ORIGINAL ARTICLE

A seabird's eye view: visual felds of some seabirds (Laridae and Procellariidae) from tropical latitudes

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Received: 6 May 2024 / Revised: 5 July 2024 / Accepted: 8 July 2024 © The Author(s) 2024

Abstract

The visual feld of a bird defnes the amount of information that can be extracted from the environment around it, using the eyes. Previous visual feld research has left large phylogenetic gaps, where tropical bird species have been comparatively understudied. Using the ophthalmoscopic technique, we measured the visual felds of seven tropical seabird species, to understand what are the primary determinants of their visual felds. The visual feld topographies of the seven seabird species were relatively similar, despite the two groups of Terns (Laridae) and Shearwaters (Procellariidae) being phylogenetically distant. We propose this similarity is due to their largely similar foraging ecology. These fndings support previous research that foraging ecology rather than relatedness is the key determining factor behind a bird's visual feld topography. Some bird species were identifed to have more limited binocular felds, such as Brown Noddies (*Anous stolidus*) where binocularity onsets lower down within the visual feld, resulting in a larger blind area about the head.

Keywords Binocularity · Foraging ecology · Ophthalmoscope · Seychelles

Introduction

Diferences in the optical structure, size, and movement of eyes infuence the quality of images formed, and the extent of the world seen from moment to moment (Martin [2007,](#page-5-0) [2012](#page-5-1)). The visual feld of an eye, and how the felds of the two eyes are combined, defne the extent of the world from which an animal can extract information at any instant, and

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is regarded as a key component in describing the sensory ecology of an animal (Martin [1994\)](#page-5-2). The visual feld of an animal has three main components: the monocular feld describes the extent of the world seen by each eye, the binocular feld describes the region where the monocular felds overlap, and the blind areas describe the region in which no vision is provided (Waldvogel [1990\)](#page-6-0). The cyclopean feld describes the total region about the head from which information can be extracted at any one moment, by either monocular or binocular vision (Fernández-Juricic et al. [2004](#page-5-3); Martin [1994\)](#page-5-2).

Among birds, the topography of the binocular feld (the size, shape, and position of the region of binocular overlap) shows considerable interspecifc variation (Burton [2008](#page-5-4); Martin [2017](#page-5-5)). These variations are dependent on the extent of the monocular feld of each eye and the positioning of the eyes in the skull (Martin [2017](#page-5-5)). In general, eyes that are placed more laterally in the skull provide smaller binocular felds but larger cyclopean felds. It has been hypothesized that interspecifc diferences in binocular feld topography among birds are primarily infuenced by species-specifc sensory requirements of foraging, rather than factors such as shared ancestry or the guidance of locomotion (Cantlay et al. [2019](#page-5-6), [2023](#page-5-7); Martin [1999](#page-5-8); Potier et al. [2018\)](#page-6-1).

Visual felds have been determined in over 180 bird species (Martin [2017](#page-5-5)), including 14 species of seabirds: two species of penguins (Spheniscidae), two species of Albatrosses (Diomedeidae), three species of petrels and shearwaters (Procellariidae), one species of cormorant (Phalacrocoracidae), and two species of auks (Alcidae). In this study, we determined the visual feld in a further seven species of seabirds which inhabit tropical regions (between latitudes 23.5° north and south): fve species of terns (Laridae), and two species of shearwaters (Procellariidae). These two taxa are distant from each other in the phylogenetic tree, but share a generally similar habitat and foraging technique, and we use data on their visual felds to test further the hypothesis that foraging technique, rather than phylogenetic relatedness, is the primary infuence on visual feld topography in birds (sensu Cantlay et al. [2023](#page-5-7)).

Materials and methods

Visual felds were measured in seven species of tropical seabirds from two avian families, Laridae (Charadriiformes) and Procellariidae (Procellariiformes) (Table [1](#page-1-0)). All species are described as foraging by dipping or foraging by plunging (Tobias et al. [2022](#page-6-2)) (Supplementary Information Table S2). Dipping is where food items are taken from the surface of the water or just below while plunging (or plunge diving) is where the bird enters the water to chase or retrieve prey underneath the water's surface (Dufy [1983](#page-5-9); Haney and Stone [1988](#page-5-10)). The estimated Laridae and Procellariidae pairwise divergence was 71 million years ago, and these families are not considered close relatives (Slack et al. [2006](#page-6-3)).

Inca Terns were studied at Birdworld, Farnham, Surrey, UK (51.813,−0.840°) in May and July 2023, and the other species were studied at Aride Island Nature Reserve, Seychelles (−4.214, 55.668°) in June 2023. All Inca Terns were adults that had been held in captivity for several years. Individual birds were caught in their holding enclosure using hand nets and placed in a cloth bag and carried to a nearby building where measurements were made. On completion of measurements, each bird was released immediately into its enclosure. Of the six species studied on Aride Island the Sooty Terns and Brown Noddies were caught using hand nets, Tropical Shearwaters, and Wedge-tailed Shearwaters were caught by hand, and White Terns and Lesser Noddies were caught with mist nets. All birds were adults. In both locations, measurements of visual felds were conducted in a darkened room, and on the completion of measurements, each bird was released within 65 min near to its capture location. Birds were walked from the site of capture to the darkened room for a maximum of 10 min.

Visual feld measurements

The ophthalmoscopic refex technique was used to measure the visual feld characteristics, following the standard procedure described in previous studies (Martin et al. [2007](#page-5-11); Martin and Wanless [2015](#page-5-12); Cantlay et al. [2020\)](#page-5-13). Each bird was held with its body immobilised in a foam rubber cradle and its bill placed in a holder specially designed for each species, with the head of the bird adopting its natural resting position. This arrangement fxed the at-rest head position with respect to the co-ordinate system used to characterise the visual feld (Martin et al. [2007\)](#page-5-11). This technique has been consistently applied across a wide taxonomy of avian species and provides a reliable method for interspecifc comparisons of visual feld topography (Martin [2017](#page-5-5)). The UK Animals (Scientifc Procedures) Act 1986 is not applicable due to the procedure being non-invasive and the short period of time (generally 30 min) for bird restraint (Martin and Portugal [2011](#page-5-14)). Spontaneous eye movements were observed in some of the species. These refer to the observation that some bird species have complex rotational eye movements, and the translational efect of these movements can alter the limits of the visual feld recorded at each elevation (Martin et al. [2008](#page-5-15)). Visual feld measurements were taken for the positions that the eyes spontaneously adopted when fully rotated forward, hence converged for the front feld, and provides an estimate of maximum binocular feld width (Potier et al. [2018\)](#page-6-1). In each species, detailed measurements were made only throughout the anterior portion of the visual feld with a single set of measurements made at−90° (directly behind the head; see Fig. [1](#page-2-0) for the explanation of co-ordinate system) to determine the width of the blind area and thus allow

Fig. 1 The mean angular separation of the retinal feld margins in the anterior portion of the visual feld as a function of elevation in the median sagittal plane of the head. Positive values indicate the width of the binocular feld, negative values indicate the width of the blind area. The coordinate system is such that the horizontal plane is defned by the elevations−90° (behind the head) and $+90^\circ$ (in front of the head), and 0° is directly above the head. The drawing shows a bird's head in profle with key coordinates indicated and the visual projection of the eye–bill tip axis. The head position shown is approximately that spontaneously adopted by an Inca Tern when held in the hand and indicates the head position at which visual feld parameters were measured. The species studied were; *Anous stolidus* (Brown Noddies)*, Anous tenuirostris* (Lesser Noddies), *Gygis alba* (White Terns), *Onychoprion fuscatus* (Sooty Terns), *Larosterna inca* (Inca Terns), *Ardenna pacifca* (Wedge-tailed Shearwaters), and *Pufnus bailloni* (Tropical Shearwaters). An alternate black and white version of Fig. 1 can be found in the supplementary information

characterisation of the monocular felds in the horizontal plane. Mean visual feld data for each species were determined and used to create vertical (Fig. [2a](#page-3-0), left panel) and horizontal (Fig. [2](#page-3-0)b, centre panel) sections through the visual felds and topographical maps of the anterior felds (Fig. [2](#page-3-0)c, right panel) for each species.

The small number of species measured in this study precluded a full phylogenetic generalised least squares analysis of the kind employed in the investigation of the relationship between visual felds and foraging in Anatidae (Cantlay, et al. [2023](#page-5-7)) and Strigidae (Potier et al. [2023](#page-6-4)), and thus our results are descriptive.

Fig. 2 Visual felds of seven tropical seabird species. **A Left panel** vertical sections of the binocular feld of each species in the median sagittal plane of the head. **B Middle panel** horizontal sections of the visual feld of each species, the position of the horizontal plane is indicated by the horizontal lines shown in the vertical sections and in the schematic head and co-ordinate diagram shown in Fig. [1](#page-2-0). **C Right panel** orthographic projection of the visual felds of each species. These fgures use the conventional latitude and longitude coor-

Results

General features of the visual felds

The general topography of the binocular visual feld of all seven species is relatively similar (Figs. [1](#page-2-0) and [2\)](#page-3-0). All species have a vertically long and narrow binocular feld, with the visual projection of the eye bill-tip occurring within the ventral portion of the binocular feld below the point of its maximum width. All species have a blind area which starts in the dorsal quadrant of the frontal feld and extends above and behind the head down to the horizontal.

Interspecifc diferences in visual felds

Within these shared general features of the visual felds, there are clear interspecifc diferences: (1) the binocular region, from the elevation at which it could be measured below the bill, extends vertically through between 90° (White Terns) and 125° (Inca Terns), with a mean vertical extent of 104° (mean of all species), (2) the maximum width of the binocular felds difered between 15° (Brown Noddy) and 32° (Inca Tern) and

dinate system where the equator is vertically aligned with the head's median sagittal plane, the grid is at 20° intervals. It should be imagined that the birds' head is placed at the centre of the sphere with the visual feld regions projected outwards onto the sphere's surface. Colours are used to indicate monocular (orange), the binocular (green), and blind (blue) portions of the visual felds. The white and black triangles indicates the directions of the eye bill-tip projections. See Table [1](#page-1-0) for species Latin names

the mean maximum width across all species equalled 26°, (3) the maximum width of the binocular region projects above the horizontal and lies at a mean value of 17° above the direction of the eye bill-tip projections, (4) laterally, there are extensive regions on monocular vision ranging from 134° (Inca Tern) to 149° (Brown Noddy) with a mean width across all species of 140°, and (5) in all species there is a blind region that projects into the dorsal anterior sector of the visual feld, it starts at an elevation of 55° in Fairy Terns and 25° in Inca Terns and has a mean width across all species of approximately 35° directly above the head, (6) the width of the blind sector directly behind the head (elevation−90°) varies between 49° (Wedge-tailed Shearwater) and 61° (Inca Tern) with a mean width across all species of approximately 53°.

Discussion

General characteristics of the visual felds

The visual felds of the seven seabird species show the characteristics of birds which rely on visual cues to guide the accurate placement and accurate timing of arrival of their bill at a target (Martin [2009](#page-5-16)). These visual field characteristics are driven particularly by the visual tasks associated with foraging for discrete targets, as opposed to using tactile cues or flter-feeding techniques. The key visual feld characteristics that underlie this interpretation of these birds' visual feld characteristics are the vertically long and relatively narrow binocular feld within which the visual projection of the bill tip is positioned below the region of maximum binocular feld width (Fig. [2\)](#page-3-0). It has been argued (Martin [2009](#page-5-16)) that this confguration serves to provide optic fow-feld information that is necessary for both accurate placement of the bill with respect to a target and for accurate estimation of time to contact the target, which is necessary for precise timing of bill opening when seizing prey, along with playing a pivotal role in obstacle avoidance (Bhagavatula et al. [2011](#page-5-17)). These kinds of visually guided tasks are practiced by all species in the present sample regardless of whether they forage using dipping for items at the sea surface or plunging for items taken at or below the sea surface. Previous studies on Procellarids documented binocular feld widths of 30°, 27°, and 32° for White-chinned Petrels (*Procellaria aequinoctialis*), Grey-headed (*Diomedia chrysostoma*) and Black-browed Albatrosses (*D. melanophris*), respectively (Martin [1998;](#page-5-18) Martin and Prince [2001](#page-5-19)), compared to the 24° and 28° binocular width in Tropical and Wedge-tailed Shearwaters found in the present study.

This similarity in visual feld confguration between the Laridae and Procellariidae species in this sample is noteworthy because these taxa are not considered to be closely related, their lineages having diverged about 71 million mya (Slack et al. [2006](#page-6-3)). It reinforces the idea that, in birds in general, visual feld characteristics are driven primarily by the visual demands of foraging rather than phylogeny. This accords with similar conclusions drawn from studies of visual felds of a larger sample of species using phylogenetic generalised least squares analysis in ducks, geese, and swans (Anseriformes) (Cantlay et al. [2023\)](#page-5-7) and in owls (Strigiformes) (Potier et al. [2023\)](#page-6-4).

Interspecifc diferences in visual felds

While there is a similarity in the overall characteristics of the visual felds of this sample of seabirds, there are also clear diferences in the dimensions of specifc aspects of their visual felds. These include diferences in the maximum width and vertical extent of binocular felds, and positions of the binocular feld relative to the eye bill-tip projection. These diferences are relatively subtle compared with the wide range of visual feld confgurations recorded in birds (Martin [2017\)](#page-5-5). However, it has been shown that relatively small diferences in visual feld topography among closely related birds can be accounted for by consideration of diferences in foraging behaviour and diet, and the nature of the associated visual challenges of diferent foraging task. This has been shown in comparisons between visual felds and foraging behaviour among species of ibis (Threskiornithidae) (Martin and Portugal [2011\)](#page-5-14), ducks (Anatidae) (Guillemain et al. [2002](#page-5-20); Martin et al. [2007](#page-5-11)), auks (Alcidae) (Martin and Wanless [2015\)](#page-5-12), plovers (Charadriidae) (Martin and Piersma [2009;](#page-5-21) Cantlay et al. [2019](#page-5-6)), vultures and hawks (Accipitridae) (Portugal et al. [2017](#page-5-22); [2023\)](#page-6-5), and petrels (Procellariidae) (Martin and Prince [2001](#page-5-19)).

These studies suggest that visual felds can be fne-tuned to specifc aspects of the foraging ecology of species within the broad parameters required for successful visually guided foraging. The particular ways in which the interspecifc differences in the visual felds of the shearwaters and terns described here can be interpreted with respect to diferent foraging tasks require a detailed analysis of these species' foraging ecology and behaviours. For example, diferences between the visual fields of Inca and Noddy Terns are likely to result from diferences in the tasks of detecting and acquiring their preferred prey types. Brown Noddies are described as feeding, "mainly by hover-dipping and contact-dipping; regularly food patters at surface… usually does not plunge-dive. Captures fying-fsh in air… Forages on moonlit nights" (Gochfeld and Burger [1996\)](#page-5-23). Inca Terns are described as feeding "mainly on small anchoveta (*Engraulis ringens*), also planktonic crustaceans; offal and scraps... Forages mainly by plunge-diving, and contact – and surfacedipping; scavenges for scraps left by sea lions and avian predators" (Gochfeld and Burger [1996](#page-5-23)). Thus, Noddy Terns feed mainly in the air, but they have the exacting perceptual demand associated with taking flying prey (flying fish, Exocoetidae**)** whose appearance is intermittent and brief. On the other hand, Inca Terns feed below the water surface on small but less evasive prey (planktonic crustaceans) (Dufy [1983](#page-5-9); Hanley and Stone 1988; Gochfeld and Burger [1996](#page-5-23)). This means that although both species have to achieve accurate location and timing of prey capture for successful foraging their tasks are quite diferent and how these diferent tasks will have driven the fne-tuning and diference between their visual felds would require further detailed analysis of their foraging tasks. Amphibious vision has particular demands compared to foraging in the air due to the loss of corneal power and the narrowing of the visual felds on entering the water (Katzir and Howland 2003), and the latter may be sufficient to account for the broader binocular field in Inca Terns compared with those in Brown Noddy (Fig. [2](#page-3-0)). To be certain of this requires more detailed knowledge of the optical structure of the eyes in these species, but entering water can result in the reduction of maximum binocular feld width by approximately 50% (Martin and Young [1984\)](#page-5-24). Thus, it would be predicted that upon immersion the maximum binocular feld of Inca Terns (34°) would narrow to about 17°,

which is similar to the maximum width of the Brown Noddy 15° in air, with the results that the binocular felds of both species are of similar width when the birds are engaged in their primary foraging tasks. This is consistent with the fact that the three species that forage using dive-plunging have the longest vertical extension of binocular vision (Inca terns, 125°; Wedge-tailed shearwaters, 115°; Tropical shearwaters, 100°) compared to those that do not dive. The two species (Lesser Noddy, Inca Tern) not reported to forage at night or crepuscular hours do not appear to show any particularity in their visual felds. Clearly, a more detailed analysis of the foraging ecology and their associated perceptual challenges across all of the species (e.g., Regular et al. [2011\)](#page-6-6) are required to determine factors which may have led to the recorded diferences in their visual felds.

Supplementary Information The online version contains supplementary material available at<https://doi.org/10.1007/s00114-024-01926-4>.

Acknowledgements We would like to thank Sean Gibson for the manufacture of the bill holders. We are also thankful to all the staf and volunteers of Aride Island Nature Reserve, who provided useful information, boat transportation and logistical support during our feld work, and to the Island Conservation Society Head Office for helping to organise our visit to Aride. We thank Polly Bramham and staff at Birdworld, Surrey. We are grateful to Hana Merchant for useful discussions.

Author contribution Conceptualization, E.A.L and S.J.P.; methodology, E.A.L., G.R.M., G.R., and S.J.P; data collection, E.A.L., G.R., and S.J.P.; formal analysis, E.A.L.; interpretation, E.A.L., G.R.M., and S.J.P.; visualisation, E.A.L. and G.R.M.; resources, G.R. and S.J.P.; writing—original draft, E.A.L. and S.J.P.; writing—review and editing, E.A.L., G.R.M., G.R., and S.J.P.

Data availability All data are available in the Supplementary information.

Declarations

Conflict of interest The authors declare no competing interests.

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