# 1 Social interactions drive efficient foraging and income equality in groups of

- 2 **fish**
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# 10 Abstract

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12 The social interactions underlying group foraging and their benefits have been mostly studied 13 using mechanistic models replicating qualitative features of group behavior, and focused on 14 a single resource or a few clustered ones. Here, we tracked groups of freely foraging adult 15 zebrafish with spatially dispersed food items and found that fish perform stereotypical 16 maneuvers when consuming food, which attract neighboring fish. We then present a 17 mathematical model, based on *inferred* functional interactions between fish, which 18 accurately describes individual and group foraging of real fish. We show that these 19 interactions allow fish to combine individual and social information to achieve near-optimal foraging efficiency and promote income equality within groups. We further show that the 20 21 interactions that would maximize efficiency in these social foraging models depend on group 22 size, but not on food distribution - suggesting that fish may adaptively pick the subgroup of 23 neighbors they "listen to" to determine their own behavior.

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# 36 Impact statement

Analysis and modeling of group behavior of adult zebrafish shows that a specialized social interaction mechanism increases foraging efficiency and equality within groups, under a

- 39 variety of environmental conditions.
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## 41 Introduction

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Living in a group has clear benefits, including expansion of sensory sensitivity (1–4), sharing of responsibilities and resources (5–7), collective computation (1–4, 8–10), and the potential for symbiotic relations between members that would allow for specialization by individual members (7, 11). Understanding the interactions among individuals that give rise to macroscopic behavior of groups is therefore central to the study and analysis of collective behavior in animal groups and other biological systems.

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50 Group foraging is a prominent example of collective behavior, and social interactions among 51 members have been suggested to increase foraging efficiency (12–17) by allowing individuals 52 to combine private and social information about the location of resources and their quality 53 (see however (18, 19) for adverse effects of social information). Theoretical models have been 54 used to study social foraging using various strategies, including producer-scrounger dynamics 55 (20) and the use of `public' information (21–23). These models explored the efficiency of the 56 underlying strategies, their evolutionary stability, as well as the effects of group composition, 57 and of the distribution of resources (20, 22, 24–27). Experimental work, in the field and in the 58 lab, aimed to identify interactions between foraging individuals (10, 28-32) and their 59 dependence on factors such as the distribution of resources, phenotypic diversity among 60 foragers, animal personality, and foraging strategies of mixed species (33–37). The interaction 61 rules studied with these theoretical models and the emerging group behavior had mostly 62 qualitative correspondence to that of real groups, as the predictions of theoretical models 63 were usually not tested against experimental data of groups at the individual level (20, 38-64 41). 65

Moreover, most models studying how schooling or flocking may improve foraging efficiency have explored the case of single sources, or clumped food patches (8–10, 27, 32, 36, 37). Yet, in many real-world situations, animals are likely to encounter distributed food sources, where maintaining a tight group may not be beneficial for all group members. Indeed, schooling and shoaling species have been shown to disperse when confronted with distributed resources (33, 42). A characterization of group foraging for multiple sources with complex spatial distribution is therefore needed.

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74 The ability to track the behavior of groups of individuals with high temporal and spatial 75 resolution under naturalistic conditions (43–47) makes it possible to explore foraging models 76 and behavior quantitatively in groups and individual fish. Here, we studied food foraging by 77 groups of adult zebrafish in a large arena, where multiple food items were scattered and 78 individuals could seek these items freely. We inferred social interaction rules between fish 79 and compared the accuracy of several mathematical models of group foraging based on these 80 rules and the swimming statistics of individual animals. We explored the performance of 81 these models in terms of foraging dynamics and efficiency of food consumption by the group

and of individuals for various resource distributions and group sizes. Our data driven approach 82 allowed us to analyze foraging strategies through the local dependencies between 83 84 conspecifics, and to show that social interactions increase foraging efficiency in real groups of fish. We further used these models to study the effect of social interactions on income 85 86 equality between members of the group. Finally, we use our models to explore the 87 implications of social interactions on the efficiency of collective foraging of larger groups 88 under different distributions of resources, and ask how animals could choose an optimal 89 foraging strategy under varying conditions. 90

### 92 Results

#### 93

94 We studied free foraging of single adult zebrafish and of groups of 3 or 6 fish in a large circular arena with shallow water, where small food flakes were scattered on the surface (Figure 1, 95 96 Figure 1-figure supplement 1A, and Methods). The trajectories and heading of individual fish, 97 the position of flakes, and food consumption events were extracted from video recordings of 98 these experiments using a tracking software that was written in house (48). Tracking of fish identities in the videos were verified and corrected using *IdTracker* (43). Fish were highly 99 100 engaged in the foraging task and consumed all flakes in less than 2 minutes in most cases (Figure 1B, Figure 1-figure supplement 1C, Video 1). The number of recorded consumption 101 102 events varied between groups, since in some cases fish ate only a part of a flake or flakes disintegrated into smaller parts (Figure 1-figure supplement 1B). The time difference between 103 104 flake consumptions reflects the nature of foraging and its efficiency, and so we found that the time it took a group of k fish to consume n flakes,  $T_k(n)$ , was accurately predicted using a 105 106 simple exponential model,

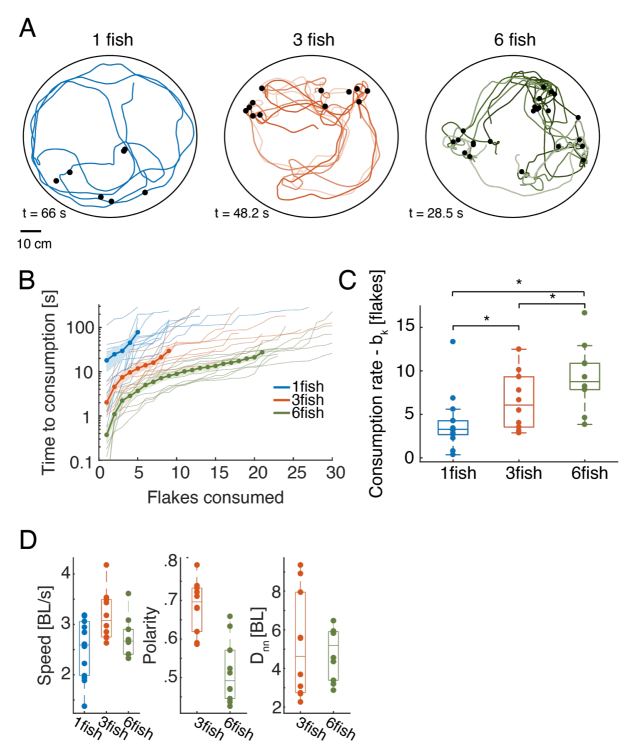
$$\log \hat{T}_k(n) = \frac{n}{b_k} + a_k \tag{1}$$

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109 where parameters  $b_k$  (the "consumption rate" of the fish) and  $a_k$  (time to detection of the 110 first flake) were found numerically to minimize the mean squared error between the model 111 predictions and the data. The correlation between the observed  $T_k(n)$  and model predictions  $\hat{T}_k(n)$  was very high, with median  $r^2$  values: 0.94 [0.09], 0.96 [0.04], 0.96 [0.05] for groups 112 113 of 1, 3, and 6 fish, respectively (brackets show interquartile range). Larger groups consumed the flakes faster than smaller ones (Figure 1B-C), and the feeding rates were higher than those 114 115 predicted just from having a larger number of foragers, which we tested using a model of 116 independent foragers (Figure 1-figure supplement 1D, see models below). Fish in groups of 117 different size differed also in their average swimming statistics, with groups of 3 fish exhibiting higher swimming speeds, higher polarization, and larger variation in group cohesion (nearest 118 119 neighbor distances) (Figure 1D). We also found that group polarity was highly correlated with 120 group cohesion for both groups of 3 and 6 fish, but was not correlated with swimming speed 121 (Figure 1-figure supplement 1E). We therefore asked what social interactions between fish 122 may underlie group foraging and swimming trajectories.

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**Figure 1: Individual and group foraging in adult zebrafish. A.** Example trajectories of groups of 1, 3, and 6 fish foraging for food flakes. Colored lines show the trajectories of individual fish, black dots show the location of flakes consumed by the fish. **B.** Time of consumption of the i<sup>th</sup> flake in the arena is shown for each of the groups tested (thin lines show N=14, 10, and 10 groups of 1, 3, and 6 fish) overlaid with the mean of all groups for every group size (thick line); light shadings represent SEM. (Because individual groups of the same size did not always consume the same total number of flakes, averages were calculated over the first 5, 9, and 21 flakes consumed by the groups of 1, 3, and 6 fish, respectively, and the number of consumed flakes is truncated at 30 for clarity; the full curves are shown in Figure 1-figure supplement 1C; We emphasize that all analyses were

conducted on the full curves). **C.** Boxplots show the rate of flake consumption  $b_k$  that was fitted for each of the groups shown in B. Middle horizontal lines represent median values and box edges are the 1<sup>st</sup> and 3<sup>rd</sup> quartiles; asterisks denote P<0.05 under Wilcoxon's rank-sum test, N = 14,10,10 fish). **D.** Average speed, polarity, and nearest neighbor distance for individual fish and groups of fish. Horizontal lines represent median values and box edges are the 1<sup>st</sup> and 3<sup>rd</sup> quartiles.

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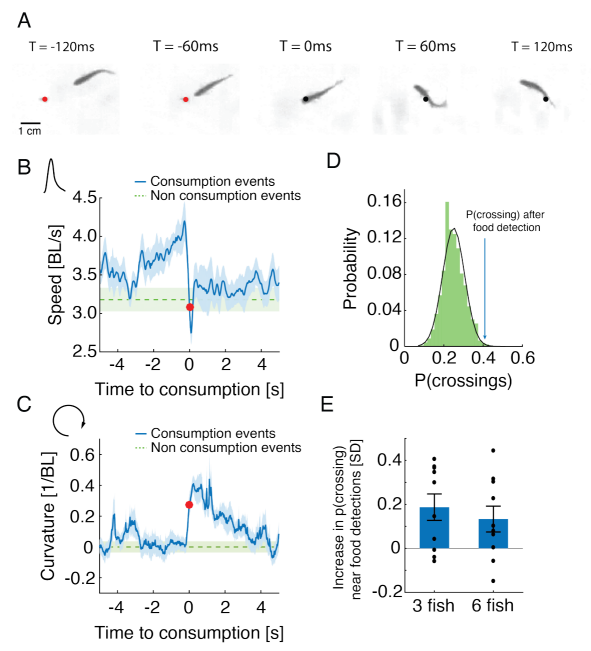
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## 126 Characterizing social interactions during foraging

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128 To explore the nature of interactions between foraging fish, we analyzed individual swimming 129 behavior before and after flake consumption events, and found that fish performed salient 130 and stereotypic maneuvers around the consumption of food items: They increased their 131 speed when approaching food and then abruptly turned in the process of consuming it (Figure 132 2A-C and Video 2). This maneuver was characterized by a decrease in speed and an increase 133 in the curvature of the trajectory (Figure 2B-C, Figure 2-figure supplement 1A). We found that for most groups of 3 and 6 fish analyzed (75%), neighboring fish responded to these salient 134 135 behaviors and were more likely to visit areas of recent flake consumption within 1-4s (Figure 2D-E), much more than expected by chance (P<0.05 for 3 and 6 fish, Wilcoxon's signed rank 136 137 test). Fish were less attracted to the location of a neighbor's consumption maneuver if flakes were more abundant in the arena (Figure 2-figure supplement 1B). To confirm that these 138 139 salient behavioral changes attracted fish to previous consumption areas, and not a physical trace of the flake, we also analyzed ``pseudo consumption events", where fish performed 140 speed changes similar to those seen near flake consumption, but with no food present (see 141 142 Methods). Neighboring fish were indeed attracted to such pseudo-consumption events, 143 affirming that fish responded to the specific behavior of their neighbors (Figure 2-figure 144 supplement 1C-D).

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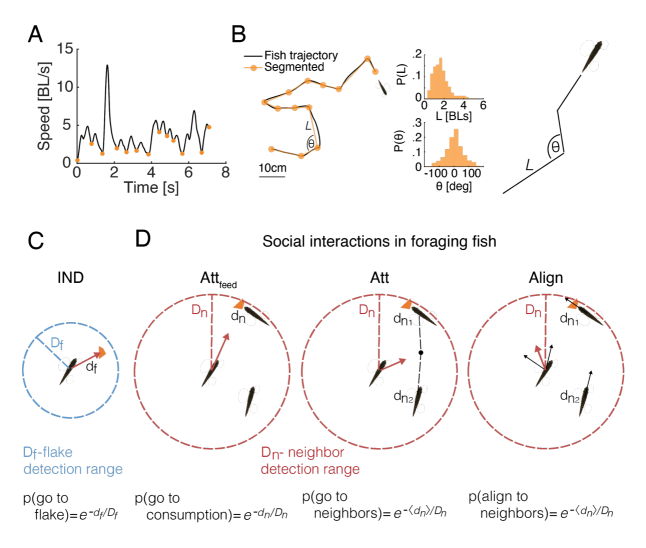
**Figure 2: Stereotypical maneuvers before and during flake consumption by one fish attracts its neighbors. A.** An example of the stereotypical behavior of one fish (in a group of 3) showing flake detection and consumption. Flake position is shown by a red circle before consumption and by a black circle after it has been eaten. **B-C.** Stereotypical behaviors around flake consumption (which we set as time 0 and mark by a red dot) include a transient increase in speed (shown in body length (BL) per second), followed by a sharp decrease (B); this is accompanied by an increase in the curvature of the trajectory (C). Bold blue lines are mean speed and curvature profiles over all detection events of all groups of 3 fish, and dotted green lines show a reference value calculated from random points along the trajectories not related to consumption events (curvature values were normalized such that the average curvature is zero). Light blue and green shadings show SEM. **D.** Probability of neighbors to cross within 2 BLs from the location of a previous flake consumption, within 1-4 s following the consumption event, for one group of 3 fish (blue arrow), compared to the distribution of such neighbor crossing events, if flake consumption events are ignored (Methods). This reference distribution of crossings is well fitted by a Gaussian distribution (mean =

0.25, SD = 0.056), which is shown by a black overlaid line. **E.** Crossing probabilities for groups of 3 and 6 fish show significant increase from the baseline neighbor crossing distribution of each group, similar to C; 0 represent the mean of the baseline crossing distributions, error bars represent SEM.

### 148 Social interaction models of group foraging

To study the implications of attraction to locations of feeding by other fish, we simulated foraging groups of fish with various social interactions and without them (see Methods). Simulations were based on the swimming characteristics of real fish and the empirical spatial distributions of flakes (Figure 1A, Figure 3A-B). The swimming trajectory of each fish was simulated by successive drawing from the distribution of step sizes (the length of the path traveled on discrete 'bouts' according to our segmentation of real fish trajectories; Figure 3A,B) and turning angles (change of heading angle between two discrete bouts) of a specific fish in the real group (Figure 3A-B, Figure 3-figure supplement 1A-B). However, if a flake was within the `range of detection' by a fish  $(D_f)$ , then that fish oriented itself directly towards the flake with a probability that monotonically decreased with the distance to the flake (see Figure 3C). The independent foraging model (IND) is based on a collection of such fish. In addition, we considered 6 social interaction models that combine attraction to neighbors' feeding events and attraction and alignment between fish regardless of feeding (Attraction to feeding events- Att<sub>feed</sub>; Attraction to neighbors- Att; Alignment with neighbors- Align; and their combinations: Att<sub>feed</sub>+Align, Att+Align, Att<sub>feed</sub>+Att+Align). In all these models, the direction of motion of each fish was modulated by the behavior of neighboring fish within the `neighbor detection range'  $D_n$  (Figure 3D and Methods) (22, 25, 49–51). These different models allowed us to tease apart the specific contribution of attraction to neighbor's previous flake consumptions and of the general schooling and shoaling tendencies of the fish (see Figure 3-figure supplement 1C and Methods for a description of all models).

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Figure 3: Comparing social and independent foraging using model simulations. A. An example of the speed profile of an individual fish in a group. Orange dots mark local minima and are used to segment the continuous motion into discrete events. B. Left: A snippet of a fish trajectory, corresponding to the speed profile in A and its segmentation into discrete steps (orange line). Middle: Distributions of step size L and angle change  $\theta$ , between discrete steps over 3 fish in one of the groups (Methods). Right: sketch of a simulated fish trajectory composed of successive drawings of  $\theta$  and L from the empirical distributions. **C.** A sketch of the independent model of fish foraging: At each time step, if a flake was present within a fish's detection range ( $d_f < D_f$  depicted by the blue circle), the fish oriented towards that flake with a probability p(go to flake). D. Sketches of the different social interactions between fish. Each fish may detect consumption of flakes by another fish (left), if that fish was within the neighbor detection range ( $d_n < D_n$  red circle). The observing fish was then attracted to the consumption point with probability p(go to consumption). Additionally, fish may respond to the swimming behavior of neighbors within  $D_n$ , regardless of flake consumption, by swimming towards the average position of their neighbors (middle) with probability p(go to neighbors) or by aligning their swimming direction (right) with neighbors within  $D_{n_{r}}$  p(align to neighbors). Different combinations of these possible social interactions comprise the 6 different social models tested here (Figure 3-figure supplement 1C, Methods).

185 For each of the real groups in our data, we simulated each of the models using a range of possible values of the model's parameters  $D_f$  and  $D_n$  (Figure 3-figure supplement 1C, 186 Methods). For each set of  $D_f$ ,  $D_n$  values, we estimated the accuracy of the models in 187 predicting the sequence of consumption events as well as two swimming statistics of the 188 groups: the average polarity of the group (or alignment between fish) and the average 189 190 cohesion of the group (average distance to the nearest neighbor  $-D_{nn}$ ) (Figure 4A-B, Methods). We evaluated the performance of each model on each of these measures (Figure 191 192 4C), and their combination (Figure 4D). We found that the IND model did not describe well the consumption times of the groups, or their swimming statistics (Figure 4C). Simple 193 attraction to neighbors (Att model) also failed to accurately represent the consumption times 194 or the polarity of the group, yet it accurately described distances between fish and slightly 195 196 improved overall accuracy over the IND model (Figure 4C-D). Alignment interactions among 197 fish (Align model) was significantly better than the Att model, specifically in describing the 198 polarity of the groups and the sequence of consumptions, indicating that fish respond to their neighbors direction while foraging (P<0.005, for groups of 3 fish with N=10, and P<0.01 for 199 200 groups of 6 fish with N=10; Wilcoxon's signed rank test)(Figure 4C-D, Figure 4-figure 201 supplement 1A). The most accurate model of fish swimming and foraging behavior was the 202 one that included both alignment and attraction to neighbors' previous consumption events (Att<sub>feed</sub>+Align), showing high accuracy in describing both the swimming statistics of the groups 203 204 and the sequence of consumptions, significantly better than the competing models (P<0.005 when comparing the Att<sub>feed</sub>+Align to all other models shown, N=10,10 for groups of 3 and 6 205 206 fish, Wilcoxon's signed rank test )(Figure 4C-D, Figure 4-figure supplement 1A). The inferred range of social interactions of the best fitted model (Att<sub>feed</sub>+Align) were ~4 and 8 times larger 207 208 than the range of flake detection (median  $D_n$  values: 21.5 [11], 11.5 [6] and median  $D_f$ 209 values: 2.5[2], 3[3] for groups of 3 and 6 fish, interquartile range in parenthesis; see Figure 4-210 figure supplement 1C).

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Importantly, the observed improvement in accuracy was not a result of increased model complexity. First, the number of model parameters is the same for all social models. Second, models that include attraction to neighbors regardless of flake consumption (Att+Align, Att<sub>feed</sub>+Att+Align models) were less accurate than the Att<sub>feed</sub>+Align model (Figure 4-figure supplement 1B, D). We conclude that fish continuously respond to the swimming direction of their neighbors, but also exhibit a specific attraction to neighbors' previous flake consumptions during foraging.

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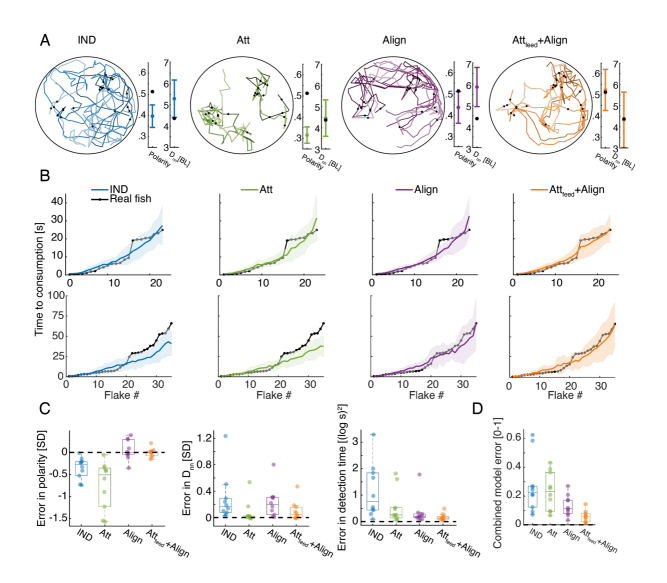


Figure 4: Social models incorporating attraction to neighbors' flake consumptions give the best fit to real foraging groups. A. Example trajectories from simulations of foraging of a group of 6 fish, for the IND, Att, Align, and the Att<sub>feed</sub>+Align models that use the parameters that gave the best fit to real group foraging. Colored lines show different individual fish and black dots are flake positions. Next to the simulated trajectories we plot the average group polarity and nearest neighbor distance in the simulations (colored dots), and the experimental values of the real foraging group (black dots); Error bars represent SD in the simulation. B. Flake detection times (black dots) of two groups of 6 fish (Top row shows the group whose trajectories are shown in A) and the average and standard deviation of the best-fit models (bold colored lines represent averages; shaded areas represent SD). C. Errors of best fit models for groups of 6 fish are shown for three statistics of interest: the polarity of the group  $E^{polarity} = \frac{P^{data} - P^{model}}{P^{data} - P^{model}}$ the nearest neighbor distance  $E^{D_{nn}} = \frac{D_{nn}^{data} - D_{nn}^{model}}{SD(D_{nn}^{data})}$ , and the SD(Pdata) consumption times  $E^{consumptions} = \frac{1}{N} \sum_{n} [log(t^{data}(n)) - log(t^{model}(n))]^2$ , where N is the number of flakes consumed. Dots represent different experimental groups; horizontal lines are median values and boxes represent the 1st and 3rd quartiles. Dotted line represents 0 error in prediction or a perfect fit to the data. D. Combined error of each of the models presented in C.  $E^{combined} = (E^{polarity} + E^{D_{nn}} + E^{consumptions})/3$ , where all error measures were scaled to be between 0 and 1, by dividing by the largest observed error for that measure). The Att<sub>feed</sub>+Align model is significantly more accurate than the IND, Att and Align models (P < 0.005 for all, Wilcoxson's signed rank test, N=10 groups, Methods). Dots represent different groups; horizontal lines are median values and boxes represent the 1st and 3rd quartiles.

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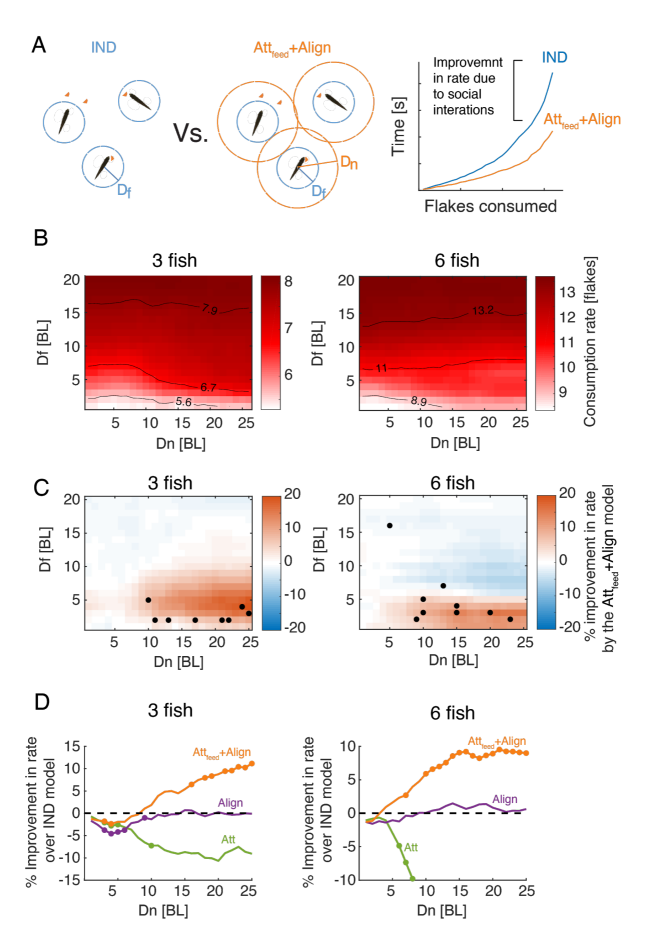
### Increased foraging efficiency is predicted by attraction to neighbor's flake consumptions 227

228 To understand the impact of social interactions on foraging efficiency, we compared the 229 feeding rates of foraging fish in the best fit social model (Att<sub>feed</sub>+Align model) and the 230 reference IND model (Figure 4), for a wide range of model parameter values  $D_f$ ,  $D_n$  (Figure 231 5A). The feeding rates were accurately approximated by an exponential function (as in Eq. 1; 232  $R^2 > 0.98$  for all simulations). As expected, the simulated groups consumed flakes faster as  $D_f$  increased (Figure 5B). For relatively short flake detection range ( $D_f \leq 6$  BL), flake 233 consumption rates increased with  $D_n$ , reflecting the effect of directly responding to 234 neighbors' foraging behavior. For  $D_f > 6$  BL, increasing  $D_n$  had very little effect on 235 236 consumption rates (Figure 5B).

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238 Social interactions in the Att<sub>feed</sub>+Align model resulted in a significant increase in consumption 239 rates, compared to the IND model, only for simulations with low  $D_f$  and high  $D_n$  values (red 240 areas in Figure 5C). Importantly, most groups of real fish were best matched by simulated 241 groups with parameter values that were well within the area of the parameter space were 242 social interactions improve foraging efficiency (low  $D_f$  and high  $D_n$ ), approaching the peak of 243 the expected improvement in foraging performance (Figure 5C). The observed improvement 244 due to social interactions was model specific - social interaction models that did not include 245 attraction to neighbors' previous flake consumptions (e.g. the Align, Att, or Att+Align models) did not show a similar improvement over the independent model (Figure 5D, Figure 5-figure 246 supplement 1A-B). In fact, social foraging strategies that included attraction to neighbors' 247 248 positions (not specifically related to flake consumptions) were less efficient than independent foragers (Figure 5D, Figure 5-figure supplement 1A-B). 249

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**Figure 5:** Attraction to neighbors' feeding results in increased foraging efficiency. A. *Left*: Sketch of two groups of 3 fish foraging, with their different interaction ranges  $D_f$ ,  $D_n$  overlaid. For  $D_n = 0$ , the group is composed of independent foragers (*IND* model). *Right*: foraging efficiency was estimated by comparing the slope (*b*; see eq. 1) of the exponential function fitted to the rate of flake consumption of socially interacting agents (Att<sub>feed</sub>+Align model) and independent (*IND*) foragers. **B.** Average consumption rates, *b*, for different combinations of  $D_f$  and  $D_n$ , the first column on the left ( $D_n = 0$ ) represent independent foragers. Contours denote 10, 50, and 90% of the highest observed rate. **C.** Difference in foraging efficiency for groups that utilize social interactions (Att<sub>feed</sub>+Align) compared to groups of independent foragers (*IND*) for all model parameters. Dots represent  $D_f$  and  $D_n$  values of simulated groups that best fitted real foraging groups. **D.** Average improvement in the rate of flake consumption by socially interacting individuals compared to independent foragers (IND); results were averaged over all simulations with  $D_f \leq 5$  which was the parameter range where real groups were matched by simulations. Dots represent statistically significant differences (P<0.05, Wilcoxon's signed rank test).

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### 252 Individual efficiency and income equality in socially interacting fish

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254 We next explored how the Att<sub>feed</sub>+Align foraging strategy might affect the foraging success of 255 individual members of the group. We simulated groups in which only a fraction of the foragers 256 used social interactions, while the others foraged independently (Figure 6A illustrates these 257 mixed strategy groups). Comparing foraging success of the social and non-social individuals 258 within the same group, we found that individuals using social information consumed up to 20% more flakes than their non-social companions, and this advantage decreased as the 259 260 number of interacting agents in the group increased (Figure 6B-C, Figure 6-figure supplement 261 1A). These effects were most pronounced in models that used the same parameter range that matched real foraging groups, namely low  $D_f$  and high  $D_n$  (Figure 6-figure supplement 1A). 262

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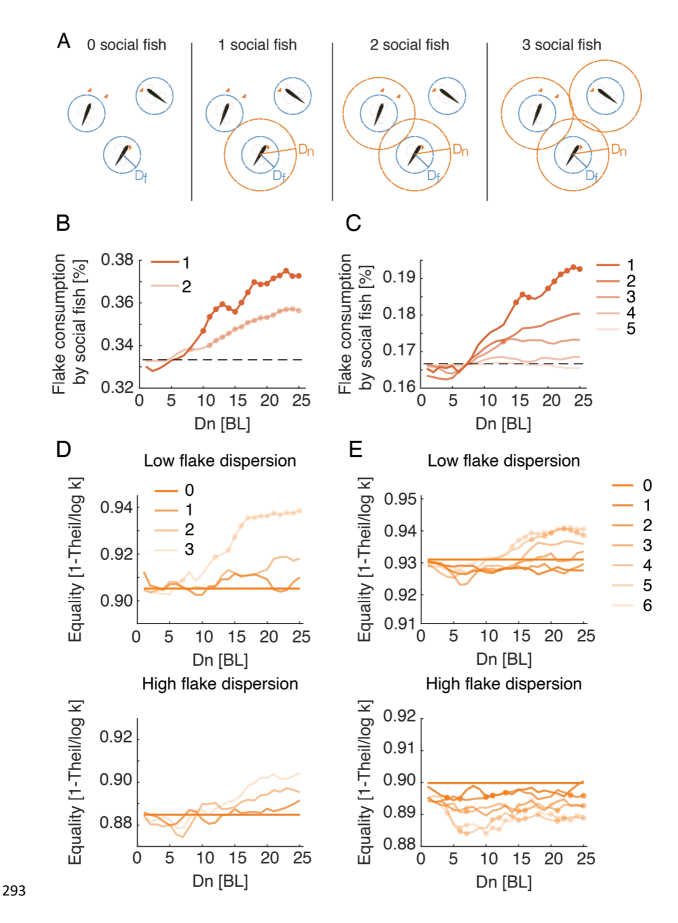
We further assessed the equality of food distribution among individuals in real groups and simulated groups using the Theil index of inequality (52):

$$I_{Theil}(k,n) = \frac{1}{k} \sum_{i=1}^{k} \frac{n_i}{\mu} \log\left(\frac{n_i}{\mu}\right),$$
(2)

where k is the number of fish in the group,  $n_i$  is the number of flakes consumed by the  $i^{th}$ 266 fish, and  $\mu = \frac{n}{k}$  is the average number of flakes consumed by a fish in the group. We 267 normalized  $I_{Theil}$  by its maximal possible value, log k (the case where one fish consumes all 268 flakes), and measured equality using  $1 - \frac{I_{Theil}}{\log k}$ , which ranges between 0 (a single fish who 269 consumed all flakes) and 1 (full equality). Real groups showed high equality values, with 270 271 median values of 0.89 [0.21] for groups of 3 fish and 0.92 [0.16] for groups of 6 fish (values in 272 brackets show the interquartile range). The corresponding simulated groups using the 273 Att<sub>feed</sub>+Align strategy exhibited similar high equality values (Median = 0.92 [0.05], 0.90 [0.1] 274 for simulated groups of 3 and 6 fish), indicating that resources were distributed relatively

equally among real and simulated fish (Figure 6-figure supplement 2A). We then compared 275 276 income equality for the different social foraging models and found that only models that 277 include specific attraction to neighbors' previous flake consumptions exhibited increased 278 equality compared to independent foraging (Figure 6-figure supplement 2C). In simulated 279 groups using the Att<sub>feed</sub>+Align strategy, equality was a function of both the number of socially 280 interacting individuals within the group and the spatial dispersion of the flakes (Figure 6-figure 281 supplement 2B). For low spatial dispersion of flakes, simulated groups composed of just independent foragers showed the highest inequality of all groups with mixed strategies, and 282 283 equality increased with the number of foragers in the group that used the Att<sub>feed</sub>+Align 284 strategy (Figure 6D-E, Figure 6-figure supplement 2D). When all individuals in the group used 285 the Att<sub>feed</sub>+Align strategy, equality was higher than in groups of independent foragers for 286 groups of 3 and 6 fish, for most parameter values that match real foraging groups (Figure 6D-287 E). In contrast, in environments with high dispersion of flakes these effects disappeared or 288 even reversed, with groups of 3 fish showing only a moderate non-significant increase in 289 equality and groups of 6 fish showing a significant decrease in equality (Figure 6D-E). Thus, 290 attraction to neighbors' consumption events increases income equality among group 291 members only when resources are hard to come by. 292

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**Figure 6: Social interactions promote individual foraging efficiency and income equality within groups. A.** Sketch showing simulated groups of 3 fish with a varying number of foragers who use

the Att<sub>feed</sub>+Align strategy during foraging. **B**, **C**. Average fraction of flakes consumed by the socially interacting individuals in the same simulated group. Colors indicate the number of social foragers in the group; dotted lines show the expected consumption values if resources were distributed equally among foragers; results were averaged over all simulations with  $D_f \leq 5$ , which was the parameter range where real groups were matched by simulations; dots represent significant increase in efficiency of social foragers over independent foragers within the group (Wilcoxon's signed rank test, N=10, 10). **D**, **E**. Equality of food distribution within groups, measured by  $1 - \frac{I_{Theil}}{\log k}$  (Eq. 2). Lighter colors represent a larger fraction of social foragers. Data is shown for both low flake dispersion environments (top), and high dispersion (bottom) (Figure 6-figure supplement 2B). Equality values were averaged over  $D_f \leq 5$ ; dots show a significant difference from independent foragers (0 social fish) (Wilcoxon's signed rank test; N=5 for each group size and dispersion level).

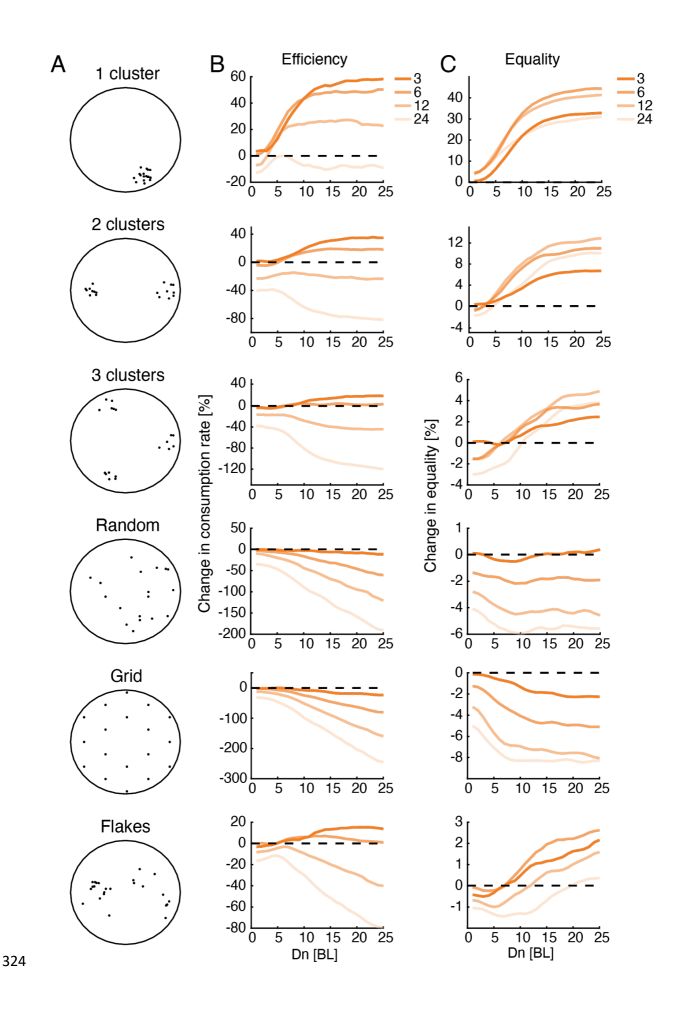
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### 295 Simulating larger groups and different distributions of food

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297 Finally, we investigated the predictions of the Att<sub>feed</sub>+Align foraging model for larger groups 298 and additional spatial distributions of food. We simulated groups of up to 24 fish in 299 environments with spatial distributions of food ranging from a single cluster of food items to 300 a uniform distribution (Figure 7A, Video 3-8). The increase in efficiency due to social 301 interactions was most pronounced when food items were highly clustered in space, whereas 302 for the extreme cases of random or uniform distributions, the models predict that social 303 interactions would hinder foraging performance (Figure 7B, Figure 7-figure supplement 304 1A)(14, 33, 53, 54). Importantly, our simulations predict that groups of 12 and 24 fish that 305 follow the Att<sub>feed</sub>+Align strategy would be less efficient than independent foragers for almost 306 all the food distributions we tested. This is mainly due to the fact that the larger groups are 307 more cohesive and disperse less in the environment, making the search less efficient (Video 8). A social foraging strategy that only includes attraction to neighbors' flake consumption 308 309 events (without a tendency to align with neighbors - Att<sub>feed</sub> model) increased foraging 310 efficiency also for the large group sizes (Figure 7-figure supplement 2A). Income equality in 311 the group also increased in clustered food distributions and decreased in non-clustered ones, 312 for all group sizes (Figure 7C, Figure 7-figure supplement 1B, Figure 7-figure supplement 2B). 313 Interestingly, simulated groups of 3 and 6 fish in the cases of clustered and real flake 314 distributions, were most efficient for intermediate  $D_n$  values (~10-12 BL, see Figure 7B). 315 Simulations of groups with longer social interaction ranges added only a small gain to foraging efficiency. In contrast, for simulated groups foraging with non-clustered food distributions 316 317 (Random and Grid, Figure 7A), increasing  $D_n$  values resulted in decreased efficiency for larger groups, but had almost no effect for groups of 3 fish. These simulations suggest that 318 319 regardless of the flake distribution, optimal interaction ranges for groups of 3 fish could be 320 long, while groups of 6 fish should use intermediate interaction ranges to balance their gains 321 at high clustered environments with their losses at distributed environments. The parameter 322 values of the best fit models to real groups conform with these predictions with median  $D_n$ values of 21.5 and 11.5 for groups of 3 and 6 fish respectively. 323

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#### 325 Figure 7: Optimal interaction range depends on group size and not on resource distribution. A.

326 Sketches showing six different resource distributions used for foraging simulations, three with varying 327 levels of clustering (1, 2 and 3 clusters), a random distribution, and uniform distribution (approximated

328 by a hexagonal grid), and an example of the distribution of flakes taken from one of the experiments.

B. Improvement in group foraging rates (mean time to consumption per flake) of the Att<sub>feed</sub>+Align

- 330 model compared to independent foragers (IND model) for groups of 3, 6, 12, and 24 fish (dark to light
- 331 colors) for the different spatial distributions (panels from top to bottom). Results were averaged over
- all simulations with  $D_f \leq 5$  (Figure 7-figure supplement 1, 100 repetition per parameter combination,
- 333 Methods). C. Same as in B but for the increase in equality of resources within groups
- 334
- 335

### 337 Discussion

#### 338

339 We studied free foraging behavior in groups of adult zebrafish and found that fish responded 340 to the salient swimming maneuvers of shoal mates that indicated the presence of food, by 341 swimming to these locations. Mathematical models of group behavior that combined the 342 tendency of fish to align with one another and to attract to the locations of previous flake 343 consumptions by other fish, accurately described fish foraging behavior and their success rates, and were superior to several other (commonly used) social interaction models. This 344 345 foraging strategy increased efficiency of groups specifically in models that best matched real 346 foraging groups, improved income equality within the groups, and was efficient under 347 different resource distribution settings. Simulations of the models also show that socially 348 interacting individuals that would rely on attraction to feeding events by other fish would 349 consume more food than shoal mates that forage independently. Our results thus present a 350 detailed social foraging heuristic that matches fish behavior in a naturalistic context, and 351 constitutes a highly efficient and robust foraging strategy.

352

353 Our modeling predicts that the inferred interaction ranges that best fit real foraging groups 354 would result in a robust foraging strategy for groups of 3 and 6 fish for various spatial 355 distributions of food. This implies that to forage efficiently fish could adjust their interaction 356 range according to the perceived group size, regardless of the (usually unknown) distribution 357 of food. Additionally, the reduction in foraging efficiency predicted by our models for larger 358 simulated groups (12 and 24 fish) predicts that these groups are more likely to break down 359 into smaller groups that will exhibit increased efficiency. This finding is consistent with the 360 observation that zebrafish both in the wild and in the laboratory are rarely found in cohesive 361 groups of 12 fish or more (48, 55). Interestingly, when simulating groups that only utilize 362 attraction to neighbors' consumption events (without the general schooling tendency 363 observed in real fish) the models predicted increased efficiency for all group sizes. We therefore hypothesize that this interaction type represents a general behavioral strategy for 364 365 individuals foraging in a social context, also for non-schooling species (16, 21, 31, 32).

366

367 Since zebrafish rely heavily on their visual system (52, 53), our modeling focused on vision as 368 the main source of social information between individuals. It is likely that other sensory 369 modalities, namely tactile or odor pathways, also play a role in information transfer during 370 foraging. However, the inferred parameters of the best fit models in our data indicated that 371 neighbor detection ranges were ~4-8 times larger than flake detection ranges - reaching up to 21 body lengths, on average. It is unclear whether odors or tactile information may be 372 373 detected from such large distances on such short time scales (54). Thus, while various 374 modalities may modulate fish behavior, we suggest that vision plays the prominent role in 375 processing social information during foraging.

377 We focused here on attraction-based strategies for the fish, since in our experiments fish 378 were attracted to areas where neighbors detected food. However, previous studies have 379 suggested other search strategies that were based on repulsion between individuals (56) or 380 on maximizing information about the location of food (8). Although our modeling framework 381 gave an excellent fit to the data, it is possible that foraging fish combine or alternate between 382 strategies in different environmental conditions, based on group composition or their internal 383 state (34, 35, 48, 57). Thus, models that incorporate additional strategies according to an 384 explicit policy, might prove to be even more accurate in explaining fish behavior.

385

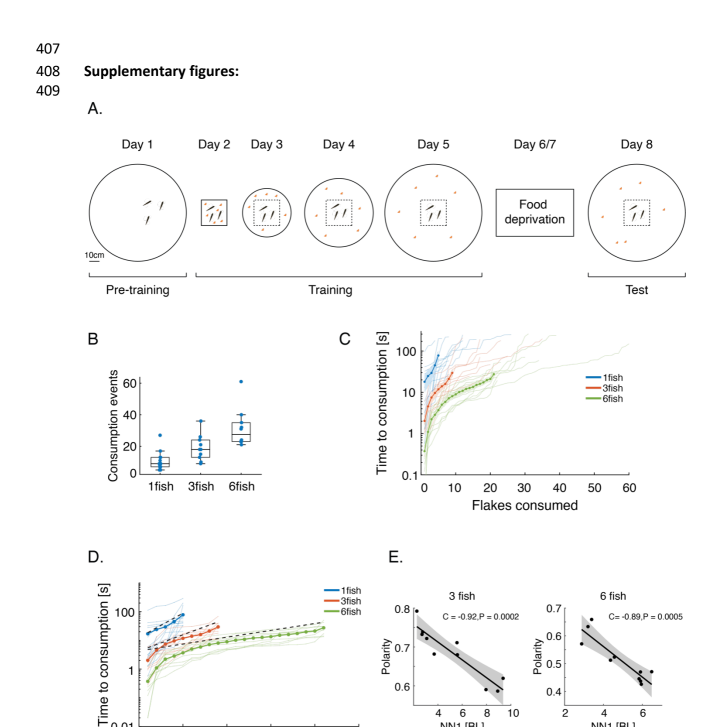
386 Observations of groups in nature, and related theoretical models, suggest that groups may 387 contain a fraction of individuals whose search is based on their personal information 388 (``producers") as well as individuals that rely mostly on social information (``scroungers") (20, 389 58, 59). Our results suggest that when individuals in the group have similar foraging 390 capabilities and a limited social interaction range (22, 60), using both individual and social 391 information is the most efficient strategy for the individual. An interesting extension of our 392 models would be to explore individual differences between members of the group and their 393 effect on individual social foraging strategies (36, 37, 61, 62), or the existence of stable sub-394 groups of individuals with higher tendencies to interact with one another in larger groups of 395 foragers.

396

397 Finally, we note that our work reflects the power of detailed behavioral analysis of individuals 398 in real groups for building accurate mathematical models of social interactions. Learning the 399 models from the data and testing them on real groups allowed us to explore the efficiency 400 and robustness of the interactions among group members in a quantitative manner. This data-driven approach would be applicable and potentially imperative in the analysis of social 401 402 behavior of large groups of individuals - where macroscopic or mean field like models would 403 not suffice to characterize the interplay between complex and possibly idiosyncratic traits of 404 individuals and emerging group behavior.

- 405
- 406

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410

0.01

0

5

10

15

Flakes consumed

20

25

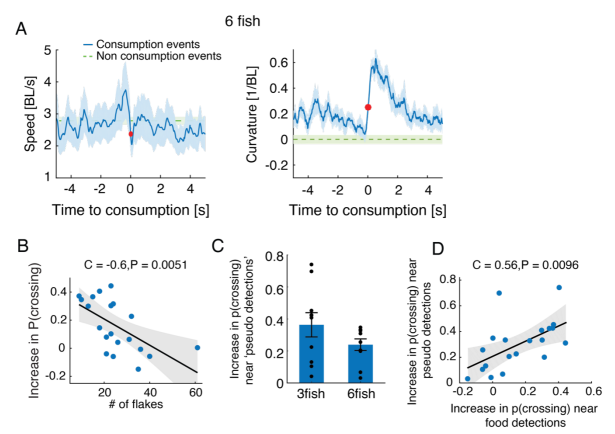
Figure 1-figure supplement 1: Experimental design for studying free foraging behavior in zebrafish groups. A. To facilitate food searching behavior we conducted a 5-day gradual acclimation procedure: After being introduced to the experimental arena with no food present (day 1), on each of the following days (2-5), fish were placed in a box in the middle of the arena, and then released to an experimental tank (which was larger on each day) where small flakes of food were scattered on the water surface (orange dots). Fish were given 5 minutes to explore the tank and consume the flakes. Next, fish were deprived of food for 2 days (days 6-7) in their home tanks, and on day 8 we tested foraging behavior of the fish in the large experimental tank; fish images in this panel are not to scale. B. Number of consumption events recorded for each of the groups tested. Horizontal lines show the median values of consumption events, and the rectangles show the 1<sup>st</sup> and 3<sup>rd</sup> quartiles.

NN1 [BL]

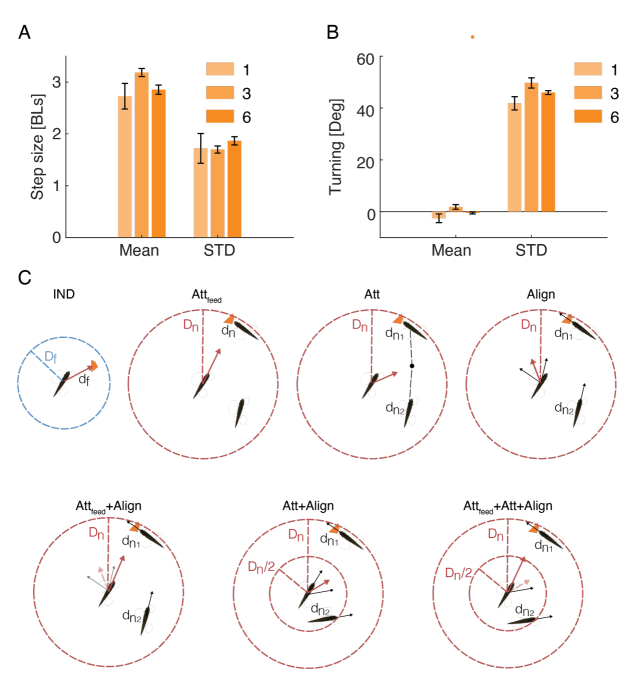
NN1 [BL]

**C.** Same data as in Figure 1B, showing all consumption events for all group sizes (soft lines) and the averages for the first 5, 9, and 21 flakes as depicted in Figure 1B (dotted lines). **D.** Same as in Figure 1B, but showing in addition the expected average rate of simulated independent groups (dotted black lines) (see Figure 3 and Methods), indicating that social interactions increase foraging rates in real groups. **E.** Average group polarity is strongly correlated with average nearest neighbor distance in groups of 3 and 6 fish (Pearson's correlation coefficient, for 10 groups in each case). Dark lines represent the best fit linear model and shaded area is the 95% confidence interval of the model.

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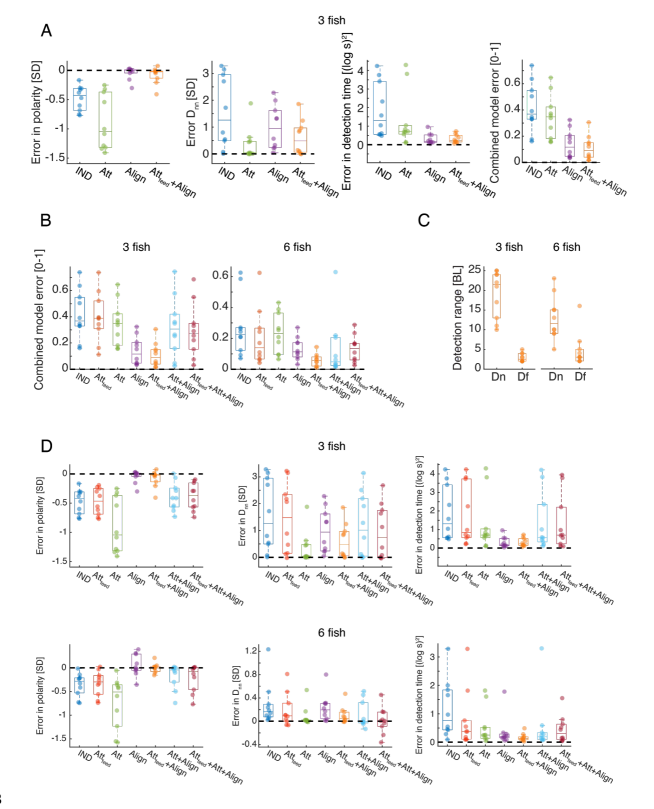


**Figure 2-figure supplement 1: Stereotypical maneuvers before and during flake consumption by one fish attracts its neighbors. A.** Average speed and curvature near flake consumption of groups of 6 fish, similar to Figure 2B-C. **B.** Increase of probability to attract to the location of neighbors' previous flake consumption decreased as flakes were more abundant in the arena. Pearson's correlation coefficient, N=20 (since no significant difference in this relationship was found between groups of 3 and 6 fish, both group sizes are included in the same analysis); The shaded area shows the 95% confidence interval of the linear regression model in black. **C.** Fish showed a significant increase in tendency to swim towards neighbors when the latter changed their speed in a way that resembled flake consumption events ('pseudo consumptions')(P<0.005 for both group sizes, N=10,10 for groups of 3 and 6 fish; Wilcoxon's signed rank test ). **D.** The tendency to swim towards flake consumptions was significantly correlated over groups, N=20 groups (10 groups of 3 fish and 10 groups of 6 fish, no differences between group sizes); the shaded area shows the 95% confidence interval of the linear regression model in black.



451 452

**Figure 3-figure supplement 1: Models of fish foraging behavior. A.** Averages of the mean and standard deviation of the step size distributions estimated for all fish of the same group size (Figure 3B); error bars represent SEM. **B.** Same as in A but for the distributions of turns. **C.** Sketches of all models used in the study: In the IND model fish swims towards flakes that were closer than the flake detection range  $d_f < D_f$ . In all social models, if flakes were not detected within  $D_f$ , fish responded to the behavior of conspecifics that were within the neighbor detection range  $d_n < D_n$ . In the Att<sub>feed</sub> model, fish oriented towards neighbors' flake consumptions. In the Att model, fish oriented towards the average swimming direction of their neighbors. In the Att<sub>feed</sub>+Align model, fish responded first to neighbors' previous consumptions. If no consumption events were detected, fish responded as in the Align model. In the Att+Align model, fish responded as in the Align model to neighbors within  $D_n/2 < d_n < D_n$ . If there were other fish in both zones, the focal fish responded according to the average of the two response vectors. In the



 $Att_{feed}+Att+Align model$ , fish responded first to neighbors' previous consumptions. If no consumption events were detected, fish responded as in the Att+Align model.

Figure 4-figure supplement 1: Social models incorporating attraction to flake consumption by neighbors show the best fit to real foraging groups. A. Calculated error for the best fit models for groups of 3 fish, for three statistics of interest - nearest neighbor distance, group polarity and consumption times (See Figure 4C-D) and the combined error based on these three measures (right). Dots represent different experimental groups; horizontal lines are median values and boxes represent the 1<sup>st</sup> and 3<sup>rd</sup> quartiles. Dotted lines represent 0 error in prediction or a perfect fit to the data (Methods). B. Combined error values for all 7 models tested for groups of 3 and 6 fish (Figure 4D, Methods). The Att<sub>feed</sub>+Align model is significantly more accurate than all other models for both group sizes (P < 0.005, Wilcoxson's signed rank test, N=10 groups) except for the Att+Align model for groups of 6 fish (P=0.21). Dots represent different groups; horizontal lines are median values and boxes represent the 1<sup>st</sup> and 3<sup>rd</sup> quartiles. **C.** The inferred detection ranges for both the flake detection range  $(D_f)$  and the neighbor detection range  $(D_n)$  for the Att<sub>feed</sub>+Align model. Horizontal lines are median values and boxes represent the 1<sup>st</sup> and 3<sup>rd</sup> quartiles. **D.** Calculated error for all models for groups of 3 (top row) and 6 (bottom row) fish, for three statistics of interest - nearest neighbor distance, group polarity and consumption times (See Figure 4A-B). Dots represent different experimental groups; horizontal lines are median values and boxes represent the 1<sup>st</sup> and  $3^{rd}$  quartiles. Dotted lines represent 0 error in prediction or a perfect fit to the data (Methods).

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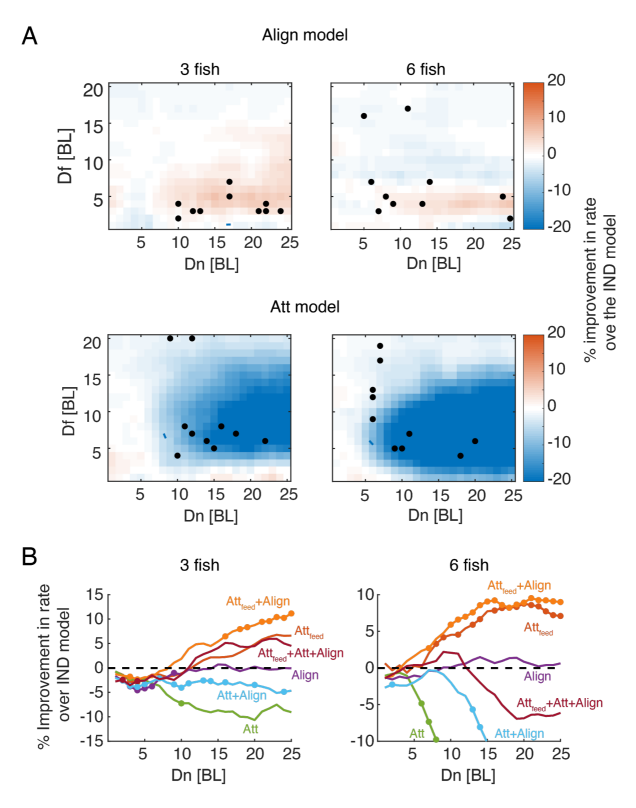
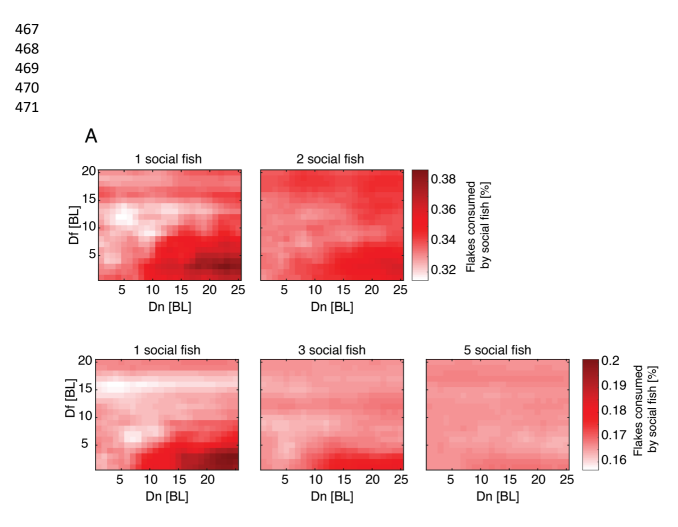




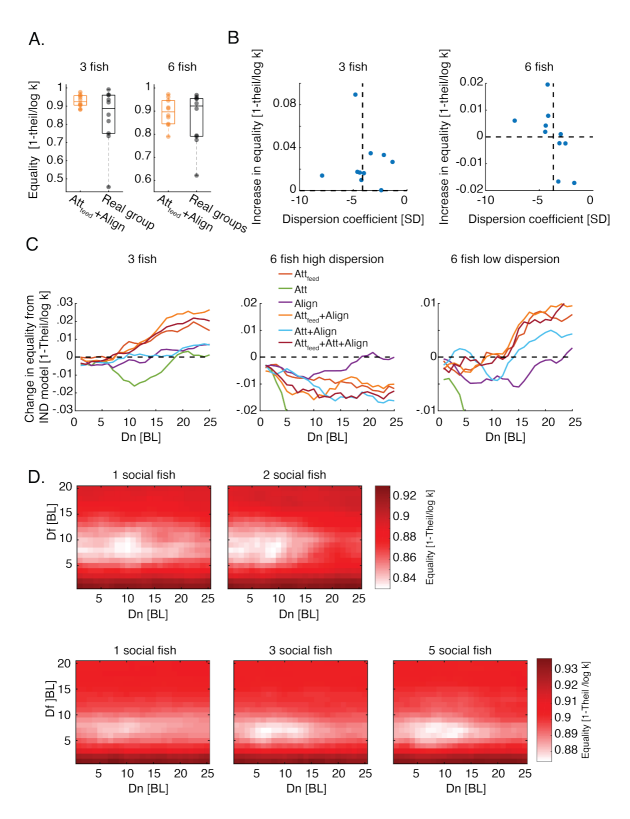
Figure 5-figure supplement 1: Foraging efficiency and attraction to neighbors' consumption events. A. Difference in foraging efficiency for groups that perform alignment (Align model, Top) or attraction (not related to neighbors' flake consumptions, Att model, Bottom) compared to groups of independent foragers (*IND*) for all model parameters. Dots represent  $D_f$  and  $D_n$  values of simulated groups that best fitted real foraging groups. B. Average improvement in the rate of flake consumption by socially interacting individuals compared to independent foragers. Colors indicate different social foraging strategies; dotted line represent no change compared to independent foragers (IND); results were averaged over all simulations with  $D_f \leq 5$  which was the parameter

range where real groups were matched by simulations; colored dots represent significant differences from the IND model (Wilcoxon's signed rank test, N=10 groups for each group size). Note that only models that incorporate specific attraction to neighbors' previous flakes consumptions (Att<sub>feed</sub> and Att<sub>feed</sub>+Align models) show increased foraging efficiency.



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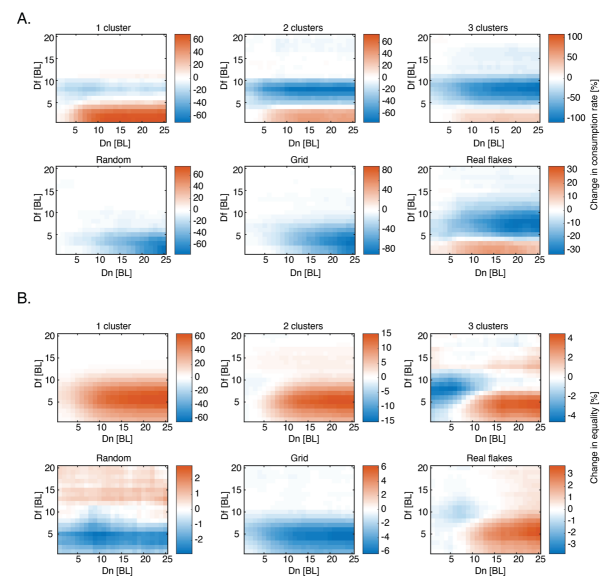
Figure 6-figure supplement 1: Social interactions improve foraging efficiency of individuals in groups. A. The mean fraction of flakes per fish that were consumed by foragers that used the Att<sub>feed</sub>+Align strategy of the total flakes consumed by all fish in the group. Mean values are shown for all simulated values of  $D_f$  and  $D_n$ . Different panels show different fractions of social foragers out of the group. Results are shown for groups of 3 fish (Top row) and 6 fish (Bottom row).



**Figure 6-figure supplement 2: Social interactions improve income equality in groups. A.** Equality of food consumption measured by  $1 - \frac{I_{Theil}}{logk}$  (see eq. 2) is shown for real foraging groups (black) and simulated groups using the Att<sub>feed</sub>+Align strategy (orange). Horizontal lines show median values, boxes represent the 1<sup>st</sup> and 3<sup>rd</sup> quartiles; no statistical differences were found between equality values in simulations and those of real groups of 3 fish (P=0.23, N=10) and 6 fish (P = 0.77, N = 10); Wilcoxon's signed rank test. B. Average Increase in equality for simulated groups of foragers using the Att<sub>feed</sub>+Align strategy, compared to independent foragers for the maximal social interaction

range tested ( $D_f \leq 5, D_n = 25$ ) plotted against the estimated dispersion of flakes in the arena (Methods). Horizontal dotted line represents 0 increase in equality and the vertical dotted line is the median dispersion value. Note that for groups of 6 fish, foraging for flakes with lower dispersion values (or high clustering) resulted in an increase in equality while foraging for flakes with higher dispersion showed a decrease in equality. **C.** Average difference in equality values between different social foraging strategies and independent foragers (IND model). Equality values were averaged over  $D_f \leq 5$  and over simulations corresponding to different groups (N = 10 groups for 3 fish and N=5,5 for low and high flake dispersion in groups of 6 fish). Note that the largest increase in equality was for models that incorporate specific attraction to neighbor's flake consumptions. **D.** Equality of food distribution among foragers for all  $D_f$ ,  $D_n$  values for all mixed strategy groups of 3 and 6 fish. Different panels show different fractions of individuals using the Att<sub>feed</sub>+Align strategy out of the total group; results are shown for groups of 3 fish (top row) and for groups of 6 fish (bottom row).

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**Figure 7-figure supplement 1: Simulating different distributions of food. A.** The change in rate of 6 socially interacting fish compared to independent ones (rate is calculated as number of flakes consumed / total search time) for all  $D_f$  and  $D_n$  values, for six different spatial distributions (Figure 7A); red colors represent increase in rate compared to independent foragers (100 repetitions for every parameter combination in every simulated distribution, Methods). **B.** Same as in A. only for the change in equality index  $1 - \frac{I_{Theil}}{\log k}$  (eq. 2).

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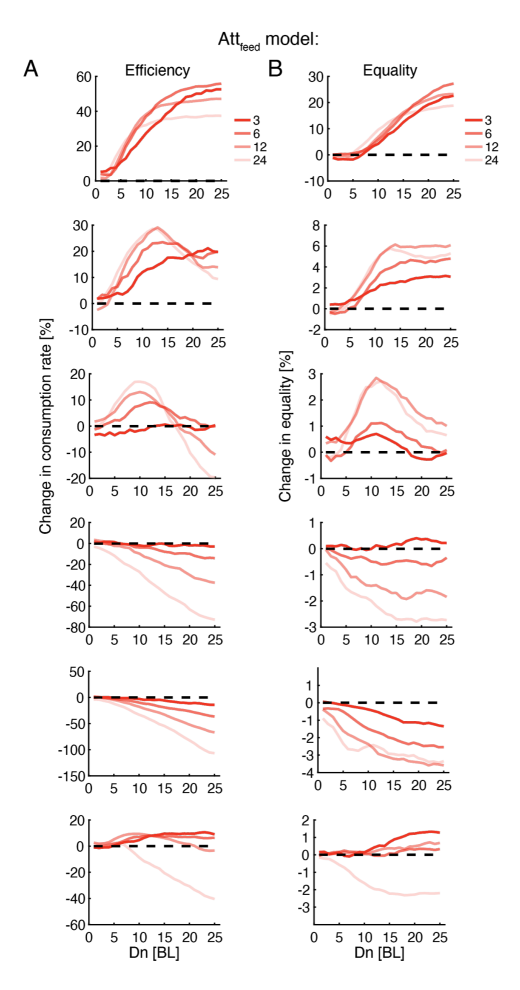


Figure 7-figure supplement 2: Optimal interaction range depends on group size and not on resource distribution. A. Improvement in group foraging rates (mean time to consumption per flake) of the Att<sub>feed</sub> model compared to independent foragers (IND model) for groups of 3, 6, 12, and 24 fish (dark to light colors) for the different spatial distributions (panels from top to bottom - Figure 7A). Results were averaged over all simulations with  $D_f \leq 5$  (Figure 7-figure supplement 1, 100 repetition per parameter combination, Methods). B. Same as in A but for the increase in equality of resources within groups. Note that the effects observed when only attraction to neighbors' previous consumptions is included in the model are very similar to those in the Att<sub>feed</sub>+Align model (see Figure 7), but also span the larger group sizes (12 and 24 fish). 

# 550 Videos

551

552 **Video 1: Foraging behavior of fish in a group.** Example group of 6 fish foraging together for 553 small flakes of food scattered on the water surface. Red circles represent flakes not eaten yet 554 by the fish, colors represent individual fish in the group.

555

556 **Video 2: Stereotypical maneuvers of fish during flake detection and consumption.** Zoom in 557 on a single fish (swimming in a group of 3 fish) showing two consecutive events of flake 558 detection and consumption. The fish speeds up towards the flake and turns sharply during 559 consumption.

560

Video 3-7: Groups of 6 Simulated fish foraging in various spatial distributions. Comparison
of simulated groups using social foraging (left: Att<sub>feed</sub>+Align) and independent foragers (right
- IND), for various spatial distributions of flakes: a single cluster (Video 3), two clusters (Video
4), three clusters (Video 5), Uniform distribution (Video 6) and a random distribution (Video
7). In all simulations, all fish start at the center of the arena, and simulations end when all
flakes are consumed (Methods).

567

## 568 Video 8: Groups of 12 Simulated fish foraging in a clustered environment.

569 Comparison of simulated 12 fish groups using social foraging (left: Att<sub>feed</sub>+Align) and 570 independent foragers (right - IND), foraging for flakes arranged in three clusters. all fish start 571 at the center of the arena, and simulations end when all flakes are consumed (Methods).

- 572
- 573

# 574 **Materials and Methods:**

575

576 **Experimental setup.** Detailed description of the setup is given in (48). Briefly, adult fish were 577 purchased from a local supplier (Aquazone LTD, Israel) and housed separately in their 578 designated groups for more than a month prior to behavioral experiments. Fish were housed 579 in a standard fish holding system consisting of a recirculating multistage filtration system 580 where temperature, conductivity, PH, and light- dark cycle were monitored.

581 Imaging of fish foraging behavior was done using an industrial recording system composed of 582 a *Vieworks VC-2MC-M340* camera with an 8 mm lens, a *Karbon-CL* frame grabber, and a 583 recording server. Camera was suspended 150 cm above the experimental tank. During 584 experiments we changed the effective experimental tank size by using different size arenas 585 (Figure 1-figure supplement 1A and see below). All water conditions were similar between 586 the holding tanks and the experimental tank.

587

588 Fish acclimation and behavioral experiments. To facilitate food searching behavior and to 589 lower fish anxiety, the following acclimation procedure was followed: On day 1, all fish were 590 transferred to the designated experimental tank (D = 95cm; water depth of 5 cm) and were 591 allowed to explore the tank for 5 minutes. On days 2-5, all groups and individual fish were 592 transferred from their home tanks to test tanks of increasing size (start box - 25x25cm, 593 D=47.5, cm D=67.2 cm, D=95 cm, Figure 1-figure supplement 1A) where 6, 12, or 18 flakes were 594 randomly scattered over the water in the area outside the start box (for groups of sizes 1, 3, 595 and 6 respectively, Figure 1-figure supplement 1A). The number of flakes used in the 596 experiments for individual fish and for groups were chosen based on preliminary 597 experiments, as the amount of food that would entice single fish to engage in the task yet not 598 to overcrowd the arena with flakes (especially for the larger groups). Fish were first placed in 599 a small starting box (25x25 cm) that was inside a larger arena. The small box was raised after 600 5 minutes and the fish were allowed to forage and consume the flakes in the larger arena for 601 an additional 5 minutes. After the allotted time ended the fish were netted and returned back 602 to their home tanks, keeping their original groups. Over the 4 days of training, we increased 603 the size of the test tank from the small start box itself (day 2) to the largest arena with D =604 47.5, 67.2, 95cm on days 3-5 (Figure 1-figure supplement 1A). On days 6-7 fish were deprived 605 of food and kept in their home tanks. Foraging was then tested on day 8. During training, no 606 food was administered to the fish outside of the experimental arena. In total, n = 106 adult 607 fish (3 months old or older) were used at approximately 1:1 male to female ratio. 16 single 608 individuals were tested, 10 groups of 3 fish (30 fish in total), and 10 groups of 6 fish (60 fish 609 in total). Two single individual fish were excluded from analysis as they did not swim when 610 transferred to the experimental tank.

611

612 **Data extraction.** Recorded videos were analyzed off-line to extract the size, position, speed, 613 and orientation of individual fish, and the position of food flakes. Position data was then used 614 to estimate fish trajectories using a designated tracker. All image processing and tracking 615 were done using MatLab<sup>©</sup> with software written in house; the details of these procedures 616 are described in (48). Fish identities were further corrected using IdTracker (43). Fish 617 trajectories were smoothed using a Savitzky-Golay filter (63) spanning 17 frames which 618 constituted ~1/3 of a second (all videos were recorded at 50 fps). Fish positions were defined as the coordinates of the center of each fish,  $\vec{x}_i(t)$ , and fish velocity was calculated as: 619  $\vec{v}_i(t) = \frac{\vec{x}_i(t+dt) - \vec{x}_i(t-dt)}{2dt}$ , with dt = 1 video frame or 20ms. Direction of motion of the fish was 620 defined as:  $\vec{d}_i(t) = \frac{\vec{v}_i(t)}{|\vec{v}_i(t)|}$  and the trajectory curvature at time t was given by c(t) = 1/R(t), 621 622 where R is the radius of the circle that gave the best Euclidean fit to a trajectory segment of 623 length 600 ms, centered on time t.

624

**Tracking flakes.** Flakes' locations were tracked with the same software that we have developed and used for tracking the fish (48). Flakes that were larger than 4 pixels (which correspond to a radius of about 1.15 mm) were reliably detected. Flakes typically disappeared when eaten, but when consumed flakes broke into smaller pieces, new (sub)flakes appeared. Consumption events were defined at times when a fish made contact with a flake and the flake disappeared from the camera's field of view. The resolution of our camera did not allow us to confirm whether the fish digested the flake entirely.

632

633 Flake consumption events and pseudo consumption events. We estimated the probability 634 of fish to cross near locations of consumption events by another fish, P(crossing), by 635 counting all events where at least one fish passed within 2 BL of that location within 1-4 s 636 after a neighbor's flake consumption, and dividing it by the total number of consumption 637 events. Since zebrafish tend to swim in groups, regardless of the presence of food, we 638 compared this number to the probability of one fish to cross near a neighbor's position within 639 1-4 s when no food was recently consumed by that neighbor (within the last 4 s) or will be consumed in the near future (within the next 4s). We therefore estimated  $P_{null}(crossing)$ 640 by drawing k random fish positions (mimicking k flake detection events) 10000 times, from 641

times when no flake was detected for at least 8 s (Figure 2D). The tendency to attract to flake position was then given by:  $P(crossing) - \langle P_{null}(crossing) \rangle$ , where angle brackets represent the average over random drawings for a given group.

645 We defined ``pseudo flake consumptions" at times when fish exhibited a similar speed profile 646 to that of a fish during real consumption events, namely gradual increase in speed followed 647 by a sudden sharp decrease back to baseline (Figure 2B). To detect such events, we convolved the speed profile of individual fish in the group at all times when no flakes were present near 648 649 the fish (for at least 8 s) with the calculated average speed profile near all real flake 650 consumption events of that group (Figure 2B) and obtained a correlation measure for each 651 point in time. We then treated the top 2.5 percent of this distribution as pseudo consumption 652 events. The average number of events was  $12.6 \pm 4$  and  $21.8 \pm 6.6$  for groups of 3 and 6 fish, 653 respectively. We compared the probability of neighbors to cross near the locations of such events,  $P(pseudo\ crossing)$  to  $P_{null}(crossing)$ . The tendency of neighboring fish to cross 654 655 near pseudo consumption events was high, and was correlated with their tendency to cross 656 near real flake consumption events over groups (Figure 2-figure supplement 1C-D).

657

**Segmentation of fish trajectories**. Segmentation of trajectories into discrete steps was based on the detection of local minima in the speed profile of the fish (Figure 3A-B). We characterized discrete steps by the total distance traveled between two minima points, *L*, and the change in angle between successive steps, or turning angle,  $\theta$ . These distributions were estimated for each fish and used to simulate their swimming behavior (Figure 3-figure supplement 1A-B).

664

# 665 Simulating swimming behavior.

666 **a.** Fish motion. We modeled fish swimming behavior as a correlated random walk, defined 667 by the distribution of step sizes, L, and of the turning angles between consecutive steps  $\theta$  (see 668 above). Thus, at each time point, we drew for each simulated fish a step size and a turning 669 angle, which determined the direction of motion and the length of the next step (Figure 3B, 670 Figure 3-figure supplement 1A-B). Average step size estimated from fish data was  $2.85 \pm 1.7$ 671 and  $3.1 \pm 1.9$  body lengths (BL) for groups of 3 and 6 fish, and turning angle distributions were nearly symmetric with an SD of 50° and of 46° for groups of 3 and 6 fish (Figure 3-figure 672 673 supplement 1A-B). The initial conditions of each simulation, i.e. starting positions of individual 674 fish, matched those of the real fish groups. All simulations were conducted within bounded 675 arenas, identical to those used for testing real fish. If a simulation step was expected to end 676 outside of the arena boundaries, that movement was discarded and a new movement was 677 chosen that did not exceed the arena boundaries. To compare simulated foraging time 678 (counted as discrete steps) to real foraging experiments (Figure 4B) we divided the length of 679 each simulated step by the calculated average speed of real foraging fish.

680

**b. Simulated Flakes**. In all simulations we used the true positions of flakes as extracted from fish foraging experiments. If a flake drifted during an experimental epoch (due to water turbulence), we only used its final position in the simulations since the speed and distance were negligible compared to the motion of the fish. Since real flakes sometimes disintegrated into smaller bits after a consumption event, we copied that in the simulations. I.e. if flake *i* at position  $\vec{x}_i$  has appeared after flake *j* was (partially) consumed, so did the corresponding flakes in the simulation.

689 **c. Sensory range of flake detection.** Each simulated fish had a circular range  $D_f$ , within which, 690 it could detect a flake, and orient towards it with probability  $p(go to flake) = e^{-d_f/D_f}$ , where 691  $d_f$  is the distance to the flake (Figure 3C). If the fish oriented itself towards the flake, its next 692 step size was drawn from the empirical distribution. If the fish reached the flake (or passed 693 it) during this movement, that flake was considered as consumed. If the simulated fish did not 694 reach the flake, the procedure was repeated. In the IND foraging model, k such fish were 695 simultaneously simulated, independent of one another.

696

697 **d. Sensory range of neighbor detection**. Since foraging fish in real groups were found to be 698 attracted to areas of previous flake consumptions, and since zebrafish are known to exhibit 699 schooling and shoaling tendencies, we allowed agents in our simulated social models to 700 detect and respond to neighbors' swimming and foraging behavior within the sensory range 701 of neighbor detection,  $D_n$ . Specifically, agents in the social foraging models could combine 702 various types of social interactions (Figure 3D, Figure 3-figure supplement 1C, and see below):

- 7031. Attraction to neighbors' previous flake consumptions ('Att<sub>feed</sub>'): if a neighbor of fish i704was within the sensory range of neighbor detection,  $D_n$ , and found a flake in the705previous  $\tau$  time steps, then fish i oriented towards the position of the previous706consumption with probability p(go to consumption) =  $e^{-d_n/D_n}$ , where  $d_n$  is the707distance to the position where a flake was consumed by a neighbor. In case a708movement towards a consumption position was successfully drawn, the step size was709drawn from the empirical distribution.
- 710 2. Neighbor attraction ('Att'): if neighbor(s) were found within the sensory range of 711 neighbor detection,  $D_n$  in the previous times step, fish oriented towards the center of 712 mass of these neighbors with probability P(attract) =  $e^{-\langle d_{n_j} \rangle/D_n}$  where  $\langle d_{n_j} \rangle$  is the 713 distance to the center of mass of the neighbors, such that the new direction of fish i 714 is  $\vec{d}_i(t+1) = \frac{\langle x_j(t) \rangle - x_i(t)}{|\langle x_j(t) \rangle - x_i(t)|}$ , where  $\langle x_j(t) \rangle$  is the center of mass of neighbors within  $D_n$
- 715 at time t.
- 3. Neighbor alignment ('Align'): if neighbor(s) were found within the sensory range of neighbor detection,  $D_n$  in the previous times step, fish adopted the average direction of these neighbors with probability P(align) =  $e^{-\langle d_n \rangle/D_n}$  where  $\langle d_n \rangle$  is the distance to the center of mass of the neighbors, such that the new direction of fish i is  $\vec{d}_i(t+1) = \frac{1}{I}\sum_{j \in d_{ij} < D_n} \vec{d}_j(t)$  where J is the number of neighbors within  $D_n$ .
- 721 722

723 e. Hierarchical nature of the social models. In the simulations, fish actions were given by the 724 following hierarchy: If a flake was within the  $D_f$  range of a fish, that fish would turn towards it with p(go to flake). If a flake was not detected (i.e. no flake was within  $D_f$ ), and a neighbor 725 consumed a flake at a distance smaller than  $D_n$ , then the fish would move towards that 726 location with the appropriate probability (for the cases where the model included response 727 728 to food consumption by neighbors). If neither a flake nor a neighbor consumption event was 729 detected, but neighbors were within a distance shorter than  $D_n$ , the fish responded to the 730 position/orientation of these neighbors (given that the model includes response to neighbors' 731 swimming). If no flakes or neighbors were detected, or if motion towards these areas was not 732 successfully drawn, then the next direction of motion of the fish was randomly chosen from 733 the empirical turning angle distribution.

### 734

**f. Mixed strategy groups.** In simulations of groups with mixed individual strategies, fish that
 used social interactions followed the procedure described above and the rest used
 independent foraging (Figure 6A).

738

**g. Simulating larger groups of foragers for various flake distributions.** We simulated group
 foraging for various group sizes: 3, 6, 12 and 24 fish in a range of flake distributions with a
 constant number of 18 flakes (Figure 7A):

- 742
- Single cluster. Flakes were randomly distributed within a small circle with a diameter
   of 9.5 cm (within the 95cm diameter arena). Circle center was situated at a distance
   of 20 cm from the arena walls.
- 746
   2. *Two clusters.* Flakes were equally distributed between two circles as described for the single cluster. Circle centers were situated on the diameter of the large arena to ensure maximal distance between clusters.
- 749
   3. *Three clusters.* Flakes were equally distributed between three clusters as described above. Circle centers were positioned on the vertices of an equilateral triangle to ensure maximal distance between clusters.
- 7524. Random. To achieve a random, non-clustered distribution we positioned flakes753randomly in the arena and assessed their clustering level based on the nearest754neighbor distance of flakes, using the clustering coefficient given in (64):  $C = \frac{NN_1}{0.5 \cdot \sqrt{\rho'}}$
- 755 Where  $NN_1$  is the average nearest neighbor distance of all flakes, and  $\rho$  is the density 756 of flakes. Values of |0 - C| < 0.01 were taken to show no spatial clustering and were 757 chosen for simulations.
- 5. Uniform (Hexagonal grid). We positioned N+1 flakes in a hexagonal grid within the
  arena boundaries. We then removed the center flake to allow fish to start in the center
  position.
- 761 6. *Empirical flake distributions.* We used the empirical flake distribution used in
   762 experiments (see above).
- 763

In all simulations, fish started foraging in the center of the arena (see Video 3-8). We repeated these simulations 100 times for all  $D_f$  and  $D_n$  values, for all group sizes in all simulated flake distributions (1-5). For the empirical flake distribution (see 6 above) we repeated simulations 20 times for all  $D_f$  and  $D_n$  values, for all group sizes and for every empirical flake distribution used in the experiments. We later averaged simulation results over the different flake distributions for each group size to achieve a single representative performance map similar to the results of the simulated flake distributions (Figure 7-figure supplement 1).

771

# 772 h. model classes and parameters.

	Independent models	Social models
Df – flake detection range	1-20 body length, increments of	1-20 body length
	1	increments of 1
Dn – neighbor detection range	0 body length	1-25 body length increments of
		1
$\tau$ – memory of a neighbor		5 time steps (where applicable)
detecting flakes		
Arena Radii	32 BL (95 cm)	32 BL (95 cm)

Turning angles	Estimated from swimming data of each fish	
Step sizes	Estimated from swimming data of each fish	
Flake positions	According to the starting position and appearance and disappearance dynamics in experiments or according to the simulated flake distributions (see G).	
Agents starting positions	According to the starting positions in experiments or in the center of the arena for simulated flake distributions.	
Model types and names	Description (see Figure 3-figure supplement 1C)	
Independent model (IND model)	fish only respond to flakes within $D_f$ around them ( $D_n$ =0)	
Attraction to consumption events (Att <sub>feed</sub> )	fish respond to flakes within $D_f$ , otherwise respond to neighbors' previous flake detections within $D_n$	
Attraction to neighbors regardless of consumption (Att model)	fish respond to flakes within $D_f$ , otherwise attract to the center of mass (average position of the group) of their neighbors within $D_n$ , regardless of consumption events	
Alignment with neighbors (Align model)	fish respond to flakes within $D_f$ , otherwise align with their average direction of neighbors within $D_n$	
Attraction to consumption events and alignment (Att <sub>feed</sub> +Align model)	fish respond to flakes within $D_f$ , otherwise respond to neighbors' previous flake detections within $D_n$ , otherwise align with neighbors within $D_n$	
Attraction and alignment with neighbors (Att+Align)	fish respond to flakes within $D_f$ , otherwise align with neighbors within $D_n/2$ , and attract to the center of mass of neighbors within $D_n/2 < d_n < D_n$ regardless of consumption events. If neighbors exist in both zones, fish average the attraction and alignment responses.	
Attraction to consumption events and attraction and alignment with neighbors (Att <sub>feed</sub> +Att+Align model)	fish respond to flakes within $D_f$ , otherwise respond to neighbors' previous flake detections within $D_n$ , otherwise align with neighbors within $D_n/2$ , and attract to the center of mass of neighbors within $D_n/2 < d_n < D_n$ regardless of consumption events. If neighbors exist in both zones, fish average the attraction and alignment responses.	

773

774

775 **Model Fitting.** Models were fitted to data by finding the parameter values ( $D_f$  for the 776 independent model, and  $D_f$ ,  $D_n$  for the social models) that maximized the log-likelihood of 777 the simulated consumption times given the empirical consumption times of a given group 778 (see below) and minimized the normalized squared error between the swimming statistics 779 (average polarity and nearest neighbor distance) of real groups and simulated groups.

780

781 *Consumption times:* we use the distribution of consumption times of flake i over the 782 simulations (100 repetitions), to assess the probability of observing a sequence of 783 consumption events on real data traces. Thus, the probability of the i-th consumption event 784 to occur at time *T* is given by its value from the simulations of the model,  $P_{D_f,D_n}(T_{data}(i))$ , 785 and the probability of a sequence of consumption times is given by the product of the 786 probabilities of the individual events  $\prod_i P_{D_f,D_n}(T_{data}(i))$ . The log-likelihood of the model is 787 then given by

789 
$$\log P_{D_{f},D_{n}}(T_{data}(1),T_{data}(2),...) = \sum_{i} \log P_{D_{f},D_{n}}(T_{data}(i))$$

790

791 where  $T_{data}(i)$  is the actual consumption time of the  $i^{th}$  flake, and the probability *P* is the 792 distribution of consumption times of the  $i^{th}$  flake over all simulations for a specific set of 793 model parameters  $D_f$ ,  $D_n$ . We used kernel density smoothing to estimate a continuous 794 probability from the discrete distribution obtained from simulations.

795

Polarity and nearest neighbor distance: we calculated the normalized mean squared error between the statistics obtained from simulations and the ones observed in the data:  $E(D_f, D_n, S) = \frac{((\hat{S}) - (S))^2}{SD(\hat{S})}$  where S is the relevant statistic measured (polarity or nearest neighbor distance),  $\hat{S}$  is the value obtain from simulations and  $\langle \dots \rangle$  is the average operation. polarity is defined *Polarity* =  $\frac{1}{N} \sum_i \vec{d}_i$ , which is the average direction vectors of the N fish in the group.

802

803 *Combined accuracy measure.* To assess the accuracy of the model we combined these three 804 separate measures into a single error function to be minimized by searching over the  $D_f$ ,  $D_n$ 805 values of the simulations:  $E^{combined} = E^{consumptions} + E^{polarity} + E^{D_{nn}}$ 

where E<sup>consumptions</sup> was taken as minus the log likelihood of consumptions (see above), and
 all three error surfaces were standardized by subtracting their mean value and dividing by the
 standard deviation, such that all errors will have similar units.

809

810 811 Measuring flake clustering. To quantify the spatial clustering of flakes, we simulated the 812 random positions of N flake consumptions (corresponding to the number of actual flake consumptions of each of the real groups). We then calculated the average nearest neighbor 813 distance of the simulated flakes  $D_{nn1}^{rand}$  and repeated this analysis 100,000 times to obtain a 814 distribution of average nearest neighbor distances expected at random. We then compared 815 816 the actual average nearest neighbor distance  $D_{nn1}$  of flake consumptions to the random distribution and assessed how likely it is to obtain  $D_{nn1}$  if flakes were randomly distributed in 817 space. The level of clustering of a group of n flakes is then defined by:  $\widehat{D}(n) =$ 818

819  $\frac{nn1(n) - \mu_{D_{nn1}}}{\sigma_{D_{nn1}}}$ , where  $\mu_{D_{nn1}}$  and  $\sigma_{D_{nn1}}$  are the mean and standard deviation of the

820 distribution of nearest neighbor distances of randomly distributed flakes.

821

822 Inequality measure. To quantify the inequality of flake consumptions between members of 823 the same group, we calculated the *Theil* index of inequality (52). This is an information theory-824 based measure, assessing the difference in the entropy of a distribution from the maximum 825 entropy expected if consumption rate was equal for all agents. This is given by

826 
$$I_{Theil}(k,n) = \frac{1}{k} \sum_{i=1}^{k} \frac{n_i}{\mu} \log\left(\frac{n_i}{\mu}\right)$$

where k is the number of agents,  $n_i$  is the number flakes consumed by fish i and  $\mu$  is the mean number of flakes consumed by a fish in the group. In addition, we normalized  $I_{Theil}$  by 829  $\log k$ , the maximum possible value if one fish consumed all flakes. We then quantified 830 equality by  $1 - \frac{I_{Theil}}{\log k}$ , where 1 indicates full equality and 0 is full inequality.

831 832

Sample sizes and power estimation. As the current research tests novel effects of social 833 behavior on group foraging, precise estimation of sample sizes and statistical power could not 834 835 be conducted a-priori. Instead, we have based our choice of sample sizes on previously 836 published studies of collective behavior and social foraging behavior of zebrafish (42, 48). In 837 addition, we chose to include more than one group size in the study design (groups of 3 and 838 6 fish) to support the generality of our findings. Finally, as the study is focused on groups, 839 sample sizes were chosen to be large enough to conduct parametric and nonparametric 840 statistical testing (e.g. Pearson's correlation coefficient, Wilcoxon's rank-sum test), while 841 minimizing the total number of animals used in the experiments.

- 842
- 843
- 844

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846

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855

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