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This is a preprint of:

Gutiérrez, J.S. & Piersma, T. (2016). Ecological context determines the choice between prey of different salinities. *Behavioral Ecology*, 27(2), 530-537

Published version: dx.doi.org/10.1093/beheco/arv185

Link NIOZ Repository: www.vliz.be/nl/imis?module=ref&refid=251202

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3 1 **Ecological context determines the choice between prey of different**
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6 2 **salinity**
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14 4 **Lay summary**
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17 5 Seawater is too salty for most land animals, but many marine birds and reptiles can cope with it
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19 6 owing to flexible cephalic “salt” glands that excrete excess salt from the bloodstream. We show that
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21 7 red knots without access to freshwater prefer prey with relatively low salt content when their salt
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23 8 glands are small, but this preference is lost after they enlarge their salt glands and regain access to
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25 9 freshwater.
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31 11 **Summary**
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34 12 Food choice has profound implications for the relative intakes of water and salts, and thus for an
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36 13 animal’s physiological state. Discrimination behaviors with respect salt intake have been documented
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38 14 in a number of vertebrate species, but few studies have considered the ecological context in which
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40 15 they occur. Here, we report on the results of a two-choice experiment designed to examine the
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42 16 influence of dietary salt content and freshwater availability in food discrimination behaviors in red
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44 17 knots *Calidris canutus* (Aves: Scolopacidae) that feed on mud snails *Peringia ulvae* (Gastropoda:
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46 18 Hydrobiidae) whose body fluids have either relatively low (25‰) or high (42‰) salinity. Birds ate
47
48 19 more and spent longer time foraging on low-salinity mud snails when their salt gland sizes—an
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50 20 indicator of excretory capacity—were relatively small and when they were deprived of freshwater.
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52 21 However, as they enlarged salt glands—following a prolonged exposure to salty diet without access
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54 22 to freshwater—and regained access to freshwater their preference for low-salinity prey disappeared.
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3 23 Such a change of preference illustrates the context-dependency of discrimination. As the birds were
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5 24 able to maintain salt-water balance—inferred from plasma sodium concentration—under all
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7 25 conditions, changes in salinity preferences may occur without measurable physiological signs of
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9 26 osmotic stress. Our results highlight the importance of ecological context for understanding foraging
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11 27 responses. We argue that areas with high salinities could act as refuges for euryhaline invertebrates
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13 28 and fish from top vertebrate predators.
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For Review Only

30 INTRODUCTION

31 Seawater is toxic to most terrestrial vertebrates due to its high salt content. However, many
32 secondarily marine vertebrates such as snakes, turtles, birds and mammals live in marine and
33 estuarine environments where they typically feed on food that is in osmotic and ionic equilibrium
34 with the surrounding water (Schmidt-Nielsen 1997; McNab 2002). To cope with the excess salt and to
35 maintain fluid homeostasis, these animals possess specialized organs (e.g. reticulate kidneys,
36 cephalic 'salt' glands, gills or urinary bladder) that can adjust in size and/or function to cope with
37 changes in environmental salinity (Peaker and Linzell 1975; Hildebrandt 2001; Ortiz 2001; Bentley
38 2002; McNab 2002). Among them, cephalic salt glands are one of the best-documented examples of
39 physiological adaptation to marine life in non-mammalian vertebrates. Most birds and reptiles from
40 marine environments have cephalic salt glands that extract salt ions from the bloodstream,
41 producing a highly concentrated salt solution that is discarded through ducts that open into the
42 nostrils (birds and lizards), eye (turtles), or tongue (snakes and crocodiles) (Peaker and Linzell 1975;
43 Schmidt-Nielsen 1997; Bentley 2002; McNab 2002). This affords them the capacity to eat salty food
44 and retain osmotically-free water (Schmidt-Nielsen 1960; Peaker and Linzell 1975; Schmidt-Nielsen
45 1997; McNab 2002).

46 Although various facets of vertebrate osmoregulation have been investigated exhaustively
47 (Peaker and Linzell 1975; Skadhauge 1981; Schmidt-Nielsen 1997; Goldstein and Skadhauge 2000;
48 Ortiz 2001; Bentley 2002), behavioral mechanisms leading to a decrease in salt intake have received
49 only limited attention (Wolcott and Wolcott 2001; Brischoux et al. 2012; Gutiérrez 2014). For
50 instance, it has been suggested that toothed whales whose diet consist mainly of hyperosmotic prey
51 (osmoconforming invertebrates) derive a 'water bonus' by also eating (the osmoregulating) bony fish
52 whose osmotic concentration resembles their own (Wolcott and Wolcott 2001). Likewise, it was
53 recently found that captive Australian pelicans *Pelecanus conspicillatus* consumed pieces of
54 elasmobranchs and squid (both osmoconformers) at substantially lower frequencies than bony fish
55 (osmoregulators) (Troup and Dutka 2014). Moreover, coastal ducks *Aythya* spp. that forage in

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3 56 energy-rich and salty estuaries regularly move to inland freshwater ponds to rest and re-hydrate
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5 57 (Woodin 1994; Adair et al. 1996). In reptiles, it has been suggested that the abilities of sea kraits
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7 58 *Laticauda* spp. to acquire fresh water on land and tolerate dehydration at sea, determine their
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9 59 environmental tolerances and geographic distributions (Brischoux et al. 2013). Clearly, behavioral
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11 60 osmoregulation plays a large part in the maintenance of the osmotic balance in many marine and
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13 61 estuarine air-breathing vertebrates.

16 62 Shorebirds (Charadriiformes, suborders Charadrii and Scolopaci) provide excellent material to
17
18 63 investigate how osmotic concentration of prey affects food discrimination behaviors in different
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20 64 environmental contexts. In estuarine and intertidal environments, both shorebirds and their prey
21
22 65 may be subjected to abrupt changes in the osmotic environment. For these organisms, fast and
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24 66 flexible behavioral responses are essential in meeting osmotic challenges (Gutiérrez et al. 2011;
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26 67 Gutiérrez et al. 2012; Gutiérrez et al. 2013; Gutiérrez 2014; Gutiérrez et al. 2015). In particular,
27
28 68 sandpipers of the genus *Calidris* have extensive arrays of taste buds (Gerritsen et al. 1982; Nebel et
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30 69 al. 2005), and several species (red knots *C. canutus*, purple sandpipers *C. maritima*, sanderlings *C.*
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32 70 *alba*, and dunlin *C. alpina*) can discriminate between 'clean' sand and sand that had contained prey,
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34 71 which suggests that they are able to use taste substances excreted by a particular prey for food
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36 72 detection (Gerritsen et al. 1982; van Heezik et al. 1983). For these reasons it seems plausible that
37
38 73 they can use the salinity of prey and surrounding water to adjust their salt intake and avoid osmotic
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40 74 stress. Indeed, NaCl-sensitive taste buds found in chickens, pigeons and parrots react to 0.2 M and
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42 75 higher concentrations of NaCl (Kitchell et al. 1959; Duncan 1962; Matson et al. 2000). A recent study
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44 76 based on relevant gene sequences associated with taste buds showed that penguins (order
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46 77 Sphenisciformes) have evolutionarily lost receptors for detecting sweet, umami, and bitter tastes,
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48 78 but still possess those for detecting salty tastes (Zhao et al. 2015); this would enable them to adjust
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50 79 their salt intake.
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3 80 In this study, we investigated whether prey salt content and freshwater accessibility influence
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5 81 food-discrimination behaviors using captive red knots that fed on mud snails *Peringia ulvae* whose
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7 82 body fluids had either relatively low (25‰) or high (42‰) salinity. Molluscivore shorebirds face the
8
9 83 dilemma of having to conserve free water while consuming hard-shelled prey with high seawater
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11 84 content and relatively little flesh (Gutiérrez et al. 2012; Gutiérrez et al. 2015). Specifically,
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13 85 maintaining the osmotic balance is a major challenge for red knots, as they may process several
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15 86 times their body mass in seawater each day (Visser et al. 2000) with limited or no access to
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17 87 freshwater in some of their main nonbreeding areas (Wolff and Smit 1990; van de Kam et al. 2004).
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19 88 To establish whether birds really display foraging preferences, and whether these depend on the
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21 89 ecological context, birds were simultaneously offered low-salinity and high-salinity diets with and
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23 90 without previous access to freshwater. Then, the choice between diets was recorded as the food
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25 91 intake and time spent foraging from each diet. To assess whether their choice pattern was related to
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27 92 their physiology, we also measured different indices of osmoregulatory state. (i) Hematocrit is known
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29 93 to increase with dehydration in birds (Hannam et al. 2003; Fair et al. 2007) and might indicate
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31 94 whether our treatments affected hydration state. (ii) Plasma sodium concentration is another
32
33 95 extensively studied hydration state parameter that serves as a good indicator of salt-water balance
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35 96 (Skadhauge 1981). (iii) The size of the salt glands positively correlates with the concentration and
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37 97 rate of their secretion, which in turn determines the amount of osmotically-free water they can
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39 98 retain for other physiological processes (Schmidt-Nielsen 1960; Staaland 1967).

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44 99 We predicted that birds would prefer the low-salinity diet over the high-salinity diet to minimize
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46 100 salt intake and avoid osmotic stress; our null hypothesis was a lack of preference. Additionally, we
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48 101 predicted that preference for low-salinity food would be stronger when birds have small salt glands,
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50 102 and when they are deprived of freshwater since under such conditions the birds would not be able to
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52 103 deliberately 'dilute' dietary salt; our null hypotheses would be lack of differences.
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105 MATERIALS AND METHODS

106 Subjects and Housing

107 Eight adult (four male and four female) red knots of the *islandica* subspecies (Nebel et al. 2000) were
108 caught in the western Dutch Wadden Sea (53°31'N, 6°23'E) in August-September 2013 and kept in
109 outdoor free-flight aviaries (4.5 m × 1.5 m surface × 2.5 m height) with unlimited access to trout
110 pellets (Vézina et al. 2006; Gutiérrez et al. 2015). Birds had free access to a freshwater tray (60 cm x
111 40 cm surface x 5 cm height) for drinking and an artificial mudflat flooded with running seawater for
112 probing. The floor of the aviaries was also flushed with running seawater to help prevent infections
113 and skin lesions caused by dry feet (see Milot et al. 2014). In January 2015, birds were transferred to
114 two separate indoor 'group' aviaries (4 birds per aviary) with similar characteristics to the outdoor
115 aviaries and fed a diet composed exclusively of 2-4 mm mud snails *Peringia ulvae* collected by
116 dredging in the Wadden Sea (Vézina et al. 2006; Gutiérrez et al. 2015). Outside the experiments, mud
117 snails were presented to the birds in two trays (60 cm x 40 cm surface x 5 cm height) with running
118 seawater taken directly from the sea (salinity ≈25‰; temperature ≈ 12°C).

119 In the intertidal zone, mud snails frequently dominate the benthic fauna numerically and in terms
120 of biomass, and form an important constituent of the diet of shorebirds (Evans et al. 1979; Britton
121 1985; van Gils et al. 2003). Indeed, this gastropod species is one of the main prey for red knots along
122 the East Atlantic flyway (Moreira 1994; van Gils et al. 2003; van den Hout 2010). *P. ulvae* can live in a
123 salinity range of 6–85‰ (Komendantov and Smurov 2009) and is isosmotic to seawater (Todd 1964),
124 meaning that red knots inevitably consume large amounts of salt when they ingest *P. ulvae* whole
125 (Gutiérrez et al. 2015). Captive red knots were kept in these indoor aviaries with water available
126 every day (salinity depending on the sessions; see below). The housing conditions were maintained
127 under a 10:14 light-dark cycle with a 20-min period of dawn:dusk ramp, similar to ambient conditions
128 during this period, and under indoor ambient temperature (12±0.5 °C). After the experiment, the
129 birds were released at the same site from which they were caught.

130 **Preparation of prey**

131 Freshly collected mud snails were stored frozen. Frozen mud snails remained in their shells, so the
132 birds had to crush the shells in their gizzard in order to digest the flesh (Vézina et al. 2006; Gutiérrez
133 et al. 2015). Unlike alive mud snails, dead (frozen-thawed) mud snails cannot moderate their
134 exposure to variation in salinity by withdrawing and closing their operculum (Berger and Kharazova
135 1997). Observations on dead snails from the stock offered to the birds showed that most (96%; N =
136 200) individuals had opened or lost their operculum when presented to birds. Therefore, we could
137 easily modify the salt concentration of their body fluids (Gutiérrez et al. 2015). To do this, freshly
138 (thawed) portions were placed in 90-L plastic containers with seawater of high ($41.51 \pm 0.22\%$, N =
139 18) or low ($24.92 \pm 0.48\%$, N = 18) salinity and maintained at $12 \pm 0.5^\circ\text{C}$ for approximately 12 h,
140 which ensured that snails had enough time to become isosmotic with the surrounding seawater (see
141 Supplementary Figure S1). At the end of this period, snails were removed from their tanks and visible
142 water was removed using a sieve (1 mm mesh).

143 Throughout the experiment the body water content of high-salinity ($50.86 \pm 0.23\%$) and low-
144 salinity ($51.22 \pm 0.28\%$) snails was similar (paired-*t* test: $t_{71} = -1.20$, $P = 0.23$; Supplementary Figure
145 S2). Water salinity was measured in the tanks daily with a portable multi-parameter instrument
146 (Delta Ohm, HD2156.1, Benelux B. V.). The salt concentration of snails' body fluids was determined
147 by inductively coupled plasma mass spectrometry (Thermo Scientific iCAP Q ICP-MS, Thermo Fisher
148 Scientific GmbH, Bremen, Germany) following Gutiérrez et al. (2015).

149 **Experimental protocol**

150 After three weeks of acclimation to mud snails, which ensures that red knots had enough time to
151 adjust to a diet of hard-shelled mollusc prey (Piersma et al. 1993), we moved on to the experimental
152 sessions. The phase consisted of three experimental sets on the basis of freshwater availability,
153 starting with 'access to freshwater' (sessions 1–2), followed by 'no access to freshwater' (sessions 3–
154 6) and finally 're-access to freshwater' (sessions 6–9). This sequence was used to manipulate the size

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3 155 of the glands (they are enlarged in birds fed salty diets and deprived of freshwater; Gutiérrez et al.
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5 156 2015, JSG pers. obs.) and thus ensured that birds encountered different environmental contexts (i.e.,
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7 157 access/no access to freshwater) with a range of salt gland sizes. As it was logistically impossible to
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9 158 measure and record all the individuals simultaneously, each individual was given one session every
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11 159 other day; therefore, the experimental period lasted 18 consecutive days. Birds were starved
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13 160 overnight before each experimental session (i.e., every other day) to get them motivated and eager
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15 161 to eat. Despite this regular fasting period, birds maintained a constant body mass throughout the
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17 162 experiment (Supplementary Figure S3). To avoid repetitive blood sampling and its potential effects
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19 163 on hematocrit and plasma sodium, we only took blood samples at the end of each experimental set
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21 164 (i.e., sessions 2, 6 and 9). Seawater (salinity $\approx 25\%$; temperature $\approx 12^\circ\text{C}$) was available at all times
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23 165 throughout the study, except during the 3-h experimental sessions; freshwater trays were available
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25 166 during both the pre-experimental period and the 'access to freshwater' and 're-access to freshwater'
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27 167 experimental sets (see above) when not in experimental procedures.
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32 168 Every day of the 18-day experimental period, we removed four birds from of one of the two
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34 169 indoor 'group' aviaries just before the start of the trials at 10:00 hours, weighed them (to the nearest
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36 170 0.1 g) and scored their salt glands (Gutiérrez et al. 2015; see below). Then, we transferred each bird
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38 171 to identical indoor 'individual' aviaries (same characteristics as the indoor 'group' aviaries) where two
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40 172 trays containing the same amount (c. 200 g) of low- and high-salinity mud snails were offered (Fig.
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42 173 1a, b). Wet snails were offered in excess in identical plastic trays with no water to prevent birds from
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44 174 making a choice based only on water salinity without tasting the prey (Fig. 1a, b). It is important to
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46 175 note a similar situation can be encountered in the wild when red knots intercept prey near the
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48 176 receding water line of mudflats.
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52 177 After each session, birds were returned to their indoor 'group' aviary and the food trays were
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54 178 removed and reweighed to determine the amount of food eaten (see below). In order to avoid the
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56 179 presence of potential visual and olfactory cues, the trays and floor of the aviaries were thoroughly
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3 180 cleaned after each trial. Moreover, the trays were reversed on a daily basis to avoid position-effect
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5 181 biases. All sessions were videotaped to code birds' behavior (see below).
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8 182 **Food intake**

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10 183 We measured food intake over each 3-hr session. Using food intakes in indoor aviaries during the
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12 184 pre-experimental period, we estimated average daily food intake at c. 250 g of wet mud snails per
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14 185 bird —this crude estimate represents the average food intake consumed in the 'group' aviaries per
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16 186 day divided by the number of birds in the aviaries. To minimize food depletion-related issues, birds
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18 187 were provided with 200 g of wet mud snails in each tray, which represents eight times the average
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20 188 amount of food a single bird would eat assuming a constant intake rate from only one tray (31.25 g;
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22 189 estimated as 250 g of daily food intake divided by 24 h of food access and multiplied by 3 h of
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24 190 experimental session). Every day, we sieved freshly thawed mud snails to remove all visible water
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26 191 and we then took two subsamples (10 g each) of food from this stock. In each session, we gave a pre-
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28 192 weighed amount of food from the same stock to the birds in the two trays. After the 3-hr session, the
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30 193 birds were returned to their 'group' aviaries and the food trays were removed from the aviaries.
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32 194 Droppings were carefully removed from the trays if present and mud snails adhered to them were
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34 195 separated and returned to their respective trays before weighing them for the second time to
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36 196 calculate food intake. Control (uneaten) portions of diet, weighed before and after trials, showed
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38 197 that water loss was negligible (on average 1%) and did not differ by diet (paired-*t* test: $t_{13} = -0.38$, $P =$
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40 198 0.71). Nonetheless, we corrected for water losses because even such small mass losses can bias the
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42 199 results (i.e. with respect to the response ratios and relative food intake rates).
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48 200 Birds occasionally fed onto the feeding trays (instead of walking around and taking food from
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50 201 them) and kicked out some snails from the tray onto the aviary floor —after correcting for water
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52 202 losses they consumed an average of 25.37 ± 2.22 g of food per session and spilled only 0.21 ± 0.09 g
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54 203 (0.84%). We assumed that snails spilled on the aviary floor belonged to the tray placed on that half of
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56 204 the aviary. Video recordings corroborated that birds did not transported food from one half of the
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3 205 aviary to the other. Therefore, we feel confident that food transport bias did not affect the results of
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5 206 this study.

207 **Hematocrit and plasma sodium concentration**

208 At the end of each set of trials, blood was taken from a wing vein into two heparinized capillary tubes
209 (75 μ l) per bird and centrifuged immediately for 10 min at 10,000 rpm. Hematocrits were read
210 immediately after centrifugation using a microhematocrit capillary tube reader. The value reported
211 herein for each bird was the mean of the two tubes. Plasma was saved to determine sodium
212 concentration, which was determined by inductively coupled plasma mass spectrometry (Thermo
213 Scientific iCAP Q ICP-MS, Thermo Fisher Scientific GmbH, Bremen, Germany) using a standardized
214 procedure (Long and Vetter 2002). All samples were collected after food deprivation to ensure a
215 post-absorptive condition.

216 **Salt gland scores**

217 We estimated salt gland scores for each individual using sensory evaluation (Gutiérrez et al. 2015).
218 Briefly, we scored the thickness of the salt glands at the postorbital ridge by sliding a finger across a a
219 smooth polyvinylchloride plate prepared with five increasing thicknesses (0–0.8 mm) at regular
220 distances from each other, to then compare these thicknesses with those of the postorbital salt gland
221 ridge.

222 **Videotaping**

223 Videocameras were placed outside the cages and were focused through a one-way mirror so that so
224 that they did not interfere with the birds' activity (Fig. 1). We coded behavior using the software
225 CowLog 2.0 (Hänninen and Pastell 2009). Behavior of each animal was categorized into foraging
226 (from left or right tray), moving (i.e., walking and flying), and resting (i.e., standing, sleeping and
227 preening). We then calculated the frequencies, bout durations, and total durations of the coded
228 behaviors. Finally, we calculated the proportion of time each bird spent foraging from each tray and

229 also noted which tray was visited first. Videos were examined by one person (J.S.G.) who was blind to
 230 the position (left/right) of the low-salinity and high-salinity trays.

231 **Statistical analyses**

232 Data were analyzed using linear mixed models (package 'nlme') in R (Team 2013). In choice trials, the
 233 intake of the two diets may not be independent, so response ratios of individual birds were used as a
 234 measure of preference (Martin and Bateson 1983). Response ratios were calculated as:

$$235 \frac{\text{amount of high-salinity snails eaten}}{\text{amount of high-salinity snails eaten} + \text{amount of low-salinity snails eaten}}$$

236
 237 If an individual only ate the high-salinity mud snails, its response ratio would be 1.0; conversely, if it
 238 only ate the low-salinity mud snails, its score would be 0.0. The chance level of response is 0.5. Linear
 239 mixed models were then performed on the response ratio to analyze whether individuals
 240 differentiated between, and showed any preference for, low-salinity or high-salinity diets. The
 241 responses ratio was also calculated for the time foraging in the two diets. The response ratio
 242 (proportional non-binomial data) was logit-transformed prior to analyses in order to fulfill linear
 243 assumptions (Warton and Hui 2010). Freshwater availability (access/no access) was included in the
 244 model as a fixed factor, salt gland scores were included as a covariate, and individual and session
 245 were included in the model as random factors. Body mass remained stable during the experiment
 246 (session effect: $t_{49} = -0.517$, $P = 0.61$; Supplementary Figure S3), so we did not consider it as a
 247 covariate. We always started with the full model and simplified it using backwards elimination based
 248 on ANOVA test with $P < 0.05$ as the selection criterion until reaching the minimal adequate model.
 249 Model assumptions were checked using the residuals of the final model. In addition, we used paired
 250 t -tests to test whether the mean difference in food consumption was significantly different between
 251 diets.

252 We performed linear regressions to explore relationships between food intake and foraging time
 253 (both for overall and diet-specific intakes). To test for potential diet-specific differences in food

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3 254 intake rate (g of mud snails min⁻¹), we also performed a linear mixed model with food intake rate as a
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5 255 response, diet and freshwater access as factors, and individual and session as random factors.
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8 256 We further explore the relationship between response ratios and salt gland scores. To do this, we
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10 257 first considered all data pooled together and then separately pooled by sets on the basis of
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12 258 freshwater availability and time: 'access to freshwater' (sessions 1-2), 'no access to freshwater'
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14 259 (sessions 3-6) and 're-access to freshwater' (sessions 7-9). We distinguished between the two sets
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16 260 with access to freshwater as previous experience with saline treatments could affect osmoregulatory
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18 261 abilities (e.g. salt gland size; see below) and yield different outcomes (Gutiérrez et al. 2011). Potential
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20 262 differences in hematocrit and plasma ion concentration recorded at the end of each set of sessions
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22 263 (sessions 2, 6 and 9) were examined using repeated measures analyses with hematocrit or plasma
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24 264 osmolality as response, session as a fixed factor, and individual as a random effect.
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28 265 One individual refused to eat during all the experimental sessions and was excluded from food
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30 266 preference analyses. In addition, six cases where another bird (always the same individual) refused to
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32 267 eat were excluded from these analyses.
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35 268 **RESULTS**

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38 269 Food intake was positively correlated with foraging time, both when considering overall intake and
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40 270 time ($F_{1,55} = 64.52$, $P < 0.001$; Fig. 2a) and when they were pooled by diets (high salinity: $F_{1,55} = 78.69$,
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42 271 $P < 0.001$; low salinity: $F_{1,55} = 104.80$, $P < 0.001$; Fig. 2b). Diet salinity had no significant effect on food
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44 272 intake rate ($F_{1,56} = 2.61$, $P = 0.11$; see Fig. 2b), indicating that birds ate high-salinity and low-salinity
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46 273 prey just as fast.
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49 274 Salt gland scores had significant effects in the response ratio of both food intake and foraging
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51 275 time (Table 1a, b; see also Fig. 3), whereas freshwater availability only marginally affected food
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53 276 intake (Table 1b; see also Fig. 3). Salt gland scores did not interact significantly with freshwater
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3 277 availability and thus their interaction was not included in the minimum adequate models (Table 1a,
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5 278 b).

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8 279 When considering all data pooled, the relationship between food intake's response ratio and salt
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10 280 gland scores was not significant ($F_{1,55} = 2.69$, $P = 0.107$). Neither did we find any significant
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12 281 relationship between these traits for the periods of 'access to freshwater' ($F_{1,12} = 0.68$, $P = 0.424$) or
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14 282 'no access to freshwater' ($F_{1,23} = 0.001$, $P = 0.98$). However, food intake's response ratio and salt
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16 283 gland scores positively correlated during the 're-access to freshwater' period ($F_{1,16} = 15.77$, $P = 0.001$),
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18 284 meaning that birds with higher salt gland scores showed a lower preference for high-salinity diet.

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21 285 Hematocrit did not differ between experimental sets (access to freshwater = $52.08 \pm 0.67\%$; no
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23 286 access to freshwater = $51.88 \pm 0.86\%$; and re-access to freshwater = $50.56 \pm 0.42\%$; set effect: $F_{2,14} =$
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25 287 2.18 , $P = 0.15$). Neither did we find any significant changes of plasma sodium concentration (access
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27 288 to freshwater = 155.36 ± 5.36 mmol L⁻¹; no access to freshwater = 161.36 ± 4.37 mmol L⁻¹; and re-
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29 289 access to freshwater = 156.41 ± 3.20 mmol L⁻¹; treatment effect: $F_{2,14} = 0.54$, $P = 0.59$). In addition, we
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31 290 found no correlation between these two blood parameters and salt gland scores either when data
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33 291 were pooled or when data were analyzed for each of the sets separately (always $P > 0.45$).

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38 39 40 293 **DISCUSSION**

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44 294 Our study demonstrates that red knots prefer prey with relatively low salt content when their salt
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46 295 glands are small (following a prolonged access to freshwater) and when they have no access to
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48 296 freshwater. This preference is lost after they enlarge their salt glands (following a prolonged
49
50 297 exposure to salty diet without access to freshwater) and regain access to freshwater. This finding is
51
52 298 consistent with the notion that behavior is dependent upon an animal's state (Houston and
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54 299 McNamara 1999), and that foraging responses are context-specific (Hurly and Oseen 1999; Chatelain
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56 300 et al. 2013; Halpin et al. 2014).

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3 301 Salt glands are the major organs for salt excretion in many birds and reptiles (Peaker and Linzell
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5 302 1975), including red knots (Staaland 1967; Gutiérrez et al. 2012). There is ample evidence that their
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7 303 size and activity change over short time-spans (Peaker and Linzell 1975; Shuttleworth and
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9 304 Hildebrandt 1999; Hildebrandt 2001; Gutiérrez et al. 2015). These rapid and flexible changes in the
10
11 305 salt glands could be correlated with short-term changes in behavior. The negative relationship
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13 306 between preference for low-salinity prey (i.e. food intake's response ratio) and salt gland scores
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15 307 observed at the end of the experiment indicates that birds became less selective; that is, birds with
16
17 308 larger salt glands, and thus higher concentrating ability (Schmidt-Nielsen 1960; Staaland 1967),
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19 309 consumed more high-salinity prey than birds with smaller salt glands. However, as larger salt glands
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21 310 require larger maintenance costs, these should also increase with salt gland size (Gutiérrez et al.
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23 311 2011). Why then did birds not minimize energy expenditure by choosing low-salinity prey under all
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25 312 conditions?
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29 313 We can think of two explanations. On one hand, captive birds with nearly unlimited access to food
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31 314 and freshwater might have offset osmoregulatory costs with no difficulty. This would explain why
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33 315 birds of several species manifested indifference to saline solutions at low concentrations in two-
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35 316 bottle drinking preference tests conducted after unlimited access to freshwater (Harriman and Kare
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37 317 1966; Harriman 1967). A second, non-exclusive, explanation is related to potential trade-offs with
38
39 318 osmoregulation; that is, higher salt loads result in larger osmoregulatory costs but also more efficient
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41 319 salt glands, so eating high-salinity prey could protect individuals from short-term physiological costs
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43 320 under variable osmotic environments (Gutiérrez 2014). This could partly explain why some pelagic
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45 321 seabirds show preference for saltwater over freshwater (Harriman and Kare 1966). In any case, birds
46
47 322 did not show signs of osmotic stress or dehydration during the experiment. Both the hematocrit and
48
49 323 plasma sodium concentration values reported varied little and sit comfortably within the range of
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51 324 values expected for mollusk-eating captive red knots (Piersma et al. 2000; Gutiérrez et al. 2015).
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3 325 Interestingly, although red knots are able to quickly develop a preference for the less salty diet
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5 326 when they had to (i.e., when they had small salt glands and no freshwater source), they normally
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7 327 continue probing the high-salinity diet (Supplementary Figure S4). This could be interpreted either as
8
9 328 rapid forgetting of food information or as natural inclination to make strategic 'mistakes' to explore
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11 329 and verify whether alternative reward rules have come into fashion (Piersma et al. 1998). Because
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13 330 probe-feeding red knots mainly feed in tight flocks on notoriously variable and patchy intertidal flats
14
15 331 (van Gils et al. 2005; van Gils et al. 2006), they are expected to share public information about
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17 332 resource quality (van Gils et al. 2006; Bijleveld et al. 2015) rather than to remember the precise
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19 333 locations within intermittently available patches.
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23 334 These findings suggest that free-ranging animals experiencing varying salinities can use
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25 335 discriminatory behaviors to adjust salt intake. For instance, they may select among
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27 336 microenvironments differing, spatially or temporally, in osmotic characteristics. Where osmotic
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29 337 characteristics of food are spatially variable, food-selection and handling behaviors can contribute to
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31 338 osmoregulation by maximizing input of required water and/or minimizing salt intake (Mahoney and
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33 339 Jehl 1985; Nyström and Pehrsson 1988; Brischoux et al. 2013; Troup and Dutka 2014). Moreover,
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35 340 they may exploit diel differences in water potential, restricting activity (e.g. foraging) to times when
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37 341 temperature is lowest (Zwarts et al. 1990). Such behaviors could be especially important for many
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39 342 bird species, including the red knot, that spend the winter in (sub)tropical intertidal sites without
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41 343 regular access to freshwater (Wolff and Smit 1990; van de Kam et al. 2004). Likewise, coping with salt
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43 344 may become particularly severe for tropical marine snakes during (and following) periods of high
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45 345 oceanic salinity with very limited access to freshwater (Brischoux et al. 2012; Brischoux et al. 2013) as
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47 346 well as for other marine and estuarine reptiles (e.g. turtles and crocodiles) that rely on the extraction
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49 347 of osmotically-free water (*via* salt glands) from food items of relatively low salt content or on
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51 348 periodic access to fresh or brackish drinking water (Schmidt-Nielsen and Fänge 1958; Taplin and
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53 349 Grigg 1981; Mazzotti and Dunson 1989; Cramp et al. 2008). Under these circumstances, behavioral
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55 350 osmoregulation may be crucial to maintaining osmotic balance.
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3 351 In addition, salinity has important direct and indirect effects on habitat structure and predator-
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5 352 prey interactions in aquatic systems (Ysebaert et al. 2000; Herbst 2001; Ravenscroft and Beardall
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7 353 2003; Ysebaert et al. 2003). Though not without cost, more salt-tolerant invertebrates and fish might
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9 354 escape potential predators by using areas of high salinity. One might expect that, if birds and other
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11 355 secondarily marine air-breathing vertebrates avoid areas of high salinity, potential prey would reduce
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13 356 predation risk in what effectively would be saline 'refuges'. In this vein, it has been suggested that
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15 357 red knots do not feed extensively on brine shrimps *Artemia* spp. at supratidal salinas (salinity: 100–
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17 358 150‰) to avoid of osmotic stress (Masero 2002). Thus, it is plausible that selection of saline refuges
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19 359 by euryhaline species enable individuals to better survive than individuals at lower salinities but
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21 360 higher predation risk. Ultimately, osmoregulatory costs may affect selection pressures acting on both
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23 361 predators and prey. Reduced salinity tolerance at high ambient temperatures has been reported in
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25 362 red knots feeding on mud snails (Gutiérrez et al. 2015); this could lead to selection for less salty prey
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27 363 in environments where osmoregulatory costs would increase substantially: in warm climates.
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31 364 In summary, discrimination behaviors with respect salt intake are a function of ecological context
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33 365 and physiological state, meaning that the decisions that birds make when they are osmotically
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35 366 challenged will be different from when they have an efficient osmoregulatory machinery and/or
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37 367 access to freshwater. We suggest that under osmotically stressful environments dietary salt may act
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39 368 as discriminative stimuli for foraging responses in birds and other secondarily marine vertebrates
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41 369 such as snakes, turtles, birds and mammals. Studies investigating how foraging decisions change with
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43 370 salinity and temperature should help us understand how climate change could affect predator-prey
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45 371 dynamics and animal populations.
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51 52 53 373 **REFERENCES**

54
55
56
57 374 Adair SE, Moore JL, Kiel WHJ. 1996. Wintering diving duck use of coastal ponds: An analysis of
58 375 alternative hypotheses. *J. Wildl. Manage.* 60:83–93.
59
60

- 1
2
3 376 Bentley PJ. 2002. Endocrines and osmoregulation: A comparative account in vertebrates. Springer-
4 377 Verlag Berlin Heidelberg.
- 5
6 378 Berger VJ, Kharazova AD. 1997. Mechanisms of salinity adaptations in marine molluscs. *Hydrobiologia*
7 379 355:115–126.
- 8
9 380 Bijleveld AI, van Gils JA, Jouta J, Piersma T. 2015. Benefits of foraging in small groups: An
10 381 experimental study on public information use in red knots *Calidris canutus*. *Behav. Processes* 117:74–
11 382 81.
- 12
13
14 383 Brischoux F, Rolland V, Bonnet X, Caillaud M, Shine R. 2012. Effects of oceanic salinity on body
15 384 condition in sea snakes. *Integr. Comp. Biol.* 52:235–244.
- 16
17 385 Brischoux F, Tingley R, Shine R, Lillywhite HB. 2013. Behavioral and physiological correlates of the
18 386 geographic distributions of amphibious sea kraits (*Laticauda* spp.). *J. Sea Res.* 76:1–4.
- 19
20 387 Britton RH. 1985. Life cycle and production of *Hydrobia acuta* Drap. (Gastropoda: Prosobranchia) in a
21 388 hypersaline coastal lagoon. *Hydrobiologia* 122:219–230.
- 22
23
24 389 Chatelain M, Halpin CG, Rowe C. 2013. Ambient temperature influences birds' decisions to eat toxic
25 390 prey. *Anim. Behav.* 86:733–740.
- 26
27 391 Cramp RL, Meyer EA, Sparks N, Franklin CE. 2008. Functional and morphological plasticity of crocodile
28 392 (*Crocodylus porosus*) salt glands. *J. Exp. Biol.* 211:1482–1489.
- 29
30 393 Duncan CJ. 1962. Salt preferences of birds and mammals. *Physiol. Zool.* 35:120–132.
- 31
32 394 Evans PR, Herdson DM, Knights PJ, Pienkowski MW. 1979. Short-term effects of reclamation of part
33 395 of Seal Sands, Teesmouth, on wintering waders and Shelduck - I. Shorebird diets, invertebrate
34 396 densities, and the impact of predation on the invertebrates. *Oecologia* 41:183–206.
- 35
36
37 397 Fair J, Whitaker S, Pearson B. 2007. Sources of variation in haematocrit in birds. *Ibis* 149:535–552.
- 38
39 398 Gerritsen AFC, Van Heezik YM, Swennen C. 1982. Chemoreception in two further *Calidris* species (*C.*
40 399 *maritima* and *C. canutus*) with a comparison of the relative importance of chemoreception during
41 400 foraging in *Calidris* species. *Netherlands J. Zool.* 33:485–496.
- 42
43 401 Goldstein D, Skadhauge E. 2000. Renal and extrarenal regulation of body fluid composition. In:
44 402 Whittow GC, editor. *Sturkie's avian physiology*. Elsevier. p. 265–297.
- 45
46
47 403 Gutiérrez JS. 2014. Living in environments with contrasting salinities: A review of physiological and
48 404 behavioural responses in waterbirds. *Ardeola* 61:233–256.
- 49
50 405 Gutiérrez JS, Abad-Gómez JM, Villegas A, Sánchez-Guzmán JM, Masero JA. 2013. Effects of salinity on
51 406 the immune response of an 'osmotic generalist' bird. *Oecologia* 171:61–69.
- 52
53 407 Gutiérrez JS, Dietz MW, Masero JA, Gill RE, Dekinga A, Battley PF, Sánchez-Guzmán JM, Piersma T.
54 408 2012. Functional ecology of saltglands in shorebirds: Flexible responses to variable environmental
55 409 conditions. *Funct. Ecol.* 26:236–244.
- 56
57
58
59
60

- 1
2
3 410 Gutiérrez JS, Masero JA, Abad-Gómez JM, Villegas A, Sánchez-Guzmán JM. 2011. Understanding the
4 411 energetic costs of living in saline environments: Effects of salinity on basal metabolic rate, body mass
5 412 and daily energy consumption of a long-distance migratory shorebird. *J. Exp. Biol.* 214:829–835.
6
7 413 Gutiérrez JS, Soriano-Redondo A, Dekinga A, Villegas A, Masero JA, Piersma T. 2015. How salinity and
8 414 temperature combine to affect physiological state and performance in red knots with contrasting
9 415 non-breeding environments. *Oecologia* 178:1077–1091.
10
11 416 Halpin CG, Skelhorn J, Rowe C. 2014. Increased predation of nutrient-enriched aposematic prey.
12 417 *Proc. R. Soc. B Biol. Sci.* 281:20133255.
13
14 418 Hannam KM, Oring LW, Herzog MP. 2003. Impacts of salinity on growth and behavior of American
15 419 avocet chicks. *Waterbirds* 26:119–125.
16
17
18 420 Hänninen L, Pastell M. 2009. CowLog: open-source software for coding behaviors from digital video.
19 421 *Behav. Res. Methods* 41:472–476.
20
21 422 Harriman AE. 1967. Laughing gulls offered saline in preference and survival tests. *Physiol. Zool.*
22 423 40:273–279.
23
24 424 Harriman AE, Kare MR. 1966. Tolerance for hypertonic saline solutions in herring gulls, starlings, and
25 425 purple grackles. *Physiol. Zool.* 39:117–122.
26
27
28 426 Herbst DB. 2001. Gradients of salinity stress, environmental stability and water chemistry as a
29 427 templet for defining habitat types and physiological strategies in inland salt waters. *Hydrobiologia*
30 428 466:209–219.
31
32 429 Hildebrandt J-P. 2001. Coping with excess salt: Adaptive functions of extrarenal osmoregulatory
33 430 organs in vertebrates. *Zool.* 104:209–220.
34
35
36 431 Houston A, McNamara JM. 1999. Models of adaptive behaviour: An approach based on state.
37 432 Cambridge: Cambridge University Press.
38
39 433 Hurly T, Oseen M. 1999. Context-dependent, risk-sensitive foraging preferences in wild rufous
40 434 hummingbirds. *Anim. Behav.* 58:59–66.
41
42 435 Kitchell RL, Ström L, Zotterman Y. 1959. Electrophysiological studies of thermal and taste reception in
43 436 chickens and pigeons. *Acta Physiol. Scand.* 56:133–151.
44
45
46 437 Komendantov AY, Smurov AO. 2009. Salinity tolerance polygon of *Hydrobia ulvae* (Pennant, 1777)
47 438 (Mollusca: Hydrobiidae). *Russ. J. Ecol.* 40:543–546.
48
49 439 Long SE, Vetter TW. 2002. Determination of sodium in blood serum by inductively coupled plasma
50 440 mass spectrometry. *J. Anal. At. Spectrom.* 17:1589–1594.
51
52 441 Mahoney SA, Jehl JR. 1985. Adaptations of migratory shorebirds to highly saline and alkaline lakes:
53 442 Wilson's phalarope and American avocet. *Condor.*
54
55
56 443 Martin P, Bateson P. 1983. Measuring behaviour: An introductory guide. Second edition, Cambridge
57 444 University Press: Cambridge.
58
59
60

- 1
2
3 445 Masero JA. 2002. Why don't Knots *Calidris canutus* feed extensively on the crustacean *Artemia*? Bird
4 446 Study 49:304–306.
5
6 447 Matson KD, Millam JR, Klasing KC. 2000. Taste threshold determination and side-preference in
7 448 captive cockatiels (*Nymphicus hollandicus*). Appl. Anim. Behav. Sci. 69:313–326.
8
9 449 Mazzotti FJ, Dunson WA. 1989. Osmoregulation in crocodilians. Am. Zool. 29:903–920.
10
11 450 McNab BK. 2002. The physiological ecology of vertebrates: A view from energetics. Ithaca: Cornell
12 451 University Press.
13
14 452 Milot E, Cohen AA, Vézina F, Buehler DM, Matson KD, Piersma T. 2014. A novel integrative method
15 453 for measuring body condition in ecological studies based on physiological dysregulation. Methods
16 454 Ecol. Evol. 5:146–155.
17
18
19 455 Moreira F. 1994. Diet and feeding rates of Knots *Calidris canutus* in the Tagus estuary (Portugal).
20 456 Ardea 82:133–133.
21
22 457 Nebel S, Jackson DL, Elnor RW. 2005. Functional association of bill morphology and foraging
23 458 behaviour in calidrid sandpipers. Anim. Biol. 55:235–243.
24
25
26 459 Nebel S, Piersma T, Gils J Van, Dekinga A, Spaans B. 2000. Length of stopover, fuel storage and a sex-
27 460 bias in the occurrence of Red Knots *Calidris c. canutus* and *C. c. islandica* in the Wadden Sea during
28 461 southward migration. Ardea 96:286–292.
29
30 462 Nyström K, Pehrsson O. 1988. Salinity as a constraint affecting food and habitat choice of
31 463 mussel-feeding diving ducks. Ibis 130:94–110.
32
33 464 Ortiz RM. 2001. Osmoregulation in marine mammals. J. Exp. Biol. 184:1831–1844.
34
35
36 465 Peaker M, Linzell JL. 1975. Salt glands in birds and reptiles. Cambridge University Press, Cambridge.
37
38 466 Piersma T, Aelst R van, Kurk K, Berkhoudt H, Maas LRM. 1998. A new pressure sensory mechanism
39 467 for prey detection in birds: the use of principles of seabed dynamics? Proc. R. Soc. B Biol. Sci.
40 468 265:1377–1383.
41
42 469 Piersma T, Koolhaas A, Dekinga A. 1993. Interactions between stomach structure and diet choice in
43 470 shorebirds. Auk 110:552–564.
44
45 471 Piersma T, Koolhaas A, Dekinga A, Gwinner E. 2000. Red blood cell and white blood cell counts in
46 472 sandpipers (*Philomachus pugnax*, *Calidris canutus*): effects of captivity, season, nutritional status,
47 473 and frequent bleedings. Can. J. Zool. 78:1349–1355.
48
49
50 474 R Development Core Team (2013) R Foundation for Statistical Computing. R Development Core
51 475 Team, Vienna, Austria.
52
53 476 Ravenscroft NOM, Beardall CH. 2003. The importance of freshwater flows over estuarine mudflats
54 477 for wintering waders and wildfowl. 113:89–97.
55
56 478 Schmidt-Nielsen K. 1960. The salt-secreting gland of marine birds. Circulation 21:955–967.
57
58
59
60

- 1
2
3 479 Schmidt-Nielsen K. 1997. Animal physiology: Adaptation and environment. Fifth edition, Cambridge.
4
5 480 Schmidt-Nielsen K, Fange R. 1958. Salt glands in marine reptiles. Science 182:783–785.
6
7 481 Shuttleworth TJ, Hildebrandt J. 1999. Vertebrate salt glands: Short- and long-term regulation of
8 482 function. J. Exp. Zool. 701:689–701.
9
10 483 Skadhauge E. 1981. Osmoregulation in birds. Hoar WS, Hoelldobler B, Johansen K, Langer H, Somero
11 484 G, editors. Springer-Verlag Berlin Heidelberg.
12
13 485 Staaland H. 1967. Anatomical and physiological adaptations of the nasal glands in Charadriiformes
14 486 birds. Comp. Biochem. Physiol. 23:933–944.
15
16 487 Taplin LE, Grigg GC. 1981. Salt glands in the tongue of the estuarine crocodile *Crocodylus porosus*.
17 488 Science 212:1045–1047.
18
19 489 Todd ME. 1964. Osmotic balance in *Hydrobia ulvae* and *Potamopyrgus jenkinsi* (Gastropoda -
20 490 Hydrobidae). J. Exp. Biol. 41:665–677.
21
22 491 Troup G, Dutka TL. 2014. Osmotic concentration of prey affects food discrimination behaviour in the
23 492 Australian pelican. J. Zool. 294:170–179.
24
25 493 van de Kam J, Ens B, Piersma T, Zwarts L. 2004. Shorebirds. An illustrated behavioural ecology.
26 494 Utrecht: KNNV Publishers.
27
28 495 van den Hout P. 2010. Struggle for safety: adaptive responses of wintering waders to their avian
29 496 predators. PhD Thesis, University of Groningen, Groningen, The Netherlands.
30
31 497 van Gils JA, Dekinga A, Spaans B, Valhl W, Piersma T. 2005. Digestive bottleneck affects foraging
32 498 decisions in red knots *Calidris canutus*. II. Patch choice and length of working day. J. Anim. Ecol.
33 499 74:120–130.
34
35 500 van Gils JA, Piersma T, Dekinga A, Dietz MW. 2003. Cost-benefit analysis of mollusc-eating in a
36 501 shorebird II. Optimizing gizzard size in the face of seasonal demands. J. Exp. Biol. 206:3369–3380.
37
38 502 van Gils JA, Spaans B, Dekinga A, Piersma T. 2006. Foraging in a tidally structured environment by red
39 503 knots (*Calidris canutus*): ideal, but not free. Ecology 87:1189–1202.
40
41 504 van Heezik YM, Gerritsen AFC, Swennen C. 1983. The influence of chemoreception on the foraging
42 505 behaviour of two species of sandpiper, *Calidris alba* and *Calidris alpina*. Netherlands J. Sea Res.
43 506 17:47–56.
44
45 507 Vézina F, Jalvingh KM, Dekinga A, Piersma T. 2006. Acclimation to different thermal conditions in a
46 508 northerly wintering shorebird is driven by body mass-related changes in organ size. J. Exp. Biol.
47 509 209:3141–54.
48
49 510 Visser GH, Dekinga A, Achterkamp B, Piersma T. 2000. Ingested water equilibrates isotopically with
50 511 the body water pool of a shorebird with unrivaled water fluxes. Am. J. Physiol. Regul. Integr. Comp.
51 512 Physiol. 279:R1795–R1804.
52
53
54
55
56
57
58
59
60

- 1
2
3 513 Warton DI, Hui FK. 2010. The arcsine is asinine: the analysis of proportions in ecology. *Ecology* 92:3–
4 514 10.
5
6 515 Wolcott GT, Wolcott DL. 2001. Role of behavior in meeting osmotic challenges. *Am. Zool.* 41:795–
7 516 805.
8
9 517 Wolff WJ, Smit CJ. 1990. The Banc d’Arguin, Mauritania, as an environment for coastal birds. *Ardea*
10 518 78:17–38.
11
12 519 Woodin MC. 1994. Use of saltwater and freshwater habitats by wintering redheads in southern
14 520 Texas. *Hydrobiologia* 279-280:279–287.
15
16 521 Ysebaert T, Herman PMJ, Meire P, Craeymeersch J. 2003. Large-scale spatial patterns in estuaries :
17 522 estuarine macrobenthic communities in the Schelde estuary , NW Europe. *Estuar. Coast. Shelf Sci.*
18 523 57:335–355.
19
20 524 Ysebaert T, Meininger PL, Meire P, Devos K, Berrevoets CM, Strucker RC, Kuijken E. 2000. Waterbird
21 525 communities along the estuarine salinity gradient of the Schelde estuary , NW-Europe. *Biodivers.*
22 526 *Conserv.* 9:1275–1296.
23
24 527 Zhao H, Li J, Zhang J. 2015. Molecular evidence for the loss of three basic tastes in penguins. *Curr.*
25 528 *Biol.* 25:R141–R142.
26
27
28 529 Zwarts LEO, Blomert A, Hupkes R. 1990. Increase of feeding time in waders preparing for spring
29 530 migration from the Banc d’Arguin, Mauritania. *Ardea* 78:237–256.
30
31 531
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532 Table 1. Statistics and coefficients of models for the response ratios of (a) food intake and (b)
 533 foraging time. Predictors included in the final model are in bold; values for excluded predictors refer
 534 to the step before their exclusion.

Response variable	Predictors	Coefficients	s.e.m.	<i>d.f.</i>	<i>t</i> -value	<i>P</i> -value
(a) Food intake	intercept	-4.222	1.067	47	-3.958	0.0003
	Freshwater access ^a	0.833	0.397	7	2.099	0.074
	Salt gland scores	0.809	0.330	47	2.454	0.018
	FW access x SGS	0.986	0.706	46	1.396	0.168
(b) Foraging time	intercept	-4.590	1.329	47	-3.453	0.001
	Freshwater access ^a	0.563	0.495	7	1.136	0.293
	Salt gland scores	0.921	0.411	47	2.242	0.029
	FW access x SGS	0.362	0.894	46	0.404	0.688

535 ^aReference category is 'no access'

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3 537 **Legends to figures**

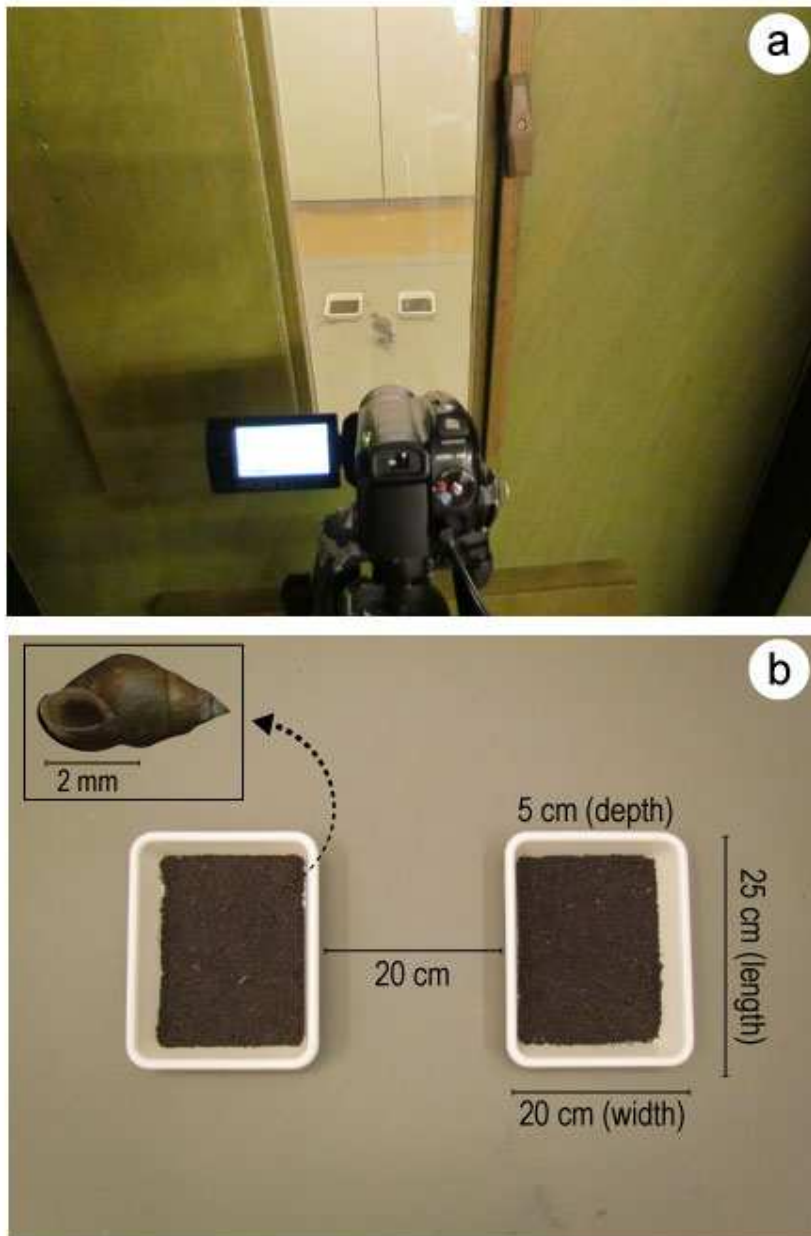
4 538 **Figure 1.** The experimental arena. (a) Frontal view showing aviary, video camera and tray locations,
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7 539 and a focal bird; (b) plan view showing tray dimensions and offered food (inset box in top left corner
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9 540 shows a prey item under a zoom binocular microscope).

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11 541 **Figure 2.** The relationships between (a) overall food intake and foraging time; and (b) between diet-
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14 542 specific food intake and foraging time. Note that individual data points refer to individual sessions
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16 543 and birds (all pooled) and thus show the between individual and experimental variation (see text
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18 544 for further details).

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21 545 **Figure 3.** (a) The mean \pm SE amount (in grams) of high- and low-salinity mud snails eaten by red knots
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23 546 during the nine 3-h experimental sessions; asterisks indicate significant differences between diets at
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25 547 each session (paired *t*-tests; **P*<0.05; ***P*<0.01). (b) The response ratio for the same sessions; the
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27 548 horizontal dashed line depicts the chance level of response (0.5), so that values < 0.5 indicates
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29 549 preference for low-salinity diet and values > 0.5 indicate preference for high-salinity diet. (c) The salt
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31 550 gland scores during the experiment. The shaded areas depict access to freshwater prior to the
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33 551 experimental session. Note that each individual was given one session every other day (see text for
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35 552 further details).

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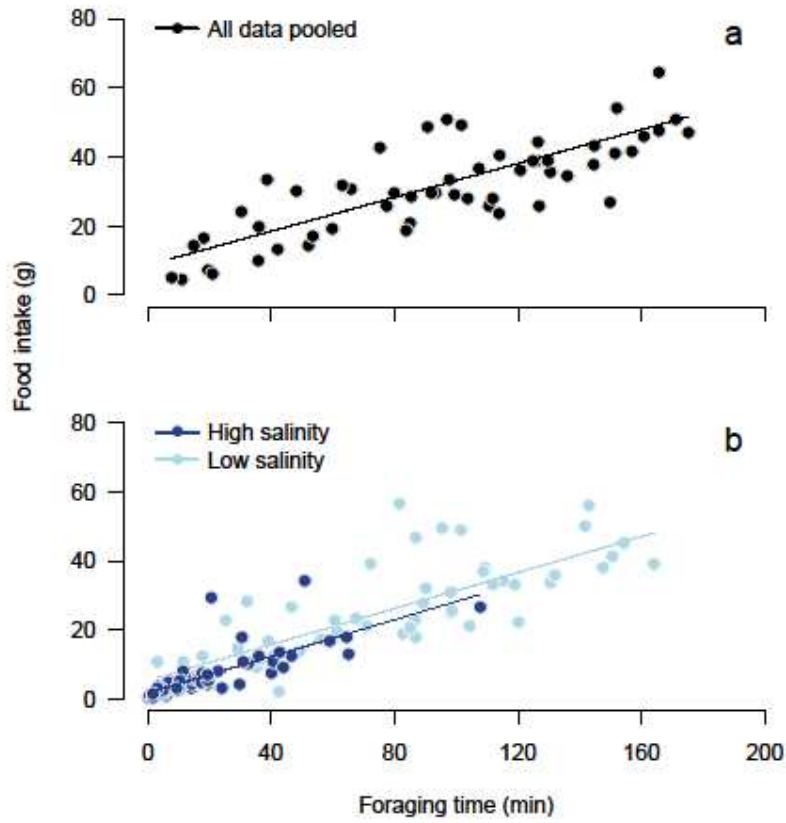
554 **Figure 1**



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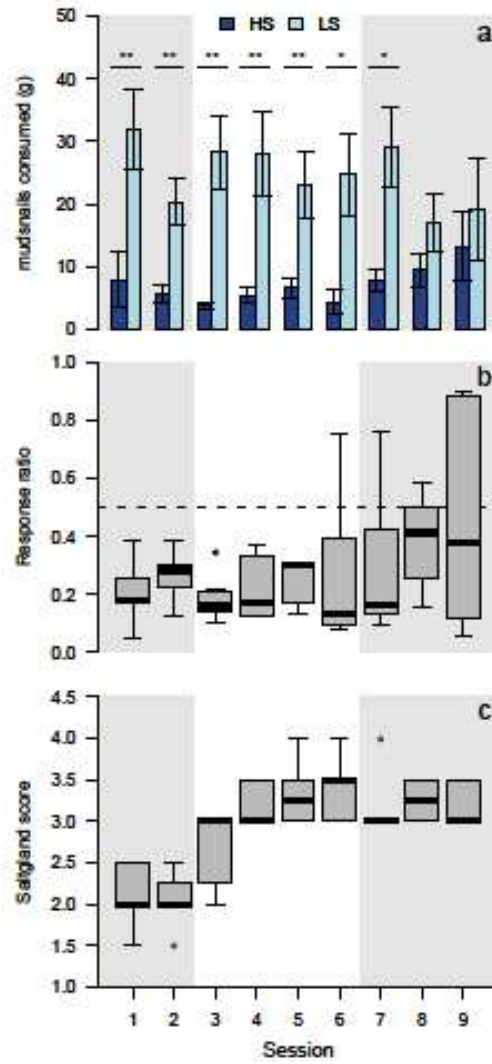
557 Figure 2.



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Only

559 **Figure 3.**



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Lay summary

Seawater is too salty for most land animals, but many marine birds and reptiles can cope with it owing to flexible cephalic “salt” glands that excrete excess salt from the bloodstream. We show that red knots without access to freshwater prefer prey with relatively low salt content when their salt glands are small, but this preference is lost after they enlarge their salt glands and regain access to freshwater.

For Review Only

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