW Choice	logit log	ZIP	7.40	9.48	Fig. 4B
Trimmed size	2_7	-	CW	logit log	ZIP	10.05	12.13	_
Trimmed size	2_8	-	_	logit log	ZIP	12.55	14.63	_
Cap cavity size	3_1	intercept	CW	log	gamma	4.45	0.00	Figs. 5A
Cap cavity size	3_2	-	CW	log	gamma	4.54	0.08	Figs. 5B
Cap cavity size	3_3	-	CW Gender	log	gamma	4.69	0.24	_
Cap cavity size	3_4	intercept	_	log	gamma	4.71	0.26	_
Cap cavity size	3_5	-	CW	identity	normal	4.75	0.30	_
Cap cavity size	3_6	intercept cw	CW	log	gamma	6.18	1.73	_
Latency for making	4_1	intercept_2	CW	logit log	ZIP	1.10	0.00	Fig. 6A
Latency for making	4_2	intercept_2	_	logit log	ZIP	1.28	0.18	_
Latency for making	4_3	-	_	logit log	ZIP	1.28	0.19	Fig. 6B
Latency for making	4_4	-	Choice	logit log	ZIP	1.30	0.20	_
Latency for making	4_5	-	CW	logit log	ZIP	1.38	0.28	_
Latency for making	4_6	-	CW Choice	logit log	ZIP	1.72	0.62	_

Table 1 Summary of model structures and the predictive performances in WAIC.

Notes.

intercept_L, intercept in the linear predictor (LP) for the choice of L; intercept_1, intercept in the LP for the decision of trimming; intercept_2, intercept in the LP for the mean of the trimmed size of the sponge; CW, carapace width; Leg, degree of the leg lack; _L and _NO, parameters for L sponge and skipping, respectively; Choice, choice of whether to cut the sponge or not; Gender, gender of the animal; intercept_2, intercept in the LP for the mean of the days to carrying; Choice, choice of sponge size; ZIP, Zero-inflated Poisson distribution; WAIC, value of Widely-Applicable Information Criterion per sample; dWAIC, the difference of the WAIC of the model against the best-performed model.

On the contrary, the WAIC for a non-hierarchical model is defined for a statistical model $p_{model}(x|w)$:

$$WAIC_{non-h} = -\frac{1}{N} \sum_{i=1}^{N} \log \mathbb{E}_{w}[p_{model}(X_{i}|w)] + \frac{1}{N} \sum_{i=1}^{N} \mathbb{V}_{w}[\log p_{model}(X_{i}|w)]$$
(3)

where $\mathbb{E}_{w}[]$ and $\mathbb{V}_{w}[]$ are the average and variance operators, respectively, of the posterior distribution of *w*. Note that the often used conditional-level WAIC is described in the Discussion.

Choice of material size (model 1_1)

To provide an overview of the specified models, we describe only the best-performing models in terms of WAIC here. The other models are summarized in Table 1.

We formulate a tendency toward a choice as $\mu[n,m]$ (m = 1,2,3 for M, L, skip, respectively):

$$\mu[n,1] = 0, \tag{4}$$

$$\mu[n,2] = a_{choice_L}[ID[n]] + b_{choice_L} \cdot CarapaceWidth[n] + c_{choice_L} \cdot LegLack[n], \tag{5}$$

$$\mu[n,3] = d_{choice_0} + e_{choice_0} \cdot CarapaceWidth[n] + f_{choice_0} \cdot LegLack[n], \tag{6}$$

$$n = 1, \dots, N_{act} \tag{6}$$

where $N_{act} = 68$ is the total number of behavioral acts, and *ID* represents animal identity (from 1 to $N_{animal} = 38$). μ is linked to the linear predictor in terms of the carapace width, *CarapaceWidth* and the level of absence of leg, *LegLack*. The choice of an M size is fixed to zero. a_{choice_L} is for each individual, thus it is hierarchized. d_{choice_0} is not hierarchized. The distribution of a_{choice_L} is defined as the normal distribution with the mean $a_{choice_L_0}$ and standard deviation $a_{choice_L_s}$:

$$a_{choice_{L}}[k] \sim Normal(a_{choice_{L0}}, a_{choice_{Ls}}), \quad k = 1, \dots, N_{animal}.$$
⁽⁷⁾

The actual choice *Choice* is defined as the categorical distribution with the softmax function:

$$Choice[n] \sim Categorical(softmax(\mu[n,])), n = 1, ..., N_{act}.$$
(8)

Thus, in this case, a statistical model $p_{model}(x|w_0)$ is set using the parameters:

$$w_0 = (a_{choice_{L_0}}, a_{choice_{L_s}}, b_{choice_L}, c_{choice_L}, d_{choice_0}, e_{choice_0}, f_{choice_0}).$$
(9)

Note that $w = a_{choice_L}$ is marginalized out when we build the predictive distribution, so that it is not included in w_0 . The choice *Choice*[*n*] is modeled as a random variable X_n . *CarapaceWidth*[*n*] and *LegLack*[*n*] are the conditioning variables.

Trimming (model 2_1)

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The probability of making the decision on whether an animal is cut off the sponge is written as ϕ_{cut} linked to the linear predictor with the carapace width *CarapaceWidth* and the selected sponge size *Choice*[*n*]:

$$\phi_{cut}[n] = InverseLogit(a_{cut}[ID[n]] + b_{cut} \cdot CarapaceWidth[n] + c_{cut} \cdot Choice[n])$$
(10)

$$u = 1, \dots, N_{act}. \tag{11}$$

 a_{cut} is assigned for each individual. $N_{act} = 51$ and *ID* is from 1 to $N_{animal} = 30$. The distribution of a_{cut} is defined as the normal distribution with the mean a_{cut_0} and standard deviation a_{cut_s} :

$$a_{cut}[k] \sim Normal(a_{cut_0}, a_{cut_s}), \quad k = 1, \dots, N_{animal}.$$
(12)

The prior distribution of a_{cut_s} is defined as the half t distribution:

$$a_{cut_s} \sim Student_t^+(4,0,10).$$
 (13)

The mean area of a sponge trimmed by the crab λ is linked to the linear predictor with the log link function:

$$\log(\lambda_{cut}[n]) = d_{cut}[ID[n]] + e_{cut} \cdot CarapaceWidth[n] + f_{cut} \cdot Choice[n],$$

$$n = 1, \dots, N_{act}.$$
(14)

 d_{cut} is assigned for each individual. The distribution of d_{cut} is defined as the normal distribution with the mean d_{cut_0} and the standard deviation d_{cut_s} :

$$d_{cut}[k] \sim Normal(d_{cut_0}, d_{cut_s}), \quad k = 1, \dots, N_{animal}.$$
(15)

The prior distribution of d_{cut_s} is defined as the half t distribution:

$$d_{cut_s} \sim Student_t^+(4,0,10).$$
 (16)

Altogether, the area of the trimmed sponge is modeled as the variable *Trimmed*. Its distribution of it is defined as the zero-inflated Poisson distribution (ZIP) with the parameters ϕ_{cut} and λ_{cut} :

$$Trimmed[n] \sim ZIP(\phi_{cut}[n], \lambda_{cut}[n]), \quad n = 1, \dots, N_{act}.$$
(17)

When a crab skips the trimming behavior, *Trimmed* is set to zero even if the sponge size is smaller than the defined M and L sizes owing to measurement errors. Note that *Trimmed* is rounded off to an integer. We assume that the rounding process has no significant impact on the data distribution.

Cap cavity making (model 3_1)

To examine the relationship between the cap cavity size *CavitySize* and the carapace width *CarapaceWidth*, gamma distribution is chosen to represent non-negative values of the cavity size. The mean of the distribution is specified by *lambda_{cavity}* with shape and rate parameters:

$$CavitySize[n] \sim Gamma(shape, \frac{shape}{\lambda_{cavity}}),$$
(18)

$$\log(\lambda_{cavity}) = a_{cavity}[ID[n]] + b_{cavity} \times CarapaceWidth[n], \quad n = 1, \dots, N_{act}.$$
 (19)

where the rate parameter was given as the shape over the log-linked linear predictor and a_{cavity} is the intercept for each individual. $N_{animal} = 30$, and $N_{act} = 51$. The a_{cavity} is taken from the normal distribution with the mean a_{cavity_0} and the standard deviation a_{cavity_0} :

$$a_{cavity}[k] \sim Normal(a_{cavity_0}, a_{cavity_s}), \quad k = 1, \dots, N_{animal}.$$
⁽²⁰⁾

Latency (model 4_1)

We assume that the latency to produce a cap, *Days*, fits the ZIP distribution, which is similar to the *Trimmed* case:

$$\phi_{day}[n] = InverseLogit(a_{day}), \tag{21}$$

$$\log(\lambda_{day}[n]) = b_{day}[ID[n]], \tag{22}$$

$$b_{day}[k] \sim Normal(b_{day_0}, b_{day_s}), \quad k = 1, \dots, N_{animal},$$
(23)

$$Days[n] \sim ZIP(\phi_{day}[n], \lambda_{day}[n]), \quad n = 1, ..., N_{act}.$$
(24)

where $N_{animal} = 32$, $N_{act} = 56$. Note that b_{day} is into this model to construct a hierarchical structure.

All the data and codes are available from the Supplemental Information 3.

RESULTS

We measured and modeled the four variables: the choice of sponge size, trimmed size, cavity size, and latency for completing making sponge, as random variables. Furthermore, we evaluated the model predictability by WAIC (see Materials and Methods).

Cap making using an artificial sponge

The behaviors of the two crabs were video recorded to confirm the Cap making behavioral sequence when using an artificial sponge (Fig. S1). The crabs grasped either side of the sponge using their second and third pereiopods, and trimmed small pieces of the sponge using their chelae (Fig. 2A upper left, upper right, Video S1). They first visited the two sides of the sponge. To make a cavity, the crabs rotated their bodies backward and grasped the sponge by the fourth and fifth pereiopods. By repeating these behaviors, the crabs made a



Figure 2 Cap making and carrying behavior. (A) Cap making behavior. (B–C) Carrying behavior of a crab. The drawing is by Keita Harada. Full-size DOI: 10.7717/peerj.9036/fig-2

groove to cut off a portion of a sponge. In 9 of the 10 trials conducted, it took about 50 min to cut the portion, and the crabs started excavating as soon as they finished the trimming behavior. In the other trial, it took 19 min.

Next, the crabs made cavities by tearing off small pieces of a sponge (Fig. 2A bottom, Video S2). It took 11 min on average to excavate the cavity. Then, the crabs rotated their bodies backward in order to catch the excavated sponges with the fourth and fifth pereiopods while they kept the portion grasped by the second and third pereiopods. Finally, the crabs released the second and third pereiopods from the cap and carried it off (Figs. 2B, 2C). In terms of behavior, the crabs often rotated their bodies forward, dorso-ventrally, to enlarge the cavity. It is rare for them to move laterally. They repeated the excavation activities up to eleven times per night and it took up to 4 h. When the crabs rotated their bodies, the direction of rotation was maintained along with the sponge. While the crabs cut the sponge, they actively moved around the sponge. In contrast, they persistently stayed under the sponge during excavation.

Sponge choice

None of the 38 animals chose the S size sponge, and 7 animals skipped the cap making behavior (Fig. 3A). Therefore, we defined the choice as a random variable taking either of the three values: M, L, or skipping. The hierarchical model assuming behavioral types 1_1 (Figs. 3A, 3B) outperformed the non-hierarchical one in terms of WAIC (2.99 nat/sample in the difference, Figs. 3A–3D, Table 1). The posterior probability of the behavioral choices was more widely variable on model 1_1 than in model 1_6 depending on the individual difference specified as a_{choice_1} (Fig. 3B). To show the variability, the probability of choice



Figure 3 Sponge choice. (A) Predictive distribution on the hierarchical model 1_1 with data points of the behavioral choices, which are M or L size choices or skipping the behavior. The points connected by dotted lines represent data from the same individual. The white curved lines are ten samples from the posterior distribution in decreasing order from the highest density of a parameter representing the probability of a choice. (B) Structure of the model 1_1 in a graphical diagram. a_{choice_L} is a parameter assigned to each individual. The variables in the black and white ellipses represent observed data and parameters to be estimated, respectively. (C) Predictive distribution of a choice on the non-hierarchical model 1_6 in a graphical diagram.

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sampled from the posterior distribution from the highest density is visualized in white lines (Figs. 3A, 3C). Note that the variability of the choice probability in the white curved lines is smaller than the model 1_1 even if the number of lines are the same. Although the body size of the animal indicated with the white arrowhead (Fig. 3A) is small, it preferably selected the size L. This indicates a large inter-individual difference. In the case of either the hierarchical or the non-hierarchical model, the behavioral choice of the sponges was better explained by the carapace width (Figs. 3A, 3C; Table 1). The estimated information gained by the model 1_1 against model 1_3 is 1.35 nat/sample (Table 1). This suggests that larger crabs tend to choose L size sponge rather than M size. However, the crabs larger than about nine cm carapace width did not choose the sponges.

Trimming

After a choice of either the M or L size sponges, the crabs decided whether to trim the extra parts of the sponges (Figs. 4A–4D). Here, we modeled the size of an area in a sponge that was trimmed ($N_{animal} = 30$). The trimming behavior for the sponge followed three patterns (Fig. 4D). They cut off (1) all four corners of a sponge, (2) one corner of the sponges elliptically, or (3) two corners of the sponge linearly. The crabs trimmed the white area



Figure 4 Trimming. (A) Predictive distribution on the hierarchical model 2_1. The white dotted lines connect the data points from the same individual. (B) Predictive distribution visualized by re-scaling the color density of the expanded area in the upper plot except for the zero in the y-axis. (C) Predictive distribution on the non-hierarchical model 2_6. (D) Outline of the trimming process from a choice of a sponge (animals larger than about nine cm skipped the whole behavior); some of the animals skipped the trimming behavior and went directly to cavity making. The drawing is by Keita Harada.

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(Fig. 4D) and started excavating a cavity (Fig. 4C Twenty-three crabs skipped the trimming behavior in 33 trials.

After trimming or skipping, they started excavating. For the behavioral act of trimming, a non-zero data point indicating a trimmed size of the sponge was recorded (Figs. 4A–4C). The size decreased with the increase of the carapace width. If a crab skipped trimming, a data point was recorded at zero, meaning no trimming. Almost all the crabs that chose the M size sponges decided not to trim the sponge except for one individual. Meanwhile, they trimmed less amounts of the sponges relative to the increase of their body sizes when they chose the L size sponges.

The WAIC of the hierarchical model 2_1 was -2.08 and that of comparable nonhierarchical model 2_6 was 7.40 (9.48 nat/sample in difference, Figs. 4A, 4D, Table 1), indicating that the hierarchical model is significantly better than the non-hierarchical one.

Cavity size

Six crabs just cut the sponge and did not excavate the sponge. We modeled the size of a cavity in a cap ($N_{animal} = 30$) as a random variable taken from the gamma distribution



Figure 5 Excavated cavity in a cap. (A) Predictive distribution of a cavity size on the model 3_1. The white points connected by dotted lines are from the same individual. (B) Predictive distribution on the model 3_1. The drawing is by Keita Harada. Full-size DOI: 10.7717/peerj.9036/fig-5

with the log link function (Fig. 5). The size increased with the carapace width, and the model considering individual behavioral types performed best (Table 1). The WAIC of the hierarchical model 3_1 is slightly smaller than that of the comparable non-hierarchical model 3_2 (0.08 nat/sample in the difference; Figs. 5A, 5B, Table 1). The individual with the arrowhead (Fig. 5A) made relatively large cap cavities, indicating large inter-individual differences. As expected, larger crabs made larger cavities. The difference of the WAIC was about 0.1 (Fig. 5B). The predictability improvement is relatively small against that of sponge choice, suggesting that the individual behavioral type would have a lower effect in the determination of cavity size.

Latency

We modeled the latency for cap making (from the choice of sponge to carrying) by 32 crabs as a random variable taken from the zero-inflated distribution (Fig. 6). No obvious relation was found between the carapace width and the latency, and the number of crabs that had carried the cap by the next day. However, the hierarchical model 4_1 outperformed the non-hierarchical model 4_2 (WAIC values, 1.10 and 1.28 respectively, thus 0.18 nat/sample in the difference).

DISCUSSION

First, we provide an explicit rationale and mathematical basis of our statistical approach for the problem of quantification of the behavioral variability within and among individuals. Second, we discuss the empirical data of the sponge crab through the lens of our statistical framework.

Statistical modeling from the predictive point of view

First, as a general theory, we state that the model construction for prediction is appropriate in our case because of the difference between prediction and discovery of true distribution.





Second, from a mathematical point of view, we explain the maximum likelihood (ML) estimation and Bayesian inference, which are typical methods of statistical model construction, and discuss the validity of Bayesian inference and WAIC. Lastly, we consider the novelty of our statistical modeling in terms of the type of prediction and the difference between conditioned-level WAIC and marginal-level WAIC.

Statistical models are broadly divided into those for prediction and those for discovering the true distribution. The two situations are as follows; (a) when there is no distribution that generates data in the finite set of models under consideration, (b) when there is a true distribution generating data in the finite set of models considered. Each situation is formulated as follows. Let n be the sample size.

In case (a), the model is constructed by minimizing the generalization loss G_n :

$$G_n = -\log \int q(x) \log p^*(x) dx, \qquad (25)$$

where q(x) is the true distribution of the data and $p^*(x)$ is the predictive distribution. Predictive distribution is a probability distribution of the unknown new data based on the training data. For instance, the predictive distribution by ML estimation $p^*_{ML}(x)$ is defined by

$$p^*(x) = p(x|\hat{w}(Data)), \tag{26}$$

the likelihood model p(x|w) whose parameter *w* is plugged in the ML estimator (MLE) $\hat{w}(Data)$. Bayesian predictive distribution is defined by the expectation of p(x|w) overall a

posterior distribution *p*_{posterior}(*w*|*Data*):

$$p^{*}(x) = \int p(x|w)p_{posterior}(w|Data)dw = \mathbb{E}_{w}[p(x|w)].$$
⁽²⁷⁾

The generalization loss G_n is decomposed into two terms as below. The first term is independent from the model and the second one is the difference from the true distribution and the predictive one, called Kullback-Leibler divergence:

$$G_n = -\log \int q(x) \log p^*(x) dx \tag{28}$$

$$= -\log \int q(x)\log q(x)dx + \int q(x)\log \frac{q(x)}{p^*(x)}dx.$$
(29)

Thus, the generalization loss quantifies how far a predicted distribution is from the true distribution.

In the case (b), the model is constructed by minimizing the negative log marginal likelihood F_n :

$$F_n = -\log Z_n,\tag{30}$$

where $p_{likelihood}(Data|w)$ is a likelihood, $\varphi(w)$ is a prior distribution, and $Z_n = \int p_{likelihood}(Data|w)\varphi(w)dw$ is the marginal likelihood. Note that Z_n is equal to the normalizing constant of the posterior distribution since

$$p_{posterior}(w|Data) = \frac{p_{likelihood}(Data|w)\varphi(w)}{Z_n}$$
(31)

holds by the definition of conditional probability. In statistical mechanics, F_n is called 'free energy'. Because of $Data = (x_1, ..., x_n)$, the true distribution of a dataset is $Q(x_1, ..., x_n) = \prod_{i=1}^n q(x_i)$. For simplicity, we put $x^n = (x_1, ..., x_n)$. We rewrite the marginal likelihood $Z_n = Z(x^n)$ to emphasize that it is a probability distribution of a dataset. We consider the expectation of F_n overall $Q(x^n)$. Just like the generalization loss, it is decomposed into the model independent term and the Kullback-Leibler divergence from $Q(x^n)$ to $Z(x^n)$:

$$\int Q(x^n)F_n dx^n = -\int Q(x^n)\log Z(x^n)dx^n$$
(32)

$$= -\int Q(x^n)\log Q(x^n)dx^n + \int Q(x^n)\log \frac{Q(x^n)}{Z(x^n)}dx^n.$$
(33)

Thus, the expected free energy quantifies how different the marginal likelihood is from the true distribution. By definition, $Z(x^n)$ is the distribution of a dataset, constructed by the model. In a practical application, a dataset is only an obtained set, hence F_n is to be minimized.

For the above formulation, the following facts are known in statistics. In case (a), the 'effective method' would be a model selection method which can select the model while

minimizing the generalization loss if $n \to \infty$. Obviously, minimizing G_n is the 'effective method'. Moreover, minimizing WAIC is also 'effective'.

On the other hand, in case (b), a model selection method that can select the true distribution if $n \to \infty$, is called 'consistent method'. Although statistical modeling in case (b) is formulated by minimizing the expected free energy $\int Q(x^n)F_n dx^n$, minimizing the free energy F_n is 'consistent'.

Our statistical analysis considers (a). We argue that no distribution generates data in the finite set of models, because our models are descriptive; they are not mechanistic models which represent individual behaviors of the crabs. Hence, it is appropriate to construct predictive models.

Next, we explain estimation methods. ML estimation and Bayesian inference are the typical methods used. However, they can be understood in a unified framework.

An analyst arbitrarily designs the simultaneous distribution of a pair (x, w)

$$p(x,w) = p(x|w)\varphi(w), \tag{34}$$

where x is an observable variable and w is a latent variable. In the ML estimation, the existence of the MLE \hat{w} is assumed and $\varphi(w)$ is set to $\delta(w - \hat{w})$ whose realization is limited to the MLE \hat{w} . As a formality, this can be interpreted as the parameter w, which is the MLE \hat{w} .

Let *w* be a real number. For an arbitrary real number *a*, the function $\delta(w - a)$ satisfies

$$\delta(w-a) = \begin{cases} \infty & w = a \\ 0 & w \neq a \end{cases} \quad \text{and} \quad \int \delta(w-a)dw = 1. \tag{35}$$

Thus, it is clear that $p(Data|w)\delta(w - \hat{w})/Z_n = \delta(w - \hat{w})$ holds.

In Bayesian inference, $\varphi(w)$ is a prior distribution. ML estimation can be described in the same way as the Bayesian inference; ML estimation is the case when the prior distribution is fixed to $\delta(w - \hat{w})$. Accordingly, ML estimation and Bayesian inference can be understood in a unified way for the parameter estimation. The method to be used depends on the purpose of statistical modeling.

In the construction of a predictive distribution, the following theorem has been proven (*Watanabe, 2018*). Let G_n^{ML} be the generalization loss of ML estimation and G_n^{Bayes} be the one of Bayesian inference. The overall expectation symbol across the dataset is denoted by $\mathbb{E}[\cdot] = \int Q(x^n)[\cdot]dx^n$. There are the constants μ, λ ($0 < \lambda < \mu$), which are dependent on a model and the true distribution such that

$$\mathbb{E}[G_n^{ML}] = S + \frac{\mu}{n} + o\left(\frac{1}{n}\right),\tag{36}$$

$$\mathbb{E}[G_n^{Bayes}] = S + \frac{\lambda}{n} + o\left(\frac{1}{n}\right)$$
(37)

hold, where $S = -\int q(x)\log q(x)dx$ is the entropy. Especially, when a prior distribution in Bayesian inference is strictly positive and bounded $(0 < \varphi(w) < \infty)$,

$$0 < \lambda \le d/2 \le \mu \tag{38}$$

holds, where *d* is the dimension of the parameter. If the likelihood and the posterior distribution can be approximated by a normal distribution, the equal signs hold. In this study, we use hierarchical models. For them, the likelihood and posterior distribution cannot be approximated by any normal distribution. Therefore, the Bayesian inference can make the generalization loss smaller than that of ML estimation, i.e., it is appropriate for constructing a predictive model:

$$\mathbb{E}[G_n^{Bayes}] + o(1/n) < \mathbb{E}[G_n^{ML}].$$
(39)

Third, we discuss the evaluation criterion used in our analysis. We consider the case (a) as appropriate, so we are to minimize the generalization loss. In our case, the model selection should be 'effective', thus neither using ML estimation nor the coefficient of determination R^2 is appropriate. However, unfortunately, the generalization loss cannot be computed since the true distribution q(x) is unknown.

When a model is set in which the likelihood can be approximated by a normal distribution, the Akaike information criterion (AIC) can estimate the generalization loss G_n^{ML} with the theoretical proof (*Akaike*, 1974). Moreover, minimizing AIC is the 'effective method' if the above assumption is satisfied. Hence, let A_n be AIC

$$A_n = -\frac{1}{n} \sum_{i=1}^n \log p(x_i | \hat{w}) + \frac{d}{n},$$
(40)

we have

$$\mathbb{E}[G_n^{ML}] = \mathbb{E}[A_n] + o\left(\frac{1}{n}\right).$$
(41)

Note that we write the AIC in the scale of the generalization loss, not the deviance scale $(2nA_n)$.

As mentioned above, our model set includes a hierarchical model, thus it is not appropriate to minimize AIC in the ML estimation. We chose inference in order to decrease the generalization loss G_n^{Bayes} . In addition, there are other advantages of the inference. The widely applicable information criterion (WAIC) *Watanabe* (2010a) can estimate G_n^{Bayes} with mathematical proof even if the likelihood and the posterior cannot be approximated by any normal distribution. Moreover, minimizing WAIC is "effective". Let W_n be WAIC

$$W_n = -\frac{1}{n} \sum_{i=1}^n \log \mathbb{E}_w[p(x_i|w)] + \frac{1}{n} \mathbb{V}_w[\log p(x_i|w)],$$
(42)

we have

$$\mathbb{E}[G_n^{Bayes}] = \mathbb{E}[W_n] + o\left(\frac{1}{n^2}\right),\tag{43}$$

without the assumption of normality for the posterior distribution (Watanabe, 2010a).

Another model evaluation criterion in Bayesian inference is well-known: widely applicable Bayesian information criterion (WBIC) *Watanabe* (2013). Let W_n^B be WBIC. WBIC approximates the free energy: $W_n^B = F_n + O_p(\sqrt{\log n})$ *Watanabe* (2013). It is useful for statistical modeling of case (b). However, in our case, we do not suggest that the mechanistic model is included in the considered model set; instead, we perform for the case (a). Therefore, we concluded that the model evaluation using WAIC was more appropriate than using WBIC. Indeed, WAIC is useful for real data even if we are limited to behavioral data (*Wakita, Kagaya & Aonuma, 2020; Barrett, McElreath & Perry, 2017*).

Lastly, we discuss the difference between conditioned-level WAIC and marginal-level WAIC. Although WAIC is beginning to be used for evaluating models with empirical data, we should be careful to compute the value of a hierarchical model. *Watanabe (2018)* introduces two different definitions of WAIC depending on two different predictions. The often-used definition of WAIC for a hierarchical model is the first case in the book:

$$p_{posterior}(w_k | (x_k)^l) \propto \varphi(w_k | w_0) \prod_{l=1}^{N_{act}} p_{model}(x_{lk} | w_k)$$

$$WAIC_l = -\frac{1}{N_{act}} \sum_{k=1}^{N_{act}} \log \mathbb{E} \left[p_{model}(x_{lk} | w_k) \right] + \frac{1}{N_{act}} \sum_{k=1}^{N_{act}} \mathbb{E} \left[\log p_{model}(x_{lk} | w_k) \right]$$
(44)

$$VAIC_{k} = -\frac{1}{N_{act}} \sum_{l=1}^{1} \log \mathbb{E}_{w_{k}}[p_{model}(x_{lk}|w_{k})] + \frac{1}{N_{act}} \sum_{l=1}^{1} \mathbb{V}_{w_{k}}[\log p_{model}(x_{lk}|w_{k})]$$
(45)

$$WAIC_{conditioned} = \sum_{k=1}^{N_{animal}} WAIC_k$$
(46)

$$k = 1, \dots, N_{animal}; l = 1, \dots, N_{act}$$
 (47)

where $(x_k)^l = (x_k^1, ..., x_k^{N_{act}})$ represents all given data for an individual. Note that the number of behavioral acts of the *k*-th animal is the same (balanced) for simplicity here (The number is unbalanced in our data). It should be noted that the statistical model $p_{model}(x_{lk}|w_k)$ is conditioned upon the w_k assigned to each individual. In other words, in prediction, this model focuses on new acts of the already obtained individuals, whereas our focus is usually on a new act of another individual in order to compare models. In many cases, when studying 'animal personality', we are not usually interested in how our models specifically explain the sampled animals personalities. Instead, we are interested in the distribution of a focused species. This is the reason why we did not use this conditioned-level WAIC. However, it is generally more convenient to use conditional-level likelihood within most Bayesian software, with the unfortunate consequence that conditional-level WAIC is often used. For example, *Mitchell et al. (2016)* uses conditioned-level WAIC to compare a hierarchical model with a non-hierarchical model. Furthermore, the use of the measure R^2 for the evaluation of the model in terms of 'variance explained' is based on how we can minimize the variation in the sample obtained, and not focused on the prediction of the true distribution (*Nakagawa & Schielzeth, 2010*).

In summary, we took 'a' best approach from the predictive point of view and explored 'a' best model rather than a 'correct' model, because it is a natural assumption that the distribution we build would never be correct in the empirical modeling of a natural behavior of the sponge crab.

Making cost and size choice: why did the crabs skip carrying the sponge?

The crabs in our experiments that did not carry caps were larger than those that carried caps. One possibility for the reason would be that when they grow up to some extent, the predators might avoid the crab and the relative energetic cost to make caps might increase. We speculate that this might be a reason why the large crabs did not make nor carry the caps.

Another possibility is that the sponges used in this experiment were smaller than those of the necessary size for the crabs. *Dembowska (1926)* reported that the proportion of caps that fit the size of *D. personata* tended to decrease with the increase of the size of the crabs, and we considered that the decision to skip carrying the caps was because there were few sponges that fit the large crabs. Similarly, the large crabs that skipped cap making and carrying, would carry a cap if a sponge size would be larger than the L size sponge. In addition, no individuals carried the S sponge, because it was too small for all of the crabs to carry. It is likely that a younger and smaller crab than those used in this experiment would carry the S sponge. However, we can not exclude the possibility that the variations of the morphology of dactyls or the molt stage affect the behavior.

CONCLUSION

We focused on the body extending behavior of the sponge crab, since the sponge crabs seemed to rely on the behavioral plasticity to make the living or non-living materials suitable to the animal body. Another crustacean that exhibits the body extending behavior is the hermit crab that is well known to prefer specific shells (*Bertness, 1980; Hazlett, 1981; Wilber, 1990*). Although hermit crabs cannot modify the shells by themselves, they are suggested to recognize and learn the shape of extended shells and the surrounding terrain (*Sonoda et al., 2012*). Therefore, the hermit crabs also might have behavioral types.

McLay (1983) showed the relationship of the body and cap size of the *Cryptodromia hilgendorfi* using a log link function and Gaussian distribution. As shown in the crab, we conditioned the variables on the carapace width of the crab *Lauridromia dehaani*. To consider 'animal personality', it is important to appropriately condition out the variables assumed to have much information about the behavioral variable. The body size is presumed to be an influential variable for the cap making behavior.

Therefore, we conditioned all behavioral variables upon body size and found that the predictability improved by adding an assumption that all behavioral aspects pertained to

the 'individual'. The improvement was larger in the sponge choice than in the cavity size. Because the cavity size was determined by repeated excavation and body rotation, the crab might have used the carapace as a 'measure'. However, in the choice task, the information processing to measure an appropriate size would be less dependent on the measure. We speculate that this makes a room for the emerging individual behavioral types dependent on behavioral plasticity that is unique to an individual.

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Competing Interests

The authors declare there are no competing interests.

Author Contributions

- Keita Harada conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Naoki Hayashi analyzed the data, authored or reviewed drafts of the paper, and approved the final draft.
- Katsushi Kagaya conceived and designed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.

Data Availability

The following information was supplied regarding data availability:

Data, source code, and supplementary videos are also available at Figshare: Harada K., Hayashi N., & Kagaya K. (2020). Supplementary Information for the paper "Individual behavioral type captured by a Bayesian model comparison of cap making by sponge crabs". Figshare. Dataset. https://doi.org/10.6084/m9.figshare.8796119.

Supplemental Information

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