

Among-year variation in the repeatability, within- and between-individual, and phenotypic correlations of behaviors in a natural population

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Abstract When mean behaviors correlate among individuals, they form behavioral syndromes. One way to understand the evolution of such a group-level phenomenon is to compare horizontally patterns of correlations among populations (or species) or follow longitudinally the same population over years in the light of parallel differences in the environment. We applied the longitudinal approach to 8-year field data and analyzed phenotypic correlations, and their within- and between-individual components, among three behaviors (novelty avoidance, aggression, and risk-taking) in male collared flycatchers, *Ficedula albicollis*, in a meta-analytic framework. The phenotypic correlation between novelty avoidance and aggression varied heterogeneously (in some years, it was positive, in others it was negative), while the other pair-wise correlations were consistently positive over the study period. We investigated four potential socio-ecological factors and found evidence that the among-year alterations in the demographic structure of the population (density, age composition) can be

responsible for the heterogeneous effect sizes. Comparing within- and between-individual correlations across pairs of traits, we found that the correlation between aggression and risk-taking at the among-individual level was the strongest suggesting that this relationship has the highest potential to form a behavioral syndrome. Within-year repeatabilities varied among traits, but were systematically higher than between-year repeatabilities. Our study highlights on an empirical basis that there can be several biological and statistical reasons behind detecting a phenotypic correlation in a study, but only few of these imply that fixed behavioral syndromes are maintained in a natural population. In fact, some correlations seem to be shaped by the environment.

Keywords Boldness · Effect size · Flight initiation distance · Personality · Phenotypic correlation · Temperament

Introduction

A striking recognition of recent day's evolutionary behavioral ecology is that, although one would expect individual animals to adaptively adjust each of their behaviors depending on the prevailing environmental conditions, apparently many behaviors cannot vary with unlimited flexibility and in isolation from others (Réale et al. 2007). Linked behaviors form behavioral syndromes, in which the non-independence of traits constrains the evolutionary trajectories that are available for particular behaviors (Dochtermann and Dingemans 2013). A fundamental question is, therefore, why and how such syndromes are maintained over generations (Dall et al. 2004; Dingemans and Wolf 2010; Wolf and Weissing 2010).

Behavioral syndromes can be defined as the between-individual correlation of functionally independent behaviors (Sih et al. 2004a, b; Dingemans and Wolf 2010; Herczeg and

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62 Garamszegi 2012). Therefore, to study behavioral syndromes, 115
63 it is inevitable to obtain repeated measures on the same 116
64 behavior from the same individuals that allow discriminating 117
65 between the within-individual and the between-individual 118
66 correlations (Dingemanse et al. 2012; Garamszegi and 119
67 Herczeg 2012; Brommer 2013; Dingemanse and Dochtermann 120
68 2013). The former type of correlation can emerge if correlative 121
69 behavioral responses to the same environmental factor occur 122
70 plastically at the within-individual level (e.g., within-individual 123
71 correlations between exploration and aggression can develop if 124
72 at low temperatures individuals are rather inactive, generally less 125
73 aggressive, and less explorative, while at higher temperatures 126
74 they become more aggressive and explorative). Only between- 127
75 individual correlations reflect links between individual-specific 128
76 attributes and are relevant for behavioral syndromes. Practically, 129
77 if one collects a single measurement for each trait from each 130
78 individual, correlations between behaviors will provide phenotypic 131
79 correlations, which combine the between-individual and 132
80 within-individual components with unknown magnitudes. 133
81 Making inferences from such phenotypic correlations for behav- 134
82 ioral syndromes (as done in many studies) relies on the assump- 135
83 tion that within-individual variation is negligible (Brommer 136
84 2013), which is a strong interpretive step as behaviors are typi- 137
85 cally very plastic traits (Bell et al. 2009).

86 Given that behavioral syndromes represent between- 139
87 individual correlations (or phenotypic correlations as a surro- 140
88 gate), such a phenomenon is inevitably a population-specific 141
89 attribute thus is manifested only at a higher group level. 142
90 Therefore, one way to investigate how behavioral syndromes 143
91 can evolve is to compare correlation structures across different 144
92 populations or species that experience different selection re- 145
93 gimes, and to determine the socio-ecological factors that gen- 146
94 erate differences in the strength and direction of these corre- 147
95 lations (Conrad et al. 2011; Herczeg and Garamszegi 2012; 148
96 Sih et al. 2012; Carvalho et al. 2013). Embracing such a 149
97 framework focusing on groups of individuals as the unit of 150
98 analysis, Bell (2005) and Dingemanse et al. (2007) investigat- 151
99 ed the activity-aggression-boldness syndrome in different 152
100 populations of the three-spined stickleback (*Gasterosteus* 153
101 *aculeatus*) that inhabit different selective environments and 154
102 genetically differentiated from each other, and found that cer- 155
103 tain types of correlations are population specific, which could 156
104 have resulted from population-level adaptations to presence or 157
105 absence of predation. Similar patterns concerning the 158
106 population-specific correlations have also been described for 159
107 other taxa (Scales et al. 2011; Bengtson et al. 2014; Martins 160
108 and Bhat 2014), but evidence at conflict with the between- 161
109 population divergence of behavioral syndromes has also been 162
110 reported (Brydges et al. 2008; Herczeg et al. 2009; Pruitt et al. 163
111 2010). At a wider scale, meta-analyses comparing a larger 164
112 number of populations of different species also detected inter- 165
113 specific differences in behavioral syndromes as inferred from 166
114 phenotypic correlations across individuals, which can be 167

115 interpreted, at least in part, as the consequence of the dissim- 116
117 ilarities in adaptation processes that species underwent during 117
118 their phylogenetic history (Garamszegi et al. 2012a, 2013). 118
119 However, the comparisons of entities that have been isolated 119
120 over a phylogeographic time scale do not allow discriminating 120
121 whether differences in the correlation structure that are ob- 121
122 served among particular populations (or species) are the result 122
123 of (i) long-term adaptation processes that generate genetic 123
124 differentiation and that stably couples or uncouples behaviors, 124
125 or (ii) phenotypic plasticity by which phenotypic correlations 125
126 are enforced by the specific environments in which popula- 126
127 tions occur, or (iii) both.

127 An alternative approach to the horizontal comparison be- 127
128 tween population/species would be to perform a longitudinal 128
129 analysis of correlations of the same population over much 129
130 smaller time scales. Monitoring concurrent changes in the 130
131 environment would allow understanding how rapidly and un- 131
132 predictably altering environmental components can affect the 132
133 correlation structure of behaviors independently of processes 133
134 due to genetic adaptation (e.g., Sinn et al. 2010; Kazama et al. 134
135 2012). In such a longitudinal framework, detected phenotypic 135
136 correlations could vary among years (or other time scales) 136
137 both for statistical and biological reasons. Statistically, detect- 137
138 ed correlations can be different because (i) between-individual 138
139 correlations vary (i.e., due to differences in genetic or perma- 139
140 nent environment correlations), because (ii) within-individual 140
141 correlations vary, because (iii) correlations due to measure- 141
142 ment error vary, or because (iv) the combinations of these vary 142
143 among samples (Dingemanse et al. 2012; Garamszegi and 143
144 Herczeg 2012; Dingemanse and Dochtermann 2013).

145 The biological reasons behind temporal alterations in the 145
146 correlation structure can include processes due to phenotype- 146
147 dependent selection and phenotypic plasticity. For example, 147
148 yearly shifts in, e.g., predation pressure, food supply, or/and 148
149 social constraints can impose differential selection pressures 149
150 on the reproductive success or survival of different pheno- 150
151 types (Dingemanse et al. 2004). As a consequence, the struc- 151
152 ture of the population will be affected in a way that the yearly 152
153 samples of individuals will represent different genetic or 153
154 permanent environment correlations. On the other hand, 154
155 differences in phenotypic correlations can be attributed to 155
156 differences in within-individual correlations if variation in 156
157 environmental conditions makes individuals to change their 157
158 behaviors from one reproductive event to the next (Bell and 158
159 Sih 2007; Shimada et al. 2010; Sih et al. 2011; Dingemanse 159
160 and Wolf 2013). Such phenotypic plasticity would allow fine 160
161 adaptation at the individual level, in which the prevailing 161
162 environmental conditions elicit the most beneficial display from 162
163 the individuals' behavioral repertoire. These two extreme sce- 163
164 narios are certainly mixed in natural populations, as multiple 164
165 biological processes can be in effect simultaneously for the 165
166 same behavioral correlation, and processes due to both 166
167 phenotype-dependent selection and phenotypic plasticity can 167

168 be in action in parallel. To make it more complex, different
 169 mechanisms may be applied to different pairs of behaviors.
 170 Therefore, it would be desirable to obtain deeper insights from
 171 wild populations of animals in how behavioral correlations
 172 vary among years and to uncover the statistical and biological
 173 causes of such variations by partitioning the within- and
 174 between-individual correlations and also by identifying paral-
 175 lel changes in the socio-ecological environment.

176 When the purpose is to compare patterns of correlations
 177 between traits, the meta-analytic framework offers a powerful
 178 tool to obtain a quantitative summary over a suite of studies
 179 that provide information on different groups of individuals
 180 (Wilson and Lipsey 2000; Borenstein et al. 2009; Ellis
 181 2010). Such an approach can estimate the overall strength
 182 and direction of any biological association in the form of an
 183 effect size by accounting for the underlying sample size, as-
 184 sess the degree of heterogeneity that arises among the findings
 185 of the source studies, and to statistically evaluate how meth-
 186 odological or biological factors shape such differences in the
 187 study results. One can borrow the meta-analytic methodology
 188 to deal with the among-year variation in a biological associa-
 189 tion that occur within the same population, as different years
 190 can be treated as separate studies. This focus differs from that
 191 of the classical ecological application in that the former covers
 192 variation in short temporal scales while the latter typically
 193 targets larger-scale variation across different populations/
 194 species that are separated by geographic distances; thus, the
 195 results have different biological implications. The benef-
 196 it of applying the meta-analysis to the same system that
 197 is consistently studied by the same standards is that it is
 198 not loaded with heterogeneity due to methodology and
 199 publication bias (Kotiaho 2002).

200 Here, our goal was to uncover whether the phenotypic cor-
 201 relations that can be detected in certain years in a natural
 202 population are the result of long-term processes that generate
 203 stable links between different behaviors, or vary more sensi-
 204 tively, as a potential response to the prevailing environmental
 205 conditions. The former mechanism predicts that the strength
 206 and direction of the phenotypic correlations between repeat-
 207 able behaviors are caused by between-individual correlations
 208 and remain consistent and similar across years. However, the
 209 latter scenario predicts considerable between-year variation in
 210 the correlation structure (that is potentially caused by within-
 211 individual correlations) if the environment also fluctuates. We
 212 tested these predictions in a Hungarian population of the col-
 213 lared flycatcher, *Ficedula albicollis*, in which we routinely
 214 monitor different behaviors in males (novelty avoidance,
 215 aggression, risk-taking) during courtship (e.g., Garamszegi
 216 et al. 2006, 2009, 2012b). We used field data from 8 years,
 217 in which we scored the focal behavioral traits upon the arrival
 218 of males from the wintering grounds to calculate phenotypic
 219 correlations. In 5 years, we also collected repeated measure-
 220 ments from the same individuals, which permitted us to

calculate within- and between-individual correlations as 221
 well as repeatabilities in these seasons. Furthermore, we 222
 characterized among-year variation in some environmen- 223
 tal factors by estimating year-specific predation pres- 224
 sure, mean daily temperature (potentially affecting the 225
 availability for food) density (potentially affecting the 226
 availability for breeding opportunities), and age compo- 227
 sition. As an explorative, hypothesis-generating exercise, 228
 we related these environmental variables to among-year 229
 variation in correlation structures. Our investigations relied on a 230
 meta-analytic framework that enabled us to rigorously compare 231
 year-specific correlations among behavioral and ecological 232
 traits. 233

Materials and methods 234

General behavioral measurements to obtain phenotypic correlations 235
 236

Our fieldwork for this study was carried out in a nest-box 237
 population of the collared flycatcher in the Pilis Mountains 238
 close to Budapest, Hungary (47°43'N, 19°01'E). In the breed- 239
 ing seasons 2007 to 2015, we applied non-invasive (i.e., with- 240
 out capturing individuals) methods to characterize three be- 241
 havioral traits in males. From the expected date of the first 242
 birds returning from the wintering sites, we regularly visited 243
 the field site for newly arrived, unpaired males showing the 244
 typical courtship behavior on their territory during the most 245
 active morning period (usually between 6.00 to 12.00 h). 246
 Once these males were localized at a nest-box, we performed 247
 behavioral assays based on standardized protocols that have 248
 been described in detail and validated elsewhere (e.g., 249
 Garamszegi et al. 2006, 2009, 2012b). We excluded year 250
 2008, as we assayed less than five males in that breeding 251
 season and did not screen all behaviors (Table 1). Here, we 252
 only provide information that is important for the interpreta- 253
 tion of the results. 254

255 We first estimated novelty avoidance, defined as the laten- 256
 cy needed to resume a key element of courtship activity in the 257
 presence of a novel object. We assessed baseline courtship 258
 activity by placing a caged stimulus female on top of the 259
 nest-box and measuring the time interval between the male's 260
 appearance on the territory (based on the conspicuous color- 261
 ation and behavior of males, we assumed that that we can spot 262
 them immediately when they arrive on the territory) and its 263
 first landing on the entrance hole of the nest-box (by this 264
 behavior, male flycatchers aim at eliciting a nest-box visit 265
 from the female). Then, we attached a novel object (white 266
 A6 sheet with small random drawings of variable colors) on 267
 the front side of the box and took the same measurements (if a 268
 male did land in the presence of novelty, we recorded 301 s for 269
 this observation based on the duration of the assay). Novelty

Table 1 Summary statistics for the three behavioral variables of males that were collected in eight breeding seasons in a Hungarian population of the collared flycatcher to study between-year variation in phenotypic correlations in a meta-analysis. Sample size, mean, and standard errors are based on the sample of males that were assayed for their behaviors at least once upon their arrival to the breeding ground. Due to the very low sample size, data for 2008 was not used further

Year	Novelty avoidance (latency to land in seconds)			Aggression (latency to fight in seconds)			Risk-taking (flight initiation distance in meters)		
	<i>N</i>	Mean	SE	<i>N</i>	Mean	SE	<i>N</i>	Mean	SE
2007	21	113.2	36.8	23	50.4	21.8	21	11.8	1.5
2008	2	121.5	154.5	0	–	–	3	10.0	5.5
2009	33	12.5	23.1	34	29.7	12.7	32	13.0	1.4
2010	28	108.5	27.5	31	50.3	17.2	31	14.0	1.2
2011	40	195.6	17.6	54	55.8	13.9	51	10.3	0.7
2012	17	201.1	26.9	25	92.3	24.4	22	13.5	1.9
2013	44	138.6	22.8	56	44.5	12.8	54	9.8	0.8
2014	45	119.1	18.3	53	40.0	11.7	52	12.6	1.1
2015	40	110.6	24.1	46	17.4	7.8	47	7.5	0.7

avoidance was calculated as a difference between the latency scores from the two situations, and is the inverse estimate of how individuals tolerate the presence of a novelty stimulus.

After the novelty avoidance test, we scored aggression by exposing the focal bird to a caged stimulus male, with which we stimulated aggressive response from the territory owner. To describe aggression, we timed the latency to the first attack (i.e., the first touch on the cage of the decoy), as elapsed since the appearance of the resident on the territory. Latency to fight predicts several other behavioral variables that describe aggression (Garamszegi et al. 2006). If the male did not attack, we assigned a score of 301 s (our observations lasted 5 min).

When the subject was localized touching the decoy's cage and being engaged in a territorial dispute, or was observed on another frequently visited position (nest-box, nearby branch), we initiated our assessment of risk-taking by measuring flight initiation distance (FID, Blumstein 2003). The observer started to walk towards the focal bird until it noticed the presence of a potential predator and interrupted its current display. The observer continued walking if the resident returned to the decoy's cage (or another focal position) within at least 1 min. This sequence was repeated until the resident bird did not return anymore to this reference position (each individual returned at least once). The closest distance between the decoy and the last standing point of the observer was measured as the number of steps of approximately 1 m to reflect flight initiation distance. By our approach, we aimed at eliminating the confounding effect of very aggressive males not noticing the approaching human (by allowing the focal male to return, we ascertained that it had noticed the observer).

We captured males after the behavioral assays with a conventional nest-box trap for identification and to perform standardized ringing protocols and measurements. We were unable to capture and subsequently identify some birds (95 out of 337) after the behavioral assays. We have previously shown that such between-individual variation in trappability is associated with the differences in the screened behaviors, and the

elimination of non-captured birds from the sample introduces bias when assessing behavioral correlations (Garamszegi et al. 2009). Such tendencies showing that individuals displaying shy behaviors are generally more difficult to capture were also prevalent in the current data covering eight field seasons (novelty avoidance: $t_{268}=2.652$, $P=0.008$; aggression: $t_{320}=2.290$, $P=0.022$, risk-taking: $t_{311}=3.359$, $P<0.001$). Therefore, to avoid such bias and a considerable loss in sample size, we did not exclude unidentified males from our analyses. However, such a strategy may potentially lead to the risk of generating partially non-independent observations, as unidentified males may be repeatedly present in different samples. We assume that the problem posed by the partial non-independence of data should be minor, as based on the list of successfully ringed individuals we estimate that the chance of assaying an individual in 2 or more years is 7.7 % (due to the modest return rate of the species—<15 % in adult males—and the fact that we can only monitor the behavior of a subsample of the population in each year).

Repeated behavioral measurements to estimate within- and between-individual correlations

In five field seasons (2009, 2011, 2013, 2014, 2015), we made efforts to relocate the birds that had been previously assayed upon their arrival to obtain subsequent behavioral measurements until they established pair bounds (birds when caught after the first set of assays were individually marked on their belly with unique combinations of three colors by water-resistant pens). By doing so, we were able to repeat the behavioral tests for about the half of the males (see Table 1 for exact sample sizes) on average 2.74 times (range, two to six occasions). We used these multiple measurements to differentiate statistically between the within-individual and the between-individual correlations within years (see below). We note that repeated measurements could only be acquired

342 for males that had been captured successfully after the first
 343 assay; thus, we could not eliminate biases due to differences in
 344 trappability (and in the probability of re-sights) in this sub-
 345 sample of males. Therefore, caution is needed when compar-
 346 ing phenotypic correlations with within- and between-
 347 individual correlations, as these correspond to different samples
 348 (see more details below).

349 **Socio-ecological variables**

350 We described each breeding season by four types of ecological
 351 variables at the population level for each year. To characterize
 352 year-specific weather conditions, we estimated the mean of
 353 daily temperature observed over the period between 15th
 354 April and 15th May (when the birds arrive and form pairs,
 355 i.e., when we took the behavioral measurements), as measured
 356 at a nearby meteorological station and supplied to the NOAA's
 357 National Climatic Data Center ([ftp://ftp.ncdc.noaa.gov/pub/
 358 data/gsod](ftp://ftp.ncdc.noaa.gov/pub/data/gsod)). This indirect climatic variable appeared to be a
 359 strong predictor of the average temperature that could be
 360 obtained directly via a small meteorological station that
 361 operated for some years in our field station ($r=0.972$, $N=12$,
 362 $P<0.001$). Furthermore, we have found a strong correlation
 363 between the mean daily temperature and the estimated
 364 caterpillar biomass ($r=0.853$, $N=12$, $P<0.001$; caterpillar
 365 biomass was estimated by collecting and weighting the
 366 produced caterpillar frass in a standard way, see Török and
 367 Tóth 1988). Given that caterpillars are one of the main items
 368 on the flycatchers' diet (Löhr 1976), we could reasonably
 369 assume that our climatic variable was a good predictor of
 370 yearly food supply.

371 Predation rate in each year was estimated as the proportion
 372 of nests that were found fully or partially predated from the
 373 egg laying to the chick-feeding period (breeding efforts were
 374 monitored in each nest-box based on regular checks). The
 375 most typical predator of the species is the Pine Marten
 376 *Martes martes* that leaves clear signatures upon their activity
 377 (heavily disturbed nest material, remainings of the chicks, or
 378 incubating females on the top of the nest box). Based on our
 379 long-term data, nest predation rate varies from 0 to 48 %
 380 among years, which mostly involved chick mortality. Given
 381 that such predation events occur *after* the behavioral assays,
 382 we assumed that, if it applies at all, the predation pressure
 383 estimated in 1 year during the period between egg laying
 384 and chick-feeding should only affect behavioral performance
 385 of males during the courtship period *in the next year*. Increase
 386 in predation rate in a given year can have considerable influ-
 387 ence on several demographic parameters in the subsequent
 388 year thus rise differences in the composition of the population
 389 (for example, predation rate in 1 year determines the propor-
 390 tion of immigrant males: $r=-0.721$, $N=18$ years, $P<0.001$).
 391 Furthermore, the degree of predation can affect individual
 392 experience, which can determine risk-taking decisions during

the future reproductive events. Therefore, we matched year- 393
 specific behavioral correlations with predation rate that corre- 394
 sponds to the previous year. 395

The degree of competition for nest boxes among males due 396
 to density effects was determined by considering the number 397
 of potential breeding opportunities estimated from the number 398
 of available nest boxes relative to the number of breeding 399
 pairs. For each year, we counted the total number of nest 400
 boxes that were available for the collared flycatcher for breed- 401
 ing (i.e., the number of nest boxes that were finally occupied 402
 by the collared flycatchers plus the number of empty boxes, 403
 i.e., that were left uninhabited by other hole nesting species 404
 that typically start breeding before flycatchers arrive). Relative 405
 density was then calculated as the number of breeding efforts 406
 of flycatchers/available nest boxes. We further corrected this 407
 estimate for synchrony effects because the level of competi- 408
 tion should be higher when most birds compete for resources 409
 at the same time. Therefore, we determined the time interval 410
 (in days) within which the 90 % of breeding efforts occurred 411
 and with which we further divided the above density index to 412
 express average competition per day. 413

Given that age may affect individual experience, we also 414
 characterized the age structure of the male population. Upon 415
 the ringing protocols (as well as through the binocular obser- 416
 vations of non-captured individuals), we assigned males into 417
 juvenile and adult age categories based on the typical colora- 418
 tion of the wing (Svensson 1984). Then, age structure was 419
 calculated for each year as the number of juvenile individuals 420
 relative to the total number of individuals by using the sample 421
 of males that were assayed for their behaviors. 422

423 **General statistical approaches**

All analyses were carried out in the R statistical environment 424
 (R Development Core Team 2015). Due to various con- 425
 straints, information on some behaviors was not be available 426
 in few cases causing slight variation in sample size both within 427
 and among years (see summary statistics for the yearly sam- 428
 ples in Table 1). The distribution of novelty avoidance and 429
 aggression showed strong deviation from being normal even 430
 after trying various transformations. Therefore, to obtain stan- 431
 dardized and comparable estimates for the strength of different 432
 relationships, we calculated Fisher's Z-transformed Spearman 433
 rank correlations between the three behavioral variables in 434
 each year separately to describe group-level patterns (see also 435
 Dingemanse et al. 2007 for a similar approach in a between- 436
 population context). Previously (Garamszegi et al. 2008, 437
 2009, 2012b), we have assessed the role of several potentially 438
 confounding factors (such as age and other attributes of males, 439
 territory quality, date of measurement, etc.) on these correla- 440
 tions and concluded that, except trappability, none of these 441
 seriously affected the focal relationships. Therefore, for sim- 442
 plicity, we did not consider additional covariates in this study 443

444 and proceeded with raw correlations instead of building complex linear models with several covariates with minor effect. 445 For illustrative purposes (Fig. 1), we present the rank-transformed raw data. The socio-ecological predictors that 446 were calculated as proportions (predation rate, competition index, age structure) were square-root transformed. 447 448 449

450 To process repeated measurements on the same individuals and to calculate the within- and between-individual components of (co-)variances, we used univariate and bivariate mixed modeling (Dingemanse and Dochtermann 2013), available in the *MCMCglmm* (Hadfield 2010) R package. *MCMCglmm* relies on Markov Chain Monte Carlo processes for parameter estimation, for which we defined a relatively uninformative prior specification equivalent to an inverse gamma prior with shape and scale equal to 0.001 and with a belief parameter (ν) set to 1.002 (alternative prior settings, e.g., the use of the default of *MCMCglmm* do not affect qualitatively the results). Each model was run for 1.3 million iterations, sampling every 1000 (thinning interval) after discarding the first 300,000 (burnin). We checked models for convergence and mixing by examining the Gelman–Rubin statistics (Gelman and Rubin 1992; the potential scale reduction factor <1.1 for all parameters) among chains, and for autocorrelation within chains (Hadfield 2010). We also

468 visually assessed the traces of all parameters for independence and consistency of the posterior distributions over iterations. 469 To check the stability of results, each model was fitted at least three times, and we also verified if longer runs (i.e., based on 5 million iterations) gave similar results. 470 471 472

473 As for model definition, to assess the repeatability of traits, we created models assuming normally distributed errors, in which one of the behavioral variables was the response, the corresponding date of observation was the predictor (see the importance of controlling for date effects in Biro and Stamps 2015), and the identity of males was added as random effect term (only random intercept was modeled). From these models, we extracted the estimated variance components and calculated repeatability as the proportion of the between-individual variance relative to the total variance (Nakagawa and Schielzeth 2010; Dingemanse and Dochtermann 2013). The 95 % confidence interval of this metric was determined from the 95 % credibility interval of the posterior distribution of the *MCMCglmm* output. To calculate within-year repeatability, we repeated this procedure for each focal variable separately for each of the 5 years, in which multiple measurements for the same individuals were available. In the between-year context, we relied on males that were scored for their behaviors in more than 1 year over the 8-year period 481 482 483 484 485 486 487 488 489 490 491

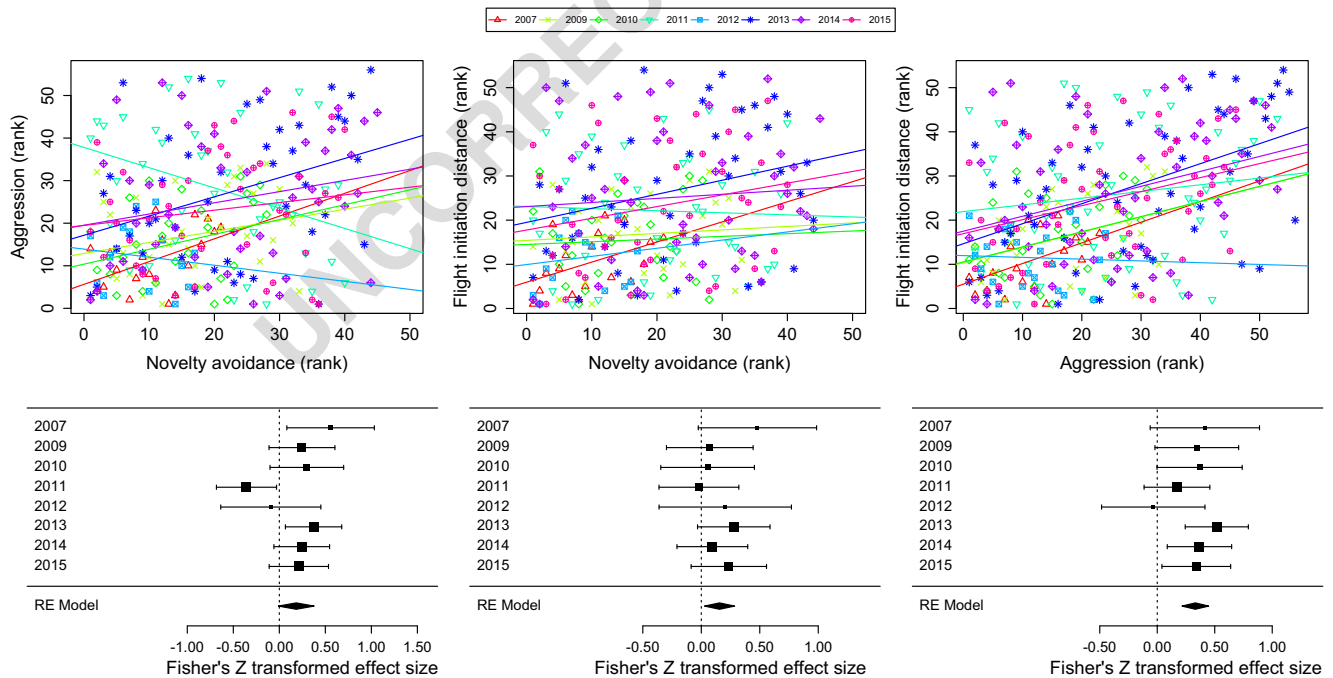


Fig. 1 Year-specific phenotypic correlations among three behavioral traits of male collared flycatchers assayed during the courtship period of eight breeding seasons (2007–2015 with 2008 excluded). *Upper panels* show the pooled ranked raw data and the fitted regression lines using different colors and symbols for different years (individuals were ranked along their behaviors in each year in a way that lower ranks systematically signify bolder behaviors, i.e., lower novelty avoidance and higher aggression and risk-taking). *Lower panels*

present the meta-analysis of the above data relying on years as unit of the analysis. *Black squares* represent year-specific effect size calculated from the corresponding Spearman rank correlation of traits, with a size proportional to the underlying sample size. *Horizontal error bars* represent the 95 % confidence intervals. *Diamonds* are the overall mean effect sizes, as calculated from a random-effect meta-analytic model over the whole 7-year sample, with a width showing 95 % confidence intervals. For exact sample sizes, see Table 1

492 (we only used the first observation, i.e., the one that corre- 543
 493 sponds to the arrival date, from 1 year if repeated measure- 544
 494 ments were available within that year to control for potential 545
 495 date effects). To analyze patterns of variation in repeatability 546
 496 estimates, we used *t* tests, in which *t* values were calculated 547
 497 based on weighted means and weighted variances (where the 548
 498 weights are the years-specific sample sizes, i.e., the number of 549
 499 individuals, see Table 3). Accordingly, we applied weighted 550
 500 univariate *t* tests to check if the within-year repeatabilities of 551
 501 traits are systematically different from their between-year re- 552
 502 peatabilities, and weighted paired *t* tests to compare within- 553
 503 year repeatabilities between pairs of traits. 554

504 For the assessment of within- and between-individual cor- 543
 505 relations, we constructed models (with normal error distribu- 544
 506 tions) by using the pair-wise combination of behavioral traits 545
 507 as bivariate response and identity as random term. We used 546
 508 procedures described in Dingemans and Dochtermann 547
 509 (2013) to obtain the two components of correlation for each 548
 510 relationship for each year. Above, we noted that our subsam- 549
 511 ples of males that have been used for this variance partition 550
 512 might be biased because we could only obtain multiple mea- 551
 513 surements for individuals that had been successfully captured 552
 514 and re-assayed. To evaluate the reliability of the estimates, 553
 515 we calculated the expected phenotypic correlations from 554
 516 them following the mathematical equation presented in 555
 517 Dingemans and Dochtermann (2013), to which we also 556
 518 supplied the estimated within-year repeatabilities. Then we 557
 519 related these expected correlations to the phenotypic cor- 543
 520 relations that we actually observed in the entire datasets 544
 521 also including all non-captured males (note that within- 545
 522 and between-individual correlations could only be derived 546
 523 for birds that had been successfully re-assayed). We found a 547
 524 strong relationship between the two sets of estimates ($r=0.764$, 548
 525 $N=15$, $P<0.001$) implying that the acquired within- and 549
 526 between-individual correlations are reliable. 550

527 **Meta-analyses**

528 In a meta-analysis, first, the outcome of each study (yearly 543
 529 samples in the current context) is converted to a common 544
 530 currency so-called effect size, which is thus comparable across 545
 531 studies (see a comprehensive description about the method in 546
 532 Nakagawa and Santos 2012). Then, an overall effect size is 547
 533 calculated across studies, which is weighted by the precision 548
 534 of the study, with a confidence interval to reflect the precision 549
 535 of the estimate. We used the Fisher's *Z*-transformed Spearman 550
 536 rank correlations as effect sizes, for which we derived confi- 551
 537 dence intervals based on their variance calculated as $1/(N-3)$, 552
 538 where *N* is the corresponding sample size (number of individ- 553
 539 uals). To calculate weighted mean effect sizes over the whole 554
 540 8-year sample, we performed random-effect meta-analytic 555
 541 models assuming that each study year has its own effect size 556
 542 and allowing that they can be different from each other due to 557

biological reasons. We particularly dealt with this degree of 543
 this dissimilarity across findings by performing tests of het- 544
 erogeneity (DerSimonian and Laird 1986). If we found evi- 545
 dence for such strong variance in effect sizes, we further ex- 546
 amined if the detected heterogeneity can be attributed to the 547
 between-year variance in any socio-ecological factor by ap- 548
 plying meta-regression (testing for the effect of moderators in 549
 a meta-analysis only makes sense, when the effect sizes truly 550
 vary across study samples). We relied on the package *metafor* 551
 (Viechtbauer 2010) for the meta-analytic procedures. For in- 552
 terpretations with regard to the magnitude of the effect, we 553
 followed the widely followed benchmarks from evolutionary 554
 ecology and other disciplines, in which untransformed $r\approx 0.1$ 555
 is a small effect, $r\approx 0.3$ is a moderate effect, and $r\approx 0.5$ is a 556
 strong effect (Cohen 1988; Møller and Jennions 2002). 557

Results

Phenotypic correlations

560 The upper panels of Fig. 1 show the relationships as estimated 560
 561 from phenotypic correlations between the ranks of the three 561
 562 behavioral traits separately for each of the 8 years (note that 562
 563 ranks corresponding to latency scores or distances are all in- 563
 564 verse estimates of exploration, aggression, and risk taking, 564
 565 respectively; thus, positive correlations between ranks system- 565
 566 atically imply that bolder individuals in one test are also bold 566
 567 in the other test). The visual inspection of these graphs sug- 567
 568 gests that although there seems to be a general tendency for a 568
 569 positive relationship between behaviors across individuals, 569
 570 there is also considerable variation among pairs of traits and 570
 571 years. In fact, in some years, some relationships can turn neg- 571
 572 ative (e.g., aggression and novelty avoidance in 2011). 572

573 When entering these correlations as effect sizes into a meta- 573
 574 analysis (lower panels of Fig. 1), we found that mean effect 574
 575 size for the relationship between novelty avoidance and 575
 576 aggression cannot be differentiated statistically from zero 576
 577 (untransformed $r=0.182$, $CI_{95\%}=-0.011/0.361$, $N=264$, $P=$ 577
 578 0.065). The other two relationships were generally significant 578
 579 and positive (novelty avoidance and risk-taking: untransformed 579
 580 $r=0.155$, $CI_{95\%}=0.027/0.278$, $N=255$, $P=0.018$; aggression 580
 581 and risk-taking: untransformed $r=0.320$, $CI_{95\%}=0.211/0.420$, 581
 582 $N=307$, $P<0.001$). A comparison of the effect sizes for the two 582
 583 significantly positive relationships yielded a statistically 583
 584 distinguishable, twofold difference in their magnitude 584
 585 ($z=2.06$, $P=0.039$). Another remarkable difference in 585
 586 the between-year patterns of phenotypic correlations of 586
 587 behaviors was that the relationship between novelty 587
 588 avoidance and aggression was heterogeneous (including 588
 589 both positive and negative correlations) among study years 589
 590 ($I^2=56.01\%$, $Q_{df=7}=15.95$, $P=0.026$), but we could not de- 590
 591 rive such evidence for the other two relationships (novelty 591

592 avoidance and risk-taking: $I^2=0\%$, $Q_{df=7}=4.058$, $P=0.773$;
 593 aggression and risk-taking: $I^2=0\%$, $Q_{df=7}=5.746$, $P=0.570$).

594 **Within- and between-individual correlations**

595 We performed some simple analyses to explore patterns of
 596 among-year variation in the within- and between-individual
 597 correlations for those five study years when repeated measure-
 598 ments for the same individuals were available. When pooling
 599 correlations across years and the type of relationships, we
 600 found that year effects did not raise any heterogeneity
 601 either in the between-individual correlation effect sizes
 602 ($Q_{df=1}=0.212$, $P=0.645$) or in the within-individual corre-
 603 lation effect sizes ($Q_{df=1}=0.285$, $P=0.594$). However, we
 604 discovered that the type of the relationship was a signifi-
 605 cant predictor of the between-individual correlations, as
 606 the relationship between aggression and risk-taking was
 607 generally stronger and more consistent than the other rela-
 608 tionships ($Q_{df=1}=9.826$, $P=0.007$, Fig. 2a). Similar con-
 609 clusions could not be made for the within-individual com-
 610 ponents ($Q_{df=1}=0.373$, $P=0.830$; Fig. 2b). However, it is
 611 noteworthy that the among-year variance in the within-
 612 individual correlation for the novelty avoidance/aggression
 613 relationship is the highest. A visual inspection of the data
 614 revealed that the between- or within-individual correlations
 615 covered similar ranges mostly in the positive direction
 616 (Fig. 2), which were also comparable with the variation in
 617 the phenotypic correlations (Fig. 1).

618 **The role of ecological factors**

619 We examined if between-year variance in certain ecological
 620 factors can cause heterogeneity in the detected within-year
 621 patterns of phenotypic correlations between novelty avoid-
 622 ance and aggression (we explored the role of ecological pre-
 623 dictors only for this particular correlation because only this
 624 covered a considerable variation among year-specific effect
 625 sizes that could be explained by a moderator variable).
 626 Corresponding meta-regressions revealed that the age compo-
 627 sition of the population significantly affected the correlation
 628 between the two behaviors when they were entered in a pair-
 629 wise fashion in the model (Table 2 and Fig. 3). However,

when we included the moderators simultaneously into the
 same model, we found that both demographic parameters
 (competition index and age structure) became significant pre-
 dictors (Table 2).

Within- and between-year repeatabilities

The repeatability of behaviors in different contexts is summa-
 rized in Table 3. Focusing on the within-year patterns, repeat-
 ability for risk-taking appeared to be consistently higher than
 for the other two traits (weighted paired t tests, novelty avoid-
 ance vs. aggression: $t_4=0.784$, $P=0.477$; novelty avoidance
 vs. risk-taking: $t_4=-2.532$, $P=0.065$; aggression vs. risk-
 taking: $t_4=-2.964$, $P=0.041$). Furthermore, there was a sys-
 tematic tendency for within-year repeatabilities being higher
 than between-year repeatabilities (weighted one-sample t
 tests, novelty avoidance: $t_4=2.352$, $P=0.078$; aggression:
 $t_4=1.807$, $P=0.145$; risk-taking: $t_4=2.564$, $P=0.062$;
 Fisher's combined significance for the three tests: $P=0.024$).

Discussion

Here, we studied among-year variation in repeatability and
 different types of correlations between three behavioral traits
 in collared flycatcher males from a free-living population. The
 major findings were the following. First, we found that phe-
 notypic correlations for the novelty avoidance/risk-taking and
 for the aggression/risk-taking relationships remained system-
 atically positive across years, while for the novelty avoidance/
 aggression relationship, they varied considerably between
 years in terms of both magnitude and sign. Second, we were
 able to demonstrate that such heterogeneous variation in effect
 sizes for the latter relationship could be mediated by the
 among-year alterations in the studied demographic factors
 determining the level of competition for breeding opportu-
 nities and age composition of the population. Third, within-
 year repeatability of traits varied among the assayed
 behaviors (it was the highest for risk-taking) and tended to
 be considerably higher than their between-year repeatability.
 Finally, we observed that the within-year between-individual
 correlations differed among the considered pairs of traits, as

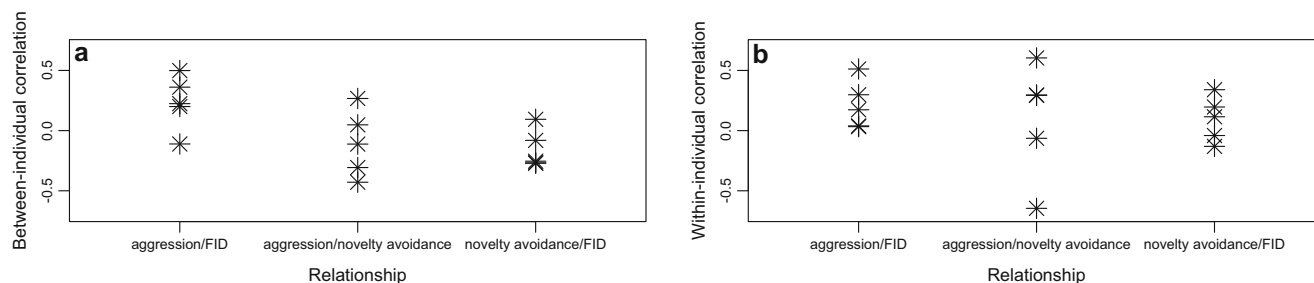


Fig. 2 The effect of the type of correlation on within- and between-individual correlations. Asterisks are year-specific point estimates of effect sizes

t2.1 **Table 2** The effects of four moderator variables on year-specific phenotypic correlations between novelty avoidance and aggression when assessed via meta-regression approaches. On the left side, statistics are given for the cases when moderator variables were tested one by one in different meta-analytic models. On the right side, the effects

correspond to a single multivariate regression model, in which the moderators were entered simultaneously (predation pressure was not included in this multivariate model because it strongly correlated with competition index: $r=-0.887$, $N=9$, $P=0.001$). Lower and upper 95 % confidence intervals for the correlation are given in brackets

t2.2	moderator	Pair-wise model			Multivariate model		
		$Q_{df=1}$	r	P	$Q_{df=3}$	r	P
t2.4	Mean daily temperature	0.692	0.322 (-0.418/0.752)	0.406	0.224 (-0.600/0.771)	0.645	
t2.5	Predation pressure in previous year	2.452	-0.539 (-0.821/0.159)	0.117	Not included		
t2.6	Competition index	1.833	0.484 (-0.240/0.804)	0.176	0.746 (0.138/0.903)	0.025	
t2.7	Age structure	4.671	0.662 (0.082/0.860)	0.031	0.767 (0.221/0.909)	0.016	
t2.8	Full model				12.353	0.006	

667 the aggression/risk-taking relationship was consistently stronger than the other relationships.

668
669 The difference in the mean and variance in effect sizes across
670 pairs of behaviors may question the existence of an overwhelming
671 applicable explanation for phenotypic correlations among
672 repeatable behavioral traits that are often interpreted as evidence
673 for behavioral syndromes (Dingemanse et al. 2012; Garamszegi
674 et al. 2012a; Brommer 2013). The novelty avoidance/aggression
675 and novelty avoidance/risk-taking relationships can be characterized
676 by similarly small overall effect size ($r<0.2$), but the former includes
677 much larger heterogeneity in terms of both magnitude and direction
678 of effect sizes (which causes that the

mean effect size cannot be statistically differentiated from zero in the current sample) than the latter. However, when we focus on phenotypic correlations that homogeneously appear positive in different years, we can still observe twofold obvious differences in their means. In fact, the aggression/risk-taking relationship reached a magnitude that represents moderate effect size, while the novelty avoidance/risk-taking relationship could only be interpreted as being a small effect size. Furthermore, the largest phenotypic correlation between aggression and risk-taking was accompanied by the largest between-individual correlation indicating that each pair-wise relationship was loaded with different within- and between-

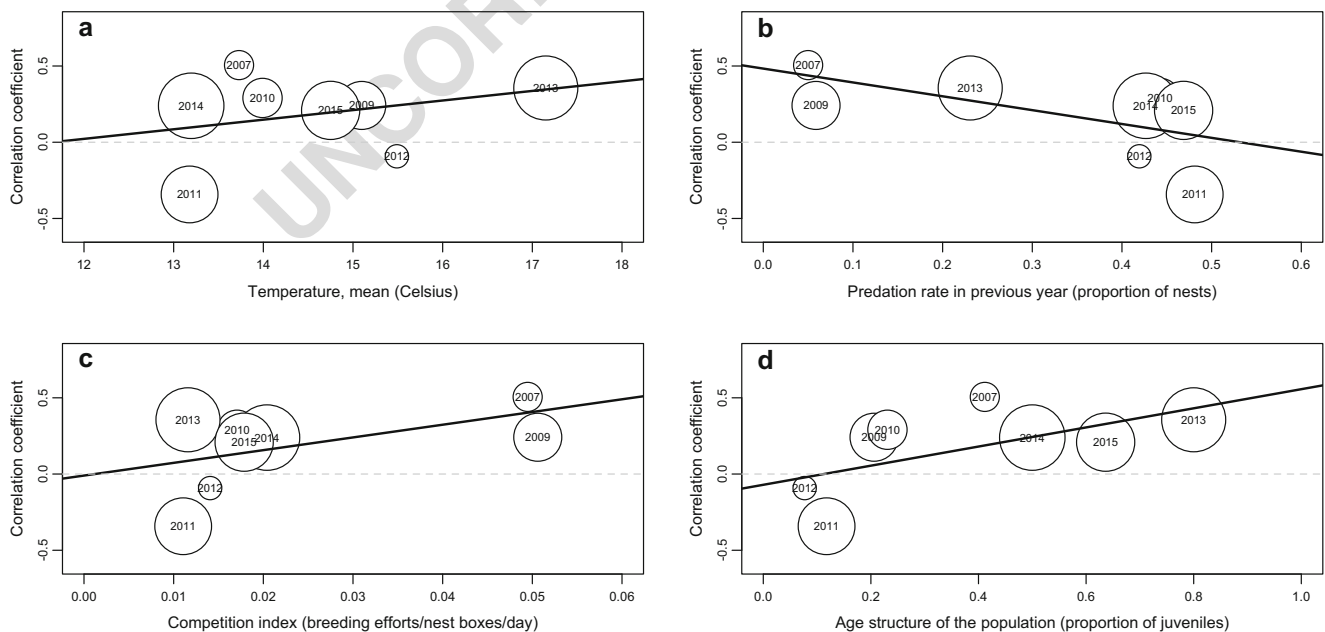


Fig. 3 Meta-regressions demonstrating the effects of four socio-ecological variables on the phenotypic correlation between novelty avoidance and aggression in male collared flycatchers. Each circle represents a correlation that was observed in the designated year with a size that is proportional to the underlying sample size (see Table 1). For

the definition and calculation of the ecological predictors, see the “Materials and Methods” section. Solid lines are the regression lines as were derived from the underlying meta-analyses using the given socio-ecological variable as mediator. Dashed grey lines represent $r=0$ correlations and are shown for guidance

t3.1 **Table 3** Within- and between-year repeatabilities of traits. Within-year repeatabilities are given for 5 years and are based on individuals that were successfully scored for their behaviors at least two times during the courtship period of the same breeding season. The corresponding samples were also used to calculate within- and between-individual

correlations (see Fig. 3). Between-year repeatabilities originate from the entire database covering the 8-year period and were calculated by using males that were tested in at least two different breeding seasons (but only the first measurement was taken from 1 year). Lower and upper 95 % confidence intervals are given in brackets

t3.2	Year	Novelty avoidance (latency to land)		Aggression (latency to fight)		Risk-taking (flight initiation distance)	
t3.3		<i>N</i>	Repeatability	<i>N</i>	Repeatability	<i>N</i>	Repeatability
t3.4	2009	27	0.449 (0.003/0.774)	27	0.345 (0.032/0.631)	26	0.652 (0.406/0.837)
t3.5	2011	16	0.047 (0.000/0.482)	16	0.037 (0.001/0.218)	16	0.116 (0.011/0.432)
t3.6	2013	25	0.235 (0.000/0.629)	28	0.061 (0.002/0.232)	28	0.414 (0.153/0.646)
t3.7	2014	16	0.046 (0.000/0.403)	17	0.185 (0.002/0.565)	17	0.517 (0.070/0.820)
t3.8	2015	18	0.104 (0.000/0.525)	19	0.147 (0.002/0.535)	19	0.109 (0.006/0.402)
t3.9	Between-year	19	0.021 (0.000/0.251)	21	0.058 (0.001/0.314)	21	0.117 (0.009/0.450)

691 individual components. Therefore, even if the studied phe-
692 notypic correlations appear positive in overall, the differ-
693 ences in their strengths and the heterogeneity they cover
694 should signify differences in their biological meaning. We
695 infer that only some of these correlations fulfill criteria for
696 behavioral syndromes.

697 Behavioral syndromes can be maintained in a population
698 if there are rigid genetic, maternal, or early environmental
699 effects that build up developmental or physiological con-
700 straints that finally keep behaviors linked together over lon-
701 ger evolutionary time scales (Sih et al. 2004a, b; Bell 2005;
702 Dochtermann and Dingemanse 2013). Such mechanisms
703 would raise stable between-individual correlations that are
704 independent of the short-term and unpredictable changes in
705 the environment, and could be potentially responsible for
706 the detected patterns in association aggression/risk-taking
707 relationship in the among-year context. In a previous study
708 focusing on the proximate effects of two functionally dif-
709 ferent genes (dopamine receptor *D4* gene and the major
710 histocompatibility complex), we found that flight initiation
711 distance was the variable that depicts the strongest relation-
712 ships with the genetic profile at these regions (Garamszegi
713 et al. 2014, 2015). These findings may imply that observed
714 among-individual variation in this behavioral phenotype is
715 mediated by genetic differences among individuals. The
716 current observation that within-year repeatability is the
717 highest for this behavior is also in line with this interpre-
718 tation. We also note that between-year repeatability for this
719 trait, although it was small, was also the highest and could
720 be differentiated from zero suggesting that between-
721 individual differences in risk-taking remain preserved, at
722 least to some degree, on a longer time scale.

723 The heterogeneous phenotypic correlation between novelty
724 avoidance and aggression, on the other hand, may have result-
725 ed from year to year changes in either the between-individual

or in the within-individual component. Under this scenario,
detected syndromes would not be stabilized by strict mecha-
nistic constraints but would be sensitive to fluctuations in the
environment (Bell 2005, see also Fig. 2 and Table 2 in the
current study) through plasticity or phenotype-dependent se-
lection (Bell and Sih 2007; David et al. 2014). Accordingly,
between-individual correlations for the same relationships
could vary among years if, as a consequence of a socio-
ecological factor, individuals alter their behavioral phenotypes
in a between-year context, even though they maintain
individual-specific correlation structures within the same
breeding season. For example, one can imagine that trait com-
binations that are expressed in a given breeding season were
shaped by experience early in that season/previous winter but
are reshuffled in the next year when new information about the
socio-ecological conditions is gathered. Given that (i) our
between-individual correlations concern with the within-
season context and does not say anything about between-
individual correlations on a longer time scales, and that (ii)
the between-year repeatability of traits was generally low,
between-year changes in the between-individual correlation re-
mains a plausible explanation for the results in association with
the novelty avoidance/aggression relationship. If this applies,
we can preclude that strong genetic (such as in Dochtermann
2011) or long-lasting early environmental effects (such as in
Sweeney et al. 2013; Bengtson et al. 2014; Urszán et al. 2015)
shape the between-individual correlations. On the other
hand, the mediator effects of the demographic parameters
(age-structure and degree of competition) may imply that
individual experience and/or year-specific adjustments to
the available breeding opportunities play more important
roles. Alternatively, we can also imagine that among-year
variation in the correlation patterns emerged, not because of
between-year adjustments within individuals but because of
the yearly shifts in the composition of individuals in the

761 population. Therefore, along the sequence of the study, we
762 would have sampled different groups of individuals that
763 could be characterized by different between-individual cor-
764 relations, which is also a scenario to be considered given
765 the minimal overlap between our yearly samples. This
766 could have occurred, for example, if certain environmental
767 factors had an effect on the survival, reproductive output,
768 and/or dispersal of individuals (Bell and Sih 2007; Logue
769 et al. 2009), and fluctuations in the age-structure and levels
770 of competition have reflected such year-specific phenotype-
771 dependent selection pressures.

772 We cannot exclude the possibility that short-term within-
773 individual effects mediate phenotypic correlations at least in
774 some years (see theory in the "Introduction," empirical exam-
775 ples can be found in Araya-Ajoy and Dingemanse 2014;
776 Brommer et al. 2014; Fresneau et al. 2014; Dösmann et al.
777 2015). For example, the statistically significant negative rela-
778 tionship between novelty avoidance and aggression that ap-
779 peared in 2011 had a very strong within-individual component
780 (Fig. 3). Between-year differences in the within-individual
781 correlations can occur, for instance, if particular socio-
782 ecological factors affect the within-season plasticity of behav-
783 iors in a year-specific way. Hence, there might be years (e.g.,
784 when there are many competitors in the population that is also
785 shifted toward juvenile bias, Fig. 2) when specific within-
786 individual correlations are enforced leading to that if an indi-
787 vidual changes its level of novelty avoidance due to some
788 reasons it also alters its level of aggression in the same direc-
789 tion. In another year, such linked plastic responses may be
790 relaxed or even go in the opposite direction resulting in the
791 situations of no or negative within-individual correlation be-
792 tween the same traits.

793 We must note that our study has certain limitations; thus,
794 certain interpretations should be made with caution. The most
795 important constraints arise from the available sample size.
796 First, although we have assayed more than 300 individuals
797 altogether (Table 1), our framework relied on year-specific
798 focal units (correlation structures) that inherently limits sam-
799 ple size to $N=8$. Meta-analyses can powerfully exploit such
800 samples by accounting for within-year sample sizes, but the
801 effect of particular years remains influential, and the estimated
802 effects all correspond to very broad confidence intervals.
803 Therefore, we cannot reject the hypothesis that we were un-
804 able to deliver statistical evidence for weaker effects that
805 remained non-significant in the current study, or that the in-
806 clusion of additional years with influential effects to the anal-
807 yses can change some of the results. Second, we also relied on
808 modest sample size for the partition of variances and correla-
809 tions into the within- and between-individual component. We
810 could use two to six within-individual repeats for these esti-
811 mations, which also raises statistical issues about precision
812 and bias (Martin et al. 2011; Garamszegi and Herczeg 2012;
813 van de Pol 2012; Dingemanse and Dochtermann 2013). At

814 least, based on the derived within- and between-individual
815 components, we were able to reconstruct the detected pheno-
816 typic correlations and delivered biologically meaningful re-
817 sults suggesting that our estimates were reliable. Third, we
818 should also consider that some of the detected heterogeneities
819 were mediated by variance in measurement errors and not by
820 variance in a biological predictor. In any case, we believe that
821 our study can be definitely expanded to alleviate the above
822 limitations.

823 In summary, our pioneer effort focusing on the temporal
824 variation in the correlation structure of behaviors brings atten-
825 tion to the often-neglected phenomenon that finding a corre-
826 lation between phenotypes in a given study year does not
827 necessarily mean that the same correlation exists in another
828 year. For the study of behavioral syndromes, this implies that
829 finding non-significant correlation between behavioral traits
830 in a narrow study period does not necessarily preclude that
831 syndromes can be formed and detected in other environmental
832 circumstances and based on a larger sample. Furthermore, we
833 can also highlight on an empirical basis that variation in phe-
834 notypic correlations can be due to variation in both the within-
835 individual and between-individual components. This empha-
836 sizes the possibility that different biological explanations are
837 responsible for different phenotypic correlations that are de-
838 tected in a study system, and only few of these are in conform-
839 ity with the definition for behavioral syndromes. We suggest
840 that at least some of the phenotypic correlations appearing in
841 wild animals are ecologically or contextually enhanced phe-
842 nomena that may supersede genetically enforced rules and
843 render within- and/or between-individual correlations spatial-
844 ly and temporally structured. Future research would benefit
845 from the identification of additional socio-ecological factors
846 that mediate long-term among-year variance in the correlation
847 between pairs of behaviors, and also from deeper studies
848 on within- and between-individual correlations that are
849 manifested on longer time scales (e.g., among years).
850 Our meta-analytic framework can be fruitfully applied
851 along these directions, and it can be easily accommodated
852 to deal with questions in relation to changes in the corre-
853 lation structure in space and time.

854 In a wider context, our results point to the importance of the
855 replicability and generalization of findings. Studies are very
856 rare that are able to demonstrate that a relationship that is
857 detected in 1 year is also persistent in other years when envi-
858 ronmental conditions are different (van Noordwijk 1998). To
859 make strong conclusions about general patterns from field
860 studies is only straightforward if the same findings can be
861 delivered in a set of independent studies (coming from differ-
862 ent years or populations), and a statistical summary over these
863 repetitions unanimously reveals evidence for homogeneous
864 patterns. When heterogeneity is detected, it is of scientific in-
865 terest to identify the sources of such heterogeneity (that can be
866 either ecological or methodological).

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879
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 881 national, and/or institutional guidelines for the care and use of animals
 882 were followed. This study was done in agreement with the Hungarian
 883 legislation (Middle-Danube-Valley Inspectorate for Environmental
 884 Protection, Nature Conservation and Water Management, ref. no.’s:
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 886 KTVF 45116-2/2011, KTVF 21664-3/2011, KTVF 12677-4/2012,
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896 **References**

898 Araya-Ajoy YG, Dingemans NJ (2014) Characterizing behavioural
 899 ‘characters’: an evolutionary framework. *Proc R Soc B* 281:
 900 20132645
 901 Bell AM (2005) Behavioural differences between individuals and two
 902 populations of stickleback (*Gasterosteus aculeatus*). *J Evol Biol*
 903 18:464–473
 904 Bell AM, Sih A (2007) Exposure to predation generates personality in
 905 threespined sticklebacks (*Gasterosteus aculeatus*). *Ecol Lett* 10:
 906 828–834
 907 Bell AM, Hankison SJ, Laskowski KL (2009) The repeatability of
 908 behaviour: a meta-analysis. *Anim Behav* 77:771–783
 909 Bengtson SE, Pruitt JN, Riechert SE (2014) Differences in environmental
 910 enrichment generate contrasting behavioural syndromes in a basal
 911 spider lineage. *Anim Behav* 93:105–110
 912 Biro PA, Stamps JA (2015) Using repeatability to study physiological and
 913 behavioural traits: ignore time-related change at your peril. *Anim*
 914 *Behav* 105:223–230
 915 Blumstein DT (2003) Flight-initiation distance in birds is dependent on
 916 intruder starting distance. *J Wildl Manag* 67(4):852–857
 917 Borenstein M, Hedges LV, Higgins JPT, Rothstein HR (2009) Introduction
 918 to meta-analysis. Wiley, West Sussex, UK
 919 Brommer JE (2013) On between-individual and residual (co)variances
 920 in the study of animal personality: are you willing to take the
 921 “individual gambit”? *Behav Ecol Sociobiol* 67:1027–1032
 922 Brommer JE, Karell P, Ahola K, Karstinen T (2014) Residual correla-
 923 tions, and not individual properties, determine a nest defense bold-
 924 ness syndrome. *Behav Ecol* 25:802–812
 925 Brydges NM, Colegrave N, Heathcote RJP, Braithwaite VA (2008)
 926 Habitat stability and predation pressure affect temperament
 927 behaviours in populations of three-spined sticklebacks. *J Anim*
 928 *Ecol* 77:229–235

Carvalho CF, Leitaó AV, Funghi C, Batalha HR, Reis S, Mota PG, Lopes 929
 RJ, Cardoso GC (2013) Personality traits are related to ecology 930
 across a biological invasion. *Behav Ecol* 24:1081–1091 931
 Cohen J (1988) Statistical power analysis for the behavioural sciences, 932
 2nd edn. Lawrence Erlbaum Associates, Hillsdale 933
 Conrad JL, Weinersmith KL, Brodin T, Saltz JB, Sih A (2011) 934
 Behavioural syndromes in fishes: a review with implications for 935
 ecology and fisheries management. *J Fish Biol* 78:395–435 936
 Dall SRX, Houston AI, McNamara JM (2004) The behavioural ecology 937
 of personality: consistent individual differences from an adaptive 938
 perspective. *Ecol Lett* 7:734–739 939
 David M, Salignon M, Perrot-Minnot M-J (2014) Shaping the antipred- 940
 ator strategy: flexibility, consistency, and behavioral correlations 941
 under varying predation threat. *Behav Ecol* 25:1148–1156 942
 DerSimonian R, Laird NM (1986) Meta-analysis in clinical trials. *Control* 943
Clin Trials 7:177–188 944
 Dingemans NJ, Dochtermann NA (2013) Quantifying individual varia- 945
 tion in behaviour: mixed-effect modelling approaches. *J Anim Ecol* 946
 82:39–54 947
 Dingemans NJ, Wolf M (2010) A review of recent models for adaptive 948
 personality differences. *Philos T Roy Soc B* 365:3947–3958 949
 Dingemans NJ, Wolf M (2013) Between-individual differences in be- 950
 havioural plasticity within populations: causes and consequences. 951
Anim Behav 85:1031–1039 952
 Dingemans NJ, Both C, Drent PJ, Tinbergen JM (2004) Fitness conse- 953
 quences of avian personalities in a fluctuating environment. *Proc R* 954
Soc Lond B 271:847–852 955
 Dingemans NJ, Wright J, Kazem AJN, Thomas DK, Hickling R, 956
 Dawnay N (2007) Behavioural syndromes differ predictably be- 957
 tween 12 populations of three-spined stickleback. *J Anim Ecol* 76:
 1128–1138 959
 Dingemans NJ, Dochtermann NA, Nakagawa S (2012) Defining behav- 960
 ioural syndromes and the role of “syndrome deviation” in under- 961
 standing their evolution. *Behav Ecol Sociobiol* 66:1543–1548 962
 Dochtermann NA (2011) Testing Cheverud’s conjecture for behavioral 963
 correlations and behavioral syndromes. *Evolution* 65:1814–1820 964
 Dochtermann NA, Dingemans NJ (2013) Behavioral syndromes as evo- 965
 lutionary constraints. *Behav Ecol* 24:806–811 966
 Dosmann AJ, Brooks KC, Mateo JM (2015) Within-individual correla- 967
 tions reveal link between a behavioral syndrome, condition, and 968
 cortisol in free-ranging Belding’s ground squirrels. *Ethology* 121:
 125–134 969
 Ellis PD (2010) The essential guide to effect sizes: an introduction to 970
 statistical power, meta-analysis and the interpretation of research 971
 results. Cambridge University Press, Cambridge, UK 972
 Fresneau N, Kluehn E, Brommer JE (2014) A sex-specific behavioral 973
 syndrome in a wild passerine. *Behav Ecol* 25:359–367 974
 Garamszegi LZ, Herczeg G (2012) Behavioural syndromes, syndrome 975
 deviation and the within- and between-individual components of 976
 phenotypic correlations: when reality does not meet statistics. 977
Behav Ecol Sociobiol 66:1651–1658 978
 Garamszegi LZ, Rosivall B, Hegyi G, Szöllösi E, Török J, Eens M (2006) 980
 Determinants of male territorial behavior in a Hungarian collared 981
 flycatcher population: plumage traits of residents and challengers. 982
Behav Ecol Sociobiol 60:663–671 983
 Garamszegi LZ, Eens M, Török J (2008) Birds reveal their personality 984
 when singing. *PLoS ONE* 3:e2647 985
 Garamszegi LZ, Eens M, Török J (2009) Behavioural syndromes and 986
 trappability in free-living collared flycatchers, *Ficedula albicollis*. 987
Anim Behav 77:803–812 988
 Garamszegi LZ, Markó G, Herczeg G (2012a) A meta-analysis of corre- 989
 lated behaviours with implications for behavioural syndromes: mean 990
 effect size, publication bias, phylogenetic effects and the role of 991
 mediator variables. *Evol Ecol* 26:1213–1235 992
 Garamszegi LZ, Rosivall B, Rettenbacher S, Markó G, Zsebök S, 993
 Szöllösi E, Eens M, Potti J, Török J (2012b) Corticosterone, 994

995 avoidance of novelty, risk-taking and aggression in a wild bird: no
 996 evidence for pleiotropic effects. *Ethology* 118:621–635

997 Garamszegi LZ, Marko G, Herczeg G (2013) A meta-analysis of
 998 correlated behaviors with implications for behavioral syndromes:
 999 relationships between particular behavioral traits. *Behav Ecol* 24:
 1000 1068–1080

1001 Garamszegi LZ, Mueller JC, Markó G, Szász E, Zsebők S, Herczeg G,
 1002 Eens M, Török J (2014) The relationship between DRD4 polymor-
 1003 phism and phenotypic correlations of behaviors in the collared fly-
 1004 catcher. *Ecol Evol* 4:1466–1479

1005 Garamszegi LZ, Zagalska-Neubauer M, Canal D, Markó G, Szász E,
 1006 Zsebők S, Szöllösi E, Herczeg G, Török J (2015) Malaria parasites,
 1007 immune challenge, MHC variability, and predator avoidance in a
 1008 passerine bird. *Behav Ecol* 26:1292–1302

1009 Gelman A, Rubin DB (1992) Inference from iterative simulation using
 1010 multiple sequences. *Stat Sci* 7:457–511

1011 Hadfield JD (2010) MCMC methods for multi-response generalized
 1012 linear mixed models: the MCMCglmm R package. *J Stat Softw*
 1013 33:1–22

1014 Herczeg G, Garamszegi LZ (2012) Individual deviation from behavioural
 1015 correlations: a simple approach to study the evolution of behavioural
 1016 syndromes. *Behav Ecol Sociobiol* 66:161–169

1017 Herczeg G, Gonda A, Merila J (2009) Predation mediated population
 1018 divergence in complex behaviour of nine-spine stickleback
 1019 (*Pungitius pungitius*). *J Evol Biol* 22:544–552

1020 Kazama K, Niizuma Y, Watanuki Y (2012) Consistent individual varia-
 1021 tions in aggressiveness and a behavioral syndrome across breeding
 1022 contexts in different environments in the Black-tailed Gull. *J Ethol*
 1023 30:279–288

1024 Kotiaho JS (2002) Meta-analysis, can it ever fail? *Oikos* 96:551–553

1025 Logue DM, Mishra S, McCaffrey D, Ball D, Cade WH (2009) A
 1026 behavioral syndrome linking courtship behavior toward males
 1027 and females predicts reproductive success from a single mating
 1028 in the hissing cockroach, *Gromphadorhina portentosa*. *Behav*
 1029 *Ecol* 20:781–788

1030 Löhr H (1976) Studies of less familiar birds. 179. Collared flycatcher.
 1031 *Brit Birds* 69:20–26

1032 Martin JGA, Nussey DH, Wilson AJ, Reale D (2011) Measuring individ-
 1033 ual differences in reaction norms in field and experimental studies: a
 1034 power analysis of random regression models. *Methods Ecol Evol* 2:
 1035 362–374

1036 Martins EP, Bhat A (2014) Population-level personalities in zebrafish:
 1037 aggression-boldness across but not within populations. *Behav Ecol*
 1038 25:368–373

1039 Møller AP, Jennions MD (2002) How much variance can be explained by
 1040 ecologists and evolutionary biologists. *Oecologia* 132:492–500

1041 Nakagawa S, Santos ESA (2012) Methodological issues and advances in
 1042 biological meta-analysis. *Evol Ecol* 26:1253–1274

1043 1092

Nakagawa S, Schielzeth H (2010) Repeatability for Gaussian and non-
 Gaussian data: a practical guide for biologists. *Biol Rev* 85:935–956

Pruitt JN, Riechert SE, Iturralde G, Vega M, Fitzpatrick BM, Avilés L
 (2010) Population differences in behaviour are explained by shared
 within-population trait correlations. *J Evol Biol* 23:748–756

R Development Core Team (2015) R: a language and environment for
 statistical computing. R Foundation for Statistical Computing,
 Vienna, Austria. <http://www.R-project.org>

Réale D, Reader SM, Sol D, McDougall PT, Dingemanse NJ (2007)
 Integrating animal temperament within ecology and evolution.
Biol Rev 82:291–318

Scales J, Hyman J, Hughes M (2011) Behavioral syndromes break down
 in urban song sparrow populations. *Ethology* 117:887–895

Shimada M, Ishii Y, Shibao H (2010) Rapid adaptation: a new dimension
 for evolutionary perspectives in ecology. *Popul Ecol* 52:5–14

Sih A, Bell A, Johnson JC (2004a) Behavioral syndromes: an ecological
 and evolutionary overview. *Trends Ecol Evol* 19:372–378

Sih A, Bell AM, Johnson JC, Ziemba RE (2004b) Behavioral syndromes:
 an integrative overview. *Q Rev Biol* 79:241–277

Sih A, Ferrari MCO, Harris DJ (2011) Evolution and behavioural res-
 sponses to human-induced rapid environmental change. *Evol Appl*
 4:367–387

Sih A, Cote J, Evans M, Fogarty S, Pruitt J (2012) Ecological implica-
 tions of behavioural syndromes. *Ecol Lett* 15:278–289

Sinn DL, Moltchanivskyj NA, Wapstra E, Dall SRX (2010) Are behav-
 ioral syndromes invariant? Spatiotemporal variation in shy/bold beh-
 avior in squid. *Behav Ecol Sociobiol* 64:693–702

Svensson L (1984) Identification guide to European passerines, 3rd edn.
 British Trust for Ornithology, Stockholm

Sweeney K, Gadd RDH, Hess ZL et al (2013) Assessing the effects of
 rearing environment, natural selection, and developmental stage on
 the emergence of a behavioral syndrome. *Ethology* 119:436–447

Török J, Tóth L (1988) Density dependence in reproduction in the col-
 lared flycatcher (*Ficedula albicollis*) at high population levels. *J*
Anim Ecol 57:251–258

Urszán TJ, Török J, Hettyey A, Garamszegi LZ, Herczeg G (2015)
 Behavioural consistency and life history of *Rana dalmatina* tad-
 poles. *Oecologia* 178:129–140

van de Pol M (2012) Quantifying individual variation in reaction norms:
 how study design affects the accuracy, precision and power of ran-
 dom regression models. *Methods Ecol Evol* 3:268–280

van Noordwijk A (1998) The absence of evidence and the evidence for an
 absence. *Acta Oecol* 19:U1–U1

Viechtbauer W (2010) Conducting meta-analyses in R with the metafor
 package. *J Stat Softw* 36:1–48

Wilson DB, Lipsey MW (2000) Practical meta-analysis. Sage, Thousand
 Oaks

Wolf M, Weissing FJ (2010) An explanatory framework for adaptive
 personality differences. *Philos T Roy Soc Lond B* 365:3959–3968

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