1	Effect of food abundance on aggressiveness and territory size of juvenile rainbow trout,
2	Oncorhynchus mykiss
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4	Asra Toobaie [*] , James W.A. Grant
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6	Department of Biology, Concordia University, Montréal, Qc, Canada
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^{*} Correspondence : A. Toobaie, Department of Biology, Concordia University, 7141 Sherbrooke street West, Montreal, QC H2B 1R6, Canada

E-mail addresses: <u>a_toob@live.concordia.ca</u>(A. Toobaie), <u>grant@alcor.concordia.ca</u>(J. Grant).

How food abundance affects (1) whether or not individuals defend territories and (2) what size of 15 territory is defended, are crucial to understanding the role that territoriality plays in regulating 16 the population size of stream-dwelling salmonids. The threshold model of feeding territoriality 17 predicts that territorial defence will be most intense at intermediate levels of food abundance. 18 19 whereas optimal territory-size models predict that territory size will decrease with increasing 20 food abundance. While the latter prediction has been supported by several studies of stream salmonids, too few levels of food abundance were typically used to describe the exact 21 relationship between territory size and food abundance. Hence, to quantify the relationship 22 between the intensity of defence and territory size in relation to food abundance, we manipulated 23 food abundance over a broad range of values (0.62, 1.25, 2.5, 5, 10, and 20% of the fish wet 24 25 body weight in dry food per day) in artificial stream channels (1.92 x 0.77m), while monitoring the territorial behaviour of juvenile rainbow trout, *Oncorhynchus mykiss*. As predicted by the 26 27 threshold model of feeding territoriality, the frequency of territorial aggression was highest at intermediate levels of food abundance, but fish never ceased defending territories entirely, 28 particularly at food densities likely found in the wild. Also as predicted, the aggressive radius of 29 rainbow trout decreased significantly, but only by 30% over a 32-fold increase in food 30 abundance, equivalent to a 51% decrease in territory size. Our results suggest that territory size 31 was relatively insensitive to changes in food abundance, and may have a regulatory effect on 32 population density. 33

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Keywords: juvenile salmonids, food density, territoriality, optimal territory size, aggressive
behaviour

37 The threshold model of feeding territoriality originally predicted that animals defend 38 territories between a lower and upper threshold of food abundance in an all-or-none fashion (Carpenter & MacMillen 1976; Wilcox & Ruckdeschel 1982; Carpenter 1987). However, when 39 individual variation was incorporated into resource defense theory, a more continuous change in 40 41 aggression was predicted (Wolf 1978; Craig & Douglas 1986; Grant 1997). As food abundance increases, the frequency or intensity of territorial aggression initially increases continuously from 42 low-intensity scramble competition to the infrequent defence of non-exclusive territories, to the 43 vigorous defence of exclusive territories, (Craig & Douglas 1986; Grant 1993) and then 44 decreases continuously as food becomes superabundant. Therefore, the threshold model of 45 territoriality, or its more continuous version, predict either an upper and lower threshold for 46 defence or a dome-shaped relationship between the frequency or intensity of territorial 47 aggression and food abundance, respectively (Grant 1993). 48 Considerable evidence of the variability in how frequently individuals defend territories 49 or whether or not territories are defended at all has now been documented in various taxa 50 (Wilcox & Ruckdeschel 1982; Monaghan & Metcalfe 1985; Lott 1991; Goldberg et al. 2001; 51 Grant et al. 2002(). While many studies show a decrease in aggression as food abundance 52 increases (i.e. the right side of the dome-shaped relationship: Magnuson 1962; Slaney & 53 Northcote 1974; Johnson et al. 2004), fewer studies have shown an increase in aggression as 54 food abundance initially increases (i.e. Newman 1956; Keenleyside & Yamamoto 1962), and 55 even fewer have documented the dome-shaped relationship (i.e. Wyman & Hotaling 1988; Grant 56 57 et al. 2002). Given that an individual defends a territory, optimality models (Hixon 1980; Schoener 1983) predict that territory size will decrease with an increase in food abundance and 58

intruder pressure (Schoener 1983). These predictions have now been supported by many studies
in a variety of taxa defending both ephemeral and more permanent territories (Grant 1997;
Adams 2001).

Juvenile stream dwelling salmonids defend feeding territories against conspecifics under 62 63 laboratory and field conditions (Slaney & Northcote 1974; Dill et al. 1981; Keeley 2000; Imre et al. 2004). Because territory size can limit the abundance of juvenile (Grant & Kramer 1990; 64 Elliott 1994) and adult (Jonsson et al. 1998) salmon, any environmental factor that reduces 65 territory size should increase their population density (Grant et al. 1998). Hence, describing the 66 exact relationship between territory size and important environmental variables, such as food 67 abundance, are crucial to understanding the role that territoriality plays in regulating population 68 size. The territory size of juvenile salmonids is inversely related to food abundance in both 69 observational field (Keeley & Grant 1995; Keeley & McPhail 1998) and experimental laboratory 70 (Slaney & Northcote 1974; Keeley 2000; but see Imre et al. 2004) studies. However, laboratory 71 studies typically use only three levels of food abundance, making it difficult to describe the 72 quantitative relationship between territory size and food abundance. If fish vary the size of their 73 74 territory to maintain a constant abundance of food, then territory size is predicted to be inversely proportional to food abundance (a slope of -1.0), and salmonid abundance will be regulated by 75 food abundance rather than territorial behaviour (Grant et al. 1998). For example, a doubling of 76 food abundance would cause a halving of territory size and a doubling of salmonid abundance. 77 However, a review of the available data indicated that a doubling of food abundance causes 78 territories to shrink by only 25% (Grant et al. 1998). 79

80	Our study had two primary goals. First, we tested the competing predictions of the
81	threshold model of territoriality and the more continuous version of the model: (1) territoriality
82	will cease entirely when food abundance is low or in excess; or (2) the frequency of territorial
83	aggression will follow a dome-shaped relationship over a broad range of food abundance.
84	Second, we quantified the relationship between territory size and food abundance in juvenile
85	rainbow trout over a 32-fold increase in food abundance. Specifically, we tested these
86	competing predictions: with increasing food abundance territory size (3) does not change; (4)
87	decreases with a slope of -1.0; or (4) decreases with a slope that is shallower than -1.0.
00	MATERIALS AND METHODS
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89 90	Experimental Subjects
01	Experimental Subjects
92	We obtained 400 young-of-the-year (YOY) rainbow trout from Pisciculture Arpents
93	Verts, Ste-Edwidge-de-Clifton, Québec, Canada. The trout were kept in holding tanks filled with
94	filtered, dechlorinated water on a 12-hour light: 12-hour dark cycle. Water temperature varied
95	throughout the experimental period from 14 to 22°C in May-July and from 14 to 18.7°C in
96	September-November, within the preferred temperature range for rainbow trout (Kerr & Lasenby
97	2000). The fish were fed daily with Vigor # 4 floating feed (Corey Feed Mills) while in holding
98	tanks. Three hundred and sixty trout were used in the experiments. After the experiments were
99	completed, all fish were euthanized with MS-222 as per animal care protocol AREC-2010-
100	GRAN issued to JWA Grant by the Concordia University Animal Research Ethics Committee in
101	accordance with the guidelines of the Canadian Council on Animal Care.
102	

105	Experiments were carried out in artificial stream channels (1.92 x 0.77 m) filled with
106	continuously recirculating (approximately 10% new water per day), filtered, dechlorinated tap
107	water to a depth of approximately 20 cm (mean \pm SD = 19.7 \pm 2.1 cm) on a 12 hour light: 12
108	hour dark cycle (09:00-21:00). Current velocity was recorded at three different points in each
109	tank at the end of each trial (mean \pm SD = 0.032 \pm 0.017 m/s). Water temperature varied with the
110	outdoor temperature and was measured daily for each trial (mean \pm SD temperature = 17.23 \pm
111	2.43 °C). The bottom of each tank was covered with a layer of aquarium gravel overlaid by a
112	four by eight grid of cobbles (mean diameter = 7.84 cm; range = $5.7-10.5$ cm). The cobbles were
113	placed with their centres 15.3 cm apart along the width and 21.3 cm apart along the length of the
114	tank and were used as visual markers to facilitate estimating the relative position of individuals
115	as well as to aid fish in establishing territories (La Manna & Eason 2003). Each stream channel
116	was stocked with 12 fish (mean \pm SD fork length = 4.20 \pm 0.22 cm; mean \pm SD mass = 1.18 \pm
117	0.28 g), equivalent to 8 YOY/m^2 , corresponding to a high density for fish of this size in the wild
118	(Grant & Kramer 1990). The experimental treatments consisted of six different food levels: 0.62,
119	1.25, 2.5, 5, 10, and 20% of the fish wet body weight in dry food per day. Each food level was a
120	multiple of the highest amount of food typically encountered by juvenile salmonids in the wild
121	(Keeley 2000), equivalent to 5% of the wet body weight in dry food (Imre et al. 2004). Each of
122	the food treatments was replicated 5 times, with each trial lasting 7 days. The daily ration of food
123	(Optimum 0.7 granulated fish feed, Corey Feed Mills) was spread evenly on an automatic belt
124	feeder that dispensed the food at the upstream end of the channel at a constant rate over the 12-
125	hour "daylight" period.

Trials were carried out from May-July and September-November 2010. Prior to each 128 129 trial, the initial fork length (to the nearest mm) and body weight (to the nearest 0.01g) of individuals were recorded. Individuals were given a unique mark with fluorescent elastomer tags 130 in the dorsal and/or caudal fins to allow for individual identification. Fish were given a 131 conditioning period of approximately 24 hours before the beginning of each trial to allow them 132 to acclimate to the new environment and to the method of food delivery. On the final day of each 133 trial, a scan sample was conducted in order to record the position of each fish on a schematic 134 135 map. Then, each individual was observed independently for 15 minutes to record the distance and frequency of aggressive acts, such as charges, chases, and nips (sensu Keenleyside & 136 Yamamoto 1962), and the location of each station (sensu Steingrímsson & Grant 2008) from 137 138 which aggressive acts were initiated. The grid of labeled cobbles acted as a simple x-y coordinate system, with the 0.0 (x, y) position in the downstream left corner of stream channel. Following 139 the observation, the final fork length and weight of all individuals were recorded. Specific 140 growth rate was calculated for all individuals as $G = (\log_e W_{\text{Final}} - \log_e W_{\text{Initial}}) / t$, where G is the 141 specific growth rate of weight (W) increase and t is the duration of the trial in days (Ricker 142 1975). 143

In each tank one dominant individual defended a large portion of the stream channel. Of the remaining individuals, called subordinates, some were aggressive while being subordinate to the dominant fish. Aggressive subordinates included territorial individuals that defended a fixed area (sensu Maher & Lott 1995) and "floaters" (sensu Puckett & Dill 1985) that did not defend fixed home ranges or maintain a station, but occasionally displayed aggressive behaviour.

Territories were relatively exclusive areas, except that subordinate territorial individuals did not
chase the dominant. The aggressive rate was obtained for all individuals in each tank by counting
the number of aggressive acts during focal animal observations.

In each tank, the defended area of all territorial fish was estimated by calculating the 152 individual's mean aggressive radius, the distance between the location of the chasee and the 153 position of the chaser when the aggressive act was initiated. We did not estimate territory size 154 using the minimum convex polygon technique because it is sensitive to sample size (i.e. 155 frequency of aggression; Schoener 1981), which varied considerably among treatments. 156 However, mean aggressive radius will be unbiased by differences in the frequency of aggression 157 between trials. The number of territorial subordinates per trial varied from 2 to 11, so we 158 calculated two territory size measurements for each trial: the average chase radius of the 159 dominant fish and the mean of average chase radii for each territorial subordinate. At the lowest 160 161 food abundance, the dominant fish did not show any aggressive behaviour over the course of observation in 2 of the trials, and were excluded from the calculation of aggressive radius. To 162 compare our data with those from the literature, we estimated territory area by first calculating 163 the average aggressive radius for all the fish in a trial and then assuming that territories are 164 circular (see Keeley & Grant 1995). 165

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167 *Statistical Analysis*

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Because food levels increased in a geometric series, we log₁₀-transformed the food abundance before fitting regressions. To meet the assumptions of parametric tests, aggressive radius was also log₁₀-transformed. Because the dominant individual defended a relatively large

portion of the stream compared to other fish, a two-way ANOVA was used to determine whether
there was a main effect of dominance status (dominant versus subordinates) and food treatment
on aggressive radius and growth rate. Only significant interactions between dominance status
and food treatment were reported in the results.

Average water temperature and initial fork length of individuals were added as covariates in all analyses. Initial fork length of the fish did not have a significant effect on any variable, whereas water temperature had a significant effect on aggressive radius. Thus, we reported the results of the two-way ANCOVA in the section on aggressive radius. SPSS ver. 12.0.1 was used for all statistical tests ($\alpha = 0.05$).

- 181 **RESULTS**
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At the beginning of trials, fish initially gathered at the downstream end of the streamchannel. 185 Within 1-2 days, a dominant fish moved upstream and defended an area of the stream channel 186 directly downstream of the feeder. Subordinate territorial individuals (2-11 per trial) defended 187 territories immediately downstream of the dominant's defended area, whereas floaters existed in 188 189 the spaces between territories of other fish. Non-aggressive fish occupied undefended home ranges at the downstream end of the stream channel. While dominant fish were initially larger 190 than subordinates (dominants: mean \pm SD mass = 1.30 ± 0.30 g; subordinates: mean \pm SD mass 191 = 1.17 \pm 0.26; paired t-test: t_{29} = 6.82, P < 0.001), dominant fish were the largest in their groups 192 in only 13 of 30 cases. At the end of the 7-day trials, dominants were still larger than 193 subordinates (dominants: mean \pm SD mass = 2.28 \pm 0.60 g; subordinates: mean \pm SD mass = 194 1.58 ± 0.67 g; paired t-test: $t_{29} = 10.37$, P < 0.001). and were the largest fish in 21 of 30 cases 195 196

¹⁸³ General Behaviour

197 Frequency of Aggression

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The total number of aggressive acts over the course of the 15-minute observation period 199 differed significantly across the food treatments (Fig. 1a; ANOVA: $F_{5, 24} = 11.10$, P < 0.001). 200 201 The total rate of aggression initially increased with increasing food abundance and then declined when food was abundant, as indicated by a significant quadratic term in a polynomial regression 202 after the linear term was first entered in the model ($t_{27} = 4.51$, P < 0.001). On average, the fish 203 engaged in 130 aggressive acts per 15-minute observation during the 5% treatment, whereas 204 aggression rate dropped to only 45 acts per 15 minutes when food was scarce (i.e. 0.63%) or 205 abundant (i.e. 10 and 20%). Fish were satiated and did not eat all the food during the 10 and 20% 206 treatments. 207

The dome-shaped pattern of total aggression was related to two other measures of 208 209 aggression. The percentage of individuals engaging in aggression also differed among food treatments (ANOVA: $F_{5, 24} = 4.70$, P = 0.004). About 40% of individuals were aggressive at low 210 and high levels of food abundance compared to 60-70% at intermediate levels of food abundance 211 (Fig. 1b). The per capita rate of aggression by the aggressive fish also differed significantly 212 among the levels of food abundance (ANOVA: $F_{5, 24} = 11.03$, P < 0.001). As with the other 213 measures, the per capita rate of aggression initially increased with increasing food abundance, 214 peaked at intermediate level of food abundance, and then decreased (Fig. 1c). Both relationships 215 (Figs. 1b, c) were confirmed by a significant quadratic term in a polynomial regression (t_{27} = 216 3.18, P = 0.004 and $t_{27} = 4.92$, P < 0.001, respectively). 217

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219 Territory Size

221	Water temperature had a significant positive effect on aggressive radius (two-way
222	ANCOVA: $F_{1,43} = 10.07$, $P = 0.003$), and hence was included as a covariate in all analyses.
223	Mean aggressive radius of the focal fish differed significantly across food treatments (two-way
224	ANCOVA: $F_{5,43} = 3.46$, $P = 0.01$) and decreased with increasing food abundance for both
225	dominant and subordinate fish (Fig. 2; two-way ANCOVA, linear contrast: $P = 0.005$). Overall,
226	dominant fish had a larger aggressive radius than subordinate fish (two-way ANCOVA: $F_{1, 43}$ =
227	12.50, $P = 0.001$).

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229 Growth

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As food abundance increased, the growth rate of all subordinates increased at a faster rate 231 than dominant individuals as indicated by a significant interaction between dominance status and 232 food treatment (Fig. 3; two- way ANOVA: $F_{5, 48} = 19.56$, P < 0.001). In spite of the significant 233 interaction, the specific growth rate of focal fish increased with increasing food abundance for 234 both dominant (ANOVA, linear contrast: $F_{1,24} = 10.05$, P = 0.004) and subordinate individuals 235 (ANOVA, linear contrast: $F_{1,24} = 504.20$, P < 0.001). Growth rate appeared to decline at the 236 highest food level, but the difference between the 10 and 20% treatments was not significant 237 (Tukey post hoc tests: dominant fish: P = 0.966; subordinate fish: P = 0.422). Dominant fish 238 grew faster than subordinate individuals in all food treatments except the 10 (Tukey post hoc 239 test: P = 0.998) and 20% (Tukey post hoc test: P = 0.999) treatments. 240

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242 *Territory Size versus Food Abundance*

244	We compared the effect of food abundance on territory size in our study to that of Slaney &
245	Northcote (1974) and Keeley (2000); both studies introduced young rainbow trout to three levels
246	of food abundance (Fig. 4). Because the absolute amount of food could not be easily compared
247	between studies, territory size and food abundance were transformed to multiples of the mean
248	territory size and food abundance for each study. Like previous studies, territory size in our study
249	changed inversely with food abundance (Fig. 2), confirming the prediction of models of optimal
250	territory size (Schoener 1983). However, the relationship in our study had a shallower slope
251	compared to those of Slaney and Northcote (1974) and Keeley (2000) (Fig. 4); the mean slopes
252	in their studies were above the 95% C.I. about the slope in our analysis. All the observed slopes
253	were shallower than the slope of -1.0, suggesting that territory size changed less than one would
254	expect if fish were maintaining a constant amount of food in their territories.

255 **DISCUSSION**

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As predicted by the continuous version of the threshold model of territoriality, there was a 257 dome-shaped relationship between the frequency of aggression and food abundance, indicating 258 259 that fish modified their aggressive behaviour to conspecifics in response to changes in food abundance. Despite the broad range of food abundance used, dominant rainbow trout never 260 ceased defending territories in our experiment. As mentioned in the Introduction, many studies 261 show a decrease in aggression when food is superabundant (i.e. Magnuson 1962; Slaney & 262 263 Northcote 1974), mainly due to satiation. In natural streams, however, food abundance is variable and tends to be lower than the rations provided in laboratory experiments (see Keeley 264 2000). Thus, only large temporary increases in invertebrate production, such as caused by heavy 265

266	rainfall or an emergence of insects (Waters 1966), would likely cause satiation of fish in natural
267	streams. On the other hand, in the absence of satiation, fish increase their territorial aggression
268	when food abundance increases - i.e. the left side of the dome-shaped relationship (Keenleyside
269	& Yamamoto 1962; Newman 1956). Hence, our study suggests that the intensity of territoriality
270	(e.g. percent of intruders chased) will increase monotonically over the range of food densities
271	typically found in the wild. However, we cannot rule out the possibility that the increase in
272	aggression rate was due to an increase in the rate of intrusion on territories (see Grant & Guha
273	1993). The only other studies to document a dome-shaped relationship between aggression and
274	food abundance also used a broad range of food abundance with aggression decreasing when
275	food was in excess (Wyman and Hotaling 1989; Grant et al. 2002).
276	As predicted by models of optimal territory size (Hixon 1980; Schoener 1983), the
277	aggressive radius of dominant fish decreased with increasing food abundance. However, a
278	doubling of food abundance caused territories to shrink by only 13% of their initial size, which is
279	much less than the 50% decrease in defended area that would be expected if fish varied the size
280	of their territories to maintain a constant supply of food (Grant et al. 1998). Both Slaney and
281	Northcote (1974) and Keeley (2000) reported a steeper decrease in territory size with increasing
282	food abundance. The steeper slopes in their studies may have been related to fish being allowed
283	to emigrate, which resulted in a decrease in fish density when food abundance was low. Thus,
284	changes in territory size in their studies were caused by changes in both intruder pressure and
285	food abundance. By contrast, in our study fish density was held constant and changes in territory
286	size were the result of changes in food abundance alone.

Stream fertilization has been considered as a habitat management option to increase 287 288 salmonid abundance in nutrient deficient streams (Slanev et al. 1994). The 32-fold increase in food abundance in our experiment caused territories to shrink to only 51% of their initial size, 289 suggesting that an increase in food abundance would have a relatively small effect on territory 290 291 size and fish population density in mesotrophic streams (Grant et al. 1998). However, the 32fold increase in food abundance caused the growth rate of individual fish to increase by 9 fold, 292 suggesting that an increase in food abundance may increase salmonid abundance primarily by 293 affecting the individual growth rate rather than the population density. Similar to our results, a 294 review of four stream fertilization experiments (Grant et al. 1998) revealed that most of the 295 increase in salmonid abundance resulted from an increase in growth rate rather than an increase 296 in population density. Increased growth rate can be beneficial to fish populations, because it can 297 increase the overwinter survival of juveniles (e.g. Hunt 1969) and decrease the age of fish 298 299 migrating to sea (Hutchings & Jones 1998). However, an increased growth rate can also cause a higher proportion of sexually mature male parr (Hutchings & Jones 1998), which may reduce the 300 number of anadromous males returning to spawn. Hence, any attempt to enhance salmonid 301 populations by increasing the productivity of the stream should carefully consider the potential 302 benefits and costs of increasing the growth rate of individuals. 303

As expected, dominant individuals occupied a central upstream foraging station, and had a priority of access to food items as they fell from the feeder. While dominant fish were not initially always the largest individual, they had a larger aggressive radius and a higher growth rate than subordinates over the course of the experiment. Consistent with our results, the dominant status of juvenile Atlantic salmon is influenced more by an individual's aggression

than its size (Huntingford et al. 1990; Thorpe et al. 1992). Dominant fish often have a higher
growth rate than subordinates (Yamagishi 1962; Li & Brocksen 1977), mainly due to differential
access to food (Yamagishi 1962). In addition, the social stress caused by dominant fish may
result in a higher metabolic rate and lower growth rate of subordinates (Millidine et al. 2009),
even if the food consumption by dominant and subordinates is equal (Abbott & Dill 1989; Lee et
al. 2011).

315 Laboratory studies may overestimate the fitness advantage of dominant individuals in the wild (Höjesjö et al. 2002). Natural environments are more complex than simple laboratory 316 environments and may weaken the strength of social interactions. For example, structural 317 complexity can reduce territory size (Imre et al. 2002), lower aggression, reduce the 318 monopolization of food (Basquill & Grant 1998; Sundbaum & Naslund 1998) and decrease the 319 growth advantage of dominant individuals (Höjesjö et al. 2004). In complex habitats, aggressive 320 321 fish may not be able to monopolize food as efficiently, due to a decrease in their ability to detect (Savino & Stein 1982) and chase intruders (Stamps 1984). Therefore, the differences between 322 dominants and subordinates observed in our experiment may be less extreme in a natural stream. 323 In summary, in our experiment the frequency of territorial defence increased and territory 324 size decreased over the range of food abundance expected to be found in the wild. Because 325 territory size was relatively insensitive to changes in food abundance, territoriality should have a 326 regulatory effect on population density. 327

328

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- 471 Figure Captions

473	Figure 1. Effect of food abundance on mean (\pm SE, N = 5) (a) number of aggressive acts
474	$(Y = -117.36 \log_{10}X^2 + 126.57 \log_{10}X + 69.37, r^2 = 0.43, P < 0.001)$, (b) proportion of
475	aggressive individuals (Y = $-40.96 \log_{10}X^2 + 39.52 \log_{10}X + 55$, r ² = 0.29, P = 0.009), and (c) per
476	capita aggressive rate of aggressive fish (chases/fish/min) (Y = $-0.71\log_{10}X^2 + 0.74\log_{10}X + 0.$
477	0.41, $r^2 = 0.48$, P < 0.001). The solid lines represent the quadratic regressions. Note the
478	logarithmic scale on the x-axis.
479	
480	Figure 2. Mean (\pm SE, N = 5) aggressive radius of dominant (\bullet) and aggressive subordinate fish
481	(\blacksquare) in relation to food abundance. The lines represent linear regressions (solid line: $log_{10}Y = -$
482	$0.10 \log_{10}X + 1.55$, $r^2 = 0.19$; dotted line: $\log_{10}Y = -0.11 \log_{10}X + 1.46$, $r^2 = 0.26$). Note that N =
483	3 for dominant fish in the 0.63% treatment and the logarithmic scale on both the axes.
484	
485	
486	Figure 3 . Effect of food abundance on mean (\pm SE, N = 5) specific growth rate of dominant (\bullet)
487	and the average subordinate fish (\blacksquare) in relation to food abundance. Note the logarithmic scale on
488	the x-axis.
489	
490	Figure 4. Effect of food abundance (multiples of the mean) on territory size (multiples of the
491	mean). The solid line is the predicted territory size if fish defend a constant amount of food.
492	Note the logarithmic scale on both axes.
493	Figure 1



Food abundance (% body mass)







