# <sup>1</sup> Sex-specific body mass ageing trajectories in adult Asian

## elephants

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### 21 Abstract

In species with marked sexual dimorphism and where one sex undergoes stronger 22 intrasexual competition, that sex is expected to age earlier or guicker. Here, we utilise 23 a unique, longitudinal dataset of a semi-captive population of Asian elephants 24 25 (Elephas maximus), a species with marked male-biased intrasexual competition, with males being larger and living shorter, and test the hypothesis that males show earlier 26 27 and/or faster body mass ageing than females. We show sex-specific body mass ageing trajectories: adult males gained weight up to the age of 48 years old, followed 28 29 by a decrease in body mass until natural death. In contrast, adult females gained body mass with age until a body mass decline in the last year of life. Our study shows 30 that sex-specific life-histories shape ageing patterns, consistent with the predictions 31 of the classical theory of ageing. 32

Keywords: Elephas maximus, long-term data, senescence, sex-differences, terminal
 decline

### 35 Introduction

Ageing – a decline in organismal functioning with age (Monaghan et al., 2008) – has been observed in many species (Jones et al., 2014). However, the onset and rates of ageing differ both between (Jones et al., 2014) and within species (Nussey et al., 2007) and between sexes (Clutton-Brock & Isvaran, 2007; Lemaître et al., 2020; Tidière et al., 2015). A main challenge in ageing research is to quantify and explain such differences in the onset and rates of ageing (Rando & Wyss-Coray, 2021).

42 In species with sex-specific intrasexual competition, classical theory of ageing 43 predicts that the sex with the highest intrasexual competition has a shorter lifespan and an earlier onset and/or higher rate of ageing (Williams, 1957). The rationale is 44 45 that high intrasexual selection often results in one sex showing conspicuous displays 46 or aggressive intrasexual behaviours, leading to increased mortality and a live fast, 47 die young pace of life (Bonduriansky et al., 2008; Clutton-Brock & Isvaran, 2007; Maklakov & Lummaa, 2013). For that sex, antagonistically pleiotropic genes or 48 49 deleterious mutations are not counter selected due to a weakened force of selection in late-life (Williams, 1957). Accordingly, in polygynous species with male-biased 50 intrasexual competition, males often die earlier (Lemaître et al., 2020) and age earlier 51 or faster than females (Beirne et al., 2015; Clutton-Brock & Isvaran, 2007; Douhard et 52 53 al., 2017; Nussey et al., 2009; Tidière et al., 2015). However, recent conceptual developments have shown that this association can be disrupted. This can occur for 54 55 example because of condition-dependent extrinsic mortality selecting particularly 56 high-performing individuals in the population (Chen & Maklakov, 2014) or 57 canalisation (*i.e.* the more a trait contributes to fitness, the less it should deviate from

optimal trait value, with respect to environmental variation (Flatt, 2005)), thereby
contradicting the theoretically expected earlier or faster ageing in males. The extent
to which such phenomena occur in nature remains unknown.

61 Here, we used a unique long-term dataset to describe sex-specific body mass ageing trajectories in a nutritionally unsupplemented semi-captive timber population of Asian 62 63 elephants (*Elephas maximus*) living in their natural environment in Myanmar. Body 64 mass is of interest in the study of ageing because it is positively associated with key 65 life-history traits such as reproduction and lifespan in many non-human species 66 (Briga et al., 2019; Gaillard et al., 2000; Hämäläinen et al., 2014; Pelletier et al., 67 2007). Therefore, the study of body mass ageing fits into the evolutionary framework 68 of ageing. Accordingly, in Asian elephants, seasonal variation in body mass was 69 positively associated with survival the following month (e.g. low body mass was 70 associated with low survival during dry season) (Mumby, Mar, Thitaram, et al., 2015). 71 Moreover, male Asian elephants benefit from being heavy during intrasexual 72 competition for dominance and mating (Sukumar, 2003).

However, we know almost nothing about body mass ageing in elephants despite the 73 74 interest in studying ageing in such a long-lived, social and sexually dimorphic non-75 human species. While females live in kin groups, adult males often roam solitarily, 76 undergo a more intense intrasexual competition for dominance and mating (Sukumar, 77 2003) and hence are bigger, heavier (Mumby, Chapman, et al., 2015), more 78 aggressive and less sociable (Seltmann et al., 2019) and shorter-lived than females 79 (respective median lifespans in this population: 30.8 and 44.7 years) (Lahdenperä et 80 al., 2018). Based on this male-biased intrasexual competition and shorter lifespan, 81 and following the classical theory of ageing (Williams, 1957), we tested the prediction

that males experience an earlier and/or faster body mass loss than females (Bonduriansky et al., 2008; Maklakov & Lummaa, 2013).

#### 84 Material and methods

#### 85 Study population

86 We studied the world's largest semi-captive Asian elephant population 87 consisting of around 3,000 individually-marked elephants owned by the government-88 run Myanma Timber Enterprise (MTE) (Leimgruber et al., 2008). Their birth, death, 89 maternal-lineage pedigree, and morphological measurements have been recorded 90 for almost a century by local veterinarians. These elephants are distributed across 91 Myanmar in forest camps and used as riding, transport and drafting animals. 92 Elephants work during the day and, at night, they socialise, mate and forage freely, 93 unsupervised in forests (Oo, 2010; Zaw, 1997). There are no husbandry procedures 94 and timber elephants are never culled. Calves born in captivity are cared for and 95 nursed by their biological mother and allomothers (Lahdenperä et al., 2016; Lynch et 96 al., 2019). Therefore, breeding rates are natural with no reproductive management. 97 Moreover, there is minimal food provisioning, but elephants benefit from veterinary care that consists of treatment of simple injuries and monitoring of working 98 conditions. 99

Both males and females are used in the workforce, and each working group of six elephants is composed of both sexes. Males and females follow the same government set limitations on taming age, working and retirement age, working-days per year, hours of work per day and tonnage pulled annually apply to both sexes, although it is possible that males might be used for somewhat different working tasks

105 at times (e.g. when use of tusks is required; only males can possess long tusks in Asian elephants). Pregnant females are given a rest period from mid-pregnancy 106 107 (around 11 months into gestation) until the calf is 1-year-old (Toke Gale, 1974), while 108 they and their calf are being monitored by their mahouts (individual caretakers and 109 riders) throughout this period. Following this break, mothers are used for light work 110 but are kept with calves at heel and able to suckle on demand until the calf is four or 111 five years old (Oo, 2010) at which point calves are assigned a rider, name, logbook and registration number. After the training period, elephants are used for light work 112 113 duties until the age of 17, when they enter the full workforce until retirement around 114 age 50. The MTE maintains their care and logbooks until death.

#### 115 Data collection and selection

Our analyses focused on age- and sex-specific variation in adult body mass 116 117 from age 18 onwards, in order to omit the phase during which elephants grow in height (Mumby, Chapman, et al., 2015) and to focus only on adult body mass age-118 specific variations. We compiled a total of 3,886 body masses on 493 individuals 119 120 (2,570 body masses on 322 females, and 1,316 body masses on 171 males). These data came from two sources: (i) body masses were either measured on elephants on 121 122 the field or (ii) estimated using height to the shoulder and chest girth (method in 123 Supplementary Information 1). For the first source, we extracted 1,901 body masses 124 of 347 elephants (1,297 measurements on 230 females, and 604 measurements on 125 117 males) and for the second source we estimated 1,985 body masses on 342 126 individuals (1,273 estimations on 226 females, and 712 estimations on 116 males - a 127 same individual can have both measured and estimated body masses). For all elephants of the 325 working localities ('township') sampled, sex, year of birth 128 129 (YOB), alive or dead status at the moment of the study, origin (captive-born or wild-

caught) and measurement season (hot: Feb-May, monsoon: Jun-Sep, cold: Oct-Jan 130 (Mumby, Mar, Thitaram, et al., 2015)) were known. The alive or dead status was 131 132 used to test for potential terminal decline. Among the 493 individuals considered, 5 133 males (63 observations) and 18 females (185 observations) were known to be dead. We had measurements during the last year of life for 2 males (7 observations) and 10 134 135 females (54 observations). Study elephants were aged 18 - 72 years (mean = 39.3) 136 and born 1941 – 1999. Age and cohort information were comparable between sexes, with 171 males (n = 1,316) born 1954 - 1999 and aged 18 - 64 years (mean = 37.4), 137 138 and 322 females (n = 2,570) born 1941 - 1999 and aged 18 - 72 years (mean = 139 40.2).

140 Most elephants of this semi-captive population get at least occasionally measured for height and chest girth, with no selection with respect to their age, sex 141 142 or condition. Body mass is measured only in camps provided with measurement 143 scales (mainly in regions with the highest concentrations of elephants and the best 144 accessibility). All elephants within the reach of those camps get weighed, again without any bias regarding their age, sex or condition. The logbooks containing these 145 146 measurements have thus far been translated from Burmese to English mainly from 147 the Sagaing region for logistic reasons, but again without any bias or pre-selection of 148 certain individuals.

In total, we obtained a median of 4.0 measurements/individual  $(2.5 - 97.5^{\circ})$ percentiles: [1.0; 36.4], followed during a median period of 2.8 years  $(2.5 - 97.5^{\circ})$ : [0.0; 36.6] on 493 elephants (n = 3,886). Two influential observations measured at age 18 and 23 were removed for one male because of particularly low  $\Delta$ age (Cook's distance = 0.61 and 0.25, mean of 0.001 on all males). Other observations for this male, all after age 50, were included.

#### 155 Statistical analyses

We investigated the age- and sex-specific variation in body mass in R version 156 4.1.1 (R Core Team, 2021), using the body mass (log-transformed to reach normality 157 158 of the variable and because of the allometric relationship between body mass and size) as a dependent variable with a normal error distribution. First, we tested in a 159 160 single model, whether there were sex-specific ageing trajectories using an interaction 161 term (Table S2). Given that this interaction was statistically significant, we compared the sex-specific ageing trajectories for both sexes separately. We did these analyses 162 163 using both general additive mixed models (GAMMs) with cubic regression splines 164 (but note that other spline functions gave similar conclusions as those shown here) 165 and general linear mixed models (GLMMs) with respectively the functions 'gamm' of the package 'mgcv' (v. 1.8-36, Wood, 2011) and the function 'Imer' of the package 166 'Ime4' (v. 1.1-27.1, Bates et al., 2015)). GAMMs allow more flexible ageing 167 168 trajectories than GLMMs, but the more constrained ageing trajectories in GLMMs 169 allow a less descriptive identification of differences in ageing trajectories (Fig. S1) 170 and both approaches gave consistent conclusions (see results section). We identified 171 the best fitting models using the model selection approach based on the second order Akaike Information Criterion (AICc) as implemented in the package 'MuMIn' (v. 172 173 1.43.17, Bartoń, 2021). In brief, the best fitting model has the lowest AICc value, with 174 other models within 4  $\Delta$ AICc being plausible and models becoming increasingly 175 equivocal up to 14  $\Delta$ AlCc, after which they become implausible (Burnham et al., 176 2011). Visual inspection of model residuals confirmed that these fulfilled all 177 assumptions of distribution and homogeneity without any influential data points or outliers (see above). 178

179 Within- vs. between-individual change

180 In all models, we accounted for non-independence of data due to repeated 181 measurements from the same individual by including elephant identity ('ID') as a 182 random intercept. The composition of the population can change with age for 183 example due to selective disappearance of certain (e.g. lighter or heavier) individuals, which can affect the age trajectory. In order to alleviate as much as possible this 184 185 problem in such a long-lived species, we decomposed body mass changes with age 186 into between- and within-individual changes following the approach developed by van de Pol & Verhulst, (2006) and van de Pol & Wright, (2009) using two terms: i) the age 187 188 at last measurement for each individual, which captures the between-individual 189 variations and *ii*) a ' $\Delta$ age' term (age at measurement minus the individual's mean age 190 for all measurements) capturing the within-individual changes with age. We mean-191 centered and standardised ' $\Delta$ age' so that *i*) individuals measured once all get a  $\Delta$ age 192 = 0 and hence contribute to the variance of the  $\Delta$  age intercept but not to its slope and 193 ii) to avoid collinearity and to have comparable variance for  $\Delta$  age and  $\Delta$  age<sup>2</sup> (Bolker, 194 2008; Zuur et al., 2009). Among the 493 individuals of our dataset, 105 individuals 195 had only one measurement. We included these individuals by giving them  $\Delta age = 0$ 196 (*i.e.* mean-centered) so they do not contribute to the coefficient but do contribute to 197 the variance along the Y axis on  $\Delta age = 0$ , diminishing the likelihood of a false 198 positive, and ii) do contribute to the coefficient of the age at last measurement term, 199 thereby avoiding a bias in the dataset from selecting only the longer-lived or most 200 monitored individuals.

201 Testing ageing trajectories

202 We tested several within-individual ageing trajectories, first using GAMMs, 203 which can provide curvilinear relationships and allow to describe trends, and using 204 GLMMs, able to detect breaking points if necessary, by testing linear, quadratic,

205 threshold and terminal models (Fig. S1) and we selected the ageing trajectory with 206 the lowest AICc. For GAMMs, we identified the age at which maxima occurred based 207 on the first-order derivative (= 0) using the function 'fderiv' of the package 'gratia' (v. 208 0.6.0, Simpson & Singmann, 2021). For threshold models (Fig. S1C), we followed the 209 approaches previously developed in Briga et al., (2019) and Douhard et al., (2017). 210 In brief, we first identified the best-fitting threshold age in a series of models, varying 211 the threshold in the ' $\Delta$ age' term between -35 to 22 years with intervals of one  $\Delta$ age (1 mean-centered  $\Delta age = 4.4$  and 4.5 years for males and females respectively) and 212 213 estimated the threshold and its confidence intervals using  $\pm 4 \Delta AIC$  age range. Then 214 we compared the best-fitting threshold model with the other ageing trajectories. 215 Sometimes, declines in trait value appear shortly before death (terminal decline). We 216 coded a 'terminal' change (Fig. S1D) as a binomial factor for whether an individual 217 died during the year following the measurement. We used a one-year-window to 218 avoid a possible seasonal covariation in weight and because it was the best fitting 219 time-window, but note that models with other time-windows gave consistent 220 conclusions (Fig. S3).

#### 221 Accounting for temporal and spatial variation in body mass

222 As body mass variation can be influenced by seasonal, spatial and within-223 individual factors, we tested whether body mass values were affected by (i) 224 measured or estimated, (ii) individuals were alive or dead, (iii) captive- or wild-born 225 and (iv) the measurement season. To this end we used a model selection approach, 226 performing a dredge on the best-fitting ageing trajectories for each sex to test for 227 confounding factors (Table S3). In our models, we included as random intercepts 228 individual identity to account for the repeated measurement of the same individual. 229 We also included 'township' to account for the spatial clustering of individuals across

230 Myanmar, although actually adding township worsened the model fit in most cases 231 (male GAMM:  $\triangle AICc = +1.5$ , GLMM:  $\triangle AICc = +1.8$ ; female GAMM:  $\triangle AICc = +0.6$ , 232 GLMM:  $\triangle AICc = -21.2$ ).

#### 233 **Results**

At the measurements' starting age of 18 years, males were on average 235 kg heavier than females, weighing respectively 2,541 kg [95%CI: 2,406; 2,683] and 2,306 kg [95%CI: 2,258; 2,355] and this difference was statistically significant ( $\Delta$ AICc = -122.6 in a GLMM with vs without sex as a fixed effect).

We identified the elephant's body mass ageing trajectories using general additive mixed models (GAMMs) and general linear mixed models (GLMMs) and both approaches gave consistent results. Both analyses showed that sexes have different body mass ageing trajectories (interaction term, GAMM:  $\Delta$ AICc = -65.7, Fig. S4, Table S2; GLMM:  $\Delta$ AICc = -47.0, Fig. 1, Table S2) and hence, we identified the ageing trajectories for both sexes separately.

244 For males, both GAMMs and GLMMs indicated a body mass gain from age 18 years 245 until their late-forties or early fifties (GAMM maximum: 54 years [95%CI: 53; 56], Fig. 246 S5; GLMM maximum: 48.2 years [4 AICc CI: 47.1; 51.6], Fig. 1A), followed by a 247 decline until death (Fig. S4A, Fig. 1A, Table 1, Table S4). In GLMMs, this maximum was shown through the best fit of a threshold model ( $\Delta AICc = -30.4$  compared to a 248 linear trajectory): males gained mass at a rate of 22 kg/year [95%CI: 19.4 23.7] or 1% 249 250 [95%CI: 0.9; 1.1] of males' mean body mass and then lost mass at a rate of 29 kg/year [95%CI: 14.9; 41.9] or 1.3% [95%CI: 0.7; 1.9] of males' mean body mass 251 252 (Fig. 1A, Table 2). Regarding the decline, neither GAMMs nor GLMMs showed

253 confident statistical support that it was terminal (*i.e.* determined by time before death 254 rather than age): in GAMMs, models with and without the terminal term were almost 255 equivalent ( $\Delta$ AICc = -0.01, Table S4) and in GLMMs adding a terminal term 256 worsened the model fit ( $\Delta$ AICc = +5.0, Table 1, Fig. S3A).

For females, both GAMMs and GLMMs indicated a body mass gain throughout their 257 lives until a terminal decline during their last year of life (Fig. S4B, Fig. 1B, Table 1, 258 259 Table S4). GLMMs indicated a mass gain of 9 kg/year [95%CI: 7.5; 10.4] or 0.35% [95%CI: 0.3; 0.4] of females' mean body mass ( $\Delta AICc = -6.7$ , Table 1, Table 2). Loss 260 261 of body mass occurred in the last year of life (GAMM:  $\Delta AICc = -12.2$ , Table S4, Fig. 262 S4B; GLMM:  $\triangle$ AICc = -4.9, Table 1, Fig. 1B, Fig. S3B) and consisted of 173 kg [95%Cl: 80; 263] or 6.8% [95%Cl: 3.2; 10.4] of their mean body mass (Table 2). For 263 comparison, the extent of the terminal decline in males, if any, is just over half that in 264 265 females at 96 kg [95%Cl: -19; 205] or 4.3% [95%Cl: -0.9; 9.2] of males' mean body 266 mass (quantified in the aforementioned best-fitting threshold model). Note that for 267 females and for both GAMMs and GLMMs, the second best model confirms the linear age trajectory, but excludes the terminal effect (GAMM:  $\Delta$ AICc = +12.2, Table S4; 268 269 GLMM:  $\triangle$ AICc = +4.9, Table 1). For males, the second best model (for GAMMs and 270 GLMMs) conserves the same threshold trajectory, but includes the terminal effect (GAMM:  $\triangle$ AICc = -0.01, Table S4; GLMM:  $\triangle$ AICc = +5.0, Table 1, Fig. S3B). Model 271 272 averaging on ageing trajectories within 7  $\Delta$ AICc (Burnham et al., 2011) confirmed the 273 ageing trajectories found, *i.e.* a threshold trajectories for males with a significant 274 decline of body mass from 48 years old onwards (GLMM:  $\beta = -0.06$  [95%CI: -0.09; -0.03]) and a non-significant terminal decline (GLMM:  $\beta = -0.04$  [95%CI: -0.10; 0.01], 275 276 GAMM:  $\beta$  = -0.04 [95%CI: -0.09; 0.01]). Similarly, model averaging performed on 277 models within 7  $\Delta$ AICc for females confirmed the ageing trajectory found, *i.e.* a body mass gain throughout life (GLMM:  $\beta = 0.015$  [95%CI: 0.01; 0.02]) with no significant decline before the last year of life (GLMM:  $\beta = -0.07$  [95%CI: -0.11; -0.03]). Also, we found no effect of none of the temporal and spatial confounding variables tested (Table S3).



Figure 1. Body mass ageing trajectories of (A) males (n = 1,316 measurements on 283 171 individuals) and (B) females (n = 2,570 measurements on 322 individuals) with 284 predictions of the best-fitting GLMMs (Table 1) with grey areas 95%CI. For males, 285 the thick dashed-line shows the threshold age at onset of the body mass decline (1.9 286 287 or 48.3 years) with thin dashed-lines the 4  $\Delta$ AlCc-Cl [46.6, 52.3]. For females, 288 measurements in the terminal year (red) are significantly lower (intercept) than measurements at other ages (blue). Note 1: the terminal slope is for illustration 289 purposes only and was not statistically tested. Note 2: the original x-axis is  $\Delta$ age, but 290 291 for simplicity, we presented here x-axis as age. For the original figure, please see 292 Fig. S2.

Table 1. Best fitting body mass ageing trajectories (bold) for males and females, using GLMMs for each model ageing trajectories ranked from the least to the most complex. AICc: second-order Akaike Information Criterion; ΔAICc: change in AICc relative to the best fitting model; k: degrees of freedom.

Model type	Model		Males			Females	
		к	AICc	ΔAICc	k	AICc	∆AICc
null	log(bm) ~ 1	4.0	-2,828.3	364.2	4.0	-5,442.9	160.7
linear	log(bm) ~ Δage + age-last	6.0	-3,162.0	30.4	6.0	-5,586.1	17.5
+terminal	log(bm) ~ Δage + age-last + terminal	7.0	-3,156.9	35.6	7.0	-5,592.3	11.3
age-last <sup>2</sup>	$\log(bm) \sim \Delta age + age-last + age-last^2$	7.0	-3,155.5	37.0	7.0	-5,598.7	4.9
+terminal	log(bm) ~ $\Delta$ age + age-last + age-last <sup>2</sup> + terminal	8.0	-3,150.4	42.1	8.0	-5,603.6	0.0
∆age²	$log(bm) \sim \Delta age + \Delta age^2 + age-last$	7.0	-3,176.8	15.7	7.0	-5,571.6	32.0
+terminal	$log(bm) \sim \Delta age + \Delta age^2 + age-last + terminal$	8.0	-3,171.2	21.3	8.0	-5,577.9	25.7
quadratic	$log(bm) \sim \Delta age + \Delta age^2 + age-last + age-last^2$	8.0	-3,170.1	22.3	8.0	-5,584.3	19.3
+terminal	$log(bm) \sim \Delta age + \Delta age^2 + age-last + age-last^2 + terminal$	9.0	-3,164.6	27.9	9.0	-5,589.3	14.3
threshold	log(bm) ~ Δage1 + Δage2 + age-last	8.0	-3,192.5	0.0	8.0	-5,580.5	23.1
+terminal	$log(bm) \sim \Delta age1 + \Delta age2 + age-last + terminal$	9.0	-3,187.5	5.0	9.0	-5,586.0	17.6
threshold (age-last <sup>2</sup> )	$log(bm) \sim \Delta age1 + \Delta age2 + age-last + age-last^2$	9.0	-3,185.8	6.6	9.0	-5,592.6	11.0
terminal	$log(bm) \sim \Delta age1 + \Delta age2 + age-last + age-last2 + terminal$	10.0	-3,180.9	11.6	10.0	-5,596.9	6.7

Table 2. Parameter estimates of linear mixed-effect models including individual body mass beyond 18 years of age as the response variable (in kg, log-transformed) for male and female Asian elephants. V: variance, SD: standard-deviation, SE: standarderror. Marginal and conditional R<sup>2</sup> give the variance explained by fixed effects, and both fixed and random effects, respectively.

	Males		Females					
Random effects	V	SD	Random effects	V	SD			
Individual identity	0.019	0.137	Individual identity	0.011	0.103			
Township	0.0005	0.023	Township	0.003	0.056			
Fixed effects	Estimate	SE	Fixed effects	Estimate	SE			
Intercept	7.707	0.021	Intercept	7.840	0.019			
Age at last measurement	0.087	0.012	Age at last measurement	0.025	0.007			
∆age1	0.041	0.002	Age at last measurement <sup>2</sup>	-0.028	0.006			
∆age2	-0.057	0.014	Δage	0.016	0.001			
			Terminal (1)	-0.071	0.020			
Marginal R <sup>2</sup>	0.24		Marginal R <sup>2</sup>	0.10				
Conditional R <sup>2</sup>	0.89		Conditional R <sup>2</sup>	0.77				

302

### 303 Discussion

We tested whether in a species with marked male-biased intrasexual competition, 304 males showed an earlier and/or faster rate of body mass ageing than females. Both 305 306 sexes gained mass during early adulthood. However, the onset of body mass ageing 307 differed between both sexes: males began to lose mass from 48.3 years old 308 onwards. In contrast, females lost body mass generally at an older age, namely in their last year of life. Compared to a previous study on growth curves of this 309 310 population (Mumby, Chapman, et al., 2015), we are now using a larger dataset, including older and more numerous retired individuals. This allowed us to evidence 311 312 body mass ageing in this species, which was not possible until now. Here we discuss

the implications of our results in the light of the classical theory of ageing (Williams,

1957) and of the management of Asian elephants.

315 We describe for the first time a sex-specific pattern of body mass ageing in this 316 species. Body mass ageing is often used in mammals as it may underpin actuarial and reproductive ageing (Beirne et al., 2015; Bérubé et al., 1999; Nussey et al., 317 318 2011). In our population, sex-specific mortality ageing has been already shown 319 (Lahdenperä et al., 2018) and males display higher mortality than females at all ages. 320 However, reproductive ageing has only been described in females of this population 321 due in part to the difficulty of recording paternity in male elephants (Hayward et al., 322 2014; Mumby, Mar, Hayward, et al., 2015; Robinson et al., 2012). Our results provide 323 valuable insights on how body condition declines with age and offer another aspect of the multifaceted ageing, often referred to as a mosaic (Walker & Herndon, 2010), 324 325 in this long-lived and highly social species.

326 Asian elephants show male-biased intra-sexual competition, with males being more 327 aggressive (Seltmann et al., 2019), fighting more for dominance and showing higher 328 rates of mortality at all ages than females, including during early development, as calves and during adulthood (Lahdenperä et al., 2018). In such species, classical 329 theory of ageing predicts that males should show an earlier onset or accelerated 330 331 ageing (Williams, 1957). Indeed, in several polygynous mammals, males display 332 earlier onset or higher rates of ageing than females, suggested to be due to their 333 stronger intrasexual competition (Clutton-Brock & Isvaran, 2007 but see also Camus 334 et al., 2012; Tower, 2006). For example, in European badgers (Meles meles, Beirne 335 et al., 2015) and Soay sheep (Ovis aries, Hayward et al., 2015), males systematically showed stronger or earlier body mass ageing compared to females. Conversely, in 336 monogamous species, males and females' onsets and rates of ageing tend to be 337

similar (Bronikowski et al., 2011; Clutton-Brock & Isvaran, 2007; Thorley et al., 2020).
Our results are consistent with those studies and with the prediction of the classical
theory of ageing. Moreover, our results are inconsistent with later alternatives that
suggest that the prediction of the classical theory of ageing can be disrupted by high
early-life condition-dependent mortality in males (Chen & Maklakov, 2014) or by
canalisation (Flatt, 2005).

344 However, as mentioned above, previous work on this population showed both reproductive and survival age-related decline in females (Hayward et al., 2014; 345 346 Robinson et al., 2012). Hence, the ageing trajectories do not synchronise between 347 traits in females. Empirically, this heterogeneity of ageing patterns is more the rule 348 than the exception as found in other species (Briga & Verhulst, 2021; Hayward et al., 2015; Walker & Herndon, 2010). In our population, this mismatch can be explained 349 350 by the fact that body mass is a poor predictor of reproductive success, number of 351 offspring produced or raised up to independence (5 years old), and that no 352 relationship between height and survival has been found in females (Crawley et al., 2017). Our results that females do not show age-dependent body mass decline 353 354 combined with previous results are at odds with study on other vertebrates. Asian 355 elephants reproduce all year long (Brown, 2014), despite living in a seasonal 356 environment, meaning that females finance reproduction through energy stored 357 before reproduction. This is contrary to, say roe deer (Capreolus capreolus), an 358 income breeder financing reproduction concurrently to gestation as this ungulate 359 does not store reserves (Andersen et al., 2000). In the latter case, reproductive 360 success therefore depends on body condition and available resources, while in the 361 former case, elephants reproduce when they have stored sufficient resources to finance gestation. This might in part explain the absence of relation between female 362

body mass ageing and reproductive senescence, contrary to males, benefiting more
 from being heavy than females during intrasexual competition (Sukumar, 2003).

365 Our study is subject to a number of limitations when it comes down to identifying why 366 the sexes may differ in their ageing trajectories. First, it is possible that male elephants in our timber elephant population are used more for tasks requiring 367 368 strength or tusks, thereby causing an earlier onset of body mass declines in males 369 than in females. However, both sexes fall under the same government-set workload, 370 care and retirement regulation, except for females' maternity leave. One substantial 371 difference between sexes is that parental care is concentrated on females, with for 372 example only females being given 'parental leave' following reproduction (Toke Gale, 373 1974). However, since maternity is energetically expensive and no more favourable than timber working, this is unlikely to lead to the delayed onset of body mass 374 375 declines in females. An ideal test would be to analyse the effect of timber work and 376 maternity leave on body mass dynamics.

377 Second, elephants have a specific dentition that consists of molar teeth that 378 eventually wear down at the end of their lives, and pathologic malocclusions or lack 379 of molars can lead to weight loss and death by starvation. In sexually dimorphic species of ungulates, males generally display smaller molar teeth size compared to 380 381 females, relatively to body size. This results in teeth wearing down faster and 382 depleting earlier for males than females (Carranza & Pérez Barbería, 2007) with 383 potential consequences on male senescence compared to females. In Asian 384 elephants, although both sexes have the same molar dental anatomy, it is possible 385 that the earlier onset of body mass declines in males reflects sex-specific differences 386 in tooth wear. Indeed, in captive species, dental problems have been described well

before the last year of life (Gaillard et al., 2015) and, if there is sex-specific tooth
wear, this could be associated with the earlier onset of body mass ageing in males.

Third, male elephants have recurring periods of physiological "musth" throughout their adult lives, which can temporarily but profoundly impact the body mass of individual males (Eisenberg et al., 1971) thereby affecting the body mass ageing trajectory. Unfortunately, recording morphological measurements is difficult during the musth period during which males display highly aggressive behaviours, although accounting for musth would improve future analyses.

395 Fourth, in our study, we did not find any evidence for body mass-based selective 396 disappearance, but, as it is often the case in long-lived species, the average 397 longitudinal individual monitoring is short relative to the lifespan of this species (e.g. Global BMI Mortality Collaboration et al., 2016; Prospective Studies Collaboration, 398 399 2009), and hence we only have limited power to detect such association. It is 400 possible that there are sex-specific dynamics of selective disappearance, but whether 401 that is the case in Asian elephants remains to be shown. An analysis with more 402 longitudinal data would be useful to tackle this question.

403 Fifth, for both sexes, it is possible that maximum body mass is set by physiological 404 and ecological constraints as indicated by the weight growth curves found earlier in 405 this population (Mumby, Chapman, et al., 2015). These constraints could be to some 406 extent sex-specific, although at this point, we can only speculate as to why these 407 constraints may drive sex-specific ageing trajectory. Finally, we found a maximum 408 body mass in males but not in females. This sex-specific differences could be driven 409 by the fact that male elephants benefit more than females from being heavy during 410 intrasexual competition (Sukumar, 2003).

Another factor to take into account is that retirement occurs at around 50 years in 411 both sexes, which likely diminishes physical exercise and allows more time for 412 413 foraging, thereby continuing the weight gain. The reduced intrasexual competition in 414 females relative to males, together with this retirement, could lead to the continued 415 mass gain of females. One way of disentangling the effect of senescence and 416 retirement on body mass trajectories would be to know whether muscle is lost over 417 fat. Unfortunately, we do not have the data to know this. However, it seems that muscle function does not decrease with age in this semi-captive population in neither 418 419 sex (Reichert et al., 2022). On the contrary, fat storage as measured by levels of 420 circulating triglycerides remained constant up to adulthood, decreasing afterwards in 421 senior elephants (from retirement onwards) in both sexes (Reichert et al., 2022). 422 Also, all elephants officially retire at age 55, but most elephants enter pre-retirement 423 around the age of 50 because of decreased strength. These results, taken together 424 with the onset of body mass decline we found in males (*i.e.* 48 years old), suggest 425 that retired individuals lose fat in both sex rather than muscle and that body mass ageing is rather a cause than a consequence of retirement in males. Nevertheless, 426 427 given that elephants in the wild do not experience timber labour and retirement, we cannot exclude that the sex-specific body mass ageing trajectories could be different 428 429 in a wild (non-working) population of Asian elephants compared to those found in our 430 study.

We found that females experienced a terminal body mass decline in the last year of life. Our data contain both males and females among the oldest ages (>50), hence sex-specific terminal decline is unlikely to emerge from differences in lifespan. In European badgers, a species in which females outlive males, both sexes displayed terminal body mass declines (Beirne et al., 2015). It is possible that the sex-specific

terminal declines in our study resulted from differences in power, with 5 dead males 436 437 and 18 dead females. Indeed, for both males and females, the coefficient and effect size of the terminal terms were negative, but the effect size in males remained about 438 half of that in females (Cohen's  $d_{mates} = -0.045 [95\%Cl: -0.10; 0.01] = a$  decline of 96 kg 439 440 [95%CI: -19; 205], Cohen's d<sub>femates</sub> = -0.071 [95%CI: -0.11; -0.03] = a decline of 173 kg 441 [95%CI: 80; 263]). Hence, it is possible that the sex-specific terminal effect is driven 442 by power issues and we look forward to testing that with several more years of 443 monitoring.

444 Terminal declines emphasise that the chronological age is rarely a perfect estimation 445 of the biological age which can better describe the 'true biological state' of an 446 organism (Klemera & Doubal, 2006). In that sense, terminal decline is a biomarker of health and remaining lifespan. The 'terminal illness' hypothesis refers to the age-447 448 independent decrease of a trait value, related to the imminent death of the individual 449 (Coulson & Fairweather, 2001). Such terminal effects were shown for example for 450 body mass in mammals (stronger in males than females in European badgers (Beirne et al., 2015), in both sexes in Soay sheep (Hayward et al., 2015) and in male 451 452 but not female Alpine marmots (Tafani et al., 2013)) and for sexual signals in birds (Simons et al., 2016). For which traits or under which conditions to expect terminal 453 454 declines remains yet poorly understood but our study highlights the importance of studying sex-specific differences in ageing and illustrates the need to improve our 455 456 understanding of the mechanisms driving the diversity of ageing patterns in the wild.

#### 457 Acknowledgements

We thank the Myanma Timber Enterprise (MTE) and the Myanmar Ministry of Natural Resources and Environmental Conservation for their collaboration and all of the MTE staff enabling this study by recording elephant life events and morphology for so long. Particularly, we thank Mu Mu Thein and Nina Aro for their patience in helping

translating countless elephant morphological measurements from burmese to
english. We also thank Carly Lynsdale for ensuring our english was fluent. We thank
the Academy of Finland, the European Research Council and the Ella & Georg
Ehrnrooth Foundation for fundings.

### 466 Ethics

The study was performed with the permission from the Myanmar Ministry of Natural Resources and Environmental Conservation and following the University of Turku

469 ethical guidelines.

## 470 **Conflict of interest**

The authors have no conflict of interest to declare.

## 472 **Funding**

- This work was supported by the Academy of Finland [292368], the European
- 474 Research Council [ERC-2014-CoG 648766] and the Ella & Georg Ehrnrooth
- 475 Foundation.

### 476 **Authors' contribution**

HHA, WH and UKN performed field work and data collection. VL, VB and MB
designed the study and LL selected, extracted and translated data. LL carried out all
statistical analyses with contributions from VB and MB. LL wrote the manuscript and
all authors critically reviewed it. All authors approved the manuscript for publication
and agree to be held accountable for the content therein and approve the final
version of the manuscript.

### 483 Data accessibility

The datasets and codes supporting the conclusions of this article have been uploaded on Dryad digital repository and will be made publicly available upon acceptance.

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