# **Energetics of diving in macaroni penguins**

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### Summary

Heart rate ( $f_{\rm H}$ ), abdominal temperature ( $T_{\rm ab}$ ) and diving depth were measured in thirteen free-ranging breeding female macaroni penguins. Measurement of these variables allowed estimation of the mass-specific rate of oxygen consumption  $(\dot{V}_{O_2})$  while diving and investigation of the physiological adjustments that might facilitate the diving behaviour observed in this species. In common with other diving birds, macaroni penguins showed significant changes in fH associated with diving, and these variables accounted for 36% of the variation in dive duration. When  $\dot{V}_{O_2}$  was calculated for dives of different durations, 95.3% of dives measured were within the calculated aerobic dive limit (cADL) for this species. Mean fH for all complete dive cycles was  $147\pm 6$  beats min<sup>-1</sup>. When this fH is used to estimate  $\dot{V}_{O_2}$  of 26.2±1.4 ml min<sup>-1</sup> kg<sup>-1</sup> then only 92.8% of dives measured were within the cADL. Significant changes in abdominal temperature were not detected within individual dives, though the time constant of the measuring device used may not have been low

enough to record these changes if they were present. Abdominal temperature did decline consistently during bouts of repeated diving of all durations and the mean decrease in  $T_{ab}$  during a diving bout was  $2.32\pm0.2^{\circ}$ C. There was a linear relationship between bout duration and the magnitude of this temperature drop. There was no commensurate increase in dive duration during dive bouts as  $T_{ab}$  declined, suggesting that macaroni penguins are diving within their physiological limits and that factors other than  $T_{ab}$  are important in determining the duration of dives and dive bouts. Lowered  $T_{ab}$  will in turn facilitate lower metabolic rates during diving bouts, but it was not possible in the present study to determine the importance of this energy saving and whether it is occurs actively or passively.

Key words: energetics, diving, macaroni penguin, heart rate, abdominal temperature, rate of oxygen consumption, calculated aerobic dive limit.

#### Introduction

Penguins are among the most accomplished of divers. Numerous studies of their diving behaviour have shown that penguins have remarkable dive performances (Kooyman et al., 1992a; Williams, T. D. et al., 1992; Bengston et al., 1993). The emperor penguin *Aptenodytes forsteri*, the largest species at 25–30 kg, can reach depths of 524 m (Kooyman and Kooyman, 1995) for durations of up to 22 min (Robertson, 1994). Even the considerably smaller (3–4 kg) Adélie penguin *Pygoscelis adéliae* can dive to 98 m for up to 160 s (Wilson et al., 2002).

Further investigations examined how physiological and behavioural adjustments might permit such impressive diving behaviour (Butler and Jones, 1997; Kooyman and Ponganis, 1998). The extent to which diving animals balance the use of aerobic and anaerobic metabolism during natural dives is unclear. The majority of evidence suggests, however, that most dives are essentially aerobic (Butler and Jones, 1997). Anaerobic metabolism may be used in some circumstances (Kooyman et al., 1980; Ydenberg and Clark, 1989; Carbone and Houston, 1996; Mori, 1998; Butler, 2001), but within any dive there must be oxygen available for the central nervous system (CNS), heart and active muscles, even after lactate begins to accumulate. Observations of diving behaviour confirm that most dives are within bouts of repeated diving with relatively low ratios of post-dive surface interval duration to dive duration (dive:pause ratio).

The aerobic dive limit (ADL), the diving duration beyond which post-dive blood lactate levels increase above resting values, was first determined experimentally in Weddell seals (Kooyman et al., 1980) and defined by Kooyman et al. (1983). Since then, ADL or diving lactate threshold (DLT; Butler and Jones, 1997) has been determined in two more species of seal (Ponganis et al., 1997a,c) under captive conditions and in freely diving emperor penguins (Ponganis et al., 1997b) and bottlenose dolphins (Williams, T. M. et al., 1999). In emperor penguins the DLT was 5–7 min, which agreed quite closely with an ADL of 8 min estimated from observations of natural diving behaviour (behavioural ADL; Kooyman and Kooyman, 1995). This behavioural ADL was calculated as the dive

duration above which recovery times at the surface were proportionately longer in duration, suggesting that dives had a substantial anaerobic component. Only 4% of natural dives exceeded this behavioural ADL, therefore it was concluded that most diving was aerobic.

ADL has also been calculated (cADL) for many diving animals, including several penguin species, by dividing an estimate of usable body oxygen stores by an estimate of the rate of oxygen consumption ( $\dot{V}_{O_2}$ ) while submerged (Butler and Jones, 1997). When compared to observed patterns of diving in different penguin species, these studies have found that 2-50% of dives exceed the cADL (Culik et al., 1994, 1996a; Boyd and Croxall, 1996; Bethge et al., 1997; Bevan et al., 2002; Wilson et al., 2002). In these studies, examination of the dive:pause ratio suggests that it is unlikely that so many dives use predominantly anaerobic metabolism. In order for a large proportion of natural dives by many species of penguins to be aerobic, the cADL must be greater. Both usable oxygen stores and  $\dot{V}_{O_2}$  are difficult to measure while submerged, and other pathways such as the metabolism of phosphocreatine might provide energy under these conditions (Butler and Jones, 1997). Submerged  $\dot{V}_{O_2}$  is particularly difficult to measure (Costa, 1988). If estimates of the usable oxygen stores for penguins are approximately correct, then  $\dot{V}_{O_2}$  during diving needs to be as low as that recorded from penguins at rest on the water surface for most dives to be within the cADL (Butler, 2000).

In the present study we measured heart rate ( $f_{\rm H}$ ), abdominal temperature  $(T_{ab})$  and depth in macaroni penguins Eudyptes chrysolophus diving freely while foraging in their natural environment, using purpose-built implantable data loggers (Woakes et al., 1995). Heart rate can be used to estimate  $\dot{V}_{O_2}$ in diving animals (Fedak, 1986; Bevan et al., 1992; Butler, 1993) and a relationship between heart rate and  $\dot{V}_{O_2}$  has been established for macaroni penguins (Green, J. A. et al., 2001). This approach allows us to consider the effects of the suite of physiological and behavioural adaptations that have been found to contribute to the maximising of cADL while submerged. These adaptations include variation of heart rate and circulation (Butler and Woakes, 1979; Fedak et al., 1988; Kooyman et al., 1992b; Davis and Kanatous, 1999), regional hypothermia (Bevan et al., 1997, 2002; Handrich et al., 1997) and the use of passive gliding during the ascent and descent phases of dives (Williams, T. M. et al., 1999, 2000). Thus these measurements enabled us to relate the energetic costs and physiological responses to diving with the observed patterns of diving behaviour.

The present study, therefore, had four main aims: (1) to estimate from heart rate the energy cost of free-ranging diving behaviour in macaroni penguins, (2) to determine if macaroni penguins dive within their cADL and establish therefore whether they predominantly use aerobic respiration, (3) to examine heart rate changes on a fine scale (measured every 2 s) in order to assess whether circulatory adjustments made during diving might extend dive duration (Butler and Jones, 1997; Davis and Kanatous, 1999), (4) to measure abdominal temperature and investigate the hypothesis that lowered body temperature contributes to the extension of diving duration (Culik et al., 1996b; Handrich et al., 1997; Bevan et al., 2002).

# Materials and methods

# Study animals

The study was undertaken at the British Antarctic Survey (BAS) base on Bird Island, South Georgia during the austral summer of 1998/99. We followed the requirements of the UK Animal (Scientific Procedures) Act 1986, especially those set out by the Home Office in the Official Guidance on the operation of the Act. As our benchmark, we followed guidance to researchers using similar methods in the UK. Our procedures also conformed to the Code of Ethics of Animal Experimentation in Antarctica. The macaroni penguins used in the study were breeding females from the colony at Fairy Point on the north side of the island. The population at this colony has been monitored for many years (Williams, T. D. and Croxall, 1991) and has also been the subject of more intensive studies (Davis et al., 1983, 1989; Croxall et al., 1988, 1993, 1997; Williams, T. D., 1989). 15 penguins Eudyptes chrysolophus Brandt were used in the present study, all of which were engaged in provisioning a growing chick. Where possible, birds were caught for implantation away from the nesting area of the colony after they had fed their chick. After capture, the birds were removed to the surgical facility and kept in an outdoor enclosure for 2-3h before the surgery to allow digestion of food.

# Implantation procedure

Implantation of the data logger into the abdominal cavity allows data to be recorded without compromising the swimming, foraging and breeding performance of animals, as has been observed with the use of externally mounted devices on the morphometrically identical royal penguin (Hull, 1997). The implantation procedure was basically the same as described for similar studies (Bevan et al., 1995a). Briefly, the sterilised data logger was implanted into the abdominal cavity via a mid-line incision made in the skin and body wall muscle in the brood patch while the bird was anaesthetised with halothane. The logger design incorporates a low power radio frequency transmitter, which emits a short pulse on each QRS wave of the electrocardiogram (ECG). Detection of this signal on a radio receiver was used to indicate when the data logger was in the correct position. Once in position, the body wall and skin were sutured, antibiotic powder (Woundcare, Animalcare Ltd, York, UK) applied to the wound and a long-acting antibiotic (LA Terramycin, Pfizer, Sandwich, UK) and analgesic (Vetergesic, Reckitt and Colman Products Ltd, Hull, UK) injected intramuscularly. Aseptic conditions were maintained wherever possible. The time at which the data logger was implanted was noted to the nearest second.

All birds were weighed immediately before surgery using a spring balance ( $10\pm0.1$  kg, Pesola, Switzerland) and a passive implantable transponder (PIT) tag, mounted on a plastic cable

tie, was secured around their ankle. Birds were put into a large darkened box to recover from the surgery. Once the birds were alert and responsive, usually after 1–2 h, they were returned to the colony where behaviour varied between individuals. Some would go swimming within a few hours, whereas others made their way to the nest site or stood alone elsewhere in the colony. Around the time at which the data logger memory was predicted to be full, implanted birds were recaptured after returning from a foraging trip and having fed their chicks. The data logger was removed using the same procedure as during implantation, and the bird was released back in to the colony once it had recovered.

### Heart rate data loggers

The data loggers could record heart rate, hydrostatic pressure (diving depth) and abdominal temperature every 2s and, at this sampling rate, could store data over 30.3 days. Before use, the devices were encased in paraffin wax and encapsulated in silicon rubber to provide waterproofing and biocompatability. The hydrostatic pressure sensor in the data logger could detect diving depth to within 1.2 m. The temperature sensor of the encapsulated data logger was calibrated by immersing the device in water baths of known temperature. This procedure was also used to determine the time constant ( $\tau$ ) of the temperature sensor, which was 74 s. Unfortunately, given the relatively short dive durations of macaroni penguins, this meant that changes in abdominal temperature could only be analysed within diving bouts, not within individual dives. The time of removal of the data logger was noted and the precise times of implantation and removal were later used to establish the time base of the data downloaded from the data logger. The heart rate, abdominal temperature and depth data from within the data logger memory were downloaded onto a computer (Acorn RISC PC) using purpose-designed software.

## Data analysis

The data were prepared and analysed using purpose-written computer programs within the SAS statistical package (version 6.11, SAS institute) on a UNIX workstation. Further analyses were performed with the statistical packages Minitab 12 (Minitab Inc.), SPSS 10.0.8 (SPSS) and Excel 97 (Microsoft). The recovery period following the implantation procedure (Bevan et al., 2002) was excluded from the analysis by ignoring data collected during the period from implantation to the start of the first foraging trip. In the present study the duration of this period was  $55.5\pm5$  h (mean  $\pm$  S.E.M.).

Time at-sea on foraging trips was estimated from the depth data, supported with data from field observations and a PIT tag recorder (FSI Ltd, Cambridge, UK) situated in a gate at the edge of the colony. Each record of heart rate, abdominal temperature and dive depth was also marked with the daylight conditions (light or dark). These were calculated using the times for civil sunrise and sunset calculated for the longitude and latitude of Bird Island (54°00'S, 38°02'W). In examining dive records, dives with maximum depths of <2.4 m were

ignored during analyses, since wave action and recorder noise degraded depth accuracy for shallower dives. In all analyses, dives were treated as independent events. While accepting that this assumption may not be strictly correct, it is necessary in order to perform further statistical analyses.

A dive cycle was defined as a dive and the following interval spent at the water surface prior to the next dive. Bouts of dives were defined following the iterative statistical method of Boyd et al. (1994), which relies on searching the dive sequence for a change in behaviour that differs significantly from the previous set of behaviours since the last significant change. A minimum dive bout was formally defined as a group of at least three dives occurring within a period of 10 min. The dive record for each penguin was searched sequentially from the start, and once a group of dives had satisfied this minimum requirement, a search was made through the subsequent dives to find the end of the diving bout. This was done by calculating the mean and standard deviation (S.D.) of the surface intervals between dives, within the diving bout, and comparing these with the next surface interval in the sequence. If the next surface interval was significantly greater than the previous surface intervals in the bout (*t*-test, P < 0.01) then the bout was deemed to have ended. If the duration of the surface interval was not significantly different from those in the current bout, then the dive was included within the bout, the mean  $\pm$  s.D. of the surface intervals for the bout were recalculated, and the analysis then moved onto the next dive in the sequence.

The *f*H data were used to estimate mass specific rate of oxygen consumption,  $\dot{V}_{O_2}$ , using the relationship obtained from macaroni penguins walking on a treadmill (Green, J. A. et al., 2001). For breeding female penguins, which were the subjects of the present study, the equation was:

$$\dot{V}_{O_2} = (0.297 \times f_{\rm H}) - 17.40,$$
 (1)

 $r^2 = 0.84$ ,  $\dot{V}_{O_2}$  is in ml min<sup>-1</sup> kg<sup>-1</sup> at standard temperature and pressure, dry (STPD), and *f*H is in beats min<sup>-1</sup>.

This technique is normally calibrated when the animals' metabolism is in steady state and hence cannot be used to estimate  $\dot{V}_{O_2}$  while the animal is submerged. However, if *f*H and  $\dot{V}_{O_2}$  are averaged over a number of complete dive/surface cycles, then *f*H is an accurate and reliable predictor of  $\dot{V}_{O_2}$  in aquatic birds and mammals (Fedak, 1986; Bevan et al., 1992; Butler, 1993). The s.D. of an estimate made using Equation 1 was calculated using equation 11 of Green et al. (2001), which includes the variability within and between calibration and field animals, and is quoted in the text where estimates have been made.

Oxygen stores have not been measured in macaroni penguins, or indeed any of the crested penguins, but have been calculated for other species of penguins (Kooyman, 1989; Kooyman and Ponganis, 1990; Chappell et al., 1993; Bethge et al., 1997), usually following the assumptions of Stephenson et al. (1989) and Croll et al. (1992). These studies have detected differences between species and within species between different studies. However, the range of estimates is not large, varying from  $45 \text{ ml O}_2 \text{ kg}^{-1}$  in little blue penguins

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	Mass	Duration of	Proportion	Proportion of time	Number of
Bird	(kg)	data record (days)	of time at-sea	submerged while at-sea	dives recorded
H02	3.6	8.26	0.61	0.61	2656
H15	3.3	12.95	0.80	0.58	6367
H17	3.1	4.65	0.78	0.58	1926
H25	3.9	11.56	0.69	0.56	4466
H29	3.6	6.17	0.79	0.47	2341
H53	3.3	28.01	0.49	0.52	7182
H59	4.0	1.29	0.83	0.48	466
H61	3.4	6.29	0.70	0.50	2339
H69	3.8	4.18	0.73	0.17	639
H73	3.8	3.48	0.54	0.42	857
H79	4.1	26.81	0.51	0.58	6932
H93	3.6	9.09	0.54	0.62	3316
H95	3.4	13.26	0.58	0.64	4535
Mean ± S.E.M.	3.6±0.3	10.46±2.32	0.67	0.52	3386±649

 Table 1. Deployment details and simple parameters of diving for 13 breeding female macaroni penguins from which data were obtained

(Bethge et al., 1997) to  $63 \text{ ml O}_2 \text{ kg}^{-1}$  for Adélie penguins (Culik et al., 1994). In the present study it was not possible to collect the data necessary to calculate oxygen stores for macaroni penguins, so a value of  $58 \text{ ml O}_2 \text{ kg}^{-1}$  was used, which is in the middle of the range of most of the calculated values for other species and has been used previously as an estimate to compare different penguin species (Butler, 2000). Stephenson et al. (1989) discuss the influence of training on

the composition of oxygen stores but there is no reason to assume that the birds in the present study were not fit and acclimated for intensive diving.

Data were analysed using analysis of variance (ANOVA) with Tukey *post-hoc* testing, linear regression and stepwise multiple linear regression. Results were considered significant at P<0.05 and the significance level is quoted in the text. Unless stated otherwise, mean values are the grand mean of

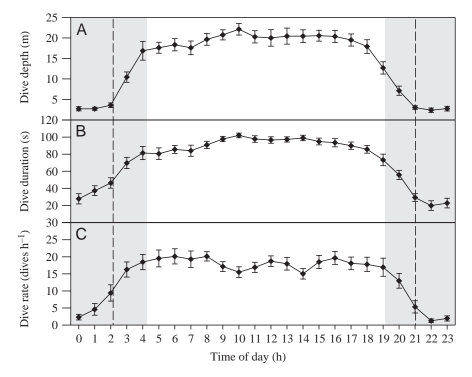


Fig. 1. Influence of time of day on (A) dive depth, (B) dive duration and (C) dive rate, recorded from 13 breeding female macaroni penguins. All values are means  $\pm$  s.E.M. Shaded areas indicate hours of darkness, and broken vertical lines indicate dawn and dusk at the beginning of the season.

the mean value for each penguin and are  $\pm$  1 s.E.M. Percentage values were arcsinetransformed before comparisons were made (Zar, 1999). All times are given in local time (GMT –3 h) unless otherwise stated.

# Results

# Deployments, diving behaviour and dive bouts

Data were obtained from 13 penguins. Failure in the encapsulation led to battery failure in the other two deployments. Table 1 shows details of the 13 birds from which data were obtained. Diving activity was greater during daylight (Fig. 1), when dives were deeper (two-way ANOVA,  $F_{23,276}$ =45.19, P<0.001), more frequent (two-way ANOVA,  $F_{23,276}$ =15.43, P<0.001) and of longer duration (two-way ANOVA,  $F_{23,276}$ =51.38, P<0.001).

When dives were classified into bouts, 98% of all dives were part of a bout consisting of at least three dives (Table 2). Only dives within bouts were considered for further analyses. When considering post-dive surface intervals, the last dive of a bout was discarded. Individual distributions of both dive depth and duration were not normal, so Kruskal-Wallis tests with Dunn's multiple comparisons were used to examine differences between individuals. There were significant differences between individuals in both dive depth (Kruskal-Wallis statistic<sub>(13)</sub>=964.3, *P*<0.001) and duration (Kruskal-Wallis statistic(13)=1088, *P*<0.001) (Table 2). Fig. 2 shows the mean frequency distributions of dive depth and duration, calculated by taking an average of the individual frequencies of occurrence of each dive depth or duration interval from all 13 penguins. These distributions were not substantially different from those of all dives from all penguins but this approach treats all individuals equally, despite large differences in the number of dives recorded from individual penguins (Table 2). 21% of all dives were to a maximum depth of 4.8 m (Fig. 2A), with declining frequencies to 94.8 m, the maximum dive depth recorded. This dive was recorded by penguin H79, which was responsible for most of the deeper and longer dives, including all those deeper than 70 m. Dive durations were more normally distributed (Fig. 2B), though slightly negatively skewed.

# Abdominal temperature during diving

The mean  $T_{ab}$  while on-shore was 40.1±0.9°C, and the mean  $T_{ab}$  during

diving bouts and while at-sea but not diving were  $34.8\pm1.2^{\circ}$ C and  $38.2\pm1.0^{\circ}$ C, respectively. Two-way analysis of variance with Tukey *post-hoc* testing ( $F_{2,38}=31.6$ , P<0.001) revealed significant differences between all three measurements of  $T_{ab}$ . Further analyses were performed to investigate the decrease in  $T_{ab}$  associated with diving and what effect it might have in improving diving performance. Average diving temperature ( $DT_{ab}$ ) was calculated for each dive as the mean temperature while submerged. Linear regressions were used to determine whether  $DT_{ab}$ , dive duration and mean diving fH varied progressively during the course of each diving bout (Table 3). 63.4% of all dive bouts showed a significant change in  $DT_{ab}$  through the course of the bout and 76.2% of these (i.e. 48.3% of all dive bouts) were significant declines, with a mean  $r^2$  of

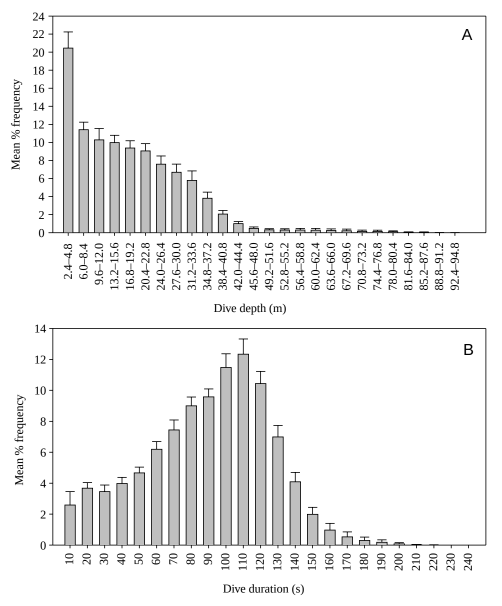


Fig. 2. Mean frequency distributions from 13 breeding female macaroni penguins of (A) dive depth and (B) dive duration. All dives were within bouts of three or more dives. Values are means + S.E.M.

0.76 (Table 3). However, only 35.0% and 35.4% of bouts showed a significant change in dive duration and *f*H, respectively, over the course of the bout, and the average  $r^2$  of these relationships was only 0.37 and 0.34, respectively. The decline in  $T_{ab}$  ( $\Delta T_{ab}$ ) during each dive bout was calculated as the difference between the maximum and minimum values of  $DT_{ab}$  from that bout. Mean  $\Delta T_{ab}$  from all 13 penguins was 2.32±0.20°C, range 0–13.51±1.1°C.  $\Delta T_{ab}$  increased with the duration of the diving bout for each individual (mean  $r^2$ =0.55, all *P*<0.001) and for all diving bouts pooled ( $r^2$ =0.46, *P*<0.001, Fig. 3).

*Heart rate and rate of oxygen consumption while diving* Mean heart rates while the penguins were on-shore and at-

	Dive	depth (m)			Dive dur	ation (s)	
 Mean	Mode	Median	Maximum	Mean	Mode	Median	Maximum
21.7±0.3	7.2	19.2	69.6	100.3±0.7	124	102	180
17.3±0.1	2.4	14.4	57.6	$81.4\pm0.4$	78	82	224
18.1±0.2	15.6	16.8	45.6	95.4±0.7	108	102	164
19.0±0.2	2.4	18.0	60.0	88.1±0.5	112	96	168
18.3±0.2	2.4	20.4	42.0	85.6±0.6	102	94	142
14.9±0.1	2.4	12.0	69.6	84.5±0.4	109	86	178
13.0±0.4	2.4	10.8	54.0	95.4±1.4	109	98	186
16.2±0.2	2.4	14.4	51.6	83.5±0.7	114	90	156
18.1±0.5	2.4	19.2	44.4	68.6±1.6	104	82	180
$14.2\pm0.4$	2.4	12.0	44.4	80.3±1.1	100	84	172
23.4±0.3	2.4	13.2	94.8	95.4±0.6	58	92	232

79.8±0.6

 $87.8\pm0.5$ 

86.6±2.4

68

112

82

92

99.8±5.4 90.9±2.0 179.2±6.8

174

174

Table 2. Characteristics of diving be

Number of dives

Within bouts

2615

6251

1879

4371

2281

7091

2262

456

618

845

6814

3277

4484

3326±640

Not

within bouts

41

116

47

95

60

91

10

77

21

12

118

39

51

 $59.85 \pm 10.24$   $35.77 \pm 2.46$ 

Average

per bout

49.34

36.56

43.70

31.00

34.56

38.12

19.00

34.27

30.90

24.14

31.55

46.15

45.76

Number of

diving bouts

53

171

43

141

66

186

24

66

20

35

216

71

98

91.54±19.20

Bird H02

H15

H17

H25

H29

H53

H59

H61

H69

H73

H79

H93

H95

 $Mean \ \pm$ 

S.E.M.

Table 3. Characteristics of the linear regressions describing changes in abdominal temperature, heart rate and dive duration within diving bouts

16.1±0.2

 $17.5 \pm 0.1$ 

 $17.5 \pm 0.8$ 

2.4

2.4

3.8±1.0

15.6

18.0

15.7±0.9

46.8

58.8

 $56.9 \pm 4.1$ 

	Dive bouts w temperature, l			Dive bouts wa temperature, l	U		Bouts with r temperature, h	no change in a neart rate or di	
Variable	Mean proportion of bouts (%)	Mean r <sup>2</sup>	Number of dives/bout (mean ± S.E.M.)	Mean proportion of bouts (%)	Mean r <sup>2</sup>	Number of dives/bout (mean ± s.E.M.)	Mean proportion of bouts (%)	Mean r <sup>2</sup>	Number of dives/bout (mean ± S.E.M.)
Abdominal temperature	48.3	$0.76 \pm 0.04$	54 <u>+</u> 4	15.1	$0.64 \pm 0.04$	27±3	36.6	0.33±0.03	14±1
Heart rate	12.0	0.31±0.04	72±9	23.31	$0.36 \pm 0.02$	60±7	64.6	$0.16\pm0.01$	21±1
Dive duration	16.1	$0.36 \pm 0.02$	80±6	18.9	$0.38 \pm 0.03$	45±4	65.0	$0.02\pm0.01$	22±2

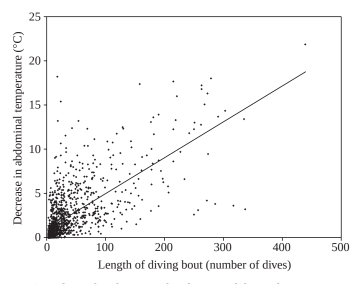


Fig. 3. Relationship between the drop in abdominal temperature  $(\Delta T_{ab})$  and duration of diving bouts recorded from 13 breeding female macaroni penguins. *y*=0.04*x*+0.91.

sea were  $116\pm 6$  and  $148\pm 7$  beats min<sup>-1</sup>, respectively. While the penguins were at-sea, mean heart rate during diving bouts (DfH) was  $147\pm6$  beats min<sup>-1</sup>, whereas mean heart rate while the birds were at-sea but not diving (NDfH), calculated from fH between diving bouts, was  $154\pm8$  beats min<sup>-1</sup>. Two-way ANOVA with Tukey post-hoc testing ( $F_{2,38}=38.1$ , P<0.001) showed that DfH was not significantly different from NDfH, but both were significantly greater than *f*H while on-shore. During the dive cycle, macaroni penguins showed increases and decreases in fH associated with dives of all durations. The extent of these changes in fH associated with diving were related to dive duration. Table 4 shows mean, maximum and minimum fH at different stages of the diving cycle for dives of different durations and for dives of all durations, while Fig. 4 shows how heart rate varied during dives lasting 102–110 s, the most frequently observed category of dive duration (Fig. 2B). A similar pattern was observed in dives of both longer and shorter durations and can be described as follows. (1) Prior to diving, fH was elevated above DfH and started to decrease just before submergence. (2) Upon submerging, fH immediately decreased before recovering slightly. fH then decreased more slowly to a level below  $Df_{H}$ . (3) At the bottom of the dive fH tended to stabilise. (4) As the penguin started to ascend to the surface, fH increased slowly. (5) After the penguin surfaced, fH then increased more rapidly to a level above  $Df_{H}$ . (6) This high heart rate was usually followed immediately by another dive, if the dive was part of a dive bout, otherwise fH declined to DfH. ANOVA showed that if dives of all durations were averaged together, there were significant differences between DfH and fH at different stages of the dive cycle (two-way ANOVA,  $F_{4,64}=97.8$ , P<0.001). Further Tukey *post-hoc* tests showed that mean pre-dive and post-dive  $f_{\rm H}$  values were significantly higher than mean  $Df_{\rm H}$ , mean fH while submerged and minimum fH while submerged.

					breeding femal	breeding female macaroni penguins	guins				
						Ψ	<i>f</i> H (beats min <sup>-1</sup> )				
							During first			Pre-dive –	Pre-dive –
Dive	Number of nonming	Number	Pre-dive	Pre-dive	During dive	During dive	10 s of dive	Post-dive	Post-dive	during dive	during dive
duranon (s)	or penguins	OI DIVES	(mean)	(max)	(mean)	(min, mean)	(umu)	(mean)	(max)	(max, min)	(mean)
0-20	13	1913	$185\pm11$	$242\pm9$	$156\pm 8$	$105 \pm 9$	$107\pm 8$	$179\pm10$	$263\pm10$	$30\pm7$	$137\pm 8$
22–40	13	3048	$194{\pm}12$	$255 \pm 10$	$143\pm 8$	$89 \pm 9$	$107{\pm}8$	$186 \pm 11$	$264{\pm}10$	$51 \pm 7$	$166\pm 8$
42–60	13	5253	$203\pm 15$	$269\pm11$	$139 \pm 7$	$81\pm 8$	$109\pm 8$	$200 \pm 14$	$274{\pm}12$	$65 \pm 11$	$188\pm 8$
62-80	13	7111	$209{\pm}16$	$273\pm12$	$133\pm 5$	$71 \pm 7$	$105 \pm 7$	$207{\pm}16$	$276\pm 12$	$76\pm12$	$203\pm 9$
82 - 100	13	8206	$214{\pm}17$	$278\pm12$	$129\pm 4$	$62\pm 6$	95±7	$212\pm 17$	$278{\pm}12$	$84{\pm}14$	$215\pm 9$
102 - 120	13	9005	$216{\pm}18$	$281\pm12$	$127\pm 4$	$56\pm 5$	$89\pm6$	$214{\pm}18$	$281 \pm 12$	$89\pm 15$	$225\pm10$
122 - 140	13	5162	$217\pm 17$	$289\pm 12$	$125\pm3$	53±5	$93\pm 8$	$215\pm 16$	$288{\pm}10$	$91\pm 15$	$236\pm 11$
142 - 160	12	1562	$209\pm19$	$284{\pm}11$	$122\pm 4$	$51\pm 6$	$91{\pm}11$	$208{\pm}19$	$282 \pm 12$	$87{\pm}19$	$233\pm 9$
162 - 180	7	554	$208\pm 26$	$264\pm 26$	$119\pm 11$	$53\pm 8$	$93 \pm 14$	266±35	$319\pm 25$	$89\pm 22$	$211\pm 20$
182 +	2	240	$210 \pm 30$	$248\pm 8$	$153\pm 54$	$47{\pm}11$	$156\pm60$	$182 \pm 11$	$283\pm 23$	57±24	$201 \pm 19$
All dives	13	42054	$208{\pm}15$	$273\pm10$	$132\pm 5$	67±6	97±7	$206\pm 15$	$275{\pm}10$	76±12	206±7
Values are	Values are means $\pm$ S.E.M.										
Max, maxii	Max, maximum; min, minimum	mum.									

Table 4. Mean, maximum and minimum values of heart rate (fH) associated with different stages of the diving cycle, for dives of different duration, recorded from 13

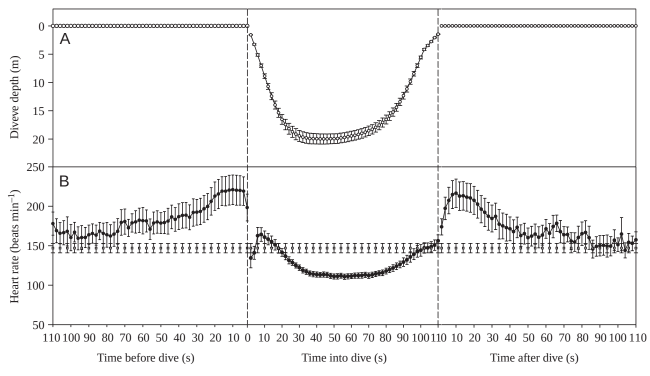


Fig. 4. (A) Changes in depth during dives of 102-110 s and (B) associated changes in heart rate (filled circles) with overall heart rate during diving bouts (open circles), recorded from 13 breeding female macaroni penguins. Values are means  $\pm$  S.E.M.

Furthermore, minimum  $f_{\rm H}$  while submerged was significantly lower than D $f_{\rm H}$  and mean  $f_{\rm H}$  while submerged. There was no significant difference between mean  $f_{\rm H}$  while submerged and D $f_{\rm H}$ .

In order to investigate further how the marked changes in heart rate during the dive cycle might be related to dive duration, stepwise multiple linear regression analysis was used. Two multiple regressions were performed. In the first analysis, the dependent variable was dive duration and the independent variables were measurements of fH made during the corresponding dive cycle, thought to be those that characterised the major features of the fH changes during a dive. These variables were: mean diving fH, minimum diving fH, minimum fH within the first 10s of submersion, pre-dive mean fH, pre-dive maximum fH, post-dive mean fH and postdive maximum fH. The analysis was performed for each penguin using all of its dives, and for all dives from all the penguins pooled (Table 5). The analysis indicated that, on average, 36% of the variation in dive duration could be predicted by the adjustments in fH. There was considerable variation between individuals but, as shown in Table 5, the most consistent influences on dive duration of the individual penguins were minimum fH while submerged, followed by minimum fH shortly after submersion and mean post-dive fH. When all the dives from all penguins were pooled, the three most important influences were minimum fH when submerged, mean pre-dive *f*H and minimum *f*H shortly after submersion.

The second analysis used differences in fH between different phases of the dive cycle, as the magnitude of these

changes also appeared to vary with changes in dive duration. In this regression, the dependent variable was again dive duration and the independent variables were: the difference in fH from mean pre-dive to mean during diving, the difference in fH from maximum pre-dive to minimum during diving and the difference in fH from maximum pre-dive to minimum within 10 s of submersion. Again, the analysis was performed for each penguin using all of its dives, and for all dives from all the penguins pooled (Table 6). This analysis explained on average 22% of the variation in dive duration, and for each individual and all dives pooled the  $r^2$  value was lower than in the corresponding first analysis. This analysis was clearly of less value than the first and was not considered further.

The resulting multiple regression equations for each individual penguin (Table 5) could be used to predict dive duration from measurements of heart rate for that animal. Though all of the individual relationships were significant (Table 5), the reliability of such a prediction would vary considerably from individual to individual as there was considerable variation in the  $r^2$  values of the relationships (0.05–0.71). The relationship for all of the penguins pooled could be used to predict dive duration for an individual from outside this study, from measurements of *f*H. However, the  $r^2$  value of this relationship was relatively low (0.20, Table 5), meaning that the confidence intervals around such a prediction would be large and the prediction of limited value.

 $\dot{V}_{O_2}$  while on-shore and at-sea, estimated using Equation 1, was  $16.9\pm1.4$  and  $26.3\pm1.4$  ml min<sup>-1</sup> kg<sup>-1</sup>, respectively.  $\dot{V}_{O_2}$ 

was not calculated from D/H and ND/H since these were not significantly different from each other or *f*H while at-sea. Equation 1 was, however, used to estimate  $\dot{V}_{O_2}$  and the 95% confidence limits of these estimates, using *f*H from completed dive cycles. Since *f*H varied with dive duration (Table 4), it was necessary to estimate  $\dot{V}_{O_2}$ and the confidence limits at each different dive duration for the full range observed by macaroni penguins (Table 7). As mean *f*H decreased with dive duration, then so did estimated  $\dot{V}_{O_2}$  (Table 7).

# Discussion

these two to give more detailed dive profiles. Despite macaroni variation in diving behaviour. studies, there was considerable individual to approximately 40-50 m. In all four to less than 5 m and other longer dives depth and duration, with many short dives were observed in distributions of dive observed in the present study. Similarities these, in turn, are similar to the patterns shown by these studies were similar and location, the patterns in diving behaviour depth recorders measuring depth every 3 s Island (Green et al., 1998), and used time breeding males and females at Heard comprehensive study was completed on was heavy, bulky and was only used on sophisticated dive depth recorder, but this breeding males. The second used a more simple depth histogram recorder on eight 1993). The first of these studies used a mounted devices breeding the diving behaviour of macaroni penguins In all Two previous studies have investigated female differences in methodology and at Bird Island, using externally four of the above penguins Diving behaviour penguins. (Croxall et al., tended Another more ð studies, 1988,

diurnal more widely dispersed through the water predominantly in daylight. Dives at night column the top of the water column at night but are (Croxall et al., 1993). Krill are found near penguins foraging in waters around Bird of shorter duration (Fig. 1). For macaroni were less frequent, to shallower depths and feeding near Heard Island, the reasons are Island, a suggested cause for this is the during daylight. For penguins migration of Antarctic dive krill

Table 5. Multiple linear regression equations describing how heart rate (fH) in female macaroni penguins during different stages of their dive cycle is related to dive

					bles and parameters				
		fH during dive	(beats min <sup>-1</sup> )	<i>f</i> H post-d	live	Min <i>f</i> H during	<i>f</i> <sub>H</sub> pre-	dive	
Bird	Intercept	Mean	Min	Mean	Max	first 10 s	Mean	Max	$r^2$
H02	$-1.47{\pm}6.16^{ns}$	ns	<sup>1</sup> -0.660±0.029***	<sup>2</sup> 0.418±0.031***	ns	<sup>4</sup> -0.088±0.022***	<sup>3</sup> 0.235±0.030***	ns	0.45
H15	67.26±1.67***	3 0.246±0.018***	<sup>1</sup> -0.727±0.022***	<sup>2</sup> 0.126±0.018***	$^{6}$ -0.018±0.008*	<sup>5</sup> 0.042±0.016**	4 0.081±0.013***	ns	0.27
H17	106.72±1.98***	ns	<sup>1</sup> -0.638±0.035***	ns	4 0.026±0.012*	<sup>3</sup> 0.060±0.024*	ns	<sup>2</sup> 0.032±0.011**	0.18
H25	100.78±4.53***	<sup>4</sup> -0.274±0.034***	<sup>1</sup> -0.707±0.024***	<sup>7</sup> 0.090±0.018***	<sup>5</sup> 0.029±0.012*	<sup>3</sup> 0.120±0.015***	<sup>2</sup> 0.090±0.018***	6 0.062±0.012***	0.43
H29	105.38±2.47***	ns	<sup>1</sup> -0.135±0.021***	4 0.039±0.013**	<sup>2</sup> -0.252±0.012***	$^{3}-0.045\pm0.014$ **	ns	ns	0.05
H53	31.47±3.32***	<sup>2</sup> -0.362±0.023***	<sup>5</sup> -0.102±0.011***	<sup>3</sup> 0.222±0.010***	ns	<sup>4</sup> -0.165±0.011***	<sup>1</sup> 0.285±0.010***	ns	0.40
H59	43.71±13.13***	ns	<sup>1</sup> -0.402±0.046***	<sup>2</sup> 0.182±0.056***	ns	ns	<sup>3</sup> 0.142±0.053**	ns	0.28
H61	95.08±2.13***	4 0.115±0.023***	<sup>1</sup> -0.577±0.033***	ns	<sup>5</sup> -0.024±0.008**	<sup>2</sup> 0.144±0.017***	<sup>3</sup> -0.092±0.015***	ns	0.18
H69	24.28±5.46***	<sup>5</sup> -0.099±0.035**	<sup>1</sup> -0.497±0.032***	<sup>2</sup> -0.195±0.017***	ns	<sup>3</sup> 0.186±0.020***	4 0.116±0.018***	ns	0.71
H73	42.66±5.93***	<sup>5</sup> -0.200±0.052***	<sup>1</sup> -0.495±0.033***	<sup>2</sup> 0.205±0.023***	ns	4 0.133±0.025***	<sup>6</sup> 0.063±0.027*	<sup>3</sup> 0.081±0.014***	0.59
H79	87.94±1.31***	ns	<sup>2</sup> -1.133±0.033***	ns	<sup>5</sup> 0.061±0.010***	4 0.199±0.025***	<sup>3</sup> 0.126±0.016***	<sup>1</sup> 0.078±0.012***	0.27
H93	66.19±4.14***	<sup>5</sup> -0.085±0.012***	$^{1}-0.492\pm0.022***$	<sup>2</sup> 0.296±0.023***	<sup>3</sup> -0.179±0.016***	4 0.185±0.015***	60.200±0.023***	<sup>7</sup> -0.079±0.015***	0.49
H95	130.61±2.75***	$^{1}-0.637\pm0.025***$	<sup>3</sup> -0.373±0.019***	<sup>5</sup> -0.090±0.013***	<sup>4</sup> 0.116±0.009***	ns	$^{6}$ -0.030±0.013*	<sup>2</sup> 0.151±0.010***	0.38
Mean ±									0.36±
S.E.M.									0.04
All dives	92.40± 0.71 ***	<sup>5</sup> -0.078±0.007***	<sup>1</sup> -0.388±0.006 ***	4 0.073±0.005***		<sup>3</sup> -0.076±0.005***	<sup>2</sup> 0.107±0.006***	<sup>6</sup> 0.011±0.003***	0.20

duration

Max, maximum; min, minimum.

 $The equations are of the format: dive duration = intercept + (parameter_1 \times variable_1) + \ldots + (parameter_n \times variable_n).$ 

\*A significant factor at P<0.05 (Wald test); \*\*a significant factor at P<0.01; \*\*\*a significant factor at P<0.001; ns, a non-significant factor.

A superscript number before the parameter describes the order in which that variable was found to improve the fit of the stepwise regression (*F*-test) and added to the model.

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		V	ariables and parameters		
Bird	Intercept	Max pre-dive f <sub>H</sub> -min f <sub>H</sub> during first 10 s	Max pre-dive f <sub>H</sub> -min f <sub>H</sub> during dive	Mean pre-dive f <sub>H</sub> -mean f <sub>H</sub> during dive	$r^2$
H02	-22.63±3.95***	ns	<sup>1</sup> 0.349±0.020***	<sup>2</sup> 0.407±0.024***	0.29
H15	64.01±1.08***	ns	1 0.133±0.008***	<sup>2</sup> -0.066±0.008***	0.06
H17	77.09±2.24***	<sup>3</sup> -0.149±0.026***	<sup>1</sup> 0.248±0.027***	<sup>2</sup> -0.130±0.017***	0.06
H25	17.80±1.89***	$^{3}-0.084\pm0.017***$	1 0.375±0.020***	<sup>2</sup> 0.056±0.010***	0.28
H29	ns	ns	ns	ns	0.00
H53	23.49±1.51***	<sup>2</sup> 0.053±0.010***	<sup>3</sup> 0.023±0.009**	$^{1} 0.377 \pm 0.008 ***$	0.33
H59	9.78±8.41 <sup>ns</sup>	ns	<sup>1</sup> 0.297±0.039***	<sup>2</sup> 0.215±0.047***	0.22
H61	83.19±1.95***	$^{1}$ -0.151±0.016	<sup>2</sup> 0.125±0.017***	$^{3}$ 0.041±0.019*	0.05
H69	19.81±2.45***	<sup>3</sup> -0.172±0.026***	<sup>1</sup> 0.262±0.026***	<sup>2</sup> 0.221±0.025***	0.56
H73	21.82±2.66***	<sup>2</sup> -0.219±0.025***	1 0.351±0.026***	<sup>3</sup> 0.211±0.025***	0.48
H79	61.22±1.35***	<sup>2</sup> -0.210±0.027***	$^{1}$ 0.412±0.028	$^{3}$ -0.082±0.013***	0.13
H93	33.14±1.92***	$^{3}$ -0.178±0.017***	<sup>2</sup> 0.222±0.020***	$^{1}$ 0.291±0.010***	0.35
H95	49.75±1.75***	<sup>3</sup> -0.048±0.017**	<sup>1</sup> 0.245±0.018***	<sup>2</sup> -0.110±0.009***	0.11
Mean ± s.e.M	I.				0.22±0.05
All dives	64.79±0.50***		<sup>1</sup> 0.109±0.003***	<sup>2</sup> 0.020±0.003***	0.07

 Table 6. Multiple linear regression equations describing how changes in heart rate (fH) in female macaroni penguins during different stages of their dive cycle are related to dive duration

Values are means  $\pm$  s.e.m.

Max, maximum; min, minimum.

The equations are of the format: dive duration=intercept+(parameter\_xvariable\_1) + ....+ (parameter\_xvariable\_n).

\*A significant factor at P<0.05 (Wald test); \*\*a significant factor at P<0.01; \*\*\*a significant factor at P<0.001; ns, a non-significant factor. A superscript number before the parameter describes the order in which that variable was found to improve the fit of the stepwise regression (*F*-test) and added to the model.

			ν̈́c	$D_2$ estimate (ml min <sup>-1</sup>	kg <sup>-1</sup> )	
Dive duration (s)	Percentage of dives	<i>f</i> <sub>H</sub> during dive cycle (beats min <sup>-1</sup> )	During dive cycle	Lower 95% confidence limit	Upper 95% confidence limit	cADL, calculated using $\dot{V}_{O_2}$ estimate (s)
2-10	2.2	172±10	33.6±1.5	30.7	36.5	104
12-20	3.0	169±9	32.6±1.4	29.8	35.5	107
22-30	3.1	161±9	30.4±1.4	27.6	33.2	114
32–40	3.8	158±9	29.4±1.4	26.6	32.2	118
42-50	4.6	158±9	29.4±1.4	26.6	32.1	118
52-60	6.2	153±8	28.1±1.4	25.3	30.8	124
62–70	7.4	151±8	27.3±1.4	24.6	30.1	127
72-80	9.1	148±7	26.5±1.4	23.8	29.2	131
82–90	9.8	146±6	26.0±1.4	23.3	28.7	134
92-100	11.8	146±6	26.0±1.4	23.3	28.7	134
102-110	12.7	145±6	25.6±1.4	22.9	28.3	136
112-120	10.8	144±6	25.2±1.4	22.5	27.9	138
122-130	7.2	143±5	25.0±1.4	22.3	27.7	139
132-140	4.2	$144 \pm 4$	25.3±1.4	22.6	28.0	138
142-150	2.0	141±6	24.4±1.4	21.7	27.1	143
152-160	1.0	137±5	23.2±1.4	20.4	26.0	150
162-170	0.5	133±13	22.2±1.5	19.2	25.2	157
172 +	0.6	137±12	23.3±1.9	19.6	27.0	149
All dives	100	149±7	26.8±1.4	24.1	29.6	130

Table 7. Mean heart rate (fH), mass-specific rate of oxygen consumption ( $\dot{V}_{O_2}$ ) with 95% confidence intervals and calculated aerobic dive limit (cADL) of macaroni penguins for dives of different durations

less clear, though little is known about the myctophid icefish on which the penguins feed and a reliance on visual foraging was suggested as the explanation for decreased diving at night (Green et al., 1998). Such a reliance on daylight for successful foraging has also been proposed in other penguin species feeding on a variety of prey in different locations (Wilson et al., 1993).

### Heart rate changes within dives

Fig. 4 shows the average change in heart rate associated with dives of 102-110 s duration. Heart rate during diving has been recorded previously in diving birds, but only within laboratory conditions (Butler and Woakes, 1979, 1984; Stephenson et al., 1986), semi-natural conditions (Culik, 1992; Kooyman et al., 1992b) or in the field at a lower resolution (Bevan et al., 1997, 2002). These studies showed similar patterns in the change of heart rate to those of the present study, with *f*H higher than the resting level before and after dives, then falling to a level close to or lower than the resting level during dives. Such a response is now widely accepted to be a trade-off between the 'classic dive response', which conserves oxygen stores while the animal is deprived of access to air, and the 'exercise response', which prioritises blood flow and oxygen uptake to active muscles when exercising (Butler, 1988).

In the present study, the mean NDfH was not significantly different from the mean heart rate during bouts of diving (DfH). It is not possible to state exactly what activities the penguins were engaged in when not diving, but it seems likely that they were travelling between the feeding sites and the colony. Swimming or porpoising while travelling is energetically more expensive than resting either in water or air (Culik and Wilson, 1991; Bevan et al., 1995b), and hence NDfH cannot necessarily be considered to be the *f*H while resting on water. In gentoo penguins, *f*H while resting on the water in a respirometer was the same as *f*H averaged over complete free-ranging dive cycles (Bevan et al., 1995b), and we have assumed that the same is true for macaroni penguins.

Adjustments in fH allow dive duration to be extended by ensuring full loading of oxygen stores before the dive, then by reducing aerobic metabolism during the dive (Butler and Jones, 1997) and ensuring the full and effective use of oxygen stores while submerged (Davis and Kanatous, 1999). Changes in heart rate, blood flow and perfusion during diving have been proposed ever since the early physiological experiments on forcibly submerged animals (Scholander, 1940) and have subsequently been observed in freely diving penguins (Millard et al., 1973) and other diving birds (Bevan and Butler, 1992). Data on these circulatory adjustments are limited (Kooyman and Ponganis, 1998), but they could have a very great effect on reducing aerobic metabolism and maximising the effective use of oxygen stores (Davis and Kanatous, 1999). The stepwise multiple linear regression showed that minimum fH had the strongest relationship to dive duration followed by minimum fH during the first 10 s of the dive and mean *f*H after the dive. Since the minimum heart rate occurs relatively early in the dive (Fig. 4), this might suggest that the penguins are to some extent setting the duration of the dive when the minimum fH is reached, though the importance of mean fH post-dive suggests that penguins adjust fH as a response to the previous dive rather than to prepare for the next one. This idea would contradict the apparent prediction of the duration and depth of the following dive and adjustment of the volume of inhaled air (Sato et al., 2002; Wilson et al., 2002) and clearly this subject requires further investigation. Currently the multiple regression analysis is instructive, but it is difficult to determine whether, within the penguin, dive duration is dependent on the cardiac and circulatory adjustments or *vice versa*. What can be stated with certainty is that in macaroni penguins, the cardiac adjustments become more exaggerated as dive duration increases.

# Rate of oxygen consumption during diving

Heart rate cannot be used to estimate  $\dot{V}_{O_2}$  while submerged. In tufted ducks Aythya fuligula, estimation of submerged  $\dot{V}_{O_2}$ using values for mean submerged fH at mean dive duration, actually underestimated mean submerged  $\dot{V}_{O_2}$  at mean dive duration, as calculated from a multiple linear regression (Woakes and Butler, 1983). However, if fH is averaged over complete dive cycles, then it is an accurate and reliable predictor of  $\dot{V}_{O_2}$  for the dive cycle (Fedak, 1986; Bevan et al., 1992; Butler, 1993). This approach was adopted in the present study and  $\dot{V}_{O_2}$  during dive cycles was estimated using mean fH recorded from completed dive cycles. If we assume that  $V_{O_2}$ while submerged is equivalent to this mean value, then it is possible to determine the cADL for macaroni penguins. As the observed dive duration increased,  $\dot{V}_{O_2}$  decreased and hence cADL increased (Table 7). For all dive durations up to 138s (95.3%) of dives), the cADL was greater than the observed dive duration (Fig. 5). The 95% confidence limits can also be used to calculate cADLs for the potential minimum and maximum estimates of  $\dot{V}_{O_2}$ . If the upper confidence limit is used, then for a given dive duration, cADL will be lower and only dives up to 126s (89.2% of dives) would be within the cADL. In contrast, at the lower confidence interval, for a given dive duration the cADL will be higher and all dives would be within the cADL. These results imply that most natural dives within diving bouts by macaroni penguins are aerobic.  $\dot{V}_{O_2}$ calculated from DfH of 147 beats min<sup>-1</sup> would be  $26.2\pm1.4$  ml min<sup>-1</sup> kg<sup>-1</sup>, with upper and lower confidence limits of 28.9 and 23.5 ml min<sup>-1</sup> kg<sup>-1</sup>, respectively. The resultant cADL would be 133 s with limits of 120-148 s. This would translate to 92.8% of observed dives being within the cADL with 95% confidence limits of 84.5-97.6%. This approach demonstrates the importance of including the variation in heart rate associated with dives of different durations. Calculating cADL at different durations suggests that 95.3% of observed dives used aerobic metabolism, whereas the more straightforward approach using overall mean DfH to calculate cADL suggests that only 92.8% of observed dives used aerobic metabolism.

cADL has been calculated using  $\dot{V}_{O_2}$  while resting on water for three other penguin species (Butler, 2000), though in each case,  $\dot{V}_{O_2}$  was measured using respirometry, rather than

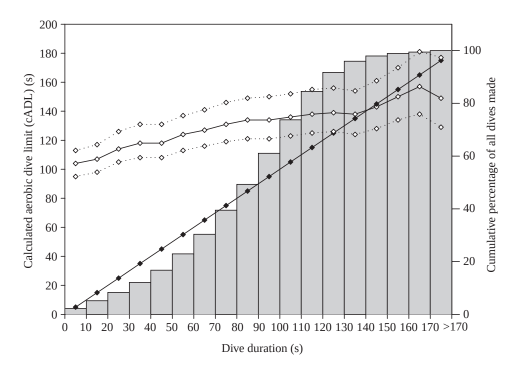


Fig. 5. Change in observed calculated aerobic dive limit (cADL, open diamonds, with 95% confidence limits) with increasing dive duration (grouped into 10s bins). The line marked by filled diamonds represents the line of equality where cADL equals dive duration. Dive durations (on the line of equality) above the observed cADLs probably involve an anaerobic component. Also shown are the cumulative percentage frequency of dive durations (grey bars).

estimated from the field. In emperor penguins, 96% of foraging dives in the field would be within the cADL, whereas in king penguins Aptenodytes patagonicus and gentoo penguins *Pygoscelis papua*, only 80% of dives in the field would be within the cADL. Given that the oxygen stores are assumed to be the same for these four species, there must be a difference in diving behaviour or  $\dot{V}_{O_2}$  while submerged between species. Food density, availability and location will fluctuate, causing variation in ecological conditions between populations and species, which are more likely to be the causes of variability in diving performance than differences in physiology. Ecological differences between gentoo and macaroni penguins breeding at Bird Island have been described previously (Croxall et al., 1997), and the breeding success of gentoo penguins is far more vulnerable than macaroni penguins to variations in their food availability (Croxall et al., 1999). Perhaps gentoo penguins are under greater pressure to gather enough food to provision their two chicks, leading to a higher proportion of anaerobic diving.

Similarly, emperor penguins are substantially larger than king penguins (approx. 25–30 and 10–15 kg, respectively) (Pütz et al., 1998), yet their diving performance is similar (Kooyman and Ponganis, 1990; Kooyman et al., 1992a; Kooyman and Kooyman, 1995). As would be expected, with their greater size and oxygen stores, emperor penguins are capable of superior maximum dive depth and duration than king penguins, but a large proportion of the foraging dives of both species are to 100–200 m depth and up to 5–6 min duration (Kooyman et al., 1992a; Kooyman and Kooyman, 1995). This implies that emperor penguins operate well within their physiological limits, whereas king penguins dive to depths and for durations that are close to the maximum of their capabilities.

# Abdominal temperature changes during dive bouts

Abdominal temperature showed a progressive decline during most dive bouts. Similar decreases in body temperature have been observed in other diving birds including king penguins (Culik et al., 1996b; Handrich et al., 1997), gentoo penguins (Bevan et al., 2002), king cormorants (Kato et al., 1996) and blue-eyed shags (Bevan et al., 1997) as well as in marine mammals (Hill et al., 1987). The mean decrease during a diving bout ( $\Delta T_{ab}$ ) in macaroni penguins was 2.32±0.2°C, similar to that in gentoo penguins of 2.6°C (Bevan et al., 2002). Mean  $\Delta T_{ab}$  was considerably less than the mean maximum  $\Delta T_{ab}$  of 13.5±1.1°C, as most individuals performed many short diving bouts where  $\Delta T_{ab}$  was low. This also explains why the mean  $T_{ab}$  during diving bouts was 4.7°C lower than the mean  $T_{ab}$  while not diving, as long bouts with large values of  $\Delta T_{ab}$ account for a large proportion of the time spent within diving bouts.

The decline in  $T_{ab}$  may be the inevitable consequence of the ingestion of cold food or of conduction to cold seawater from exposed surfaces on the feet and flippers. Local changes in circulation may effect the dissipation of heat from the abdominal region. Animals may attempt to reduce this heat loss or simply allow it to continue. Alternatively, in an effort intentionally to lose or 'dump' heat, animals may increase blood flow to the abdomen and/or exposed surfaces. These alternative mechanisms for heat loss, and determination of whether this an active or passive process, are still subject to investigation (Kooyman et al., 1980; Hill et al., 1987; Kooyman, 1989; Handrich et al., 1997; Ponganis et al., 2001). However, studies of the barnacle goose Branta leucopsis, a non-diving bird, have shown that it is possible for birds to experience anapyrexia (Cabanac and Brinnel, 1987), a resetting of their body temperature to a lower level when conservation of energy may be important, even if the animal is active and food is not scarce (Butler and Woakes, 2001).

Data from king penguins suggest that the decrease in  $T_{ab}$  is in some way facilitated and not just the consequence of ingesting cold food, as the  $T_{ab}$  of foraging king penguins was lower than that in the stomach (Handrich et al., 1997). It has been proposed that this reduction in  $T_{ab}$  leads to lowered metabolic rates in diving birds (Boyd and Croxall, 1996; Culik et al., 1996b; Butler, 2000), through the effect of cold temperatures on metabolically active tissues (Heldmaier and Ruf, 1992) and reduced thermoregulatory costs. Barnacle geese engaged in a long, energetically costly migration were found to allow their  $T_{ab}$  to fall progressively by 4.4°C, and it is proposed that if this hypothermia extended to the whole body, an amount of fat could be saved equivalent to up to 25% of that used for migration (Butler and Woakes, 2001). In diving birds, a lowering of  $T_{ab}$  and metabolic rate is suggested to be sufficient to bring most natural dives observed in the field within the cADL (Boyd and Croxall, 1996; Butler, 2000). This is not the only mechanism that might account for the discrepancies between observed diving behaviour and cADL. For example, phosphocreatine may be a source of energy that animals use while submerged (Butler and Jones, 1997) and further research into this possibility should be a priority.

In the present study, it was not possible to detect variation in  $T_{ab}$  within dives. In king penguins, fluctuations in temperature in localised parts of the body were found to vary between consecutive dives (Culik et al., 1996b). Similar experiments investigating changes in  $T_{ab}$  of emperor penguins diving from man-made holes in sea-ice using a thermistor with a much smaller time constant (0.2 s) showed that  $T_{ab}$  can drop quite considerably within individual dives (Ponganis et al., 2001). However, in the same study (Ponganis et al., 2001), another thermistor placed in the inferior vena cava, which receives blood drained from core organs such as the kidneys, liver and gastrointestinal tract, registered no significant changes in temperature during diving. The authors concluded that there was no evidence to suggest that reduction in  $T_{ab}$ facilitates diving durations greater than the cADL or DLT, as core temperature did not vary during diving and there was no relationship between the magnitude of  $T_{ab}$  fluctuation and dive duration. Further work, involving more sensitive and faster responding temperature sensors at multiple locations around the body, may cast more light on the extent of this regional hypothermia and its possible importance in extending dive durations in different species.

Though it was not possible to detect differences in  $T_{ab}$  within individual dives in the present study,  $T_{ab}$  did decline progressively during diving bouts. The shape and gradient of this temperature decline varied between individuals (which may be attributable to the position of the data logger) and between diving bouts performed by the same individual. However, in each case the decline was progressive throughout the bout, and abdominal temperature only increased after or at the very end of the bout. The magnitude of the temperature drop did increase consistently with the duration of diving bouts (Fig. 3). If diving

behaviour was determined only by physiological capacity, and lowered abdominal temperature was essential to facilitate increased diving duration, then we might expect to see dive duration increasing and/or mean *f*H decreasing progressively through bouts as abdominal temperature decreases. However, as Table 4 shows, nearly as many diving bouts showed a progressive decrease in dive duration during bouts as showed a progressive increase, and over 64% showed no significant change at all. In addition, nearly all dives were within the cADL. This supports the suggestion that, for macaroni penguins, factors other than physiological ones are likely to be more important in determining average diving behaviour. Such factors could include progressive satiation during dive bouts and the location and density of patches of food within the water column, especially since Antarctic krill are found in swarms (Everson, 2000). In gentoo and king penguins, which may be pushing the physiological limits of aerobic diving more than macaroni penguins, patterns of increasing dive duration within bouts might be observed.

The progressive decrease in  $T_{ab}$  of macaroni penguins is likely to be the result of many smaller decreases associated with individual dives. The abdomen may not have sufficient time to return to its initial temperature during the surface interval between dives, and the overall decrease in temperature may be the result of an accumulation of these cycles. This pattern was found to occur in diving emperor penguins (Ponganis et al., 2001) where  $T_{ab}$  started to decrease as soon as a dive commenced and continued to decrease until the animal surfaced. Upon surfacing,  $T_{ab}$  immediately increased until the next dive commenced. However, the increase while at the surface was not sufficient to match the decrease while diving and the net effect was a progressive decline in  $T_{ab}$ during diving bouts.

# Conclusions

The present study suggests that most dives by macaroni penguins are likely to be aerobic. Circulatory adjustments and the associated reduction of heart rate during dives permit a sufficiently low level of oxygen consumption such that even the longest observed dives performed by these animals may be supported by aerobic metabolism. Bouts of repeated diving are also associated with a reduction in abdominal temperature, which is probably a result of the accumulation of many smaller decreases during individual dive/surface cycles. Decreased temperature in the abdomen will further contribute to a reduction in metabolic rate, but further work would be required to determine the extent of cooling in the penguins' bodies and to what extent this might lead to a significant reduction in metabolic rate during dives.

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### References

- Bengston, J. L., Croll, D. A. and Goebel, M. E. (1993). Diving behaviour of chinstrap penguins at Seal Island. *Antarct. Sci.* 5, 9-15.
- Bethge, P., Nicol, S., Culik, B. M. and Wilson, R. P. (1997). Diving behaviour and energetics in breeding little penguins (*Eudyptula minor*). J. Zool., Lond. 242, 483-502.
- Bevan, R. M. and Butler, P. J. (1992). Cardiac output and blood flow distribution during swimming and voluntary diving of the tufted duck (*Athya fuligula*). J. Exp. Biol. 168, 199-217.
- Bevan, R. M., Boyd, I. L., Butler, P. J., Reid, K., Woakes, A. J. and Croxall, J. P. (1997). Heart rates and abdominal temperatures of freeranging South Georgian shags, *Phalocrocorax georgianus. J. Exp. Biol.* 200, 661-675.
- Bevan, R. M., Butler, P. J., Woakes, A. J. and Boyd, I. L. (2002). The energetics of gentoo penguins, *Pygoscelis papua*, during the breeding season. *Func. Ecol.* **16**, 175-190.
- Bevan, R. M., Butler, P. J., Woakes, A. J. and Prince, P. A. (1995a). The energy expenditure of free-ranging black-browed albatrosses. *Phil. Trans. R. Soc. Lond.* **350**, 119-131.
- Bevan, R. M., Keijer, E. and Butler, P. J. (1992). A method for controlling the feeding behaviour of aquatic birds: heart rate and oxygen consumption during dives of different duration. J. Exp. Biol. 162, 91-106.
- Bevan, R. M., Woakes, A. J., Butler, P. J. and Croxall, J. P. (1995b). Heart rate and oxygen consumption of exercising gentoo penguins. *Physiol. Zool.* **68**, 855-877.
- Boyd, I. L. and Croxall, J. P. (1996). Dive durations in pinnipeds and seabirds. Can. J. Zool. 74, 1696-1705.
- Boyd, I. L., Arnould, J. P. Y., Barton, T. and Croxall, J. P. (1994). Foraging behaviour of Antarctic fur seals during periods of contrasting prey abundance. J. Anim. Ecol. 63, 703-713.
- Butler, P. J. (1988). The exercise response and the 'classical' diving response during natural submersion in birds and mammals. *Can. J. Zool.* 66, 29-39.
- Butler, P. J. (1993). To what extent can heart rate be used as an indicator of metabolic rate in free-living marine mammals. *Symp. Zool. Soc. Lond.* 66, 317-332.
- Butler, P. J. (2000). Energetic costs of surface swimming and diving of birds. *Physiol. Biochem. Zool.* 73, 699-705.
- Butler, P. J. (2001). Diving Beyond the Limits. *News Physiol. Sci.* 16, 222-227.
- Butler, P. J. and Jones, D. R. (1997). Physiology of diving of birds and mammals. *Physiol. Rev.* 77, 837-899.
- Butler, P. J. and Woakes, A. J. (1979). Changes in heart rate and respiratory frequency during natural behaviour of ducks, with particular reference to diving. J. Exp. Biol. 79, 283-300.
- Butler, P. J. and Woakes, A. J. (1984). Heart rate and aerobic metabolism in humboldt penguins, *Spheniscus Humboldti*, during voluntary diving. *J. Exp. Biol.* **108**, 419-428.
- Butler, P. J. and Woakes, A. J. (2001). Seasonal hypothermia in a large migrating bird: saving energy for fat deposition? J. Exp. Biol. 204, 1361-1367.
- Cabanac, M. and Brinnel, H. (1987). The pathology of human temperature regulation: Thermiatrics. *Experientia* 43, 19-27.
- Carbone, C. and Houston, A. I. (1996). The optimal allocation of time over a dive cycle: An approach based on aerobic and anaerobic respiration. *Anim. Behav.* **51**, 1247-1255.
- Chappell, M. A., Shoemaker, V. H., Janes, D. N., Bucher, T. L. and Maloney, S. K. (1993). Diving behavior during foraging in breeding adélie penguins. *Ecology* 74, 1204-1215.
- Costa, D. P. (1988). Methods for studying the energetics of freely diving animals. *Can. J. Zool.* 66, 45-52.
- Croll, D. A., Gaston, A. J., Burger, A. E. and Konnoff, D. (1992). Foraging behaviour and physiological adaptation for diving in thick-billed murres. *Ecology* 73, 344-356.
- Croxall, J. P., Briggs, D. R., Kato, A., Naito, Y., Watanuki, Y. and Williams, T. D. (1993). Diving pattern and performance in the macaroni penguin *Eudyptes chrysolophus. J. Zool., Lond.* 230, 31-47.
- Croxall, J. P., Davis, R. W. and O'Connell, M. J. (1988). Diving patterns in relation to diet of gentoo and macaroni penguins at South Georgia. *Condor* 90, 157-167.
- Croxall, J. P., Prince, P. A. and Reid, K. (1997). Dietary segregation of krilleating South Georgian seabirds. J. Zool., Lond. 242, 531-536.
- Croxall, J. P., Reid, K. and Prince, P. A. (1999). Diet, provisioning and productivity responses of marine predators to differences in availability of Antarctic krill. *Mar. Ecol. Prog. Ser.* **177**, 115-131.

- Culik, B. M. (1992). Diving heart rates in Adélie penguins (Pygoscelis adeliae). Comp. Biochem. Physiol. 102, 487-490.
- Culik, B. M. and Wilson, R. P. (1991). Energetics of under-water swimming in Adélie penguins (*Pygoscelis adeliae*). J. Comp. Physiol. 161, 285-291.
- Culik, B. M., Pütz, K., Wilson, R. P., Allers, D., Lage, J., Bost, C.-A. and Le Maho, Y. (1996a). Diving energetics in king penguins (*Aptenodytes patagonicus*). J. Exp. Biol. 199, 973-983.
- Culik, B. M., Pütz, K., Wilson, R. P., Bost, C.-A., Le Maho, Y. and Verselin, J.-L. (1996b). Core temperature variability in diving king penguins (*Aptenoydytes patagonicus*) a preliminary analysis. *Polar Biol.* 16, 371-378.
- Culik, B. M., Wilson, R. P. and Bannasch, R. (1994). Underwater swimming at low energetic costs by pygoscelid penguins. J. Exp. Biol. 197, 65-78.
- Davis, R. W. and Kanatous, S. B. (1999). Convective oxygen transport and tissue oxygen consumption in weddell seals during aerobic dives. J. Exp. Biol. 202, 1091-1113.
- Davis, R. W., Croxall, J. P. and O'Connell, M. J. (1989). The reproductive energetics of gentoo (*Pygoscelis papua*) and macaroni (*Eudyptes chrysolophus*) penguins at South Georgia. J. Anim. Ecol. 58, 59-74.
- Davis, R. W., Kooyman, G. L. and Croxall, J. P. (1983). Water Flux and estimated metabolism of free-ranging gentoo and macaroni penguins at South Georgia. *Polar Biol.* 2, 41-46.
- **Everson, I.** (ed.) (2000). *Krill: Biology, Ecology and Fisheries*. Oxford: Blackwell Science. 384pp.
- Fedak, M. A. (1986). Diving and exercise in seals: a benthic perspective. In *Diving in Animals and Man* (ed. A. D. Brubakk, J. W. Kanwisher and G. Sundess), pp. 11-32. Trondheim: Tapir.
- Fedak, M. A., Pullen, M. R. and Kanwisher, J. (1988). Circulatory responses of seals to periodic breathing: heart rate and breathing during exercise and diving in the laboratory and open sea. *Can. J. Zool.* **66**, 53-60.
- Green, J. A., Butler, P. J., Woakes, A. J., Boyd, I. L. and Holder, R. L. (2001). Heart rate and rate of oxygen consumption of exercising macaroni penguins. *J. Exp. Biol.* **204**, 673-684.
- Green, K., Williams, R. and Green, M. G. (1998). Foraging ecology and diving behaviour of macaroni penguins *Eudyptes chrysolophus* at Heard Island. *Mar. Orn.* 26, 27-34.
- Handrich, Y., Bevan, R. M., Charrassin, J.-B., Butler, P. J., Pütz, K., Woakes, A. J., Lage, J. and Le Maho, Y. (1997). Hypothermia in foraging king penguins. *Nature* 388, 64-67.
- Heldmaier, G. and Ruf, T. (1992). Body temperature and metabolic rate during natural hypothermia in endotherms. J. Comp. Physiol. 162, 696-706.
- Hill, R. D., Schneider, R. C., Liggins, G. C., Schuette, A. H., Elliott, R. L., Guppy, M., Hochachka, P. W., Qvist, J. E., Falke, K. J. and Zapol, W. M. (1987). Heart rate and body temperature during free diving of Weddell seals. *Am. J. Physiol.* 253, R344-R351.
- Hull, C. L. (1997). The effect of carrying devices on breeding royal penguins. *Condor* **99**, 530-534.
- Kato, A., Naito, Y., Watanuki, Y. and Shaughnessy, P. D. (1996). Diving pattern and stomach temperature of foraging king cormorants at subantarctic Macquarie Island. *Condor* **98**, 844-848.
- Kooyman, G. L. (1989). Diverse Divers. Berlin: Springer-Verlag. 199pp.
- Kooyman, G. L. and Kooyman, T. G. (1995). Diving behavior of emperor penguins nurturing chicks at Coulman Island, Antarctica. *Condor* 97, 536-549.
- Kooyman, G. L. and Ponganis, P. J. (1990). Behavior and physiology of diving in emperor and king penguins. In *Penguin Biology* (ed. L. S. Davis and J. T. Darby), pp. 229-242. New York: Academic Press.
- Kooyman, G. L. and Ponganis, P. J. (1998). The physiological basis of diving to depth: Birds and mammals. *Annu. Rev. Physiol.* 60, 19-32.
- Kooyman, G. L., Castellini, M. A., Davis, R. W. and Maue, R. A. (1983). Aerobic dive limits in immature Weddell seals. J. Comp. Physiol. 138, 335-346.
- Kooyman, G. L., Cherel, Y., Le Maho, Y., Croxall, J. P., Thorson, P. H., Ridoux, V. and Kooyman, C. A. (1992a). Diving behavior and energetics during foraging cycles in king penguins. *Ecol. Monogr.* 62, 143-163.
- Kooyman, G. L., Ponganis, P. J., Castellini, M. A., Ponganis, E. P., Ponganis, K. V., Thorson, P. H., Eckert, S. A. and Le Maho, Y. (1992b). Heart rate and swim speeds of emperor penguins diving under sea ice. J. Exp. Biol. 165, 161-180.
- Kooyman, G. L., Wahrenbrock, E. A., Castellini, M. A., Davis, R. W. and Sinnett, E. E. (1980). Aerobic and anaerobic metabolism during voluntary diving in Weddell seals: evidence of preferred pathways from blood chemistry and behavior. J. Comp. Physiol. 138, 335-346.

- Millard, R. W., Johansen, K. and Milsom, W. K. (1973). Radiotelemetry of cardiovascular responses to exercise and diving in penguins. *Comp. Biochem. Physiol.* 46, 227-240.
- Mori, Y. (1998). The optimal patch use in divers: optimal time budget and the number of dive cycles during bout. J. Theor. Biol. 190, 187-199.
- Ponganis, P. J., Kooyman, G. L., Baranov, E. A., Thorson, P. H. and Stewart, B. S. (1997a). The aerobic submersion limit of Baikal seals, *Phoca sibirica. Can. J. Zool.* 75, 1323-1327.
- Ponganis, P. J., Kooyman, G. L., Starke, L. N., Kooyman, C. A. and Kooyman, T. G. (1997b). Post-dive blood lactate concentrations in emperor penguins, *Aptenodytes forsteri. J. Exp. Biol.* 200, 1623-1626.
- Ponganis, P. J., Kooyman, G. L., Winter, L. M. and Starke, L. N. (1997c). Heart rate and plasma lactate response during submerged swimming and trained diving in California sea lions, *Zapholus californianus*. J. Comp. Physiol. 197, 9-16.
- Ponganis, P. J., Van Dam, R. P., Knower, T. and Levenson, D. H. (2001). Temperature regulation in emperor penguins foraging under sea ice. *Comp. Biochem. Physiol.* 129, 811-820.
- Pütz, K., Wilson, R. P., Kierspel, M. A. M., Culik, B. M., Adelung, D., Charrassin, J.-B., Raclot, T., Lage, J. and Le Maho, Y. (1998). Foraging strategy of king penguins (*Aptenodytes patagonicus*) during summer at the Crozet Islands. *Ecology* 79, 1905-1921.
- Robertson, G. (1994). The foraging ecology of emperor penguins (*Aptenodytes forsteri*) at two Mawson Coast colonies, Antarctica. *ANARE report* **138**, 139pp.
- Sato, K., Naito, Y., Kato, A., Niizuma, Y., Watanuki, Y., Charrassin, J.-B., Bost, C.-A., Handrich, Y. and Le Maho, Y. (2002). Buoyancy and maximal diving depth in penguins: do they control inhaling air volume. *J. Exp. Biol.* 205, 1189-1197.
- Scholander, P. F. (1940). Experimental investigations on the respiratory function in diving mammals and birds. *Hval. Skr.* 22, 1-131.
- Stephenson, R., Butler, P. J. and Woakes, A. J. (1986). Diving behaviour and heart rate in tufted ducks *Aythya fuligula*. J. Exp. Biol. 126, 341-359.
- Stephenson, R., Turner, D. L. and Butler, P. J. (1989). The relationship between diving capacity and oxygen storage capacity in the tufted duck (Aythya fuligula). J. Exp. Biol. 141, 265-275.

- Williams, T. D. (1989). Agression, incubation behaviour and egg-loss in macaroni penguins, *Eudyptes chrysolophus*, at South Georgia. *Oikos* 55, 19-22.
- Williams, T. D. and Croxall, J. P. (1991). Annual variation in breeding biology of macaroni penguins, *Eudyptes chrysolophus*, at Bird Island, South Georgia. J. Zool., Lond. 223, 189-202.
- Williams, T. D., Briggs, D. R., Croxall, J. P., Naito, Y. and Kato, A. (1992). Diving pattern and performance in relation to foraging ecology in the gentoo penguin, *Pygoscelis papua*. J. Zool., Lond. 227, 211-230.
- Williams, T. M., Davis, R. W., Fuiman, L. A., Francis, J., Le Bouef, B. J., Horning, M., Calambokidis, J. and Croll, D. A. (2000). Sink or swim: strategies for cost-efficient diving by marine mammals. *Science* 288, 133-136.
- Williams, T. M., Haun, J. E. and Friedl, W. A. (1999). The diving physiology of bottlenose dolphins (*Tursiops truncatus*) I. Balancing the demands of exercise for energy conservation at depth. J. Exp. Biol. 202, 2739-2748.
- Wilson, R. P., Pütz, K., Bost, C. A., Culik, B. M., Bannasch, R., Reins, T. and Adelung, D. (1993). Diet dive depth in penguins in relation to diel vertical migration of prey: whose dinner by candlelight? *Mar. Ecol. Prog. Ser.* 94, 101-104.
- Wilson, R. P., Steinfurth, A., Ropert-Coudert, Y., Kato, A. and Kurita, M. (2002). Lip-reading in remote subjects: an attempt to quantify and separate ingestion, breathing and vocalisation in free-living animals using penguins as a model. *Mar. Biol.* 140, 17-27.
- Woakes, A. J. and Butler, P. J. (1983). Swimming and diving in tufted ducks, *Aythya fuligula*, with particular reference to heart rate and gas exchange. J. *Exp. Biol.* 107, 311-329.
- Woakes, A. J., Butler, P. J. and Bevan, R. M. (1995). Implantable data logging system for heart rate and body temperature: Its application to the estimation of field metabolic rates in Antarctic predators. *Med. Biol. Eng. Comput.* 33, 145-151.
- Ydenberg, R. C. and Clark, C. W. (1989). Aerobiosis and anaerobiosis during diving by western grebes: an optimal foraging approach. J. Theor. Biol. 139, 437-449.
- Zar, J. H. (1999). *Biostatistical Analysis*. New Jersey: Prentice-Hall International Inc. 718pp.