- 1 Citizen science data supports sexual dichromatism but rejects thermal melanism
- 2 in the European fire salamander (*Salamandra salamandra*)
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36 Abstract

37 Background

Conspicuous color patterns are traditionally believed to advertise the toxicity of prey to potential predators. However, many aposematic species show drastic variation in coloration, indicating the possibility of other functions of coloration such as a role in thermoregulation (e.g., thermal melanism). To study these other functions, we can investigate the influence of intrinsic (e.g., sex) and external factors (e.g., climate) on color variation. We used the aposematic European fire salamander (*Salamandra salamandra*) to study drivers of variation in the yellow proportion of their dorsal black and yellow coloration based on citizen science data available in a new online database.

## 45 Results

Our results suggest a widespread sexual dichromatism in fire salamanders in Germany with males
displaying a larger proportion of yellow than females. This dichromatism persisted even after
correcting for a sex-difference in body shape that accounts for some of the variation in the
proportion of yellow. Among six investigated putative drivers of this color variation, the proportion
of the aposematic yellow coloration increased with latitude and longitude but showed no association
with other environmental variables such as temperature or the productivity of the habitat.

52 Conclusions

Integrating citizen science data in this study enabled a comparison of fire salamander populations across large parts of their distribution and conclusively supports sexual dichromatism. Given the lack of assocations with temperature-related variables, thermal melanism appears to play no role in fire salamander coloration. Future studies should further investigate potential mechanisms of mate choice in fire salamanders and other selective factors on coloration such as differential predation pressure between both sexes or the role in crypsis. Lastly, multi-national studies supported by citizen science data should fully unravel the extent of color variation in this species.

#### 60 Keywords:

61 Aposematism, Amphibians, Coloration, Image analysis

# 62 Background

63 Color is among the most diverse traits in the animal kingdom. Different abiotic and biotic factors affect the coloration of an animal, both, via evolutionary changes and over the lifespan of an 64 65 individual [1–4]. For centuries, coloration has attracted ecologists and evolutionary biologists aiming 66 to study the drivers of color differences [1, 2]. However, since studying environmental drivers of 67 animal coloration in the wild would involve sampling across large spatial scales in order to capture 68 gradual changes of environmental variables, studies using a larger distribution area and larger 69 gradients of environmental factors are scarce. Harnessing the data available in social media or on 70 citizen science platforms can drastically increase our sampling effort and thereby, improve the 71 generalizability of the conclusions from our studies [5–7].

72 One important selective agent of coloration is predation [8]. Both predator and prey can avoid being 73 seen through background matching or disruptive coloration [9]. However, in other cases, animals 74 feature highly conspicuous colors and patterns. If this form of coloration is paired with an underlying 75 defense such as toxicity or spines, this antipredator strategy is termed "aposematism" [10, 11] and 76 can be very effective at preventing predator attacks once the association between the conspicuous 77 coloration and an underlying defense has been established in the predator [12]. In addition to 78 avoiding predation, animal coloration can play an important role in thermoregulation. By increasing 79 the density of melanin, the skin of an animal becomes darker, increasing its ability to absorb 80 radiation [13]. This thermoregulative strategy termed "thermal melanism" can be adaptive for 81 ectotherms inhabiting cooler habitats by aiding in body temperature regulation [14]. Indeed, a cline 82 in melanism along gradients of temperature or it's proxies for (e.g., geographic latitude or elevation) 83 has already been reported in several taxa [7, 15–17].

84 In addition to natural selection, sexual selection can also play an important role in the evolution of 85 animal color [18–20]. For example, a costly coloration can be selected through mate choice [18, 20, 86 21]. Costly coloration involves energetically costly pigments or highly conspicuous color patterns that 87 increase the likelihood of detection by predators, thus constituting an honest signal of mate quality 88 [20]. Furthermore, if coloration is heritable, mating with a partner that displays costly and therefore 89 attractive coloration can result in the production of attractive offspring. Therefore, highly contrasting 90 and elaborate color patterns can evolve in one or both sexes, depending on the mating system (i.e., if 91 only one of the sexes (e.g., females) is choosy, contrasting and elaborate color patterns should evolve 92 in the opposite sex (e.g., males); [22]). In aposematic species, in which coloration is traditionally 93 believed to be linked to predator avoidance only (see above), it has been hypothesized that sexual 94 selection via mate choice could play an important role in maintaining or even enhancing aposematic 95 coloration [4, 23]. Once the link between conspicuousness and prey unprofitability has been 96 established in the predator, the strength of selection by the predator on coloration could weaken. 97 Consequently, sexual selection could take over the role of the main selective agent and induce a 98 sexual dichromatism in the intensity or pattern of the aposematic coloration. In fact, evidence for the 99 role of sexual selection in shaping aposematic coloration is accumulating [4, 22–25]. In aposematic 100 amphibians in particular, it has been hypothesized that conspicuousness can act as a quality signal 101 and can thus be selected by the choosy sex [26–30].

102 The coloration of an animal can also change within the lifespan of an individual through phenotypic 103 plasticity, either during development [3, 28] or, in some species, within seconds or minutes as a fast 104 response to environmental stimuli [31]. Common drivers of color plasticity are changes in the 105 environment, such as seasonal changes that challenge individuals to exhibit drastic changes in color 106 to match their phenotype to the environment [32–34]. In times of rapid environmental changes 107 caused by human activity, the extent to which plasticity in coloration can enable organisms to adapt 108 to changes in the environment and whether plasticity can buffer a phenotype-environment-109 mismatch are open questions [35–37]. For example, given the importance of color for

thermoregulation (see above), the question of whether animal coloration changes with increasing
temperature due to plastic changes or adaptive evolution is now motivating studies across many
different taxa and geographical areas [38, 39].

113 The European fire salamander (Salamandra salamandra, hereafter referred to as "fire salamander") 114 represents an excellent study organism for drivers of color variation, as postmetamorphic individuals 115 have a conspicuous yellow-on-black coloration that is easily quantifiable due to the strong contrast 116 between the two colors [40]. While the coloration of fire salamanders is considered aposematic given 117 the potent toxic substances produced in specialized glands on the skin [41, 42], surprisingly high 118 inter- and intrapopulation variation exists in the dorsal proportion of yellow in these salamanders 119 [29, 30, 43–47]. There is strong evidence for the heritability of color-associated traits [28, 48], and 120 yellow coloration is energetically costly for an individual [41, 47]; however, the proportion of yellow 121 color does not correlate with the potency of the toxins produced by an individual [28, 29, 45, 49]. 122 Therefore, further studies concerning the emergence and maintenance of color variation in fire 123 salamanders are needed to unravel the causes of the remarkable variation in this species.

124 Previous studies have demonstrated that conditions during the larval period strongly influence 125 postmetamorphic coloration [28, 41, 47]. For example, [41] reported that fire salamanders raised 126 under poor nutritional conditions as larvae had lower proportions of yellow on their back after 127 metamorphosis compared to individuals raised under rich nutritional conditions during the larval 128 stage, indicating that yellow coloration is energetically costly to produce. Another interesting 129 hypothesis, referred to as "thermal melanism", has been postulated, which suggests that the 130 proportion of yellow color could also be mediated by environmental temperatures and, accordingly, 131 by elevation above sea level or distance from the equator (i.e., geographic latitude), resulting in more 132 black and less yellow color in cooler environments [43, 49, 50]. However, to our knowledge, no study 133 has found empirical evidence supporting the existence of "thermal melanism" in fire salamanders 134 [45, 47]. Instead, [47] reported an association between habitat productivity quantified as the 135 normalized difference vegetation index (NDVI; an indicator of photosynthetic activity and green

biomass) and the hue in the yellow color of fire salamanders but not the proportion of dorsal yellow
color. The authors argued that this relationship could indicate that greater resource availability could
facilitate the production of the costly pigments (pteridines and carotenoids stored mainly in
xanthophores; [45]) responsible for the hue of the yellow color of the skin of a fire salamander.

140 Two studies of fire salamanders in six populations in Slovakia and three populations in Germany 141 reported a sexual dichromatism with males having a greater dorsal proportion of yellow than females 142 [29, 30], but no such pattern was found in 25 Italian populations [47] (see also [44, 45], who did not 143 find sex-specific differences in the dorsal color patterns of Polish and Spanish fire salamanders). 144 However, to date, no study has considered the confounding effect of body shape, which could also 145 drive sexual differences in the dorsal yellow proportion of fire salamanders. Fire salamanders in 146 Central Europe are usually mostly black on their lateral sides [42, 43], and an individual with a wider 147 body shape (e.g., due to pregnancy) will therefore have a larger proportion of black in dorsal view, as 148 the abdomen is more stretched out and a larger proportion of the sides is visible (Fig. 1).

149 In this preregistered study (link to preregistration: https://osf.io/92x8w), we used images of fire 150 salamanders from the recently developed web-based software Amphibian and Reptile Wildbook 151 (ARW; [51]), which applies AI-based capture-recapture analysis via pattern recognition. In the ARW, 152 scientists and citizen scientists can upload photos of fire salamanders for population analyses (e.g., 153 [52, 53]). Using images of fire salamanders uploaded to the ARW, we studied variation in the fire 154 salamander dorsal black-to-yellow ratio (B/Y ratio; [28]; low values of the B/Y ratio denote large 155 proportions of yellow) on a large regional scale across wide swaths of the natural distribution of fire 156 salamanders in Germany (Fig. 2). Using 30 pictures (depicting 15 males and 15 females) per location at 20 different locations, we tested for sex-specific differences in the B/Y ratio on the entire body 157 158 viewed from above as well as on the head. We used coloration on the head specifically, as we expect 159 that entire-body coloration and head coloration will be strongly correlated, but that head coloration 160 is not influenced by sex- and season-specific differences in body shape that could confound our 161 results regarding a potential sexual dichromatism (see above and Fig. 1). Furthermore, we tested for

162 correlations between the B/Y ratio and between-site variation in six environmental variables that 163 have been shown to drive animal coloration in previous studies (e.g., [7, 15, 17, 47]). These variables 164 included elevation, average temperature within the period of adult fire salamander peak activity, 165 temperature change over the last 50 years, NDVI, and geographic latitude and longitude. We studied 166 both average temperature and temperature change, as the former represents the current thermic 167 conditions under which the individual developed and thus would affect coloration via developmental 168 plasticity [3], whereas the latter represents how the thermic conditions changed in recent times and 169 thus could represent an important selective agent leading to adaptive evolution [36, 39]. 170 We expected 1) male fire salamanders to show lower B/Y ratios (i.e., more yellow coloration) both on 171 their entire back (body B/Y ratio) and on their head (head B/Y ratio) than females. We also expected 172 2) no sex-difference in the relationship between body B/Y ratio and head B/Y ratio, indicating the 173 absence of a body shape effect on these ratios. Based on the results from testing hypothesis 1) and 174 2), we planned to select either the body B/Y ratio or the head B/Y ratio to study the relationship 175 between fire salamander coloration and environmental location-specific factors.

# 176 Methods

177 All methods were carried out in accordance with our preregistration (link to preregistration:

178 <u>https://osf.io/92x8w</u>), complemented with additional statistical tests as justified in the Statistical

analysis section for further details.

#### 180 Sampling and processing of images

181 Using the *Mapped Results* function in the ARW, we mapped fire salamander encounters color coded

according to sex. Next, we identified locations with enough fire salamander encounters (i.e., a

- 183 minimum of 15 males and 15 females). There had to be a distance of at least 5 km between locations
- 184 [47] to ensure no regular gene flow between populations [54, 55]. If a suitable fire salamander
- 185 population within a location was identified, we requested for permission to include the fire
- 186 salamander images provided by the ARW-users in our analysis and active participation in this study. If

187 the users accepted, for each population, we randomly selected and downloaded 15 images of male 188 and female fire salamanders, respectively, assuring that each fulfilled minimum quality criteria [40]. 189 In brief, acceptable images had a sufficient resolution to delineate color patterns and body parts, 190 showed the entire dorsal side of the individual, and were taken as top-view with an approximately 191 90° angle to the fire salamander. Images failing at least one of these criteria were discarded, and 192 replaced by another randomly selected image from the map. Applying this process, 598 images from 193 20 locations (15 images per sex per location; for one location, we were only able to obtain 13 images 194 of females; Table 1, Fig. 2) were downloaded. All photos included in this study were taken between 195 2017 and 2023 and depicted adult fire salamanders (i.e., snout-to-tail-length > 13 cm, as determined 196 by the person taking the photo).

197 **Table 1** Overview of the study populations with the corresponding number of photos of females  $(n_{\varphi})$ 198 and males ( $n_{\sigma}$ ) included in this study as well as the mean value of the body black-to-yellow ratio ( $\bar{\mathbf{x}}_{body}$ 199 <sub>BY ratio</sub>) and the head black-to-yellow ratio ( $\overline{x}_{head BY ratio}$ ) by population (differences between the sexes in 200 the ratios were independent of population; see Statistical analysis and Results). GPS coordinates, 201 elevation (in m above sea level), average temperature (in °C), temperature change (in °C) and the 202 Normalized Differential Vegetation Index (NDVI) for each location are given as well. See 203 Environmental data collection in the Methods section for further information on how these data 204 were obtained.

Population	nç	n <sub>đ</sub>	₹body BY ratio	Xhead BY ratio	Latitud e	Longitu de	Elevati on	Aevera ge Temper ature	Temper ature Change	NDVI
Altenau	15	15	2.18	1.25	51.806	10.438	446.1	11.004	1.161	0.710
Bad_Harzburg	15	15	1.52	0.79	51.868	10.569	349.5	11.989	1.385	0.710
BGBI	15	15	1.64	0.95	52.014	8.510	173.9	13.122	0.986	0.635
EBWOB	15	15	1.45	0.73	52.380	10.819	116	13.311	1.299	0.633
Ehrenberg	15	15	1.04	0.62	51.039	13.016	225.3	13.092	1.311	0.598
Floeha	15	15	1.83	1.05	50.863	13.081	323.7	12.841	1.387	0.629
FRD	15	15	1.42	0.85	51.918	8.876	203.1	13.011	0.999	0.634
Gartensalamander	13	15	1.34	0.63	50.173	7.621	384	13.289	1.160	0.695
Heidelberg	15	15	2.12	1.20	49.362	8.705	259	14.546	1.302	0.644
Ilsenburg	15	15	1.29	0.72	51.851	10.669	306.4	11.128	1.234	0.679
Kaesenbachtal	15	15	2.32	1.44	48.534	9.041	408.5	13.154	1.171	0.684
KF	15	15	3.33	1.25	50.667	7.083	174.8	13.979	1.176	0.594
KO-Bienhorntal	15	15	2.00	1.14	50.344	7.610	129.3	13.813	1.090	0.642
Kruppwald	15	15	2.52	1.30	51.403	6.995	99.5	14.033	0.993	0.591
Muelheimer_Wald _Rottbachtal	15	15	1.93	1.07	51.380	6.842	64.6	13.337	0.885	0.584
Ratinger_Wald	15	15	1.97	1.04	51.312	6.866	75.6	14.110	0.927	0.613
Remscheid_Eschba chtal	15	15	2.19	1.31	51.152	7.170	157.4	13.004	1.301	0.727
Wasserburgertal	15	15	1.77	1.29	47.895	8.852	639.9	12.245	1.207	0.698
Wernigerode	15	15	1.19	0.51	51.821	10.742	259.9	12.609	1.320	0.679
Wuppertal_Kothen er_Wald	15	15	2.05	0.86	51.255	7.190	259.7	13.053	1.237	0.687

<sup>205</sup> 

206 Following the instructions provided by [40], the fire salamander body, excluding the legs, was

207 cropped out and saved as a new image. Additionally, the head was cropped out and saved as a

separate image to compare overall body coloration with head coloration (Fig. 3). The B/Y ratio was

- then automatically quantified via the Python script provided in [40]. All image processing and
- analyses were conducted by two researchers (RH and MS) who were blinded to the sex and location

211 of the fire salamander in each picture.

#### 212 Environmental data collection

- 213 For each population, a centroid was determined as the approximate center of the fire salamander
- encounters. Using the GPS coordinates of the centroid for each population, the elevation above sea
- 215 level (in m) was obtained via the free web tool "Elevation Finder"
- 216 (https://www.freemaptools.com/elevation-finder.htm). The average temperature (in °C) for each
- 217 location was calculated on the basis of the centroids via a fine-scale (0.25° × 0.25°) temperature 218 dataset of daily temperatures dating back to 1950 [56]. We only considered temperatures during the 219 period of fire salamander activity between the beginning of March and the end of October, as fire 220 salamanders in Germany hibernate during the winter in underground shelters that are likely to 221 constitute constant-temperature environments [42]. Furthermore, the average temperature was 222 calculated from 2000 to 2015, as this timeframe likely covers the time the individuals were born and 223 matured (an average age of 7 to 9 years was reported for fire salamanders in different countries in 224 Europe; however, a maximum age of 25 years has also been reported in the literature [42, 57]). Thus, 225 this value represents the average temperature to which the fire salamanders were exposed during 226 ontogeny and adult life. Similarly, the average temperature between 1950 and 1965 was calculated 227 from the same dataset [56] and then subtracted from the average temperature from 2000 to 2015 to 228 approximate how much a specific location changed in average temperature over those 50 years. 229 While all locations experienced an increase in temperature (Table 1), between-site variation in this 230 temperature change could be an important driver of evolutionary change in color through it's effect 231 on previous generations of fire salamanders. NDVI data were obtained via the fine-scale  $(0.05^{\circ} \times$ 232 0.05°) MODIS vegetation index dataset (https://www.cen.uni-hamburg.de/en/icdc/data/land/modis-233 vegetationindex.html). The NDVI per location was also averaged for March to October in the period of 2000 to 2015. 234

235 Statistical analysis

All statistical analyses were carried out in R version 4.2.2 [58]. Prior to analyses, we checked our data
for unexplainable outliers and ensured that the statistical models did not include any strong
collinearity between predictor variables following the protocol outlined in [59]. We did not find any
unexplainable outliers and proceeded as follows. All figures presented in the results section were
created using the R package ggplot2 [60].

241 We used linear mixed effects models (LMMs) to study variation in body B/Y ratio and head B/Y ratio. 242 All LMMs were fitted via the function *lmer* from the lme4 package [61]. We log<sub>10</sub>-transformed the 243 body B/Y ratio and head B/Y ratio for all models to ensure that the assumptions of linearity, 244 homoscedasticity and normality of our models were met, as confirmed via the function check\_model 245 in the performance package [62]. To test the significance of the fixed effects in our models, we used 246 the summary function provided by the package ImerTest [63]. To find the most suitable random 247 effect structure for each model, we used the function ranova provided by ImerTest [63]. If our initial more complex random effect structure (i.e., a random slope for sex and random intercept for 248 249 population) was not a significantly better fit to the data (based on a p-value > 0.1) than a simpler 250 random effect structure (i.e., just a random intercept), we chose the simpler random effect structure 251 for the final model. For each model, we present the initial and the final random effect structure 252 below but in the results section we present only the results from the final model. To determine the 253 role of sex-specific differences in body shape on the black-and-yellow coloration of fire salamanders, 254 we used the body B/Y ratio and the head B/Y ratio in separate LMMs as dependent variables. As a 255 fixed effect, the models included sex (categorical: female or male). As random effects in both models, 256 we included a random intercept of *population* to reflect the paired nature of samples from the same 257 population and a random slope for sex to control for random variation in sexual color differences 258 between populations [64]. For the model on body B/Y ratio, the random intercept and slope 259 structure provided a marginally better fit (AICrandom intercept and slope model = -93.219) than the random 260 intercept model (AIC<sub>random intercept model</sub> = -91.922; p = 0.071, so we kept the former but we removed the

261 random slope for the model on head B/Y ratio (AICrandom intercept and slope model = 45.511; AICrandom intercept 262  $_{model}$  = 42.512; p = 0.606). We also studied the correlation between the body B/Y ratio and head B/Y 263 ratio (both log<sub>10</sub>-transformed) to investigate an effect of body shape (see above) by using Pearson's 264 correlation test in the package stats [58]. Finally, we studied sex-differences in the relationships 265 between the body B/Y ratio and head B/Y ratio by using a LMM. The LMM included the log<sub>10</sub>-266 transformed head B/Y ratio as dependent variable, as well as the log<sub>10</sub>-transformed body B/Y ratio, 267 sex and their interaction as fixed effects. The log<sub>10</sub>-body B/Y ratio was centered around it's mean to 268 improve the interpretability of the intercept [65]. Any sex-differences in slopes would indicate an 269 effect of sex-specific differences in body shape on the body B/Y ratio and a significant intercept 270 would additionally indicate differences in the mean head B/Y ratio compared to the mean body B/Y 271 ratio. Initially, this model included a random intercept for population, a random slope of sex by 272 population, as well as a random slope of the body B/Y ratio by population to control for population-273 specific differences in the head B/Y ratio as well as population specific differences in the sexual 274 dichromatism of the head B/Y ratio and population specific differences in the relationship of the 275 body B/Y ratio with head B/Y ratio. The model including a random intercept of population and a 276 random slope for the body B/Y ratio for each population as well as the initial model provided the best 277 fit (all p < 0.004) but the two models were not significantly better when compared with each other 278 (AICinitial model = -591.19; AICrandom intercept of population and random slope of body B/Y ratio model = -592.20; p = 0.322). 279 Therefore, we picked the simpler model (i.e., the model including a random intercept of population 280 and a random slope of the body B/Y ratio) for the final analysis.

Using the model described above, we found evidence for sex-specific differences in the relationship between the head B/Y ratio and the body B/Y ratio that are likely a consequence of sex-specific differences in body shape (for more details see the results). Therefore, in accordance with our preregistration, we used the head B/Y ratio to study the role of environmental variables on fire salamander black-to-yellow ratio. We present corresponding results from the models using the body B/Y ratio in the supplementary information. We used separate models for each environmental 287 variable (i.e., elevation, average temperature, temperature change, NDVI, latitude, longitude) with 288 the mean-centered (see above) environmental variable, sex and their interaction as fixed effects. We 289 also divided elevation by 100 in order to have all environmental variables on similar scales. As 290 random effects, we included a random intercept for population as well as a random slope of sex by 291 population. Following the same step-wise reduction of the random effect structure as described 292 above, we removed the random slope in every model as it did not improve the model slope (all p >293 0.477). Our preregistration did not include geographic latitude and longitude as potential drivers of 294 fire salamander coloration. However, based on earlier reviewer suggestions, we included these 295 variables and therefore, we explicitly explore spatial variation in coloration. Since all the 296 environmental variables showed weak to moderate pairwise correlations (tested using Spearman's 297 correlation tests in ggpairs in the GGally package [66]; all  $\rho < 0.690$ ; see Supplementary Fig. 1), we 298 also used one model that included all centered environmental variables (elevation divided by 100) as 299 well as sex and the interactions of each environmental variable with sex to study the effect of each 300 variable on the head B/Y ratio while controlling for the effect of each other environmental variable 301 [67]. As random effects, this model included a random intercept for population and a random slope 302 of population by sex. The random slope was subsequently removed as it did not improve the model 303 fit (AIC<sub>random intercept and slope model</sub> = 58.791; AIC<sub>random intercept model</sub> = 55.345; *p* = 0.758). We used the 304 function emtrends provided by the package emmeans [68] to determine if the slope for males is 305 significantly different from 0. As the study populations are not evenly distributed across Germany 306 (Fig. 2), we tested for residual spatial autocorrelation of our models using Moran's I provided by the 307 function Moran. I in the ape package [69] but found no evidence for residual spatial autocorrelation 308 (all *p* > 0.670).

309

310 Results

### 311 Sexual dichromatism

312 Male fire salamanders had both, a higher proportion of yellow on their entire body as well as on their

head only (i.e., lower body B/Y ratio and head B/Y ratio) than females (Fig. 4; Table 2). In both sexes

314 combined, the body B/Y ratio and the head B/Y ratio were strongly positively correlated (Corr =

0.792; p < 0.001). However, the slopes differed between the sexes, with males showing a greater

316 increase than females (Fig. 5; Table 3; see Supplementary Fig. 2 for regression plots by sex and

population). Overall, fire salamanders of both sexes had a higher proportion of yellow on their heads

than on their entire body (i.e., lower head B/Y ratio than body B/Y ratio; Table 1; Table 3, see

319 Supplementary Fig. 2 for population differences). For further analyses, we continue with the head

B/Y ratio.

321 Table 2 Summary of the linear mixed effects models used to investigate differences in the body B/Y 322 ratio and the head B/Y ratio of female and male fire salamanders across 20 locations in Germany. The 323 model output is on the log<sub>10</sub> scale, as the dependent variables were log<sub>10</sub>-transformed prior to 324 analysis. The model coefficient estimates ( $\beta$ ) are presented with their standard errors (SE) for the 325 fixed effects. Test statistics (t) and p values are given, and all significant effects (p < 0.05) are 326 presented in bold. Variance estimates ( $\sigma^2$ ) for the random effect and residual variance are given. For 327 categorical predictors, reference levels are presented in parentheses after the model parameter 328 name.

Model	Model Parameter		Model	Output	
	Fixed Effects	в	SE	t	p
Body B/Y ratio	Intercept (Female)	0.26	0.03	9.73	< 0.01
	Sex (Male)	-0.12	0.02	-5.46	< 0.01
	Random Effects	$\sigma^2$			
	Intercept (Population)	0.01			
	Sex (Male)	< 0.01			
	Residual	0.04			
Head B/Y ratio	Fixed Effects	в	SE	t	p

Intercept (Female)	-0.03	0.03	-0.73	0.47
Sex (Male)	-0.11	0.02	-5.43	< 0.01
Random Effects	$\sigma^2$			
Intercept (Population)	0.02			
Residual	0.06			

329

330	Table 3 Summary of the linear mixed effects model investigating differences in the relationship of the
331	head B/Y ratio with the body B/Y ratio by sex. The model output is on the $log_{10}$ scale, as the
332	dependent variable was $\log_{10}$ -transformed prior to analysis. The model coefficient estimates ( $m{ heta}$ ) are
333	presented with their standard errors (SE) for the fixed effects. Please note that $log_{10}$ body B/Y ratio
334	was centered around the mean ( $\overline{x}$ = 0.199). Test statistics (t) and p values are given, and all significant
335	effects ( $p < 0.05$ ) are presented in bold. Variance estimates ( $\sigma^2$ ) for random effects and residual
336	variance are given. For categorical predictors, reference levels are presented in parentheses after the
337	model parameter name.

Model Parameter	Model Output					
Fixed Effects	в	SE	t	р		
Intercept (Female)	-0.07	0.01	-5.11	< 0.01		
Log <sub>10</sub> body B/Y ratio	0.83	0.05	16.95	< 0.01		
Sex (Male)	0.01	0.01	0.84	0.40		
Log <sub>10</sub> body B/Y ratio : Sex (Male)	0.16	0.05	3.13	< 0.01		
Random Effects	$\sigma^2$					
Intercept (Population)	< 0.01					
Log <sub>10</sub> body B/Y ratio	0.02					
Residual	0.02					

338

# 339 Investigating environmental correlates with color variation

We found a negative correlation of the head B/Y ratio with latitude for both sexes (i.e., in the north, fire salamanders are more yellow on their heads; Table 4; Fig. 6A) and a negative correlation of the head B/Y ratio of males with longitude (i.e., in the east, males are more yellow on their heads; Table 4B; Fig. 6B) while this association was non-significant in females (p = 0.078; Table 4A). None of the 344 other environmental variables showed a significant relationship with the head B/Y ratio in single 345 predictor analyses (all p > 0.139; Table 4; Supplementary Fig. 3). All these relationships vanished in 346 the model including all environmental variables simultaneously (all p > 0.200; Table 5A) with the 347 exception of a just non-significant negative correlation of the males' head B//Y ratio with longitude 348 (i.e., in the east, males have more yellow on their heads). While the males' head B/Y ratio 349 relationship with temperature change was significantly different to the relationship of the females' 350 head B/Y ratio with temperature change (Table 5A), the slopes of both, the males' and females' head 351 B/Y ratio with temperature change were not significantly different from zero (Table 5B). In line with 352 the previous analysis, all models investigating the relationships of the environmental variables with 353 the head B/Y ratio indicated that males have a lower head B/Y ratio (i.e., more yellow on their heads) than females (all p < 0.001; Table 4; Table 5A; Fig. 6). 354

355 Table 4A Summary of the linear mixed effects models used to investigate the relationship between 356 each environmental variable with head B/Y ratio. The model output is on the log<sub>10</sub> scale, as head B/Y 357 ratio was log<sub>10</sub>-transformed prior to analysis. Please note that elevation was divided by 100 in order 358 to have it on a similar scale to the other variables. All environmental variables were centered around 359 their means in order to facilitate better model interpretation ( $\bar{X}_{Elevation} = 2.524$ ;  $\bar{X}_{Average Temperature} =$ 360 13.030;  $\overline{x}_{Temperature Change} = 1.177$ ;  $\overline{x}_{NDVI} = 0.653$ ;  $\overline{x}_{Latitude} = 50.950$ ;  $\overline{x}_{Longitude} = 9.040$ ). The model coefficient 361 estimates ( $\beta$ ) are presented with their standard errors (SE) for the fixed effects. Test statistics (t) and 362 p values are given, and all significant effects (p < 0.05) are presented in bold. Variance estimates ( $\sigma^2$ ) 363 for random effects and residual variances are given. For categorical predictors, reference levels are 364 presented in parentheses after the model parameter name. B Estimated slopes for males for each 365 environmental variable.

A) Model Summaries							
Environmental Variable	Model Parameter	Model Output					
	Fixed Effects	в	SE	t	р		
Elevation	Intercept (Female)	-0.03	0.04	-0.71	0.48		
	Sex (Male)	-0.11	0.02	-5.44	< 0.01		

	Elevation	0.02	0.03	0.68	0.50
	Sex (Male) : Elevation	-0.02	0.01	-1.63	0.11
	Random Effects	$\sigma^2$			
	Intercept (Population)	0.02			
	Residual	0.06			
	Fixed Effects	в	SE	t	р
	Intercept (Female)	-0.03	0.03	-0.74	0.47
	Sex (Male)	-0.11	0.02	-5.44	< 0.01
	Average Temperature	0.02	0.04	0.64	0.53
Average Temperature	Sex (Male) : Average Temperature	0.03	0.02	1.61	0.11
	Random Effects	$\sigma^2$			
	Intercept (Population)	0.02			
	Residual	0.06			
	Fixed Effects	в	SE	t	р
	Intercept (Female)	-0.03	0.03	-0.75	0.46
	Sex (Male)	-0.11	0.02	-5.42	< 0.01
Townsonature	Temperature Change	-0.34	0.23	-1.47	0.16
Change	Sex (Male) : Temperature Change	0.07	0.13	0.50	0.62
	Random Effects	$\sigma^2$			
	Intercept (Population)	0.02			
	Residual	0.06			
	Fixed Effects	в	SE	t	р
	Intercept (Female)	-0.03	0.04	-0.72	0.48
	Sex (Male)	-0.11	0.02	-5.43	< 0.01
	NDVI	0.17	0.82	0.21	0.84
NDVI	Sex (Male) : NDVI	-0.61	0.45	-1.37	0.17
	Random Effects	$\sigma^2$			
	Intercept (Population)	0.02			
	Residual	0.06			
	Fixed Effects	в	SE	t	р
	Intercept (Female)	-0.03	0.03	-0.82	0.42
	Sex (Male)	-0.11	0.02	-5.43	< 0.01
	Latitude	-0.06	0.03	-2.28	0.03
Latitude	Sex (Male) : Latitude	-0.00	0.02	-0.20	0.84
	Random Effects	$\sigma^2$			
	Intercept (Population)	0.02			
	Residual	0.06			

	Fixed Effects	в	SE	t	р
	Intercept (Female)	-0.03	0.03	-0.80	0.43
	Sex (Male)	-0.11	0.02	-5.43	< 0.01
	Longitude	-0.03	0.02	-1.85	0.08
Longitude	Sex (Male) : Longitude	-0.01	0.01	-0.75	0.45
	Random Effects	$\sigma^2$			
	Intercept (Population)	0.02			
	Residual	0.06			
B) Estimat	tion of Slopes for Males				
Environmental	Variable		Output		
	в	SE		t	р
Elevation	-0.01	0.03		-0.20	0.85
Average Tempe	erature 0.06	0.04		1.54	0.14
Temperature C	hange -0.28	0.23		-1.18	0.25
NDVI	-0.44	0.82		-0.54	0.60
Latitude	-0.07	0.03		-2.41	0.02
Longitude	-0.04	0.02		-2.30	0.03

366

367 Table 5A Summary of the linear mixed effects model investigating the relationship of head B/Y ratio 368 with each environmental variable simultaneously. The model output is on the log<sub>10</sub> scale, as head B/Y 369 ratio was  $log_{10}$ -transformed prior to analysis. The model coefficient estimates ( $\beta$ ) are presented with 370 their standard errors (SE) for the fixed effects. Please note that elevation was divided by 100 and that 371 all environmental variables were centered around their means in order to facilitate better model 372 interpretation (for means of each variable, see Table 4). Test statistics (t) and p values are given, and 373 all significant effects (p < 0.05) are presented in bold. Variance estimates ( $\sigma^2$ ) for the random effect 374 and the residual variance are given. For categorical predictors, reference levels are presented in 375 parentheses after the model parameter name. B Estimated slopes for males for each environmental 376 variable.

A) Model Summary							
Model Parameter	Model Output						
Fixed Effects	в	SE	t	р			
Intercept (Female)	-0.03	0.03	-0.81	0.43			

Sex (Male)		-0.11	0.02	-5.45	< 0.01
Elevation		-0.02	0.06	-0.35	0.73
Average Temperature		-0.03	0.09	-0.32	0.75
Temperature Change		-0.40	0.42	-0.97	0.35
NDVI		0.59	1.57	0.38	0.71
Latitude		-0.08	0.06	-1.34	0.201
Longitude		0.00	0.04	-0.14	0.89
Sex (Male) : Elevation		-0.05	0.04	-1.35	0.18
Sex (Male) : Average Tem	perature	-0.08	0.05	-1.48	0.14
Sex (Male) : Temperature	0.60	0.25	2.38	0.02	
Sex (Male) : NDVI		1.53	0.96	-1.60	0.11
Sex (Male) : Latitude		-0.05	0.04	-1.25	0.21
Sex (Male) : Longitude		-0.03	0.02	-1.60	0.11
Random Effects		$\sigma^2$			
Intercept (Population)		0.02			
Residual		0.06			
B) Estimation of	Slopes for Mal	es			
Environmental Variable		Output			
	в	SE	-	t	p
Elevation	-0.08	0.0	6	-1.18	0.25
Average Temperature	-0.11	0.0	9	-1.23	0.24
Temperature Change	0.20	0.4	2	0.49	0.63

1.57

0.06

0.04

-0.60

-2.11

-1.12

0.56

0.05

0.28

377

# 378 Discussion

NDVI

Latitude

Longitude

## 379 Sexual dichromatism in fire salamanders

Using a semi-automated analysis of photos of fire salamanders across Germany taken by scientists and citizen scientists, we show that males have a higher proportion of yellow on their entire dorsal surface as well as on their head only compared to females. While body B/Y ratio and head B/Y ratio were strongly positively correlated, this relationship was more positive (i.e., a steeper slope) in males than in females. This indicates that a female with a high body B/Y ratio (i.e., low proportion of yellow on her body) has a lower head B/Y ratio (i.e., higher proportion of yellow on her head) than a male

-0.94

-0.13

-0.04

386 fire salamander with an equal body B/Y ratio (Fig. 5). Most fire salamanders in Germany have only 387 very little yellow coloration on their sides, and females usually have a wider body shape than males 388 (due to pregnancy; [42, 43]). Therefore, the more stretched out bodies of females make the sides 389 appear more visible from above, reducing the proportion of yellow when viewed from above. This 390 sexual dimorphism in body shape likely influenced the sexual dichromatism in body coloration 391 observed in this study and in previous studies [29, 30]. The head coloration is less affected by 392 changes in body shape due to pregnancy or feeding status and therefore, comparing the head 393 coloration between sexes is less biased. Since the yellow proportion on the head is also higher in 394 males than in females, however, the difference in body shape can only partly explain the observed 395 sexual dichromatism. Nevertheless, we advise caution for researchers aiming to study sex-specific 396 differences in dorsal coloration (specifically: of the abdomen) of fire salamanders in the future as 397 differences in body shape will likely need to be accounted for when comparing females with males. 398 Our finding of a sexual dichromatism, with males being more yellow than females, is in line with 399 previous studies in Germany [29] and Slovakia [30]; however, fewer populations were tested (three 400 and six populations, respectively). In contrast, two studies investigating sexual dichromatism in fire 401 salamanders in northern Spain and northern Italy, respectively, did not find differences in coloration 402 between males and females [45, 47]. Therefore, our study, which used data from 20 locations, 403 spanning the largest area of the natural distribution of fire salamanders so far, provides the first 404 conclusive evidence of widespread sexual dichromatism in this species. One potential cause of this 405 sexual dichromatism is sexual selection driven by mate choice, whereby female fire salamanders 406 prefer to mate with yellower males. Fire salamanders are capable of discriminating yellow from other 407 colors even under low light conditions [70]; therefore, color-based mate choice is theoretically 408 possible in this species. When raised under rich food conditions as larvae, postmetamorphic fire 409 salamanders have a higher proportion of yellow than individuals raised under poor food conditions 410 as larvae, indicating that yellow coloration is likely energetically costly for fire salamanders [41]. 411 However, this difference in yellow proportion disappeared with time when the postmetamorphic fire

412 salamanders were maintained under identical nutritional conditions, which could indicate that the 413 maintenance of the pigments responsible for yellow coloration is also costly [41]. Therefore, the 414 proportion of yellow coloration in fire salamanders could act as a signal of a potential mate's quality, 415 especially when considering that more yellow coloration is more effective at deterring predators 416 [41], although importantly, more yellow coloration does not indicate higher levels of toxicity [28, 29, 417 45, 49]. In the spadefoot toad Scaphiopus couchii, larger males and males with a better body 418 condition have a more dissimilar body coloration to females [71]. Furthermore, clay models that 419 mimic these males in coloration were also preferred by females over clay models of males that 420 resembled females in color and dorsal pattern [71]. Similarly, natterjack toads (Epidalea calamita) 421 are sexually dichromatic, and their coloration is correlated with age, body size, hindlimb length, and 422 sprint speed [72]. Although no correlation between color and toxicity has been detected in fire 423 salamanders [28, 29, 45, 49], other factors, such as genetic quality, age or performance capacity, 424 might be identifiable for females via coloration [73]. Male fire salamanders have been reported to 425 prefer sites that are linked to female activity [74] and are frequently observed in an upright position 426 referred to as a "presenter pose" [70, 75]. This pose could provide females with the opportunity to 427 examine a male's coloration and body condition and could therefore play an important role in visual 428 communication during mate choice [75]. Future studies could either directly (e.g., [71]) or indirectly 429 (e.g., [76]) investigate the role of fire salamander coloration in mate choice and whether other traits 430 correlate with a fire salamander's coloration, which would indicate whether color acts as a quality 431 signal. Furthermore, the body parts visible for a female should be further investigated (i.e., "taking the perspective" of a female fire salamander). 432

Alternatively, this sexual dichromatism could be related to differential selection pressure (e.g., by
predators) for both sexes in fire salamanders. As described above, male fire salamanders are often
seen in "presenter pose" with their bodies raised and their forelimbs extended, staying motionless
for up to several minutes [75]. This behavior could not only be related to intraspecific
communication, as the dorsal pattern is more visible for conspecifics when the body of the male is

438 raised, but it could also expose the male to a greater threat of predation (by e.g., Natrix natrix snakes 439 and birds of the order Strigiformes; [42, 47]). Furthermore, males were more active than females in 440 behavioral assays (Mühlenhaupt et al., in preparation), which could be another reason why males are 441 exposed to a greater threat of predation [77]. Similarly, the development of fire salamander larvae in 442 a female's body might necessitate greater metabolic activity and therefore, the need for higher 443 temperatures, which could be supported by a greater proportion of black in females, i.e., "thermal 444 melanism". For example, [78] reported that males of the ground cricket Allonemobius socius had 445 darker cuticles than females did, which could be the result of, e.g., additional sex-specific selection 446 due to temperature-dependent display behavior.

#### 447 Differences in the yellow proportion of body and head

448 An interesting result from our investigations is that fire salamanders have higher proportions of 449 yellow on their head than on their overall dorsal surface. Caspers et al. (2020) used models of fire 450 salamanders with varying proportions of yellow to show that models with higher proportions of 451 yellow received fewer attacks from potential predators [41]. Since the head of a fire salamander is an 452 irreplaceable body part necessary for the survival of an individual, protecting this body part might 453 prove especially important compared with, e.g., the tail (fire salamanders with missing tails are 454 frequently observed (personal observations)). Using coloration to focus or divert predator attacks on 455 specific body parts is a well-known strategy in the animal kingdom [8]. For example, [79] used clay models of lizards with three sections-head, body and tail-that were either black or blue and 456 457 reported that models with blue sections received the majority of attacks by birds on their blue 458 sections rather than on the black sections. This is especially interesting given that lizards of multiple 459 families have evolved blue tails that can be autotomized [80]. Thereby, attacks are guided toward the 460 body part that can be sacrificed (i.e., is nonvital). In fire salamanders, yellow coloration could act in 461 the opposite way and divert predator attacks from the head as the overall proportion of yellow has 462 already been shown to reduce predator attacks on fire salamander models [41]. Furthermore, fire 463 salamanders have prominent toxin glands, called parotoid glands, on the sides of their heads that are

in almost every case at least partly yellow [42, 43], and the yellow coloration highlights the dark
glandular pores. Advertising these toxin glands might be an effective strategy for avoiding predator
attacks as fire salamanders have been reported to engage in a defensive posture by lowering their
heads and exposing their parotoid glands towards a predator [42, 90]. Similarly, in natterjack toads
(*Epidalea calamita*) that also have parotoid glands, plasticine models with larger and more
conspicuously colored parotoid glands received fewer predator attacks [81].

The head coloration might also play an especially important role in mate choice of fire salamanders.
For example, [82] showed that males of the two extant species of *Salamandrina* have more
conspicuous gular coloration. When females are nearby, this coloration is displayed in a stereotyped
throat hyperextension while being in a similar "presenter pose" to fire salamanders. Therefore, the
head coloration of fire salamanders might play an important role in mate choice. Especially the gular
coloration of fire salamanders merits further investigation given that the throat is especially visible
during the "presenter pose" (see also [70]).

## 477 Environmental correlates with fire salamander coloration

478 Several other studies have already attempted to find environmental correlates with the yellow 479 proportion in fire salamander coloration. For example, Burgon et al. (2020) found no relationship 480 between elevation and the frequency of color morphs that vary in yellow proportion and value in S. 481 s. bernadezi in northern Spain. Similarly, Barzaghi et al. (2022) found no correlation between the 482 proportion of yellow with NDVI, elevation or predator richness in fire salamanders of northern Italy 483 that likely represent a hybrid of S. s. salamandra and S. s. gigliolii [43]. In our study, we tested for 484 environmental correlates with yellow proportion in fire salamanders from the widest area of fire 485 salamander distribution studied so far and with individuals belonging to both the S. s. terrestris 486 lineage (in western Germany) and the hybrid lineage of S. s. salamandra and S. s. terrestris (in 487 eastern Germany) [43, 83]. We found evidence for variation in the yellow proportion of fire 488 salamanders along their north-to-south and west-to-east distribution. Given that in our study, fire 489 salamander populations from southwestern Germany to eastern Germany were included, previous

490 studies might have failed to find similar clines due to a smaller sampling area. Interestingly, neither 491 an east-west cline nor a south-north cline can be explained in variation due to underlying 492 environmental factors such as variation in temperature or habitat productivity (i.e., NDVI), as these 493 variables were not significantly associated with the B/Y ratios either when tested individually or in 494 models including all environmental variables. One possible driver of this variation could be 495 differences in yellow proportion between genetic lineages across Germany. For example, it is well 496 known that S. s. salamandra and S. s. terrestris and their hybrids differ in dorsal color pattern [43, 44, 497 83], and it is very likely that the color patterns also affect the yellow proportion (unpublished data). 498 However, these differences between lineages do not ultimately explain the variation in yellow 499 proportion. A potential explanation for this variation could be differences in the mate choice 500 behavior of females between these lineages, as latitude and longitude were more often significantly 501 associated with the B/Y ratios of males than of females, especially when accounting for the effects of 502 other environmental variables (i.e., using the models that included all environmental variables). 503 Clearly, further research spanning larger geographic areas and including more lineages of the widely 504 distributed and highly variable European fire salamander is needed. This research could also indicate 505 whether the sexual dichromatism that we have identified for fire salamanders in Germany is also 506 present in other lineages outside Germany (e.g., in the fire salamander diversity hotspot on the 507 Iberian peninsula [43]). Integrating citizen science approaches and even using data available through 508 apps such as iNaturalist or social media can make such a task feasible [5].

Although the putative role of thermal melanism in fire salamanders has been suggested multiple times [43, 49, 50], neither average temperature nor temperature change showed significant associations with the yellow proportion of the fire salamanders studied here. While comparing the fully black alpine salamander (*Salamandra atra*) with the European fire salamander might lead to this conclusion as alpine salamanders inhabit higher elevated habitats with lower average temperatures [43], the intraspecific variation in yellow proportion of fire salamanders could not be explained through differences or changes in temperature in this or former studies [45, 47]. While we were able 516 to cover a span of 3 °C in average temperature (Table 1) and [45] covered a span in elevation of 78 to 517 1312 m a.s.l., no associations with elevation or average temperature have been found, yet. 518 Therefore, it might be possible that other mechanisms such as behavioral adaptations could play a 519 more important role in thermal regulation of fire salamanders. For example, fire salamanders in 520 Mediterranean areas such as parts of the Iberian peninsula show drastically different activity 521 patterns (i.e., activity during winter) to fire salamanders in temperate climates (i.e., hibernation 522 during winter) [42, 43]. Furthermore, in some regions fire salamanders show diurnal activity whereas 523 in other areas, they are almost exclusively nocturnal ([42, 84], personal observations). Therefore, 524 future studies could also consider differences in activity patterns across large geographic areas. 525 We have included temperature change as there was quite some variation in how much the average 526 temperatures have increased in our study locations and as from other studies it is clear that global 527 warming will have a strong impact on the coloration of ectotherms [36, 38, 39, 85]. However, in our 528 study we did not find evidence for an effect of warming on the yellow proportion of fire salamanders. 529 Again, behavioral adaptations such as shifts in phenology (e.g., increased importance of winter and 530 decreased importance of summer for fire salamanders in Germany) might mitigate the changes in 531 temperature as for other sympatric amphibian species shifts in phenology with milder winters have 532 already been shown [86, 87]. An alternative approach to the one we have used here would be to 533 compare the coloration of fire salamanders, preferably from multiple sites, across time (e.g., by using 534 photos and/or specimen from biological collections) to correlate the change in temperature over a 535 span of several decades with the proportion of yellow.

Further alternative explanations of the variation in yellow proportion should also be considered. In other aposematic amphibians such as poison dart frogs (Dendrobatidae), a role of coloration in camouflage has already been shown [88, 89]. Locality-dependent differences in background could be associated with differences in coloration of fire salamanders in these locations. While the "greenness" of the habitat (i.e., NDVI) did not show an association with the yellow proportion, other features of the habitat could reduce the detectability of specific color morphs. It can be surprisingly easy to miss a fire salamander when actively searching for these aposematic salamanders (personal
observation). A possible role of fire salamander coloration in crypsis has already been proposed [90,
91] but remains to be thoroughly tested.

545 Conclusion

546 This study provides strong evidence for a widespread sexual dichromatism of fire salamanders in 547 Germany through the use of citizen science data. Furthermore, we find evidence for variation in the 548 coloration of the head and the body indicating as of yet underappreciated involvement in different 549 biological functions. Future studies investigating differences in dorsal coloration of fire salamanders 550 should consider differences in body shape between females and males as these will likely affect the 551 dorsal proportion of yellow. The clines in yellow proportion with latitude and longitude that we have 552 found could be the result of differences between lineages of fire salamanders in Germany but the 553 underlying mechanisms are yet to be uncovered. Clearly, more large-scale comparisons, integrating 554 multiple lineages across diverse habitats of fire salamanders, are necessary to better understand the 555 drivers of inter- and intrapopulation as well as intraindividual variation in fire salamander coloration. 556 To facilitate such extensive studies, the implementation of citizen science approaches will be 557 extremely helpful as has been shown for other animals already.

#### 558 Abbreviations

- 559 NDVI Normalized Difference Vegetation Index
- 560 ARW Amphibian and Reptile Wildbook
- 561 B/Y ratio Black-to-yellow ratio
- 562 LMM Linear mixed effects model

# 563 Supplementary Information

564 The supplementary information to this manuscript can be found at the end of this document.

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- 571 Author contributions
- 572 MM, OK, and BAC conceived the idea and designed the study. All authors with the exception of BAC,
- 573 RH, MS, and OK were involved in sampling. RH and MS conducted the image analysis. MM and NA
- 574 conducted the statistical analysis. MM and BAC interpreted the results. All authors contributed
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- 579 Data availability
- 580 Data and R code will be made publicly available on Open Science Framework (OSF) upon acceptance
- 581 of this manuscript for publication.

## 582 **Declarations**

- 583 Ethics approval and consent to participate
- 584 Download of photos from the ARW were approved by the users who uploaded these photos. No
- 585 ethics approval was necessary for this study as no fire salamanders were touched or elsewise
- 586 harmed. If animals were touched by researchers, these procedures were approved in alignment with
- 587 the specific projects.

- 588 Competing interests
- 589 The authors declare no competing interests.
- 590 References

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# 801 Figure captions



802

Fig. 1 Representative specimen of female (♀) and male (♂) fire salamanders (*Salamandra salamandra*) with a 2 €- coin as a size reference. Both individuals were photographed in the same
night in the Botanical Garden Bielefeld (BGBI), a study location. Females are usually much wider in
body shape [42, 43] and, as the lateral side of a fire salamander is usually mostly black, females might
have a lower proportion of yellow when viewed from above despite having the same relative amount
of pigment cells responsible for the yellow coloration.







*salamandra*) (according to [43]) highlighted in yellow. White circles denote the sampling locations.





Fig. 3 Schematic representation of image processing. First, the fire salamander body was cropped as described in [40]. Subsequently, the head of each fire salamander was also cropped out and saved as an additional image to compare overall body coloration with head coloration specifically. Images of the cropped body as well as the head only were then used to automatically quantify the black-toyellow-ratio (B/Y-ratio) of the fire salamander coloration using the Python script provided in [40]. All image processing and analysis was conducted by two researchers (RH and MS) who were blind to the sex and location of the fire salamander depicted in the image.



820

Fig. 4 Differences in A the body black-to-yellow ratio and B the head black-to-yellow ratio of female
(red) and male (blue) fire salamanders by population. Left to the y-axes, you can see a graphical
representation of the coloration of a fire salamander (i.e., very yellow or very black fire salamanders
or fire salamander heads, respectively). The legs of the fire salamanders in panel A were not cropped
for aesthetical reasons. The boxplots show the median (horizontal bar) and the 2<sup>nd</sup> and 3<sup>rd</sup> quartile
with the minimum and maximum range outside of the boxes. The dots represent each sample. Please
note that the black-to-yellow ratios are on the log<sub>10</sub> scale.



828

829 Fig. 5 The correlation of the head black-to-yellow-ratio with the body black-to-yellow ratio. The

regression lines as well as the dots were colored by sex (red – females, blue – males). The correlation

test statistics in the bottom right corner represent the results of the Pearson's correlation test.

832 Please see Supplementary Fig. 2 for a correlation plot by sex for each population.



833

**Fig. 6** The relationship between the head black-to-yellow ratio and **A** latitude and **B** longitude for



representation of the coloration of a fire salamander (i.e., very yellow or very black fire salamander

- heads). The dots represent each sample. Please note that the black-to-yellow ratios are on the log<sub>10</sub>
- scale and that the relationship between head B/Y ratio and longitude was not statistically significant
- 839 (Table 4).

840

## Supplementary Information for the Manuscript titled:

# "Citizen science data supports sexual dichromatism but rejects thermal melanism in the European fire salamander (*Salamandra salamandra*)"

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#### Summary

These supplementary information show additional data of the manuscript titled "Citizen science data supports sexual dichromatism but rejects thermal melanism in the European fire salamander (Salamandra salamandra)". The environmental variables used to investigate the influence of environmental factors on the yellow coloration of fire salamanders in this study showed weak to moderate correlations with each other ( $\rho < 0.690$ ; Supplementary Fig 1). Therefore, in addition to models investigating the individual effect of each environmental variable on the black-to-yellow ratio (B/Y ratio) of fire salamanders, we also used a model that integrated all environmental variables and thereby tests the contribution of each environmental variable decomposed of the effect of every other environmental variable [1]. The correlation of head B/Y ratio and body B/Y ratio was sex and population dependent (Table 3). Therefore, we show correlation plots with different colors for each sex and a panel for each population in Supplementary Fig 2. Using body B/Y ratio in separate models including one environmental variable at a time revealed a negative relationship with longitude (i.e., in the east, fire salamanders have more yellow bodies) and a marginally non-significant negative correlation of males' body B/Y ratio with average temperature but no association with any other environmental variable (all p > 0.101; Supplementary Table 1A; Supplementary Table 1B). Using one model including all environmental variables, indicated no effect of any environmental variable on females' body B/Y ratio, however, this model still indicated that in males, body B/Y ratio declines with longitude (Supplementary Table 1B and C). All models investigating the relationships of the environmental variables with body B/Y ratio indicated that males have a lower body B/Y ratio (i.e., more yellow on their bodies) than females (Supplementary Fig. 4; Supplementary Table 1).



**Supplementary Fig. 1** The correlation of the environmental variables. In the top right hand plots, the Spearman's correlation test statistics for each combination of environmental variables are shown. "Corr" denotes the correlation coefficient  $\rho$ . Asterisks represent the p-value (\* - < 0.05; \*\* - < 0.01; \*\*\* - < 0.001). The diagonal plots show the distribution of the data using density plots. In the bottom left hand plots, you can see the relationship of each combination of the environmental variables as depicted using a regression line. Each dot represents one study location.



Body black-to-yellow ratio (log-scaled)

Head black-to-yellow ratio (log-scaled)

**Supplementary Fig. 2** The correlation of the head black-to-yellow-ratio with the body black-to-yellow ratio within each population. The regression lines as well as the dots were colored by sex (red – females, blue – males).



**Supplementary Fig. 3** The relationship of the head black-to-yellow-ratio with **A** elevation, **B** average temperature **C** temperature change, and **D** the normalized difference vegetation index. The regression lines as well as the dots were colored by sex (red – females, blue – males).



**Supplementary Fig. 4** The relationship of the body black-to-yellow-ratio with **A** elevation, **B** average temperature **C** temperature change, **D** the normalized difference vegetation index, **E** latitude, and **F** longitude. The regression lines as well as the dots are colored by sex (red – females, blue – males).

Supplementary Table 1 A Results of the linear mixed effects models investigating the relationship of body B/Y ratio with each environmental variable individually as well as B the results from the emtrends [2] test of the significance (i.e., difference from 0) of slopes by sex and environmental variable. C the results of the linear mixed effects model investigating the relationship of body B/Y ratio with each environmental variable simultaneously as well as **D** the results from the *emtrends* [2] test of the significance (i.e., difference from 0) of slopes for males by each environmental variable. The output is on the log<sub>10</sub> scale, as body B/Y ratio was log<sub>10</sub>-transformed prior to analysis. The model coefficient estimates ( $\beta$ ) are presented with their standard errors (SE) for the fixed effects. Please note that elevation was divided by 100 and that all environmental variables were centered around their means in order to facilitate better model interpretation (for means of each variable, see Table 4). Test statistics (t) and p-values are given, and all significant effects (p < 0.05) are presented in bold. Variance estimates ( $\sigma^2$ ) for random effects and residual variances are given for the models in A and B. For categorical predictors, reference levels are presented in parentheses after the model parameter name. If the interactions of Sex (Male) with the environmental variables in the large model presented in B were significant, the slopes for females and males were tested with emtrends to investigate if the slopes for each sex were significantly different from 0 and are presented in C.

A) Summaries Variables	of the Models Investigating Ind	ividual Effects of	the Envir	onmental		
Environmental Variable	Model Parameter		Model Output			
	Fixed Effects	в	SE	t	р	
	Intercept (Female)	0.26	0.03	8.68	< 0.01	
Elevation	Sex (Male)	-1.12	0.02	-7.11	< 0.01	
	Elevation	0.00	0.02	0.06	0.95	
	Sex (Male) : Elevation	-0.02	0.01	-1.70	0.09	

	Random Effects	$\sigma^2$			
	Intercept (Population)	0.01			
	Residual	0.04			
	Fixed Effects	в	SE	t	р
	Intercept (Female)	0.26	0.03	9.13	< 0.01
	Sex (Male)	-0.12	0.02	-7.11	< 0.01
Average	Average Temperature	0.03	0.03	1.01	0.33
Temperature	Sex (Male) : Average Temperature	0.03	0.02	1.51	0.13
	Random Effects	$\sigma^2$			
	Intercept (Population)	0.01			
	Residual	0.04			
	Fixed Effects	в	SE	t	р
	Intercept (Female)	0.26	0.03	10.22	< 0.01
	Sex (Male)	-0.12	0.02	-5.44	< 0.01
	Temperature Change	-0.30	0.17	-1.73	0.10
Temperature	Sex (Male) : Temperature Change	0.14	0.15	0.93	0.36
Change	Random Effects	$\sigma^2$			
	Intercept (Population)	0.01			
	Sex (Male)	< 0.01			
	Residual	0.04			
	Fixed Effects	в	SE	t	р
	Intercept (Female)	0.26	0.03	8.67	< 0.01
	Sex (Male)	-0.12	0.02	-7.11	< 0.01
	NDVI	0.08	0.70	0.12	0.90
NDVI	Sex (Male) : NDVI	0.65	0.40	-1.63	0.10
	Random Effects	$\sigma^2$			
	Intercept (Population)	0.02			
	Residual	0.04			
	Fixed Effects	в	SE	t	р
	Intercept (Female)	0.26	0.03	9.99	< 0.01
Latitude	Sex (Male)	-0.12	0.02	-5.34	< 0.01
	Latitude	-0.03	0.02	-1.43	0.17
	Sex (Male) : Latitude	-0.01	0.02	-0.38	0.71
	Random Effects	$\sigma^2$			
	Intercept (Population)	0.01			
	Sex (Male)	< 0.01			
	Residual	0.04			
Longitude	Fixed Effects	в	SE	t	р

Intercept (Female)	0.26	0.02	11.57	< 0.01
Sex (Male)	-0.12	0.02	-5.23	< 0.01
Longitude	-0.03	0.01	-2.97	< 0.01
Sex (Male) : Longitude	-0.00	0.01	-0.29	0.77
Random Effects	$\sigma^2$			
Intercept (Population)	0.01			
Sex (Male)	< 0.01			
Residual	0.04			

B) Estimation of Slopes for Males for Model presented in A

Environmental Variable		Output			
	в	SE	t	р	
Elevation	-0.02	0.02	-0.92	0.37	
Average Temperature	0.06	0.03	1.93	0.07	
Temperature Change	-0.15	0.23	-0.67	0.51	
NDVI	-0.57	0.69	-0.82	0.42	
Latitude	-0.04	0.03	-1.42	0.17	
Longitude	-0.04	0.02	-2.49	0.02	

# C) Summary of the Model Investigating the Effects of each Environmental Variable Simultaneously

		Model Output		
Fixed Effects	в	SE	t	р
Intercept (Female)	0.26	0.03	9.80	< 0.01
Sex (Male)	-0.12	0.02	-7.21	< 0.01
Elevation	0.02	0.05	-7.21	0.76
Average Temperature	0.01	0.07	0.13	0.90
Temperature Change	-0.12	0.35	-0.34	0.74
NDVI	0.11	1.32	0.08	0.93
Latitude	-0.01	0.05	-0.22	0.83
Longitude	-0.03	0.03	-1.00	0.33
Sex (Male) : Elevation	-0.08	0.04	-2.33	0.02
Sex (Male) : Average Temperature	-0.14	0.05	-2.92	< 0.01
Sex (Male) : Temperature Change	0.84	0.23	3.71	< 0.01
Sex (Male) : NDVI	-2.21	0.85	-2.60	< 0.01
Sex (Male) : Latitude	-0.08	0.03	-2.41	0.02
Sex (Male) : Longitude	-0.04	0.02	-2.31	0.02
Random Effects	$\sigma^2$			
Intercept (Population)	0.01			
Residual	0.04			
D) Estimation of Slopes for Males for Model presented in C				

Environmental Variable Output			put	
	в	SE	t	р
Elevation	-0.07	0.05	-1.19	0.25
Average Temperature	-0.13	0.07	-1.75	0.10
Temperature Change	0.72	0.35	2.04	0.06
NDVI	-2.10	1.32	-1.59	0.13
Latitude	-0.09	0.05	-1.77	0.10
Longitude	-0.07	0.03	-2.48	0.02

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