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Cite this article: Herbert-Read JE, Kremer L, Bruintjes R, Radford AN, Ioannou CC. 2017 Anthropogenic noise pollution from pile-driving disrupts the structure and dynamics of fish shoals. *Proc. R. Soc. B* **284**: 20171627.

http://dx.doi.org/10.1098/rspb.2017.1627

Received: 20 July 2017 Accepted: 29 August 2017

Subject Category:

Global Change and Conservation

Subject Areas:

behaviour, environmental science, ecology

Keywords:

noise, collective behaviour, global change, shoaling, pile-driving

Authors for correspondence:

James E. Herbert-Read

e-mail: james.herbert.read@gmail.com

Christos C. Ioannou

e-mail: c.c.ioannou@bristol.ac.uk

Electronic supplementary material is available online at https://dx.doi.org/10.6084/m9. figshare.c.3878416.

THE ROYAL SOCIETY

Anthropogenic noise pollution from pile-driving disrupts the structure and dynamics of fish shoals

James E. Herbert-Read¹, Louise Kremer², Rick Bruintjes³, Andrew N. Radford⁴ and Christos C. Ioannou⁴

(i) JEH-R, 0000-0003-0243-4518; ANR, 0000-0001-5470-3463; CCI, 0000-0002-9739-889X

Noise produced from a variety of human activities can affect the physiology and behaviour of individual animals, but whether noise disrupts the social behaviour of animals is largely unknown. Animal groups such as flocks of birds or shoals of fish use simple interaction rules to coordinate their movements with near neighbours. In turn, this coordination allows individuals to gain the benefits of group living such as reduced predation risk and social information exchange. Noise could change how individuals interact in groups if noise is perceived as a threat, or if it masked, distracted or stressed individuals, and this could have impacts on the benefits of grouping. Here, we recorded trajectories of individual juvenile seabass (Dicentrarchus labrax) in groups under controlled laboratory conditions. Groups were exposed to playbacks of either ambient background sound recorded in their natural habitat, or playbacks of pile-driving, commonly used in marine construction. The pile-driving playback affected the structure and dynamics of the fish shoals significantly more than the ambient-sound playback. Compared to the ambient-sound playback, groups experiencing the pile-driving playback became less cohesive, less directionally ordered, and were less correlated in speed and directional changes. In effect, the additional-noise treatment disrupted the abilities of individuals to coordinate their movements with one another. Our work highlights the potential for noise pollution from piledriving to disrupt the collective dynamics of fish shoals, which could have implications for the functional benefits of a group's collective behaviour.

1. Introduction

Human activities, such as urbanization, resource extraction, transportation, and energy production, generate considerable noise. Since the Industrial Revolution, these human-generated noise sources have resulted in major changes in sound-scapes across the globe, both due to an increase in sound levels and the addition of sounds that are different from those arising from natural sources [1–3]. Consequently, anthropogenic noise is now recognized as a pollutant of international concern, being included in legislation such as the US National Environment Policy Act and the European Commission Marine Strategy Framework Directive. To inform policymakers, to develop effective management strategies, and to design suitable mitigation methods, detailed information on the organismal impacts of anthropogenic noise are needed.

There is mounting experimental evidence that anthropogenic noise can have a variety of negative physiological and behavioural effects on individual animals, ultimately affecting their survival and reproductive success [4–7]. For example, noise from human activities can directly cause injury or hearing

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¹Department of Zoology, Stockholm University, 10691, Stockholm, Sweden

²Department of Agronomy, Agroequipments, Farming and Environment, AgroSup Dijon, Dijon, France

³Biosciences, College of Life and Environmental Sciences, University of Exeter, Exeter, UK

⁴School of Biological Sciences, University of Bristol, Bristol, UK

loss in some species [8,9], or as a consequence of masking (i.e. reducing the signal-to-noise ratio), can impair the ability of animals to communicate [10-12]. Further, noise can be perceived as a threat, be a distraction, or can cause increased stress, in turn impairing an animal's ability to forage efficiently [13,14], respond appropriately to information about predation risk [15-17], perform adaptive behaviours during habitat-selection [18,19], or reproduce successfully [20,21]. However, despite the abundance of group-living species, there has been relatively little research on how the social behaviour of animals is affected by anthropogenic noise (but see [22-26]). An understanding of noise effects in this regard is crucial because many animals rely on such behaviours for their survival and reproductive success [27,28], and often adjust their social behaviour in response to risk [29,30]. Thus, any potential impacts of noise on social behaviour could have fundamental ecological and evolutionary implications for social species.

The impacts of anthropogenic noise can be particularly prevalent in aquatic environments, where sound travels further and faster before attenuation than in air [31]. Fish, in particular, are known to be affected by noise in a variety of ways [6,32]. For example, acoustic communication between individuals may be disrupted in the presence of noise, fish may move away from noisy sound sources, and in extreme circumstances, noise can even result in injury or death [6,32]. There has also been recent experimental evidence that anthropogenic noise can negatively affect the foraging [14,33], anti-predator [16,34], and parental care [21,23] behaviour of individual fish. It is estimated that approximately 50% of fish species form shoals during their lifetimes [35], with juvenile fish regularly shoaling in inshore areas [36] which are often subject to noisy exploration and construction projects [37]. Shoaling is achieved when individuals use simple interaction rules, including speed and direction changes, to coordinate their movements with near neighbours [38-40]. Information about neighbours' movements and positions is acquired through the lateral-line and visual systems [41,42], and there are good reasons to suspect that noise generated by human activity might affect shoaling dynamics.

Noise could impact the ability of individuals to coordinate their movements by masking information about neighbours' positions that could have been detected through the lateral line (uni-modal effects). Alternatively, or in addition to masking effects, distraction or stress could impair the coordination of individuals' movements by compromising an individual's ability to process information in another sensory channel (i.e. vision or olfaction), otherwise known as 'cross-modal' effects [43]. There is some evidence that noise produced by motorboats can affect the shape and structure of bluefin tuna (Thunnus thynnus) schools [22], and this could impair two of the key benefits of shoaling. First, shoaling provides anti-predatory benefits through dilution, confusion, and selfish-herd effects, with individuals in larger, more cohesive groups having proportionally less risk than individuals in smaller, less cohesive groups [44-47]. Changes in the cohesion of groups, therefore, could act to increase predation risk. Second, shoaling provides individuals access to social information, whereby information about detected threats or resources can be gathered by copying the movement decisions of others [48–51]. Disruption to the abilities of individuals to copy these decisions could have considerable implications for how

individuals in groups detect resources or avoid predators. High-resolution data on the positions and movements of individuals in shoals are needed, therefore, to measure how individuals are interacting in groups, and hence how anthropogenic noise may affect these interactions.

Here, we use a laboratory-based experiment to ask how anthropogenic noise (specifically playback of pile-driving noise, an impulsive sound source) impacts the shape, organization, and dynamics of European seabass (Dicentrarchus labrax) shoals. Seabass are known to be affected by playbacks of anthropogenic noise [52,53], making them a model species to use in these experiments. Laboratory-based experiments cannot perfectly replicate real-world sound fields or natural behaviour [54,55], but they allow tight control of other variables [55], as well as the collection of detailed (high spatial and temporal resolution) tracking data on shoaling behaviour, which has only recently been recorded in the wild [56]. We predicted that if the additional noise was perceived as a threat [57], the seabass would form denser, more directionally ordered shoals, with increased coordination of speed and direction changes, in the pile-driving treatment compared to an ambient-sound control treatment. If, however, the additional noise masks important information, or causes stress or distraction [4,58], the seabass would be predicted to form less cohesive and directionally ordered shoals, with reduced directional and speed coordination, in the pile-driving treatment compared to times with ambient-sound playback.

2. Material and methods

(a) Experimental subjects

Juvenile sea bass were sourced from Ifremer (Plouzane, France) and transported to the University of Exeter, where they were held for two months before being transported to the University of Bristol aquarium facilities. The fish were held in $40 \times 70 \times$ $34\,\text{cm}$ and $20 \times 70 \times 34\,\text{cm}$ (width \times length \times height) $5\,\text{mm}$ glass stock tanks that contained artificial plants. Fish were generally fed daily on a uniform commercial fish food diet (Perla MP Pellet, Skretting, Norway) except during a 7-week period in February-March 2015 when half of them were only fed three times per week. In this study, fish were randomly allocated to the sound treatments regardless of this feeding regime difference. Water temperature was 15.7 \pm 0.2°C; lighting was kept on a 12 L:12 D cycle; salinity was maintained between 35 and 36 parts per thousand (ppt). Experiments were conducted in July 2015 when the fish measured 9.7 \pm 0.7 cm (mean \pm s.d.) standard body length. The size of the fish did not differ between treatments (see below; Linear Model (LM): $F_{1,118} = 0.10$, p = 0.75).

(b) Recordings and playbacks

Original field recordings of offshore pile-driving in Swansea Bay, UK, were made between 87 and 200 m from the sound source [52,53]. Pile-driving at this site involved a 1.2 m diameter monopole being driven around 25 m into the seabed at a water depth of 6.5 m. The recordings of this process were made with a Hi Tech Inc. HTI-99HF hydrophone with inbuilt preamplifier (manufacturer calibrated sensitivity $-204 \, dB \, re \, 1 \, V \, \mu Pa^{-1}$, $20-125 \, 000$ Hz frequency range) and a data logger (RTsys EASDA, 44.1 kHz sampling rate). Recordings of ambient coastal sound were made at Portsmouth, Plymouth, and Gravesend, UK, using a Hi Tech Inc. HTI 96-MIN hydrophone with inbuilt preamplifier (manufacturer calibrated sensitivity $-164\,dB$ re $1\,V\,\mu Pa^{-1}$, $20-30\,000$ Hz frequency range) and a digital recorder (Roland Edirol R09HR 24 bit, 44.1 kHz sampling rate) [52,53]. For ambient-sound recordings, the hydrophone was positioned at 1 m depth 20–40 m offshore. All recordings were made during low to moderate wind speeds.

The original recordings of pile-driving noise and ambient sound were used to create three tracks per sound treatment; a random part of the relevant recording was used in each case (as in [52,53]). Multiple playback tracks per sound treatment were used to reduce pseudo-replication issues. All ambient playback tracks were 5 min in duration, and the pile-driving playbacks were 10–30 s in duration, with each track looped for the 5 min playback period. All the pile-driving playbacks had a pile-driving rate (time between pile-driving events) of 1.7 s. All tracks were created using Audacity 1.3.13 (http://audacity.sourceforge.net/).

Tracks were played back via an underwater loudspeaker (Aqua30; frequency range 80-20000 Hz: www.dnh.no), an amplifier (Kemo Electronics GmbH; 18W; frequency response range approx. 40-20 000 Hz), and a laptop (Toshiba Portege R930-1CW), as in [34,52,53]. To measure the recordings of the playbacks and any ambient sound in the room, we placed a hydrophone (HTI 96-MIN) in the middle of the experimental tank, 5 cm above the tank bottom. Recordings of the sound during the trials were made using a digital sound recorder (Sony PMC-M10, 44.1 kHz sampling rate) connected to the hydrophone. Because of unresolved challenges in measuring particle motion in small tanks at the time of the experiment, acoustic conditions were assessed in the sound-pressure domain only. In this experiment, we do not establish absolute values for sensitivity, but rather explore the potential impact of the change in additional sound on the fish's behaviour (see also, for example, [34,52,53]).

(c) Acoustic analysis

All sound recordings were analysed in MATLAB (v. 2013a and 2017a) using pamGuide [59] and paPAM [60]. Spectrograms (see the electronic supplementary material, figure S1) were calculated for 1–2 000 Hz (the frequencies most likely to be of relevance to seabass [61]) using a Hann evaluation window, 50% overlap, 0.1 s window length over 20 s recordings. Cumulative sound exposure level (SELcum) was calculated for the whole 5 min exposure period, whereas sound-pressure level (SPL) was calculated over 20 s recordings (electronic supplementary material, table S1). Zero-to-peak level, 90% energy envelope, rise time, and single-strike sound-exposure level (SELss) were calculated using an average of five randomly selected pile strikes (electronic supplementary material, table S1).

(d) Experimental protocol

Trials took place in an octagonal arena located at one end (10 cm from the wall) of a $2.5 \times 1.25\,m$ aluminium tank lined with a white plastic PVC pond liner (electronic supplementary material, figure S2). The arena was made of white $68 \times 43.4 \, \text{cm}$ Perspex panels, so the narrowest width of the arena was 105 cm. The loudspeaker was located at the other end of the tank, 20 cm from the wall facing the arena and half way along the width of the tank. The loudspeaker was suspended using string to be 2 cm above the bottom of the tank. Water depth was 10 cm and temperature and salinity conditions matched those in the stock tanks. The whole tank was covered by a cuboid frame and white sheeting to minimize disturbance and diffuse overhead fluorescent lighting to minimize reflections on the water surface. A Panasonic X920 camcorder, positioned centrally and 193 cm above the water's surface, was used to film the arena at a resolution of 1920×1080 pixels and frame rate of 59 frames per second.

Our focus in this study was the effect of exposure to additional noise; comparisons were made to individuals that experienced control playbacks (of recordings of ambient coastal noise) but were otherwise from the same cohort and held under the same conditions. Four fish were netted from the same stock tank and transferred gently to the test arena. Juvenile seabass generally occur in small group sizes [62], and we chose groups of four fish as this is within the range of group sizes used in previous studies on these fish [63-65]. The groups of four fish were given 15 min to acclimatize, during which time no playback occurred, with the last 5 min of this period filmed (termed '1st half of trial' hereafter). One of the two sound treatments (pile-driving noise; n = 15 groups, or ambient sound; n =15 groups) was then played to the fish for 5 min (termed '2nd half of trial' hereafter), with filming continuing during this period. The trial order of sound treatments was determined by a complete random block; i.e. for each pair of trials (1st and 2nd, 3rd, and 4th, etc.), one of each treatment was given, but in a random order within that pair. Which of the three replicate recordings was used for each treatment was randomly determined. Each fish was tested only once. Fish were not fed on the day of testing until after the trials.

(e) Response measures

The 10 min videos from each trial were converted to MPEG-4 with Handbrake 0.10.5 (https://handbrake.fr/). idTracker [66] was used to track the x and y coordinates of each fish throughout each trial. All subsequent analyses were performed using MATLAB (2016a) and followed similar methods to [67–69]. The parameters associated with the spatial and directional organization, as well as the movement dynamics of the fish, are detailed below. Measures were calculated separately for each fish in the group. All variables were calculated for the 1st (no playback) and 2nd (playback) half of each trial separately.

(i) Spatial and directional organization of the shoals

We first calculated measures associated with the cohesiveness of shoals including the mean distance each individual was to the shoal's centroid, and the modal nearest-neighbour distance of each individual. The modal nearest-neighbour distance represents the distances that pairs of individuals are most commonly observed apart [39,68]. We then calculated the distance from each fish to its nearest-neighbour perpendicular to their direction of travel (i.e. how far apart side-by-side) and parallel to the direction of travel (i.e. how far apart in front-orbehind one another). We further calculated the bearing angle to a fish's nearest neighbour, which represents the direction that a neighbour was most likely to be found in relation to the focal individual [38]. We treated bearing angles to the neighbour ahead or behind of the focal fish separately for ease of interpretation in the statistical models. A bearing angle of 90° would represent a neighbour that was directly to the side of a focal individual, whereas a bearing angle of 0° or 180° would represent a neighbour that was, respectively, directly in front or behind a focal individual. We also calculated the heading difference between nearest neighbours, i.e. the angle between the direction nearest neighbours were facing. This measures how closely aligned nearest neighbours in the shoals were, effectively measuring their directional organization [70]. The heading difference ranges from 0° (individuals were facing in the same direction = high alignment) to 180° (individuals were facing in opposite directions = low alignment). Full details of these calculations can be found in the electronic supplementary materials.

(ii) Movement dynamics of individuals in the shoals

The above measures determine the spatial organization of fish shoals, but do not capture how individuals are moving and

interacting within them. To assess how individuals were moving and interacting in the shoals, we first calculated the speed and direction of each fish at each time point. From these values, we then calculated the cross-correlations between an individual's speed (or direction) and its nearest-neighbour's speed (or direction). Cross-correlations assess how strongly pairs of individuals copy each other's speed or direction changes in time, and we followed established methods described by Nagy et al. [71]. In brief, we identified the peak of the correlation in speed (or direction) and the time delay between when two individuals' speeds (or directions) were most strongly correlated. Higher peaks of the cross-correlation indicate the two fish were more strongly correlated in speed (or direction), and shorter absolute time delays indicate that the two individuals direction were more synchronized, with individuals adopting the speed or direction changes of their partner sooner. Because individuals' speeds (but not direction) were highly correlated in time (i.e. showed minimal time delays to the peak correlation) we did not analyse the time-delays between nearest-neighbours' speeds, but did analyse these delays for direction changes. Full details of these calculations can be found in the electronic supplementary materials.

(f) Statistical analysis

All measures calculated from the fish trajectories were analysed as response variables in Mixed Models. All models included the treatment (pile-driving or ambient-sound playback) as a between-subjects term, and the half of the trial (1st or 2nd) as a within-subjects term. The interaction between these two fixed terms was included in the initial models, but was removed where it was non-significant and models were re-run with main effects only. All models included fish identity nested within trial (which is equivalent to the group the individuals belonged to) as the random term. The average difference between fish perpendicular or parallel to their direction of travel, the bearing of the nearest neighbour in-front or behind, and the difference in heading between nearest neighbours were analysed for each fish using negative binomial Generalized Linear Mixed Models (GLMMs), as the data were typically right skewed. For all GLMMs, the dispersion parameter was checked to be approximately equal to 1 (more than 0.5 and less than 2) using Generalized Linear Models with the same model structure but without the random effects. The variance function for the negative binomial models is $\mu \times (1 + \mu/k)$, for k > 0 (i.e. variance is approx. equal to the mean for $\mu \ll k$ and proportional to the mean squared for $\mu\gg k$), where μ is the mean and k is the shape parameter for a negative binomial distribution.

The remaining response variables were analysed using Linear Mixed Models (LMMs). The median speed and the mean time delay that maximized the directional correlation with the nearest neighbour of each fish were analysed without transforming these response variables. The mean distance to the centroid and modal nearest-neighbour distance for each fish were log₁₀ transformed before analysis. The mean maximum speed correlation and mean maximum directional correlation with the nearest neighbour for each fish were transformed by subtracting the correlation from one and then applying a log₁₀ transformation (i.e. $log_{10}(1-correlation coefficient)$). For all LMMs, the residuals from each model were checked to ensure normality (using QQ plots) and homoscedasticity (using the residuals plotted against the fitted values). The variance function here is $Var(\mu) = 1$. Owing to correlations between speed and maximum speed and directional correlations, and between speed and the time delay that maximized the directional correlation, the models that analysed these response variables were repeated with the fish's speed as an additional main effect to control for the correlation of speed with the response variables.

3. Results

(a) Spatial and directional organization of the shoals

The spatial structure of the shoals changed in both the ambient-sound and pile-driving playbacks. There was an interaction between treatment and the half of the trial when investigating group cohesiveness (LMM: $F_{1,118}=4.44$, p=0.04; electronic supplementary material, table S2a). When the ambient-sound playback was initiated, the mean distance of individuals to the group's centroid decreased, whereas this distance increased when the pile-driving playback treatment was initiated. Similarly, the modal nearest-neighbour distance decreased in the ambient-sound playbacks, whereas this distance increased in the pile-driving playbacks (interaction between treatment and trial half: $F_{1,118}=14.88$, p<0.001; figure 1a; electronic supplementary material, table S2b).

To investigate in more detail how the distances between fish in the pile-driving playback increased, we assessed the relative positions that individuals adopted next to their nearest neighbour. Individuals tended to position themselves in a lattice formation, with nearest neighbours most frequently being found at either 43° or 133° in front or behind the focal fish, respectively, and to the left or right, rather than directly in front or behind one another (figure 1b). During both playback treatments, the bearing angle to the nearest neighbour moved closer to 90°, indicating that fish were more likely to be observed side-by-side compared to before the playbacks were initiated (GLMM, effect of trial half on angle to neighbour in-front: $\chi^2 = 11.22$, d.f. = 1, p < 0.001; electronic supplementary material, table S2c; and angles to neighbour behind: $\chi^2 = 12.78$, d.f. = 1, p < 0.001; electronic supplementary material, table S2d). The distance between fish in this direction (i.e. the distance fish were apart perpendicular to their direction of travel) increased in both the ambient-sound and pile-driving playbacks, but this effect was larger in the pile-driving playback than the ambientsound playback (interaction between treatment and trial half: $\chi^2 = 7.72$, d.f. = 1, p < 0.01; electronic supplementary material, table S2e).

The angular difference in heading between nearest neighbours also increased in both treatments, but this effect was larger in the pile-driving playback than in the ambient-sound playback (GLMM, interaction between treatment and trial half: $\chi^2 = 7.99$, d.f. = 1, p < 0.01; figure 1c; electronic supplementary material, table S2g).

(b) Movement dynamics of individuals in the shoals

The speed of fish decreased when both the ambient-sound and pile-driving playbacks were initiated, but this effect was larger in the pile-driving playback (LMM, interaction between treatment and half of trial: $F_{1,118} = 32.53$, p < 0.001; figure 2a,b; electronic supplementary material, table S3a). The maximum correlation between nearest-neighbours' speeds also decreased in both playback types, but again this effect was larger in the pile-driving playback (interaction between treatment and half of trial: $F_{1,118} = 46.43$, p < 0.001; figure 2c; electronic supplementary material, table S3b). Repeating the statistical model with speed included as a covariate did not change this finding ($F_{1,132} = 20.61$, p < 0.001; electronic supplementary material, table S3c).

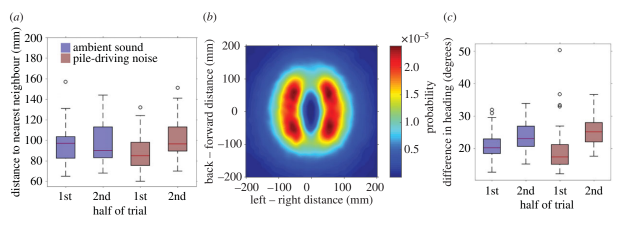


Figure 1. (a) Modal nearest-neighbour distances of fish during the 1st (no playback) and 2nd (playback) half of the trials. (b) The relative positions of a fish's nearest neighbour, combined for all fish and across both sound treatments. The focal fish is located at the origin of the plot and is facing along the positive y-axis. The heat in the plot shows the probability of finding the focal fish's nearest neighbour in locations surrounding the focal fish. 'Hotter' regions indicate a larger probability of finding a neighbour in that location. For plotting purposes, the heatmap has been smoothed with a Gaussian filter, $\sigma = 6$. (c) Mean angular difference in heading between a fish and their nearest neighbour in the 1st and 2nd half of the trial. In (a,c), blue bars represent the ambient-sound treatment and red bars represent the pile-driving treatment. Edges of each of the boxes represent the 25 and 75% percentiles, whiskers extend to all included data, black circles represent outliers.

The maximum correlation between nearest-neighbours' directions in time decreased when the playbacks were initiated, but this effect was stronger in the pile-driving playback (LMM, interaction between treatment and trial half: $F_{1.118} = 11.14$, p < 0.01; figure 2d; electronic supplementary material, table S3d). This result held when including speed as a covariate in the model ($F_{1,132} = 4.23$, p = 0.04; electronic supplementary material, table S3e), indicating that larger reductions in speed in the pile-driving treatment could not solely explain this result. The time delay between nearestneighbours' maximum directional correlations also increased more in the pile-driving playback than in the ambient-sound playback (interaction between treatment and trial half: $F_{1.118} = 14.57$, p < 0.001; electronic supplementary material, table S3f). However, this result could be explained based on the larger reductions in the speed of fish during the pile-driving playback (interaction between treatment and trial half when speed included as a covariate: $F_{1,132} = 2.65$, p = 0.11; electronic supplementary material, table S3g). In other words, larger reductions in speed in the piledriving playback also caused larger delay times between nearest-neighbours' maximum directional correlations.

4. Discussion

Both pile-driving and ambient-sound playbacks affected the spatial and directional organization, as well as the coordination of seabass shoals, but these effects were often more pronounced when there was additional anthropogenic noise. In particular, the distance between fish increased more, and the directional and speed organization of the shoals decreased more during the pile-driving playbacks compared with the ambient-sound playbacks. Noise from the pile-driving treatment therefore caused significant changes to how individuals coordinated their movements with near neighbours, ultimately affecting the structure of the shoals. While most studies have concentrated on individual behavioural responses to anthropogenic noise, this study provides conclusive evidence that the social interactions of individuals within groups are also impacted by added noise.

Our experiment using both ambient-sound and piledriving playbacks highlights that both sound types impacted the structure and dynamics of fish schools. The simple addition of sound beyond current baseline levels, therefore, impacts the shoaling behaviour of fish regardless of its source (i.e. ambient sound or pile-driving noise). Indeed, across sensory modalities, sensory systems are highly responsive to sudden changes in background conditions [72], as this reflects information about changes in the environment. Changes in behaviour in both treatments, such as reductions in speed, may therefore reflect increased alertness due to changes in environmental conditions. This highlights the importance of relevant controls that should be used during these types of playback experiments. Changes in the fish's behaviour between the 1st half of the trial (no playback) to the 2nd half of the sound (playback), however, were typically much larger in the pile-driving compared to the ambientsound playback. When the ambient-sound and pile-driving playbacks changed the behaviour of the fish in the same direction, all effect sizes of these changes (comparing changes in the behaviour of the fish between the 1st and 2nd half of the trials) except one ranged between 0.75 and 1.18 (see electronic supplementary material, table S4), indicating the differences between the playbacks were medium to strong effects [73]. Neo et al. [65] found that fish exposed to impulsive sound took longer to recover (return to swimming closer to the water's surface) compared to continuous noise [26]. This suggests that the temporal structure of a sound source, as well as the frequencies and amplitude of it, could have an important influence on behavioural responses to that source. Further work is needed on how the temporal, frequencies, and amplitude of anthropogenic noise sources affect behaviour.

Pile-driving playback decreased the cohesiveness of seabass groups, which is the opposite to what would be expected if the fish treated the additional noise as a predation threat [57]. Under predation threat, groups are expected to become more cohesive as individuals reduce risk through dilution and confusion effects [74–78]. Instead, fish in our experiment increased their distance to the group's centroid and between nearest neighbours. We also found that the

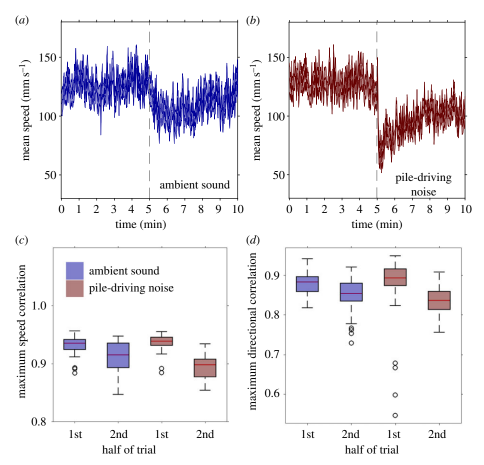


Figure 2. Mean speed (\pm s.e.) of fish in (a) ambient sound or (b) pile-driving noise treatment during the 1st half of the trial (no playback) and the 2nd half of the trial (playback). The dashed lines at 5 min separates the 'no playback' and 'playback' regions. (c) Mean maximum cross-correlation in speed between nearest neighbours in the 1st and 2nd half of the trials. (d) Maximum directional correlation between a fish and its nearest neighbour in the 1st and 2nd half of the trial. In each plot, blue bars represent the ambient-sound treatment, whereas red bars represent the pile-driving treatment. Edges of the boxes represent the 25 and 75% percentiles, whiskers extend to all included data, black circles represent outliers.

speed and directional coordination of nearest neighbours decreased more in the pile-driving playback compared to the ambient-sound playback. Again, this is the opposite to what would be expected under increased predation threat, where individuals should be highly sensitive and coordinate their movements more strongly with near neighbours [48,49]. Instead, our findings are consistent with the idea that pile-driving playback disrupts the cohesion and coordination of individuals in the shoals.

Coordination of the movements of individual fish is thought to be modulated primarily by lateral-line and visual sensory inputs [41,46]. If detection of nearestneighbours' movements through the lateral line were obstructed by the playbacks, this could explain reductions in directional and speed correlations between nearest neighbours. This would effectively be an example of masking, a uni-modal effect of additional noise, although that has mostly been considered to date with respect to vocal communication [79]. Alternatively, even if lateral-line information was not disrupted, additional noise may still have impacted the ability of individuals to process sensory information through cross-modal effects. Cross-modal effects occur when the processing and effective use of information is negatively affected by additional noise as a consequence of stress and/or distraction [15]. These cross-modal effects have recently been demonstrated in other species [15] and are well known in the cognitive sciences [80]. Therefore, cross-modal effects could also occur when attempting to coordinate movement with near neighbours, ultimately affecting the structure of these groups. It may be possible to assess whether uni-modal, cross-modal, or both effects influence the schooling behaviour of fish by knocking out the functionality of the lateral-line system using aminoglycoside antibiotics [42]. By then assessing whether the schooling behaviour of fish was impacted further by the addition of noise, this would provide evidence that noise impacts schooling behaviour even if sound could not be detected with the lateral-line system. More generally, cross-modal effects can be assessed by measuring whether behavioural responses to stimuli that have no auditory component (e.g. visual cues or olfactory cues) are impacted by sound, as has recently been investigated [15].

Disruption to how individuals interact in groups could impact some of the associated benefits of group living, including a reduced predation risk and access to social information [27,28]. Individuals in less cohesive groups are attacked more frequently than individuals in more cohesive groups [78], and we observed that individuals increased their distances between one another more in response to the pile-driving compared to the ambient-sound playback. Individual fish in shoals also gain information from others, for example, about a detected threat, by copying the speed and directional changes of near neighbours [49,81]. Similarly, the collective ability of groups to sense complex gradients in their environment is modulated by how individuals copy the speed changes of others [50]. How individuals respond to

each other's movements determines the likelihood and extent of information propagation in animal groups [82], and disruptions by anthropogenic noise could have considerable ecological implications for group-living species. Non-lethal effects such as these are crucial for our understanding of how noise pollution impacts the behaviour and survival of animals, and this will be important to consider for animals in natural conditions with real sound sources.

Our experiment focused on responses to a single relatively short-term noise exposure, as is the case with the majority of fish research to date (see [53,83] for exceptions). For a full understanding of the impacts of anthropogenic noise, longer-term studies are also needed, because animals may be able to compensate during quieter periods and responses may change with repeated or chronic exposure [53,84–86]. Ideally, those future studies should be conducted in natural conditions with real-world sound sources [16] to ensure maximum ecological and acoustic validity [55]. However, captive experiments such as ours do provide a valuable stepping stone in the study of environmental stressors, including noise [16,34,87], not least because of the ability to control tightly the conditions and to collect detailed individual-based data.

Our work highlights the potential for noise from anthropogenic sources to disrupt the coordination of shoaling fish. Whether this translates to functional consequences for fishes, such as changes in feeding success or predation risk, will now need to be assessed. Nevertheless, our results demonstrate that sound can influence the spatial and directional organizational characteristics of fish shoals, and as such, should be considered when environmental impact assessments of construction projects in marine or freshwater environments are conducted.

Ethics. All procedures regarding use of animals in research followed United Kingdom guidelines and were approved by the University of Bristol Ethical Review Group (UIN UB/10/034).

Data accessibility. All data accompanying this paper can be downloaded from: https://figshare.com/s/237fda89f8ffc265a6f6.

Author's contributions. C.I., L.K., R.B., and A.R. designed the study. L.K. and C.I. collected the data. J.E.H.-R., R.B., and C.I. analysed the data. J.E.H.-R., R.B., A.R., and C.I. wrote the manuscript. All authors gave approval for publication.

Competing interests. We have no competing interests.

Funding. J.E.H.-R. was supported by a Knut and Alice Wallenberg Foundation Grant awarded to D.J.T. Sumpter. C.C.I. was supported by an NERC Independent Research Fellowship NE/K009370/1.

Acknowledgements. We thank Mérin Broudic for the offshore pile-driving recordings, and Irene Voellmy and Sophie Nedelec for the ambient noise recordings. We thank two anonymous reviewers for useful comments.

References

- 1. McDonald MA, Hildebrand JA, Wiggins SM. 2006 Increases in deep ocean ambient noise in the Northeast Pacific West of San Nicolas Island, California. J. Acoustic. Soc. Amer. 120, 711-718. (doi:10.1121/1.2216565)
- Watts RD, Compton RW, McCammon JH, Rich CL, Wright SM, Owens T, Ouren DS. 2007 Roadless space of the conterminous United States. Science 316, 736 – 738. (doi:10.1126/science.1138141)
- Normandeau Associates I. 2012 Effects of noise on fish, fisheries, and invertebrates in the U.S. Atlantic and Arctic from energy industry soundgenerating activities. A literature synthesis for the U.S. Dept. of the Interior, Bureau of Ocean Energy Management. Contract Number: M11PC00031, p. 153.
- Kight CR, Swaddle JP. 2011 How and why environmental noise impacts animals: an integrative, mechanistic review. Ecol. Lett. 14, 1052-1061. (doi:10.1111/j.1461-0248.2011.01664.x)
- Shannon G et al. 2015 A synthesis of two decades of research documenting the effects of noise on wildlife. Biol. Rev. 91, 982-1005. (doi:10.1111/brv.
- Kunc HP, McLaughlin KE, Schmidt R. 2016 Aquatic noise pollution: implications for individuals, populations, and ecosystems. Proc. R. Soc. B 283, 20160839. (doi:10.1098/rspb.2016.0839)
- Morley EL, Jones G, Radford AN. 2014 The importance of invertebrates when considering the impacts of anthropogenic noise. Proc. R. Soc. B 281, 20132683. (doi:10.1098/rspb.2013.2683)
- Halvorsen MB, Casper BM, Matthews F, Carlson TJ, Popper AN. 2012 Effects of exposure to pile-driving

- sounds on the lake sturgeon, Nile tilapia and hogchoker. Proc. R. Soc. B 279, 4705-4714. (doi:10.1098/rspb.2012.1544)
- Popper AN, Smith ME, Cott PA, Hanna BW, MacGillivray AO, Austin ME, Mann DA. 2005 Effects of exposure to seismic airgun use on hearing of three fish species. J. Acoust. Soc. Am. 117, 3958-3971. (doi:10.1121/1.1904386)
- 10. Slabbekoorn H, Ripmeester EA. 2008 Birdsong and anthropogenic noise: implications and applications for conservation. Mol. Ecol. 17, 72-83. (doi:10. 1111/j.1365-294X.2007.03487.x)
- 11. Kunc HP, Lyons GN, Sigwart JD, McLaughlin KE, Houghton JD. 2014 Anthropogenic noise affects behavior across sensory modalities. Am. Nat. 184, 93 – 100. (doi:10.1086/677545)
- 12. Lampe U, Schmoll T, Franzke A, Reinhold K. 2012 Staying tuned: grasshoppers from noisy roadside habitats produce courtship signals with elevated frequency components. Funct. Ecol. 26, 1348 – 1354. (doi:10.1111/1365-2435.12000)
- 13. Wale MA, Simpson SD, Radford AN. 2013 Noise negatively affects foraging and antipredator behaviour in shore crabs. Anim. Behav. **86**, 111 – 118. (doi:10.1016/j.anbehav.2013. 05.001)
- 14. Voellmy IK, Purser J, Flynn D, Kennedy P, Simpson SD, Radford AN. 2014 Acoustic noise reduces foraging success in two sympatric fish species via different mechanisms. Anim. Behav. 89, 191-198. (doi:10.1016/j.anbehav.2013.12.029)
- 15. Morris-Drake A, Kern JM, Radford AN. 2016 Crossmodal impacts of anthropogenic noise on

- information use. Curr. Biol. 26, R911-R912. (doi:10.1016/j.cub.2016.08.064)
- 16. Simpson SD, Radford AN, Nedelec SL, Ferrari MCO, Chivers DP, McCormick MI, Meekan MG. 2016 Anthropogenic noise increases fish mortality by predation. Nat. Commun. 7, 10544. (doi:10.1038/ ncomms10544)
- Purser J, Bruintjes R, Simpson SD, Radford AN. 2016 Condition-dependent physiological and behavioural responses to anthropogenic noise. Phys. Behav. 155, 157 – 161. (doi:10.1016/j.physbeh.2015.12.010)
- 18. Holles S, Simpson SD, Radford AN, Berten L, Lecchini D. 2013 Boat noise disrupts orientation behaviour in a coral reef fish. Mar. Ecol. Prog. Ser. **485**, 295 – 300. (doi:10.3354/meps10346)
- 19. Simpson SD, Radford AN, Holles S, Ferarri MCO, Chivers DP, Mccormick MI, Meekan M. 2016 Smallboat noise impacts natural settlement behavior of coral reef fish larvae. In The effects of noise on aquatic life II (eds A Popper, A Hawkins), vol. 875. Advances in Experimental Medicine and Biology, pp. 1041 – 1048. Berlin, Germany: Springer.
- Blickley JL, Blackwood D, Patricelli GL. 2012 Experimental evidence for the effects of chronic anthropogenic noise on abundance of greater sagegrouse at leks. *Conserv. Biol.* **26**, 461 – 471. (doi:10. 1111/j.1523-1739.2012.01840.x)
- 21. Nedelec S. 2017 Motorboat noise impacts parental behaviour and offspring survival in a reef fish. Proc. R. Soc. B 284, 20170143. (doi:10.1098/rspb. 2017.0143)
- 22. Sarà G et al. 2007 Effect of boat noise on the behaviour of bluefin tuna Thunnus thynnus in the

- mediterranean sea. Mar. Ecol. Prog. Ser. 331, 243-253. (doi:10.3354/meps331243)
- 23. Bruintjes R, Radford AN. 2013 Context-dependent impacts of anthropogenic noise on individual and social behaviour in a cooperatively breeding fish. Anim. Behav. 85, 1343-1349. (doi:10.1016/j. anbehav.2013.03.025)
- 24. Owens JL, Stec CL, O'Hatnick A, 2012 The effects of extended exposure to traffic noise on parid social and risk-taking behavior. Behav. Process. 91, 61-69. (doi:10.1016/j.beproc.2012.05.010)
- 25. Fewtrell J, McCauley R. 2012 Impact of air gun noise on the behaviour of marine fish and squid. Mar. Poll. Bull. 64, 984-993. (doi:10.1016/j. marpolbul.2012.02.009)
- 26. Neo YY, Parie L, Bakker F, Snelderwaard P, Tudorache C, Schaaf M, Slabbekoorn H. 2015 Behavioral changes in response to sound exposure and no spatial avoidance of noisy conditions in captive zebrafish. Front. Behav. Neurosci. 9, 28. (doi:10.3389/fnbeh.2015.00028)
- 27. Krause J, Ruxton GD. 2002 Living in groups. Oxford, UK: Oxford University Press.
- 28. Ward A, Webster M. 2016 Sociality: the behaviour of group-living animals. Switzerland: Springer International Publishing.
- 29. Ioannou CC, Ramnarine IW, Torney CJ. 2017 Highpredation habitats affect the social dynamics of collective exploration in a shoaling fish. Sci. Adv. 3, e1602682. (doi:10.1126/sciadv.1602682)
- 30. Schaerf TM, Dillingham PW, Ward AJ. 2017 The effects of external cues on individual and collective behavior of shoaling fish. Sci. Adv. 3, e1603201. (doi:10.1126/sciadv.1603201)
- 31. Williams R et al. 2015 Impacts of anthropogenic noise on marine life: publication patterns, new discoveries, and future directions in research and management. Ocean Coast. Manag. 115, 17-24. (doi:10.1016/j.ocecoaman.2015.05.021)
- 32. Radford AN, Kerridge E, Simpson SD. 2014 Acoustic communication in a noisy world: can fish compete with anthropogenic noise? Behav. Ecol. 25, 1022-1030. (doi:10.1093/beheco/aru029)
- 33. Purser J, Radford AN. 2011 Acoustic noise induces attention shifts and reduces foraging performance in three-spined sticklebacks (Gasterosteus aculeatus). PLoS ONE 6, e17478. (doi:10.1371/journal.pone. 0017478)
- 34. Simpson SD, Purser J, Radford AN. 2015 Anthropogenic noise compromises antipredator behaviour in European eels. Glob. Change Biol. 21, 586 – 593. (doi:10.1111/gcb.12685)
- 35. Shaw E. 1978 Schooling fishes: the school, a truly egalitarian form of organization in which all members of the group are alike in influence, offers substantial benefits to its participants. Am. Sci. 66, 166 - 175.
- Macpherson E. 1998 Ontogenetic shifts in habitat use and aggregation in juvenile sparid fishes. J. Exp. Mar. Biol. Ecol. 220, 127-150. (doi:10.1016/S0022-0981(97)00086-5)
- 37. Maragos JE. 1993 Impact of coastal construction on coral reefs in the US-affiliated pacific islands. Coast.

- Manage. 21, 235-269. (doi:10.1080/089207593 09362207)
- 38. Herbert-Read JE. 2016 Understanding how animal groups achieve coordinated movement. J. Exp. Biol. **219**, 2971 – 2983. (doi:10.1242/jeb.129411)
- 39. Katz Y, Tunstrøm K, Ioannou CC, Huepe C, Couzin ID. 2011 Inferring the structure and dynamics of interactions in schooling fish. Proc. Natl Acad. Sci. **108**, 18 720 – 18 725. (doi:10.1073/pnas.1107583108)
- 40. Mann RP, Herbert-Read JE, Ma Q, Jordan LA, Sumpter DJT, Ward AJW. 2014 A model comparison reveals dynamic social information drives the movements of humbug damselfish (Dascyllus aruanus). J. R. Soc. Interface 11, 20130794. (doi:10. 1098/rsif.2013.0794)
- 41. Partridge BL, Pitcher TJ. 1980 The sensory basis of fish schools: relative roles of lateral line and vision. *J. Comp. Physiol.* **135**, 315 – 325. (doi:10.1007/ BF00657647)
- 42. Faucher K, Parmentier E, Becco C, Vandewalle N, Vandewalle P. 2010 Fish lateral system is required for accurate control of shoaling behaviour. Anim. Behav. 79, 679-687. (doi:10.1016/j.anbehav.2009. 12.020)
- 43. Halfwerk W, Slabbekoorn H. 2015 Pollution going multimodal: the complex impact of the humanaltered sensory environment on animal perception and performance. Biol. Lett. 11, 20141051. (doi:10. 1098/rsbl.2014.1051)
- 44. Hamilton WD. 1971 Geometry for the selfish herd. J. Theor. Biol. 31, 295-311. (doi:10.1016/0022-5193(71)90189-5)
- 45. Seghers BH. 1974 Schooling behavior in the guppy (Poecilia reticulata): an evolutionary response to predation. Evolution 28, 486-489.
- Ioannou CC, Couzin ID, James R, Croft DP, Krause J. 2011 Social organisation and information transfer in schooling fish. In Fish cognition and behavior (eds C Brown, K Laland, J Krause), pp. 217-239. New York, NY: Wiley-Blackwell. (doi:10.1002/97814 44342536.ch10)
- 47. Ioannou CC. 2017 Swarm intelligence in fish? The difficulty in demonstrating distributed and selforganised collective intelligence in (some) animal groups. Behav. Process. 141, 141-151. (doi:10. 1016/j.beproc.2016.10.005)
- Radakov DV. 1973 Schooling in the ecology of fish. New York, NY: John Wiley.
- Herbert-Read JE, Buhl J, Hu F, Ward AJ, Sumpter DJ. 2015 Initiation and spread of escape waves within animal groups. R. Soc. open sci. 2, 140355. (doi:10. 1098/rsos.140355)
- 50. Berdahl A, Torney CJ, Ioannou CC, Faria JJ, Couzin ID. 2013 Emergent sensing of complex environments by mobile animal groups. Science **339**, 574 – 576. (doi:10.1126/science.1225883)
- 51. McDonald ND, Rands SA, Hill F, Elder C, Ioannou CC. 2016 Consensus and experience trump leadership, suppressing individual personality during social foraging. Sci. Adv. 2, e1600892. (doi:10.1126/sciadv. 1600892)
- Bruintjes R, Purser J, Everley KA, Mangan S, Simpson SD, Radford AN. 2016 Rapid recovery

- following short-term acoustic disturbance in two fish species. R. Soc. open sci. 3, 150686. (doi:10. 1098/rsos.150686)
- 53. Radford AN, Lèbre L, Lecaillon G, Nedelec SL, Simpson SD. 2016 Repeated exposure reduces the response to impulsive noise in European seabass. Glob. Change Biol. 22, 3349-3360. (doi:10.1111/ acb.13352)
- 54. Rogers PH, Hawkins AD, Popper AN, Fay RR, Gray MD. 2016 Parvulescu revisited: small tank acoustics for bioacousticians. In The effects of noise on aquatic life II (eds A Popper, A Hawkins). Advances in Experimental Medicine and Biology, vol 875, pp. 933-941. New York, NY: Springer.
- 55. Slabbekoorn H. 2016 Aiming for progress in understanding underwater noise impact on fish: complementary need for indoor and outdoor studies. In The effects of noise on aquatic life II (eds A Popper, A Hawkins). Advances in Experimental Medicine and Biology, vol 875, pp. 1057 – 1065. New York, NY: Springer.
- 56. Ward AJW, Schaerf TM, Herbert-Read JE, Morrell L, Sumpter DJT, Webster MM. 2017 Local interactions and global properties of wild, free-ranging stickleback shoals. R. Soc. open. sci. 4, 170043. (doi:10.1098/rsos.170043)
- 57. Frid A, Dill L. 2002 Human-caused disturbance stimuli as a form of predation risk. Conserv. Ecol. 6, 11. (doi:10.5751/ES-00404-060111)
- 58. Chan AAYH, Blumstein DT. 2011 Attention, noise, and implications for wildlife conservation and management. Appl. Anim. Behav. Sci. 131, 1-7. (doi:10.1016/j.applanim.2011.01.007)
- 59. Merchant ND, Fristrup KM, Johnson MP, Tyack PL, Witt MJ, Blondel P, Parks SE, Hodgson D. 2015 Measuring acoustic habitats. Methods Ecol. Evol. 6, 257 - 265. (doi:10.1111/2041-210X.12330)
- 60. Nedelec SL, Campbell J, Radford AN, Simpson SD, Merchant ND. 2016 Particle motion: the missing link in underwater acoustic ecology. Methods Ecol. Evol. **7**, 836 – 842. (doi:10.1111/2041-210X.12544)
- 61. Lovell J. 2003 The hearing abilities of the bass, Dicentrarchus labrax. Technical report commissioned by ARIA Marine Ltd. for the European Commission Fifth Framework Programme. Project Reference: Q5AW-CT-2001-01896. Technical report.
- 62. Anras MB, Lagardère JP, Lafaye JY. 1997 Diel activity rhythm of seabass tracked in a natural environment: group effects on swimming patterns and amplitudes. Can. J. Fish. Aquat. Sci. 54, 162-168. (doi:10.1139/f96-253)
- 63. Herskin J, Steffensen J. 1998 Energy savings in sea bass swimming in a school: measurements of tail beat frequency and oxygen consumption at different swimming speeds. J. Fish Biol. 53, 366-376. (doi:10.1111/j.1095-8649.1998.tb00986.x)
- 64. Russell N, Fish J, Wootton R. 1996 Feeding and growth of juvenile sea bass: the effect of ration and temperature on growth rate and efficiency. J. Fish Biol. 49, 206-220. (doi:10.1111/j.1095-8649.1996. tb00017.x)
- Neo YY, Seitz J, Kastelein RA, Winter HV, ten Cate C, Slabbekoorn H. 2014 Temporal structure of sound

- affects behavioural recovery from noise impact in european seabass. Biol. Conserv. 178, 65-73. (doi:10.1016/j.biocon.2014.07.012)
- 66. Pérez-Escudero A, Vicente-Page J, Hinz RC, Arganda S, de Polavieja GG. 2014 idtracker: tracking individuals in a group by automatic identification of unmarked animals. Nat. Methods 11, 743-748. (doi:10.1038/nmeth.2994)
- 67. Herbert-Read JE, Krause S, Morrell LJ, Schaerf TM, Krause J, Ward AJW. 2013 The role of individuality in collective group movement. Proc. R. Soc. B 280, 20122564. (doi:10.1098/rspb.2012.2564)
- 68. Herbert-Read JE, Perna A, Mann RP, Schaerf TM, Sumpter DJT, Ward AJW. 2011 Inferring the rules of interaction of shoaling fish. Proc. Natl Acad. Sci. USA **108**, 18 726 – 18 731. (doi:10.1073/pnas. 1109355108)
- 69. Schaerf TM, Herbert-Read JE, Myerscough MR, Sumpter DJT, Ward AJW. 2016 Identifying differences in the rules of interaction between individuals in moving animal groups. (http://arxiv. org/abs/1601.08202)
- 70. Tunstrøm K, Katz Y, Ioannou CC, Huepe C, Lutz MJ, Couzin ID, Ben-Jacob E. 2013 Collective states, multistability and transitional behavior in schooling fish. PLoS Comput. Biol. 9, e1002915. (doi:10.1371/ journal.pcbi.1002915)
- 71. Nagy M, Ákos Z, Biro D, Vicsek T. 2010 Hierarchical group dynamics in pigeon flocks. Nature 464, 890-893. (doi:10.1038/nature08891)
- 72. Endler JA. 1992 Signals, signal conditions, and the direction of evolution. Am. Nat. 139, S125-S153. (doi:10.1086/285308)

- 73. Cohen J. 1977 Statistical power analysis for the behavioural sciences (Rev. ed.). New York, NY: Academic Press.
- 74. Foster W, Treherne J. 1981 Evidence for the dilution effect in the selfish herd from fish predation on a marine insect. Nature 293, 466-467. (doi:10.1038/ 293466a0)
- 75. Hogan BG, Hildenbrandt H, Scott-Samuel NE, Cuthill IC, Hemelrijk CK. 2017 The confusion effect when attacking simulated three-dimensional starling flocks. R. Soc. open sci. 4, 160564. (doi:10.1098/ rsos 160564)
- 76. Ioannou C, Tosh C, Neville L, Krause J. 2008 The confusion effect-from neural networks to reduced predation risk. Behav. Ecol. 19, 126-130. (doi:10. 1093/beheco/arm109)
- 77. Ioannou CC, Morrell LJ, Ruxton GD, Krause J. 2009 The effect of prey density on predators: conspicuousness and attack success are sensitive to spatial scale. Am. Nat. 173, 499-506. (doi:10.1086/
- 78. Ioannou C, Guttal V, Couzin I. 2012 Predatory fish select for coordinated collective motion in virtual prey. Science 337, 1212-1215. (doi:10.1126/ science.1218919)
- 79. Brumm H, Slabbekoorn H. 2005 Acoustic communication in noise. Adv. Stud. Behav. 35, 151 – 209. (doi:10.1016/S0065-3454(05)35004-2)
- Parmentier FB. 2008 Towards a cognitive model of distraction by auditory novelty: the role of involuntary attention capture and semantic processing. *Cognition* **109**, 345 – 362. (doi:10.1016/ j.cognition.2008.09.005)

- 81. Gautrais J, Ginelli F, Fournier R, Blanco S, Soria M, Chaté H, Theraulaz G, Levin SA. 2012 Deciphering interactions in moving animal groups. PLoS Comput. Biol. 8, e1002678. (doi:10.1371/journal.pcbi. 1002678)
- 82. Sumpter D, Buhl J, Biro D, Couzin I. 2008 Information transfer in moving animal groups. Theor. Biosci. 127, 177-186. (doi:10.1007/s12064-008-0040-1)
- 83. Nedelec SL, Mills SC, Lecchini D, Nedelec B, Simpson SD, Radford AN 2016 Repeated exposure to noise increases tolerance in a coral reef fish. Environ. *Pollut.* **216**, 428 – 436. (doi:10.1016/j.envpol.2016.
- 84. Bejder L, Samuels A, Whitehead H, Finn H, Allen S. 2009 Impact assessment research: use and misuse of habituation, sensition and tolerance in describing wildlife responses to anthropogenic stimuli. Mar. Ecol. Prog. Ser. **395**, 177 – 185. (doi:10.3354/ meps07979)
- 85. Wale MA, Simpson SD, Radford AN. 2013 Sizedependent physiological responses of shore crabs to single and repeated playback of ship noise. Biol. Lett. 9, 20121194. (doi:10.1098/rsbl.2012.1194)
- 86. Bruintjes R, Radford AN. 2014 Chronic playback of boat noise does not impact hatching success or post-hatching larval growth and survival in a cichlid fish. PeerJ 2, e594. (doi:10.7717/peerj.594)
- 87. Scott GR, Johnston IA. 2012 Temperature during embryonic development has persistent effects on thermal acclimation capacity in zebrafish. Proc. Natl Acad. Sci. USA 109, 14 247 – 14 252. (doi:10.1073/ pnas.1205012109)