

1 **Ontogeny of leukocyte profiles in a wild altricial passerine**

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20

21 **Abstract**

22 Ecophysiological studies have highlighted the relevance of the avian immune system in  
23 individual fitness prospects in the wild. However, studies on the ontogeny of avian  
24 immunity are scarce. We analyse age-related changes in the cellular constitutive  
25 immunity throughout nestling development, as well as its relationship with sex and  
26 brood size. We found that cellular constitutive immunity could be affected by age, sex,  
27 brood size or daily rhythm. Early-stage nestlings relied more on cells of the innate  
28 immunity rather than on cells linked to the adaptive immune system. Cellular immunity  
29 may not be fully mature in fledglings, as reflected by differences in phagocytic cell counts  
30 with regard to adults. Beyond the age-dependent effects, agranulocyte cell counts were  
31 affected by sibling competition while granulocyte cell counts showed a daily rhythm. We  
32 also show that the heterophil to lymphocyte ratio was negatively related to body weight  
33 when nestlings become more independent. Our study contributes knowledge to the  
34 fields of developmental immunology and ecological immunology based on essential  
35 components of the cellular immune system.

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39 nestling development; *Sturnus unicolor*

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53

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56

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62

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64

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66

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69

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73 JM and LP-R carried out fieldwork. JM performed the cell counts. JM and CV analysed  
74 and interpreted the data. JM took the lead in writing the manuscript. All authors  
75 provided critical feedback and helped shape the research, analysis and manuscript.

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## 85 **Introduction**

86 Hosts develop different immunological mechanisms to protect themselves against  
87 pathogens, but their maintenance and effective functioning is costly (Sheldon and  
88 Verhulst 1996; Hasselquist and Nilsson 2012). Therefore, from an evolutionary and  
89 ecological perspective, we may expect that a high investment in immunity must be  
90 traded off against investment in other costly traits, such as growth during the nestling  
91 period (Van Der Most et al. 2011) or reproductive traits for adults (Ardia, 2005;  
92 Colominas-Ciuró et al., 2017), which require high energetic demands, in particular  
93 during the breeding season (Kulaszewicz et al. 2017). Field studies have used white  
94 blood cells (WBC) or leukocyte profiles to assess immune function and environmental  
95 physiological stress in wild birds (Davis et al. 2008; Masello et al. 2009). Thus, leukocytes  
96 involved with the innate immune system, such as monocytes and granulocytes  
97 (heterophils, basophils and eosinophils), could offer an important measure of the non-  
98 specific host immune function and health status in birds (Davis et al. 2008; Masello et  
99 al. 2009). Additionally, the acquired immune system, which involves a high metabolic  
100 cost due to immunological memory processes, informs us about long-lasting protection,  
101 being highly specific and acting more effectively against various pathogens, including  
102 ectoparasites, viruses and bacteria (Masello et al. 2009; Kaiser 2010).

103         Birds, as with other animal species, can balance resources based on the energy  
104 requirements delimited by their trade-offs between reproduction and self-  
105 maintenance, which may vary through different life stages (Lochmiller and Deerenberg  
106 2000), and even between sexes (Klein and Flanagan 2016; Roved et al. 2017). Since  
107 immunity varies ontogenetically, younger individuals are more exposed and quite

108 vulnerable to environmental pathogens, as their immune system is not yet fully  
109 matured, as well as because they are confined to nests that typically harbour  
110 ectoparasites and other pathogens (Fellah et al. 2013; López-Rull and Macías Garcia  
111 2015). Newly hatched birds are rarely exposed to any external pathogen, so it would be  
112 expected that their innate immune system is much more developed than the acquired  
113 immunity (Bar-Shira and Friedman, 2006; but see Killpack and Karasov, 2012), which  
114 would mature throughout nestling development. Beyond maternally-transferred  
115 antibodies produced by their mothers (reviewed in Hasselquist and Nilsson, 2009),  
116 nestlings start producing their own specific antibodies in response to exposure to  
117 foreign antigens over the first few days or weeks after hatching (Staszewski et al. 2007;  
118 Hasselquist and Nilsson 2012). Adaptive immune responses are gradually developed and  
119 regulated in response to several factors, such as environment, life stage or season (Evans  
120 et al. 2016); although there is evidence that both the acquired and the innate immune  
121 systems fully mature post-fledgling (Killpack and Karasov, 2012; Stambaugh et al., 2011;  
122 respectively). Although the immune systems of adult birds (e.g. Davis et al., 2004;  
123 Palacios et al., 2011; Råberg et al., 2003) and nestlings (e.g. Brommer, 2004; Soler et al.,  
124 2003) have received considerable attention, comparisons between the two ages are  
125 more rarely reported (e.g. Stambaugh et al., 2011; Tella et al., 2002). Indeed, these  
126 studies mostly focus on measurements taken at a specific age of nestlings, without  
127 considering the possible immunological differences that arise throughout nestling  
128 development.

129         Several studies have shown that females and males may differ in both immune  
130 responsiveness and parasite load (e.g. Klein and Roberts, 2015; Klein, 2004). This sexual  
131 bias could lie in the effect of sex hormones on the functioning of the immune system

132 (Grossman 1985), where testosterone could have an immunosuppressive effect in males  
133 leading to a higher parasite load and a lower immune response than in females (Folstad  
134 and Karter 1992; Foo et al. 2017). However, it is barely known whether this sexual-  
135 dimorphism in immunocompetence is already present at the first stages of bird  
136 development.

137         Beyond age- and sex-dependent effects, innate and adaptive immune responses  
138 could be conditioned by both the nutritional status during the nestling stage (reviewed  
139 in Alonso-Alvarez and Tella, 2001) and environmental growth conditions (e.g. Ardia,  
140 2007; Christe et al., 2001). Among these external factors, some studies have highlighted  
141 that chicks raised in large broods, where sibling competition is greater, may show  
142 reduced immunocompetence (Ilmonen et al. 2003). For this reason, analysing the  
143 impact of body condition and the position occupied by the individual within the size  
144 hierarchy of the nest is required to understand the factors affecting the maturation of  
145 the cell-mediated immune function in birds.

146         Here we report a detailed study of the ontogeny of the leukocyte profile, also  
147 exploring the effect of some key factors that may influence the maturation of the  
148 immune function of developing birds. We used the spotless starling (*Sturnus unicolor*)  
149 as a study species, a medium sized altricial bird. The specific aims of the present study  
150 are to analyse: 1) age-related changes in WBC profiles and H/L ratio throughout nestling  
151 development, 2) the relationship between WBC profiles and H/L ratio and key factors  
152 that may influence immune function (i.e. brood size, body mass and sex); 3) the  
153 differences in both innate and acquired immunity between fledgling and adult  
154 individuals. We expect that the proportion of the innate cell types (e.g. heterophils,

155 eosinophils, basophils, and monocytes) would decrease throughout chick's ontogeny  
156 while those related to the acquired immunity (e.g. lymphocytes) would increase.  
157 Because of the sex-specific energetic, physiological and behavioural costs of maintaining  
158 an effective immune system (Lochmiller and Deerenberg 2000), we expect a difference  
159 in immune function between males and females throughout the nestling and  
160 independence periods, where probably the larger sex pays higher costs. Finally, we  
161 expect a lower immune cell fraction in chicks from nests with a larger brood size (Chin  
162 et al. 2005), in particular in the smaller siblings of the brood.

163

## 164 **Materials and methods**

### 165 Study area and study species

166 This study was conducted during three consecutive breeding seasons (from 2009 to  
167 2011) in a nest-box population of spotless starlings located in central Spain (Soto del  
168 Real, Madrid, ca. 40°45'N, 3°48'W, 920-940 m above sea level). The study area is  
169 covered by a deciduous woodland of oak (*Quercus pyrenaica*) and ash (*Fraxinus*  
170 *angustifolius*) with abundant open areas used by grazing cattle. It exhibits a continental  
171 Mediterranean climate [Köppen–Geiger climate classification: Csb category (reviewed  
172 in Peel et al. 2007)] with hot, dry summers. Thus, breeding conditions become harsher  
173 as the season advances, where late breeding conditions are characterized by higher  
174 temperatures and scarcer food (Muriel et al. 2015). The spotless starling is a relatively  
175 long-lived, colonial and sedentary passerine species that shows sexually dimorphic  
176 characters and exhibits a facultative polygynous breeding system (Cramp et al., 1982-  
177 1994; Monclús et al., 2017; Moreno et al., 1999b). Females can lay up to two clutches



178 per season, the first one in early April and the second one about the end of May in our  
179 study area (López-Rull et al. 2011). Incubation usually starts before the last egg is laid  
180 (modal clutch size is five eggs, López-Rull et al., 2007). The nestling period lasts about  
181 21-22 days (Cramp et al., 1982-1994).

182

### 183 Field protocols and sample collection

184 During the pre-laying period (from early March until the first egg of the colony was laid),  
185 male and female adult starlings were caught by traps placed inside nest-boxes. From  
186 every individual captured, we took a blood sample by brachial vein puncture. A drop of  
187 blood was smeared on individually marked microscope slides and air-dried. From early  
188 April onwards, nest-boxes were inspected each day to determine laying date. Broods  
189 were visited several times a day from the 10th day after the beginning of incubation to  
190 determine the exact hatching date (i.e., date first nestling hatches from egg = day 1).  
191 Chicks were labelled by distinct down cuttings to carry out an individual identification,  
192 and they were measured on different time points: 1, 3, 6, 9, 12, 15, 18 and 21 days post-  
193 hatching. At these ages, we recorded body mass with a digital balance (Ohaus Scout II  
194 SC2020, China, accuracy = 0.1 g) and tarsus length with digital calipers (Mitutoyo  
195 Absolute, Japan, accuracy = 0.01 mm). All chicks were ringed at 6 days old with  
196 numbered aluminium bands, when their tarsus was large enough for this purpose. A  
197 small blood volume (50 µl) was collected from the nestling jugular vein using a 29-ga  
198 insulin syringe (U-100, 0.5 ml, TERUMO®, Terumo Corporation, Tokyo, Japan) to perform  
199 their molecular sexing and the respective blood smears. All blood samples were  
200 collected within 2 min after capture and prior to any further manipulation of the

201 individual to avoid any leukocyte alteration due to handling time (Davis 2005). The solar  
202 time at which the blood sampling was carried out was recorded in order to control for it  
203 the statistical analyses. Most of the broods were sampled at consecutive time points.  
204 Each chick of a given nest was sampled at its real age. That is, when a brood included  
205 nestlings of three days of hatching asynchrony (i.e. nestlings of three different ages), we  
206 visited these nests three consecutive days per sampling event in order to get all the  
207 siblings sampled at their own exact age. This allowed us to avoid confounding age and  
208 size effects derived from hatching asynchrony in our analyses (Muriel et al. 2019).  
209 Overall, we analysed samples of 22 adults captured in 2009 (11 males and 11 females),  
210 60 different chicks sampled in 2010 (30 males and 30 females repeated sampled  
211 between 1 and 21 days old) and 34 different chicks sampled in 2011 (16 males and 18  
212 females repeatedly sampled between 1 to 6 days old) (see Supplementary Table 1S).  
213 These nestlings belonged to 19 different nests in 2010 and 9 different nests in 2011.

214

#### 215 Differential leukocyte count

216 For identification of white blood cells (WBC), a drop of blood was smeared on one  
217 individually marked microscope slide. Once the blood had air dried, we fixed the slide  
218 by 3 min immersion in 100 % methanol and stained it using commercial Giemsa diluted  
219 with PBS pH 6.8 (1:2). Slides were examined under the microscope using the oil  
220 immersion objective (1000× magnification) to estimate the proportion of different types  
221 of leukocytes (Campbell and Ellis, 2007). Estimates of the total WBC were calculated per  
222 approximately 10,000 erythrocytes. Total count for each type of leukocyte was  
223 calculated by multiplying the total leukocyte count by the respective differential WBC  
224 proportion, which were classified as heterophils, eosinophils, basophils, lymphocytes or

225 monocytes. We also took the ratio of heterophils/lymphocytes (H/L ratio) and the total  
226 leukocyte count as a measure of physiological stress and immunity in birds (Gross and  
227 Siegel 1983; Maxwell and Robertson 1998). One person (JM) conducted all cell counts  
228 to eliminate variation between observers. Counts were done blind to any information  
229 from the individual.

230

### 231 DNA extraction and molecular sexing

232 For sex determination, DNA was extracted from the blood samples using ammonium  
233 acetate techniques (Bensch and Åkesson 2003), and diluted to a working DNA  
234 concentration of 25 ng/ $\mu$ l. This solution was used in a polymerase chain reaction (PCR;  
235 using the primers P2 and P8) to amplify a part of the CHD-W gene in females and the  
236 CHD-Z gene in both sexes (Griffiths et al. 1998). PCR products were electrophoresed for  
237 60–90 min at 100 V in 1.5% agarose gels stained with SYBR safe (Invitrogen, Carlsbad,  
238 CA) and were visualized under UV light, where one band was scored as male and two  
239 bands as female.

240

### 241 Statistical analyses

242 All calculations were performed in the r language v. 3.5.3 (R Core Team, 2019), and the  
243 significance level was set at  $\alpha = 0.05$  for all tests. We applied Generalized Linear Mixed  
244 Models with Poisson distribution using ‘glmer’ function for count data (for all cellular  
245 variables except for the H/L ratio, which was analysed by Negative Binomial models  
246 using ‘glmer.nb’ function) and Linear Mixed Models using ‘lmer’ function for continuous  
247 data (body weight) using “lme4” package (Bates et al. 2017). These repeated measures

248 analyses were run considering nestling age as a continuous variable (from 1 to 21 days)  
249 to allow testing for different trends (age-related increases, decreases or quadratic  
250 relationships), taking into account other factors such as sex, time of the day and brood  
251 size by including them as covariates in the models. In order to control for non-  
252 independence of individuals from the same brood, nest of origin was defined as a  
253 random effect affecting the model intercept, and the identity of the individual was  
254 entered as repeated factor. In addition, we created an obs\_effect variable (observation-  
255 level random effect) with a unique value for each observation to control overdispersion  
256 in the model (Harrison 2014). All biologically meaningful double interactions were  
257 included in the original model. For thrombocyte and differential WBC counts, for which  
258 we had data from 2010 and 2011, we pooled them in the same dataset because we  
259 found no significant year effect (all  $P > 0.13$ ). In order to evaluate the effect of body  
260 weight on cell counts but avoiding collinearity problems with age, we carried out  
261 generalized linear models where we analyse the effect of body weight on WBC and  
262 thrombocyte counts at day 6, when there is an inflection point for the growth curve and  
263 nestlings become more independent, and sexual size dimorphism is not detectable yet  
264 (see Results). Finally, with the aim of analysing the degree of maturation of the cellular  
265 immune system at fledging, we also compared cell counts between fledglings (aged 21  
266 days posthatch; 12 males and 9 females) and adults (11 males and 11 females). To do  
267 so, we included ran separate models for each cell type including age as a categorical  
268 variable with two levels (fledglings vs. adults), sex, and their interaction, also controlling  
269 for sampling time as a covariate. In all cases, initial models were simplified by  
270 subsequently following a backward stepwise procedure to remove non-significant terms  
271 (i.e.  $P > 0.05$ ).

272

## 273 **Results**

274 Within broods, body weight increased throughout nestling development (from 1 to 21  
275 days old) adjusting to a negative quadratic function ( $\chi^2 = 591.94$ ,  $P < 0.001$ , estimate (SE)  
276 =  $-0.2556 \pm 0.0105$ ), where body weight reached an asymptote in the final days before  
277 fledging. Male nestlings were heavier than females ( $\chi^2 = 3.64$ ,  $P = 0.056$ , estimate (SE) =  
278  $-1.8068 \pm 0.9463$ ), although this sexual dimorphism became particularly evident at the  
279 end of the phase of linear growth of the nestlings (Fig. 1).

280

### 281 Leukocyte profiles during the nestling period

282 Age significantly affected all immune variables considered with the exception of  
283 monocyte and total leukocyte counts (see Fig. 2, all  $P > 0.110$ ), although the sign of this  
284 effect depended on the type of WBC analysed (see Fig. 2, all  $P < 0.02$ ). Relative  
285 percentage of each type of leukocyte to total white blood cells counted along the chick  
286 ontogeny as well as in adult stage is detailed in the Electronic Supplementary Material  
287 (see Supplementary Fig. 1S). Heterophil levels decreased along the ontogeny (Fig. 2a),  
288 while basophil and lymphocyte levels increased (Fig. 2c and 2d, respectively). In turn,  
289 eosinophils respond to a negative quadratic (i.e. inverse U-shaped) effect of age,  
290 reaching the highest levels at 9 days old (Fig. 2b). That maximum in eosinophil levels was  
291 concomitant with the cross of the opposite trends of heterophils and lymphocytes. In  
292 fact, that cross in the trends is responsible for the abrupt fall in the  
293 heterophil/lymphocyte (H/L) ratio at the beginning of the nestling period (Fig. 2g).

294 The results about the effect of age, time and brood size on the total and  
295 differential leukocyte count, as well as the H/L ratio, are shown in Table 1. We detected  
296 a negative effect of brood size in the interaction with age on both lymphocyte and  
297 monocyte counts, where the level of these agranulocytes decreased as sibling  
298 competition increased during early development, but increased during late  
299 development. Eosinophil level responded to an interaction between age and sex, where  
300 males had lower levels than females during the early stage of development (up to  
301 approximately 9 days old), a sex-related difference that disappeared after 12 days of age  
302 (Fig. 2b). Beyond the age-related effects, the total WBC count, and specifically  
303 granulocyte counts (heterophils, eosinophils and basophils), were affected by the time  
304 of day in which the nestlings were sampled throughout their ontogeny. Thus, the levels  
305 of these white blood cells increased throughout the day, adjusting to a daily rhythm (see  
306 Supplementary Fig. 2S).

307 Thrombocyte count was similar in both sexes ( $\chi^2 = 1.12$ ,  $df = 1$ ,  $P = 0.287$ ) but it  
308 was affected by a positive quadratic (U-shaped) effect of age throughout nestling  
309 development ( $\chi^2 = 3.971$ ,  $df = 1$ ,  $P = 0.046$ , estimate (SE) =  $0.0055 \pm 0.0027$ ), especially  
310 increased number of thrombocytes towards the last phase of the ontogeny (Fig. 2h).  
311 Circulating blood thrombocyte levels changed depending on the time of day at which  
312 blood samples were taken ( $\chi^2 = 9.50$ ,  $df = 1$ ,  $P = 0.002$ , estimate (SE) =  $-2.8015 \pm 0.9086$ ).  
313 Thus, the presence of these cells decreased as the day advanced.

314

315 Effect of body weight on differential cell count

316 At 6 days-old, we found a negative effect of body weight on H/L ratio (Table 2). This  
317 effect seems largely attributable to the negative effect of weight on heterophil levels.  
318 We also found a negative impact of body weight on total leukocyte counts, which may  
319 be due to the lower levels of heterophiles and, to a lesser extent, the lower lymphocyte  
320 levels (Table 2) in the lighter nestlings of each brood as compared to their heavier  
321 siblings. Additionally, lighter chicks also had higher thrombocyte levels (Table 2). The  
322 rest of the cell types did not show a clear pattern of variation based on body weight,  
323 with the exception of a marginally positive effect on basophile levels (Table 2).

324

#### 325 Leukocyte comparison between fledglings and adults

326 When comparing the cellular counts of fledglings vs adults in order to explore the degree  
327 of maturity of the immune function of the former, we found that adults had higher levels  
328 of total leukocytes than fledglings. This pattern could result from the differences in  
329 heterophils and monocytes (Table 3, Supplementary Fig. 3S-a and 3S-e) between adults  
330 and fledglings. However, in the case of eosinophil and lymphocyte counts, the effect of  
331 age class showed an interaction with sex (Table 3), showing that adult males had lower  
332 levels than the other age-sex combinations (Supplementary Fig. 3S-by 3S-d). For the  
333 basophil count, age class also had an effect in interaction with sex (Table 3), but in this  
334 case males maintained intermediate levels irrespective of the age, while female  
335 fledglings showed higher basophil levels than adult females (Supplementary Fig. 3S-c).  
336 Overall, adults had a higher H/L ratio than fledglings (Table 3). As found before, time of  
337 day also had an effect on total WBC count, heterophils and lymphocytes (Table 3), which  
338 showed increasing levels as the day progressed.

339           Regarding thrombocyte counts, we found similar levels for both sexes ( $\chi^2 = 0.22$ ,  
340  $P = 0.633$ ) and higher counts in adults as compared to fledglings ( $\chi^2 = 5.40$ ,  $P = 0.020$ ,  
341 estimate (SE) =  $0.012 \pm 0.005$ ). We also found a decrease in thrombocyte counts as the  
342 day progressed ( $\chi^2 = 9.35$ ,  $P = 0.002$ , estimate (SE) =  $-4.414 \pm 1.442$ ).

343

## 344 **Discussion**

345 In the context of ecoimmunology, the age-specific variation of the immune functions in  
346 wild birds has been relatively understudied. However, it is important to understand these  
347 variations when evaluating the immune system of nestlings since interpretation errors  
348 could arise if early- and late-stage chicks are pooled together, or if comparative analyses  
349 are made between adults and chicks of early or late stages. It is known that the immune  
350 system develops during ontogeny and thus both relative leukocyte quantifications and  
351 H/L ratios are likely to differ between adults and nestlings (Quillfeldt et al. 2008; Palacios  
352 et al. 2009). In this regard, our study shows that heterophil numbers decreased  
353 significantly along the ontogeny, while the levels of lymphocytes and basophils  
354 increased. This age-dependent pattern makes H/L ratios of newly hatched chicks very  
355 different from fledgling birds, highlighting a wide range of variation for one of the most  
356 used immune activation indicators in field studies on birds (reviewed in Davis et al.,  
357 2008). Taking into account this time-dependent effect, we analyse the importance of  
358 each cell type based on its relative abundance at specific moments of ontogeny,  
359 reducing possible individual variations when considering nestling sex, brood size and  
360 day-time (e.g. Chin et al., 2005; Markowska et al., 2017).



361           Leukocytes are central in effecting innate and acquired immunocompetence in  
362 birds. A number of haematological tests, among them differential in leukocyte and  
363 thrombocyte profiles, can be used as indirect methods to assess the level of stress,  
364 infection, condition and change in several immune responses in birds (Masello et al.  
365 2009). In fact, many authors have studied cellular immunity through total and  
366 differential WBC counts, where total WBC has been interpreted as a signal of increased  
367 immunocompetence (Gustafsson et al. 1994). However, most of them focus on a single  
368 measure (Lobato et al. 2005; D'Amico et al. 2016; Merrill et al. 2019), dismissing  
369 potentially relevant effects of the development phase of the individual. By analysing the  
370 variation of the thrombocyte and WBC levels every three days since hatching, our results  
371 contribute knowledge to the development of the constitutive cellular immunity.

372           We found an age-related decrease in heterophil levels, whereas the number of  
373 lymphocytes increased as chicks got older. Heterophils were highly represented in new-  
374 born chicks. This makes sense as heterophils are an essential part of the first line of  
375 defence during initial stages of pathogen infections (Maxwell and Robertson, 1998). In  
376 hatchlings, innate immunity is much more developed than the acquired immunity  
377 (reviewed in Alkie et al., 2019; Smits and Baos, 2005). After hatching, continued  
378 exposure to a high diversity of new pathogens could stimulate the chick immune system  
379 (Spencer and Garcia 1995; Killpack and Karasov 2012), leading to an increase in  
380 lymphocyte levels. This could explain the opposite trend between heterophils and  
381 lymphocytes shown in this study. Therefore, newly hatched chicks have much higher H/L  
382 ratios than fledglings, mainly due to increased lymphocytes and decreased heterophils  
383 throughout the ontogeny. Heterophilic profiles and increased H/L ratios have been  
384 interpreted as a symptom of environmental and physiological stress (e.g. ectoparasites,

385 brood reduction, inflammation, etc.) and infection. Since variations in stress hormone  
386 levels (i.e. glucocorticoids) trigger changes in these leukocyte cells (reviewed in Davis et  
387 al., 2008), many studies have used the H/L ratio as a measure of stress in birds (Gross  
388 and Siegel 1983). However, we cannot conclude from this that younger chicks have  
389 higher stress levels than older ones since both the immune system components and the  
390 Hypothalamic-Pituitary-Adrenal (HPA) axis tissues are not yet fully mature (Killpack and  
391 Karasov, 2012; Torres-Medina et al., 2019, respectively), so that these comparisons  
392 studying stress could only be made between nestlings of the same age.

393         We have found a quadratic effect of age on the number of circulating  
394 eosinophils, which is coincident with the pattern reported by Burton and Harrison (1969)  
395 in domestic chickens. By contrast Palacios et al. (2009) have showed that eosinophil  
396 levels decreased with age in tree swallow nestlings. Males had a lower eosinophil level  
397 than females during the first week and a half of development. During this early critical  
398 stage of development, trade-offs between growth and immunity could explain this  
399 pattern (Van Der Most et al. 2011), as our results showed that males were larger than  
400 females from the early stages of ontogeny. No other cell type was affected by the chick  
401 sex, which is coincident with previous studies (Wilk et al. 2007). The change in patterns  
402 in eosinophil, heterophil and lymphocyte counts (around 9 days old) coincides with a  
403 moment in nestling physiology in which eyes are newly open and feathers start to grow  
404 (see Fig. 3). Increased mobility and the protective effect of feathers from this age  
405 onwards make chicks less susceptible to common blood-sucking ectoparasites of our  
406 population, such as the carnid fly *Carnus hemapterus* (López-Rull et al. 2007) or  
407 mosquito vectors of blood parasites (Muriel et al. 2018). It is possible that the observed  
408 thrombocyte increase, especially high towards the last phase of the nestling ontogeny,

409 is also due to the increase of nest-dwelling ectoparasites and blood-sucking flies, which  
410 produce small wounds when feeding on the chicks. This age-related increase in  
411 thrombocyte levels is in line with the findings made by Fairbrother and O’Loughlin  
412 (1990), who showed that thrombocyte numbers increased from 5 days of age to a peak  
413 at 18 days of age. The changes in the composition of WBC could also be due to age-  
414 related changes in the microbiota (which changes around day 9 in house sparrows, Kohl  
415 et al., 2019) or be a consequence of the catabolism of maternal antibodies that usually  
416 disappear from the offspring within 5–14 days old (Staszewski et al. 2007).

417         Our results also showed that basophil levels increased throughout ontogeny. This  
418 pattern is opposed to data from Burton and Harrison (1969), whereas Palacios et al.  
419 (2009) did not find any age-dependent effect on this granulocyte cell. Basophil count  
420 could be used as a good biomarker of condition and health (Vinkler et al. 2010). They  
421 are one of the main cells of innate immunity (Martin et al., 2006a), so we would expect  
422 that basophiles would be much more represented in the early stages of ontogeny than  
423 in the later ones (Bar-Shira and Friedman 2006). However, the age-related increase that  
424 we have found suggests that, since they are not precocial species (Burton and Harrison,  
425 1969), starling chicks are more exposed to ectoparasites such as *C. hemapterus*, whose  
426 abundance increases along chick growth (Liker et al. 2001). This result would be  
427 consistent with an experimental study relatively higher basophil levels on cliff swallows  
428 nestlings exposed to ectoparasites (Chapman and George, 1991).

429         All granulocyte cells showed an increase with the time of day at which blood  
430 samples were taken. However, these effects were not observed in agranulocyte cells.  
431 This diel variation led to an increase of the number of both total leukocytes and

432 heterophils, together with the corresponding rise in H/L ratio. This daily rhythm is  
433 supported by previous studies reporting cyclic changes in different components of the  
434 immune system (e.g. Markowska et al., 2017; Stinson et al., 1980), which could be  
435 mediated by melatonin secretion patterns (Siopes and Underwood 2008). These  
436 dynamics could be an adaptive response to a circadian rhythm of parasites (Martin et al.  
437 2001; Navarro et al. 2003; Villanúa et al. 2006) or could be explained by a cumulative  
438 effect of sibling competition during the day (Martínez-Padilla 2006). Whatever the cause  
439 of this diel variation, our results indicate that sampling hour should be recorded and  
440 controlled for in future studies on performing blood cell counts in wild birds.

441       When comparing adults (older than 1 year-old) with fledglings, we found higher  
442 thrombocyte and total leukocyte counts in adults. Although thrombocytes are the most  
443 abundant cell (excluding erythrocytes) found in the blood of chickens (St. Paul et al.  
444 2012), their counts are rarely given in the literature because they tend to clump  
445 (Campbell and Ellis, 2007). However, in addition to their hemostatic effects intervening  
446 in blood coagulation, thrombocytes also play a potential role in innate immunity and  
447 inflammatory response (Wigley et al. 1999; St. Paul et al. 2012). Previous studies  
448 comparing adults vs. young thrombocyte levels have reported contrasting results  
449 (Alonso et al. 1991; Howlett et al. 2002; Martin et al. 2006). The fact that the levels of  
450 thrombocytes, heterophils and monocytes are higher in adults than in fledglings in the  
451 spotless starling suggests that phagocytic components may require further maturation  
452 post-fledgling, as other functions of the immune system do (Killpack and Karasov, 2012;  
453 Stambaugh et al., 2011).

454           Regarding the sex-dependent effects on immunity, our data suggests that some  
455 components of the cellular immune system of males could be less represented than that  
456 of females, as previously suggested in our study species (Muriel et al. 2017). In  
457 accordance with our results, Tschirren et al. (2003) found a pronounced sexual  
458 dimorphism in the cell-mediated immune response with males showing a reduced  
459 cellular immunity than females. In fact, according to the immunocompetence handicap  
460 hypothesis (Folstad and Karter 1992), a previous study carried out in our starling  
461 population showed that the phagocytic function in males was strongly inhibited under  
462 moderate physiological concentrations of testosterone (Gil and Culver 2011). This could  
463 also explain the differences found between adult and young males, highlighting that the  
464 immunological costs derived from the trade-offs between reproduction and immunity  
465 may be greater than those between growth and immunity (Lochmiller and Deerenberg  
466 2000).

467

## 468 **Conclusion**

469 In conclusion, our study of free-living spotless starlings provide compelling evidence that  
470 cellular constitutive immunity throughout early development is associated with key  
471 biological factors such as age, sex or degree of sibling competition. In addition, we show  
472 that body weight is negatively associated to H/L ratio at day 6 post-hatching, when  
473 nestlings are in the phase of linear growth. Taking into account that age is essential when  
474 evaluating immunity by cell counts, our findings partially support the prediction that  
475 early stage nestlings rely more on cells of the innate immune system rather than on cells  
476 linked to the adaptive immune system, as previously shown in other studies (reviewed

477 in Alkie et al., 2019). However, although we have found enormous variability in cell  
478 counts throughout ontogeny, it seems that the cellular immune system is still not fully  
479 mature days before leaving the nest, since we have found differences in phagocytic cell  
480 counts between fledglings and adults. Future studies could test whether the variability  
481 reflected in our data about the development of the immune system could depend on  
482 environmental factors, such as food availability, climatic influences or pressures of endo-  
483 and ectoparasites. Since adults can have age-dependent reproductive strategies (Rebke  
484 et al. 2010; Oro et al. 2014; Muriel et al. 2019), it would also be interesting to include  
485 different classes of adulthood (from one year onwards) to have a wider perspective of  
486 the development of the cellular constitutive immunity in wild animal populations.

487

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493

494

495 **Fig. 1** Variation of body weight (g) throughout the nestling development (age expressed  
496 in days). Values represented are means  $\pm$  SE (white squares: female and black squares:  
497 males). The continuous line represents the estimate for males and the dotted line for  
498 females.

499

500 **Fig. 2** Variation of heterophil (a), eosinophil (b), basophil (c), lymphocyte (d), monocyte  
501 (e), and total leukocyte (f) count, as well as variation of H/L ratio (g) and thrombocyte  
502 (h) count (per 10,000 erythrocytes) throughout the nestling development (age  
503 expressed in days). Values represented are means  $\pm$  SE. For the eosinophil count (b),  
504 values represented are means  $\pm$  SE (white squares: female and black squares: males),  
505 where continuous line represents the estimate for males and the dotted line for females.

506

507 **Fig. 3** Different nestling growth stages during the first two weeks of ontogeny. Picture  
508 credit: Jaime Muriel.

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757 **Table 1.** Summary of final generalized linear mixed models for repeated measures  
 758 analysis showing the effect of age (days) on total and differential nestling leukocyte  
 759 counts, taking into account sex, time and brood size. In all cases  $df = 1$ .

<b>Dependent variable</b>	<b>Fixed terms</b>	<b>Estimate <math>\pm</math> SE</b>	<b><math>\chi^2</math></b>	<b>P</b>
<b>Total leukocyte count</b>	Age	0.031 $\pm$ 0.019	2.58	0.107
	Age <sup>2</sup>	-0.001 $\pm$ 0.001	4.34	0.037
	Time	0.714 $\pm$ 0.270	6.97	0.008
<b>Heterophils</b>	Age	-0.110 $\pm$ 0.009	126.03	< 0.001
	Time	1.258 $\pm$ 0.444	8.01	0.004
<b>Eosinophils</b>	Age	0.206 $\pm$ 0.028	50.63	< 0.001
	Age <sup>2</sup>	-0.009 $\pm$ 0.001	61.80	< 0.001
	Sex	0.345 $\pm$ 0.144	5.69	0.017
	Age x Sex	-0.028 $\pm$ 0.014	4.14	0.041
	Time	0.848 $\pm$ 0.390	4.73	0.029
<b>Basophils</b>	Age	0.060 $\pm$ 0.022	7.15	0.007
	Time	2.311 $\pm$ 1.036	4.97	0.025
<b>Lymphocytes</b>	Age	0.123 $\pm$ 0.046	7.10	0.007
	Age <sup>2</sup>	-0.005 $\pm$ 0.001	19.72	< 0.001
	Brood size	-0.272 $\pm$ 0.101	7.22	0.007
	Brood size x Age	0.021 $\pm$ 0.008	6.32	0.011
<b>Monocytes</b>	Age	-0.525 $\pm$ 0.219	5.73	0.016
	Brood size	-1.743 $\pm$ 0.739	5.55	0.018
	Brood size x Age	0.169 $\pm$ 0.064	7.00	0.008
<b>H/L ratio</b>	Age	-0.427 $\pm$ 0.037	127.6	< 0.001
	Age <sup>2</sup>	0.012 $\pm$ 0.002	36.61	< 0.001
	Brood size	0.156 $\pm$ 0.082	3.60	0.057
	Time	1.740 $\pm$ 0.522	11.11	< 0.001

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768 **Table 2.** Initial generalized linear models of body weight effects on differential leukocyte  
 769 and thrombocyte counts in 6-day-old nestlings. P-values considered significant ( $P < 0.05$ )  
 770 are in bold. In all cases  $df = 1$ .

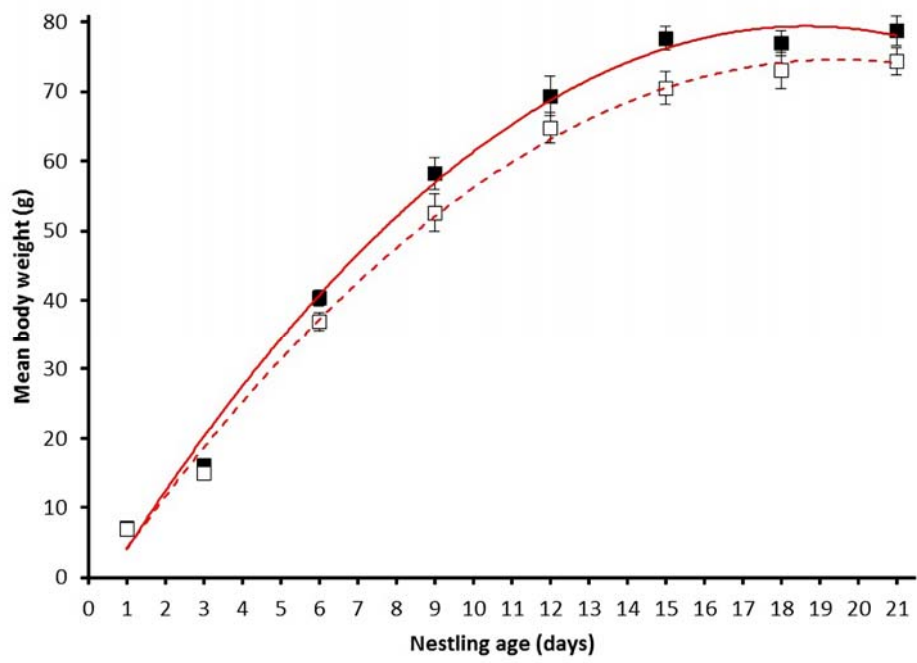
Dependent variable	Body weight effect		
	Estimate $\pm$ SE	$\chi^2$	P
Thrombocytes	-0.016 $\pm$ 0.004	11.7	< <b>0.001</b>
Heterophils	-0.015 $\pm$ 0.004	13.4	< <b>0.001</b>
Eosinophils	0.006 $\pm$ 0.004	1.50	0.221
Basophils	0.051 $\pm$ 0.029	3.23	0.072
Lymphocytes	-0.011 $\pm$ 0.004	6.78	<b>0.010</b>
Monocytes	0.022 $\pm$ 0.048	0.22	0.638
H/L ratio	-0.043 $\pm$ 0.016	6.58	<b>0.010</b>
Total leukocyte count	-0.007 $\pm$ 0.002	8.25	<b>0.004</b>

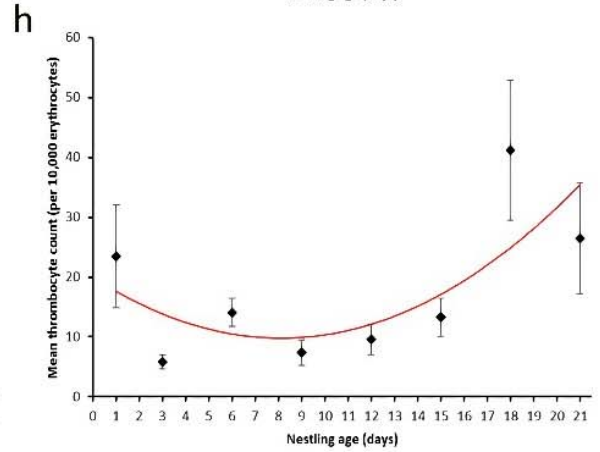
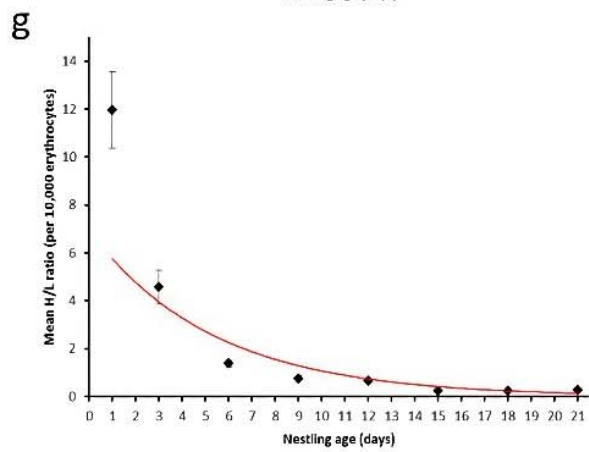
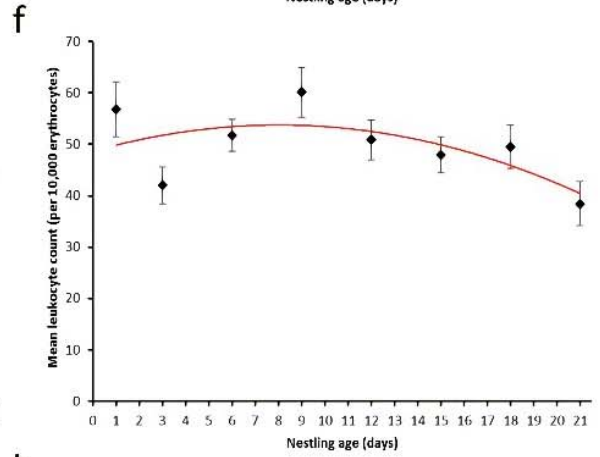
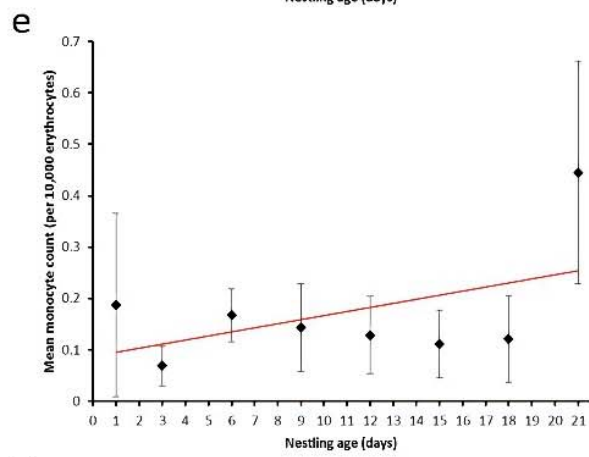
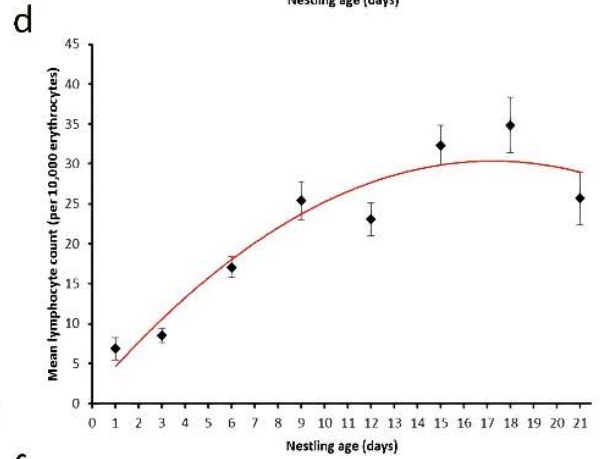
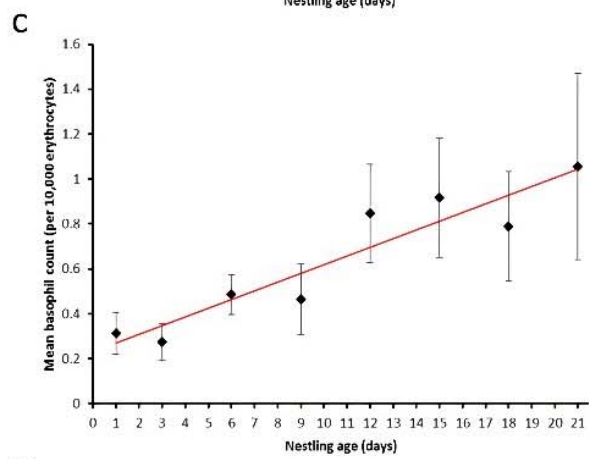
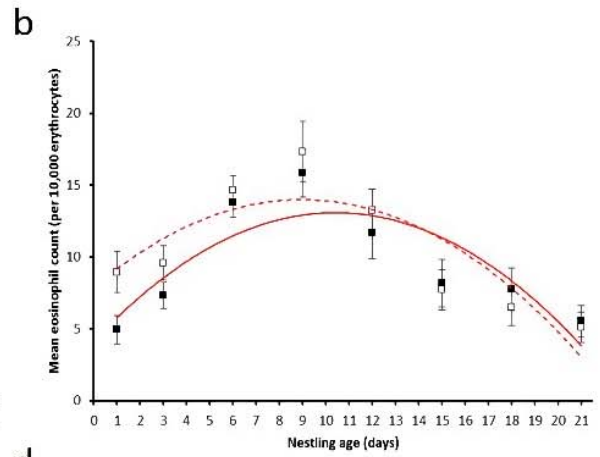
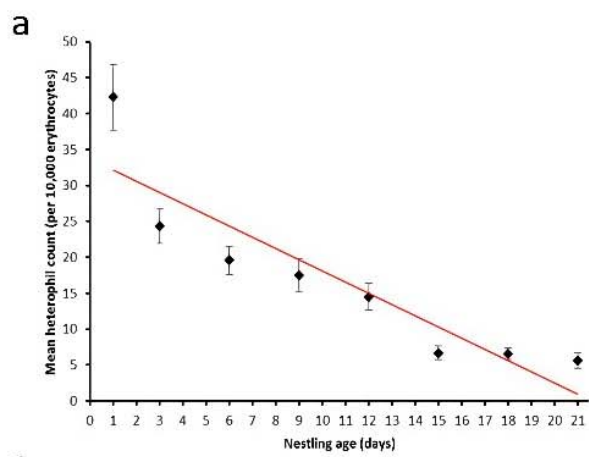
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772 **Table 3.** Summary of final generalized linear mixed models showing the effect of age  
 773 class (fledglings vs. adults) on total and differential nestling leukocyte counts, taking into  
 774 account sex and time of the day. In all cases  $df = 1$ .

Dependent variable	Fixed terms	Estimate $\pm$ SE	$\chi^2$	P
Total leukocyte count	Age (adults)	0.552 $\pm$ 0.196	7.91	0.004
	Sex (females)	0.268 $\pm$ 0.131	4.14	0.041
	Time	1.837 $\pm$ 0.671	7.49	0.006
Heterophils	Age (adults)	0.023 $\pm$ 0.004	32.18	< 0.001
	Time	2.916 $\pm$ 1.110	6.90	0.008
Eosinophils	Age (adults)	-0.025 $\pm$ 0.009	7.88	0.004
	Sex (females)	-0.471 $\pm$ 0.395	1.42	0.232
	Age (adults) x Sex (females)	0.012 $\pm$ 0.005	5.43	0.019
Basophils	Age (adults)	0.076 $\pm$ 0.819	0.008	0.926
	Sex (females)	0.984 $\pm$ 0.779	1.59	0.206
	Age (adults) x Sex (females)	-2.812 $\pm$ 1.426	3.88	0.048
Lymphocytes	Age (adults)	-0.341 $\pm$ 0.292	1.35	0.243
	Sex (females)	0.185 $\pm$ 0.227	0.66	0.414
	Age (adults) x Sex (females)	0.590 $\pm$ 0.313	3.55	0.059
	Time	1.487 $\pm$ 0.780	3.63	0.056
Monocytes	Age (adults)	0.024 $\pm$ 0.005	22.4	< 0.001
	Sex (females)	0.611 $\pm$ 0.296	4.25	0.039
H/L ratio	Age (adults)	0.021 $\pm$ 0.006	7.16	0.007

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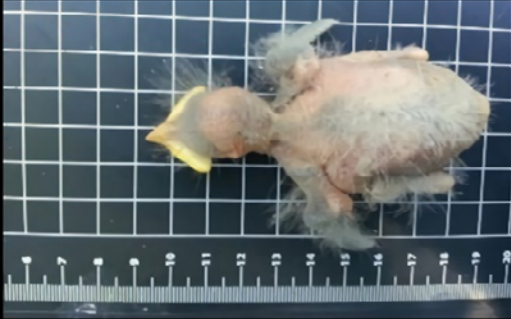




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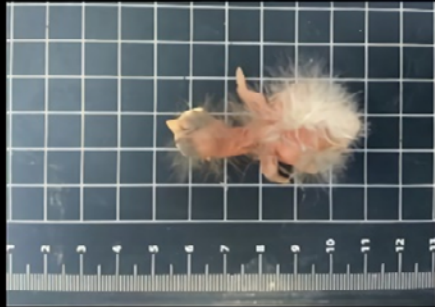
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Day 4



Day 1

1 **Electronic Supplementary Material**

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3 **Journal of Comparative Physiology B**

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5 **Ontogeny of leukocyte profiles in a wild altricial passerine**

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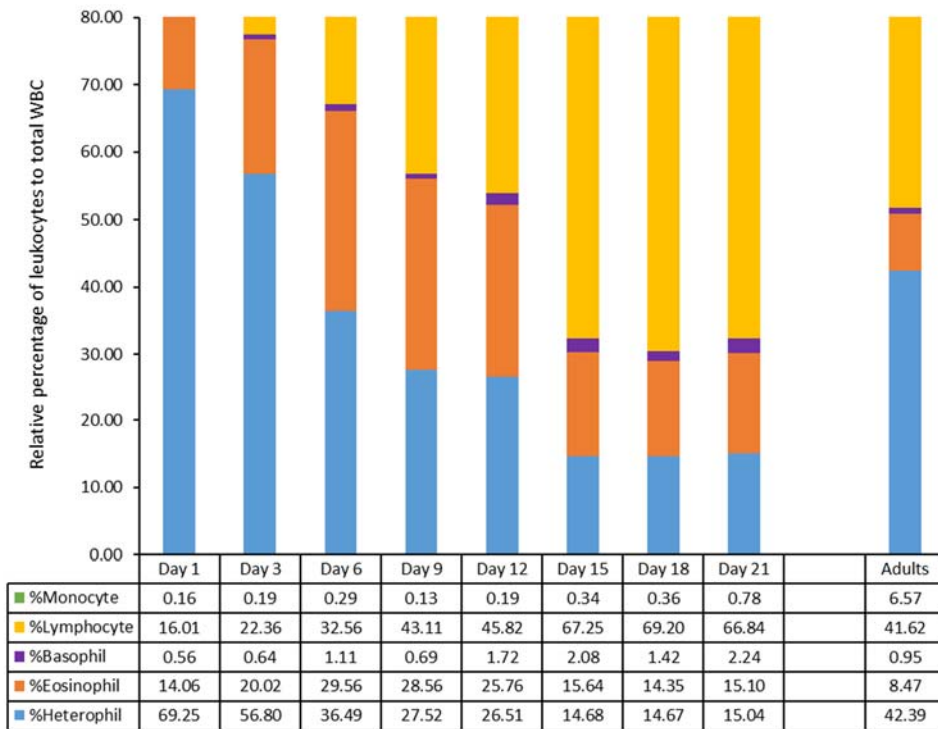
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17 **Supplementary Table 1S.** Number of blood smears analysed for each age and sex  
18 category in different years.

<b>Year</b>	<b>Age</b>	<b>Male</b>	<b>Female</b>	<b>Total</b>
2009	Adults (> 1 year old)	11	11	<b>22</b>
2010	Nestlings 1 day	2	1	<b>3</b>
2010	Nestlings 3 days	18	19	<b>37</b>
2010	Nestlings 6 days	25	28	<b>53</b>
2010	Nestlings 9 days	22	25	<b>47</b>
2010	Nestlings 12 days	18	22	<b>40</b>
2010	Nestlings 15 days	17	19	<b>36</b>
2010	Nestlings 18 days	17	16	<b>33</b>
2010	Nestlings 21 days	12	9	<b>21</b>
2011	Nestlings 1 day	16	18	<b>34</b>
2011	Nestlings 3 days	16	18	<b>34</b>
2011	Nestlings 6 days	16	18	<b>34</b>
<b>Total</b>		<b>190</b>	<b>204</b>	<b>394</b>

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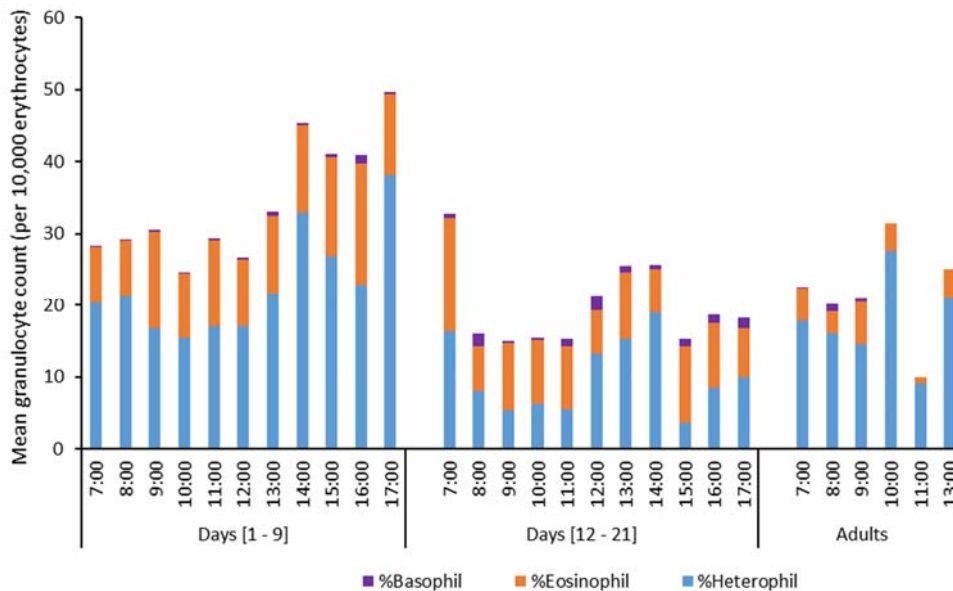




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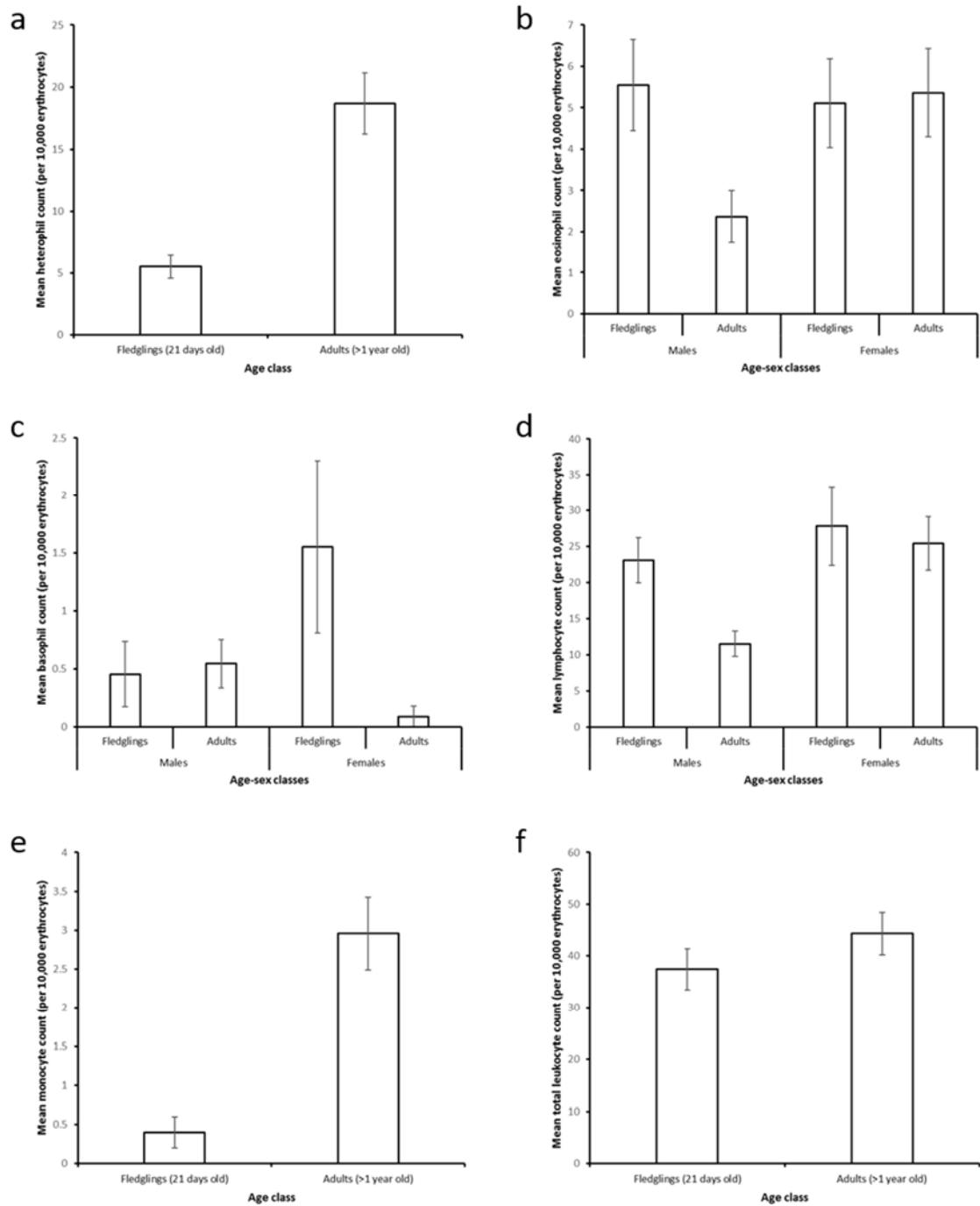
21 **Supplementary Figure 1S.** Stacked bar chart showing the relative percentage of each  
 22 type of leukocyte to total white blood cells counted along the chick ontogeny as well as  
 23 in the adult stage.

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26 **Supplementary Figure 2S.** Stacked bar chart showing the daily variation of heterophil,  
 27 eosinophil and basophil counts (per 10,000 erythrocytes) in three age classes (Days 1-9,  
 28 12-21 and adults) throughout the sampling time (expressed as solar time).



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30 **Supplementary Figure 3S.** Differences in heterophile (a), monocyte (e) and total  
 31 leucocyte (f) counts according to age class (fledglings Vs adults), and differences in  
 32 eosinophil (b), basophil (c) and lymphocyte (d) counts according to age class and sex  
 33 (fledgling-adult males vs fledgling-adult females). Values represented are means  $\pm$  SE.

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