1	A WEAK INDIVIDUAL SIGNATURE MIGHT NOT ALLOW CHICK CALL RECOGNITION
2	BY PARENT STONE CURLEWS BURHINUS OEDICNEMUS
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13	Short title: Vocal recognition in the stone curlew
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#### 1 ABSTRACT

2 Offspring recognition has been confirmed in many bird species and vocal signatures appear to be its 3 major component. Up to now most studies dealt with colonial species, but recent findings indicate that 4 similar recognition is also present in non-colonial birds. By integrating spectrographic analysis and 5 playback experiments we investigated for the first time parent-offspring vocal recognition in the stone 6 curlew (Burhinus oedicnemus), a highly vocal, crepuscular-nocturnal species that usually rears a 7 maximum of two precocial and nidifugous chicks. Even though the species is usually non-colonial and 8 non-cooperative, in Taro River Regional Park, Parma, Italy, breeding territories are densely packed with 9 distances between simultaneously active nests as close as 40 m, which creates the possibility that chicks 10 might be exchanged. Our analysis identified two main vocalizations in the chick call repertoire: (1) s-call, 11 a brief strangled soft sound often uttered when parents and chicks were at close quarter; and (2) c-call, a 12 chirping sound with a complex structure, which seems a long-distance contact vocalization. The acoustic 13 analysis showed that, even though the structure was remarkably different among chicks, c-calls were an 14 unreliable individual signature given their high intra-individual variability. This was confirmed by 15 playback experiments. Indeed, when presented simulated c-calling chicks, adults approached both to their 16 own and to foreign chicks with equal indifference. This results suggest that parent stone curlews are 17 unable to recognize their offspring by voice. Further experiments are needed to test whether the chicks are 18 able to recognize their parents' voice and, more generally, to understand the details of parent-offspring 19 communication in this species.

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21 Keywords: stone curlew, *Burhinus oedicnemus*, individual recognition, playback experiments,
22 spectrographic analysis.

23

## 24 Introduction

25 The theory of natural selection predicts that parents should not feed unrelated young, but rather should

26 invest solely in their own offspring (Hepper 1986); this explains the evolution of parent-young

27 recognition in many species whose young could intermingle while still dependent on parents (McArthur

28 1982). Many studies have demonstrated the existence of parent-offspring recognition in birds (Snow

1958; Peek et al. 1972; Miller & Emlen 1975; Burtt 1977; Balda & Balda 1978; McArthur 1982; Rowley
 1980; Beecher et al. 1981; Seddon & van Heezik 1993; Searby et al. 2004; Mulard et al. 2008). In most
 cases the mechanism of this recognition is based on individual vocal signatures, facilitating either mutual
 recognition (Lefevre et al. 1998; Insley 2003), recognition of parents by chicks (Charrier et al. 2001;
 Lengagne et al. 2001; Taylor & Perrin 2008), or recognition of chicks by parents (Stoddard & Beecher
 1983; Lessells et al. 1991).

7 In colonially breeding birds, where young are mobile and the likelihood of intermingling is high, parent-8 offspring recognition should have strong selection. Indeed, the use of individual vocal signatures has been 9 demonstrated for many colonial species (Lefevre et al. 1998; Jouventin & Aubin 2002). This phenomenon 10 has been explored deeply in swallows (Beecher et al. 1981; Stoddard & Beecher 1983; Medvin et al. 11 1993). In two colonial species -bank swallow (*Riparia riparia*) and cliff swallow (*Hirundo pyrrhonota*)-12 in which young form crèches that contain many dependent fledglings, offspring develop individually 13 distinctive vocal signatures and parents learn the calls of their young. On the other hand in two non-14 colonial species of swallows -barn swallow (*Hirundo rustica*) and Northern rough-winged swallow 15 (Stelgidopteryx serripennis)- nestlings do not develop distinctive call features as fledglings of these 16 species seldom intermingle with those of other broods. However, recent studies have found offspring 17 vocal recognition in non-colonial species. Long-tailed tits (Aegithalos caudatus) breed independently in 18 pairs, but most nests fail due to depredation. Failed breeders often re-nest, but may instead become 19 helpers later in the season and preferentially care for close relatives. This kin-biased helping occurs in 20 absence of reliable spatial cues and Sharp et al. (2005) showed that cooperatively breeding Long-tailed 21 tits can discriminate between kin and non-kin according to the individual characteristics of their contact 22 calls. The brown-headed parrot (Poicephalus cryptoxanthus) nests in hollows in trees, but upon fledging 23 the chicks are escorted to nursery areas in heavily foliated tree where they remain camouflaged. 24 Therefore, although nest site recognition may be sufficient to identify chicks prior to fledging, a 25 recognition problem arises when those chicks are mobile yet camouflaged. Taylor & Perrin (2008) 26 showed that these chicks are able to recognize their parents by voice. 27 In colonial birds the timing for development of recognition varies among species, but normally occurs

28 just before chicks are able to leave the nest area (Taylor & Perrin 2008). It is interesting to note that the

selective pressure acting in favour (or against) offspring vocal recognition seems to be modulated by
 many factors which can change also within species (Falls 1982). For example, Rautenfeld (1978)
 demonstrated that for herring gulls (*Larus argentatus*) nesting on cliffs, where intermingling is
 impossible, recognition was lacking until the chicks were 14 days old, while Tinbergen (1953) showed
 that herring gulls nesting on the ground in close proximity to other nests can recognize their chicks 5 days

6 after hatching.

7 The aim of our work is to investigate parent-offspring vocal recognition in the stone curlew (*Burhinus* 

8 *oedicnemus*), a non-colonial and non-cooperative breeding species, characterized by a small clutch size (2

9 eggs, ca. 19% of female body mass) and a relatively long fledgling period (usually more than 40 days)

10 (Cramp & Simmons 1983; Nethersole-Thompson & Nethersole-Thompson 1986; D. Giunchi et al.

11 unpubl. data), which should favour the recognition between parents and chicks. We studied the population

12 breeding in Taro River Regional Park, Parma, Italy. At this site breeding territories of stone curlews are

13 usually densely packed with distances between simultaneously active nests sometimes even less than 40

14 m (nearest-neighbour distance between breeding pairs in the year 2010: median = 115 m, interquartile-

15 range = 95-160 m, n = 91; see also Giunchi et al. 2009). Such a high density might increase the likelihood

16 of chicks swapping, also given that even young chicks (less than 10-days old) often move more than 100

17 m from the nest sites, as observed also by means of radiotracking (Giunchi et al., unpublished data).

18 Moreover selective pressure should favour acoustic recognition if the probability of misdirected care is

19 high. This might be the case for this species as the chicks are mobile (precocial and nidifugous, Cramp &

20 Simmons 1983), active mainly during the night and rather cryptic during the day. Moreover, even though

21 parents stone curlew do not directly feed their young, they actively care for chicks e.g. by brooding them

22 and by providing food items (1-3 times/h according to Barros 1994 in Vaughan & Vaughan-Jennings

23 2005), which are placed on the ground near the chicks and often pointed out by the adult with its beak

24 (Cramp & Simmons 1983). Furthermore the stone curlew is a highly vocal species and which may be an

25 additional condition favourable for parent-offspring recognition based on vocal individuality. A key

26 behavioural component for recognition is an increase in vocal signature variation among individuals and

an increased coefficient of variation of inter-individual/intra-individual signatures (Aubin et al. 2004),

thus increasing information about individual identity (Beecher 1988). Therefore we investigated both (1)

the call repertoire of chicks and their vocal individuality, and (2) the ability of adults to recognize their chick by voice. To our knowledge, this is the first in-depth analysis of chick call repertoire of the stone curlew, as previous papers (reviewed by Vaughan & Vaughan-Jennings 2005) reported only anecdotal descriptions without any quantitative results.

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#### 6 Materials and Methods

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# 8 Study area, recordings and spectrographic analysis

9 Recordings and playback experiments were carried out in the period 2008-2010 in the Taro River 10 Regional Park, Parma, Italy (44.74° N, 10.17° E) where more than 80 pairs of stone curlews breed mainly 11 in the dry gravel riverbed (Giunchi et al. 2009; Caccamo et al. 2011). Chicks were caught during the night 12 using a lamp and a dip net. Vocalizations were recorded after ringing procedures. Birds were put singly in 13 a wooden arena (35 x 35 x 37 cm) with a Sennheiser M67 shotgun microphone placed about 30 cm away, 14 connected to a Sony Mini Disc MZ-R30 (2008-2009) or to a Fostex FR2-LE digital recorder (2010). Even 15 though the distance between the microphone and the sound source was short, we used a shotgun 16 microphone in order to reduce the environmental background noise and to increase the signal to noise 17 ratio of the recordings. Each recording session lasted on average 15 min. Audio tracks were digitized at 18 44.1 kHz sampling rate, with 16 bit accuracy and stored on a PC in wav format (PCM). Spectrograms 19 were made using SEAWave 1.1 software (Pavan 1998) with parameters set as follows: FFT size = 1024, 20 overlap = 75%, window type = Gaussian, frequency resolution = 23 Hz and time resolution = 11 ms. 21

#### 22 Chick call repertoire and vocal individuality

Call repertoire was studied by analysing the spectrograms of 66 recordings on chicks between 11 and 30
days old. Power spectra were obtained using the function *meanspec* from the *SeeWave* package v. 1.6.1

25 (Sueur et al. 2008) running under R environment (R Development Core Team 2011). This function returns

26 the mean frequency spectrum (i.e., mean relative amplitude of frequency distribution) of a time wave;

- 27 parameters of the *meanspec* function were sampling frequency = 44100 Hz, FFT window length = 1024,
- 28 and range of frequency axis = 1-10 kHz.

In 2010, 14 chicks (also used for playback experiments, see below) were recorded twice on average 9.4 days apart (range 9-12 days). Recordings from seven birds were discarded because of the low number of calls (< 12) in one of these two recordings. The seven remaining chicks (age=17-22 days) were used to assess within-individual variability and are identified hereafter as the w-group. An additional 13 chicks (age = 14-29 days) recorded only once during the same year, identified hereafter as the A-group, were used to asses among-individual variability.

7 Only one call type -the c-call used for long-distance contact with adults - was considered for this analysis 8 (see Results below). Every chick was recorded for 15 min. Chicks uttered from a minimum of 25 to a 9 maximum of 300 calls, consisting of only one syllable, with an irregular call rate, rather variable among 10 and within chicks (see Results). For each recording we randomly selected 25 calls and then we cut 25 11 250-ms audio samples each containing one call. These samples were submitted to bandpass filtration 12 between 1 and 8 kHz to clean background noise, because preliminary analyses of power spectra revealed 13 that more than 95% of the vocalization was below 8 kHz. Audio tracks were silenced before and after the 14 call using Audacity v. 1.3.4 beta (Audacity Team, http://audacity.sourceforge.net/). Comparisons between 15 two call series -belonging either to the same or different chick- were performed by calculating a similarity 16 index between each call pair by means of the function covspectro from the SeeWave package with the 17 following settings: sampling frequency = 44100 Hz, window length = 1024 points, window type = 18 Hanning, number of covariance computed when sliding the two calls = 19. This function calculates the 19 covariance between two spectrograms and returns the maximal covariance value (Hopp et al. 1998). For 20 each chick from the w-group we calculated an index of intra-individual similarity (hereafter s-INTRA) by 21 randomly pairing one call from the first and one call from the second recording, which was taken on 22 average 9.4 days apart, as mentioned above. Each call was used only once. One advantage of this method 23 is that the whole call is compared and no acoustic parameter is selected a priori, therefore avoiding 24 subjective choices. Furthermore the quantitative evaluation of the function *covspectro* is strongly 25 influenced by the harmonic part of the vocalization, which has frequency as well as amplitude modulated 26 over time and which may be relevant for individual recognition (Hopp et al. 1998; Sueur et al. 2008). The 27 inter-individual similarity (hereafter s-INTER) indexes were calculated by comparing each bird of the w-28 GROUP to the remaining six chicks from the same group plus all the 13 chicks from the A-GROUP, yielding a

1 total of 19 inter-individual comparisons. Similarity indexes were obtained by the same method described 2 above, randomly pairing one of the two sets of 25 calls used in the intra-individual analysis, derived 3 either from the first or the second recording (randomly chosen) of each chick from the w-group, and 25 4 calls belonging to each other 19 chicks. In this way, for each w-group bird we got 19 sets of 25 s-INTER 5 index values to compare with the series of 25 s-INTRA indexes previously calculated. Significantly lower 6 values of s-inter indexes with respect to corresponding s-intra values would mean that the variability of 7 vocal characteristics of calls within a chick is significantly lower than the variability among different 8 chicks; clearly this is a pre-requisite for discrimination and individual recognition (Terry 2005). It is 9 important to notice that according to this analysis design, which involves two sources of random variation 10 (i.e. random choice of one of the two call series used in the intra-individual analysis and random pairing 11 of calls for each inter-individual comparison), the S-INTER indexes were always calculated with different 12 datasets, even if w-group birds were considered twice in the analysis. For this reason two pairwise 13 comparisons involving the same birds (e.g. TH0375 vs. TH0384 and TH0384 vs. TH0375, see Table 1) 14 did not necessarily yield the same results. This approach was followed in order to reduce the degree of 15 pseudoreplication (Hulbert 1984) due to repeatedly considering the same bird in different comparisons. 16 Vocal individuality was also tested by performing a multivariate analysis on the same dataset (25 calls 17 from 19 chicks) used for the calculation of the s-INTER indexes, i.e. 25 calls from each of the 13 A-GROUP 18 chicks plus 25 calls derived from one of the two recordings (randomly chosen) of each chick from the w-GROUP. 1<sup>st</sup> and 3<sup>rd</sup> frequency quartiles, peak frequency (frequency with the maximal energy) and duration 19 20 were measured from each call using the *Seewave* functions *specprop* (for quartiles), *fpeaks* (for peak 21 frequency) and *timer* (for call duration; amplitude threshold = 1).

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## 23 Playback experiments

Playback experiments were performed in 2010. 37 chicks belonging to 26 different pairs were captured (age=12-26 d) and tagged with PIP3 single-celled leg mounting tags (Biotrack Ltd., UK; weight =  $2.5 \text{ g} \le$ 3% of bird body mass). According to visual observations collected before and after the first capture, all chicks from each breeding pair were captured and tagged. After ringing chicks were recorded using the techniques detailed in the previous paragraph. These recordings were then analysed and edited. We 1 selected a 2-min audio sample with low background noise from each chick. The relative amplitude values 2 (16 bit = track saturation) of these samples were digitally manipulated with slight increases or decreases 3 to obtain samples with nearly equal maximal values, while variability of amplitude within samples was 4 not changed. The call rate was left unchanged, as it could be an individually distinctive feature. Each 2-5 min sample was digitally filtered (high pass filter at 1 kHz) and pauses between calls silenced by means 6 of Audacity software to erase all background noises. From this 2-min samples we prepared the following 7 15-min playback sequence, which was the same in all experiments:

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9 17 recordings from chicks captured in 2009, which were not offspring of tested adults, were similarly 10 processed and used as foreign playback stimuli. For each experiment a different foreign stimulus was 11 randomly selected to avoid pseudoreplication (Kroodsma 1989; McGregor 2000). The order of 12 presentation of offspring or foreign signals was systematically changed between tests. In case of sibling 13 pairs, the call sequence of only one of the two chicks was randomly selected for playback. 14 Playback experiments were performed on average 4.2 days (range = 2-8 days) after initial capture. Chicks 15 were recaptured in the early morning (within two hours after dawn) or in the late afternoon (within two 16 hours before dusk), put into cloth bags and then displaced more than 200 m under the care of one 17 researcher to avoid predation. The broadcast chain -consisting of 1) Sandisk Sansa m240 digital player; 2) 18 Sonic Impact T-Amp integrated amplifier; 3) Ciare CT 263 loudspeakers (frequency range: 250 Hz-25 19 kHz)- was placed 20-40 m from the capture site and hidden under a relatively big patch of riverine 20 bushes, as chicks often use this kind of vegetation as squatting place in response to any approaching 21 danger (Cramp & Simmons 1983; D. Giunchi et al. personal observations). Signals were played back at 22 44.1 kHz sampling rate, with 16 bit accuracy and at 80 dB spl (sound pressure level, reference pressure 2 23 x  $10^{-5}$  Pa), measured 1 m from the loudspeaker, with a AVM2050 analogue sound level meter (fast 24 response, A-weighting). A videocamera (Sony DCR-SR55) and a recording apparatus (Sennheiser M67 25 shotgun microphone connected to Fostex FR2-LE digital recorder) were put in front of the speaker at a 26 distance of more than 10 m. The speaker was placed in dense vegetation so as to prevent parents from

1 approaching within 5 m from behind. Distance thresholds (5 and 1 m from the speaker, see below) were 2 indicated by means of stones or twigs used as markers. When possible, birds were also observed by 3 means of a telescope (20x-60x) from a hide located more than 400 m from the speaker. In each 4 experiment, the researcher started the videocamera and broadcast of playback sequence, walked away at 5 least 150 - 200 m, and hid in riparian vegetation. After 15 min, the same researcher returned to the 6 speaker, started the second playback sequence and then returned to the same hiding place. Broadcasts of 7 the two sequences were thus separated by about 10 min. At the end of the experiments, the apparatus was 8 removed and chicks were returned to the parents' territory in the same location at which they were 9 captured. On the whole, breeding pairs were disturbed for less than 1 h.

Our initial sample size was 26 pairs, but results from one pair was discarded because the chick did not call at all during the first recording and three more were discarded because chicks were depredated before we could conduct the experiments. We performed a total of 17 experiments, but two tests were discarded because of technical problems. Thus 15 tests were included in the analyses, 8 in which parents were tested first with their own offspring signal and 7 with the signal of a foreign chick. As control, we performed 5 blank tests using the same procedure, but without any broadcast both during the first and the second 15min sequences.

17 Radio tags were removed on the day of the experiment, except for the 14 chicks recorded twice (see 18 above), which all survived the experiments and had tags removed the day of the second recording, on 19 average 9.4 days after the first capture date (range 9-12 days).

20 Parental response was analyzed from recorded videos by a researcher blind to the experimental protocol 21 (i.e., order of playback sequences, experimental vs control tests). Responses were recorded from the start 22 of each sequence to its end 15 min later. Birds were considered as responding when they moved at least 23 once toward the speaker to a distance of less then 5 m Given the secretive behaviour of the species 24 (Vaughan & Vaughan-Jennings 2006) and its sensitivity to disturbance (Taylor et al. 2007), it was not 25 possible to follow the birds from the start of playback, as parents hide within vegetation when disturbed. 26 For this reason we only measured the latency of first response (i.e. when birds moved for the first time 27 toward the speaker to a distance of less then 5 m) and distance of nearest approach to the speaker (<5 m) 28 or <1 m). The study was approved by the Animal Care Review Committee of the University of Pisa and

1 complied with current Italian laws on animal welfare.

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### 3 Statistical Analysis

4 Pair-wise comparisons between s-INTRA and s-INTER indexes were performed using the general symmetry 5 (GS) permutation test in the R-package *coin* 1.8-18 (Hothorn et al. 2006, 2008) which tests the symmetry 6 of a response for repeated measurements in a complete block design. We considered the 25 call pairs used 7 to calculate similarity indexes within and between chicks as blocks. All tests were one-tailed under the 8 hypothesis that s-INTRA indexes should be higher than s-INTER ones. p-values were calculated using 10,000 9 Monte-Carlo resample approximations. This approach produced one series of 19 tests (13 from the A-10 GROUP plus six from the w-group - see Table 1) for each of the seven w-group chicks. To control for the 11 family-wise error rate, p-values in each series were corrected by means of the false discovery rate (FDR; 12 Benjamini & Hochberg 1995; Pike 2010) which controls the expected proportion of false discoveries 13 among rejected hypotheses. 14 Multivariate analyses on the four parameters derived from chick calls (1<sup>st</sup> and 3<sup>rd</sup> frequency quartiles, 15 peak frequency and duration) were performed using the R-package rrcov 1.3-01 (Todorov & Filzmoser 16 2009). The comparison between inter- and intra-individual variability was done by means of the Wilks' 17 Lambda Statistic based on the Minimum Covariance Determinant (MCD) estimator, which is robust to 18 the influence of outliers (Todorov & Filzmoser 2010). Robust linear discriminant analysis based on 19 MCD were also performed (Todorov & Filzmoser 2009) and the overall probability of misclassification 20 were estimated by means of the leave-one-out cross-validation method. 21 Proportions were compared by means of chi-squared tests with p-value calculated by Monte Carlo 22 simulations of 10,000 replicates. Significance of behavioural changes obtained from the two experimental 23 signals (offspring and foreign) was examined using the exact version of the McNemar's test (Sokal &

Rohlf 1995; Fay 2010). Latencies to first approach to the stimulus were compared by means of the GS test

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#### 27 RESULTS

(see above) using adult pairs as blocks.

#### 1 Chick call repertoire

2 Spectrographic analysis revealed that stone curlew chicks have two main call types: 1) s-call, a brief 3 strangled soft sound with a broadband spectrogram often uttered when parents and chicks were at close 4 quarter (Figure 1), and 2) c-call, a chirping sound with a complex structure, characterized by a modulated 5 tonal-harmonic part and non-tonal components (Figure 2). C-calls can be selectively stimulated by adult 6 vocalizations, as indicated by the analysis of 24 audio tracks where at least one call of distant adult(s) (< 7 100 m) was recorded in the background (Figure 3). Indeed, 1-min after a 'kurlee' call (see Vaughan & 8 Vaughan-Jennings 2005 for an account of adult call repertoire), the number of c-calls increased 9 significantly with respect to the 1-min before (Z = -3.04, p << 0.001, N = 14, gs test). This is not true when 10 considering all the other types of adult vocalization (Z = 0.63, p = 1, N = 10, gs test). Spectrograms in 11 Figure 2A show a high variability of the c-call structure between different individuals, but we found 12 variability within individuals too (Figure 2B). Mean power spectrum of all chicks, calculated over 25 13 randomly selected calls for each chick, shows that the third quartile is at 3.1 kHz, the peak frequency is at 14 2.7 kHz, more than 95% of the total power is below 8 kHz (Figure 4) and the average frequency range of 15 the two first harmonics is from about 2 kHz to about 6.5 kHz (Figure 4 and see also Figure 2).

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#### 17 Vocal individuality of chick calls

Table 1 reports the values of all s-INTRA and s-INTER indexes showing the great variability in these data. In 68% of cases the s-INTER index is lower than the s-INTRA index, as would be expected if vocal individuality exists, but in 32% of cases the situation is reversed.

21 On average 44% of the s-INTER indexes between chicks are significantly lower than the correspondent s-

22 INTRA values (GS test, q < 0.05, adjusted FDR value, see Table 2). These results indicate that according to

this acoustic analysis we can successfully recognize individual chicks in less than 50% (range in

24 successful recognition from less than 6% to more than 80%, depending on the reference chick; see Table

25 2). It is interesting to note that even with chicks with a very high s-INTRA index (see for example chick

26 TJ2961, Table 1) it is possible to find cases among the s-INTER index that are equal to or even greater than

27 s-intra.

1 The results of the robust one-way MANOVA calculated on the four parameters derived from chick calls 2 indicate that inter-individual variability was significantly higher then intra-individual variability (Wilks' 3 Lambda = 0.045, chi-squared = 1124.6, df = 68.4, p << 0.001). However, the discriminating power of 4 these parameters, estimated by means of the robust linear discriminant analysis, turned out to be very low 5 (global misclassification probability = 0.63) and rather variable among chicks (range of misclassification 6 probability at individual level = 0.20-0.96).

7

# 8 Playback experiments

9 In 10 out of 15 playback experiments we recorded at least one positive response (i.e., one adult bird

10 approached within 5 m of the speaker), while results from 5 control experiments without playback

emission were all negative (chi-squared = 6.67, df = 1, p = 0.03). Chick calls do stimulate the approach of parents.

13 Table 3 shows the results of the 15 playback experiments, each with one foreign and one own offspring 14 stimulus in random sequence. In all but one successful trial, only one parent responded to the playback. 15 Both type of stimuli and order of presentation did not have a significant effect on the number of positive 16 responses (exact McNemar's test, p = 0.68 and p = 0.69 respectively). Latency to first approach recorded 17 in the 10 tests with at least one positive response is showed in Figure 5. There is no statistically 18 significant difference between own offspring and foreign chick playback (Z = -1.07, p = 0.28, Gs test) nor 19 between first and second playback calls (Z = -0.22, p = 0.83, GS test). We did not record any obvious 20 behavioural differences by parents approaching the two types of stimuli. The minimum approach distance 21 to the speaker was always less than 1 m, except in two cases (one to the foreign chick stimulus and one 22 towards their own offspring). Adults approached the speaker repeatedly and searched nearby, often 23 uttering soft calls as is typical of vocal behaviour of parents getting close to their offspring (see Figure 1). 24

### 25 **DISCUSSION**

26 We have found at least two main vocalizations in the chick call repertoire of the stone curlew. In our

27 experimental conditions the s-call is often uttered in close proximity to adults and this finding is

28 confirmed by some field observations of unrestrained chicks still receiving care from their parents (M.

1 Dragonetti & D. Giunchi, unpublished data). Instead, we have used the c-call to investigate the vocal 2 individuality of chicks and the parent-offspring recognition for the following reasons. This call is 3 stimulated by distant adult calling and therefore seems to function as a long-distance contact vocalization. 4 Spectrograms show a modulated harmonic structure which could be a good requisite for auditory 5 discrimination, because birds are quite sensitive to changes in frequency of acoustic signals and they have 6 an enhanced capacity in resolving the temporal fine structure of complex sounds, enabling discrimination 7 of subtle differences in vocalizations (Marler & Slabbekoorn 2004). Furthermore power spectrum 8 analyses revealed that bandwidth of the first two harmonics of c-call is about 4 kHz; signal bandwidth 9 exceeding 3 kHz has been shown to be critical in allowing discrimination and localization of sound 10 sources in owls (Saberi et al. 1999).

Vocal individuality was studied by comparing the acoustic structure of the c-call within and among chicks. Results showed a higher similarity within chicks than among chicks, allowing correct individual recognition in 44% of cases. However c-calls show marked intra-individual variability too, which often prevented individual distinction among chicks by acoustic features alone (see Table 2, chicks TH0400 and TJ2951). While acoustic analyses show that chick call structure is individually distinct, nevertheless it does not seem to be a reliable cue for identification, as both confirmed by cross correlation and multivariate analysis of the four considered acoustic parameters.

18 Playback experiments suggest that c-calls of stone curlew chicks stimulate an approach by adults, but 19 parent stone curlews respond to playback stimulus in the same way to calls of both their own offspring 20 and stranger chicks; all parameters considered showed no statistical difference between the two groups. 21 Since adult birds responded to both playback stimuli, a question arises about adult response to stranger 22 calls. Might it be an aggressive response against an intrusion? There are several reasons to answer this 23 negatively: we examined carefully all the videotapes of the experiments and no aggressive display (see 24 Cramp & Simmons 1983) was ever observed; the behaviour of approaching adults was exactly the same 25 in all experiments. Actually, in the Florida scrub jay Barg & Mumme (1994) found an aggressive 26 response towards unfamiliar young call with parameters clearly different from those recorded for their 27 own offspring. Stone curlew pairs that own a territory usually respond with an aggressive approach (e.g. 28 'high-upright' posture, Cramp & Simmons 1983), to the vocalizations and/or the presence of an adult

intruder (Cramp & Simmons 1983; M. Dragonetti & D. Giunchi personal observations), but this has not
 been reported towards foreign chicks. In our playback experiments chicks were caught and removed from
 their nesting territories for the whole duration of the test, therefore any stimulus for adults to search for
 their own offspring would certainly be very strong.

5 To conclude, while it is always difficult to interpret non-significant results, as birds discrimination ability 6 might not always lead to a difference in response to playback (see McGregor 1992, 2000 for references), 7 our results suggest that parent stone curlews are unable to recognize their offspring by voice, confirming 8 our findings about the unreliability of individual signatures of chick calls. This does not mean that a vocal 9 recognition mechanism is absent in this species, because many researchers have found that in some bird 10 species chicks are able to recognize their parents vocally and this phenomenon is mutual (Aubin & 11 Jouventin 2002; Mulard et al. 2008) or not (Storey et al. 1992; Charrier et al. 2001; Lengagne et al. 2001; 12 Taylor & Perrin 2008).

Therefore it is possible that stone curlew chicks are able to recognize their parent's voice and that parents use a recognition system based on different behaviour of their young. This kind of mechanism has been found in Herring Gulls by Knudsen & Evans (1986). Our finding that young stone curlews change their vocal behaviour on hearing adult calls is consistent with this hypothesis. However further field experiments are needed to test whether chicks are able to recognize their parents' voice and by what mechanism.

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# 1 LEGENDS TO FIGURES

3	Figure 1. Chick call repertoire. Spectrogram (above) and oscillogram (below) of s-call, a brief strangled
4	soft sound uttered by chicks near their parents and which was answered by parents.
5	
6	Figure 2. Chick call repertoire. Spectrogram (above) and oscillogram (below) of c-call, a chirping call
7	characterized by a complex structure with modulated tonal-harmonic part and non-tonal components. A.
8	Seven samples from different chicks; note the very high inter-individual variability of these vocalization.
9	B. Eight samples from the same chick; note the high intra-individual variability.
10	
11	Figure 3. c-calls are stimulated by adult 'kurlee' call. Boxplot of the number of c-calls 1-min before and 1-
12	min after a distant (< 100 m) adult uttered a 'kurlee' ( $N = 14$ ) or another call type ( $N = 10$ ). 'Hinges' = first
13	and third quartile; horizontal line = median; 'whiskers' = $\pm 1.5$ times the interquartile range.
14	
15	Figure 4. Power spectrum of c-call. Mean power spectrum of 25 c-calls randomly selected among all
16	chicks ( $N = 66$ ). y-axis = relative amplitude of the frequency distribution. Q25 = 1st quartile of the power
17	distribution; Q75 = 3rd quartile. Numbers are peak values in kHz.
18	
19	Figure 5. Latency of first response to playback. Box-plots of latency to first approach recorded in the 10
20	tests with at least one positive response. See Figure 3 for other details.

3 Table 1. Vocal similarity within and among chicks. Average similarity index values (SD) of 25 c-calls randomly selected from 20 recordings belonging to different 4 chicks. Bold data are reference values calculated using two subsequent recordings of the same chick (s-INTRA indexes), all other values are calculated on recordings 5 belonging to different chicks (s-INTER indexes). Symbols refer to results of statistical comparison between s-INTRA and s-INTER indexes calculated by means of the GS

6 test (see Methods for further details). +, q < 0.05; ++, q < 0.01 (adjusted FDR-values).

Group	Chick tags	TH0375	TH0384	TH0400	TJ2951	TJ2959	TJ2961	TJ2970
W-GROUP	TH0375	0.66 (0.10)	0.57 (0.07)++	0.58 (0.10)	0.66 (0.07)	0.46 (0.1)++	0.53 (0.10)++	0.59 (0.09)++
	TH0384	0.51 (0.08)++	0.67 (0.10)	0.52 (0.12)	0.61 (0.14)	0.55 (0.16)	0.66 (0.11)	0.50 (0.17)++
	TH0400	0.57 (0.10)++	0.60 (0.10)+	0.45 (0.13)	0.71 (0.08)	0.70 (0.11)	0.70 (0.10)	0.46 (0.11)++
	TJ2951	0.60 (0.10)+	0.65 (0.15)	0.57 (0.13)	0.62 (0.11)	0.54 (0.20)	0.62 (0.13)++	0.66 (0.13)
	TJ2959	0.54 (0.07)++	0.42 (0.07)++	0.38 (0.08)	0.57 (0.11)	0.61 (0.11)	0.41 (0.10)++	0.67 (0.09)
	TJ2961	0.60 (0.08)+	0.69 (0.09)	0.46 (0.14)	0.71 (0.08)	0.38 (0.14)++	0.71 (0.11)	0.61 (0.11)+
	TJ2970	0.64 (0.08)	0.76 (0.05)	0.71 (0.07)	0.54 (0.14)	0.62 (0.07)	0.47 (0.11)++	0.69 (0.13)
	TJ2967	0.44 (0.14)++	0.50 (0.14)++	0.46 (0.14)	0.58 (0.14)	0.57 (0.12)	0.45 (0.14)++	0.63 (0.08)+
	TJ2974	0.41 (0.07)++	0.32 (0.06)++	0.26 (0.05)++	0.34 (0.05)++	0.59 (0.06)	0.25 (0.05)++	0.45 (0.06)++
	P14_03A	0.53 (0.08)++	0.68 (0.12)	0.46 (0.21)	0.63 (0.08)	0.53 (0.18)	0.61 (0.14)++	0.60 (0.10)++
	TJ2955	0.62 (0.09)	0.64 (0.11)	0.60 (0.13)	0.67 (0.09)	0.50 (0.14)+	0.59 (0.13)++	0.68 (0.09)
	TJ2956	0.60 (0.09)++	0.66 (0.12)	0.57 (0.16)	0.63 (0.10)	0.67 (0.07)	0.58 (0.14)++	0.65 (0.11)
	P17_01A	0.62 (0.08)	0.65 (0.08)	0.69 (0.10)	0.66 (0.09)	0.66 (0.11)	0.59 (0.11)++	0.63 (0.10)
A-GROUP	TH0395	0.43 (0.07)++	0.63 (0.04)	0.60 (0.08)	0.49 (0.15)++	0.22 (0.07)++	0.55 (0.06)++	0.45 (0.08)++
	TJ2962	0.59 (0.08)++	0.70 (0.06)	0.72 (0.06)	0.46 (0.09)++	0.73 (0.07)	0.40 (0.06)++	0.64 (0.07)
	TJ2965	0.51 (0.05)++	0.54 (0.07)++	0.76 (0.05)	0.62 (0.12)	0.31 (0.06)++	0.76 (0.04)	0.40 (0.06)++
	TJ2966	0.60 (0.05)++	0.63 (0.11)	0.58 (0.13)	0.69 (0.09)	0.66 (0.10)	0.59 (0.12)++	0.74 (0.10)
	TJ2969	0.61 (0.07)+	0.76 (0.08)	0.63 (0.09)	0.64 (0.08)	0.67 (0.09)	0.61 (0.11)+	0.65 (0.09)
	TJ2975	0.58 (0.08)++	0.64 (0.11)	0.62 (0.13)	0.66 (0.11)	0.47 (0.15)++	0.61 (0.14)++	0.66 (0.11)
	P14_04A	0.52 (0.09)++	0.56 (0.11)++	0.67 (0.06)	0.69 (0.06)	0.32 (0.11)++	0.71 (0.07)	0.42 (0.11)++

- 1 Table 2. Summary of number and percent of statistically significant comparisons (as test, q < 0.05,
- 2 adjusted FDR-values) showing vocal similarity to be lower among different chicks (as expected if vocal
- 3 individuality exists) than within chick.

Chick tags	Number	Percent
TH0375	16	84
TH0384	7	37
TH0400	1	5
TJ2951	3	16
TJ2959	7	37
TJ2961	15	79
TJ2970	10	53
Mean	8	44

- 1 Table 3. Responses to playback stimulation. Responding birds = adult(s) approaching within 5 m of
- 2 speaker.

Experiment	Tested birds	Responding birds
Answer to own chick	15	6
Answer to foreign chick	15	8
Answer to first test	15	8
Answer to second test	15	6



Fig. 1



Fig. 2



Fig. 3



Fig. 4



Fig. 5