1	Erythrocytes nuclear abnormalities and leukocyte profile of the
2	immune system of Adélie penguins (<i>Pygoscelis adeliae</i>) breeding at
3	Edmonson Point, Ross Sea, Antarctica
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5 6	Silvia Olmastroni ^{1,2} , Giulia Pompeo ¹ , Awadhesh N. Jha ³ , Emiliano Mori ⁴ , Maria Luisa Vannuccini ¹ , Niccolò Fattorini ⁴ , Nicoletta Ademollo ⁵ and Ilaria Corsi ¹
7 8 9	¹ Dipartimento di Scienze Fisiche, della Terra e dell'Ambiente, Università di Siena, Via Mattioli 4 53100 Siena, Italia
10	² Museo Nazionale dell'Antartide "Felice Ippolito" Via del Laterino 8 53100 Siena, Italia
11	³ School of Biological and Marine Sciences, University of Plymouth, Plymouth, PL4 8AA, UK,
12	⁴ Dipartimento di Scienze della Vita, Università di Siena, Via Mattioli 4, 53100 Siena, Italia
13 14	⁵ Istituto di Ricerca sulle Acque, Consiglio Nazionale delle Ricerche (IRSA-CNR), Via della Mornera, 25, 20047 Brugherio, Italia
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24 25	Corresponding author: <u>silvia.olmastroni@unisi.it;</u> tel. +39 0577 233775; <u>ORCID 0000-0002-9319-9914</u>

26 Abstract

Antarctic seabirds well adapted to extreme environments often deal during their life cycle with sub-optimal 27 conditions and occasionally with severe environmental stress. Climate changes, pollution, habitat loss, 28 increasing human presence can all significantly affect organism's health status from molecular to individual 29 up to population level. In the present study, erythrocytes nuclear abnormalities (ENAs) and white blood cells 30 31 (WBC) differential were investigated in 19 adults of Adélie penguin (Pygoscelis adeliae) breeding at 32 Edmonson Point, Antarctic Specially Protected Area (ASPA n. 165) in the Ross Sea. Micronuclei (MN) 33 accounted for 10.50% of observed abnormalities in penguin erythrocytes while kidney-shaped nucleus 34 (KSN) was the most abundant (20.88%). Heterophils (HE) were the most common WBC (36.93%) in agreement with the generic avian leukocytes profile while eosinophils (EO) were the lowest (7.45%). A low 35 number of lymphocytes were detected resulting in a higher heterophils to lymphocytes ratio. ENAs and H:L 36 ratio are confirmed as reliable indexes of penguin's health status since they reflect their individual adaptation 37 during breeding season. These baseline data will be useful for future studies as indicators of penguin's health 38 39 status mainly as response to environmental changes. 40 41 42

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- 44

45 Keywords: Adélie penguin, Antarctica, genotoxic damage, immune response, Ross Sea

47 Introduction

Organisms rarely experience optimal state in their natural habitat but for the most of their life they deal with variable conditions and occasionally with severe environmental stress. A variety of intrinsic factors can influence organism's physiological stress response such as reproductive status, age, sex, developmental or/and recent experiences. Whatever the source, physiological stress is a relevant parameter to consider when assessing animal welfare in both captive and wild populations (Davis et al. 2008).

Blood cells counts and classification, in particular erythrocytes' nuclear abnormalities (ENAs) and white 53 54 blood cells (WBC), are considered efficient tools for assessing genomic instability and immune status in 55 wildlife (Kursa and Bezrukov 2008). Although the mechanisms of the formation of ENAs are still little investigated in birds (Clark 2015), Van Ngan et al. (2007) promote the use of ENAs count for detecting 56 57 genomic damage caused by prolonged exposure to several physico-chemical stressors during organism's 58 lifespan. According to Kursa and Bezrukov (2008), occurrence of both micronucleus (MN) and in general 59 nuclear abnormalities (NA) should be considered useful tools for assessing genome instability also in 60 Antarctic birds since they represent a cellular reaction to natural and environmental stressors. MN occurrence may document what happen during erythrocytes' lifetime thus reflecting possible chronic effects. 61 62 Furthermore, the white blood cell count (WBC) reflects animal's immune status and response to stressful 63 conditions. The use of blood smears for detection of ENAs and WBC have several advantages such as low amount of blood needed with consequent low impact on animal health and a quick sampling procedure 64 which can be readily used also in extreme environmental conditions as for instance with polar birds (Dantzer 65 et al. 2014). Heterophils/ lymphocytes ratio (H:L) is considered a suitable indicator of organism's stress 66 67 associated to reproductive cycle, seasonal changes, injury and also to pathogens and parasites (Dufva and 68 Allander 1995; Krams et al 2012). Moreover, it reflects food and water deprivation, extremes temperature, constant light, long-distance migration and social disruption too. All these stressors result in an increased 69 70 level of heterophils (innate immune system), decreased number of lymphocytes (acquired immune system) and a high H:L ratio (Vleck et al, 2000). 71

Commonly, birds exhibit low level of spontaneous blood cell anomalies such MN, therefore it might be
 rather easy to detect any alteration due genotoxicants exposure or other environmental stressors (Zúñiga González et al. 2000, 2001).

Antarctic seabirds feed over wide geographical areas at different trophic level and therefore they are studied to monitor health conditions across large aquatic ecosystems and at different trophic levels. In turn they are able to reflect both natural and anthropogenic stressors (Mallory et al. 2010). Their health and physiological tolerance to stressors is closely influenced by their adaptation capability necessary to survive in their natural environment.

ENAs and immune status have been investigated in seabirds and in pygoscelid species breeding in the SubAntarctic and Antarctic Peninsula (Vleck et al. 2000; Kursa and Bezrukov 2008; D'Amico et al. 2014; De
Mas et al. 2015; Barbosa 2013). ENA and immune status have been linked to contaminants exposure
augmenting stress on penguin populations (D'Amico et al. 2014; Colominas-Ciuró et al. 2017) but also to

different stages of the breeding cycle, sex and individual condition and activities (Vleck et al. 2000; Morenoet al. 1998).

It is well known that climate changes are affecting the bioavailability of toxic contaminants in the wildlife altering the toxicokinetics due to an increase in temperature and salinity and leading to changes in organism' homeostasis and other physiological defence mechanisms (Noyes et al. 2009). Thus, during penguin's lifetime, contaminants exposure may vary according to the ecosystem changes. Different diets and foraging areas have been recognized also as major drivers for genome instability of penguin species from Antarctic Peninsula (De Mas et al. 2015).

In the background of above information, the present study investigates for the first time the occurrence of ENAs and WBCs in blood cells of an Adélie penguin (*Pygoscelis adeliae*, Hombron and Jacquinot 1841) population breeding at Edmonson Point, an Antarctic Specially Protected Area (ASPA n. 165) localized in the Ross Sea. The Adélie penguin is considered a keystone species of the Antarctic environment and currently most affected by environmental changes such as sea ice extent anomalies in different Antarctic regions (Ainley 2002; Olmastroni et al. 2004; Emmerson and Southwell 2008; Ropert-Coudert et al. 2013; Ducklow et al. 2013; Ballerini et al. 2009, 2015; Cimino et al. 2016).

99 In comparison with other Antarctic territories as for instance Antarctic Peninsula, the Ross Sea is still considered a pristine area (Halpern, 2008) and recently partially included in a Marine Protected Area (SC-100 101 CAMLR, 2016) to be preserved from increasing human activities. On the other hand, human pressure has 102 increased significantly in the last twenty years mainly due to increase in fishing, tourism and number of scientific bases (De Mas et al. 2015; Tin et al. 2009). Scientific research communities strongly required 103 104 protection for Antarctica from which the designation of the Ross Sea' MPA with the aim to preserve the marine ecosystem and biodiversity, as well as to limit and regulate current and future human impact. The 105 106 Ross Sea is the home of 38% of the global population of Adélie penguin, therefore it is mandatory to address 107 the current health status of population living in this territory in order to prove the efficacy of the MPA and to 108 monitor any potential impact in the future. While ecology of Adélie penguin breeding at the Edmonson Point 109 colony has been the focus of studies in the last 20 years (Olmastroni et al 2001, 2004; Pezzo et al 2007; 110 Ballerini et al. 2009, 2015), genome and immune stability have not been investigated so far. This issue inspired our study on the occurrence of ENA and leukocyte profile of the immune system, with the aim to 111 112 provide a baseline of health status of penguin living in the area.

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114 Materials and Methods

115 Study area and samples collection

The penguin colony is located at Edmonson Point (74 ° 20' S, 165° 08' E), Ross Sea, an ice-free area of about 6 km² along the Eastern slopes of Mt. Melbourne and *c*. 50 Km NW from Mario Zucchelli Italian Research Station (Fig. 1). The area has been occupied by Adélie penguin (*Pygoscelis adeliae*) from almost 3000 years BP (Baroni and Orombelli 1994) and breeding population size consisted of 3066 pairs in 2014/15 summer season. Since 1994 Edmonson Point is a monitoring site to carry out scientific research on the

Adélie penguin's ecology and to collect data for the Ecosystem Monitoring Program (CEMP) lead byCCAMLR Commission for the Conservation of Antarctic Marine Living Resources.

123 All data were collected during 2014-15 austral summer, following protocols approved by SCAR (SCAR's

124 Code of Conduct for the Use of Animals for Scientific Purposes in Antarctica, 2011) and under permission

125 from PNRA for working in an ASPA.

All blood and feather samples (19) were collected from adult penguins at the beginning of the breeding season (incubation period) when mostly males occupy colony. Blood was collected from apparently healthy penguins (i.e. not showing any sign of illness or injuries). No ectoparasites, feather or skin changes or emaciation were observed.

- 130 In order to reduce stress induced by capturing and handling each bird was restrained for the minimum time necessary (max 5 minutes) to carry out blood and feathers sampling and to record biometrics (Vleck et al. 131 2000). After sampling each bird was then released in front of its nest and observed until it returned to regular 132 breeding activity. Blood samples (one drop) were collected by venipuncture of the brachial vein using a 133 134 heparinized syringe with sterilized needle (22 gauge) according to Owen (2011). Up to five feathers per individual were sampled from the chest area. Feathers were conserved in sealed plastic bags at -20°C. 135 136 Penguins were weighted with a Salter scale to the nearest 50 g, and bill depth and bill length measured using 137 a calliper. Blood smears were prepared in the field immediately after collection using a drop of blood on a clean slide (15 min 10% HCl and rinsed with MilliQ water and oven-drying at 100°C). Slides were then 138 139 stored at +4°C in a slides' box.
- 140

141 Genome and immune analysis

Slides were processed at the University of Plymouth Ecotoxicology Lab for the analysis of genome instability. The following procedure was used: slides were fixed using (100% v/v) cold methanol for 30 min, stained with 10% Giemsa stain modified solution (Giemsa buffer tablets, pH 6.4 – BDH), and DPX Mounting Media (Leica Biosystems). They were then observed under a light microscope equipped with Digital Microscope Leica DMD108 Digital Microimaging Device with 40x objective. The images were acquired, stored and processed by using LAS program (Leica Application Suite).

Areas with a clear distribution of erythrocytes were identified for each slide as a well-defined and separatecytoplasm. Areas which presented overlapping cells were not taken into consideration.

150 Cell counting was carried out by taking the reference coordinates x, y and progressively moving from the left

to the right margin. Upon selecting the best images per slide, 1,000 erythrocytes were counted for each slide

according to Clark (2015) and the number of leukocytes and thrombocytes localized between them recorded.

153 A total amount of 19,000 cells was analysed.

154 In order to increase the identification of all known abnormalities in the nucleus of the erythrocytes of avian

species, in the present study blood cells of penguins were analysed according to the established method of

156 Kursa and Bezrukov (2008) already used for pygoscelid species by D'Amico et al (2014) and De Mas et al.

157 (2015). Erythrocytes nuclear abnormalities were determined as follows: (a) micronucleus, (b) lobed nucleus,

- (c) tailed nucleus, (d) two-lobed nucleus, (e) budding nucleus, (f) nucleus with cavity, (g) kidney-shaped
 nucleus, (h) unknown nuclear malformation. Their sum as ENAs was also calculated.
- White blood cells were classified along the five types of leukocyte according to Samour (2006): (a) heterophils, (b) lymphocytes, (c) monocytes, (d) basophils, (e) eosinophils were identified based on morphologic and staining characteristics according to the Table 22.10 reported in the chapter by Samour (2006). Erythrocytes and WBC were counted using ImageJ 1.6.0 24 (NIH, USA).

164 Sex determination

Sex of penguins (12 males and 7 females) was determined by molecular analysis on feathers except for one 165 166 individual in which blood was used. DNA was extracted using the PureLinkTM DNA Mini Kit (Invitrogen, by Thermo Fisher Scientific), following the manufacturer's instructions. The reliability of DNA extraction was 167 168 monitored through a negative control (no tissue added), and the DNA content determined through an Eppendorf Ultraviolet Spectrophotometer (AG Eppendorf). The chromo-helicase-DNA-binding-1 gene 169 (CHD1), found on sex chromosomes, was amplified which length varies among male (sex-chromosomes: 170 171 ZZ) and female (ZW) penguins (Zhang et al. 2013). The following specific primers for penguins were used: PL (5'-CCC AAG GAT GAT AAA TTG TGC-3') and PR (5'-CAC TTC CAT TAA AGC TGA TCT GG-172 3'). PCR was run through a 2720 Thermal Cycler (Applied Biosystems), following this profile: 3 min 94°C, 173 30 cycles of 35" at 94°C, 45" at 55°C and 3' at 72°C, followed by 7 min at 72°C. PCR reactions were 174 175 prepared with 0.5 μ L of Taq Polymerase, 1 μ L of each primer, 6 μ L of PCR Master Mix (with PCR buffer, 176 MgCl₂ and dNTPs: Genaid Biotech Ltd.) and about 20 ng of each DNA template. The electrophoresis was 177 run for 45' on a 3% agarose gel (Zhang et al. 2013).

178 Statistical analyses

- Descriptive statistical analyses including average, standard error (SE), range (minimum-maximum) of blood 179 180 smear parameters were carried out with R Studio software (Version 0.99.902 - © 2009-2016 RStudio, Inc). Differences between sexes were determined through the nonparametric Monte Carlo exact permutation test 181 182 for the equality of means, which computed all the possible permutations and uses the absolute difference in 183 means as test statistic (Anderson 2001). The Monte Carlo exact permutation test assumes that the two 184 samples are equal in distribution if the null hypothesis is true (Anderson 2001). Thus, we checked that each variable, both for male and female, followed the same distribution through a Kolmogorov-Smirnov test for 185 equal distributions if the null hypothesis was true. Significance level was set at $\alpha = 0.05$. Analyses were 186 performed through the software Past (Hammer et al. 2001). 187
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189 Results

- 190 Erythrocytes nuclear abnormalities in the penguin's blood smears from Edmonson point colony are listed in
- 191 Table 1. ENAs was found in 4.31 % over 19,000 mature erythrocytes analysed. Mean values of ENA varies
- from the lowest number of TLN (1.74 ± 0.40), which account for 4.03% of total ENAs to the highest of KSN
- 193 (9.0 ± 1.14) accounting for 20.88% (Table 1).

- Mean values of MN (4.53 \pm 0.52) resulted similar to that found for lobed nucleus (LN) (4.79 \pm 1.64) and budding nucleus (BN) (4.79 \pm 0.95) and account for 10.5% of total ENAs (LN and BN 11.11 % respectively).
- 197 Therefore, the most recurring ENA are KSN, NWC and TN, followed by LN, MN and BN. Amongst those,
- 198 MN is the lowest abnormality occurring with \leq 5 over 1,000 mature erythrocytes while NWC, TN, LN e
- 199 KSN exhibited higher variability. In addition, a small percentage showed unknown nuclear malformation
- 200 (UNM) (Table 1). The figure 2 shows all ENAs detected in Adélie penguin's blood smears classified
- according to Kursa and Bezrukov (2008) and De Mas (2015).
- Table 2 summarizes WBC identified in Adélie penguin's blood samples. White blood cells were 658 over
- 203 19,000 cells scored in penguin's blood smears. Heterophils (HE) were the most common WBC, followed by
- 204 lymphocytes (LY), basophils (BA), monocytes (MO) and eosinophils EO (Table 2 and shown in Fig. 2).
- Although not significant, toxic HE (THE) resulted higher than normal HE (NHE) in the total HE found (243
- over 19,000 erythrocytes scored). LY resulted lower than total HE (23.70% compared to 36.90%) while BA
- were 19.45% of the total WBC (Table 2 and Fig. 3).
- Total MO resulted 12.46% of the total leukocytes over 19,000 cells scored. Mean TMO numbers resulted higher than NMO even though not significantly different. The lowest WBC (Fig. 3) detected were EO
- 210 (7.45%). Heterophil: Lymphocyte ratio (H:L) was calculated and the mean value was 3.08 ± 0.87 .
- 211 The number of HE (*Monte Carlo* exact permutation test: p = 0.016) and the number of NHE (*Monte Carlo*
- exact permutation test: p = 0.012) were approximately four times greater in males (n = 12) than in females (n
- = 7 (Fig. 4), the rest of parameters analysed showed not gender differences.
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215 Discussion

The present study investigates for the first time the occurrence of ENAs and WBCs in blood cells of Adélie

penguin (*Pygoscelis adeliae*, Hombron and Jacquinot 1841) breeding at Edmonson Point, an Antarctic
Specially Protected Area (ASPA n. 165) localized in the Ross Sea.

- The most frequent ENAs described for bird populations including penguins have been observed in blood smears of Adélies from Edmonson Point (Kursa and Bezrukov 2008 and De Mas 2015); in particular peculiar nuclear anomalies as TN, KSN and TL were observed (Lucas and Jamroz, 1961) as well as BN and LN which are considered in interphase as precursors of MN formation and associated to cell death, genomic
- instability, or cancer development (Webster et al. 2009).
- MN frequency is also in the range of natural values reported for birds (from 0.40 to 4.30 over 1000 erythrocytes scored) (Zúñiga-González et al. 2001), thus suggesting a low genome instability of individual nesting at the Edmonson Point colony.
- The analysis of immune parameters also reveals that total number of WBC are within the normal range reported for birds (Kursa and Bezrukov 2008). By comparing ENAs and H:L ratio observed in Adélies from Edmonson Point with those documented in penguins breeding in Antarctic Peninsula (i.e. more anthropogenically impacted: Tin et al. 2009; SCAR 2010), some considerations can be made.
- ENA values result similar to those reported by De Mas et al. (2015) in Adélie penguin from Torgensen and Avian Islands (43.11 ± 27.59 ; 46.90 ± 46.50 ; 41.20 ± 40.10 respectively) whereas those of penguins from Yalour and King George Island result far higher (109.90 ± 80 and 72 ± 35.3). Lower values are on the contrary reported by D'Amico et al. (2014) in penguins from Potter Peninsula at Stranger Point (26.20 ± 3.20) scoring 40,000 mature erythrocytes out of 20 individuals.
- MN values (4.53 ± 0.52) are similar to the range reported in penguins breeding in colonies located in the Antarctic Peninsula (Yalour Island, 5.2 ± 4.1 ; Avian Island, 3.25 ± 3.7) but higher than those recorded in penguins from Torgensen Island (1.3 ± 1.5) and King George (1.9 ± 1.4) (De Mas et al. 2015).
- Interspecific comparison among *Pygoscelis* genus, shows lower MN values in individuals of *Pygoscelis papua* (Gentoo penguin) and *Pygoscelis antarcticus* (Chinstrap penguin) (De Mas et al. 2015 and reference
- 241 within) compared to Adélies from Edmonson Point (this study).
- Several ecological and environmental factors, such as species-specific sensitivity, diet, wintering areas and 242 243 exposure to toxic pollutants, could affect penguin's genome and immune stability (Bargagli 2005; Barbosa et 244 al. 2013; De Mas et al. 2015). According to De Mas et al. (2015), a different sensitivity to environmental disturbance of Gentoo and Chinstrap penguins compared to the strictly sea ice dependent Adélie penguin, 245 might have resulted in the development of a physiological defence mechanism able to cope better with 246 247 genotoxic agents. In addition, it has been hypothesized that some of the observed differences among species 248 could be related to the diet spectrum, which is wider in Gentoo penguins compared to Adélies (D'Amico et al. 2016). D'Amico et al. (2016) address also anthropic sources as responsible of observed ENAs recorded in 249 250 Adélie penguins from Stranger Point where high levels of heavy metals (Ni, Cu, Zn ad Se) have been
- detected in their feathers. Ancora et al. (2002) reported heavy metals (Cd, Pb and Hg) in stomach contents,

excreta, and feathers of Adélie penguins breeding at Edmonson Point. At that time a natural occurrence has been hypothesized for Cd and, to a lesser extent, for Hg, but not a direct anthropogenic impact of local sources. In fact the nearest scientific stations are far (*c*. 50 Km) from the Edmonson Point colony.

255 Concerning other source of anthropic pollution, contaminants stored in pack ice during years (via global 256 distillation process), could be released as a result of the seasonal melting also amplified by increasing 257 temperatures of surface waters as a consequence of climate changes (SCAR 2010). For instance, Persistent 258 Organic Pollutants (POPs) have been documented to cause alteration on immune system (Jara et al. 2018) 259 and to correlate with alterations in ENAs and WBC in penguin's species (Jara-Carrasco et al. 2015). In 260 Adélie penguin population breeding at Edmonson Point, legacy POPs have been reported in stomach 261 contents, blood samples and unhatched eggs by Corsolini et al. (2003, 2011, 2017), but overall toxicity was estimated to be low compared to other Antarctic areas. Emerging contaminants like PBDEs (Corsolini et al. 262 263 2017) and PFAS (Ademollo, unpublished data) were also detected in Adélie penguin blood samples and eggs 264 from Edmonson Point. Therefore exposure to contaminant in penguins breeding at Edmonson Point cannot be considered negligible; Antarctic penguin's colonies are also considered a secondary source of POPs 265 266 (Roosens et al. 2007). Nonetheless the impact of human activities that determines local inputs need further 267 investigations (Wang et al. 2017). A small seasonal field camp (average of 2 personnel unit) located 600 m far from the breeding groups represents so far the only local source of contamination at Edmonson Point 268 (Olmastroni 2002). 269

270 As far as immune status parameters, mean H:L value results higher compared to those reported by D'Amico 271 et al. (2014; 2016) in Adélie penguin from Stranger Point (1.10 ± 0.20 and 1.07 ± 0.11 respectively). In 272 particular, the percentages of LY, HE and EO result lower than those reported by D'Amico et al. (2014) 273 while MO and BA are 37% higher. MO and EO can be used to make a distinction among factors that alter 274 the leukocyte profile: stress, disease and infection. In fact, MO number increases in case of infections and 275 diseases since their main role is to phagocyte foreign particles. On the opposite, a reduction in EO number is 276 commonly a measure of stress reaction and rarely a response to disease. Early studies in human and mammals confirmed that glucocorticoids induced by stress often carry out a reduction on EO numbers 277 (Davis et al. 2008 and references within). THE were also detected and described by Jara-Carrasco et al. 278 279 (2015) as a cytological alteration consequent to exposure to various stress agents. THE in Adélie penguins 280 accounted to 53.50% of total HE and may suggest a bird's response to stress. The presence of THE associated with the abnormal high number of BA identified in Adélie penguin's blood smears may indicate 281 282 some disease occurring in the population under study. Mild lymphocytosis and moderate basophilia have 283 been associated with feather loss in penguins population from the Ross Sea (Grimaldi et al., 2014) which has been lately observed also in individuals from Edmonson Point in a similar percentage of occurrence 284 (Olmastroni personal. observation, 2018-19 Antarctic expedition). 285

286 Concerning WBC, higher values are reported by D'Amico et al. (2016) in Adélie penguin from different287 Islands around Antarctic Peninsula. However, among them, similar values as those measured in our study

288 were reported in Adélie penguins from Stranger Point in which in a comparable number of individuals was 289 analysed (n = 20). HE shows the highest percentage and this type of WBC are phagocytic cells that increase when the organism needs to cope with infections causing an increase in the level of H:L ratio. For instance, 290 291 HE are the first line of defence that an organism uses as immune response against gastrointestinal parasites 292 incorporated through the diet (D'Amico et al. 2016). An organism affected by heterophilia and lymphopenia presents the same leukocyte profiles as one who is experiencing infection and/or diseases. In addition, 293 294 despite anthropogenic pressure may have a strong influence on H:L ratio, this factor might have had less 295 influence in penguins monitored in the present study since penguins from Edmonson Point colony seem less 296 affected by organic pollutants compared to other colonies (Schiavone et al. 2009).

297 Although difficult at this stage to connect to any contamination or stress sources, this information will be 298 helpful for future investigation for comparison with different seasons, colonies and breeding stages. In 299 addition, some aspects of the breeding ecology need to be considered for assessing the health status of the penguin population. During the breeding stage, females usually arrive later at the breeding colonies (Ainley 300 2002), and fasting period and intraspecific competition are reduced if compared to mates. At the time of 301 sampling adults were incubating eggs or attempting to breed, according to Edmonson Point breeding 302 chronology (Olmastroni et al. 2000; Pezzo et al. 2007). Consequently males were fasting from their arrival at 303 304 the breeding colony (late October) and underwent competition with conspecifics for territory occupation, 305 nest building and mating. Thus, reproductive cycle, seasonal changes, fasting, long-distance migration, 306 competition for resources and injuries can all affect health status e.g. H:L ratio (Moreno et al. 1998; Vleck et 307 al. 2000; Minias 2019). Seasonal changes may influence organism's stress levels forcing individuals to use 308 more energy for thermoregulation. Vleck et al. (2000) reported that injured birds during fights for defending 309 their territory and/or nest, exhibit higher H:L ratio level than healthy birds. In addition, pathogens, ecto and 310 endoparasites are known to affect immune status. Individuals sampled in the present study were healthy 311 penguins, as their weights ranged 3100-5650 g, no sign of illness or injuries, and no ectoparasites, feather or 312 skin changes or emaciation were observed. There are no studies available on pathogens or parasites on 313 Edmonson Point population. We cannot exclude potential influence of disease or parasites hampering health 314 status in the studied population, but no evidence of blood pathogens was detected in the current study. In addition, studies on pygoscelids suggested absence of blood parasites and a low richness of ecto and 315 316 endoparasites for wild sub-Antarctic and Antarctic species (Jones and Shellam1999; Diaz et al. 2016; 317 Vanstreels et al. 2014, 2016).

Environmental natural stressors and increasing anthropogenic impact on wildlife are expected to grow in Antarctica in the near future, potentially by altering individual's level of stress and immune status. The present results depict a preliminary overall assessment of the health status of Adélie penguin's colony at Edmonson Point since it reflects the different components of an organism's response to its environment. ENAs and H:L ratio parameters represent a first baseline for future monitoring and assessment of genome and immune stability of Adélie penguin population in the mid Victoria Land area. Because high H:L ratio may represent a corticosterone-mediate response of organism to various exogenous stressors and an adaptive

- evolutionary trait (Minias 2019) future investigation and sampling will be carried out in the framework of the
- 326 ongoing research program PNRA2016 AZ1.11 (PenguinERA). Blood parameters such as estimations of
- 327 ENAs, WBC and H:L could be useful physiological and ecological indicators in monitoring and conservation
- 328 studies to assess population and ecosystem health in a changing environments.
- 329

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 mesopredator sensitive to environmental changes.

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340 Author's contribution

341 SO and IC conceived of this study and wrote up the manuscript, IC and ANJ planned genome and immune 342 lab analyses, SO and NA collected data, GP performed genome and immune lab analyses, EM and MLV 343 performed molecular lab analyses, NF performed statistical analyses. All authors helped to draft the 344 manuscript, read and approved the final manuscript.

345

346 **Ethical approval**

All applicable international, national and /or institutional guidelines for the care and use of animals were
followed. All procedures performed in studies involving animals were in accordance with the ethical
standards of SCAR's Code of Conduct for the Use of Animals for Scientific Purposes in Antarctica.

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351 Conflict of interest

We declare that we have no conflict of interest with the data presented in this scientific contribution.

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354 Figure and table captions

Fig. 1 Adélie penguin colony at Edmonson Point (74°20' S, 165°08' E), Victoria Land, Ross Sea.

Fig. 2 Erythrocytes nuclear abnormalities (ENAs) in Adélie penguin blood samples according to Kursa and
Bezrukov (2008): (a) micronucleus (MN), (b) lobed nucleus (LN), (c) tailed nucleus (TN), (d) two-lobed
nucleus (TLN), (e) budding nucleus (BN), (f) nucleus with cavity (NWC), (g) kidney-shaped nucleus (KSN),
(h) unknown nuclear malformation (UNM).

Fig. 3 Differential white blood cells (WBC): (a) Heterophil (HE), (b) Toxic Heterophil (THE), (c)
Lymphocyte (LY), (d) Monocyte (MO), (e) Toxic Monocyte (TMO), (f) Basophil (BA), (g) Eosinophil (EO).

- **Fig. 4** Mean ± standard error of n. of heterophils and n. of normal heterophils counted in males and females
- 364 Adélie penguin (females: n = 7; males: n = 12)
- 365 Table 1 Number of micronucleus (MN) and other erythrocytes nuclear anomalies (ENA) analysed per
- 366 19,000 mature erythrocytes of Adélie penguin's blood smears according to Kursa and Bezrukov (2008), and
- to De Mas (2015)
- **Table 2** White blood cells (WBC) per 19,000 mature erythrocytes in Adélie penguin's blood samples and H:
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