**Distribution and habitat use of a cryptic small cetacean, the Burmeister’s porpoise, monitored from a small-scale fishery platform**

Running title: Habitat use of a cryptic small cetacean

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**Abstract**

There is widespread evidence that small-scale fisheries (SSF) bycatch threatens many populations of small cetaceans, yet conservation efforts are often limited by a lack of basic knowledge regarding their abundance, distribution and habitat use. Here, we used passive acoustic monitoring from an SSF platform-of-opportunity to better characterize the distribution and habitat use of small cetaceans in northern Peru, focussing on the little known Burmeister’s porpoise *Phocoena spinipinnis.* From 2009 to 2012, acoustic click detectors (C-PODs) were attached to fishing nets for the duration of 116 fishing sets (30 fishing trips). Dolphins (unspecified delphinids) and porpoises were recorded around 71% and 22% of fishing sets, respectively. The probability of occurrence and buzzing activity (a proxy of foraging), and time spent were linked to both static and dynamic environmental variables to examine the drivers of habitat use. Dolphin activity was spread evenly throughout the fishing area and was not linked to any habitat variables. In contrast, porpoises were detected in neritic waters and habitat models performed well, identifying preferences for shallow (< 200 m depth) and cooler (17-18 °C) waters, close (< 50 km) to shore. The high bycatch rate of small cetaceans in the Peruvian SSF gave us the unique opportunity to investigate the link between bycatch and cetacean acoustic activity. We found a positive relationship between the likelihood of a bycatch event and acoustic presence for both dolphins and porpoises, however as we did not know the timing of entanglement, we could not link vocalization rates to mortality events. Nonetheless, as small cetaceans (particularly dolphins) frequently encounter fishing boats, the likelihood of entanglements may be reduced through effective efforts to alert animals to the presence of the net, either acoustically (through acoustic alarms) or visually. This study demonstrates that passive acoustic monitoring from a fisheries platform can provide insights into the distribution and habitat use of small cetaceans at relatively low cost, and is likely to be suitable in regions with low monitoring effort and high fishing pressure.

**Introduction**

Artisanal or small-scale fisheries (hereafter SSF) play a vital role in global food production and employment, particularly in developing countries (Berkes et al., 2001; Chuenpagdee et al., 2006; Pauly, 2006). However, there is growing evidence that they have widespread impacts on non-target animals such as marine mammals, seabirds and sea turtles, which are incidentally caught as bycatch (e.g. Awkerman et al., 2006; Peckham et al., 2007; Moore et al., 2010; Alfaro-Shigueto et al., 2011). In particular, many small cetacean (dolphins and porpoises) species are subjected to both direct take and bycatch, as their coastal, estuarine or freshwater, and often restricted, ranges co-occur with large gillnet fleets (Dawson, 1991; Jefferson and Curry, 1994; Reeves et al., 2013). Indeed, bycatch in SSF is thought to have contributed to the extinction of the Baiji or Yangtze River dolphin *Lipotes vexillifer* (Turvey et al., 2007) and threatens several (sub)species of small cetaceans such as the vaquita porpoise *Phocoena sinus* (Jaramillo-Legorreta et al., 2017; Taylor et al., 2017) and Maui’s dolphin *Cephalorhynchus hectori maui* (Slooten et al. 2006). Small-scale fisheries have traditionally received relatively little attention from fisheries managers, while conservation measures are often not implemented due to lack of political will or viable fishing alternatives (Mangel et al., 2010; Read, 2008; Reeves et al., 2013). Additionally, research on the abundance and distribution of captured species is often lacking, so the impacts of SSF on cetacean populations are generally poorly understood (Read, 2008; Reeves et al., 2005).

While there is a need for focussed survey effort in regions with historically high levels of bycatch (Read, 2008), monitoring cetacean abundance and distribution can be costly, logistically challenging and labour intensive (Kyhn et al., 2012; Mellinger et al., 2007). Passive acoustic techniques are a lower cost alternative to visual surveys (Mellinger et al., 2007) and are effective at sampling the behaviour of dolphins and porpoises as they are highly vocal (Marques et al., 2013). Unlike visual surveys which are usually limited by daylight and weather conditions, passive acoustic data can be collected year-round, at night and under most sea states (Kyhn et al., 2012). Passive acoustic methods have proven particularly effective at detecting coastal species that are hard to observe, are known to avoid boats or exist at extremely low densities, providing novel insights into their habitat use and abundance (Gallus et al., 2012; Jaramillo-Legorreta et al., 2017; Rayment et al., 2011).

Generally, passive acoustic devices are towed behind a survey vessel or moored in one location (usually to buoys) for extended time periods (Mellinger et al., 2007; Sousa-Lima et al., 2013). Indeed, static acoustic arrays enable continuous measurements to be collected over long time periods, enabling the quantification of fine-scale temporal patterns in behaviour (e.g. Leeney et al., 2011). However, their limitations are that moorings can be challenging to set-up and maintain (Sousa-Lima et al., 2013), and as the maximum detection ranges of devices are usually a few hundred metres (e.g. Philpott et al., 2007), in the majority of cases, the spatial coverage of the sampled area is extremely limited (Mikkelsen et al., 2016; except see Pirotta et al., 2014). While this limitation may be offset to some degree by the highly mobile nature of many cetacean species, the use of mobile platforms-of-opportunity, such as passenger ferries (e.g. Kiszka et al., 2007; MacLeod et al., 2008) or fishing vessels (e.g. Lopez et al., 2004), provide a means to sample large areas. Data collected from platforms-of-opportunity are not spatially or temporally randomized and so can present analytical challenges (Isojunno et al., 2012); nonetheless, in regions where traditional monitoring methods are prohibitively costly or logistically challenging, platforms-of-opportunity may provide novel insights into relative spatial distributions of cetaceans (e.g. Kiszka et al., 2007; MacLeod et al., 2008). As SSF are ubiquitous in many of the world’s coastal regions (Chuenpagdee et al., 2006; Pauly, 2006), fishing vessels could presumably be used as platforms for monitoring marine species, particularly in data deficient regions (Braulik et al., 2018).

Here, we take advantage of the large spatial extent of the Peruvian driftnet fleet (described in Alfaro-Shigueto et al., 2010; Mangel et al., 2010), to examine the distribution and habitat use of small cetaceans off northern Peru. We focus on the cryptic Burmeister’s porpoise *Phocoena spinipinnis*, a predominantly coastal Phocoenid endemic to South America, with a range spanning from northern Peru to southern Chile in the Pacific and from Tierra del Fuego, Argentina to southern Brazil in the Atlantic (Brownell Jr. and Praderi, 1984; Goodall et al., 1995a). While it is unclear whether it has a continuous distribution throughout its range due to a lack of survey effort (Brownell Jr. and Clapham, 1999), the Peruvian population appears to be genetically differentiated from populations in Chilean and Argentine waters (Rosa et al., 2005). Observations are challenging in anything but calm conditions due to the low, posterior located dorsal fin, and its shy and elusive swimming behaviour (Brownell Jr. and Praderi, 1982, 1984), and there have been remarkably few sightings, with most of them occurring off the coast of Peru (reviewed in Van Waerebeek et al., 2002). While there are no abundance estimates (Jefferson and Curry, 1994), information about the species’ biology has largely been obtained from incidentally or directly captured animals (García-Godos et al., 2007; Reyes and Van Waerebeek, 1995); these studies indicating that it is not uncommon (Brownell Jr. and Praderi, 1982; Rosa et al., 2005; Van Waerebeek et al., 1997; Van Waerebeek and Reyes, 1994).

Burmeister’s porpoises are caught in gillnets throughout its range (Jefferson and Curry, 1994); in particular, large numbers have been captured over the last four decades by SSF in Peru (Majluf et al., 2002; Mangel et al., 2010; Read et al., 1988; Van Waerebeek et al., 1997; Van Waerebeek and Reyes, 1990, 1994). Despite a national law introduced in 1996 banning the capture and trade of dolphins and porpoises, the Peruvian small-scale driftnet fishery predominantly targeting elasmobranchs still has one of the highest rates of small cetacean bycatch in the world, due in part to its vast capacity, with an estimated 10,000-20,000 animals killed per year (Read et al., 1988; Van Waerebeek and Reyes, 1990, 1994; Alfaro-Shigueto et al., 2010; Mangel et al., 2010). Market surveys and on-board observer schemes have indicated that porpoises are caught less frequently than they once were, possibly suggesting population declines in Peruvian waters (Rosa et al., 2005; Tzika et al., 2010). Burmeister’s porpoise is listed as “Data Deficient” by the International Union for the Conservation of Nature (IUCN) (Hammond et al. 2012), while the IUCN Cetacean Specialist Group and International Whaling Commission (IWC) have listed the Peru population as a priority for bycatch reduction, and recommended increased research on its distribution and population status (Reeves et al., 2005).

In this study, acoustic click detectors or C-PODs (Chelonia Ltd., Mousehole, Cornwall, UK) were attached to fishing nets over a three year period, providing information on the acoustic activity of dolphins and porpoises in the vicinity of SSF vessels. Over the duration of the study, the fishery sampled a large geographic area, encompassing a range of marine habitats, from coastal to oceanic waters (Mangel et al., 2010). The aim of the study was to relate acoustic detections of vocalizing small cetaceans (Burmeister’s porpoise and unspecified delphinids) to habitat variables to better understand their space use. Specifically, we 1) briefly describe the acoustic characteristics of Burmeister’s porpoise off the coast of Peru, and 2) identify areas of dolphin and porpoise activity and their associated habitat characteristics. As the fishery has naturally high bycatch rates, particularly of dolphins (Mangel et al., 2010, 2013), we also 3) link acoustic activity of dolphins and porpoises to bycatch events, in order to better understand how small cetaceans become entangled in fishing gear.

**Materials and methods**

*Data collection*

Between April 2009 and December 2012, fishing trips were monitored by observers on SSF vessels from the port of Salaverry (8°14’S, 78°59’W) in northern Peru. SSF are defined according to Peruvian fisheries regulations as boats of <15 m in length, with a maximum of 32.6 m3 of storage capacity and principally based on manual fishing techniques throughout fishing operations (El Peruano, Ley General de Pesca, 2001). Fishing vessels set multifilament driftnets at the sea surface to capture sharks and rays, but they also incidentally capture small cetaceans including Burmeister’s porpoise (Mangel et al., 2010). Vessels depart on fishing trips for approximately one week at a time with a set of nets laid and hauled every 24 hr. Nets are typically 1.5–2 km long and are set during the late afternoon and hauled the following morning with an average soak time of 13 hr (Mangel et al., 2013).

C-PODs are autonomous loggers used to detect echolocation clicks of odontocetes at distances of up to 400 m for porpoises and 500–1,000 m for dolphins (Philpott et al., 2007; Rayment et al., 2009; Tougaard et al., 2006). A C-POD was deployed on each fishing set, in order to detect dolphin and porpoise activity in the vicinity of fishing nets. A device was attached to the lead-line of the net, at approximately 12 m depth (in order to reduce noise interference at the sea surface), and recorded acoustic activity for the duration of the fishing set, and was recovered at the start of the following haul. Five devices were used throughout the study by different fishing vessels, but no more than one was used at the same time by the same vessel. Fisheries observers were trained how to maintain and deploy C-PODs, as well as relevant data collection methods including marine mammal identification, gear used, the timings of net setting and hauling as well as bycatch events. Additionally, at the start of each set, observers recorded the GPS position of the vessel, the sea surface temperature (hereafter SST) using a hand-held Enviro-Safe© thermometer, and sea state using the Beaufort scale.

*Data processing*

Clicks produced by echolocating animals are logged by devices if they show a sufficiently high peak sound pressure level and distinct spectral peak in the frequency range (Tregenza, 2014), and the accompanying C-POD.exe software (*v* 2.032) applies a click train detection algorithm that detects and classifies trains into categories based on how likely they are to be of cetacean origin. The classification software can only distinguish between species groups; narrow-band high frequency (NBHF) echolocation characteristic of all porpoise and some small dolphin species, and broad-band echolocation of all other toothed whale species; boat sonars are also identified. We used the standard KERNO classifier to categorize click trains as dolphin or porpoise in origin, however initial visual screening indicated that some broadband click trains had high modal frequencies and were being wrongly identified as NBHF (Nick Tregenza, *pers. comm.*). To remedy this, we also used an additional classifier (GENENC) developed specifically for this project; its use is recommended both to reduce false positive NBHF clicks in areas where porpoises coexist with broadband species, and to reduce false positive broadband clicks in noisy environments (Robbins et al., 2016; Tregenza, 2014). Only acoustic detections classified in the top reliability categories (“Cet Hi” and “Cet Mod”) were used (Pirotta et al., 2014; Rayment et al., 2009). All click trains were also inspected visually on screen and validated using guidelines provided by the CPOD manufacturer (Gallus et al., 2012; Jaramillo-Legorreta et al., 2017; Tregenza, 2014).

For each fishing set, data were exported as detection positive minutes (DPM), the number of minutes in which click trains were detected, separately for dolphins and porpoises. We also exported the time series of clicks and calculated the inter-click interval (ICI) for each logged click. During approach and capture of prey items, odontocetes produce clicks with shorter and decreasing ICIs, known as feeding “buzzes” (Akamatsu et al., 2005). As such, clicks associated with low ICIs (usually <10 ms; Carlström, 2005) can be used as proxies of foraging behaviour (Leeney et al., 2011; Pirotta et al., 2014). Following Pirotta et al. (2014), we used Gaussian mixture-models fitted to log-transformed ICIs to identify multimodal peaks, representative of different behaviours: 1) low ICIs associated with buzzing behaviour, 2) regular clicks, and 3) intervals between click trains (see Supplementary Material for details; Pirotta et al., 2014).

*Environmental data*

 Previous studies have indicated that small cetaceans have preferences for habitats influenced by bathymetry and distance to the shore (Embling et al., 2010; Isojunno et al., 2012; Kiszka et al., 2007). As a result, for each fishing set locations, we extracted the following variables in ArcGIS 10.1: 1) ocean floor depth (hereafter Depth) was obtained from the General Bathymetric Chart of the Oceans (GEBCO, [www.gebco.net](http://www.gebco.net); IOC 2003) and 2) ocean floor slope (Slope) was calculated using the *slope* function in the Spatial Analyst toolbox. We also calculated 3) the Euclidean distance from the nearest coast (Distance). The spatial resolution of the bathymetry layer was one arc second, corresponding to c. 30 m at the equator.

*Statistical analysis*

 Analyses were conducted separately for porpoises and dolphins to determine the influence of habitat variables on variation in their presence-absence (1 = present, 0 = absent), and once present (DPM > 0), the drivers of time spent in the local area. The data were modelled in a two-step process using hurdle generalized linear mixed models (GLMMs) in the R package *glmmADMB* (Fournier et al., 2012): for the first (zero) part, we estimated the probability of presence using a binomial error structure and logit link function; and for the second (count) part we modelled DPM as the response variable using only positive values, and specifying a zero-truncated negative binomial error distribution and log link function (Zuur et al., 2009). Negative binomial was chosen over a Poisson distribution due to the presence of a few extremely large and many small observations. Each data point corresponded to one fishing set, which we considered a discrete monitoring unit in space and time. However, as sets within fishing trips were non-independent, trip identity was included as a random effect. Also, as recording duration varied with each set, we initially included the log-transformed duration of time recorded as an offset in count models, and as a fixed effect in presence-absence models as it is not possible to include offsets in binomial error models. In separate models we also investigated the drivers of presence-absence of buzzing activity using GLMMs with a binomial error structure and logit link in the R package *lme4* (Bates et al., 2015).

 The following habitat variables were considered as covariates (summarized in Table 1): 1) SST, 2) Depth, 3) Slope, 4) Distance, 5) year and 6) season (quarter 1 = January – March, 2 = April – June, 3 = July – September, and 4 = October – December), to account for disparities in sampling effort. We also included the following operational factors: 7) bait use, as the presence of bait may influence cetacean behaviour, 8) C-POD identity to control for variation in detection capabilities among devices, and 9) sea state. Finally, we also included both 10) a factor corresponding to the bycatch (entanglement) events during the set, and 11) the number of individuals captured. These variables were tested to determine a relationship between acoustic activity and bycatch, however we cannot determine causality between the two as we do not have precise information on the timing of entanglement events.

Prior to modelling, Depth and Slope were log-transformed to improve data spread, while in the porpoise count model, Distance was square-root-transformed. We checked for collinearity between explanatory variables using variance inflation factors (VIF). Due to the small number of observations, we initially limited the number of explanatory variables included by testing the significance of all variables (and where appropriate, their quadratic terms), in standalone models, and compared them to null (intercept only) models using likelihood ratio tests (LRTs, Table S1). For several models, this process still yielded a large number of covariates; as such, the most parsimonious models were selected by the sequential stepwise addition of significant variables, in order of decreasing importance (ranked using $X\_{}^{2}$ values; Table S1). Significance was assessed with LRTs and variable importance was determined as the proportional deviance explained by calculating the percentage reduction in residual deviance upon removal from the most parsimonious model (Zuur et al. 2009). We evaluated the performance of binomial GLMMs by computing the area under the Receiver Operator Characteristic (ROC) curve (AUC) in the R package *pROC* (Robin et al., 2017). Values of 0.5–0.7, 0.7–0.9, and >0.9 represent poor, reasonable and very good model performance, respectively. Unless otherwise specified, all analyses were conducted in the statistical program R 3.3.1 (R Core Team, 2014) and means are given + standard error (SE).

**Results**

Between April 2009 and December 2012, 116 sets were observed across 30 fishing trips (a mean of 3.9 ± 1.6 sets per trip). Logistical constraints prevented monitoring in the first half of 2010 and for parts of 2012, and so a greater number of sets were observed in 2009 and 2011 (Fig. 2). Effort was also not spread evenly across seasons with a greater number of sets observed in quarters 2 and 4 (autumn and spring, respectively). C-PODs attached to fishing nets logged 1,279 hours (53.3 d) of data, with a mean of 11.0 ± 3.6 hours per set. While the recording duration was occasionally cut short by device malfunction, there did not appear to be any spatial pattern in the monitoring duration (Fig. S1).

*Acoustic characteristics*

The acoustic presence of small cetaceans (dolphins or porpoises) was recorded by C-PODs in 76% (n = 88) of fishing sets; dolphins (71%, n = 82) were recorded in a higher number of sets than porpoises (22 %, n = 25). Over the study duration, 56,222 and 190,976 clicks were classified in 4,687 and 3,231 click trains, as belonging to dolphins and porpoises, respectively. Dolphins were detected in 4,101 min, a mean of 50 ± 50 (range: 0–463) per set and 4.6 ± 6.2 h-1, and porpoises detected in 1,002 min, a mean of 40 ± 50 (0–206) per set and 3.0 ± 4.1 h-1. The average modal frequency of click trains was 133.9 ± 4.4 (121–144) kHz and 117.6 ± 23.6 (30–146) kHz for porpoises and dolphins, respectively. Dolphin and porpoise ICIs ranged from *ca.* 1 µs to *ca.* 30 ms and from *ca.* 8 µs to *ca.* 15 ms, respectively. Buzz ICIs were identified by Gaussian mixture models with an average ICI of 46 ± 0.2 μs and 98.2 ± 0.5 μs for dolphins and porpoises respectively, and buzzing activity was recorded in 57% (n = 66) and 8% (n = 9) of sets for dolphins and porpoises, respectively, constituting 26% (n = 49,182) and 17% (n = 9,413) of total clicks detected, respectively.

*Distribution and habitat use*

The minimum convex polygon encompassing the distribution of fishing effort covered an area of 61,557 km2, which included a range of marine habitats (coastal, continental shelf, shelf-break and oceanic waters; Fig. 2) and a large range in SST values (15 – 25°C). Burmeister’s porpoises were predominantly detected in coastal regions slightly offshore from Salaverry and a near-shore region around 50 km northwards (Fig. 2a). In contrast, dolphins were detected throughout the sampled area (Fig. 2b).

For some sets, observers did not record certain variables, such as bait use, sea state, SST or the GPS position, and so sample sizes varied depending on which explanatory variables were included (Table S1). The variables Depth, Slope and Dist. were correlated, so if more than one was deemed significant, only that which resulted in the highest deviance explained, was chosen. A similar procedure was used for the categorical variables C-POD ID and year, which were correlated in porpoise count models. Initial fitting of standalone models for each habitat variable indicated that several were important predictors of porpoise but not dolphin activity (Table S1). Final models were then assessed through forwards selection of significant variables; performance was very good for both the porpoise occurrence and buzzing occurrence models (AUCs = 0.94 and 0.98 respectively), while performance was poor and reasonable for dolphin occurrence (0.65) and buzzing occurrence (0.77) models, respectively.

The most parsimonious models included the effect of depth on porpoise occurrence ($X\_{1}^{2}$ = 36.9, *p* < 0.001; Table 2), SST on time spent ($X\_{1}^{2}$ = 12.2, *p* < 0.001; Table 2) and distance from the shore on the occurrence of buzzing activity ($X\_{1}^{2}$ = 16.4, *p* <0.001; Table 3). In particular, depth and distance explained a large percentage of deviance (> 20%) in their respective models (Tables 2 and 3). Porpoises were detected in shallow waters up to 200 m depth, and predicted probability of presence was highest in waters of 50 m depth or shallower (Fig. 3a). Porpoises spent longer periods of time in cooler waters of 17 - 18°C, and appeared to avoid areas with SSTs > 19°C (Fig. 3b). Porpoise buzzing behaviour was detected within 50 km from the shore, but occurrence was greatest within the first 20–30 km, indicating that porpoises preferentially forage in shallow waters close to the shore (Fig. 3c).

No habitat variable explained dolphin occurrence, time spent or buzzing occurrence (Tables 2 and 3). The C-POD recording duration was a significant predictor of dolphin occurrence ($X\_{1}^{2}$ = 4.8, *p* = 0.029; Table 2) and buzzing activity ($X\_{1}^{2}$ = 4.4, *p* = 0.035; Table 2), such that with longer durations, there was an increased probability of presence and buzzing activity, which appeared to plateau at around 800 min (Fig. 3c, d). The recording duration was not important for porpoises, while the use of bait, C-POD ID, sea state, slope, quarter and year were not significant predictors of dolphin or porpoise activity (Tables 2 and 3, Table S2).

*Acoustic activity and bycatch*

Dolphins were incidentally captured in 19% of sets (n = 22) and a total of 30 individuals were captured, consisting of 18 common dolphins *Delphinus capensis*, six dusky dolphins *Lagenorhynchus obscurus*, five bottlenose dolphins *Tursiops truncatus* and one Risso’s dolphin *Grampus griseus* (Fig. 2d). The dolphin bycatch per unit effort (BPUE) was 1.4 ± 0.5 animals per set for sets with bycatch and 0.3 ± 0.6 individuals per set for all sets. Burmeister’s porpoises were caught in 2% of sets (n = 2) (Fig. 2c), with a mean BPUE of 0.02 ± 0.1 individuals per set for all sets; both these sets registered acoustic activity of porpoises. Of those 22 sets with dolphin bycatch, 20 recorded acoustic presence. Porpoise bycatch was associated with increased acoustic time spent ($X\_{1}^{2}$ = 20.8, *p* < 0.001; Table 2, Fig. 4a), but was not linked to the probability of presence or occurrence of buzzing (Tables 2, 3). Similarly, the bycatch of dolphins was significantly linked to higher probability of presence ($X\_{1}^{2}$ = 5.5, *p* = 0.019; Table 2, Fig. 4c) and occurrence of buzzing activity ($X\_{1}^{2}$ = 7.0, *p* = 0.008; Table 3, Fig. 4d), but not to time spent around fishing vessels (Table 2). Dolphins and porpoises were still detected in sets where no bycatch was recorded (53% and 17% of sets, respectively). As we do not know the timing of bycatch in relation to the acoustic detections, or the number of animals around the net, we cannot determine whether some bycatch is of silent animals. Finally, bycatch as a categorical variable explained activity metrics better than the continuous variable, suggesting no link between the number of individuals caught and acoustic activity.

**Discussion**

Knowledge of the distribution and habitat use of small cetaceans in crucial for informing conservation efforts, particularly in regions where they overlap or interact with human activities such as fisheries. Our study demonstrates that passive acoustic monitoring from a SSF platform can provide novel insights into the distribution and habitat use of small cetaceans. We identified a coastal region that is likely to be important for the little studied Burmeister’s porpoise in the northern extent of its range. In contrast, as dolphins were detected in equal measure throughout the fishery area, we were unable to link occurrence or time spent to habitat variables. The latter results are likely to be confounded as passive acoustic techniques make distinguishing between different broadband echolocating species extremely challenging (see Robbins et al., 2016). Dolphin activity was likely to belong to a number of sympatric species, including those recorded as bycatch (common, bottlenose, dusky and Risso's dolphins; Mangel et al., 2010, 2013).

It is generally recommended that acoustic recordings be supplemented with visual sightings (Leeney et al., 2011; Rayment et al., 2011), however this was not possible as nets were set predominantly during darkness. Nonetheless, observations of Burmeister’s porpoise are difficult in anything but calm conditions due to its posterior located dorsal fin and neophobic behaviour (Brownell Jr. and Praderi, 1982, 1984), and as a result of this, as well as scarce monitoring, there are fewer than 20 documented sightings in Peru (reviewed in Van Waerebeek et al., 2002). Porpoises are characterised by their similar vocalizations: high frequencies centred around 125-140 kHz narrow-band (10kHz) clicks (Akamatsu et al., 1994). A recent study has provided the first characterization of Burmeister’s porpoise sounds, and confirmed they use NBHF clicks, with peak frequency of 135 ± 2 kHz (Reyes Reyes et al., 2018), which is similar to the average modal frequency of click trains documented here (133.9 ± 4.4 kHz; range 121 – 144 kHz). While verification of NBHF detections is an important future step, we are confident that NBHF clicks belonged to Burmeister’s porpoise as it is the only NBHF species in Peruvian coastal waters, unlike in other parts of the species range such as southern Chile and Argentina, where it co-occurs with *Cephalorhynchus* dolphins (which also use NBHF echolocation; Heinrich, 2006, Morisaka and Connor 2007). This is supported by the fact that the two fishing sets in our study to have porpoise bycatch, also recorded NBHF activity.

*Distribution and habitat use of Burmeister’s porpoise*

 Through collaboration with small-scale fishers, we were able to acoustically sample a range of habitats from near-shore and continental shelf to shelf-edge and oceanic regions, at relatively low cost. However, we acknowledge that like other opportunistic surveys, our methodology had associated limitations (Isojunno et al., 2012); for example, the activity of small cetaceans around fishing vessels may not be considered natural behaviour. In other regions, dolphins are known to depredate from fishing nets (e.g. Read et al., 2003), however Peruvian dolphins and porpoises do not appear to be attracted to bait and have never been observed by fishers depredating from or feeding in the nets (Pro Delphinus, *pers. comm.*). Indeed, dietary analyses of bycaught cetaceans indicate their main prey species are forage fish such as the Peruvian anchovy *Engraulis ringens* and squid (Garcia-Godos et al., 2007); in contrast, the drift-net fishery captures large fish (predominantly sharks and rays) and the large mesh size of nets prevents capture of smaller fish (Alfaro-Shigueto et al., 2010; Mangel et al., 2010). It is more likely that they are detected while passing through the area or while feeding on nearby prey, however we acknowledge that dolphins (but probably not porpoises) may be attracted to fishing boats due to their inquisitive nature, which may bias our estimates of habitat use. As this bias is likely to be similar across fishing vessels and habitats, our main conclusion that dolphins (unspecified delphinids) were detected throughout the fishing area and porpoises were limited to a neritic and near-shore waters, remains unchanged.

 Past research on the foraging ecology of Burmeister’s porpoise has largely been anecdotal, with the majority of information derived from studies of bycaught individuals (García-Godos et al., 2007; Reyes and Van Waerebeek, 1995; Rosa et al., 2005; Tzika et al., 2010); nonetheless, our results corroborate the limited number of documented sightings (reviewed in Van Waerebeek et al., 2002) and studies of SSF bycatch (Mangel et al., 2010, 2013) that suggest it is predominantly a neritic species. We found several habitat variables to have a significant influence on porpoise activity: porpoises were detected in regions with <200 m bottom depth and <50 km from the shore, with greater time spent in cooler waters (17–18 °C). Important relationships with topographic variables such as bottom depth and distance to the shore have been observed in many other coastal cetaceans (e.g. Embling et al., 2010; Garaffo et al., 2007). While no habitat variables predicted dolphin occurrence, there appeared to be subtle partitioning by distance from the shore, whereby dolphin activity was lower in sets closest to the coast where porpoise activity was highest. Burmeister’s porpoise use similar depths in other parts of its range, however the manner in which habitats are partitioned varies between regions (Heinrich, 2006).; For example, in southern Chile spatial segregation with Chilean *Cephalorhynchus eutropia* and Peale’s dolphins *Lagenorhynchus australis* has resulted in porpoises using slightly deeper waters (20-50 m compared to 0-25m for the two dolphin species) (Heinrich, 2006), whereas in Golfo San Jose, Argentina, porpoises use the intermediate depths (5-15 m) in between inshore bottlenose dolphins (< 10 m) and dusky dolphins (35 m) (Goodall et al., 1995b).

 For both species groups we found evidence of buzzing activity, which is generally thought to represent foraging behaviour (Akamatsu et al., 2005; Pirotta et al., 2014) In particular, the higher probability of porpoise buzzing activity in near-shore habitats likely represents less time spent travelling (or other activities) and more time spent feeding. This could reflect the distribution of their preferred prey; despite similarity between diets of Peruvian small cetaceans, Burmeister’s porpoises were found to have a higher reliance on anchovy than other species (occurrence of 88% in stomachs of bycatch individuals, Garcia-Godos et al. 2007). The Humboldt Current upwelling is one of the most productive marine ecosystems in the world, and in particular, the continental shelf off Salaverry hosts large densities of anchovy, particularly in December – April (Bertrand et al., 2004b), and a large industrial fishery, based out of numerous ports in central and northern Peru (Fréon et al., 2008). Anchovy aggregations exhibit large spatial variability depending on the year and season (Fréon et al., 2008), yet while our sampling effort was uneven across years and seasons, we did not find any evidence of temporal variation in porpoise or dolphin activity. Visual observations and catch records indicate little seasonality in movements (Van Waerebeek et al., 2002; Van Waerebeek and Reyes, 1990, 1994), however our limited sampling for a given season prevents a robust comparison. Anchovy distribution is tightly linked to cold and productive coastal waters (Bertrand et al., 2004a), which may explain why porpoise time spent was in the coolest waters sampled. Indeed, this preference for cooler waters in the northern part of their distribution indicates that porpoises might be at the limits of their thermal tolerance, and are likely to be particularly affected by increases in SSTs and shifts in anchovy distributions during El-Nino years (García-Godos et al., 2007), as demonstrated by high numbers of stranded individuals on Peruvian shores during these periods (Van Waerebeek et al., 1997).

*Links between small cetacean acoustic activity and bycatch*

 Little is known about how small cetaceans become entangled in gillnets, due to the difficulty of underwater observations (Martin and Crawford, 2015; Read et al., 2003), and their high bycatch rate in the Peruvian driftnet fishery gave us the unique opportunity to investigate links between entanglement events and acoustic activity. Firstly, we found small cetaceans to be in the vicinity of gillnets for a large proportion (76%) of fishing sets. As bycatch is both the product of exposure (driven by distribution of animals and fishing activities and magnitude of effort) and vulnerability of a species (ecological characteristics of a species, e.g. behaviour) (Lewison et al., 2014), the high encounter rate, particularly of dolphins (71% of sets had dolphin activity), goes some way to explaining why tens of thousands of individuals are likely caught by this fleet annually (Mangel et al., 2010; Van Waerebeek and Reyes, 1990, 1994). While the bycatch rate of this fishery is among the highest globally (Mangel et al. 2010), much fewer sets (<20%) recorded bycatch of small cetaceans than acoustic presence. These results indicate that small cetaceans are present in the vicinity of gillnets for long periods of time without becoming entangled, mirroring the observations of Read *et al.* (2003) that bottlenose dolphins frequently encounter and interact with nets without becoming entangled.

 We also found a positive relationship between the likelihood of a bycatch event and acoustic activity for both dolphins and porpoises. Although we only recorded two porpoise bycatch events, DPM in these sets was significantly higher. We were not able to record the timing of entanglement events and so the temporal relationship between vocalization and entanglement lacks precision. Also, it is important to note that gillnets were generally longer than the likely detection range of small cetaceans by C-PODs (Philpott et al., 2007; Rayment et al., 2009; Tougaard et al., 2006), and so there were likely to be portions of the net that were not detected. Our findings support previous suggestions that entanglement results either from travelling animals not detecting the net as a barrier, or from entanglements while feeding in the vicinity of the net (Cox and Read, 2004). As dolphins and porpoises have a sophisticated sonar, entanglement in nets is generally thought to occur as a result of mistakes made, rather than their sonar not detecting the net (Dawson, 1991; Nielsen et al., 2012). However, a monofilament net may be hard to detect at some distance (Kastelein et al., 2000). Dolphins are not always actively engaged in echolocation, and may become entangled when not vocalizing (Dawson et al., 2013), particularly in low light levels such as at night (Akamatsu et al., 1991) or in particularly turbid or noisy environments (Martin and Crawford, 2015; Northridge et al., 2017). Indeed, there is anecdotal evidence from Peruvian fishers that dolphin bycatch is higher in windier conditions (Pro Delphinus, *pers. comm.*), which may be the result of increased mixing of sediments or reduced acoustic detectability of nets due to high levels of ambient noise, however this requires further investigation.

*Conservation implications*

 Coastal cetaceans are exposed to a range of human activities including fisheries, aquaculture, shipping, pollution and climate change (Mann, 2000), yet for many species, still little is known about their abundance and distribution (Read, 2008; Reeves et al., 2005). Using a relatively low-cost monitoring tool (Braulik et al. 2018), our study provides a quantitative assessment of the distribution and habitat use of Burmeister’s porpoise, adding substantially to the knowledge base of a “Data Deficient” species (Hammond et al., 2012). One of our key findings was that porpoises use a narrow coastal region (< 50km from the shore); indeed, their distribution studied here in the northern extent of their range completely overlaps the extent of a small-scale gillnet fishery. Given historically large capture rates of small cetaceans and the large capacity of the gillnet fleet (Alfaro-Shigueto et al., 2010; Mangel et al., 2010; Read et al., 1988; Van Waerebeek and Reyes, 1994), our finding that dolphins and porpoises regularly encounter gillnet vessels (recorded in 71% and 22% of monitored sets, respectively), raises serious concerns about the long-term impact of this fishery on local cetacean populations (Read et al. 1988, Reeves et al. 2005, Rosa et al. 2005, Van Waerebeek et al. 1997), and highlights the chronic nature of these interactions. While no abundance estimates exist, information gathered from onboard observer schemes and in-port and market surveys suggests Burmeister’s porpoises used to be more common in Peruvian waters, but their numbers relative to other small cetaceans have decreased over the last 30 years (Mangel et al., 2010; Tzika et al., 2010). The Peruvian population is not thought to undergo long-range movements (Reyes and Van Waerebeek, 1995) and is reproductively isolated from the Chilean population (Rosa et al., 2005). Furthermore, Burmeister’s porpoise is believed to be particularly susceptible to changes in the abundance and distribution of anchovy, its main prey (García-Godos et al., 2007). Given the lack of information on movements and population trends, we strongly encourage formal monitoring efforts, including the establishment of passive acoustic arrays, ideally in combination with boat-based surveys or satellite tracking (e.g. Mikkelsen et al., 2016).

 The results of our study suggest that the protection of key habitats for porpoises may be more straightforward than for dolphins, given their more restricted range. Successful mitigation of SSF bycatch is likely to be achieved through multiple approaches; for example, restricting fishing activities in near-shore waters, or use of acoustic alarms (pingers) on nets (e.g. van Beest et al., 2017). Indeed, a previous study has shown that pingers reduce bycatch of Peruvian small cetaceans (Mangel et al., 2013), and future work should refine the efficacy of using pingers to deter porpoises and dolphins from nets (Dawson et al. 2013). These approaches do not come without social, economic or logistical constraints, and the cost of pingers remains a major impediment to their implementation (Dawson et al., 2013, Mangel et al., 2013). As such, widespread use of pingers or other bycatch mitigation solutions will likely require concerted political will or market-based solutions that reward fishers for responsible practices (Jacquet and Pauly 2008, Jose Alava et al., 2017).

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**Tables**

**Table 1.** Explanatory variables included in models of Burmeister’s porpoise *Phocoena spinipinnis* and dolphin (unspecified delphinids) activity around fishing vessels in northern Peru. Abbreviations of variable names and units referred to in the text are shown in parentheses.

|  |  |  |  |
| --- | --- | --- | --- |
| **Explanatory variable** | **Type** | **Units** | **Description** |
| **Habitat** |  |  |  |
|  | Sea floor depth (Depth) | Continuous | Metres (m) | Extracted from GEBCO layer in ArcGIS 10.1 |
|  | Sea floor slope (Slope) | Continuous | Degrees (°) | Extracted from GEBCO layer in ArcGIS 10.1 |
|  | Distance from the coast (Dist) | Continuous | Kilometres (km) | Extracted in ArcGIS 10.1 |
|  | Sea surface temperature (SST) | Continuous | Degrees Celsius (°) | Recorded at start of set. |
| **Temporal** |  |  |  |
|  | Season | Categorical | n/a | Quarter: 1 (Jan–Mar), 2 (Apr–Jun), 3 (Jul–Sep), 4 (Oct–Dec) |
|  | Year | Categorical | n/a | Calendar year: 2009 - 2012 |
| **Operational** |  |  |  |
|  | Bait | Categorical | n/a | Presence of small cetacean bait |
|  | C-POD ID | Categorical | n/a | Identity of C-POD |
|  | Sea state | Continuous | Beaufort scale | Recorded at start of set, ranging 0 (calm) to 6 (strong breeze).  |
|  | Recording duration (Duration) | Continuous | Minutes (min) | Time recorded by C-POD in each fishing set |
| **Bycatch** |  |  |  |
|  | Bycatch occurrence (Bycatch) | Categorical | n/a | Presence of cetacean bycatch |
|  | Bycatch number | Continuous | - | Number individual cetaceans captured |

**Table 2.** Explanatory variables retained by the Burmeister’s porpoise *Phocoena spinipinnis* and dolphin occurrence (presence-absence; binomial) and time spent (detection positive minutes [DPM]; count) components of hurdle models. Important variables are given separately for each component of the hurdle model. The percentage of deviance explained were calculated on removal from the minimum adequate model. No variables were considered important predictors of dolphin DPM.

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| Species group | Response variable | Explanatory variable | Estimate ± SE | *X2* | *p* | Dev. Explained (%) |
| Porpoise | Occurrence | Depth | -2.94 ± 0.74 | 36.9 | <0.001 | 36.6 |
| DPM | Bycatch | 3.43 ± 0.45 | 20.8 | <0.001 | 9.8 |
| SST | -0.84 ± 0.23 | 12.2 | <0.001 | 6.0 |
| Dolphin | Occurrence | Bycatch | 1.81 ± 0.89 | 5.5 | 0.019 | 4.4 |
| Recording duration | 1.27 ± 0.62 | 4.8 | 0.029 | 3.8 |
| DPM | - | n/a | n/a | n/a | n/a |

**Table 3.** Explanatory variables retained by the Burmeister’s porpoise *Phocoena spinipinnis* and dolphin buzzing occurrence (presence-absence; binomial) models. Important variables are given separately for each component of the model. The percentage of deviance explained were calculated on removal from the minimum adequate model.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| Species group | Explanatory variable | Estimate ± SE | *X2* | *p* | Dev. Explained (%) |
| Porpoise | Distance from the shore | -0.10 ± 0.07 | 16.4 | <0.001 | 33.1 |
| Dolphin | Bycatch | 1.53 ± 0.65 | 7.0 | 0.008 | 4.8 |
|  | Recording duration | 1.05 ± 0.53 | 4.4 | 0.035 | 3.1 |

**Figures**

**Fig. 1.** The distribution of gillnet fishing sets observed by a) year and b) quarter, from the port of Salaverry in northern Peru, over the study duration. Quarter 1 = January-March, 2 = April-June, 3 = July-September, 4 = October-December.

**Fig. 2.** Map of the study area, (a-b) acoustic activity recorded by C-PODs attached to gillnets and (c-d) bycatch, of Burmeister’s porpoise (left panels) and dolphins (unspecified delphinids; right panels) in fishing vessels operating from the port of Salaverry, northern Peru. For plots of acoustic activity and bycatch, the locations of sets where small cetaceans were present and absent are shown by circles and crosses, respectively, with the size of circles representing time spent (detection positive minutes [DPM] per hour) and the number of individuals captured, respectively. The position of the 100, 200, 500, 1,000 and 2,000 m isobaths are shown with grey lines.

**Fig. 3.** Responses of small cetaceans to important variables included in models: a) bathymetry depth on Burmeister’s porpoise *Phocoena spinipinnis* probability of presence, b) sea surface temperature on porpoise time spent (detection positive minutes [DPM]), c) distance from the shore on the probability of porpoise buzzing activity, and the C-POD recording duration on the d) probability of presence and e) buzzing activity of dolphins (unspecified delphinids). For a) the depth range is much greater (c. 2,000 m), but only depths up to 500 m are plotted. Observed values are shown by grey dots, and the predicted relationship by a black line.

**Fig. 4**. The modelled relationship between bycatch of Burmeister’s porpoises *Phocoena spinipinnis* and dolphins (unspecified delphinids) and their acoustic activity recorded by C-PODs attached to fishing nets: for porpoises, a) time spent (detection positive minutes [DPM]), shown on the natural log scale, and for dolphins, c) the probability of presence and d) buzzing activity. Error bars represent the 95 % confidence intervals around modelled estimates. b**)** A Burmeister’s porpoise *Phocoena spinipinnis* hauled onto the deck of an SSF vessel after becoming entangled in a driftnet in northern Peru. Photo credit: ProDelphinus.

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**Fig. 1.**

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**Fig. 2.**

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**Fig. 3.**

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**Fig. 4**.