University of Windsor Scholarship at UWindsor

Biological Sciences Publications

Department of Biological Sciences

3-24-2018

Phenotypic integration of behavioural and physiological traits is related to variation in growth among stocks of Chinook salmon

Mitchel G.E. Dender

Pauline M. Capelle

Oliver P. Love University of Windsor

Daniel D. Heath University of Windsor

Follow this and additional works at: https://scholar.uwindsor.ca/biologypub

Part of the Biology Commons

Recommended Citation

Dender, Mitchel G.E.; Capelle, Pauline M.; Love, Oliver P.; and Heath, Daniel D., "Phenotypic integration of behavioural and physiological traits is related to variation in growth among stocks of Chinook salmon" (2018). *Canadian Journal of Fisheries and Aquatic Sciences*. https://scholar.uwindsor.ca/biologypub/1230

This Article is brought to you for free and open access by the Department of Biological Sciences at Scholarship at UWindsor. It has been accepted for inclusion in Biological Sciences Publications by an authorized administrator of Scholarship at UWindsor. For more information, please contact scholarship@uwindsor.ca.

Phenotypic integration of behavioural and physiological traits is related to 1 variation in growth among stocks of Chinook salmon 2 3 MITCHEL G.E. DENDER¹, PAULINE M. CAPELLE², OLIVER P. LOVE², DANIEL D. 4 HEATH¹, JOHN. W. HEATH³, CHRISTINA A.D. SEMENIUK^{1,*} 5 6 ¹ Great Lakes Institute for Environmental Research, University of Windsor, Windsor, Ontario 7 N9B 3P4, Canada 8 ² Department of Biological Sciences, University of Windsor, Windsor, Ontario N9B 3P4, Canada 9 ³Yellow Island Aquaculture Ltd., Heriot Bay, BC, Canada V0P 1H0 10 11 * Corresponding author. Great Lakes Institute for Environmental Research, University of 12 Windsor, 401 Sunset Ave., Windsor, Ontario, Canada N9B 3P4. Tel.: +1 519 253 3000 ext. 3763 13 14

15 *E-mail address:* semeniuk@uwindsor.ca (C.A.D. Semeniuk)

16

17 **Running headline:** Phenotypic integration benefits salmon growth

Page 2 of 32

Can. J. Fish. Aquat. Sci. Downloaded from www.nrcresearchpress.com by UNIV WINDSOR on 04/02/18 For personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record.

18 Abstract

19

20

21

22

23

24

25

26

27

28

29

30

31

32

The selection for a single organismal trait like growth in breeding programs of farmed aquaculture species can counter-intuitively lead to lowered harvestable biomass. We outbred a domesticated aquaculture stock of Chinook salmon (*Oncorhynchus tshawytscha*) with 7 wild stocks from British Columbia, Canada. We then examined how functionally related traits underlying energy management - diel variation in cortisol; foraging, social, and movement behaviours - predicted stock-level variation in growth during the freshwater life history stage, a performance metric under aquaculture selection. Outbreeding generated significant variation in diel cortisol secretion and behaviours across stocks, and these traits covaried, suggesting tight integration despite hybridization. The coupling of nighttime cortisol exposure with daytime behavioural phenotype was the strongest predictor of stock-level variation in body mass. Our results suggest selecting for an integrated phenotype rather than on single mechanistic traits alone can generate the greatest effect on aquaculture fish growth under outbreeding practices. Furthermore, selecting for these traits at the stock level may increase efficiency of farming methods designed to consistently maximize fish performance on a large scale.

33

Key words: Aquaculture, Behaviour, Diel cortisol, Phenotypic integration, Functional traits,
Growth, Chinook salmon,

Page 3 of 32

36 Introduction

37 For the first time on record, people are consuming more farm-raised than wild-caught fish 38 (FAO 2016), and urbanization is simultaneously creating a greater demand for both improved 39 quantity and quality protein due to its recognition in healthy diets (Clapp & Cohen 2009: 40 Henchion et al. 2017). Aquaculture production has been relieving the pressure on wild fish 41 populations by attempting to provide a more sustainable and economic source of protein (FAO 42 2016). However, aquaculture faces the same challenges as other farming practices: to harvest the greatest amount of product at the minimum operator cost. As with any domestication process, 43 44 fish stocks have the tendency to inadvertently become inbred when only selecting the largest, 45 fastest growing individuals as broodstock (Bentsen & Olesen 2002), thereby becoming susceptible to disease (Arkush et al. 2002), experiencing slower growth (Kincaid 1983), or 46 47 having low fecundity (Su et al. 1996). (Re)introducing alleles at fertilization is the classic 48 method of lessening the effects of inbreeding depression via the outbreeding of domestic 49 broodstock with wild individuals (Lehnert et al. 2014). First generation (F1) hybrid offspring of 50 wild-caught and domesticated parents may experience benefits associated with outbreeding such as increased size (i.e., hybrid vigour: Gharrett et al. 1999; Whitlock et al. 2000). However, stocks 51 52 may also experience outbreeding depression (Allendorf et al. 2001; Neff et al. 2011), whereby a shift in mean phenotype occurs and causes a reduction in growth (Tymchuk et al. 2006) and 53 survival (Gharrett et al. 1999; Tallmon et al. 2004; Edmands 2007). These discrepancies can be 54 55 explained through the evolutionary concept of an integrated phenotype (Murren 2012), which 56 recognizes that optimal functioning of the organism requires multiple traits to work in unison. As such, when genetic recombination disrupts the evolutionary orchestration of integrated traits, 57 58 individual performance and fitness can be compromised (Lancaster et al. 2010).

59 To enhance the performance of captive stocks, aquaculture producers should ideally be 60 able to assess whether the genotypes selected for outcrossing will maintain the expression of 61 maximal performance metrics such as growth through the continued integration of key 62 phenotypic traits (e.g., traits related to energetic management) that ultimately increase harvest 63 biomass (Tallmon et al. 2004). One of these metrics is the ability to manage energetic 64 homeostasis, both physiologically and behaviourally, since energetic management is widely 65 recognized as critical for maximizing growth in aquaculture (Boisclair and Sirois 1993; Silverstein et al. 1999). In particular, diel management of baseline levels of glucocorticoids is 66 important for overall energetic balance via the optimal timing of fuel mobilization (e.g., glucose, 67 68 lipids, and fatty acids; Gregory and Wood 1999). Likewise, behavioural traits related to 69 energetics such as aggression, schooling, neophilia, and coping style all directly affect the ability 70 of an individual to acquire, consume, and efficiently convert food to tissue (Gregory and Wood 1999; Almazán-Rueda et al. 2005; Martins et al. 2006). Collectively, physiological and 71 72 behavioural traits related to energy management are expected to impact fitness (Gilmour et al. 2005), yet the linkages among these traits in novel environments remain largely unexplored. 73

74 Here we apply the framework of phenotypic integration (Ketterson et al. 2009; Lancaster 75 et al. 2010; Murren 2012) to assess the impacts of outbreeding on performance at an important 76 early life-history stage within an emerging model of Pacific salmonid aquaculture (Chinook 77 salmon; Oncorhynchus tshawytscha). Specifically, our objectives are to determine whether: 1) 78 outcrossing a domesticated stock with multiple wild populations generates variability in traits 79 expected to impact juvenile growth (e.g., diel cortisol patterns, exposure to cortisol during 80 nighttime and daytime, and behavioural phenotype); 2) physiological and behavioural traits are 81 tightly coupled; and 3) taking an integrative, compared to a traditional, single-trait approach,

Page 5 of 32

82 better predicts variation in growth in an aquaculture setting. We examine these questions using a 83 domesticated stock outbred with wild genotypes from 7 regional populations in coastal British 84 Columbia, Canada. We focused these questions on an early-life stage given the recently 85 highlighted importance of identifying mechanisms controlling early development since these are 86 considered significant for the determination of developmental windows that most impact growth 87 variation and survival (Valente et al. 2013). Given the high degree of local adaptation of 88 semelparous Pacific salmon (Taylor 1991), reflecting the environment-specific functional 89 properties of specific loci (Carlson & Seamons 2008), our chosen system also provides the 90 unique opportunity to examine whether physiology and behaviours associated with energy 91 management become decoupled in a hybrid F1 generation, and whether the expression of energy-92 management phenotypes is optimally related to growth in a novel, captive environment. Given 93 that adaptive covariation between physiology and behaviour should be ultimately linked to 94 performance (McGlothlin et al. 2007), selection for suitable captive phenotypes expressing 95 optimal cortisol-behavioural trait combinations should be critical for producing the ideal farmed fish. As such, we expected hybridized stocks expressing an optimal combination of low daytime 96 97 baseline cortisol (representing low energetic demand), intermediate nighttime baseline cortisol 98 (representing the ability to mobilize energy without entering stress-induced levels; Mommsen et 99 al. 1999), and favorable energy management behaviours (e.g., high foraging rates, high sociality, 100 and lower movement) to exhibit the largest mean body mass.

102

101

103

104

105 Methods

106 Animal husbandry and growth

107 Research was conducted at Yellow Island Aquaculture Ltd. (YIAL) on Quadra Island, British 108 Columbia (latitude: N 50° 7' 59.124"; longitude: W 125° 19' 51.834"). The YIAL Chinook 109 salmon population originated from crosses made in 1985 from two nearby hatcheries: Robertson Creek (latitude: N 49° 18' 37.8792"; longitude: W 124° 57' 36.4392") and Big Qualicum River 110 (latitude: N 49° 21' 56.3616"; longitude: W 124° 39' 6.2964"). The domesticated YIAL stock has 111 112 been maintained in captivity for seven generations. In 2013, sires from 7 wild stocks originating 113 from tributaries on Vancouver Island and the lower mainland of British Columbia (Figure 1) 114 were selected for generating outbred crossed stocks with YIAL dams, with an additional YIAL x 115 YIAL stock serving as a control (Semeniuk et al. submitted). Gamete collection and fertilization 116 occurred during October and November of 2013, with milt of 10 males from each stock 117 fertilizing mixed eggs from 17 YIAL dams, who were the offspring of self-fertilization in one 118 functional hermaphrodite, to control for known maternal effects on growth (Heath et al. 1999). 119 After incubation in vertical-stack trays, fry were transferred on March 14, 2014 to 200 L barrels 120 (mean: 122 ± 5 fish per barrel; range: 120-136) split by stock and duplicated to account for barrel effects (n=16 barrels), with a flow through water system maintained between 10-12 °C and water 121 122 turnover at 1 L/min. All barrels were cleaned approximately every 7 days and dissolved oxygen 123 was monitored once a week and maintained above 80% saturation. Fish were fed to satiation 8 124 times daily between 8:00am to 5:00pm, and mortalities removed every two days (mean mortality rate (March 26-June 4, 2014): $2.52 \pm 3.9\%$ per barrel; range: 0-16 fish). For this study, barrels for 125 126 each stock consisted of a mixture of all 10 families. On June 6, 2014, a random subset of 10-13 127 individual fish were removed from every barrel, individually weighed to the nearest 0.01g, and

fish were returned to their original barrel. Body mass at this stage was used as our performance metric to represent early growth from the fry stage, a critical freshwater life-history stage under strong selection in aquaculture (Piper et al. 1982).

131

132 Blood sampling and cortisol assays

We sampled fish for diel cortisol patterns at 8 months post fertilization, over a three-day period 133 134 beginning on June 25, 2014. Specifically, six fish from each stock were sampled (alternating 135 between barrel replicates) at 6 time periods across the entire diel cycle for a total sample size of 136 36 fish per stock (Figure 2A). These specific sampling times were chosen to represent both key 137 periods over the 24-hour period where fish have been shown to be active behaviourally and physiologically (Thorpe et al. 1987; Gries et al. 1997; Pavlidis et al. 1999; Bremset 2000), to 138 139 capture as much finer-scale variation with day- and night-time periods as possible. Fish were captured via dip net, immediately euthanized in a clove oil solution, and blood was collected via 140 141 caudal severance using heparinized hematocrit tubes. All sampling was completed within 5 142 minutes of the initial disturbance to account for increases in cortisol due to capture and handling, 143 and a given barrel was only sampled once every 24 hours to ensure that individuals had not been 144 disturbed from a previous capture session. Packed red blood cells were then separated from 145 plasma via centrifugation at 14,000 rpm (13,331 g) and plasma was stored at -80°C until further analyses. Baseline plasma cortisol was determined using a cortisol enzyme-linked 146 147 immunosorbent assay (Cayman Chemical, MI, USA) optimized in-house for use in juvenile 148 Chinook salmon (Capelle et al. 2016). Samples were run in triplicate across fourteen assay plates 149 at 1:20 dilution. Individual assay plates included standard curves and common control samples,

Page 8 of 32

and were read at a wavelength of 412nM using a spectrophotometer (Biotek Inc). Inter- and
intra-assay coefficients of variation were 11.0% and 4.6%, respectively.

152

153 Behavioural analysis

All barrels (2 barrels per stock) were each filmed once over a span of seven days (June 3–10, 154 155 2014). GoPro cameras (Woodman Labs, USA) were placed on the bottom center of the barrels at 156 ~6:00am and behaviours were recorded for 1.5 hours. At 1 hour post camera deployment, fish in the barrels were fed ~4.2g of feed. Feed amount was calculated using Taplow Feeding charts 157 158 (Chilliwack, BC, Canada), with daily amounts representing $\sim 2\%$ of the biomass in a barrel to 159 match a satiated growth promotion diet. Behavioural analyses consisted of: assigning a score to 160 the degree of group cohesion, calculating individual swimming velocity, and scoring uniformity of gregarious swimming direction fifteen minutes prior and post the addition of food, at one-161 162 minute intervals. During food presentation, position and movement of a subset of fish (~ 60) were noted, as well as latency (in seconds) to resume pre-feeding behaviours (see supplementary 163 164 materials, Tables S1 and S2). Behavioural videos of the Capilano-River stock were highly 165 distorted due to unforeseen technical issues and thus this population was excluded from all 166 behavioural analysis.

167

168 Statistical analysis

All analyses were conducted using JMP version 12 (SAS Institute Inc.), except where indicated.
 Prior to analysis, model assumptions of normality and homogeneity of variances were tested by
 visual inspection of residual versus predicted plots, and normality was confirmed using the

Page 9 of 32

182

172 Shapiro–Wilk test. Non-normally distributed data were \log^{10} transformed where indicated. In all 173 cases, results were evaluated for significance at $\alpha = 0.05$.

174 To describe diel cortisol variation across all stocks, a general linear model with stock, time of day, and their interaction included as fixed effects was used. Cortisol data were log¹⁰ 175 transformed to achieve normality. Next, the area under the diel cortisol curve (AUC) was 176 177 calculated for daytime (6am-6pm) and nighttime (6pm-6am) cortisol (see Figure 2A), following 178 Pruessner et al. (2003) as an estimate of the amount of hormone fish were exposed to 179 (aggregated at the stock level), and as a means to capture the potential different roles of cortisol 180 across the 24-hour cycle. Daytime and nighttime cortisol exposures were pooled separately for 181 each stock but differences were examined using a paired t-test.

183 Behavioural variables from video recordings were grouped a priori into three categories with known implications for fish growth and consisting of distinct variables: Foraging, Sociality, and 184 185 Movement (Fernö et al., 1988; Fernö et al., 1995; Oppedal et al., 2011). A principal component analysis (PCA) with varimax rotation was used for initial exploration of each behavioural 186 187 category independently to reduce redundancies in the data and to create more homogenous 188 groupings. Each 'factor' extracted explained a minimum of 35% of the variance in behaviour 189 based on the Kaiser criterion (eigenvalue >1) and visual inspection of variance plots (Table S1), 190 and were retained for the second stage of PCA. In the second stage, the rotated factors resulting 191 from each behavioural category were loaded into a PCA to produce an overall behavioural 192 phenotype. This PCA produced only a single component with an eigenvalue score that was >1193 (eignevalue = 2.3), explained 38.3% of the overall variance, and was used in subsequent models. 194 High positive scores were associated with fish that spent longer time feeding, retained group

Can. J. Fish. Aquat. Sci. Downloaded from www.nrcresearchpress.com by UNIV WINDSOR on 04/02/18 For personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record.

cohesiveness, and collectively exhibited consistent swimming directions, suggestive of moreefficient energetic management.

197 To investigate the effects of cortisol and behaviour on growth, analyses were conducted 198 using daytime and nighttime cortisol exposure separately. To first confirm the coupling of 199 cortisol during daytime and nighttime directly with behaviour, we used a linear regression with the behavioural phenotype score regressed against cortisol exposure (AUC_{night} or AUC_{day}). 200 201 Next, we assessed whether each trait in isolation or in combination influenced size using linear 202 regressions with mean body mass per population as the response variable (from the sub-sample 203 of individuals weighed prior to the experiment and averaged across replicate barrels), and population-level behavioural phenotype with nighttime cortisol exposure (AUC_{night}) or daytime 204 cortisol exposure (AUC_{dav}) as the explanatory variables. Both linear and quadratic terms for 205 206 cortisol and behaviour were used in a subset of models to account for any hormetic relationships 207 (Schreck 2010). For model selection, we used a combination of fit, significance, and AIC 208 modified for small sample size (AICc) using the package "AICcmodavg" in R (Mazerolle and 209 Mazerolle 2017).

210

217

All analyses were aggregated to the level of the population since screening at the individual level is not necessarily a feasible practice within aquaculture (Castanheira et al. 2013; Pigliucci, 2003). Instead, determining whether integrated phenotypes exist at the population level is not only practical, but since intraspecific variability influences fundamental evolutionary processes of adaptation (Bennett et al. 2016), it also increases the potential that the desired phenotypes will remain robust under the stresses of captivity in subsequent generations.

218 **Results**

219 Variation in diel cortisol and behaviour among stocks

Stock and time of day interacted to influence baseline cortisol levels ($F_{47,279} = 14.01$, n = 280, p < 0.001). While each stock showed similar diel rhythms in their baseline cortisol (Figure 2A), they differed in overall exposure to cortisol across the full 24-hour cycle (Figure 2B). In addition, daytime cortisol exposure was significantly lower than nighttime exposure (*paired t(6) = 5.59*, p < 0.001). The behavioural phenotype scores for different stocks ranged from -1.88 (Nitinat) to 2.23 (Chilliwack), and were additionally composed of differences in the contributions of each independent behavioural trait (Figure 2C).

227

228 Integrated effects of cortisol and behaviour on growth

Daytime cortisol exposure had a negative linear relationship with behavioural phenotype (R^2 = 229 0.64, p = 0.031; Figure 3), while nighttime cortisol exposure showed no such relationship ($R^2 =$ 230 0.12, p = 0.44). As univariate predictors of performance, nighttime cortisol exposure and 231 232 behavioural phenotype each showed a non-linear, hormetic relationship with body mass (cortisol: $R^2 = 0.94$, p = 0.003, AICc = 6.1; quadratic regression: body mass = (4.44 - 0.002*(AUC_{night} -233 75.31) 2 ; (behaviour: $R^{2} = 0.95$, p = 0.003, AICc = 5.8, guadratic regression: body mass = 234 $(4.03 - (0.12 * behaviour^2))$. AUC_{dav} had no effect on body mass (linear or quadratic 235 relationships; $R^2 < 0.4$, *p*-values > 0.36, AICc's > 11.0). However, the inclusion of both 236 237 nighttime cortisol exposure and behaviour increased the model's predictive ability and its weighting to explain variation in body mass, with stocks exhibiting mid-range in nighttime 238 cortisol exposure and high energy-management behaviours having the largest body masses (R^2 = 239 0.998, p = 0.07, AICc = -143.3; Figure 4A). There was no meaningful or significant integration 240

Can. J. Fish. Aquat. Sci. Downloaded from www.nrcresearchpress.com by UNIV WINDSOR on 04/02/18 For personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record.

of daytime cortisol exposure with behavioural phenotype on body mass. All models tested areincluded in Table S3.

244 **Discussion**

243

245 Rapid human population growth and increases in anthropogenic stressors on wild ecosystems 246 require improved efficiency in alternative food-production systems such as aquaculture to keep 247 pace with world demand for protein (Green et al. 2005). However, aquaculture practices such as 248 selecting for fast growing individuals can inadvertently decrease stock genetic diversity and/or 249 co-select for undesirable traits that will ultimately impact production (Castanheira et al. 2017). 250 We sought to determine whether outbreeding a domesticated Chinook salmon stock with 251 multiple wild populations resulted in the optimal integration of physiological and behavioural 252 traits at the nexus of energetic management, and if this integration lead to maximized 253 performance at an early-life freshwater stage. Outbreeding generated significant variability in 254 physiological and behavioural traits that are associated with energetic management and known to 255 impact growth. As predicted by the theory of phenotypic integration (Murren 2012), growth was 256 influenced by a coupling of nighttime cortisol exposure and behavioural phenotype, specifically 257 with intermediate cortisol values and high energy-management behaviours combining to 258 generate an optimized hormetic relationship to maximize body size. Our results also suggest that 259 examining the effects of the co-variation of multiple functionally related traits on body mass is a 260 stronger approach than examining these traits in isolation. Moreover, relationships that exist 261 between phenotypic traits such as daytime cortisol exposure and behaviour do not imply 262 integration (and by extension maximized performance), highlighting that only within the context 263 of a performance measure can phenotypic integration be revealed. While phenotypic correlations

264 among traits have been extensively studied, fewer have been examined in light of performance-265 or fitness outcomes, and fewer still across multiple populations. This is the first study to show 266 how among-population coupling of physiological and behavioural traits can influence early-life 267 metrics such as growth rate (here measured as body mass), considered important in assessing 268 long-term success in aquaculture (Valente et al. 2013). This work therefore suggests that 269 researchers and producers may benefit from determining how phenotypic integration impacts 270 early-life performance when employing methods such as outbreeding to enhance or rescue 271 breeding programs.

272

273 Impacts of outcrossing on variability in performance-mediated traits

274 Optimal management of energy budgets over both short- and long-term periods within the 275 constraints of local environmental variation is expected to maximize fitness (Fong 1975; Taylor 276 1991; Sanford & Kelly 2011), a critical consideration for production output in aquaculture 277 (Akvaforsk 2005). Outbreeding in our study produced substantial variation in diel physiological 278 and behavioural traits central to daily energetic management. We focused on diel variation in 279 baseline cortisol because it can differentially affect growth by regulating metabolic processes and 280 promoting homeostasis in fish (Mommsen et al. 1999). In diurnal vertebrates, an optimal diel 281 cortisol cycle consists of high levels early in the morning to provide the energy via 282 gluconeogenesis following night fasting (Dallman et al. 1993), to initiate foraging and other 283 activities (Astheimer et al. 1992; Breuner et al. 1999; Breuner and Hahn 2003). Levels then tend 284 to decline for the remainder of the day to avoid the "high cost of living" associated with 285 maintaining high baseline circulating glucocorticoid levels (Sapolsky et al. 2000; Bernier et al. 286 2004). Although our data confirm that all populations showed strong diel rhythms in baseline

cortisol, populations nonetheless differed significantly in patterns of cortisol exposure during
night and day. Outbreeding also generated a range in behavioural traits associated with energetic
management, and when combined with differences in cortisol exposure, revealed certain stocks
to be less ideally suited to captive environments as evidenced by the resultant low body size.

291 The differences observed among stocks are presumably driven by large sire effects that 292 result from genetic differences among the source stocks that have arisen through local adaptation 293 and potentially genetic drift (Fraser et al. 2011). The sire effect would have dominated possible 294 dam effects among stocks given that we used a common inbred line of domestic dams to both 295 minimize maternal effects and equalize any influences across all stocks (as eggs were also 296 pooled). Within our study, not all hybrid stocks showed high performance. While there are many 297 possible causes of this pattern of reduced performance, one may be the poor performance of the 298 F1 generation is due to the intermediate phenotypes generated by hybridization which are 299 unsuited to the domestic parental environments (Frankham et al. 2002; McClelland & Naish 300 2007). Another mechanistic cause of lowered performance in hybrid stocks may be intrinsic 301 outbreeding depression due to the disruption of epistatic interactions in large co-adapted gene 302 complexes (Edmands 1999); however, we do not believe this to be the case since previous 303 studies have shown Chinook salmon F1 hybrids of genetically different parental lines 304 demonstrate no evidence of hybrid breakdown (Lehnert et al. 2014). Instead, suboptimal 305 performance of juvenile Chinook salmon of certain stocks most likely reflects local adaptation of 306 performance traits within freshwater systems that are very important for the success of early life-307 history stages of anadromous, semelparous salmonids (Waples 1991). When different genotypes 308 are raised in common environments, the impacts of these locally adapted phenotypes persist and 309 are known to generate significant variation in performance outcomes (McClelland & Naish

314

310 2007), and so deliberate selection of these tightly integrated traits will be robust past the F1 311 generation and persist through multiple generations. However, the potential for non-additive 312 genetic effects coupled with residual epigenetic effects may interfere with the predictable 313 inheritance of some of the traits measured.

Effects of outcrossing on the outcome of phenotypic integration under novel captive conditions

317 Traditionally, researchers have focussed on single traits rather than multi-trait combinations to 318 predict fitness or performance of wild populations (Gilmour et al. 2005; Killen et al. 2013). 319 However, since traits that depend on common mechanisms for their expression may evolve as a unit (McGlothlin & Ketterson 2008), ignoring the potential for synergistic effects of integrated 320 321 phenotypes on performance can limit our appreciation of their adaptive value (Pigliucci 2003; 322 Ketterson et al. 2009; Laughlin & Messier 2015). In our study, examining interactions between 323 physiology and behaviour suggested that daytime cortisol exposure predicted behavioural 324 phenotype in a simple linear fashion, setting up the expectation that both would interact to 325 impact growth. However, an integrated phenotypic approach revealed it was the combination of 326 exposure to nighttime cortisol and behaviour that was the best predictor of maximized growth 327 across the 7 stocks, with stocks having mid-range nighttime cortisol exposure and maximal 328 energetics behaviour exhibiting the highest body mass (Figure 4A).

Chronically high cortisol levels are known to negatively impact behavioural traits such as foraging and competitive ability due to a reduction in appetite (Gregory & Wood 1999) and can inhibit aggression and locomotion in salmonids (Øverli et al. 2002). As such, prolonged exposure can lead to tertiary responses such as slow growth, reduced immune function, and compromised

333 survival (Pickering & Pottinger 1983; Barton 2002). In our study, stocks with individuals that 334 expressed elevated daytime cortisol exposure may have incurred these behavioural costs that led 335 to reduced growth. During nighttime, when fish were not fed and feeding activity was much 336 reduced (Fraser & Metcalfe 1997), cortisol plays a more direct role in growth: elevated baseline 337 levels maintain energetic homeostasis, allowing stored glycogen in the liver (generated from 338 food consumed and digested during the day) to be mobilized as glucose to fuel the maintenance 339 and growth of tissues (see Dallman et al. 1993). Behaviours that promote energy gain while 340 reducing energetic costs of locomotion (including sociality) are generally assumed to lead to the 341 highest growth (e.g., foraging efficiency: Eklöv, 1992; high cohesiveness: Johnsson 2003). 342 Interestingly, as a sole (univariate) measure, these maximized behaviours seemingly had a 343 counter-intuitive, negative effect on growth (i.e., significant hormetic relationship), and yet as an 344 integrated unit with physiology, these same behaviours conferred the greatest performance. We 345 therefore argue that an integrated approach situated within a performance context is needed to 346 understand how underlying traits will ultimately impact performance in aquaculture. We also 347 demonstrate that with this integrated relationship - visualized as a contour plot (Figure 4B), there 348 exists the potential for further selection of integrated traits to maximize production. For instance, 349 actively selecting for specific behaviours (i.e., greater locomotory-, foraging efficiency, and 350 sociality) - should these traits remain co-varied within a physiological range, can result in even 351 greater mass gains than those observed. This integrated selection approach at the stock level can 352 be of value for other performance traits (such as disease resistance) and in other breeding 353 programs, outside of salmon culture: it is not only feasible for farming methods that must 354 routinely document production data of functional traits and performance on a large scale, but can 355 be used to possibly increase efficiency as well.

Page 17 of 32

356 Conclusions

357 While phenotypic integration can increase organismal performance by optimizing the adaptive 358 phenotype over evolutionary time, it may decrease flexibility in a changing environment 359 (Schlichting 1989). This apparent trade-off is an important realization for evolutionary biologists 360 and applied producers since both robust and flexible phenotypes carry costs due to (1) the 361 potential for a "mismatched" phenotype to a given environment (Hendry 2004; Nosil et al. 2005) 362 and (2) the associated energetic costs of flexibility (DeWitt et al. 1998; Snell-Rood 2013), 363 respectively. In fact, with regards to food production, abrupt changes in environment (e.g., from 364 the wild to captivity, or fresh water to salt water) may better suit a less flexible and instead a 365 more integrated phenotype, as the temporal lag in endocrine activity and its associated 366 behavioural response may be too slow (and take several generations) to adaptively respond to 367 changing conditions (Taff & Vitousek 2016). Since the management of cortisol is intimately 368 involved in the osmoregulatory transition of fish from fresh to salt water (rev. in McCormick 369 2012), we would predict that (above and beyond energetic management) a tighter integration 370 between cortisol and behaviour could also result in a less costly transition. Taken together, 371 appreciating population-level effects associated with integrated energetic management has the 372 ability to better inform researchers and aquaculture producers about early-life stage mechanisms 373 that influence variation in growth and survival (Valente et al. 2013), and that can also undergo 374 intentional selection for increased food production long-term.

376

377 Acknowledgements

378 We thank Yellow Island Aquaculture Ltd, especially Dr. Ann Heath, J. Drown and the YIAL

379 staff (N. Antoniolli, K. Jones, E. Haugen and D. Meester) for fish care and providing the

380 facilities for the experiments. We also thank C. Harris and K. Janisse for lab assistance, and all

field volunteers. This manuscript was improved by the input of three anonymous reviewers. We

382 thank the Natural Sciences and Engineering Research Council (NSERC) of Canada for Strategic

383 Partnership Grant and Discovery Grant funding to DDH, OPL, and CADS, and the Canada

384 Research Chairs program for funding to OPL. All research followed the University of Windsor

Animal Care Committee guidelines (AUPP #13-22) for the care and use of animals and complied

386 with the Canadian Council on Animal Care guidelines.

387	References
388	Akvaforsk Å. 2005 Selection and breeding programs in aquaculture. Gjedrem T, editor.
389	Dordrecht: Springer.
390	Allendorf FW, Leary RF, Spruell P, Wenburg JK. 2001 The problems with hybrids: setting
391	conservation guidelines. Trends Ecol. Evol. 16, 613-622. (doi:10.1016/S0169-
392	5347(01)02290-X)
393	Almazán-Rueda P, Van Helmond ATM, Verreth JAJ, Schrama JW. 2005 Photoperiod affects
394	growth, behaviour and stress variables in Clarias gariepinus. J. Fish Biol. 67, 1029-
395	1039. (doi:10.1111/j.0022-1112.2005.00806.x)
396	Arkush KD, Giese AR, Mendonca HL, McBride AM, Marty GD, Hedrick PW. 2002 Resistance
397	to three pathogens in the endangered winter-run chinook salmon (Oncorhynchus
398	tshawytscha): effects of inbreeding and major histocompatibility complex genotypes.
399	Can. J. Fish Aquat. Sci. 59, 966–975. (doi:10.1139/f02-066)
400	Astheimer LB, Buttemer WA, Wingfield JC. 1992 Interactions of corticosterone with feeding,
401	activity and metabolism in passerine birds. Ornis Scand. 23, 355-365.
402	(doi:10.2307/3676661)
403	Barton BA. 2002 Stress in fishes: a diversity of responses with particular reference to changes in
404	circulating corticosteroids. Integr. Comp. Biol. 42, 517-525. (doi:10.1093/icb/42.3.517)
405	Bennett MG, Whiles MR, Whitledge GW. 2016 Population level responses of life history traits
406	to flow regime in three common stream fish species. <i>Ecohydrology</i> 9,1388-1399.
407	Bentsen HB, Olesen I. 2002 Designing aquaculture mass selection programs to avoid high
408	inbreeding rates. Aquaculture 204, 349-359. (doi:10.1016/S0044-8486(01)00846-8)

430

409 Bernier NJ, Bedard N, Peter RE. 2004 Effects of cortisol on food intake, growth, and forebrain 410 neuropeptide Y and corticotropin-releasing factor gene expression in goldfish. Gen. 411 Comp. Endocrinol. 135, 230-240. (doi:10.1016/j.ygcen.2003.09.016) 412 Boisclair D, Sirois P. 1993 Testing assumptions of fish bioenergetics models by direct estimation of growth, consumption, and activity rates. T. Am. Fish Soc. 122, 784-796. 413 (doi:10.1577/1548-8659) 414 415 Bremset, G. 2000 Seasonal and diel changes in behaviour, microhabitat use and preferences by 416 young pool-dwelling Atlantic salmon, Salmo salar, and brown trout, Salmo trutta. Env. Biol. Fishes. 59, 163–179. (doi:10.1023/A:100769131) 417 Breuner CW, Hahn TP. 2003 Integrating stress physiology, environmental change, and behavior 418 in free-living sparrows. Horm. Behav. 43, 115-123. (doi:10.1016/S0018-506X(02)00020-419 420 X) Breuner CW, Wingfield JC, Romero LM. 1999 Diel rhythms of basal and stress-induced 421 422 corticosterone in a wild, seasonal vertebrate, Gambel's white-crowned sparrow. J. Exp. 423 Zool. 284, 334-342. (doi:10.1002/(SICI)1097-010X(19990801)284:3<334::AID-424 JEZ11>3.0.CO;2-#) 425 Capelle PM, Semeniuk CAD, Sopinka NM, Heath JW, Love OP. 2016 Prenatal stress exposure 426 generates higher early survival and smaller size without impacting developmental rate in a Pacific salmon. J. Exp. Zool. Part A 325, 641-650. (doi:10.1002/jez.2058) 427 428 Carlson SM, Seamons TR. 2008 A review of quantitative genetic components of fitness in 429 salmonids: implications for adaptation to future change. Evol. Appl. 1, 222-238.

(doi:10.1111/j.1752-4571.2008.00025.x)

ord.		
of rec	431	Castanheira MF, Herrera M, Costas B, Conceição LE, Martins CIM. 2013 Can we predict
version	432	personality in fish? Searching for consistency over time and across contexts. PLoS One.
ficial v	433	8, e62037. (doi:10.1371/journal.pone.0062037)
final of	434	Castanheira MF, Conceicao LEC, Millot S, Rey S, Bégout M-L, Damsgard B, Kristiansen T,
m the	435	Höglund E, Øverli Ø, Martins CIM 2017 Coping styles in farmed fish: consequences for
in 04/0 ffer fro	436	aquaculture." Reviews in Aquaculture 9: 23-41.
may di	437	Clapp J, Cohen MJ, editors. 2009 The global food crisis: Governance challenges and
ion. It	438	opportunities. Wilfrid Laurier Univ. Press. 1, 1–288.
UNI V mposit	439	Dallman MF, Strack AM, Akana SF, Bradbury MJ, Hanson ES, Scribner KA, Smith M. 1993
com by age co	440	Feast and famine: critical role of glucocorticoids with insulin in daily energy flow. Front.
and p	441	Neuroendocrinol. 14, 303-347. (doi:10.1006/frne.1993.1010)
wintoaded from www.incresearchpress.com by $ONIY$ with DSOK on $04/02/18$ of manuscript prior to copy editing and page composition. It may differ from the final official version of record	442	DeWitt TJ, Sih A, Wilson DS. 1998 Costs and limits of phenotypic plasticity. Trends Ecol. Evol.
w.mrcr to copy	443	13 , 77–81. (doi:10.1016/S0169-5347(97)01274-3)
om ww t prior	444	Edmands S. 1999 Heterosis and outbreeding depression in interpopulation crosses spanning a
aded ir nuscrip	445	wide range of divergence. Evolution 53, 1757–1768. (doi:10.2307/2640438)
ownlos ted mai	446	Edmands S. 2007 Between a rock and a hard place: evaluating the relative risks of inbreeding
accep	447	and outbreeding for conservation and management. Mol. Ecol. 16, 463-475.
Aquat. it is the	448	(doi:10.1111/j.1365-294X.2006.03148)
. FISD. nuscrip	449	Eklöv P. 1992 Group foraging versus solitary foraging efficiency in piscivorous predators: the
Can. J	450	perch, Perca fluviatilis, and pike, Esox lucius, patterns. Anim. Behav. 44, 313-326.
is Just-	451	(doi:10.1016/0003-3472(92)90037-A)
Can. J. Fish. Aquat. Sci. Do se only. This Just-IN manuscript is the accepte	452	FAO. 2016 The State of World Fisheries and Aquaculture. Rome. 1–204.
se oi		

Can. J. Fish. Aquat. Sci. Downloaded from www.nrcresearchpress.com by UNIV WINDSOR on 04/02/18 For personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record.

462

473

453 Fernö A, Furevik DM, Huse I, Bjordal Å, 1988 A multiple approach to behaviour studies of 454 salmon reared in marine net pens. ICES.

455 Fernö A, Huse I, Juell J-E, Bjordal Å. 1995 Vertical distribution of Atlantic salmon (Salmo salar

456 L.) in net pens: trade-off between surface light avoidance and food attraction. 457

Aquaculture 132, 285–296. (doi:10.1016/0044-8486(94)00384-Z)

- 458 Frankham R, Briscoe DA, Ballou JD. 2002 Introduction to conservation genetics. Cambridge 459 University Press.
- Fraser NH, Metcalfe NB. 1997 The costs of becoming nocturnal: feeding efficiency in relation to 460 light intensity in juvenile Atlantic salmon. Funct. Ecol. 11, 385-391. 461
 - (doi:10.1046/j.1365-2435.1997.00098.x)
- 463 Fraser DJ, Weir LK, Bernatchez L, Hansen MM, Taylor EB. 2011 Extent and scale of local 464 adaptation in salmonid fishes: review and meta-analysis. Heredity 106, 404-420. (doi:10.1038/hdy.2010.167) 465
- Fong H. 1975 Animal physiology: adaptation and environment. Yale J. Biol. Med. 48, 262-263. 466
- 467 Gharrett AJ, Smoker WW, Reisenbichler RR, Taylor SG. 1999 Outbreeding depression in
- 468 hybrids between odd-and even-broodyear pink salmon. Aquaculture 173, 117–129. 469 (doi:10.1016/S0044-8486(98)00480-3)
- 470 Gilmour KM, Wilson RW, Sloman KA. 2005 The integration of behaviour into comparative physiology. Physiol. Biochem. Zool. 78, 669-678. (doi:10.1086/432144) 471

472 Green RE, Cornell SJ, Scharlemann JP, Balmford A. 2005 Farming and the fate of wild nature.

Science 307, 550-555. (doi:10.1126/science.1106049)

Page 23 of 32

ord.		
of recc	474	Gregory TR, Wood CM. 1999 The effects of chronic plasma cortisol elevation on the feeding
ownloaded from www.nrcresearchpress.com by UNIV WINDSOR on 04/02/18 ted manuscript prior to copy editing and page composition. It may differ from the final official version of record	475	behaviour, growth, competitive ability, and swimming performance of juvenile rainbow
	476	trout. Physiol. Biochem. Zool. 72, 286–295. (doi:10.1086/316673)
final of	477	Gries, G, Whalen, KG, Juanes, F, Parrish, DL. 1997 Nocturnal activity of juvenile Atlantic
(2/18 m the	478	salmon (Salmo salar) in late summer: evidence of diel activity partitioning. Can. J. Fish.
on 04/0 ffer fro	479	Aquat. Sci. 54, 1408–1413. (doi: 10.1139/f97-046)
SOR of may di	480	Heath DD, Fox CW, Heath JW. 1999 Maternal effects on offspring size: variation through early
WINE ion. It	481	development of chinook salmon. Evolution 53, 1605–1611. (doi:10.2307/2640906)
- UNIV mposit	482	Henchion M, Hayes M, Mullen AM, Fenelon M, Tiwari, B. 2017 Future protein supply and
com by page co	483	demand: strategies and factors influencing a sustainable equilibrium. Foods, 6, 53.
npress. g and j	484	Hendry AP. 2004 Selection against migrants contributes to the rapid evolution of ecologically
esearch / editin	485	dependent reproductive isolation. Evol. Ecol. Res. 6, 1219-1236.
/w.nrci to cop	486	Johnsson JI. 2003 Group size influences foraging effort independent of predation risk: an
om wv ot prior	487	experimental study on rainbow trout. J. Fish Biol. 63, 863-870. (doi:10.1046/j.1095-
aded fr nuscrip	488	8649.2003.00187.x)
Jownlo ted ma	489	Ketterson ED, Atwell JW, McGlothlin JW. 2009 Phenotypic integration and independence:
. Sci. Ľ e accep	490	hormones, performance, and response to environmental change. Integr. Comp. Biol. 49,
. Aquat pt is th	491	365-379. (doi:10.1093/icb/icp057)
J. Fish anuscri	492	Killen SS, Marras S, Metcalfe NB, McKenzie DJ, Domenici P. 2013 Environmental stressors
Can. t-IN mi	493	alter relationships between physiology and behaviour. Trends Ecol. Evol. 28, 651-658.
his Jus	494	(doi:10.1016/j.tree.2013.05.005)
only. T	495	Kincaid HL. 1983 Inbreeding in fish populations used for aquaculture. <i>Aquaculture</i> 33 , 215–227.
al use (496	(doi:10.1016/0044-8486(83)90402-7)
Can. J. Fish. Aquat. Sci. D For personal use only. This Just-IN manuscript is the accept		
For		

497	Lancaster LT, McAdam AG, Sinervo B. 2010 Maternal adjustment of egg size organizes
498	alternative escape behaviors, promoting adaptive phenotypic integration. Evolution 64,
499	1607–1621. (doi:10.1111/j.1558-5646.2009.00941)
500	Laughlin DC, Messier J. 2015 Fitness of multidimensional phenotypes in dynamic adaptive
501	landscapes. Trends Ecol. Evol. 30, 487-496. (doi:10.1016/j.tree.2015.06.003)
502	Lehnert SJ, Love OP, Pitcher TE, Higgs DM, Heath DD. 2014 Multigenerational outbreeding
503	effects in Chinook salmon (Oncorhynchus tshawytscha). Genetica 142, 281-293.
504	(doi:10.1007/s10709-014-9774-5)
505	Martins CIM, Schrama JW, Verreth JAJ. 2006 The effect of group composition on the welfare of
506	African catfish (Clarias gariepinus). Appl. Anim. Behav. Sci. 97, 323–334.
507	(doi:10.1016/j.applanim.2005.07.003)
508	Mazerolle MJ, Mazerolle MM. 2017. Package 'AICcmodavg'.
509	McClelland EK, Naish KA. 2007 What is the fitness outcome of crossing unrelated fish
510	populations? A meta-analysis and an evaluation of future research directions. Conserv.
511	Genet. 8, 397–416. (doi:10.1007/s10592-006-9178-x)
512	McCormick, SD. 2012 Smolt physiology and endocrinology. Fish Physiol. 32, 199-251.
513	(doi: <u>0.1016/B978-0-12-396951-4.00005-0</u>)
514	McGlothlin JW, Jawor JM, Ketterson ED. 2007 Natural variation in a testosterone-mediated
515	trade-off between mating effort and parental effort. Am. Nat. 170, 864–875.
516	(doi:10.1086/522838)
517	McGlothlin JW, Ketterson ED. 2008 Hormone-mediated suites as adaptations and evolutionary
518	constraints. Philos. T. Roy. Soc. B 363, 1611-1620. (doi:10.1098/rstb.2007.0002)

Page 25 of 32

ġ

1 of reco	519	N
ersion e	520	
ficial v	521	
final of	522	N
2/18 m the 1	523	
on 04/0 ffer fro	524	١
DSOR of may di	525	
/ WINI tion. It	526	
v UNIV inposi	527	N
com by page cc	528	
hpress. ig and	529	(
researc y editir	530	
ww.nrci to cop	531	Ç
rom wy	532	
aded fi nuscrij	533	F
Jownlo ted ma	534	
. Sci. I e accep	535	
. Aquat pt is the	536	F
J. Fish. nuscrij	537	
Can. -IN ma	538	
nis Just	539	F
nly. TI	540	
or personal use only. This Just-IN manuscript is the accept		
persona		
For]		

Mommsen TP, Vijayan MM, Moon TW. 1999 Cortisol in teleosts: dynamics, mechanisms ofaction, and metabolic regulation. *Rev. Fish Biol. Fisher.* **9**, 211–268. (doi:10.1023/A:1008924418720)

Murren CJ. 2012 The integrated phenotype. *Integr. Comp. Biol.* **52**, 64–76.

- (doi:10.1093/icb/ics043)
- Neff BD, Garner SR, Pitcher TE. 2011 Conservation and enhancement of wild fish populations:
 preserving genetic quality versus genetic diversity. *Can. J. Fish Aquat. Sci.* 68, 1139–
 1154. (doi:10.1139/F2011-029)
 - Nosil P, Vines TH, Funk DJ. 2005 Reproductive isolation caused by natural selection against
 immigrants from divergent habitats. *Evolution* 59, 705–719. (doi:10.1554/04-428)
 - Oppedal F, Dempster T, Stien LH. 2011 Environmental drivers of Atlantic salmon behaviour in
 sea-cages: a review. *Aquaculture* 311, 1–8. (doi:10.1016/j.aquaculture.2010.11.020)
 - Øverli Ø, Kotzian S, Winberg S. 2002 Effects of cortisol on aggression and locomotor activity in
 rainbow trout. *Horm. Behav.* 42, 53–61. (doi:10.1006/hbeh.2002.1796)
- Pavlidis 1999. The effect of photoperiod on diel rhythms in serum melatonin, cortisol, glucose,
 and electrolytes in the common dentex *Dentex dentex. Gen. Comp. Endocrinol.* 113, 240250. (doi: 10.1006/gcen.1998.7190)
- Pickering AD, Pottinger TG. 1983 Seasonal and diel changes in plasma cortisol levels of the
 brown trout, *Salmo trutta* L. *Gen. Comp. Endocrinol.* 49, 232–239. (doi:10.1016/0016-6480(83)90139-9)
- Pigliucci M. 2003 Phenotypic integration: studying the ecology and evolution of complex
 phenotypes. *Ecol. Lett.* 6, 265–272. (doi:10.1046/j.1461-0248.2003.00428.x)

	of
	n
	Sic
	vei
	a
	al official
	off
	al
	Ë,
	le 1
by UNIV WINDSOR on 04/02/18	담
05	uo
4	Ē.
n C	fer
0	dif
Q	<u>y</u>
S	ũ
E	It
M	л.
5	Ξ
F	os
5	du
<u>S</u>	. <u>5</u>
В	ē
ō	pag
sss.com by UNIV WINDSOR on 04/	- p
t. Sci. Downloaded from www.nrcresearchpress.com by U	e accepted manuscript prior to copy editing and page composition. It may differ from the final official version of
chp	ng
arc	ΞΞ.
ese	ec.
Ľ.	py
n.	3
M	5
M	or
t. Sci. Downloaded from w	ED
ē	b
df	CI.
de	sn
oa	an
<u>vn</u>	8
00	ted
ci. D	eb
.2	S
ړ <i>ن</i>	ea
ua	th
Åq	IS .
	ĮĮ.
isi	SCL
<u> </u>	nu
	na
Car	z
0	Ξ.
	nsı
	SJ
	EFi.
	nly
	0
	ust
	al
	on
	ers
	ď
	ō
	<u> </u>

551

557

558

559

record.

541 Piper, RG, McElwain, IB, Orme, LE, McCraren, JP, Fowler, LG, Leonard, JR. 1982) Fish
542 hatchery management. United States Department of of the Interior Fish and Wildlife
543 Service, Washington, D.C., USA.

Pruessner JC, Kirschbaum C, Meinlschmid G, Hellhammer DH. 2003 Two formulas for
computation of the area under the curve represent measures of total hormone
concentration versus time-dependent change. *Psychoneuroendocrino*. 28, 916–931.
(doi:10.1016/S0306-4530(02)00108-7)

Sanford E, Kelly MW. 2011 Local adaptation in marine invertebrates. *Annu. Rev. Mar. Sci.* 3,
509–535. (doi:10.1146/annurev-marine-120709-142756)

550 Sapolsky RM. 2000 Stress hormones: good and bad. *Neurobiol. Dis.* 7, 540–542.

(doi:10.1006/nbdi.2000.0350)

Schlichting CD. 1989 Phenotypic integration and environmental changes. *BioScience* 39, 460–
464.

Schreck CB. Stress and fish reproduction: the roles of allostasis and hormesis. 2010 *Gen. Comp. Endocrinol.* 165, 549–556. (doi: 10.1016/j.ygcen.2009.07.004)

556 Semeniuk CAD, Capelle P, Dender M, Devlin R, Dixon B, Drown J, Heath J, Hepburn R, Higgs

D, Janisse K, Lehnert S, Love O, Mayrand J, Mickle M, Pitcher T, Neff B, Semple S,

Smith J, Toews S, Wellband K, Venney C, Heath D. submitted Domestic-wild

hybridization to improve aquaculture performance in Chinook salmon. Aquaculture

Silverstein JT, Shearer KD, Dickhoff WW, Plisetskaya EM. 1999 Regulation of nutrient intake
and energy balance in salmon. *Aquaculture* 177, 161–169. (doi:10.1016/S00448486(99)00076-9)

Page 27 of 32

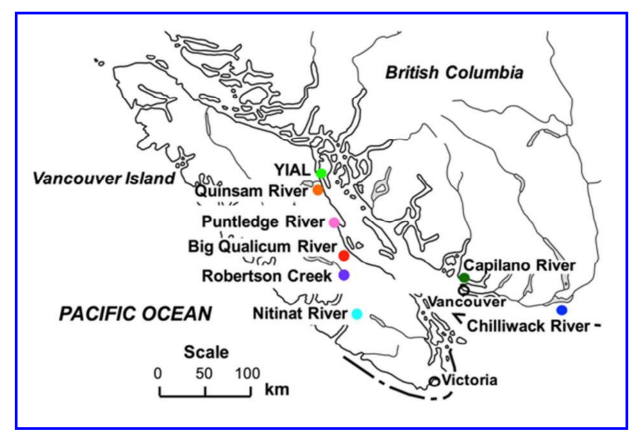
585

563 Snell-Rood EC. 2013 An overview of the evolutionary causes and consequences of behavioural 564 plasticity. Anim. Behav. 85, 1004-1011. (doi:10.1016/j.anbehav.2012.12.031) 565 Su G-S, Liljedahl L-E, Gall GAE. 1996 Effects of inbreeding on growth and reproductive traits 566 in rainbow trout (Oncorhynchus mykiss). Aquaculture 142, 139-148. (doi:10.1016/0044-567 8486(96)01255-0) Taff CC, Vitousek MN. 2016 Endocrine flexibility: optimizing phenotypes in a dynamic world? 568 569 Trends Ecol. Evol. 31, 476–488. (doi:10.1016/j.tree.2016.03.005) Tallmon DA, Luikart G, Waples RS. 2004 The alluring simplicity and complex reality of genetic 570 571 rescue. Trends Ecol. Evol. 19, 489-496. (doi:10.1016/j.tree.2004.07.003) 572 Taylor EB. 1991 A review of local adaptation in Salmonidac, with particular reference to Pacific and Atlantic salmon. Aquaculture 98, 185-207. (doi:10.1016/0044-8486(91)90383-I) 573 574 Thorpe, JE, McConway, MG, Miles, MS, Muir, JS. 1987 Diel and seasonal changes in resting plasma cortisol levels in juvenile Atlantic salmon, Salmo salar L. Gen. Comp. 575 576 Endocrinol. 65, 19-22. (doi:10.1016/0016-6480(87)90217-6) 577 Tymchuk WE, Biagi C, Withler R, Devlin RH. 2006 Growth and behavioral consequences of 578 579 introgression of a domesticated aquaculture genotype into a native strain of coho salmon. 580 T. Am. Fish Soc. 135, 442–455. (doi:10.1577/T05-181.1) Valente, LMP, Moutou, KA, Conceição, LEC, Engrola, S, Fernandes, JMA, Johnston, IA. 2013. 581 582 What determines growth potential and juvenile quality of farmed fish species? Rev. 583 Aquacul. 5, S168–S193. (doi:10.1111/rag.12020) Waples RS. 1991 Pacific salmon, Oncorhynchus spp., and the definition of "species" under the 584

Endangered Species Act. Mar. Fish. Rev. 53, 11-22.

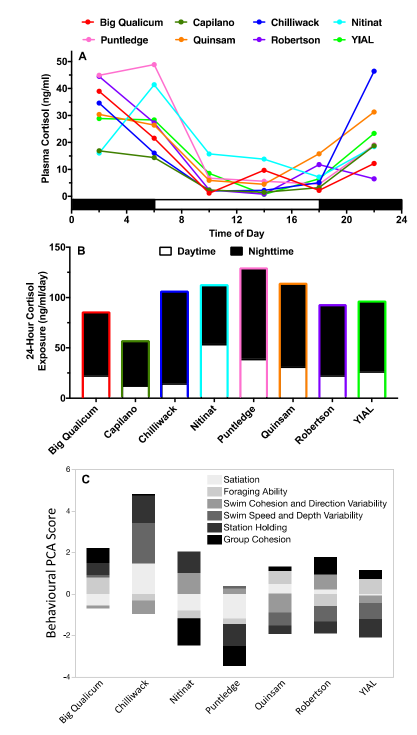
586	Whitlock MC, Ingvarsson PK, Hatfield T. 2000 Local drift load and the heterosis of
587	interconnected populations. Heredity 84, 452-457. (doi:10.1046/j.1365-
588	2540.2000.00693.x)

Figure 1. Map of British Columbia, Canada showing the source of wild Chinook salmon stocks
used for outbreeding in the study. A captive, inbred stock was used as the basis for all crosses
and is held at Yellow Island Aquaculture Ltd. Colours: Bright green – YIAL; purple – Robertson
Creek; orange – Quinsam River; pink – Puntledge River; light blue –Nitinat River; dark blue –
Chilliwack River; dark green – Capilano River; red –Big Qualicum River.



597 daytime and nighttime exposure to baseline cortisol, and (C) variation in behavioural traits of

598 Chinook salmon parr.



Can. J. Fish. Aquat. Sci. Downloaded from www.nrcresearchpress.com by UNIV WINDSOR on 04/02/18 For personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record.

Figure 3. Exposure to daytime cortisol predicts overall behavioural phenotype (combination of foraging, sociality, and movement behaviours) as a linear relationship across 6 outbred populations of Chinook salmon and the inbred control population.

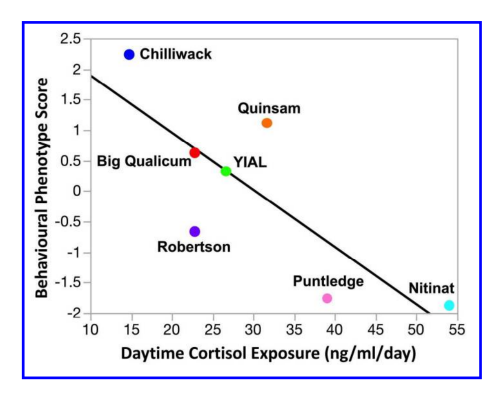


Figure 4. Body mass outcome of optimal phenotypic integration between exposure to cortisol during the night and overall behavioural phenotype (see Methods) across 6 outbred populations of Chinook salmon and the inbred control population (YIAL), represented as both (A) a three-dimensional plot and (B) as a contour map.

А

