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High definition video loggers provide new insights into behaviour, physiology, and the oceanic habitat of marine top predators

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animal-borne video loggers, marine top-predators, diving behaviour, benthic habitat, predator-prey interactions, penguins

Summary Statement

The use of full HD animal-borne video loggers can provide information about a wide range of behavioural, physiological and environmental aspects of marine top-predators' biology.

18 Abstract

- 19 1. Camera loggers are increasingly used to examine behavioural aspects of free-ranging
20 animals. However, often video loggers are deployed with a focus on specific behavioural
21 traits utilizing small cameras with a limited field of view, poor light performance and video
22 quality. Yet rapid developments in consumer electronics provide new devices with much
23 improved visual data allowing a wider scope for studies employing this novel methodology.
- 24 2. We developed a camera logger that records full HD video through a wide-angle lens,
25 providing high resolution footage with a greater field of view than other camera loggers.
26 Main goal was the analysis of foraging behaviour of a marine top-predator, the Yellow-eyed
27 penguin in New Zealand, in the context habitat characteristics. Frame-by-frame analysis
28 allowed accurate timing of prey pursuits and time spent over certain seafloor types. Similarly,
29 it was possible to time breathing intervals between dives and quantify exhalation events
30 during prey events, a previously undescribed behaviour. Screen overlays facilitate analysis of
31 flipper angles and beat frequencies throughout various stages of the dive cycle.
- 32 3. The recorded video footage showed that prey species were associated with certain seafloor
33 types, revealed different predator evasion strategies by benthic fishes, and highlighted
34 varying energetic consequences for penguins pursuing certain types of prey. Flipper
35 movement analysis confirmed decreasing effort during descent phases as the bird gained
36 depth, and that ascent was principally passive. Breathing episodes between dives were short
37 (<1 s) while the majority of the time was devoted to subsurface scanning with a submerged
38 head.
- 39 4. Video data recorded on free-ranging animals not only provides a wealth of information
40 recorded from a single deployment but also necessitates new approaches with regards to
41 analysis of visual data. Here, we demonstrate the diversity of information that can be
42 gleaned from video logger data, if devices with high video resolution and wide field of view
43 are utilized.

44 Introduction

45 Examining the at-sea behaviour of marine animals has long been a challenging endeavour. Direct
46 visual observations of behaviour are almost impossible, especially when most of it happens under
47 the ocean's surface. In recent decades, advances in telemetry technologies and the emergence of
48 bio-logging hardware have provided the means to track marine animals and reveal their foraging
49 behaviour in great detail. Starting in the 1970s with rather crude location estimates and limited data
50 quality recorded by unwieldy devices that could only be used on large animals, advancements in
51 micro-electronics have resulted in ever smaller and more accurate loggers to pinpoint an animal's
52 position to within a few metres and record their diving depths with oceanography-grade precision
53 (Wilmers *et al.* 2015). New technologies such as accelerometers and gyroscopes further refined
54 methods to study marine habitat use (e.g. Noda *et al.* 2014). Yet, placing dive metrics into a complex
55 behavioural and environmental context can be difficult; ideally a reference framework based on
56 direct observations is used to match up dive metrics and actual behaviours (e.g. Moreau *et al.* 2009;
57 Volpov *et al.* 2016). So the original dilemma of having to make direct observations of marine animal
58 behaviours still persists. Animal-borne video recorders offer the means to overcome this problem.

59 In recent years animal-borne camera systems have made it possible to log in situ observations of
60 behaviour from the animal's point of view (Moll *et al.* 2007). For example, deployment of light-
61 weight video cameras on flying birds provided new perspectives on prey pursuit in falcons (Kane &
62 Zamani 2014) and revealed how albatrosses use the presence of killer whales to locate prey
63 (Sakamoto *et al.* 2009). No other animal group has been more subject to deployment of video
64 recording devices in recent years than marine animals. By overcoming the observational barrier at
65 sea, video loggers are providing copious amounts of novel data that range from identification of
66 feeding strategies (Takahashi *et al.* 2008) and previously unknown food sources (Thiebot *et al.*, in
67 review), to social interactions such as group foraging (Sutton *et al.* 2015) or kleptoparasitism
68 (Handley & Pistorius 2015). Video data also offers the means to calibrate other bio-logging data
69 (Watanabe & Takahashi 2013; Gómez-Laich *et al.* 2015).

What most of these studies have in common is their focus on specific behavioural traits while providing limited information about the environment the behaviours occurred in. This is principally due to limitations of the video hardware used, which has to be small and light-weight so as to not overly impede the study animal's movement capabilities (Ludynia *et al.* 2012) and hence behaviour. As a result, video quality (i.e. image resolution, field of view/FOV) is sacrificed in favour of smaller cameras. However, with the rise in popularity of action cams on the consumer market, new video devices have recently become available with high definition video capabilities and wide-angle optics, suitable for deployment even on smaller marine animals such as penguins. This leap in quality has significant implications for the study of marine animals as it not only allows monitoring of wide-ranging aspects of behaviour, but also provides new opportunities for the visual analysis of the environment the animals use. This is particularly relevant in species that forage at the seafloor where video data can provide extensive information about the benthic habitat (Watanuki *et al.* 2008).

The yellow-eyed penguin (*Megadyptes antipodes*) in New Zealand is known to be a benthic forager (Mattern *et al.* 2007) that feeds primarily on demersal fish species (van Heezik & Davis 1990; Moore *et al.* 1995). It has been suggested that this strategy might come at the expense of reduced behavioural flexibility, with subsequent vulnerability to changes in the marine environment (Mattern *et al.* 2007). In particular, degradation of seafloor ecosystems in the wake of commercial bottom fisheries are suspected to influence yellow-eyed penguin foraging success and population developments (Browne *et al.* 2011; Mattern *et al.* 2013). While the species' at-sea movement and diving behaviour has been subject to a number of studies in the past decades (Moore *et al.* 1995; Mattern *et al.* 2007, 2013), information about their benthic habitat is scarce.

To be able to assess the extent to which penguin behaviour and foraging success correlates with the composition of the benthic habitat, we developed a camera logger that records full HD videos through wide-angle lenses. The main focus of our study was to assess the suitability of the device for the visual analysis of penguin prey pursuit behaviour and characteristics of the benthic ecosystem. However, the deployment revealed far more information than was anticipated. The video data

provided novel insights into physiological aspects of the penguin's diving activities and allowed us to draw conclusions about prey capture techniques. In this paper, we summarise our findings, demonstrate analytical approaches to evaluate animal-borne video data, and highlight the multi-disciplinary potential of full HD video loggers.

Materials and methods

Study site and species

The Yellow-eyed penguin, classified as "Endangered" by the IUCN Redlist (BirdLife International 2012), is one of five penguin species endemic to the New Zealand region and occurs on the sub-Antarctic Auckland and Campbell Islands as well as the south-eastern coastlines of New Zealand's South Island and Stewart Island (Seddon *et al.* 2013). This study was carried out at the Boulder Beach complex, Otago Peninsula, South Island, New Zealand (45.90°S, 170.56°E). Penguins from this site have been subject to foraging studies in the past decade that have suggested substantial impact of bottom trawling activities on the yellow-eyed penguins' at-sea movements (Mattern *et al.* 2013).

The research was approved by the University of Otago's Animal Ethics Committee (#11/14) and was permitted by the NZ Department of Conservation (45799-FAU).

Video logger & deployment

We developed a high-definition video logger (dimensions LxWxH, 80x40x23mm; weight: 48g) which is combined with a time-depth recorder (TDR, 12x31x11mm, 6.5g; AXY-depth, Technosmart Ltd. Italy) and a GPS logger (modified i-gotU, GT-120, Mobile Action Technology Inc., Taiwan, 22x31x11mm, 12g). The latter two devices were combined into a single unit by gluing the AXY-depth to the longer side of the GPS device. Camera and logger combination were then attached individually in line to the lower back of the penguin using adhesive tape (Wilson *et al.* 1997). Additional drag of the devices was principally limited to the cameras frontal area (Bannasch *et al.* 1994). The camera logger consists of a modified Mobius action-cam with a 130° wide-angle lens (www.mobius-actioncam.com). In order to achieve the smallest and lightest device possible, the camera electronics, video sensor and

lens were removed from the casing and the battery replaced with a 1200 mAh Lithium Polymer battery to extend recording time. A small bespoke timer board was developed to allow the camera to be fired at a pre-determined time. The recording then ran until the battery fell below the minimum operating voltage of the camera (ca. 2-4 hours). Connections were provided to allow programming the alarm and also to access the camera's USB port for managing camera setting, extracting the video data and recharging the battery. The board was isolated electrically to prevent the contacts from shorting as sea-water is conductive. Activation of the interface was achieved using a Hall-effect device. An Arduino-based interface was developed to allow the current date/time and alarm time to be set. The camera was programmed to record video data at a resolution of 1920x1080 pixels (1080p) at a frame rate of 30 frames per second. Video data was recorded in H.264 MPEG4 format and stored on a 32GB MicroSD card. The camera was encased in epoxy resin to prevent water damage and provide the device with a hydrodynamic shape to reduce additional drag (Culik *et al.* 1994). After device recovery, data was downloaded through the camera's USB interface.

Since the logger stores video data as a series of full frame images ('progressive scan'), it was possible to conduct a frame-by-frame analysis to accurately time components of the bird's behaviour – i.e. breathing intervals, flipper beat frequencies and amplitudes – as well as time spent over certain benthic habitats. Video analysis was conducted in professional editing software (Adobe Premiere Pro CS 6, Adobe Systems Inc., San Jose, CA, USA) which allows video scrubbing and provides the option to display frame number in the preview timer.

The video logger was deployed on a breeding male Yellow-eyed penguin tending two chicks on 17 December 2015. The bird left on a single foraging trip on 18 December before the device was recovered on 19 December.

Analysis of behaviours

Prey pursuits. We defined the beginning of a prey pursuit as the moment when the penguin markedly accelerated while swimming along the seafloor; the end was reached when the penguin

decelerated again to its previous cruise speed (if no prey was caught), or when the prey item was swallowed completely. Acceleration and deceleration were associated with temporary blurring of the video footage due to irregular body movement, allowing for exact timing of prey pursuits. Where possible, prey species were identified from frames providing a clear view of the prey item.

Beyond prey interactions, the video data offered the opportunities to analyse physiological aspects of the penguin's behaviour.

Surface breathing. We timed breathing events when the penguin was at the surface following a dive. Noting frame numbers when the bird raised its head out of the water before lowering below the surface again made it possible to determine time the penguin was able to respire (<https://vimeo.com/179414575#t=145>).

Flipper movements. During dives, flipper beat frequencies (beats per minute, BPM) were determined by counting the number of frames required to complete one flipper beat cycle, beginning the count when the flipper angle reached its maximum upward inclination and ending with the frame prior to the subsequent maximum upward inclination. In the video editing software we overlaid a template indicating 10, 30, 50, 70 and 90 degree angles radiating from the base of the flippers on the video data (<https://vimeo.com/179414575>). This allowed us to visually determine maximum amplitude of each flipper beat to the nearest 5°.

Analysis of benthic habitat

For all dives, the benthic habitat was classified according to sediment type (fine sand, coarse sand with shell fish fragments, gravel), sediment structure (flat, sediment ripples) and composition of the epibenthic communities. For the latter, we used a presence/absence approach in which the occurrence of brittlestars (*Ophiuroidea*), anthozoans (anemones and soft corals), and horse mussels (*Atrina zelandica*) within a 30-frames time window. Future deployments with a functional GPS logger can be used for more elaborate analysis of the benthic habitat, e.g. the creation of biodiversity indices.

A selection of edited video clips demonstrating the various behaviours and habitat types described above can be accessed via <https://vimeo.com/album/4103142>.

Dive data analysis

Dive data recorded by the TDR was analysed following methods described in detail in Mattern et al. (2007). Key dive parameters determined were maximum depth reached, duration of the dive event and its three main phases (i.e. descent, bottom phase, ascent) as well as vertical velocities during descent and ascent. Dives were classified as pelagic or benthic dives using dive profile characteristics, where near horizontal bottom phases with little vertical variance as well as consistent maximum dive depths on consecutive dives were used as cues for diving along the seafloor. This approach was validated by recorded video data.

Statistical analysis was carried out in R (R Development Core Team 2008).

Results

Foraging trip length, diving events and video coverage.

The day following camera deployment, the penguin performed a 10.7 hour-long foraging trip. The first dive event was recorded at 5:30 hrs and the last event concluded at 16:10 hrs. The bird performed 286 dives of which 159 dive profiles matched the criteria for benthic dives (Figure 1). Median dive depth reached during benthic dives was 54.4m (range: 4.8-62.1 m, n=159) whereas the majority of pelagic dives occurred in the upper 10m of the water column (median: 7.8m, range: 0.5-31.7 m, n=127): camera footage confirmed these to be principally travelling behaviour (<https://vimeo.com/179414642>). For the first 3½ hours of the foraging trip (05:30-09:00 hrs) the bird performed mainly pelagic dives, indicating primarily travelling behaviour towards its main foraging grounds; the remaining hours (09:00-16:10 hrs) the bird principally devoted its time to benthic diving (Figure 1). The camera operated continuously from 11:00:22 hrs to 13:01:43 hrs. Due to occasional frame loss when data were written to memory, total length of the recorded footage amounted to 2

hours 8 seconds). 46 complete dives were video recorded which corresponds to 16% of all dive events; of these 32 dives were benthic dives. However, dives were longer during the middle of the day so that camera footage covered 25% of the trip's cumulative dive time.

Prey pursuits & capture

A total of 20 prey pursuits was recorded at the seafloor; 14 of these resulted in successful capture of either opalfish (*Hemerocoetes monopterygius*, 10 specimens) or blue cod (*Parapercis colias*, 2 specimens); prey species could not be identified during two captures, but the penguin's searching behaviour and ease of ingestion suggested these were opalfish (Figure 2). All of these prey pursuits occurred at the sea floor with the penguin swimming very close to the bottom (<https://vimeo.com/179414724>). During the camera operation time, the penguins spent 5.7 minutes on prey pursuit, which corresponds to 19% of the total time the bird foraged along the seafloor (29.9 minutes) and 6% of its total dive time (89.9 minutes); 3.8 minutes were devoted to pursuing and capturing opalfish; 46 seconds were used for the two blue cod captures, and the remaining 1.2 minutes were unsuccessful prey pursuits (Figure 2).

Two main prey pursuit strategies became apparent that were associated with prey species. When catching opalfish, the penguin would glide closely above the seafloor, sometimes briefly accelerating before starting to hover over a certain spot while repeatedly pecking at the substrate until the prey item was captured (<https://vimeo.com/179414724>). During encounters with blue cod prolonged pursuits ensued during which fish zigzagged at a fast pace along the seafloor (<https://vimeo.com/179414724#t=2m46s>). In one instance the fish was caught as it appeared to seek shelter at the base of a horse mussel protruding from the substrate (<https://vimeo.com/179414724#t=2m55s>). An unsuccessful prey pursuit of blue cod ended with the fish escaping under what appeared to be a half-buried back plate of a dishwasher (<https://vimeo.com/179414777>). A third blue cod encounter occurred just seconds after a successful capture of an opalfish; it seems likely that the resulting prolonged bottom time and oxygen-demanding prey pursuits drove the penguin to carry the fish to the surface at an almost vertical angle

as indicated sun disc's central position in the frame; the fish was ultimately dropped at the surface (<https://vimeo.com/179414724#t=3m07s>).

Benthic habitat

During the video logger's operating time, the penguin spent 29.9 minutes foraging along the seafloor. The majority of the penguin's bottom time (90%) was spent over coarse sand, whereas time spent over fine sand (7%) and gravel (0.9%) was negligible (Figure 2). Two thirds of the bottom time (65.9%) was spent over sand ripples, the remaining time (34.1%) the bird foraged over flat ground. Brittle stars and anthozoans were present in most areas visited by the penguin with the former being present in 22.5 mins (75%) of the benthic video footage while the latter occur for a total of 17.9 mins (60%). Horse mussels were present for a total of 9.3 minutes (31%) of the bottom time.

Prey encounters were associated with certain benthic habitat types. All prey encounters occurred over coarse sand although the sediment structure differed depending on prey species. Opalfish were principally encountered on sediment ripples (93.6% of the total prey pursuit time, <https://vimeo.com/179414724>), while flat bottom habitat played a more important role during blue cod pursuits (52.8% of pursuit time, <https://vimeo.com/179414724#t=2m32s>). With regards to epibenthic characteristics, brittle stars and anemones were present during the majority of the prey pursuit times for both fish species (Figure 3). However, horse mussels were present only during blue cod pursuits (81.4% of pursuit time).

Flipper movement

When descending to the sea floor the penguin propelled itself with fast, strong flipper strokes that got progressively slower and less pronounced with time and, thus, increasing depth (flipper amplitude: $\rho=-0.83$, $F_{1,363}=791.8$, $p<0.001$, BPM: $\rho=-0.36$, $F_{1,363}=55.2$, $p<0.001$, Figure 4a&b; <https://vimeo.com/179414575>). In contrast, ascending was principally passive with the penguin using its natural buoyancy to return to the surface, occasionally aided by a few strokes in the early stages of the ascent (flipper amplitude: $\rho=-0.08$, $F_{1,74}=0.5$, $p=0.488$; BPM: $\rho=-0.52$, $F_{1,74}=0.5$, $p<0.001$, Figure

4c&d) and no observable flipper movements towards the end of the dive (<https://vimeo.com/179414575#t=1m49s>). Despite differences in flipper movement between the two transit phases of a dive, the vertical velocities recorded by the TDR did not differ significantly (mean descent velocity: 1.45 ± 0.28 m/s, mean ascent velocity: 1.36 ± 0.57 m/s, $n=159$ dives, Welch's t -test: $t_{232}=1.73$, $p=0.09$).

During the bottom phase flipper amplitudes and beat frequencies showed no correlation with relative bottom time (flipper amplitude: $\rho=-0.08$, $F_{1,74}=0.5$, $p=0.488$; flipper BPM: $\rho=-0.52$, $F_{1,74}=0.5$, $p<0.001$, Figure 4c&d). This is owing to the fact that bottom phases consisted of a mix of searching behaviour and high speed prey pursuit (<https://vimeo.com/179414575#t=0m33s>). While searching the penguin showed lower flipper beat frequencies (133 ± 48 BPM, $n=809$) paired with greater flipper amplitudes ($53^\circ \pm 14^\circ$) when compared to prey pursuit (BPM: 162 ± 44 , $n=113$, $t_{232}=-13.37$, $p<0.001$; amplitude: $45^\circ \pm 7^\circ$, $t_{152}=6.39$, $p<0.001$).

Surface breathing & underwater exhalation

Frame counts of the video footage during 10 random selected surface periods between dives showed that the penguin lifted its head out of the water to breathe for only brief moments (average duration: 0.77 ± 0.22 s, $n=193$); for the majority of the time at the surface the bird kept its head under water (1.53 ± 1.19 s, $n=182$) (<https://vimeo.com/179414575#t=2m25s>). Duration of breathing intervals increased with ongoing duration of the surface period (Pearson correlation: $\rho=0.45$, $F_{1,191}=47.4$, $p<0.001$, Fig 3) indicating increased respiration activity in preparation for the next dive (Figure 3).

During the dive, exhalation regularly occurred at the onset of phases with increased acceleration (i.e. prey pursuit). Such exhalations were brief but performed with substantial force; air was jetted from the nostrils as a fine gas spurt (<https://vimeo.com/179418254>). During the passive phase of the ascent, the penguin frequently exhaled as indicated by a stream of large bubbles released from the nostrils. The bird released substantial amounts of air on the last few meters immediately prior to

reaching the surface (<https://vimeo.com/179414575#t=2m18s>). While some of this air may have been released from the plumage (c.f. Davenport *et al.* 2011) bubbles seem principally to originate from the frontal head region; there was no visible major gas release from the penguin's back region.

Discussion

The high-quality video footage provided a substantial amount of new insights into the foraging behaviour of Yellow-eyed penguins and their benthic habitat, while the device did not appear to substantially affect the penguin's underwater mobility.

Device effects

Attaching external recording devices to diving animals always comes at the cost of compromising their streamlined body shape (e.g. Ludynia *et al.*, 2012), a problem that can be mitigated via device shape, size and attachment position (Bannasch *et al.* 1994). At the surface there were no indications that the penguin was negatively affected by the device; the bird did not exhibit balancing problems which externally attached devices can cause in smaller species (Chiaradia *et al.* 2005), nor did it peck at the device frequently which suggests aberrant behaviour (Wilson & Wilson 1989). Moreover, the number of successful prey captures further suggests that the bird's foraging capabilities were not drastically affected by the video logger. The bird was one of the few breeders that raised two chicks to fledging in an otherwise poor breeding season.

Predator-prey interactions & prey species importance

In line with previous descriptions of yellow-eyed penguins as primarily benthic foragers (Mattern *et al.* 2007), the penguin's prey pursuit and captures recorded during the camera operation indeed all occurred at the sea floor. Swimming very close to the seafloor could serve several purposes. It could be a strategy to flush out benthic prey that blends in with the substrate, but it could also mean the penguin has a greater chance to see its prey from the side, and thus reduce the effect of prey camouflage. Opalfish, for example, are very well camouflaged and very difficult to make out from above (Roberts *et al.* 2015). This species seems to principally rely on its camouflage as means of

predator avoidance since none of the opalfish captures involved a chase. In contrast, during both successful blue cod encounters extended high-speed chases ensued before the fish was ultimately captured. Blue cod and opalfish differ significantly in their anatomy with the small, slender opalfish presumably lacking the physical prowess for prolonged swimming when compared to muscular blue cod (Roberts *et al.* 2015). When facing an air breathing predator, the latter strategy is likely advantageous as the predator's increased energy requirements for pursuit make escape a more likely outcome for the prey. The penguin's hasty ascent and subsequent failure to consume a blue cod it captured after a 22-second-long chase demonstrates the efficacy of this evasion strategy.

Both opalfish and blue cod have previously been found to be among the most important prey items in the Yellow-eyed penguin's diet (van Heezik 1990b; Moore & Wakelin 1997). While both fish species have comparable energetic values ($\sim 20 \text{ kJ g}^{-1}$, Browne *et al.*, 2011), the body mass of opalfish is considerably lower when compared to blue cod (van Heezik 1990a,b). So it is possible that the energy gain from catching blue cod justifies the expenditure to catch it, while the easier-to-catch opalfish might need to be caught in larger quantities. However, recent studies suggest that blue cod might be suboptimal prey for chick-rearing yellow-eyed penguins due to their size (Browne *et al.* 2011; Mattern *et al.* 2013) so that the penguins ability to locate prey such as opalfish might be a decisive factor with regards to reproductive success.

Benthic environment

Judging from the total time the bird spent over a benthic environment dominated by coarse sand and sediment ripples (65.9% of total bottom time) as well as almost exclusive encounters of opalfish over such habitat (Figure 2), it can be assumed that the penguin focussed principally on this species. Blue cod encounters were associated with the presence of horse mussels. These large bivalves protrude from the seafloor and provide hard substrate for other epibenthic taxa, thereby increasing local benthic biodiversity (Cummings *et al.* 1998). Benthic habitat with increased benthic biodiversity is generally more attractive to a variety of benthic fish species, most likely due to enhanced feeding

conditions (Cranfield *et al.* 2001). Our video data also suggests that the fish use the bivalves as shelter to avoid capture (<https://vimeo.com/179414777>).

The majority of prey pursuits occurred in areas that featured anthozoans, principally sea anemones (Figure 2). Anemones are known to play an important role as refugia and feeding habitats for small fish (Elliott 1992) and could therefore be another indicator for locally increased biodiversity. Brittle stars on the other hand, although equally abundant, seemed to be of lesser relevance with regards to prey encounters. So it appears that examining the composition of the benthic habitat alone might enable assessment of which prey types penguins are foraging for, though more data is required before conclusions can be drawn. However, this already hints at the potential for wide-ranging habitat analysis of at-sea movements in benthic top predators, provided that spatial distribution of the different benthic habitats can be obtained. While in our specific case, no such habitat maps exist, planned further deployments of video loggers are expected to provide the necessary environmental information.

Deploying video loggers on penguins could enable detailed mapping of the benthic habitat within the species home ranges. Yellow-eyed penguins are known to have preferred individual foraging areas often with little overlap between birds (Moore 1999). Moreover, the birds tend to often dive along the seafloor when swimming towards their foraging grounds (Mattern *et al.* 2007) so that camera logger data in combination with GPS information can be used to establish spatial biodiversity indices and benthic habitat maps.

The outer ranges of the marine habitat of Yellow-eyed penguins from the Otago Peninsula is subject to bottom fisheries which have a profound effect on benthic ecosystems (e.g. Hinz *et al.*, 2009; Queirós *et al.*, 2006; Schratzberger and Jennings, 2002). Yellow-eyed penguins have been found to forage in the wake of trawl fisheries, potentially to the detriment of their reproductive success (Mattern *et al.* 2013). Changes in sediment structure and epibenthic biodiversity as a result of bottom trawl disturbance likely negatively affect the penguins' foraging success (Browne *et al.* 2011).

Camera loggers can help to determine how much of the penguins' foraging habitat has been compromised by fishing activities and what the consequences are for this species' foraging behaviour and success.

Beyond investigations of behaviour in a wider environmental context, our study also shows the potential application of camera loggers for the investigation of physiological aspects of marine animals.

Flipper movements

Our observations of flipper movements, i.e. strong flipper movements at the beginning of a dive that decrease with depth, and cessation of flipper movements during ascent, align with findings reported in other penguins. Using accelerometers, Sato *et al.* (2002) found that King penguins showed vigorous flipper beating at the beginning of a dive to counter positive buoyancy. With increasing depth, air volume in the penguin's body becomes compressed, reducing its buoyancy so that fewer flipper beats are required. That this also applies to flipper amplitude (Fig 4) was not detectable by using body acceleration as the only measure. A more elaborate system of sensors and magnets attached to flippers was used on Magellanic penguins which allowed the recording of both flipper amplitudes and beat frequencies (Wilson & Liebsch 2003). However, the system proved to be prone to failure, rendering the use of back-mounted wide-angle cameras a much more reliable alternative. Flipper beat frequencies and amplitudes are directly related to energy expenditure (Kooyman & Ponganis 1998; Sato *et al.* 2011). They provide the means for the quantification of energy budgets (Wilson & Liebsch 2003) and subsequently can be used to assess individual fitness in relation to foraging success and subsequent reproductive performance (Kooyman & Ponganis 1998).

We provide proof that the ascent phase in penguins is largely passive, as has been suggested using both accelerometers and magnets (Sato *et al.* 2002; Wilson & Liebsch 2003). Sato *et al.* (2002) concluded that during ascent penguins benefit from expanding air volume in their body which increases their buoyancy as they get closer to the surface. Penguins also actively slow down their

ascent and it was argued that this could be achieved by increasing the attack angles of their flippers to increase drag (Sato *et al.* 2002). Judging from body movements apparent in the video data during the ascent phases we suggest that the yellow-eyed penguin indeed adjusted flipper attack angles while ascending, although this seems to be more for steering. Based on the video footage it appears that the birds might have used controlled exhalation towards the end of the ascent to control speed (<https://vimeo.com/179414575#t=2m18s>).

Respiration

The video data provides new insights into the respiration of Yellow-eyed penguins. To date it was unclear whether penguins exhale regularly while diving. Various studies estimated diving air volume via a penguin's buoyancy calculated from its ascent speeds at the end of dives (Sato *et al.* 2002, 2011). However, the accuracy of this approach is compromised if the penguins were to exhale prior to their final ascent (Ponganis *et al.* 2015). The video data clearly showed that the penguin generally exhaled when accelerating during prey pursuit so that models estimating diving air volume via the proxy buoyancy must take acceleration into account. The fact that the penguin exhaled when accelerating probably serves the purpose of reducing blood CO₂ and mobilizing O₂ from oxygen stores for prey pursuit. Such pursuits must be costly in terms of oxygen consumption as is evident from the observed consecutive prey encounters during one single dive, which resulted in the penguin letting go of the second fish after a rapid ascent to the surface (<https://vimeo.com/179414724#t=3m07s>). Unlike seals that have been found to exhale when ascending from deep dives, most likely to reduce drop in blood oxygen (Hooker *et al.* 2005), the penguin principally exhaled during the second half of the ascent possibly indicating adjustment of buoyancy and ascent speed (but see also Davenport *et al.* 2011). Reoxygenation during the surface period in penguins is highly optimized (Wilson *et al.* 2003). Inhalation events at the surface are brief so that the bird can frequently lower its head into the water, presumably in an effort to look out for potential predators (e.g. sharks, sea lions; Seddon *et al.*, 2013).

Conclusions

The deployment of a full HD video logger on a Yellow-eyed penguin resulted in a versatile visual data set that provided a variety of information well beyond what was initially intended. Enhanced video quality allows detailed analysis of the benthic environment as well as prey encounter rates and prey composition. In combination with GPS data, the potential for a comprehensive survey of benthic ecosystems is substantial highlighting the multi-disciplinary potential of such data.

A large field of view achieved through wide-angle lenses furthermore allows detailed analysis of flipper movements, which to date could only be achieved through elaborate modelling of accelerometer data (Sato *et al.* 2002, 2011) or use of complicated magnetic logger setups (Wilson & Liebsch 2003). Neither of these setups provided information about exhalation, which appears to play a much more important role during diving than previously thought. When comparing video data recorded here with videos from previously published studies (e.g. Watanabe and Takahashi, 2013) it becomes clear that greater visual fidelity of full HD cameras comes along with a much wider range of quantifiable data. This creates a new opportunity for a more holistic approach to study the diving behaviour of marine animals that integrates behaviour, physiology and their environment.

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Competing Interest

The authors declare no competing or financial interests.

Author contribution

T.M. designed the study; M.M. and T.M. developed the camera loggers; T.M., P.J.S. and J.v.H. conducted the field work; T.M. and U.E. analysed the data; T.M., M.M., U.E., Y.v.H. and P.J.S. wrote the manuscript.

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Figure 1. Proportion of benthic and pelagic dives throughout the Yellow-eyed penguin's foraging trip while fitted with a camera logger. Numbers at the top end of bars indicate number of dives performed during the corresponding hour. Red box indicates the hours during which continuous camera footage was recorded.

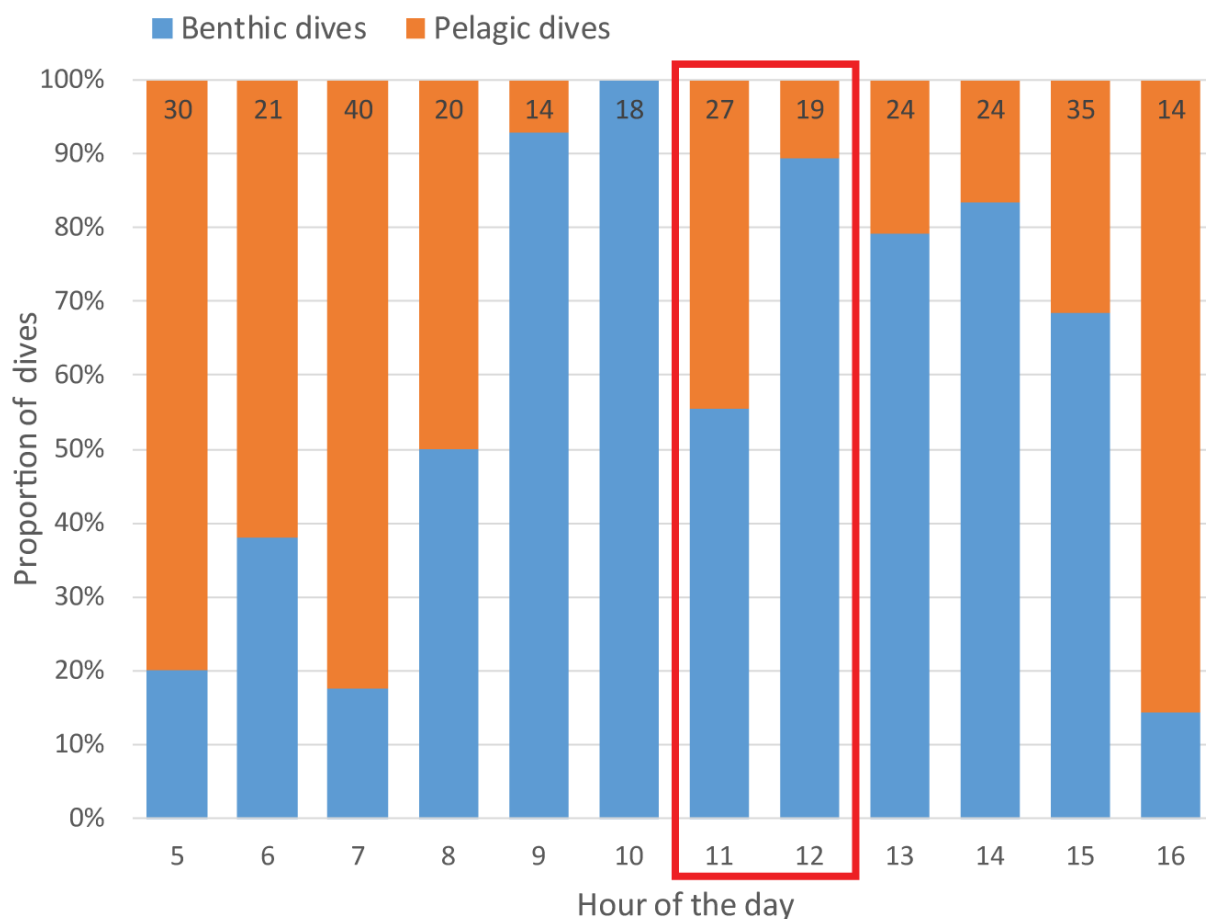


Figure 2. Composition of sediment structure, type and epibenthic communities during the bottom phases of 32 dives performed by a Yellow-eyed penguin fitted with a video camera logger. The x-axis indicates the cumulative time the penguin spent at the seafloor (29.9 minutes) during the 2 hours of camera operation. Coloured horizontal bars indicate duration of periods during which the penguin foraged over certain sediment structures and types, certain epibenthic taxa were present, as well as the length of prey pursuits (including pursuit outcome and prey species).

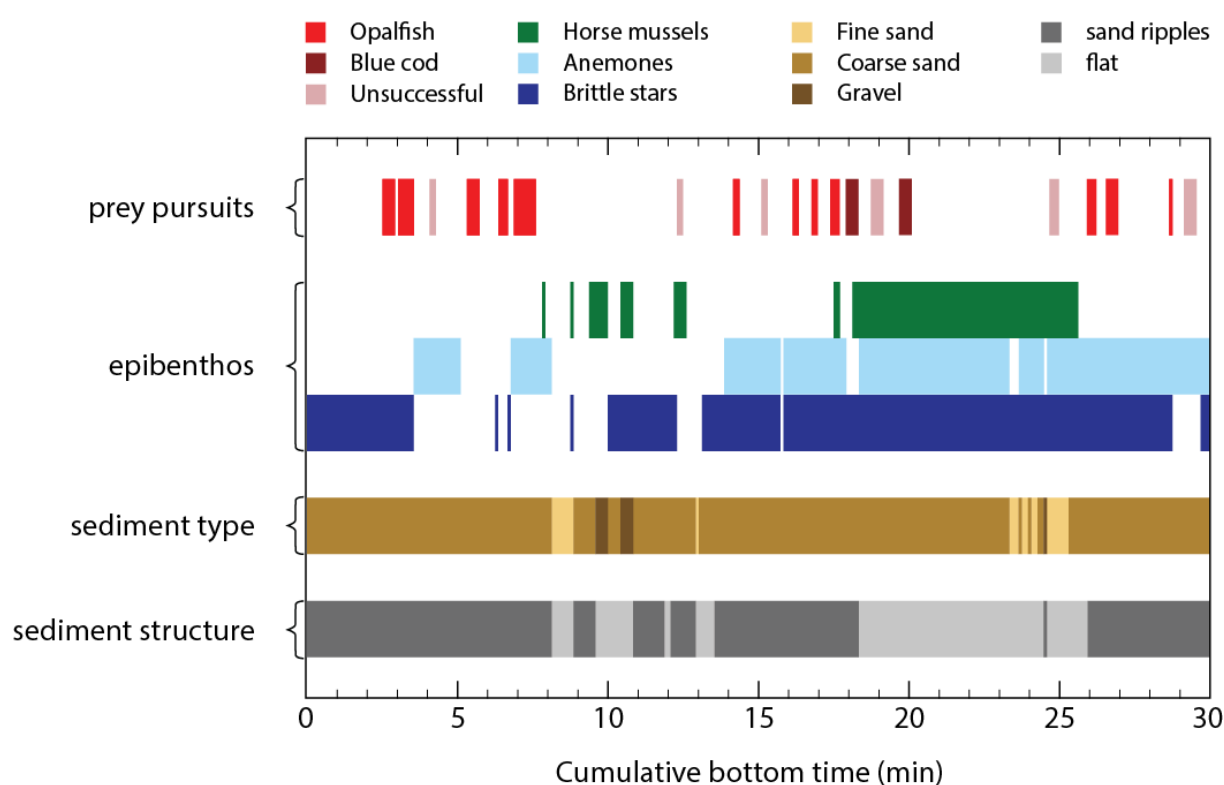


Figure 3. Increasing duration of breathing intervals (n=193) during the surface period after 10 randomly selected dives performed by a Yellow-eyed penguin. Note that the x-axis shows relative time to account for varying surface period durations. Red line indicates regression of data (see Results for details).

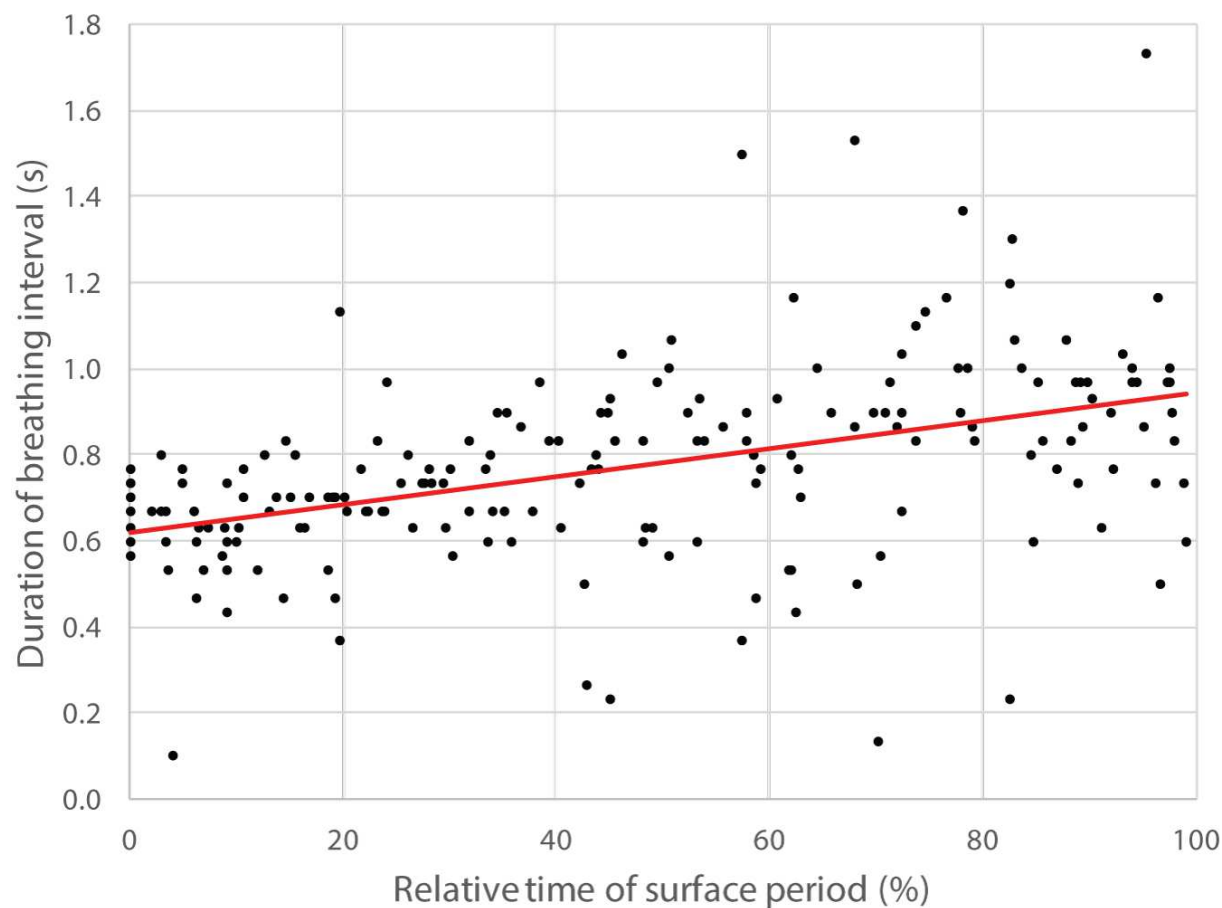


Figure 4. Flipper movements in a Yellow-eyed penguin during the descent, bottom and ascent phases of 10 randomly chosen benthic dives. Graphs in the upper row depict changes in flipper beat frequencies while the lower row consists of graphs showing flipper amplitude (i.e. maximum angle). Red lines indicate regression of the corresponding data (see Results for details). Note that x-axis shows relative durations of the dive phases to account for dive dependent time variations.

