1 2	Come dine with me: food-associated social signalling in wild bottlenose
3	dolphins (<i>Tursiops truncatus</i>)
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13	Abstract
14	The occurrence of food-related signaling is prolific in the animal kingdom with some food-
15	associated vocalizations considered functionally referential. Food calls can, however, vary
16	greatly in the type of information they convey. Thus, there are a multitude of purposes for which
17	food calls are used, including social recruitment, caller spacing, the indication of type, quantity,
18	quality, divisibility of food and/or the caller's hunger level and even as tools to manipulate prey
19	behaviour. Yet little work has focused on the social aspect of food calling in animals. We
20	investigated the association of social signals in wild bottlenose dolphins with foraging behaviour
21	where context specific food-associated calls are commonly produced. Our data showed that
22	specific social signals were significantly correlated with food call production and these calls
23	rarely occurred in the absence of food calls. We suggest that animals are sharing additional
24	information on the food patch itself with their social affiliates.

25 Keywords: food calls, bottlenose dolphin, signature whistles, vocal learning, call matching.

- 27 Introduction
- 28 29

Food calls are found throughout the animal kingdom when animals are trying to attract others to 30 31 a foraging site (Clay et al. 2012). In many cases, these vocalizations function in attracting related 32 animals to increase the caller's inclusive fitness. Animals may also call to attract non-related 33 partners, which may help enhance the caller's social standing (Slocombe et al. 2010). Alternatively, the presence of others may increase the caller's food intake by helping to herd 34 prey (Brown et al. 1991) or protecting against predators (Radford and Ridley 2007) or competing 35 36 conspecifics (Heinrich 1988). Animals may also communicate more detailed information such as the type, quality or quantity of food available (Clay et al. 2012). Thus much interest has been 37 placed on understanding how food-related signaling benefits the caller. Yet little focus has been 38 placed on the other types of social signals that may accompany the production of food-associated 39 calls. 40

In cetaceans, for example, food-associated vocalizations have been described for several species 41 in the wild, including killer whales (Orcinus orca, Ugarte et al. 2006; Deecke et al. 2011), 42 humpback whales (Megaptera novaeangliae, Stimpert et al. 2011; Parks et al. 2014), and 43 bottlenose dolphins (Janik 2000a). While these calls may be used to attract conspecifics, reports 44 45 suggest that their primary function may be one of manipulating prey to facilitate capture (Janik 46 2000a). While this is an interesting possibility, there is so far no experimental confirmation. One study looked at the effect of high, mid and low frequency dolphin clicks on fish behaviour but 47 found no effect (Benoit-Bird et al. 2006). Many of these food-related burst-pulsed calls, 48 49 however, are stimulus specific and therefore particular calls may have evolved to modify the behaviour of particular prey species. If food associated calls do play a role in facilitating prey 50 51 capture in wild animals then the attraction of conspecifics may be a by-product that does not

actually benefit the caller. However, animals may use other social calls concurrently with food-associated vocalizations in order to share information on the food patch.

Ridgway et al. (2014) reported calls in a food context in captive bottlenose dolphins and beluga whales that were structurally different from other dolphin food calls, such as the bray call (Janik 2000a). Ridgway et al. (2014) believed these calls signalled the emotional state of the animal where by an animal is anticipating a food reward. Thus, these calls should not be specific to a food context and would therefore not qualify as food calls (Clay et al. 2012). Interpreting calls in training contexts can be difficult and further work on wild animals is required to explore the distinction between such calls.

To investigate the possibility of added communication through other social calls when food calls are given, we investigated the association of whistle interactions with bray calls in bottlenose dolphins. Bray calls have a frequency spectrum that makes them suitable for manipulating prey, but also for attracting conspecifics to the foraging location (Janik 2000a). We investigated whether there was a correlation between food calling and a certain sign of social interaction, the matching of whistles between different animals (Janik 2000b).

67 Materials and Methods

During May to September 2010, we collected acoustic recordings from the dolphins that frequent
St. Andrews Bay, Scotland. These animals are members of a resident population of
approximately 195 bottlenose dolphins that range between the Moray Firth and the Firth of Forth
(Cheney et al. 2013). To identify foraging, we looked for the presence of food-related bray calls
in our acoustic recordings. Animals produce bray calls at depth (20m-30m) (Hastie et al. 2006)

and often return to the surface visibly chasing and foraging on large fish (Janik 2000a; Hastie etal. 2006). We then explored if whistle matching occurred in close proximity to bray calls.

Group follows were conducted from a 6-m boat at Beaufort sea state three or less and animals
were photographed with a Canon Digital 30D SLR camera with a Sigma 100-300mm, F4 zoom
lens. Acoustic recordings were taken using two HTI-96 MIN hydrophones (frequency response:
0.002 to 30 kHz ±1 dB) towed at 2-m depth and a Toshiba Satellite Pro Laptop with an Edirol
UA-25 sound card (sampling rate: 96 kHz, 16 bit).

80 We analysed acoustic recordings by screening the spectrographic displays (FFT length 1024, 87.5% overlap, Hanning window) in Adobe Audition v2.0 (Adobe Systems) by eye for 81 82 occurrences of low-frequency bray calls and frequency modulated whistles. Those recording 83 segments where engine noise exceeded 2 kHz were discarded from the analysis. Bray calls are 84 highly distinctive and were identified by detecting their two-part structure consisting of a long, 85 low frequency (peak at less than 2 kHz) pulsed sound followed by a short down-sweep (Janik 86 2000a). Each bray call is 500 to 600 milliseconds in length (dos Santos et al. 1995; Janik 2000a) 87 and can be recorded from over a km from the calling animal. Bray calls are usually produced in bouts (Janik 2000a) and therefore a series of brays in close proximity was treated as one braying 88 event (Figure 1). Bray sequences can last up to 30 s (dos Santos et al. 1995). Frequency 89 90 modulated whistles were identified and could either be continuous in their frequency contour pattern or could be multi-loop whistles. Multi-loop whistles were defined as a repeated 91 modulation pattern that could be separated by periods of stereotyped silence up to 250ms in 92 length (Esch et al. 2009). Whistles with periods of silence of less than 250ms but not overlapping 93 94 were treated as one whistle. A total of 23 hours and 51 minutes of recordings were inspected spanning 18 days from June to August 2010, of which 12 hours and 08 minutes from 15 days 95

96 were of sufficient sound quality for further analysis. A total of 81 braying events were identified97 in this recording time.

98 The simple two-element recording system used in this study meant we were unable to localise 99 whistles to individual animals. Instead we used overlapping whistle interactions as an indicator 100 of two animals communicating with each other. We focused on those overlaps in which both 101 animals were using the same whistle type, an indicator of animals addressing each other (Janik 102 2000b; King and Janik 2013; King et al. 2014).

103 An 'overlap match' was defined as two whistles of the same contour pattern (change of

frequency over time) where the end point of one whistle finished after the start of the other
whistle (Figure 2). All identified occurrences of overlap whistle matching were separately
checked and then agreed by both authors who have been shown to agree with classifications by
groups of independent observers (Janik 1999; Deecke and Janik 2006; Janik, King, Sayigh,
Wells 2013).

To be sure the overlapping of whistles was not merely a result of an increase in call rate, we also looked at the occurrence of different whistle types that overlapped but did not match. An 'overlap no match' was defined as two whistles with different contour patterns where the end point of one whistle finished after the start of the second whistle.

Permutation tests were performed to test whether overlap matching and bray calls occurred in close temporal proximity above chance levels. The times between each 'overlap match' and the nearest bray call (n=30) were shuffled with the times between each 'overlap no match' and the nearest bray call (n=55). The random distribution was calculated from 10000 permutations under the null hypothesis that there was no relationship between the timing of the 'overlap matches' and bray call production. The observed test statistic (mean proximity between 'overlap matches'
and bray calls) was then compared with the random distribution. The observed difference in
mean proximity to bray calls between overlap matches and overlap non-matches was also
compared to a random distribution of differences in mean proximities between the two whistle
categories, which was calculated from 10000 permutations.

123 All analyses were performed in R (R project for statistical computing; GNU project).

124 **Results**

125 A total of 30 occurrences of overlap matching and 55 occurrences of overlap non-matching

events were identified from recordings made during 10 encounters across 9 days (Table 1).

Individual re-sightings of animals across the 10 encounters were relatively low (mean: 1.85,
range: 1-5) and group sizes were large (mean: 9.5 animals, range: 3-23); see Table 2. A total of
46 animals were identified plus 4 calves. Of those, 69% were only sighted in one or two
encounters, 22% in three encounters, and 9% in four or five encounters. No single animal was
found in all 10 encounters.

132 Overlap matches occurred in significantly closer temporal proximity to bray calls than whistles that overlapped but did not match (P < 0.0066, permutation test, Table 1 and Figure 3). Under 133 the random distribution only 57% of whistles were expected to occur within 1 minute of a bray 134 call. However, 73% of observed overlap matches occurred within 1 minute of a bray call 135 136 whereas of those cases where whistles overlapped but did not match, only 49% occurred within 1 137 minute of a bray call. The randomization test showed that overlap matches and bray calls 138 occurred in tight temporal proximity with the mean proximity between overlap matches and bray 139 calls significantly smaller than expected by chance (permutation, P < 0.0001; Figure 4). In

140 contrast, the mean proximity between whistles that overlapped but did not match and bray calls 141 was significantly larger than expected by chance (permutation, P < 0.0001; Figure 4).

142 Thus the rapid matching of call-types was closely associated with bray call production and

appears to play a significant role in bottlenose dolphin foraging behaviour while the animals

144 decrease rapid, non-matching interactions during foraging.

145 The majority (60%) of whistles used in overlap matching events were variations of one particular

146 whistle pattern (type A; Figure 2 & Table 2), with a second whistle pattern (type D) produced in

- 147 20% of events (Figure 5 & Table 2).
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- 149

150 Discussion

We have shown that bottlenose dolphins repeatedly produce specific social signals, overlap matches, alongside food-associated calls and that they rarely produce these signals independently of the food call. In addition, they appear to decrease the production of overlapped non-matched calls when foraging. This suggests that there is a social aspect of food calling in animals that has not previously been reported.

156 A few studies have reported matching exchanges (Janik 2000b; King et al. 2013) and

157 overlapping in dolphin whistles (Caldwell and Caldwell 1968; Nakahara and Miyazaki 2011).

158 Bottlenose dolphins use vocal learning to develop individually distinctive signature whistles that

allow for individual recognition (Janik and Sayigh 2013). In specific contexts, these are also

used in matching interactions (Janik and Slater 1998; King et al. 2013). In captive bottlenose

161 dolphins, signature whistle matching i.e. copying the signature whistle of an animal right after it 162 produced one, leads to further signature whistling (King et al. 2014). This does not occur when using whistles of other types as a reply. Furthermore, the animals show no sign of aggression 163 164 when matched, suggesting that signature whistle matching is affiliative and helps to maintain contact (King et al. 2013; King et al. 2014). In the wild, the matching of call types in quick 165 166 succession in coordinated foraging, as shown here, could facilitate the directing of signals to particular individuals that share a large communication network (McGregor and Dabelsteen 167 1996; Janik 2005). 168

169 The production of food-related bray calls in dolphins has previously been correlated with 170 animals swimming quickly towards the caller's location, perhaps facilitating joining events 171 between individuals or groups of animals(Janik 2000a). Animals who wish to join may therefore use overlap matching to signal their intention when approaching a caller. However, although 172 173 dolphins are known to increase whistle rates dramatically when groups join together, whistle matching has not been observed during these joining events (Quick and Janik 2012). It is thus 174 unlikely that overlap matching facilitates group joins, but it may help coordination between 175 animals within a group when rapid reactions are needed. 176

177

An example for this type of rapid signaling can be found in the pied-babbler (*Turdoides bicolor*),
a species that uses a close call, called the 'chuck' to regulate spacing between foraging
competitors (Radford and Ridley 2008). Although not used aggressively, the close call did deter
conspecifics that attempted to share an individual's foraging patch, and was effective in
indicating the forager's current position. Pied-babblers increased chuck production rate when in
larger groups, and when neighbours were closer (Radford and Ridley 2008).

185	Overlapping can occur in a chorus when animals compete over resources such as females
186	(Staicer et al. 1996) or aim at an acoustic dilution effect for predator avoidance (Gerhardt and
187	Huber 2002). We suggest a more interactive function here where overlapping of shared whistles
188	regulates the spacing of individuals during intensive feeding bouts where group sizes are large.
189	Such directed overlapping can be an aggressive signal in some species (e.g. manakins; Maynard
190	et al. 2012), and a sign of social bonding (e.g. chimpanzees; Fedurek et al. 2013) or social
191	cohesion (e.g. wolves; Mazzini et al. 2013) in others. Additional studies are needed to clarify the
192	function of overlapping during foraging in dolphins.
193	
194	As one may expect with vocal matching interactions, the number of whistles that occurred within
195	an overlap match was usually two, indicating only two animals were vocalizing. However, there
196	were two occasions where three matching whistles overlapped (Figure 2). Janik (2000b) also
197	found that matching interactions mostly involved two animals, where each produced a single
198	whistle. He used a randomisation test to show that the number of matching interactions was
199	significantly greater than expected if all animals were calling independently of each other (Janik

200 2000b). The fact that mainly pairs and sometimes trios of animals engaged in these relatively
201 isolated sharing events, makes them different from the production of food calls found in other
202 species (Clay et al. 2012). In the events we reported here, the bray call is the food call, and
203 whistles are used additionally to apparently coordinate behavior between pairs or trios of animals
204 within the larger foraging group.

Interestingly, the fact that only a small number of whistle types appear to be represented in thisdataset suggests that these calls may not be signature whistles. While we had re-sightings of

animals in encounters, our sample comprised a large number of animals. We therefore think that
these whistles may represent shared whistles, which are used by multiple animals in this very
specific context. If so, the matching of calls by bottlenose dolphins may go beyond the exchange
of signature whistles. However, since we cannot completely rule out that the whistles are not
signatures of specific animals, further investigation into whistle use during group foraging is
needed.

We do not know how many different individuals participated in these matching events. Whether it was a small or large number of animals, our results show that overlap matching is closely related to bray call production across multiple recordings. Further work is required to fully understand the function of overlap matching, now that we have shown that there is a social aspect to food calling. Future studies should investigate not only the use of the food call itself, but the social signals that accompany those calls in order to give further insight into the cognitive significance of food-associated vocalizations.

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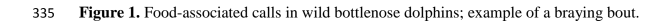
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Table 1. The number of occurrences of the two whistle categories and their mean proximity tothe nearest bray call.

		Whistle Category	# of occurrences	mean proximity to bray call (seconds)
		overlap match	30	82 (range: 0.0001-627)
		overlap no match	55	392 (range: 0.0001-3163)
326				
327				
328				
329	Table 2. The	number of different re	ecording days where o	verlap matching occurred, the number
330	of overlap ma	tches that were record	led, the group size at the	he time of recording, with the
331	additional nur	nber of calves identifi	ed provided in parentl	neses, and the whistle type (frequency
332	contour patter	m).		

Recording Day	# of overlap matches	Group Size	Whistle Type
1	1	7	А
2	1	3	В
3	1	4(1)	А
4	3	3 (1)	BC
5	4	5	А
6	6	8	AD
7	5	18 (2)	AD
8	4	8	ABD
9.1	2	21 (2)	А
9.2	3	11 (1)	А



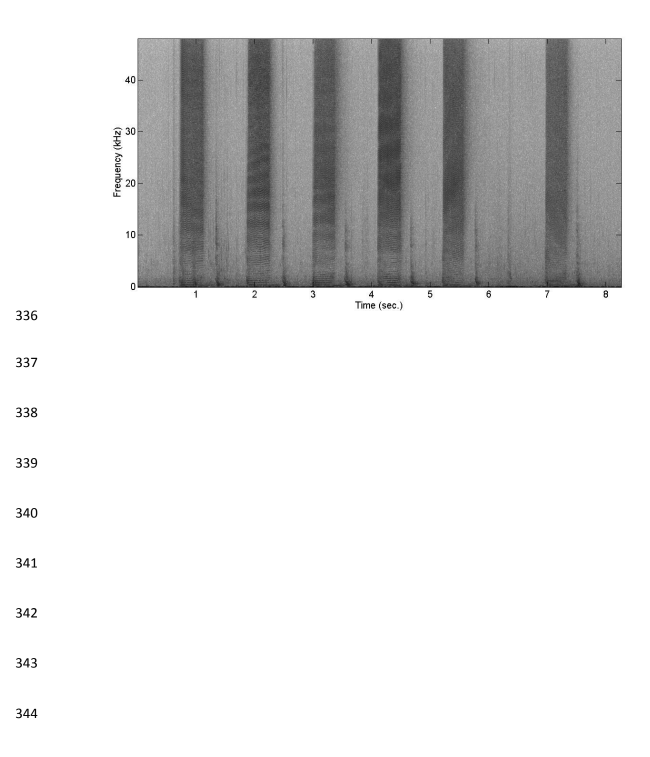
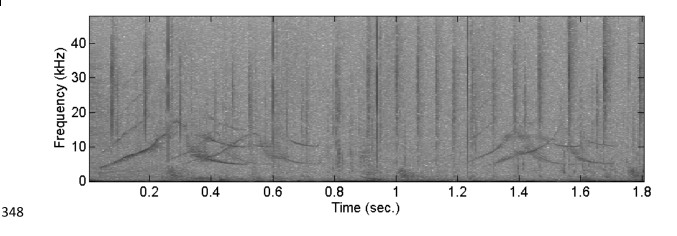


Figure 2. Examples of two occurrences of a matching overlap, where three and then two
whistles of the same type (whistle type A) overlap in time. The end of the first whistle occurs
after the start of the second and third whistle.



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Figure 3. The distribution of proximities to the nearest bray call for the two whistle categories;
the median proximity to the nearest bray call is 13.5 seconds for overlap matches and 71 seconds
for whistles that overlap but do not match.

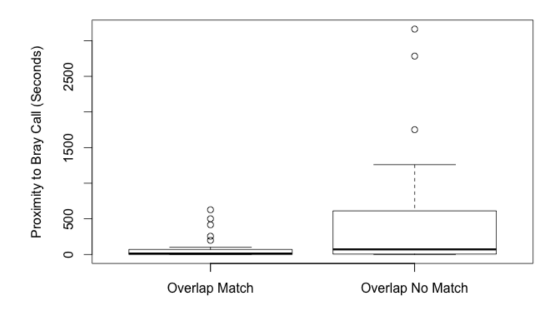


Figure 4. Histograms of the expected mean proximities, based on 10 000 randomisations, of each whistle category to the nearest bray call (seconds), the dotted line indicates the observed mean proximity.

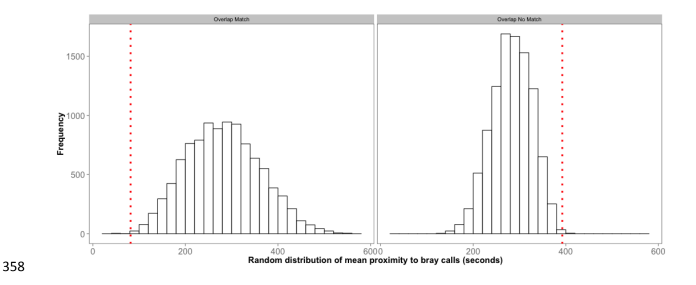


Figure 5. Example of overlap matching, where two whistles of the same type (whistle type D)
overlap in time. The end of the first whistle occurs after the start of the second whistle. A third
whistle (type D) follows later.

