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Gregory B. Cunningham
Saint John Fisher College, gcunningham@sjfc.edu

Richard W. Van Buskirk
Pacific University

Mark J. Hodges
University of California - Davis

Gabrielle A. Nevitt
University of California - Davis

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Responses of common diving petrel chicks (Pelecanoides urinatrix) to burrow and colony specific odours in a simple wind tunnel

GREGORY B. CUNNINGHAM1,2, RICHARD W. VAN BUSKIRK1,3, MARK J. HODGES1 and GABRIELLE A. NEVITT1

1Department of Neurobiology, Physiology and Behavior, One Shields Avenue, University of California, Davis, CA 95616, USA
2Current address: Department of Biology, St John Fisher College, 3690 East Avenue, Rochester, NY 14618, USA
3Current address: Environmental Studies Department, Pacific University, 2043 College Way, Forest Grove, OR 97116, USA
gcunningham@sjfc.edu

Abstract: Researchers have previously assumed that common diving petrels (Pelecanoides urinatrix) have a limited sense of smell since they have relatively small olfactory bulbs. A recent study, however, showed that adult diving petrels prefer the scent of their own burrow compared to burrows of other diving petrels, implying that personal scents contribute to the burrow’s odour signature. Because diving petrels appear to be adapted to use olfaction in social contexts, they could be a useful model for investigating how chemically mediated social recognition develops in birds. A first step is to determine whether diving petrel chicks can detect familiar and unfamiliar odours. We compared behavioural responses of chicks to three natural stimuli in a wind tunnel: soil collected from their burrow or colony, and a blank control. During portions of the experiment, chicks turned the least and walked the shortest distances in response to odours from the nest, which is consistent with their sedentary behaviour within the burrow. By contrast, behaviours linked to olfactory search increased when chicks were exposed to blank controls. These results suggest that common diving petrel chicks can detect natural olfactory stimuli before fledging, and lay the foundation for future studies on the role of olfaction in social contexts for this species.

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Introduction

Diving petrels (Procellariidae) are small, sub-Antarctic seabirds that nest in underground burrows. During the breeding season, these birds forage at sea (Bocher et al. 2000a) and periodically return to the burrow to provision a single chick after dark to avoid predation (reviewed by Warham 1990). Among birds, procellariiforms are known for their olfactory abilities (reviewed by Nevitt 2008), but diving petrels were historically thought to have a diminished sense of smell (e.g. Cunningham et al. 2003). Diving petrels have the smallest olfactory bulb of all procellariiforms (mean bulb to brain ratio for the common diving petrel Pelecanoides urinatrix (Gmelin, 1789) 18%, for all procellariiforms 29%, Bang & Cobb 1968). Moreover, the low aspect ratio of the diving petrel wing limits their ability to engage in dynamic soaring, a characteristic flight style of many procellariiforms. Instead, they rapidly flap their wings (12.3 Hz, Pennycuick 1996) and forage by diving to considerable depth (c. 33 m, maximum depth 60+ m; Chastel 1994, Bocher et al. 2000b). This foraging style suggests that they do not track surface odours to locate prey resources (reviewed by Nevitt 2008).

However, diving petrels have recently been shown to detect odour cues in contexts other than foraging. For example, during the breeding season, a variety of adult burrow-nesting procellariiforms, including both common and South Georgian diving petrels (Pelecanoides georgicus Murphy and Harper, 1916) have been shown to use scent cues to relocate their burrow when returning from foraging trips (Bonadonna et al. 2003, reviewed by Hagelin 2007) and in some species scent may play a role in individual recognition (Bonadonna & Nevitt 2004). Still, relatively little attention has been paid to investigating how these abilities develop in chicks. This is because petrel chicks do not typically leave their burrows until just before fledging and unless they are natally philopatric, these birds should not need to learn odours to relocate their burrow. However, we have recently proposed that odours associated with self/non-self recognition may be important for mate choice, and that learning self-specific odours begins before the chick leaves the nest in some species (for discussion see Nevitt 2008, O’Dwyer & Nevitt 2009). Since current evidence suggests that diving petrels use olfaction primarily in social rather than in foraging contexts, they may make an ideal model among birds for studying the development of social odour recognition.
Fig. 1. Mean distances (cm) walked by common diving petrel chicks in response to nest soil (black), colony soil (grey) and control presentations (white). Values are given as mean ± standard error. Repeated measures ANOVA revealed a significant difference among treatments during minute 2 (Greenhouse-Geisser: $F(1.53, 33.56) = 3.94, P = 0.04$). In minute 2, chicks walked farther in the presence of the empty dish compared to their own nest soil (Tukey’s test: $P = 0.05$). The data were analysed using a square root transformation, but are presented here as raw numbers. There were no other differences in the distance walked (see text).

In the present study, our goal was to determine whether common diving petrels could detect familiar, burrow related odours. Diving petrel chicks are not mobile enough to perform choice tests (e.g. O’Dwyer et al. 2008), but olfactory responses can be tested using a simple wind tunnel (Cunningham et al. 2006). Unlike Y-maze experiments, the wind tunnel does not require chicks to make a choice, but can be used to elicit behaviours related to olfactory search. Here we compared the behavioural responses of 23 common diving petrel chicks to the scents of soil from their nest chamber (nest soil), soil from the surface of the area surrounding the burrows (colony soil), and an empty Petri dish (control). Given that chicks tend to stay within the nest chamber of their burrow throughout development, we predicted that chicks would sit still and search less in response to their own burrow odour than to the other stimuli.

Materials and methods

This study was performed on Mayes Island (49°28’S, 69°57’E), Iles Kerguelen, from 21–25 February 2002. Pelecanoides urinatrix nests were identified and monitored daily during the incubation stage to determine hatch date for each chick. Chicks were then tested at 30 ± 0.9 (mean ± s.e.) days post hatching (fledging occurs at 50–55 days, Jouventin et al. 1985). All trials were conducted between 10h00 and 14h00 local time at temperatures of 13.0 ± 0.5°C.

We performed experiments in a simple wind tunnel, which has been described elsewhere (see fig. 1 and text in Cunningham et al. 2006). Briefly, the apparatus consisted of a testing arena (0.8 × 0.6 m, 0.3 m high) constructed of Plexiglas®, and was housed in a well-ventilated hut (1.5 × 1.5 m, 2.5 m high). Airflow (0.6–0.7 m sec$^{-1}$) to the arena was generated by two tubeaxial fans (Mechatronics model UF12A12, Preston, WA, USA; air volume: 3 m$^3$ min$^{-1}$) mounted to the sides of a pressure box that was located at one end of the arena. These fans were positioned at a 90° angle to an exit port that directed air into the arena. Fans were coupled to two flexible air ducts (9 cm diameter), which, in turn, ran through wall vents to draw fresh air from outside. Air passed in through these ducts, into the pressure box and then exited into the arena. To reduce turbulence, air entering the testing arena passed through an egg crate, plastic lined with organdy fabric. The wind tunnel did not provide birds with a discrete plume of odour that they could follow to a source, rather birds were exposed to the presence or absence of the odour in the test arena.

We tested 23 common diving petrel chicks for this experiment. Chicks were tested and handled one at a time. Prior to an experiment, a test subject was removed from its nest chamber located within its burrow. The chick was placed in a clean cotton bird bag and carried to the hut (10–100 m from the nest). Once inside, the chick was placed on a pre-defined start point within the wind tunnel. The start position was c. 40 cm downwind from a glass dish (radius 71 mm, height 41 mm). The glass dish held one of two test stimuli: soil (15 g) from the chick’s burrow (hereafter called nest soil), or soil (15 g) from the colony in the region where the burrow was located (colony soil). Colony soil was collected at least 10 m from any known burrow entrance and had the same look and consistency as nest soil. The third test stimulus was an empty glass dish (control). Each chick was exposed to all three stimuli. Although no order effects were ultimately observed, to control for this possibility, we balanced the order of presentation for all three stimuli such that all six possible presentation orders were equally distributed amongst the chicks. To reduce the likelihood that chicks were able to detect an odour signature from a previous trial, the wind tunnel was washed with 75% methanol between trials.

Chicks were tested one at a time in dim lighting and videotaped from above using a Sony camcorder (model DCR-TRV30) under infrared illumination. At the start of each trial, chicks were allowed to acclimate to the testing arena for 30 sec. Following acclimation, we removed the lid of the dish and recorded the chicks’ behaviours for 3 min. This duration was chosen to minimize the amount of time chicks were out of the nest, and prior work on blue petrels (Halobaena caerulea (Gmelin, 1789)) suggested that activity was greatest within this time interval (Cunningham et al. 2006). After the first 3 min trial, we introduced the next dish, and repositioned the chick at the start position. We repeated this procedure for the third stimulus. Once the experiment was completed, the chick was returned to its burrow. Chicks were away from their burrows for less than...
35 min. Video images were projected using a Sony Trinitron colour video monitor (model PVM-1351Q) and analysed by a single ‘blind’ observer in real time using Jwatcher software (http://www.jwatcher.ucla.edu/).

**Statistical analyses**

Statistical analyses were performed using IBM SPSS Statistics (v.19) software (http://www-01.ibm.com/software/analytics/spss/). Data were analysed for differences in body turns (a turn of the body 45° to the right or left) and distances travelled (quantified by recording the chick’s position at 15 sec intervals and then adding these distances together on a per minute basis) in response to the three stimuli. Because activity tended to vary over the time course of the experiment for this species, data were analysed in 1 min time bins. Prior to analysis, data were first normalized using a square root transformation. To determine if there were overall differences in the chick’s responses to the three stimuli, we used a repeated-measures ANOVA on the transformed data, employing the Greenhouse-Geisser correction if the assumption of sphericity was violated. Where a significant difference was found, we used a Tukey’s test on the transformed data to make pair-wise comparisons (Zar 1996).

**Results**

Once chicks adjusted to the wind tunnel, we found consistent, significant differences in behaviour in response to odour stimuli, both in terms of distances travelled and turning behaviour.

**Distance travelled**

In minute 1, chicks walked statistically similar distances in response to the three stimuli (Fig. 1; repeated measures ANOVA: df = 2, F = 2.14, P = 0.13). However, in minute 2, mean responses to the three stimuli were significantly different (Greenhouse-Geisser: df = 1.53, F = 3.94, P = 0.04). Similar to our prediction, chicks walked significantly less distance when they were presented with their own nest soil compared to the control (Tukey’s test: q = 3.44, P = 0.05). Mean distances walked were not significantly different between colony soil and control stimuli (q = 2.09, P > 0.20), or between nest soil and colony soil stimuli (q = 1.35, P > 0.50). In minute 3, there were no significant differences in the distance walked in response to the three stimuli (df = 2, F = 2.58, P = 0.09). This was probably due to reduced overall activity in minute 3.

**Turning behaviour**

We observed similar patterns for body turns (Fig. 2). During minute 1, there were no significant differences in turning behaviour among the three groups (df = 2, F = 1.71, P = 0.32). However, during minute 2, chicks showed significant differences in response to the three stimuli (Greenhouse-Geisser: df = 2, F = 4.85, P = 0.01). As predicted, nest soils evoked less turning than control stimuli (q = 4.40, P = 0.01). However, turning behaviour was similar when comparing responses to either colony soil and control (q = 2.24, P > 0.20), or nest soil and colony soil (q = 2.17, P > 0.20). Chicks responded similarly to the three stimuli during minute 3 (df = 2, F = 0.89, P = 0.42).

**Discussion**

Consistent with our prediction, our data suggest that common diving petrel chicks can detect odours associated with their nest material. We saw the largest effects during minute 2, probably because chicks took longer to acclimate to the wind tunnel than we anticipated from prior work with blue petrels in response to food odours (Cunningham et al. 2006). Chicks responded to burrow scents essentially by settling down and not moving. Given that chicks do not tend to move within their burrows unless an adult is present, our interpretation is that the scent of the burrow is a sufficient stimulus to trigger a “sit-still response”.

Staying still while in the nest chamber is probably adaptive for survival in colonies where chicks are at high risk of predation. Unlike other sympatric species such as blue petrels that dig complex burrows ~2 m deep, common diving petrels dig short (< 1 m), simple tunnels that lead directly to the nesting chamber. Common diving petrel chicks are also smaller than most petrel chicks (fledging...
weight of common diving petrel: 115 g, fledging weight of blue petrel: 208 g, Jouventin et al. 1985) and particularly vulnerable to predation by skuas (Catharacta skua lombergi Mathews, 1912, reviewed by Warham 1996). Chicks do not emerge from their burrows until they fledge (Warham 1990). Thus, it is not surprising that chicks tended to sit relatively still when exposed to the odours associated with the nest, since this is consistent with their normal behaviour. We did not test whether the “sit-still response” is a general response to petrel odour or more specific to the scent of an individual’s burrow, but this question should be tested.

Chicks turned more and walked greater distances during the control presentations (Figs 1 & 2). Our interpretation is that, in the absence of familiar burrow odours, chicks responded by moving around in the arena, perhaps searching for a cue associated with the nest. Chicks also tended to walk less when presented with colony soil than to control conditions. We did not analyse scented compounds in colony soil as part of this preliminary study, however, nitrogenous wastes have been shown by others to be present in petrel colonies (Erskine et al. 1998). We have recently shown that blue petrel chicks are sensitive to ammonia at picomolar concentrations suggesting a potential source of a familiar odour (Nevitt et al. 2006). Petrel derived odours may also be present in colony soil, but this has not yet been demonstrated.

Given that chicks generally decreased both of their behaviours over the duration of the experiment, an alternate explanation for our findings is that chicks were stressed by the initial disturbance of handling and responded to this by turning more and walking further distances. Over time, particularly in the presence of the familiar burrow scent, chicks became less affected by the initial disturbance. This interpretation is interesting because it suggests that a familiar odour may act to mitigate the fear response in diving petrels as it does in chickens (Gallus domesticus L., 1758, Jones & Gentle 1985). Although our experiment was not originally designed to address this possibility, it is an exciting interpretation that warrants further study in the future.

In conclusion, we previously showed that adult common diving petrels can identify their burrows by odour cues alone (Bonadonna et al. 2003). Our current findings expand on these results by suggesting that common diving petrel chicks can detect and respond appropriately to the scent of their nest material in a simple wind tunnel. The implications of this initial study are that, despite having smaller olfactory bulbs than other procellariiforms, diving petrels respond to odour cues associated with their burrows, and this behaviour is already expressed in chicks before they leave the nest. Given that diving petrels do not leave the nest prior to fledging, our study suggests that olfaction may be involved in the development of individual recognition, as it appears to be in Leach’s storm-petrels (Oceanodroma leucorhoa Vieillot, 1818, O’Dwyer et al. 2008). Our results further suggest that a simple wind tunnel can be used to examine olfactory specific behaviours without requiring birds to be motivated to make a choice between two alternatives. Thus, this methodology provides a useful alternative to Y-maze testing in the field.

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