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2	Individual variation in corticosterone and personality
3	traits in the blue tit Cyanistes caeruleus
4	
5	Kathryn E. Arnold ^a , Katherine A. Herborn ^{b,} , Lindsay J. Henderson ^b , Aileen Adam ^c , Lucille Alexander ^{d, e}
6	and Neil Evans ^c
7	
8	^a Environment Department, University of York, YO10 5DD, UK
9	^b Centre for Behaviour and Evolution, IoN, Newcastle University, Henry Wellcome Building,
10	Framlington Place, Newcastle NE2 4HH, UK
11	^c Institute of Biodiversity, Animal Health and Comparative Medicine, School of Life Sciences, College
12	of Medical, Veterinary and Life Sciences, University of Glasgow Glasgow G12 8QQ
13	^d WALTHAM [®] Centre for Pet Nutrition, Waltham-on-the-Wolds, UK
14	^e Current Address: Enterprise and Business Development Office, University of Leicester,
15	University Road, Leicester, LE1 7RH, UK
16	
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18	
19	Tel. +44 1904 322997
20	Email: Kathryn.Arnold@york.ac.uk
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1 Summary

2

3 Variation in personality traits is predicted to reflect physiology, but the extent to which variations in 4 stress hormones derive from differences in personality and/or state-dependent factors remains 5 unclear. To investigate this, wild blue tits Cyanistes caeruleus were briefly brought into captivity and 6 scored for personality and corticosterone (Cort) concentrations. More active females had lower 7 baseline Cort than less active individuals. Exploratory tendency and neophobia did not co-vary with 8 baseline Cort. Stress-induced Cort concentrations were correlated negatively with exploratory 9 tendency and haematocrit, but positively with mass gain in captivity. Baseline and stress-induced 10 Cort concentrations in wintering blue tits were associated with state-dependent variables sex, age 11 and personality traits. Key to interpreting the physiology of personality traits seems to be their 12 interactions with other traits that mediate ability to utilise resources, and thus influence an 13 individual's perception of its current and future energy balance.

1 Introduction

2

3 When exposed to the same environmental or social stimuli, under standardised conditions, the 4 behavioural (Verbeek et al., 1996; Gosling, 2001) and physiological responses (Cockrem, 2007) of 5 individuals of the same species often differ. Behavioural differences that are stable within individuals 6 measured repeatedly across a range of situations or contexts are often referred to as 'personality 7 traits' (Gosling, 2001). While the heritability of personality traits (Drent et al., 2003; Oers et al., 2004; 8 Oers et al., 2005; Martins et al., 2007) suggests that they have a genetic basis, they can also be 9 affected by early environmental conditions (e.g. (Arnold et al., 2007; Stamps & Groothuis, 2010). 10 Similarly, both genes and early environmental conditions have been shown to affect physiological 11 systems such as the stress axis of adults (Monaghan, 2008; Koolhaas et al., 2010). The consistent 12 nature of these differences between individuals suggests they may contribute towards variability in 13 fitness-related traits among individuals (Buchanan, 2000; Both et al., 2005; Blas et al., 2007; Bonier 14 et al., 2009).

15 In birds, the physiological stress response is modulated by both basal levels and the acute 16 release of the adrenal steroid corticosterone (Cort), amongst other hormones. Basal concentrations 17 of Cort have been reported to vary in response to environmental variables such as weather 18 (Astheimer et al., 1995; Rubenstein, 2007) food availability (Jenni-Eiermann et al., 2008), as well as 19 state-dependent variables such as an individual's body condition (Raja-aho et al., 2010) or health 20 status (Martin, 2009). These changes in basal Cort can trigger a series of physiological and 21 behavioural changes that should restore homeostasis. For example the increase in Cort seen 22 following food restriction simulates increased motivation to feed (Lohmus et al., 2003; Long & 23 Holberton, 2004; but see Piersma et al., 2000). Chronic elevations of basal Cort, due to long term or 24 repeated exposure to a stressor, have been reported to have negative effects, suppressing insulin 25 production hence rendering the animal prone to fatigue and disease (reviewed in Buchanan, 2000, 26 see also (Goutte et al., 2010Hau & Goymann, 2015). However, high basal Cort might not necessarily

signal challenging conditions: basal Cort also varies consistently across individuals under the same
conditions, with some individuals showing relatively high and others showing low levels of Cort. The
extent to which such differences in basal Cort reflect individual differences in personality and fitness
are still the focus of discussion (Bonier et al., 2009; Angelier et al., 2010; Dingemanse et al., 2010;
Koolhaas et al., 2010; Hau & Goymann, 2015).

6 Rapid on-set, acute stress responses evoked in response to an 'emergency' often involve an 7 immediate increase in blood glucose, to facilitate rapid behavioural reactions such as fight or flight 8 (Cockrem & Silverin, 2002a) and are generally considered to be adaptive (Buchanan, 2000). In Parids, 9 like many small passerines (Wingfield & Kitayskya, 2002), the acute stress response is generally 10 characterised by a rise in Cort that usually peaks between ten and approximately 40 minutes after 11 the stressor before declining (Cockrem & Silverin, 2002b; Henderson et al., 2014)). The level and 12 timing of the peak in Cort varies both within and between species (Bokony et al., 2009; Cockrem et 13 al., 2009; Henderson et al., 2014). Extensive blood sampling is not possible in small birds, so 14 researchers generally present data on 'stress-induced' (rather than strictly peak) Cort 15 concentrations, measured after an animal has been held for a set time under standardised restraint 16 conditions (Wingfield, 1994). As physiological responses to stress are often accompanied by changes 17 in behaviour, there is a prediction, with some supporting data, that personality types might show 18 associated consistent differences in both basal and acute Cort levels (Cockrem, 2007; Hau & 19 Goymann, 2015). These effects might be directly related, for example more neophobic juvenile rats 20 have a higher glucocorticoid response as adults (Cavigelli & McClintock, 2003), or indirect, for 21 example where personality affects the ability to acquire or control food, which might have an impact 22 on both stress-induced and baseline Cort (Koolhaas et al., 2010). Similarly, age, sex and dominance 23 status can affect resource holding potential and thus energetic balance (Gosler, 1996); Fat reserves 24 reflect a trade-off between starvation (high fat reserves are beneficial) and predation (high fat 25 reserves slow down escape flight, so are costly) resulting in great tits Parus major living under high 26 perceived predation risk carrying lower fat reserves than under low perceived predation risk.

Moreover, dominant birds (adult males) carried the least fat, presumably because they were able to
displace subordinate birds (juvenile females) at feeders, so had a low risk of starvation despite their
low fat reserves (Gentle & Gosler, 2000). Thus, age, sex and dominance status might also influence
both baseline and stress-induced Cort (Rubenstein, 2007). So although there is evidence of
correlations between an individual's personality and its glucocorticoid profile (Carere et al., 2003;
Cavigelli & McClintock, 2003; Cockrem, 2007; Martins et al., 2007; Cavigelli et al., 2009; Koolhaas et
al., 2010), causality is difficult to assign.

8 Assessments of 'personality traits' in animals result in individuals being traditionally defined 9 as falling somewhere on a linear continuum: neophobic-neophilic or bold-shy (Wilson et al., 1994); 10 aggressive-passive; (Huntingford, 1976); active-inactive (Sih et al., 1992) and fast-slow explorers 11 (Verbeek et al., 1994). Such personality traits are often highly correlated within individuals, for 12 example activity and exploration (Dingemanse et al., 2007; Martin & Reale, 2008), or boldness and 13 aggression (Bell, 2005). A commonly described trait correlation is the "proactive-reactive" syndrome 14 (Carere, 2003; Koolhaas et al., 2007). Such correlations imply proximate links between traits, via 15 genetic linkage or shared physiology (Verbeek et al., 1994). In a review, Cockrem (2007) concluded 16 that across avian species, birds with 'proactive' (bold, fast exploring, aggressive etc.) personalities 17 have relatively active behavioural responses and low stress-induced Cort concentrations, whilst birds 18 with reactive personalities have relatively passive behaviour and large stress-induced Cort 19 concentrations. However, correlations can also occur between traits that are not mechanistically 20 connected but rather subject to the same selection pressures or jointly determined by other factors 21 (Bell & Sih, 2007; Dingemanse et al., 2007; Ruiz-Gomez et al., 2008; Carere et al., 2010). 22 Furthermore, not all species show correlations between such suites of behaviours (e.g. (Bell & Sih, 23 2007; Herborn et al., 2010; Herborn et al., 2011) which would suggest that the regulation of these 24 traits may differ between species. Thus, while neophobia and exploratory tendency are correlated in 25 many but not all species (Herborn et al., 2010) they could differ in their associations with baseline 26 and stress-induced Cort concentrations within individuals and across populations. This is in line with

Cockrem et al.'s (2009) assertion that there is no optimum Cort response for all conditions that
 maximises fitness.

3 Whether or not an individual's hormonal stress response is a function of environmental and 4 state-dependent variables or associated with personality is important for understanding the 5 mechanisms underlying fitness variation. In this study, we investigated the environmental, state-6 dependent and behavioural factors that might be associated with inter-individual differences in 7 baseline and stress-induced Cort in wild blue tits Cyanistes caeruleus. In contrast to studies of the 8 closely related great tit (Verbeek et al., 1996), we have previously shown that neophobia and 9 exploratory tendency are uncorrelated in blue tits (Herborn et al., 2010). This would suggest that in 10 the blue tit these traits can potentially differ in their relationships with basal and/or stress-induced 11 Cort. Moreover, we have found that personality types differed in antioxidant defences, and it was 12 the combination of an individual's personality traits that proved important (Arnold et al., 2015). This 13 suggests that the physiology underlying personality traits is complex. Here, we specifically addressed 14 whether an individual's baseline or stress-induced plasma Cort concentrations were correlated with: 15 1) weather variables on the day of capture, as these would affect opportunities to forage and thus 16 metabolic state; 2) personality traits (neophobia and two measures of behaviour in a novel 17 environment: exploratory tendency and activity); 3) state-dependent variables (condition, 18 haematocrit, change in mass in captivity and motivation to feed) and permanent inter-individual 19 differences (or variables that would not change within a season i.e sex, age and size), as these may 20 also affect foraging opportunities in the wild and therefore hormonal status.

21

22 Material and methods

23

The study was conducted over two winters between 2007 and 2009 on a population of blue tits
living in oak dominated woodland on the east bank of Loch Lomond, UK (56°08'N 4°37'W) (Arnold et

26 al., 2007). In October 2007, eight feeding stations were established at approximately 500m intervals

1 (Herborn et al., 2010). These feeding stations were removed at the end of February 2008 and 2 reinstalled in the same positions between October 2008 and February 2009. Between November and 3 February, birds were captured as they approached the feeding stations, using mist-nets. Mist-netting 4 was conducted between dawn and noon, three times at each feeding station in the winter of 2007-8 5 and twice in 2008-9. Birds arrived in captivity generally between 10:00 and 12:00, within 15 minutes 6 journey time from their capture site. A total of 125 blue tits were caught, with 69 birds (54 male: 15 7 female) for which we had a Cort sample and a full set of behavioural data being included in this 8 study. For the other 56 individuals, in most cases we did not have a large enough blood sample to 9 run the Cort analyses as well as other planned blood assays, in a few cases the bird did not perform 10 properly in one of the behavioural trials so had to be excluded and/or a morphometric measurement 11 was missing. This sex ratio bias reflected that of the population of birds caught at the feeders 12 (Herborn et al., 2010). Birds were then taken into captivity for 2 days in 2007/8 and 3 days in 2008/9. 13 In 2007/8 roughly two hours after blood sampling, birds were then released at the site of capture. In 14 2008/9, birds were released on day 3 following the second exploration trial. 15

16

Environmental variables on the day of capture

Variables that may have affected foraging immediately prior to capture and hence metabolic state
were recorded. These included day length, rainfall (mm) and minimum and maximum temperature
for the day of capture. Weather data were collated from records for Glasgow Bishopton sent to us
by the Met office.

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Personality traits

After removal from the mist-net, birds were held in cloth bags and transported, up to 3km, to the Scottish Centre for Ecology and the Natural Environment. After arrival birds were scored for their morphometrics and housed indoors, at a temperature of 17°C±1°C on a 12:12 hour light:dark cycle (to enable us to conduct all of the tests and minimise the time the birds would be held in captivity

1 whilst standardising conditions across birds). Trials ran from mid-November until the end of 2 January, during which time the natural daylength was at maximum 8hr30 and minimum 7hr. Each 3 bird was housed individually in a 150cm x 50cm x 50cm high cage, with six cages per room arranged 4 such that birds were visually but not acoustically occluded from one another. Birds were fed peanut 5 granules, Haiths' Prosecto™ insectivorous mix and water ad libitum, supplemented with around ten 6 Tenebrio molitor and two Galleria mellonella larvae per day. All birds were observed eating within 10 7 minutes of arrival in captivity. After arrival birds were left undisturbed for a minimum of 2 hours and 8 a maximum of 3 hours before the start of personality trait assessments. As we wanted to return the 9 birds to the wild as soon as possible, we did not allow the birds a long acclimation period in captivity. 10 We know that the personality traits measured in captivity predicted neophobia and exploratory 11 tendency measured for the same individuals in the wild weeks or even months later (Herborn et al. 12 2010). So we have some evidence that the short acclimation period was not having significant 13 negative impacts on our behavioural assays. Three behavioural traits, exploratory tendency, activity 14 and neophobia were assayed as fully described in Herborn et al (2010):

15 Exploratory tendency was assessed within what would become the home cage of the focal 16 bird. Each cage was split into two halves with an opaque cage divider (Figure S1 in Electronic 17 Supplementary material (ESM)). On arrival into captivity, the bird was introduced to one side of the 18 cage, selected at random, and allowed to acclimate for two hours thereby creating a "familiar" and, 19 behind the divider, a "novel" environment. To assay exploratory tendency and not neophobia, the 20 arrangement of plastic plants and perches was the same in each cage half, so that the novel 21 environment was novel only in that it was unexplored. At the start of the trial, the food bowl and any 22 spilt food were removed and 30 mins after that the water bowl was also removed, this was to 23 ensure that the birds did not spent the whole trial eating or drinking, thus making it difficult to 24 determine their motivation for moving. After a further 30 mins, the researcher removed the opaque 25 cage divider and observed the focal bird for 10 mins. Unlike other exploration trials (e.g. (Verbeek et 26 al., 1994), individuals had the option of remaining within the familiar environment. This allowed us

1 to distinguish activity due to exploration which is related to information gathering about a novel 2 environment, from activity due to escape behaviours which is not necessarily linked to information 3 gathering (Mettke-Hofmann et al., 2009). The number of movements (hop or flight) in each side of 4 the cage was recorded, with the endpoint of each movement defining the side of the cage: novel or 5 familiar. Exploratory tendency was calculated as the number of movements in the novel 6 environment minus the number of movements in the familiar environment. In 2008/9, the 7 exploratory tendency trial was repeated between 08:00 and 011:00 on day 3 of captivity before 8 releasing the birds. This was to allow us to calculate the repeatability of this behavioural trait. To 9 create a new novel environment, the paper lining the cage was changed from white to brown or vice 10 versa. Also, the arrangement and size of perches and artificial plant material were similar between 11 these cage types, but different leaf shapes were used in the brown versus white-lined cages.

Activity was derived from the same exploration trial, as the number of movements in both the familiar and novel side of the cage. We described behaviour in the exploration trial in these two ways to determine whether patterns relating to exploration were due to individual variation in true exploratory tendency, as reflected by space use within the trial, or simply activity levels, as commonly reported in other exploration studies (Verbeek et al., 1994; Martins et al., 2007).

17 Neophobia was assessed over two trials conducted between 13:00 and 17:00 on day 1 and 18 08:00 and 11:00 on day 2. Each trial had two phases: a novel object phase and a disturbance control 19 phase. Food and water were removed for 30 mins prior to each phase. Water was removed so that 20 during the experiment only one resource, food, was available. In the novel object phase, the 21 observer then returned the food bowl with one of two novel objects (different novel objects were 22 used in the two trials) placed inside. The objects were a bright pink plastic frog and half of a purple 23 rubber ball, of similar size (approximately 4cm diameter and 2cm height) (Figure S2 in ESM). The 24 latency to land on familiar food bowl was recorded. The object was then removed and the water 25 returned. The disturbance control phase was completed so as to allow interpretation of the results 26 relative to any possible differences in an individual's motivation to feed, or their response to

disturbance by the observer returning the food bowl to the cage (van Oers et al., 2005). Thus in a
random order either one hour before or after the novel object phase, the latency to feed was
measured by food bowl removal and replacement as before but without a novel object. Neophobia
was defined as the latency in the novel object phase minus the latency in the disturbance phase.
Mean latency to approach the food bowl was significantly greater when a novel object was present
(Wilcoxon Signed ranks Z = - 3.34, N = 68, p = 0.001), than when absent. Thus the presence of the
novel object modified behaviour so could be said to induce neophobia.

8 Repeatability of neophobia, exploratory tendency or activity scores between day 1 and day 2 9 was calculated using the mean squares from an analysis of variance, following Lessells and Boag 10 (1987). Neophobia across days 1 and 2 (ANOVA; r = 0.29, F_{63, 68} = 1.811, p = 0.009), as well as 11 exploratory tendency (r = 0.27, $F_{36.68} = 1.71$, p = 0.04) and activity (r = 0.42, $F_{36.68} = 2.56$, p = 0.001) 12 across days 1 and 3 were repeatable. In the following results, only data from the first of each trial 13 per individual were analysed as they were therefore considered representative of the individual's 14 personality traits but ensured no carry-over effects through learning or habituation to the 15 experiments. This also meant that we had a larger sample size for exploratory tendency and activity 16 than if we had taken a mean of both trials.

17

18 State-dependent and permanent inter-individual differences

Next, we determined whether state-dependent (i.e. changeable within a season) variables, and permanent inter-individual differences (fixed at least within one season) predicted an individual's Cort concentration. An individual's state was assessed as its body condition (residual of mass at blood sampling against tarsus length), haematocrit and change in mass in captivity (mass at blood sampling after 24 hours of captivity – mass at capture).

The duration of the first and second feeding bouts in each disturbance trial (see below) were taken as a measure of hunger and motivation to feed. Neither of these measures differed significantly between the two years, juveniles and adults or between the sexes (Mann-Whitney U-

test p > 0.11 in all cases), so the data were pooled. The duration of the two feeding bouts were
correlated, i.e. consistent, within individuals (Kendall's tau b = 0.23, N = 56, p = 0.027), so from the
first disturbance trial we analysed just the duration of the second feeding bout as an index of hunger
and motivation to feed in case there was any disturbance associated with returning the bowl to the
cage in the first feeding bout.

6

Permanent variables (unchanged within a field season) were also assessed and recorded for each bird. Wing length was used as a measure of overall body size (not measured in one bird). Age (juvenile/adult) was determined from plumage traits (Svensson, 1992). Sex was determined from a blood sample using primers P2 and P8 (Arnold et al., 2007).

10

11 Blood sampling

12 Approximately 24 hours after capture, birds were blood sampled. Birds were sampled after 13 undergoing behavioural assays as there is some evidence that blood sampling and/or associated 14 handling can change the physiology and behaviour of birds (e.g. (Le Maho et al., 1992; Cockrem et 15 al., 2009; Dickens et al., 2009). Moreover, in captivity all the birds had received ad libitum food, 16 constant temperature and light/dark cycles and minimal social interactions, thus many of the 17 environmental factors known to affect Cort concentrations had been standardised across individuals 18 (Romero et al., 2000; Pravosudov et al., 2001). During blood sampling, two birds per room were 19 removed from their cages with the order randomly assigned. For baseline Cort, birds were blood 20 sampled within 3 mins of the researcher entering the aviary room, which is generally considered to 21 be representative of baseline Cort concentrations (Romero & Reed, 2005). For the stress-induced 22 Cort, the bird was put into a cloth bag upon capture, removed after 14 mins and blood sampled 15 23 mins after initial disturbance. Caged birds were then left undisturbed for one hour before re-entry to 24 each room to collect samples from different birds. The last birds to be sampled had therefore been 25 disturbed a maximum of twice that day, but disturbance frequency did not significantly influence 26 either baseline (ANOVA F $_{2,35}$ = 0.065, p = 0.94) or stress-induced Cort (ANOVA F $_{2,29}$ = 0.092, p =

1 0.91). To collect the blood, a sterile 25 gauge needle was used to prick the brachial vein. Blood 2 samples were collected in 75 μ l heparinized capillaries and centrifuged for 5 minutes at 14,000 g. A 3 haematocrit score (a measure of anaemia) was taken for each tube before the red cells and plasma 4 were split, red cells mixed with ethanol and samples stored at -70 °C until assayed. Less than 10% of 5 the circulating blood volume was collected, as defined by the UK Home Office. Collecting this volume 6 of is not considered to have negative impacts on the welfare of a bird. As we could only take enough 7 blood from each bird for a single Cort assay, due to ethical considerations and the limits defined by 8 the UK Home Office, we only have either baseline or stress-induced Cort concentration from each 9 individual. 10 Corticosterone assays 11 Corticosterone concentrations were measured after extraction of 5-20µl aliquots of plasma in 12 diethyl ether, using a double antibody radioimmunoassay (Wingfield et al., 1992). Primary antibody 13 Esoterix B183, Secondary antibody Sigma goat anti-rabbit and [3H]-corticosterone label (GE 14 Healthcare, UK). See Henderson et al. (2013) for full methods. Extraction efficiency was 93 ±8% 15 (mean ± SD). Recoveries were measured for samples independently and individual corticosterone 16 concentrations adjusted accordingly. The assay detection limit was 0.03 ng/ml. Samples were run 17 within two assays. Inter- and intra-assay coefficients of variation were 6.68% and 4.35±0.001%

- 18 respectively.
- 19

20 Ethical Note

21 All work was carried out in accordance with ASAB/ABS's guidelines for the treatment of animals in

22 research. Work was under license of the UK Home Office and subject to ethical review by

23 WALTHAM[®] Centre for Pet Nutrition and the University of Glasgow. Captive studies were completed

24 and feeders removed 2 months before the first record of nest building in the area. Birds had on

average gained (2.97% ± 7.3%) body mass whilst in captivity. 108 out of the total 125 birds released

26 were later recorded using the feeders or were re-trapped in the area.

2 Statistical Methods

3 Analyses were carried out using SPSS (Ver 20). Prior to analysis, data were checked for normality, 4 homogeneity of variance and co-linearity. Where these assumptions were violated, data were 5 transformed or non-parametric tests were carried out. The distribution of stress-induced Cort 6 concentrations, for example, showed a slight skew and a log transformation improved the situation. 7 In addition, Cook's distances were calculated for all co-variates and subsequently neophobia data 8 from three individuals and exploration data from one individual were excluded as extreme outliers 9 that would have distorted the outcome of the analyses. Exploratory tendency (Shapiro–Wilks test: 10 W_{120} = 0.94, p < 0.0001), activity during the exploration trial (W_{120} = 0.95, p < 0.0001) and neophobia 11 (Shapiro–Wilks test: W_{78} = 0.89, p = <0.0001) were leptokurtic and it was not possible to normalise 12 their distributions. Exploration scores (Mann-Whitney U-test Age: z = -0.38, N = 64, p = 0.71; sex: z = 13 -0.23, N = 60, p = 0.82), activity levels (Age: z = -0.39, N = 64, p = 0.70; sex: z = -0.94, N = 60, p = 0.35) 14 and neophobia scores (Mann-Whitney U-test Age: z = -1.25, N = 63, p = 0.21; sex z = -0.88, N = 59, p15 = 0.38) did not differ between adults and juveniles or males and females, therefore data were 16 pooled to analyse other sources of between-individual variation. With the Bonferroni correction 17 threshold p-value of 0.004, all other morphometric (mass at capture, mass at blood sampling, wing 18 length) and environmental variables (rainfall, minimum temperature and maximum temperature on 19 day of capture) were non-significant in relation to neophobia, exploratory tendency and activity. 20 Therefore analyses of these traits were calculated using actual scores, rather than, for example, 21 residuals or transformed data. 22 General linear models (GLMs) were run with Cort plasma concentration (basal or peak) as 23 the dependent variable, sex and age (adult or juvenile) as factors, and change in mass in captivity (mass on day 2 - mass on day1), wing length, haematocrit and personality traits as co-variates. The 24 25 interactions between sex or age and personality traits or change in mass in captivity were also

26 included in the full model. All tests were two-tailed. Means ± S.E. are presented.

2 Results

3	
4	Neither basal nor stress induced Cort concentrations differed significantly between years (Mann-
5	Whitney U-test z = -0.36, N = 69, p = 0.72) so data were pooled. Baseline Cort concentrations varied
6	among individuals, from 0.33 – 13.21 ng/ml. Stress-induced plasma Cort concentrations were
7	significantly higher (mean = 10.55 ng/ml \pm 0.94, range from $1.32 - 23.83$ ng/ml) than baseline
8	concentrations (mean = 4.17 ng/ml \pm 0.55; GLM F _{1, 66} = 30.81, p < 0.0001), although we did not have
9 10	both measures from the same individual.
11	Environmental variables on the day of capture
12	Neither baseline nor stress-induced Cort concentrations were significantly related with daily rainfall
13	(Baseline: Kendall's tau = 0.13, N = 38, p = 0.27; Stress-induced: Kendall's tau = 0.07, N = 28, p =
14	0.63), minimum daily temperature (Baseline: Kendall's tau = 0.02, N = 38, p = 0.87; Stress-induced:
15	Kendall's tau = -0.04, N = 28, p = 0.75) or maximum daily temperature on the day of capture
16	(Baseline: Kendall's tau = 0.00, N = 38, p = 1.0; Stress-induced: Kendall's tau = 0.06, N = 28, p = 0.63).
17	
18	Personality traits
19	As mentioned above, we only used data from the first of each trial per individual. In these first trials,
20	neophobia scores varied from -117s to 597s with a mean of 85.59s \pm 19.18 (six birds scored zero and
21	six birds failed to complete the trial), with positive scores indicating highly neophobic birds and
22	negative scores meaning that the birds were not neophobic. Exploratory tendency ranged from -280

- to 226 with a mean of 9.69s ± 11.79 (three birds scored zero and two birds failed to complete the
- 24 trial) and total activity level from 0 467 with a mean of 233.42s ±15.90 (two birds scored zero, i.e.
- 25 did not move, and two failed to complete the trial). High scores indicated highly exploratory and
- active birds respectively.

Neophobia scores were not correlated with either exploratory tendency (Kendall's tau = 0.04, N = 62, p = 0.69) or activity levels during the exploration trial (Kendall's tau = 0.01, N = 62, p =
0.89). Activity levels and exploratory tendency were uncorrelated (Kendall's tau = 0.051, N = 67, p =
0.54). Thus, exploratory tendency, activity levels during the exploration trial and neophobia are
distinct personality traits (i.e. were repeatable across time), not simply three measures of the same
trait or by-products of differences in activity levels. Also, we did not find evidence of a behavioural
syndrome.

8 Baseline Cort concentrations were significantly related to the interaction between sex and 9 activity ($F_{1, 36}$ = 5.46, p = 0.026; sex $F_{1, 36}$ = 8.82, p = 0.006, activity level $F_{1, 36}$ = 10.09, p = 0.003; Fig. 10 1a). Specifically, more active birds had lower baseline Cort than inactive birds. The slope of this 11 relationship was much steeper for females than males. Using a backwards stepwise process, non-12 significant interactions then age, exploratory tendency (Fig. 1b) and neophobia (Fig. 1c) were 13 removed from the model. The sample size for females in this analysis was only six, so we repeated 14 the analysis just for males. For males there was no significant correlation between activity levels and 15 baseline Cort (Pearson correlation = -0.244, N = 31, p = 0.185).

16 Stress-induced Cort was uncorrelated with activity ($F_{1, 28} = 0.02$, p > 0.8; Fig. 1d). There was a 17 non-significant tendency for more neophobic birds to have lower stress-induced Cort concentrations 18 than less neophobic birds ($F_{1, 24} = 3.40$, p = 0.079; Fig. 1e). Stress-induced Cort was significantly 19 associated with exploratory tendency ($F_{1, 24} = 5.04$, p = 0.035; Fig. 1f), with more exploratory 20 individuals having lower stress-induced Cort concentrations than slow explorers.

21 22

State-dependent and permanent inter-individual differences

Mass at capture did not differ between age classes (adult mean = $11.24 \pm 0.11g$, juvenile mean = 11.15 ± 0.11g; GLM F_{1,64} = 0.38, p = 0.54) neither did condition at capture (GLM F_{1,64} = 0.66, p = 0.43). At capture, males weighed more (male mean = $11.33 \pm 0.09g$, female mean = $10.89 \pm 0.09g$; GLM $F_{1,67} = 6.12$, p = 0.016) and were in better condition (GLM F_{1,66} = 6.46, p = 0.013) than females. Birds that were in poor condition upon capture put on significantly more mass in captivity than those in
better condition (Kendall's tau = -0.32, N = 68, p < 0.0001).

Following GLM modelling to investigate the relationship(s) between baseline Cort concentrations and state-dependent traits, all variables dropped out of the model (p > 0.2) except the interaction between age and change in mass (GLM $F_{1, 35} = 5.92$, p = 0.021; age $F_{1, 35} = 2.76$, p = 0.11; change in mass $F_{1, 35} = 0.36$, p = 0.55; Fig. 2). Baseline Cort was lower in juveniles that increased in mass whilst in captivity than compared with those that lost mass, with the opposite relationship in adults.

9 When the above model was re-run with stress-induced Cort as the dependent variable; 10 change in mass ($F_{1,28}$ = 5.26, p = 0.030; Fig. 3a) and haematocrit level ($F_{1,28}$ = 5.40, p = 0.044; Fig. 3b) 11 remained in the GLM. Across all individuals, birds that put on more mass in captivity, typically those 12 that were in poor condition at capture, had higher stress-induced Cort concentrations than those 13 that lost mass in captivity. Also, birds that had lower haematocrit levels (i.e. were more anaemic) 14 had higher stress-induced Cort concentrations than those with higher haematocrit scores. 15 Change in mass and haematocrit contributed additively to the model and were not 16 correlated with one another (Kendall's tau b = - 0.13, N= 65, p = 0.14). Age, sex, condition at blood sampling and feeding bout duration (an index of hunger and motivation to feed on day 1) dropped 17 18 out of the model.

19

20 Discussion

21

In this study, we investigated extrinsic and intrinsic correlates of baseline and stress-induced plasma Cort concentrations in wild blue tits held under standard conditions. Both baseline and stressinduced Cort concentrations varied widely despite all animals being maintained at a standard temperature with *ad libitum* food availability for the 24 hours prior to blood sampling. Weather on the day of capture, which might indicate the condition of birds on entry to the study, was not related

1 to Cort. Notably, neophobia and exploratory tendency, although significantly repeatable within 2 individuals, at least over a 2-3 day interval, did not predict and were not related to baseline Cort 3 concentrations. There was a hint that birds that were generally more active in the exploration trial 4 had lower baseline Cort than less active individuals, but this was driven by data from a small number 5 of females. Unfortunately, given the small number of females sampled, we cannot interpret this sex-6 difference further. With stress-induced Cort, more exploratory individuals had lower Cort 7 concentrations than less exploratory birds. In contrast, there was a tendency for more neophobic 8 birds to exhibit lower stress-induced Cort than less neophobic birds. There were also relationships 9 between Cort and individual state. Variation in baseline Cort concentrations was determined by a 10 significant interaction between age and change in mass in captivity. Juveniles that had gained mass 11 in captivity had a lower baseline Cort concentration than those that declined in mass. Conversely 12 adults that put on more mass in captivity had higher baseline Cort than those that lost mass. With 13 stress-induced Cort though, all birds that put on mass in captivity had higher stress-induced Cort 14 concentrations than those that lost mass, although condition at blood sampling did not contribute 15 significantly to the model. Birds with a low haematocrit level also had higher stress-induced Cort 16 concentrations than birds with higher haematocrit levels.

17 Our mixed evidence for associations between behavioural responses to novel and 18 potentially stressful situations and physiological responses to stress are intriguing but not without 19 precedent (Carere et al., 2003; Martins et al., 2007; Scheid & Noe, 2009; Schoech et al., 2009; Scheid 20 & Noe, 2010). One interpretation of our results is that our behavioural assays did not generate 21 neophobia in blue tits. This was unlikely because most birds took significantly longer to approach the 22 food bowl in the presence of the novel object than in the disturbance phase, although a few 23 individuals were clearly not neophobic, approaching the food faster in the presence than absence of 24 the novel object. This is consistent with the notion that the object created a motivational conflict 25 between desire to feed and to avoid the object which is perceived as a risk (Mettke-Hofmann et al., 26 2002). However, whether the novel objects induced 'stress' is harder to determine. The classic flight

1 or fight response to acute stress, is typically (Buchanan, 2000) but not always (Muller et al., 2006) 2 associated with an elevation in Cort in wild animals. The few studies on birds comparing Cort before 3 and after presentation of a novel object, or more broadly Cort concentrations in high neophobic 4 versus low neophobic individuals, find different responses depending on the experimental design. 5 Generally, those placing a novel object with food identify a Cort response but those presenting the 6 novel object in a neutral location do not (Daisley et al., 2005; Martins et al., 2007; Tobler & Sandell, 7 2007; but see Apfelbeck & Raess, 2008). This absence of a Cort response (although of course other 8 stress hormones might have been elevated) in neutral studies comes even when accompanied by 9 typical "stress" behaviours – e.g. aversions, tonic responses, distress call (Daisley et al., 2005). 10 Furthermore, within the same species, latency to approach a novel object in a neutral location is not 11 necessarily correlated with latency to approach a novel object with food, suggesting they stimulate 12 different behaviours (Mettke-Hofmann et al., 2002). However, stereotypical stress behaviours are 13 not necessarily evidence of physiological stress (Muller et al., 2006). Neophobia, i.e. the avoidance of 14 novelty, might be a behavioural strategy to avoid a potentially stressful situation. Therefore, we 15 cannot assume that neophobia is an index of responsiveness to stressors in general. Neophobia is 16 simply a behavioural response to something unfamiliar which may or may not be dangerous or 17 stressful. Indeed, contrary to predictions, we found a tendency for more neophobic birds to show a 18 lower hormonal stress response than low neophobia birds, but this needs to be interpreted with 19 caution. Moreover, we also have the issue that we might have captured a biased sample of the least 20 neophobic individuals in the population. Certainly we caught fewer females than males which might 21 suggest that we were lacking data on individuals who were subordinate at feeders, but also 22 potentially those that avoided novel food sources in the wild. The ecological significance of 23 neophobia in the wild is still to be fully assessed.

Exploration has been previously shown to be linked with stress-induced Cort in some but not all species studied (reviewed in Cockrem, 2007), although interpretation of the patterns can be complex. For example, in captive populations of passerines subject to selection on either

1 behavioural or hormonal responses to stressors, Cort and behaviour are sometimes related: In two 2 lines of great tits (Parus major) selected to be either fast or slow explorers, Carere et al. (2003) 3 found that slow exploring birds had higher stress-induced Cort metabolites in their faeces and less 4 impairment in activity than fast birds when stressed (Carere et al., 2003). In zebra finches 5 (Taeniopygia guttata), over five generations researchers selected for high or low stress-induced 6 Cort. In the low Cort line, stress-induced Cort concentration was positively correlated with 7 exploratory behaviour but no relationship was present in the high Cort or control lines (Martins et 8 al., 2007). Thus, the results were ambiguous, but provide some evidence of heritable coping styles 9 suggesting selective maintenance of variation in Cort secretion patterns within populations. In 10 contrast to these studies in which the measurement of exploration and activity were confounded 11 (Carere et al., 2003; Martins et al., 2007) we found no relationship between exploration and baseline 12 Cort. On the contrary, experimentally increasing circulating Cort can increase activity levels in birds 13 (Breuner et al., 1998). Thus we suggest that exploratory tendency per se is not a behavioural 14 indicator of chronic stress, or that individuals that are chronically stressed are less likely to gather 15 information about a novel environment. Comparison of our behavioural results with those of others 16 (Marchetti & Drent, 2000; van Overveld & Matthysen) suggests that interpretation of "exploratory 17 tendency", and particularly its importance in explaining differences in variation in information 18 acquisition and use, is dependent on the method by which exploration is scored. The differing 19 patterns of covariation between circulating Cort concentrations and personality traits suggests that 20 the physiological mechanisms and potentially the costs of different personality traits will vary. Thus, 21 we would emphasize the need to check the assumptions that exploration, activity and neophobia / 22 risk responsiveness are always correlated or that they represent three measures of the same trait 23 (Reale et al., 2007).

One issue to be explored is the impact of bringing birds into captivity on baseline Cort measurements and both behavioural and hormonal stress responsiveness. In order to standardise the food, temperature and day length experienced by the birds and to ensure that time spent in the

1 mist net did not influence the baseline Cort measure (Romero & Reed, 2005), we decided to blood 2 sample birds after 24 hours in captivity. However, it might be considered that this reflects a 3 disturbed state. Recent work by Dickens et al (2009) on chukars (Alectoris chukar) showed that on 4 the first day of captivity (which involved transport of 100km rather < 3km as in our study) birds lost 5 weight, haematocrit decreased and both baseline and stress-induced Cort concentrations were 6 raised. These effects took 3 to 9 days to fully attenuate. In contrast, we found that after 24 and 48 7 hours of captivity, the majority of blue tits had gained rather than lost body mass. Moreover, the 8 plasma concentrations of Cort we measured in captivity were similar to those measured in the field 9 at the capture site (basal Cort in the field in 2008 only: $3.2 \text{ ng/ml} \pm 1.8$, n = 36; in captivity: 4.2 ng/ml10 ±3.1, n = 16, t-test P = 0.22; (Henderson et al., 2013)). Thus, the elevated stress markers seen in 11 chukars (Dickens et al., 2009) might have been associated with the long distance transport, during 12 which animals often do not feed or drink, rather than captivity per se, as has been observed in 13 livestock (Fazio & Ferlazzo, 2003). More research is needed to determine whether individuals with 14 different personality traits vary in how they cope with captivity. It should be noted that we have 15 previously shown that neophobia and exploratory tendency measured in the wild have been shown 16 to be consistent over several weeks to months and was predicted by measurements taken in 17 captivity using analogous methods and the same individuals (Herborn et al., 2010). Thus, whilst only 18 reflecting a snapshot in time, we feel that our results have some ecological relevance at least for our 19 study population.

20 Our data suggest that both baseline and stress-induced Cort concentrations in wintering 21 blue tits are associated with state-dependent and permanent variables, as well as personality traits. 22 This is in line with evidence that Cort is involved in maintaining a homeostatic energy balance, 23 allowing individuals to respond to food shortages and environmental perturbations (Wingfield & 24 Kitayskya, 2002; Bonier et al., 2009; Hau & Goymann, 2015). For adults only, in the case of baseline 25 Cort, and birds (independent of age), in the case of elevated Cort, birds that put on more mass in 26 captivity had higher plasma Cort concentrations. Moreover birds that were light on entry into

1 captivity gained more mass during captivity than relatively heavy individuals. What we do not know, 2 and could only find out with manipulative experiments, is whether the poor quality birds gained 3 mass because of their high Cort concentrations (which can stimulate hyperphagia (Landys et al., 4 2004)) or the rapid increase in mass resulted in raised baseline and stress-induced Cort. There was 5 no sex difference in the relationship between basal Cort and change in mass, although on entry into 6 captivity males were in better condition than females. In contrast to adults, for juveniles the 7 relationship between baseline Cort and mass gain was negative. However, juveniles and adults did 8 not, on average, differ significantly in mass upon capture or their mass change in captivity. Thus, 9 juveniles may have a different stress physiology and/or feedback mechanisms from adults (Bonier et 10 al., 2009). Experience in coping with environmental perturbations and/or relative dominance status 11 have also been shown to influence how birds respond to changes in food supply (Gosler, 1996; 12 Grieco et al., 2002), and thus potentially Cort levels.

13 For stress-induced Cort, high concentrations were associated with low haematocrit 14 (indicating poor condition) and a greater mass increase. These data indicate that high Cort may 15 signal individuals in poor condition – which might be true for both basal and stress-induced Cort. 16 However, this needs further study, not least because one limitation of our study is that we did not 17 have basal and stress-induced Cort from the same individual for welfare reasons. Thus, we do not 18 know whether blue tits with high basal Cort also had high stress-induced Cort (which might actually 19 indicate a relatively small change in Cort in response to stress) or those with low basal Cort could 20 have high stress-induced Cort (i.e. had a relatively great physiological response to restraint). 21 Selection experiments have shown that basal and stress-induced Cort levels are not necessarily 22 linked (Martins et al., 2007), but this requires more investigation in relation to behavioural 23 responses.

To conclude, exploratory tendency, activity and neophobia in blue tits were uncorrelated and showed differing patterns of co-variation with state-dependent, permanent and physiological traits. As individuals vary along these three independent axes of behavioural variation, the

1 underlying physiological mechanisms, and indeed the costs (Herborn et al., 2011), may vary in a 2 complex fashion. More research both under captive and wild conditions is needed to determine whether both baseline and stress-induced Cort, i.e. physiological responses to chronic and acute 3 4 stress, are linked with individual variation in behaviour directly or indirectly via homeostatic control 5 of energetic balance. Of particular interest is how personality traits and other factors that affect 6 ability to control resources interact, and thus influence an individual's current and future energy 7 balance. 8 9 Acknowledgements

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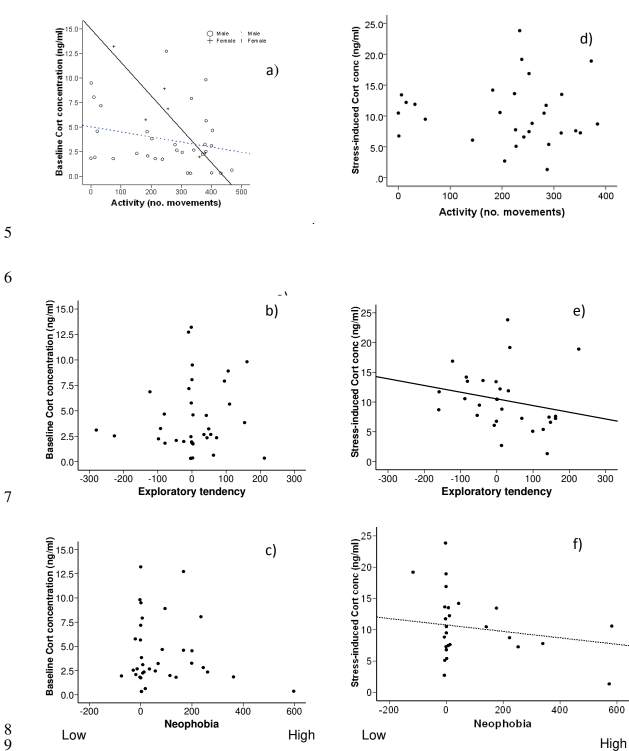
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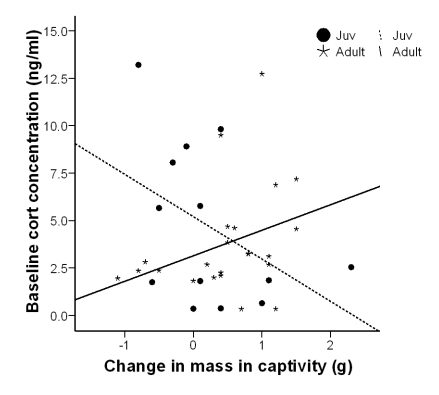
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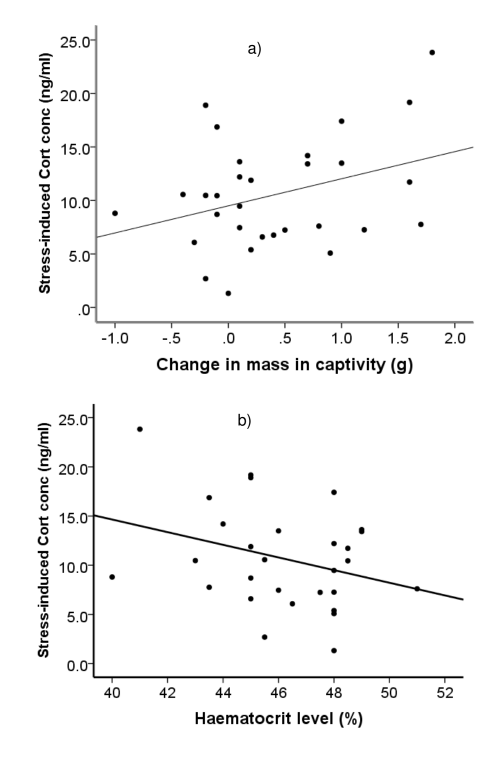
1 Figure legends

3	Fig. 1: Relationships between personality traits and plasma Cort concentrations. Baseline Cort
4	concentrations were related to a) activity level in females (crosses, solid line, N = 6), but not in males
5	(circles, dashed line, N = 31); and unrelated to b) exploratory tendency (N = 37); and c) neophobia (N
6	= 35). Stress-induced Cort concentrations were d) unrelated to activity levels (N = 29); e) marginally
7	related to neophobia (N = 24); and f) significantly related to exploratory tendency (N = 29).
8	
9	Fig. 2: Relationships between change in mass over two days of captivity (mass at blood sampling on
10	day 2 – mass at capture from wild) and baseline plasma Cort concentrations for juveniles (circles,
11	dashed line, $N = 13$) and adults (stars, solid line, $N = 23$).
12	
13	Fig. 3: Relationships between stress-induced plasma Cort concentrations and a) change in mass over
14	two days of captivity (mass at blood sampling on day 2 – mass at capture from wild)(N = 30); b)
15	Mean haematocrit level (N = 30).
16	



2 Figure 2





1 SUPPLEMENTARY MATERIAL

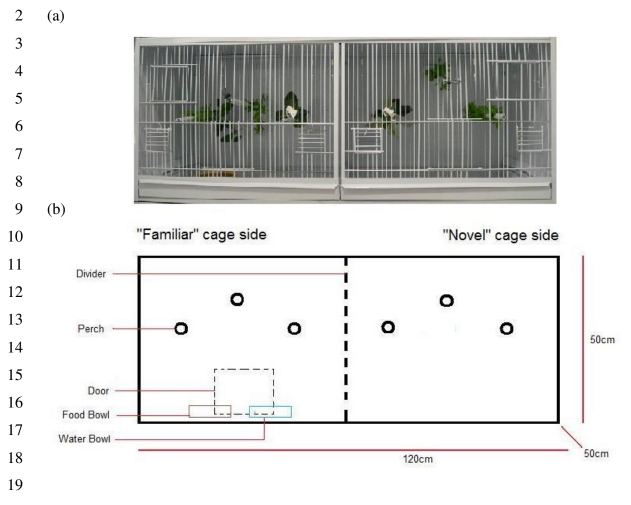
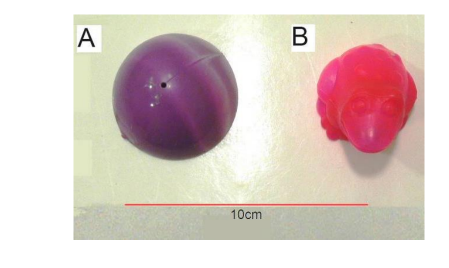


Figure S1: Apparatus used during the captive exploration trials (a) Photo of cage set up. (d) Schematic of cage set up; the food was removed 1hr and water bowl 30min before the start of the trial, via the door.

2 (a)



- 5 (b)



Figure S2: The apparatus used for neophobia trial. (a) A photo of the novel objects used in the 9 neophobia trial: A: half a purple rubber ball, B: a plastic pink frog. (b) A photo of a blue tit 10 approaching the novel object in a food bowl (photo credits: Katherine Herborn).