

Population biology and phenology of the colour polymorphic damselfly *Ischnura elegans* at its southern range limit in Cyprus

BEATRIZ WILLINK,¹ RACHEL BLOW,² DAVID J. SPARROW,³ ROSALYN SPARROW³ and ERIK I. SVENSSON⁴ ¹School of Biology, University of Costa Rica, San José, Costa Rica, ²Department of Zoology, University of Cambridge, Cambridge, U.K., ³Cyprus Dragonfly Study Group, Paphos, Cyprus and ⁴Evolutionary Ecology Unit, Department of Biology, Lund University, Lund, Sweden

Abstract. 1. Geographically widespread species provide excellent opportunities to investigate how phenotypes change across large-scale environmental gradients. Temperature is a fundamental environmental variable and an important determinant of insect fitness. However, field research is often geographically restricted, and typically concentrated in northern latitudes. Basic population biology and phenotypic clines in relation to temperature therefore remain poorly known across the entire geographic range, even in otherwise well-studied taxa.

2. We surveyed populations of the trimorphic damselfly *Ischnura elegans* in Cyprus, which is the southern range limit in Europe of this widespread insect species. Females of *I. elegans* occur in three discrete and heritable colour morphs, which vary in suites of phenotypic traits. One of these female morphs is a male-mimic that avoids excessive male-mating harassment by its male-like appearance, and which is more cold-tolerant than the two other morphs.

3. In contrast to the situation in northern Europe, these male-mimicking females are the minority morph in Cyprus, representing only about 5% of all females. Male mimics also have lower mating rates than alternative female morphs.

4. Individuals in Cyprus are relatively small in comparison to the reported European range for body size, consistent with Bergman's rule.

5. Finally, populations of *I. elegans* on the island have the longest flight period known in Europe, and there is only partial evidence for seasonality in flight activity.

6. These results underscore the benefits of considering the entire range of environmental conditions encountered by insect species when conducting evolutionary ecology research.

Key words. Body size, colour polymorphism, flight season, geographic variation, mating rate, temperature.

Introduction

The genetic structure and reproductive phenology of animal populations vary geographically, particularly in widely distributed species that occupy diverse habitats and encounter contrasting ecological conditions (Slatkin, 1987; Conover, 1992; Lenormand, 2002; Morrison & Hero, 2003). Quantifying such

geographic variation is necessary to understand how geographic differences in selection can shape genetic and phenotypic diversity. For example, latitude is one of the most intensely investigated geographic axes of genetic differentiation between animal populations (Palo *et al.*, 2003; Fabian *et al.*, 2012). Latitude is also associated with clinal variation in numerous phenotypic traits (Azevedo *et al.*, 1998; Schmidt & Paaby, 2008; Parsons & Joern, 2014), the timing of breeding periods (Schmidt & Paaby, 2008; Mazaris *et al.*, 2013; Parsons & Joern, 2014) as well as allele frequencies underlying discrete phenotypic morphs (Forsman & Shine, 1995; Phifer-Rixey *et al.*, 2008;

Correspondence: Erik Svensson, Evolutionary Ecology Unit, Department of Biology, Lund University, SE-223 62 Lund, Sweden. E-mail: erik.svensson@biol.lu.se

Gosden *et al.*, 2011; Takahashi *et al.*, 2011). A better understanding of how environmental factors such as temperature, seasonality, and humidity underpin observed latitudinal gradients can be used to capitalize on geography to infer ecological and evolutionary responses to ongoing climate change (De Frenne *et al.*, 2013). Yet, to achieve this goal, an essential first step is to build up a comprehensive record of population features across diverse species and their entire distribution ranges.

Despite the ubiquity of geographic variation in mean phenotypes, allele frequencies, and phenologies, most research on many wide-ranging species have been limited to the areas inhabited by the majority of researchers, leading to a geographically restricted understanding of their ecology (Zuk, 2016). In terms of numbers of researchers and more favourable economic conditions for basic research, there has been an ongoing bias to carry out research in northwestern Europe and North America, whereas other regions of the world remain largely understudied. For example, the Common Bluetail, *Ischnura elegans*, is a pond damselfly species (family Coenagrionidae) with a broad distribution across the Palearctic. It occurs from Spain in the West, to Japan in the East, and from Sweden in the North to Iran in the South (Boudot & Salamun, 2015). Classic studies on the life-history and demography of *I. elegans* (Parr, 1965, 1973; Parr & Palmer, 1971; Parr & Parr, 1972; Van Noordwijk, 1978; Hinnekint, 1987; Anholt *et al.*, 2001) have been all conducted in Western Europe and Scandinavia, thereby capturing only a small fraction of the environmental conditions this species encounters across its entire range. Similarly, the flight season, during which individuals may be reproductively active and subject to sexual selection, has been investigated in depth only in this same region (Parr, 1965, 1970). Although *I. elegans* has been subject to many recent ecological and evolutionary studies under natural or seminatural conditions (Takahashi *et al.*, 2014; Le Rouzic *et al.*, 2015; Lancaster *et al.*, 2017; Willink & Svensson, 2017; Willink *et al.*, 2019; Svensson *et al.*, 2020), there is a strong geographic bias in these previous studies, which limits our understanding of its reproductive and population biology, and how they vary across a larger geographic area.

During the last two decades, *I. elegans* has been subject to numerous studies focusing on the behavioural mechanisms behind, and the population consequences of, the maintenance of genetic variation (Van Gossum *et al.*, 1999, 2001; Svensson *et al.*, 2005; Le Rouzic *et al.*, 2015). This interest is motivated by the existence of a heritable and discrete female-limited colour polymorphism, which can be used as a visual genetic marker to study *I. elegans* (Sánchez-Guillén *et al.*, 2005; Svensson *et al.*, 2009). Previous studies have shown that the female colour polymorphism in *I. elegans* is maintained locally by negative frequency-dependent selection via male mating and pre-mating harassment (Svensson *et al.*, 2005, 2009; Gosden & Svensson, 2009; Le Rouzic *et al.*, 2015). Males disproportionately target their mating attempts towards a particular female morph as its frequency increases (Van Gossum *et al.*, 2001). Excessive mating in turn reduces female fecundity (Gosden & Svensson, 2007), leading to a fitness advantage to rare females (Gosden & Svensson, 2009). In this and other damselfly species with similar female-colour polymorphisms, one of the female morphs has a colour pattern nearly identical to

that of males (Robertson, 1985; Van Gossum *et al.*, 2011). Such male-like females benefit from their visual resemblance to males, which acts to further reduce superfluous male-mating attempts, and they are therefore considered to be male mimics (Robertson, 1985; Hammers & Van Gossum, 2008; Gosden & Svensson, 2009; Iserbyt *et al.*, 2011; Van Gossum *et al.*, 2011). However, due to plasticity in male search image formation, the extent of such an advantage of male mimicry is also frequency- and probably density dependent (Hammers & Van Gossum, 2008; Gosden & Svensson, 2009; Iserbyt *et al.*, 2011). These frequency-dependent male–female mating interactions will prevent the fixation of any single female morph locally. In fact, of over one hundred populations surveyed in Europe, none is monomorphic and only a few (~10%) appear to be dimorphic, with one of the three female morphs being absent or extremely rare (Gosden *et al.*, 2011). This suggests that such frequency-dependent selection is strong and operating in most populations. Moreover, in a decade-long longitudinal study of more than a dozen populations of *I. elegans* in southern Sweden, none lost any of the three morphs and all remained trimorphic during more than ten generations (Le Rouzic *et al.*, 2015).

Despite the frequency-dependent processes that maintain the polymorphism locally, populations can vary quite markedly in morph composition across large geographic scales (Gosden *et al.*, 2011). Importantly, female morphs differ in several phenotypic traits, besides colouration, that are ecologically relevant, namely resistance and tolerance against parasitic mites (Willink & Svensson, 2017), cold tolerance (Lancaster *et al.*, 2017), larval developmental time (Abbott & Svensson, 2005) and temperature sensitivity of adult sexual maturation and colour development (Svensson *et al.*, 2020). Therefore, ecological selective agents, other than male mating harassment, such as climatic factors, parasite prevalence, and virulence, are also likely to shape geographic variation in morph frequencies across the large range of *I. elegans*. For instance, the equilibrium frequency of the more cold-tolerant morph could be higher in more northerly populations, with cooler temperatures during the flight season (Svensson *et al.*, 2020). Such ecological agents shaping regional variation in morph frequencies are also likely to operate in tandem with sexual conflict dynamics. Our knowledge about how these different selection processes interact, reinforce or oppose each other can be increased with broader geographic studies of the population biology of *I. elegans*, to include the full range of the ecological conditions that this widely distributed species encounters across its range. In particular, in-depth studies at the southern range limit of this widespread species should complement recent studies on phenotypic and genomic variation at its northern range limit in Sweden (Lancaster *et al.*, 2016, 2017; Dudaniec *et al.*, 2018). In addition, the environmental and genetic factors that determine the location of species range limits is a question that is of central interest to all ecologists, evolutionary biologists and conservation biologists (Kirkpatrick & Barton, 1997; Bridle & Vines, 2007).

Here, we report the results from surveys of breeding populations of *I. elegans* throughout Cyprus, at the southern range limit of this species in Europe. We recorded female morph frequencies using standardised field sampling procedures based on previous studies (Svensson *et al.*, 2005; Gosden & Svensson, 2009; Le

Rouzić *et al.*, 2015). These data add to a growing database of standardised population samplings of *I. elegans* in Europe where morph frequencies have been quantified in a similar fashion (Svensson *et al.*, 2020). Moreover, Cyprus has a Mediterranean climate, that differs strongly from the climate in Sweden and northwestern Europe (Cornes *et al.*, 2018), where most previous studies on natural populations have been conducted.

We also report on the flight season of *I. elegans* on Cyprus and compare it to the flight season in other parts of Europe. Flight season data were derived from the Cyprus Dragonfly Study Group (CDSG) database, from records collected between January 2013 to December 2019. Population densities and the strength of mating interactions are likely to vary seasonally and geographically and are likely to be influenced by the overall length of the reproductive season. Consequently, information on reproductive phenology is important for understanding the interaction between natural and sexual selection in this and other animal species. Furthermore, by influencing the strength and opportunity for intersexual interactions, breeding phenology may influence the fitness costs and benefits of the alternative reproductive and ecological strategies in the different morphs (Machado *et al.*, 2016), thereby shaping geographic variation in the equilibrium morph frequencies among populations.

Finally, we present data on body size in both sexes and all three female morphs from Cyprus. Adult body size is a fundamental and ecologically important phenotypic trait that is almost invariably a target of natural and sexual selection in dragonflies and damselflies (Waller & Svensson, 2017). Our study therefore provides the groundwork necessary to establish *I. elegans* as a model organism for ecological studies across wider biogeographic scales.

Materials and methods

Study species

Colour development in females of *I. elegans* is controlled by a single locus or a set of tightly linked loci, with sex-limited expression and three alleles in a dominance hierarchy (Sánchez-Guillén *et al.*, 2005; Svensson *et al.*, 2009). Thus, in females, the six possible genotypes give rise to three discernible colour morphs, whereas males are always monomorphic. One of the female morphs, traditionally known as *Androchrome* and hereafter referred to as A-females, develops a colour pattern very similar to that of males, with green-blue antehumeral stripes and a bright blue patch on the eighth abdominal segment (Cordero *et al.*, 1998; Svensson *et al.*, 2009; Fig. 1a). The other two female morphs, denoted as *Infuscans* and *Infuscans-obsoleta* (and abbreviated as I- and O-females respectively), are strikingly different from males when sexually mature (Cordero *et al.*, 1998; Svensson *et al.*, 2009; Fig. 1b,c). I-females in their final, sexually mature colouration have olive-green to brown antehumeral stripes (Fig. 1b). O-females lack stripes but exhibit a uniform pink thoracic colour (Fig. 1c), that turns bronze-brown upon sexual maturity. Both I- and O-females express the bright blue abdominal patch of males and A-females early after their emergence from the larval state (Svensson *et al.*, 2009). However, these

two female morphs conceal the blue colouration during adult development, so that unlike males and A-females, sexually mature I- and O-females exhibit a dark abdominal colouration dorsally (Cordero *et al.*, 1998; Svensson *et al.*, 2009; Willink *et al.*, 2019). Immature A- and I-females have similar violet markings on the thorax, and their striking colour differences described above develop with the onset of sexual maturation (Willink *et al.*, 2019, 2020).

The Common Bluetail, *I. elegans*, breeds in diverse habitats throughout Europe. It is found in ponds, lake shores, and slow-flowing streams, primarily in open landscapes and up to an elevation of 1600 m.a.s.l. (Boudot & Salamun, 2015). *I. elegans* is considered a widespread, non-threatened species within its local range and is highly tolerant to human disturbance of aquatic habitats (Harabiš & Doln, 2012). Voltinism, the number of generations in a year, is inversely correlated with latitude, ranging from larvae which take more than a year to emerge onto land in Scotland, to multiple generations in the same season in Southern Europe (reviewed in Corbet *et al.*, 2006). Latitudinal variation in voltinism is in turn contingent on variation in the length of the reproductive season. Reproductive phenology has not been systematically investigated in *I. elegans*, yet in other widely distributed species of dragonflies and damselflies breeding seasons are longer at lower latitudes (reviewed in Corbet, 1999).

Field sampling

Field surveys of female morph frequencies of *I. elegans* were conducted in three mating seasons (2017–2019), each year over the course of one week in June. These surveys spanned a total of 45 localities throughout the island (20 in 2017, 24 in 2018, and 24 in 2019; Fig. 1d), where *I. elegans* had been previously recorded in large numbers (D.J. Sparrow and R.L. Sparrow, pers. obs.). Surveys were conducted using comparable procedures as in previous studies of *I. elegans* populations in Sweden (Svensson *et al.*, 2005; Gosden & Svensson, 2009; Le Rouzić *et al.*, 2015). In short, a single researcher walked a transect along suitable breeding habitat in each locality, collecting individuals with a hand net upon encounter. Collected individuals were temporarily housed in plastic mesh cages if single and in plastic cups, if collected while mating. After the sampling period (mean = 32.58 min; SD = 19.78), we recorded the sex and mating status of all collected individuals, and for females, the colour morph and developmental colour phase were also recorded. Females were classified to the immature colour phase if they displayed a bright blue colour patch and, in the case of A- and I-females, if they had violet markings in the thorax. Females with their final sexually mature colour pattern were classified to the mature colour phase.

Individuals were subsequently released at or near (within a few hundred meters) their site of collection. Temporarily caging individuals prevented us from double-counting any damselflies and allowed for more accurate estimates of female morph frequencies. A few sampled individuals (mean = 2.42; SD = 1.39; per sex and female morph) in each of the localities with a higher abundance of *I. elegans* ($n = 26$) were used

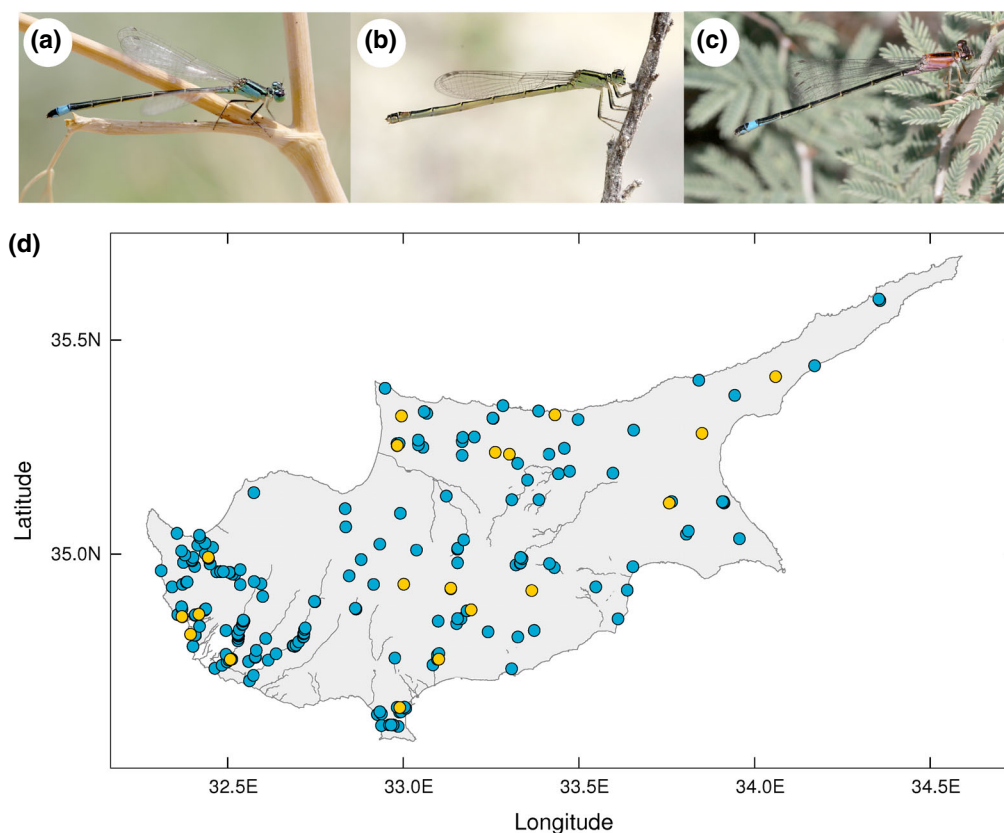


Fig. 1. Colour variation and study localities of the Common Bluetail damselfly, *Ischnura elegans*, in Cyprus. A heritable colour polymorphism controlled by a single locus or set of tightly linked loci is expressed only in females. (a) The dominant allele (p) drives the development of male-coloured A-females, thought to be male-mimics. (b) The I-female morph results from two genotypes (qq and qr) and is markedly different from males and A-females when sexually mature. (c) Only one genotype (rr) gives rise to O-females, which unlike males and the two other female morphs lack antehumeral stripes. An immature O-female is shown. I- and O-females also differ from males and A-females in that the former two conceal the abdominal blue patch over the course of adult development, so that it is no longer visible in sexually mature individuals. (d) Localities surveyed by the Cyprus Dragonfly Study Group, between January 2013 and December 2019. Localities marked in yellow were used for morph-frequency estimation. [Colour figure can be viewed at wileyonlinelibrary.com].

for body size measurements prior to their release. We used a hand-held electronic calliper to measure the total length (excluding genital appendages), abdomen length, and hind wing length of these individuals.

The CDSG is a citizen-science group founded in late 2012. Since 1st January 2013 its members have been monitoring dragonfly and damselfly communities along defined transects at over 50 sites, selected to give broad geographic coverage, and include all habitat types and known species in the island. For each species at each site, the number of single individuals is counted, where possible recording males and females separately. The numbers of pairs in tandem, pairs in copula, and females ovipositing are also recorded. These records are entered into the CDSG database along with other incidental records and those from visitors and non-CDSG members. The CDSG aims to monitor each site monthly or twice-monthly, yet as this is a citizen-science effort, data completeness varies among sites. Although several of the initially selected sites were lost for a variety of reasons, such as drying up in years of low rainfall or becoming overgrown, particularly with the Mediterranean

Reed (*Arundo donax*), new sites were found and included in the monitoring. Between 1 January and 31 December 2019, a total of 7877 site visits had been made. Here, we use these records to estimate the country-wide flight period, following the standardised procedure in Boudot and Kalkman (2015) for European Odonata. To investigate seasonal patterns in the flight period of *I. elegans* across localities, we used the sites where monthly data is available for at least 36 months (up until 25 June 2019). Both phenology and morph frequency surveys were conducted during the activity hours of *I. elegans* between 08.00 and 15.00 avoiding periods of rain and strong winds.

Statistical analysis

We used generalised mixed-effect models fitted by MCMC to estimate the frequencies and mating rates of the three female morphs of *I. elegans* in Cyprus. The analyses were conducted using the package *MCMCglmm* (Hadfield, 2010) in R (R Core Team, 2018). To estimate island-wide morph frequencies, we

used a multinomial model accounting for the random effects of the sampling locality during each mating season. The residual variance (not identified in a multinomial model) was fixed to 1 and the residual covariance was set to $1/3 * (I + J)$, where I and J are two-dimensional identity and unit matrices (Hadfield, 2018). This means that we start with *a priori* belief that every sampled female has a similar probability of being of any of the three morphs. We used parameter expanded priors on the morph-specific between-locality variances to improve mixing of the variance components (Hadfield, 2018). These priors come from a non-central scaled F-distribution, and we used a scale of 25 following Gelman (2006).

To estimate morph-specific mating rates we used another categorical model with mating as a binary (0 or 1) response variable and morph as the fixed predictor. Here, we also fixed the residual variance to 1, but there was no prior on the residual covariance because the binary response can be reduced to a single dimension. Unlike the previous model, we allowed for different and independent variance components for the three sampling seasons (2017–2019) in each locality. This way, we accommodated variation in mating rates due to factors such as weather conditions, which could vary between sampling seasons and in different ways in different localities. Also, we did not specify a random categorical interaction and instead assumed that population and season effects on mating rates were similar across female morphs. Finally, we used a normally distributed fixed effect prior with mean equal to 0 and standard deviation equal to $1 + \pi^{2/3}$, as this prior is relatively flat on the probability scale when using a logit link (Hadfield, 2018).

We present the results of analyses including only those localities in which at least 10 females were sampled. For estimation of morph frequencies, we included females of both developmental colour phases, whereas for the estimation of mating rates we considered only females in their final, sexually mature colour phase. Both models were run for 2 250 000 iterations thinning every 1000th and with a burn-in of 250 000. We report posterior means (PM) and 95% highest posterior density interval (HPD) of parameter estimates. As significance tests of mating rate differences between morphs, we report the proportion of the posterior estimates in which the female morph with the lower mean mating rate has a higher mating rate than the other morph. We consider a difference in mating rate significant if this proportion was below 5%. We performed MCMC diagnostics for both morph-frequency and mating rate analyses using the R package *coda* (Plummer *et al.*, 2006). We visually evaluated the stationarity of posterior distributions and convergence of independent runs. We also evaluated convergence using the Gelman-Rubin diagnostic and report effective sample sizes and autocorrelation between posterior samples for fixed effects (Figures S1–S2).

We estimated sex- and morph-specific body size (total length, abdomen length and hind wing length) and compared it to the range reported for Europe and the U.K. (Dijkstra & Lewington, 2006). We used a Gaussian model for each size measurement, also including random effects of the different localities and seasons sampled. In these models, we used a normal prior with mean of 0 and large variance ($10e+8$) on the size estimates and inverse Wishart priors with a low degree of belief on the

residual and random variance components. The MCMC sampling strategy for these models was the same as for the morph frequency and mating rate analyses. MCMC performance was also diagnosed as mentioned above (Figure S3). As significance tests, we report the proportion of the posterior distribution of body size estimates that fall within the European/U.K. range, for males and for each of the female morphs. We considered a difference in size significant if this proportion was below 5%.

We determined the flight period of *I. elegans* in Cyprus from the number of records (2793 in total) entered in the CDSG database for the period from January 2013 to December 2019. We followed the protocol used in Boudot and Kalkman (2015), except we grouped records half monthly (i.e. 1st to 15th and 16th to 28th–31st of each month) rather than in 10-day periods to reflect the monitoring schedule. Since the earliest and latest sightings often refer to unusual events, we define the start and end of the flight season as the first half month in which one or 99 percent of the records respectively have been made (Boudot & Kalkman, 2015). Here, a record is defined as an observation of *I. elegans* during sampling by the CDSG, independently of the number of individuals observed. Boudot and Kalkman (2015) normally define the main flight season as the first and last half month period in which 10 percent or more of the total records occur. However, *I. elegans* has a long flight season in Cyprus and the 10 percent level is not achieved in any half month. Thus, we lowered the threshold for the main flight season to five percent of the total records, as Boudot and Kalkman (2015) did with other species with a long flight season in Europe.

To investigate patterns of seasonality in the flight period of *I. elegans* in Cyprus, we focused on the sites with the most complete data over the period between January 2013 and June 2019, as well as the island-wide pattern. We selected localities with data for at least 36 months, and combined two pairs of such localities in very close proximity ('Morphou area 1' and 'Morphou area 2' into 'Morphou area' and 'Diarizos River 1' and 'Diarizos River 2' into Diarizos River 1–2, $n = 8$). However, as population surveys are conducted by volunteers, there were periods of missing data in all localities (Figure S4). A total of 750 sampling events were conducted by the CDSG in these core localities during the study period. For each site, we obtained the average number of adults recorded per observer per sampling per month. We interpolated missing values in these time series using the *na.approx* function in the R package *zoo* (Zeileis & Grothendieck, 2005). We then obtained three measures of seasonality for each site and for the island-wide data using the package *greenbrown* (Forkel *et al.*, 2013; Forkel *et al.*, 2015). Here, the 'pgram' test for seasonality is positive if the maximum frequency of a periodogram based on a fast fourier transformation equals 1 (i.e. one period every year). A positive 'acf' test indicates a minimum autocorrelation in a de-trended time series at a lag of 0.5 (i.e. the least correlation between surveys is observed at six-month intervals). Finally, the 'lm' test fits two linear models to the time series, one including just the trend and one including the trend and seasonal cycle. The test is positive if the second model is selected based on the Bayesian information criterion (BIC), that is, if inclusion of the seasonal cycle parameter substantially increases model fit. In all cases, these tests of seasonality yielded identical results if applied to either

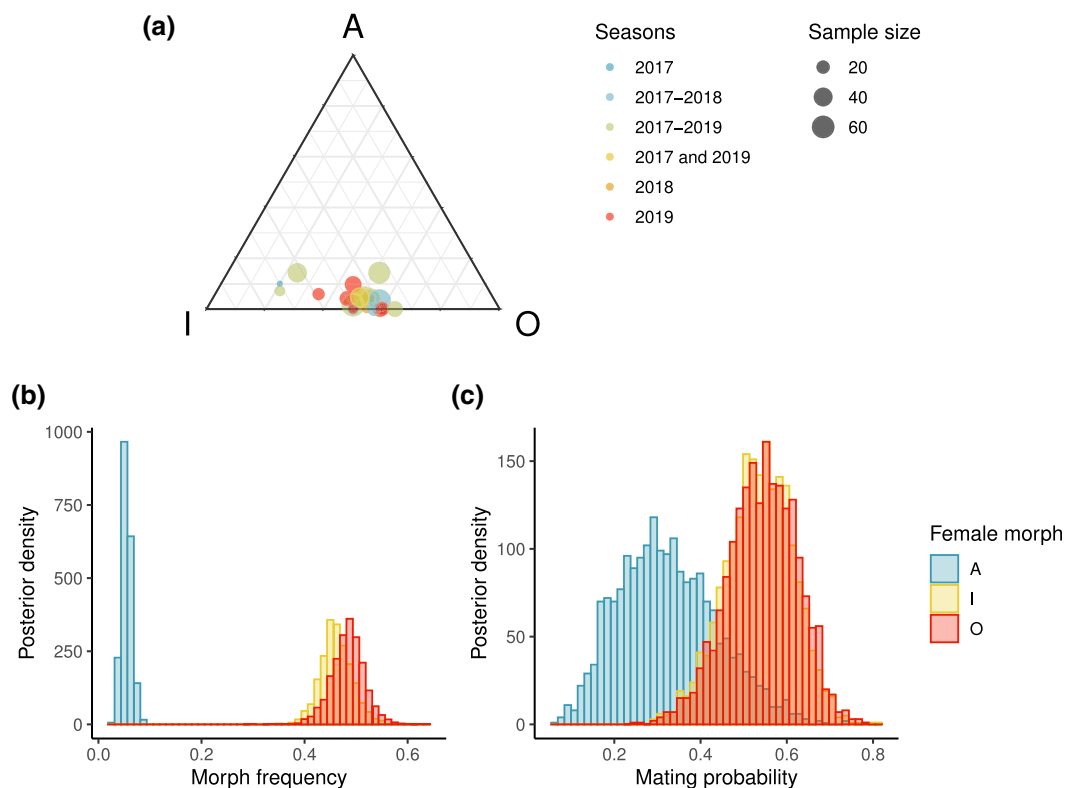


Fig. 2. Morph frequencies and mating rates of females of *I. elegans* in Cyprus. (a) Ternary plot showing observed female-morph frequencies at 19 localities between 2017–2019. Each circle corresponds to a locality sampled in 1–3 seasons. The size of the circle is proportional to female sample size [mean = 32.58; SD = 19.78; range (10, 72)], and the colour shows the mating season(s) in which each locality was surveyed. The vertices of the triangle represent hypothetical populations, fixed for a single female morph, whereas the centre of the triangle would correspond to a population with perfectly even morph frequencies. (b) Posterior distributions of island-wide morph frequency estimates. The location of the histograms along the *x*-axis indicates the estimated frequency of each morph across the sampled localities, whereas the width of the curves represents uncertainty on the estimated frequencies. Note that the posterior densities of I- and O-female frequencies are almost entirely overlapped. (c) Posterior distributions of morph-specific mating probabilities across Cyprus. Mating probability increases towards the right along the *x*-axis, and the width of the histograms represents uncertainty around each estimate. [Colour figure can be viewed at wileyonlinelibrary.com].

the original time series with missing values or the interpolated time series.

Results

Female-morph frequencies and mating rates

We visited a total of 45 localities, of which 19 had high enough female density to sample 10 or more females over the course of up to three mating seasons (Tables S1 and S2). In these populations, we sampled a total of 1682 individuals of which 619 were females. A-females were rare, if at all observed in these localities (Fig. 2a,b). The island-wide PM frequency of A-females was only 5.45% [95% HPD (0.038, 0.074)], whereas I-females [PM = 0.462; 95% HPD (0.398, 0.523)] and O-females [PM = 0.484; 95% HPD (0.419, 0.550)] occurred at similarly high frequencies (Fig. 2a,b). This pattern was mirrored across individual localities (Figure S5). The two common female morphs also had higher mating probabilities than A-females (Table 1; Fig. 2c). The PM mating probability

Table 1. Statistical contrasts of mating rates between female morphs of *I. elegans* in Cyprus. ‘PM’ represents the posterior mean of the difference in mating rate between a pair of morphs, ‘95% HPD interval’ is the 95% highest posterior density interval of the difference estimate, and ‘PMCMC’ is proportion of the posterior distribution in which the female morph with the lower mean mating rate has a higher mating rate than the other morph.

Contrast	PM	95% HPD interval	PMCMC
I versus A	0.216	[−0.009, 0.423]	0.038
O versus A	0.224	[−0.004, 0.433]	0.032
I versus O	−0.008	[−0.111, 0.125]	0.450

was 0.535 [95% HPD (0.371, 0.679)] for I-females, 0.544 [95% HPD (0.384, 0.696)] for O-females and 0.325 [95% HPD (0.122, 0.549)] for A-females.

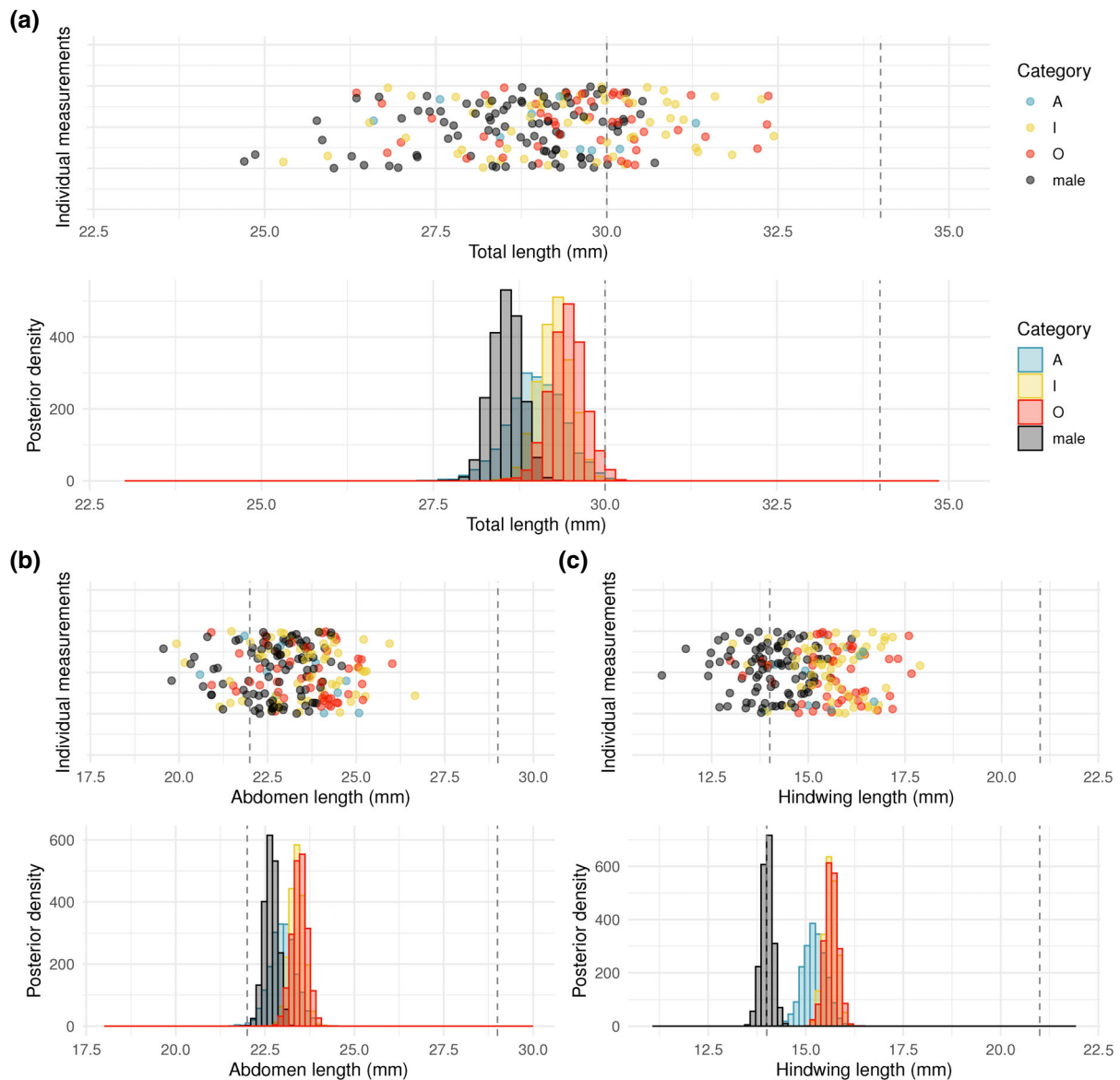


Fig. 3. Body size estimates of *I. elegans* in Cyprus and in comparison to the ranges reported in the field guide to the Odonata of Britain and Europe (Dijkstra & Lewington, 2006). For each body-size metric, the upper panel shows the individual data collected in this study, and the panel below shows the posterior distribution of size estimates from a Gaussian linear model (see Methods). The dashed grey lines indicate the size ranges reported for *I. elegans* in Britain and Europe (Dijkstra & Lewington, 2006). (a) Total length, (b) abdomen length, (c) hind wing length. [Colour figure can be viewed at [wileyonlinelibrary.com](https://onlinelibrary.wiley.com).]

Body size

We obtained body size data for 204 individuals of *I. elegans* in Cyprus ($N_{\text{males}} = 90$; $N_{\text{A-females}} = 10$; $N_{\text{I-females}} = 57$; $N_{\text{O-females}} = 47$). Both males and females in Cyprus were smaller in total length than the minimum reported for Europe and the U.K. (all PMCMC < 0.02 ; Fig. 3a). I- and O-females were both larger than males (both PMCMC ≤ 0.001 ; Fig. 3a) and similar to each other (PMCMC = 0.192; Fig. 3a), whereas A-females were of intermediate size, overlapping in total length with males, I-females and O-females (all PMCMC > 0.05 ;

Fig. 3a). Despite this overall difference in size, the abdomen length of *I. elegans* in Cyprus was within the known European/U.K. range, although within the lower half of the range (Fig. 3b). Male abdomens were also shorter than those of I- and O-females (both PMCMC < 0.001 ; Fig. 3b), while A-females were intermediate (all PMCMC > 0.05 ; Fig. 3b). Females of all morphs had longer wings than males (all PMCMC < 0.001), and were within the reported European/U.K. range (Fig. 3c). In contrast, the posterior distribution of male hind wing lengths was centred around the range minimum for Europe and the U.K. (PMCMC = 0.538; Fig. 3c).

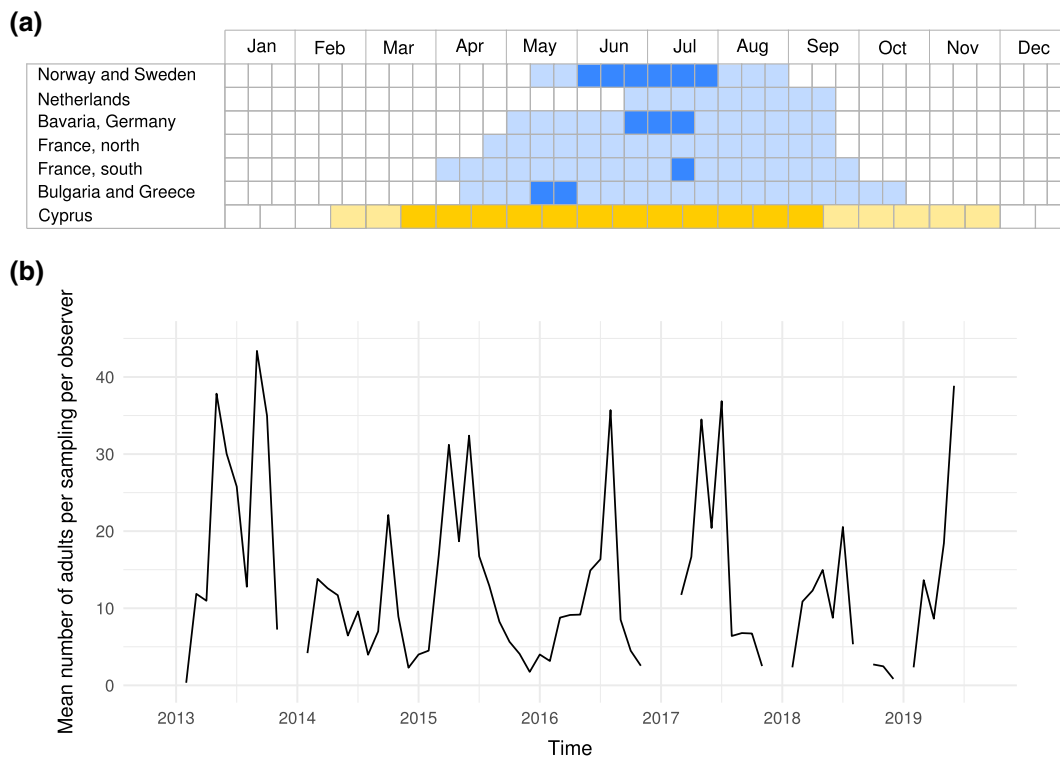


Fig. 4. Flight season and variation in flight activity of *I. elegans* in Cyprus. (a) Flight season (yellow) and main flight season (darker yellow) estimated for *I. elegans* in Cyprus, following Boudot and Kalkman (2015) and compared with more northerly countries given in the European Atlas (Boudot & Kalkman, 2015; flight season in light blue, main flight season in darker blue). (b) Mean number of adult individuals of *I. elegans*, per observer per sampling per month recorded by the CDSG across Cyprus between January 2013 and June 2019. [Colour figure can be viewed at wileyonlinelibrary.com].

Flight season

During the period between January 2013 and December 2019 members of the CDSG had made a total of 7877 site visits to around 703 localities island-wide. *I. elegans* was recorded during 2788 of these visits with an adult count of 56169 individuals. *I. elegans* is by far the most common zygopteran on Cyprus and has been recorded from 216 localities island-wide (Fig. 1d), from sea level up to the Prodomos reservoir at 1600 m.a.s.l., which is the highest water-body monitored by the CDSG.

The flight season of *I. elegans* in Cyprus starts in the second half of February and continues to the second half of November (Fig. 4a). The main flight season is from the second half of March to the first half of September (Fig. 4a). This is substantially longer than that reported for the more northerly regions (Fig. 4a). There was evidence of seasonality in the island-wide occurrence data according to the periodicity ('pgram') and autocorrelation ('acf') criteria, but not according to the linear model ('lm') criterion (Table 2, Fig. 4b), which compares the fit of models with and without a seasonal cycle via BIC. Evidence for seasonality varied among localities, with some localities showing a seasonal pattern under periodicity or both periodicity and autocorrelation criteria (Table 2). However, the 'lm' criterion of seasonality was not met in any locality with long-term data (Table 2).

Discussion

I. elegans is a geographically widely distributed damselfly species that occurs across the Palearctic (Boudot & Salamun, 2015) and has become a well-known study system in evolutionary ecology. Population studies on *I. elegans* have addressed a variety of topics, including the age-, frequency- and density-dependent dynamics of male-mating harassment (Svensson *et al.*, 2005; Gosden & Svensson, 2009; Van Gossium *et al.*, 2011; Willink *et al.*, 2019) and the maintenance of heritable morphs by balancing selection (Takahashi *et al.*, 2014; Le Rouzic *et al.*, 2015). Population studies with *I. elegans* have also been used to investigate phenotypic correlations between alternative mechanisms of defence against parasites (Willink & Svensson, 2017), and the role of environmental and social effects on temperature sensitivity and range expansion (Lancaster *et al.*, 2017). These studies rely on female colour morphs as visual genetic markers of suites of correlated traits on which natural and sexual selection operate. However, the majority of such field studies have taken place near the northern limit of the distribution range of *I. elegans*. How these phenomena are influenced by large-scale environmental variation is therefore an important question, which could be addressed with comparative studies across the geographic range of *I. elegans*. As a first step on the road towards such a broader geographic scope of studies of *I. elegans*, here we present data

Table 2. Seasonality in the flight activity of *I. elegans* across eight localities and for the island-wide data. Seasonality was tested according to three criteria (see Methods): maximum frequency of a periodogram ('pgram'), the minimum autocorrelation of a de-trended time series ('acf'), and a model comparison with and without the seasonal component ('lm'). 1 = seasonality detected; 0 = seasonality not detected.

Locality	Latitude °N	Longitude °E	Elevation	Seasonality test		
				'pgram'	'acf'	'lm'
Diarizos river 1 below Prastio	34.7858	32.6859	220	1	1	0
Diarizos river 5 near Kidasi	34.7862	32.6880	234	1	1	0
Ezousa river below Episkopi	34.8119	32.5289	162	1	0	0
Koineli	35.2336	33.3022	162	1	1	0
Kanli	35.238	33.2620	174	1	0	0
Mavrokolympo: feeder stream	34.8599	32.4180	139	0	0	0
Morphou area: reservoir	35.2545	32.9831	48	0	0	0
Zakaki: marsh	34.6413	32.9898	1	1	0	0
Island-wide	–	–	–	1	1	0

on the basic population biology and phenology of this trimorphic species in Cyprus. This is the southernmost region where populations of this widespread species have been systematically studied to date.

Three main features distinguish breeding populations in Cyprus from those in Northern Europe. Firstly, male-mimicking A-females are the minority morph in Cyprus, occurring only at ~5% frequency, while I- and O-females occur at similarly high frequencies (Fig. 2a,b). Such a low frequency of A-females is striking, given that this may be the most visually conspicuous morph to human observers. A-females display a bright blue colouration throughout their adult life, which may increase their detectability compared to the other morphs, which develop a darker and duller colour pattern during sexual maturation (Henze *et al.*, 2019; Willink *et al.*, 2020). In contrast, the frequency of A-females increases with latitude, with females in Southern Sweden typically being composed of 60–80% male mimics (Gosden *et al.*, 2011; Le Rouzic *et al.*, 2015). The increasing frequency of A-females in northern Europe is likely due to their developmental advantages at cooler temperatures, whereby A-females enjoy higher pre-reproductive survival and faster sexual maturation and colour development (Svensson *et al.*, 2020). The developmental success of I- and O-females after emergence from the last nymphal stage is in contrast more sensitive to temperature, and these two morphs are therefore expected to benefit more from the warmer Mediterranean climate of Cyprus (Svensson *et al.*, 2020).

The low island-wide frequency of A-females might imply that some populations in Cyprus are effectively dimorphic, with only I- and O-females. In fact, a previous survey across continental Europe suggested that about 10% of populations were dimorphic, although they all included A-females (Gosden *et al.*, 2011). The loss of a female morph may cause increased pre-mating and mating harassment in the other two morphs, as males would have fewer competing targets while forming a search image of potential mates (Dukas & Kamil, 2001). However, if male-mimicry is effective, the local absence of otherwise rare A-females should not dramatically alter male-mating harassment in the other two morphs, which would already account for most male-mating attempts under negative frequency-dependent selection. The

local absence of A-females might also be temporary. In *I. elegans*, the A-allele is dominant over I- and O-alleles. While dominant alleles under negative frequency-dependent selection are more likely to be lost by genetic drift, they also re-invade populations more easily due to Haldane's Sieve, the expectation that a weakly advantageous mutation will increase more rapidly in frequency if dominant (Haldane, 1924; Pannell *et al.*, 2005). Haldane's Sieve acting on migrants might thus contribute to the maintenance of the dominant A-allele at a regional scale (Roux & Pannell, 2019).

As in Cyprus, A-females are usually not the majority morph in the south of continental Europe, but typically I-females are more common than O-females (Gosden *et al.*, 2011; Svensson *et al.*, 2020). In the closely related species *Ischnura genei*, populations in the Mediterranean island of Sardinia also have relatively low frequencies of A-females, but O-females are generally more common than I-females (Sanmartín-Villar & Cordero-Rivera, 2016). Although several ecological differences between A- and both I- and O-females have previously been reported for *I. elegans* (Lancaster *et al.*, 2017; Willink & Svensson, 2017; Svensson *et al.*, 2020), ecological differences between I- and O-morphs have not been investigated to a similar extent, probably because O-females are so rare in northern Europe. The role of ecological mechanisms versus historical contingencies and 'island effects' that might shape morph-frequency variation within the Mediterranean region therefore remains an interesting question that should be addressed in the future, to get a better understanding of the ecological factors and evolutionary processes operating in this trimorphic system.

Secondly, adult individuals of *I. elegans* are relatively small in Cyprus (Fig. 3). This is particularly the case for males, which are also generally smaller than females (Abbott & Gosden, 2009), and when compared to the reported European range of total body length and hind wing length (Dijkstra & Lewington, 2006; Fig. 3b,c). This qualitative result is consistent with Bergman's rule, a pattern of increasing body size with decreasing temperature. Bergman's rule is supported in vertebrate endotherms (Ashton *et al.*, 2000; Meiri & Dayan, 2003; Salewski & Watt, 2017; but see Riemer *et al.*, 2018), but is not generally

supported across diverse clades of insects and other ectotherms (Mousseau, 1997; Blanckenhorn & Demont, 2004; Adams & Church, 2008; Shelomi, 2012; Wonglersak *et al.*, 2020). One mechanistic explanation for Bergman's rule that is applicable for *I. elegans*, and odonates in general, is that developmental rate (i.e. cell division and differentiation) increases more rapidly with temperature than does metabolism (Blanckenhorn & Demont, 2004). Therefore, higher developmental temperature at more southern latitudes should result in faster maturation at a smaller body size. Because growth in odonates occurs only during the aquatic larval stage, differences in developmental temperature should affect both the duration of the growing period (i.e. voltinism) and the adult body size (Johansson, 2003; De Block *et al.*, 2008; Hassall *et al.*, 2014). Our study did not address whether Bergman's rule is met throughout the distribution range of *I. elegans*. However, a decreasing number of generations per year at higher latitudes has been previously reported (Corbet *et al.*, 2006), and the extended flight season of *I. elegans* in Cyprus (see below) also suggests that populations on this island are multivoltine.

A-females in Cyprus were more male-like in size than either I- or O-females. This is consistent with previous analyses in Swedish populations, showing that A-females are more male-like in shape than the other morphs (Abbott & Gosden, 2009). Recent studies in female-polymorphic insects, including the widespread tropical and subtropical *I. senegalensis* (a congener of *I. elegans* that has been intensively studied in Japan), suggest that the development of male-coloured females is more masculinised, compared to the development of alternative female morphs (Takahashi *et al.*, 2019). These developmental differences between morphs may be caused by alternative splicing and differential expression patterns of the regulatory gene *doublesex* (Takahashi *et al.*, 2019, 2020), which also underlies the development of somatic sex differences across many insect taxa (Kopp, 2012). To date, there is no direct evidence of a male-like expression pattern of *dsx* in A-females of *I. elegans*. However, the locus or set of tightly linked loci that govern colour morph development in *I. elegans* seems to have pleiotropic effects during colour development and differentiation of the female morphs (Willink *et al.*, 2020). Such widespread pleiotropy may also impact the rate and duration of larval development, in turn generating size differences between morphs. In southern Sweden, the larval developmental period is shorter in the offspring of O-females, suggesting this allele is associated with a faster developmental rate (Abbott & Svensson, 2005). In contrast, the A-allele may be associated with a slower growth rate, as A-females have a similar developmental period as I-females but become mature at a smaller size (Abbott & Svensson, 2005). Whether these developmental differences occur at warmer temperatures has not been investigated.

Finally, the flight season, during which adult damselflies emerge and can potentially mate, is considerably longer in Cyprus than in any other location with comparable data (Fig. 4a; Boudot & Salamun, 2015). This is consistent with overall fast development and multivoltinism caused by warm to mild temperatures throughout the year. Flight activity in Cyprus spanned more than 9 months (Fig. 4a), resulting in partial evidence for a

seasonal pattern in the island-wide dataset (Table 2; Fig. 4b). Some core localities with long-term data, had even less support for seasonality in flight activity (Table 2). Variation in seasonality among localities could be explained by regional differences in the length of growth periods and breeding seasons, for instance due to an altitudinal temperature gradient. Although a test of this hypothesis would entail estimating seasonal strength from a larger number of localities with long-term data, such a parallel between latitudinal and altitudinal gradients in the seasonality of (potential) mating activity is known for other ectotherms (Morrison & Hero, 2003). Nevertheless, the marked contrast between the flight season of *I. elegans* in Cyprus and more Northern European sites (Fig. 4a) suggests a pervasive role of temperature influencing life-history evolution in this widely distributed damselfly.

In conclusion, species with broad distribution ranges, such as *I. elegans*, provide excellent opportunities to investigate how large-scale climatic variation shapes the phenotypic outcomes of selection driven by local interspecific and intraspecific interactions. However, for many widespread species, local field studies tend to closely match the geographic distribution of the scientists who study them, which in turn reflects economic factors and research traditions, rather than strict biological considerations, and can lead to biases in the perception of which ecological factors are most important (Zuk, 2016). Here, we have studied the population biology of the widely distributed damselfly *I. elegans* in Cyprus, which is the southern range limit of this species in Europe. Populations of *I. elegans* in Cyprus are distinguished by the rarity of male-mimicking females, reduced body size, and a long flight season. These ecological differences from northern Europe, where the majority of field studies have been conducted, underscore the importance of broadening the geographic scope of field studies in *I. elegans* and many other widespread organisms.

Acknowledgements

Many thanks go to past and present members of the CDSG for their dedicated monitoring and submission of records that made the phenology analysis possible. Jean-Pierre Boudot is thanked for agreeing to the use of the flight season data from Boudot and Kalkman (2015) and for useful discussion on the flight season of the *Ischnura* genus. Damla Beton and Kuşkor are thanked for assistance at site nos 4, 5, 10, 12, 13, 14, 16, and 17. We thank three anonymous reviewers for their comments that improved our manuscript. The authors declare that they have no conflicts of interest.

Author contribution

D.J.S., R.S., and B.W. conceived this study. D.J.S., R.S., R.B., and B.W. collected the data. B.W. and D.J.S. conducted the analyses. B.W. wrote the paper with input from E.I.S., D.J.S., R.S., and R.B.

Data availability statement

The data that support the findings of this study and all R code to reproduce the analyses are available online at <https://doi.org/10.6084/m9.figshare.13119980.v2>.

Supporting Information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Figure S1. Traces for two independent runs of a model estimating female morph frequencies of the Common Bluetail damselfly (*Ischnura elegans*) in Cyprus. Traces are shown for two morphs as the third frequency can be obtained from the other two. Autocorrelation between posterior samples is <0.05 , and effective sample sizes = 2000 in both cases. The multivariate potential scale reduction factor (mgsrf) of the Gelman-Rubin convergence diagnostic is 1.01.

Figure S2. Traces for two independent runs of a model estimating female morph-specific mating rates in the Common Bluetail damselfly (*Ischnura elegans*) in Cyprus. Autocorrelation between posterior samples is <0.05 , and effective sample sizes >1845 in all cases. The multivariate potential scale reduction factor (mgsrf) of the Gelman-Rubin convergence diagnostic is 1.01.

Figure S3. Traces for two independent runs of models estimating body size in the Common Bluetail damselfly (*Ischnura elegans*) in Cyprus. Autocorrelation between posterior samples is <0.07 , and effective sample sizes >1612 in all cases. The multivariate potential scale reduction factor (mgsrf) of the Gelman-Rubin convergence diagnostic is 1.01 in all three models.

Figure S4. Mean number of adults of *Ischnura elegans* recorded per observer per sampling per month, in 8 core populations by the Cyprus Dragonfly Study Group between January 2013 and June 2019.

Table S1. Locality information and sampling dates for 19 sites in Cyprus, where female morph frequencies of *Ischnura elegans* were recorded from 2017 to 2019.

Table S2. Number of females of three colour morphs of the damselfly *Ischnura elegans*, in 19 localities sampled in Cyprus between 2017–2019.

Figure S5. Posterior distributions of morph frequency estimates across populations of *I. elegans* in Cyprus. The location of the histograms along the x -axis indicates the estimated frequency of each morph in each of the sampled populations, whereas the width of the curves represents uncertainty on the estimated frequencies.

References

Abbott, J.K. & Gosden, T.P. (2009) Correlated morphological and colour differences among females of the damselfly *Ischnura elegans*. *Ecological Entomology*, **34**, 378–386.

- Abbott, J. & Svensson, E.I. (2005) Phenotypic and genetic variation in emergence and development time of a trimorphic damselfly. *Journal of Evolutionary Biology*, **18**, 1464–1470.
- Adams, D.C. & Church, J.O. (2008) Amphibians do not follow Bergmann's rule. *Evolution*, **62**, 413–420.
- Anholt, B.R., Vorburger, C. & Knaus, P. (2001) Mark-recapture estimates of daily survival rates of two damselflies (*Coenagrion puella* and *Ischnura elegans*). *Canadian Journal of Zoology*, **79**, 895–899.
- Ashton, K.G., Tracy, M.C. & Queiroz, A.D. (2000) Is Bergmann's rule valid for mammals? *American Naturalist*, **156**, 390–415.
- Azevedo, R.B.R., James, A.C., McCabe, J. & Partridge, L. (1998) Latitudinal variation of wing: thorax size ratio and wing-aspect ratio in *Drosophila melanogaster*. *Evolution*, **52**, 1353–1362.
- Blanckenhorn, W. & Demont, M. (2004) Bergmann and converse Bergmann latitudinal clines in arthropods: two ends of a continuum? *Integrative and Comparative Biology*, **44**, 413–424.
- Boudot, J. P. & Kalkman, V. J. (eds) (2015) *Atlas of the European Dragonflies and Damselflies*. KNNV Publishing, The Netherlands.
- Boudot, J.P. & Salamun, A. (2015) *Ischnura elegans* (Vander Linden, 1820). *Atlas of the European Dragonflies and Damselflies* (ed. by J. P. Boudot and V. J. Kalkman). KNNV Publishing, The Netherlands.
- Bridle, J.R. & Vines, T.H. (2007) Limits to evolution at range margins: when and why does adaptation fail? *Trends in Ecology & Evolution*, **22**, 140–147.
- Conover, D. (1992) Seasonality and the scheduling of life history at different latitudes. *Journal of Fish Biology*, **41**, 161–178.
- Corbet, P.S. (1999) *Dragonflies: Behavior and Ecology of Odonata*. Harley, Colchester, U.K.
- Corbet, P.S., Suhling, F. & Soendgerath, D. (2006) Voltinism of Odonata: a review. *International Journal of Odonatology*, **9**, 1–44.
- Cordero, A., Carbone, S.S. & Utzeri, C. (1998) Mating opportunities and mating costs are reduced in androchrome female damselflies, *Ischnura elegans* (Odonata). *Animal Behaviour*, **55**, 185–197.
- Cornes, R.C., van der Schrier, G., van den Besselaar, E.J.M. & Jones, P.D. (2018) An ensemble version of the E-OBS temperature and precipitation data sets. *Journal of Geophysical Research-Atmospheres*, **123**, 9391–9409. <https://doi.org/10.1029/2017JD028200>.
- De Block, M., Slos, S., Johansson, F. & Stoks, R. (2008) Integrating life history and physiology to understand latitudinal size variation in a damselfly. *Ecography*, **31**, 115–123.
- De Frenne, P., Graae, B.J., Rodriguez-Sánchez, F., Kolb, A., Chabrierie, O., Decocq, G. *et al.* (2013) Latitudinal gradients as natural laboratories to infer species' responses to temperature. *Journal of Ecology*, **101**, 784–795.
- Dijkstra, K.D.B. & Lewington, R. (2006) *Field Guide to the Dragonflies of Britain and Europe*. British Wildlife Publishing, Gillingham, U.K.
- Dudaniec, R.Y., Yong, C.J., Lancaster, L.T., Svensson, E.I. & Hansson, B. (2018) Signatures of local adaptation along environmental gradients in a range-expanding damselfly (*Ischnura elegans*). *Molecular Ecology*, **27**, 2576–2593.
- Dukas, R. & Kamil, A.C. (2001) Limited attention: the constraint underlying search image. *Behavioral Ecology*, **12**, 192–199.
- Fabian, D.K., Kapun, M., Nolte, V., Kofler, R., Schmidt, P.S., Schlötterer, C. *et al.* (2012) Genome-wide patterns of latitudinal differentiation among populations of *Drosophila melanogaster* from North America. *Molecular Ecology*, **21**, 4748–4769.
- Forkel, M., Carvalhais, N., Verbesselt, J., Mahecha, M.D., Neigh, C.S. & Reichstein, M. (2013) Trend change detection in NDVI time series: effects of inter-annual variability and methodology. *Remote Sensing*, **5**, 2113–2144.
- Forkel, M., Migliavacca, M., Thonicke, K., Reichstein, M., Schaphoff, S., Weber, U. *et al.* (2015) Codominant water control on global interannual variability and trends in land surface phenology and greenness. *Global Change Biology*, **21**, 3414–3435.

- Forsman, A. & Shine, R. (1995) The adaptive significance of colour pattern polymorphism in the Australian scincid lizard *Lampropholis delicata*. *Biological Journal of the Linnean Society*, **55**, 273–291.
- Gelman, A. (2006) Prior distributions for variance parameters in hierarchical models. *Bayesian Analysis*, **1**, 515–534.
- Gosden, T.P. & Svensson, E.I. (2007) Female sexual polymorphism and fecundity consequences of male mating harassment in the wild. *PLoS One*, **2**, e580.
- Gosden, T.P. & Svensson, E.I. (2009) Density-dependent male mating harassment, female resistance, and male mimicry. *American Naturalist*, **173**, 709–721.
- Gosden, T.P., Stoks, R. & Svensson, E.I. (2011) Range limits, large-scale biogeographic variation, and localized evolutionary dynamics in a polymorphic damselfly. *Biological Journal of the Linnean Society*, **102**, 775–785.
- Hadfield, J.D. (2010) MCMC methods for multi-response generalized linear mixed models: the MCMCglmm R package. *Journal of Statistical Software*, **33**, 1–22.
- Hadfield, J. (2018) *MCMCglmm course notes*. URL <http://cran.r-project.org/web/packages/MCMCglmm/vignettes/CourseNotes.pdf> [accessed on March 1, 2019].
- Haldane, J.B.S. (1924) A mathematical theory of natural and artificial selection, part I. *Proceedings of the Cambridge Philosophical Society*, **23**, 19–41.
- Hammers, M. & Van Gossum, H. (2008) Variation in female morph frequencies and mating frequencies: random, frequency-dependent harassment or male mimicry? *Animal Behaviour*, **76**, 1403–1410.
- Harabiš, F. & Doln, A. (2012) Human altered ecosystems: suitable habitats as well as ecological traps for dragonflies (Odonata): the matter of scale. *Journal of Insect Conservation*, **16**, 121–130.
- Hassall, C., Keat, S., Thompson, D.J. & Watts, P.C. (2014) Bergmann's rule is maintained during a rapid range expansion in a damselfly. *Global Change Biology*, **20**, 475–482.
- Henze, M.J., Lind, O., Wilts, B.D. & Kelber, A. (2019) Pterin-pigmented nanospheres create the colours of the polymorphic damselfly *Ischnura elegans*. *Journal of the Royal Society Interface*, **16**, 20180785.
- Hinnekin, B.O.N. (1987) Population dynamics of *Ischnura E. elegans* (Vander Linden)(Insecta: Odonata) with special reference to morphological colour changes, female polymorphism, multiannual cycles and their influence on behaviour. *Hydrobiologia*, **146**, 3–31.
- Iserbyt, A., Bots, J., Van Dongen, S., Ting, J.J., Van Gossum, H. & Sherratt, T.N. (2011) Frequency-dependent variation in mimetic fidelity in an intraspecific mimicry system. *Proceedings of the Royal Society of London B: Biological Sciences*, **278**, 20110126.
- Johansson, F. (2003) Latitudinal shifts in body size of *Enallagma cyathigerum* (Odonata). *Journal of Biogeography*, **30**, 29–34.
- Kirkpatrick, M. & Barton, N.H. (1997) Evolution of a Species' Range. *American Naturalist*, **150**, 1–23.
- Kopp, A. (2012) Dmrt genes in the development and evolution of sexual dimorphism. *Trends in Genetics*, **28**, 175–184.
- Lancaster, L.T., Dudaniec, R.Y., Chauhan, P., Wellenreuther, M., Svensson, E.I. & Hansson, B. (2016) Gene expression under thermal stress varies across a geographical range expansion front. *Molecular Ecology*, **25**, 1141–1156.
- Lancaster, L.T., Dudaniec, R.Y., Hansson, B. & Svensson, E.I. (2017) Do group dynamics affect colour morph clines during a range shift? *Journal of Evolutionary Biology*, **30**, 728–737.
- Le Rouzic, A., Hansen, T.F., Gosden, T.P. & Svensson, E.I. (2015) Evolutionary time-series analysis reveals the signature of frequency-dependent selection on a female mating polymorphism. *American Naturalist*, **185**, E182–E196.
- Lenormand, T. (2002) Gene flow and the limits to natural selection. *Trends in Ecology & Evolution*, **17**, 183–189.
- Machado, G., Buzatto, B.A., Garcia-Hernández, S. & Macías-Ordóñez, R. (2016) Macroecology of sexual selection: a predictive conceptual framework for large-scale variation in reproductive traits. *American Naturalist*, **188**, S8–S27.
- Mazaris, A.D., Kallimanis, A.S., Pantis, J.D. & Hays, G.C. (2013) Phenological response of sea turtles to environmental variation across a species' northern range. *Proceedings of the Royal Society of London B: Biological Sciences*, **280**, 20122397.
- Meiri, S. & Dayan, T. (2003) On the validity of Bergmann's rule. *Journal of Biogeography*, **30**, 331–351.
- Morrison, C. & Hero, J.M. (2003) Geographic variation in life-history characteristics of amphibians: a review. *Journal of Animal Ecology*, **72**, 270–279.
- Mousseau, T.A. (1997) Ectotherms follow the converse to Bergmann's rule. *Evolution*, **51**, 630–632.
- Palo, J.U., O'Hara, R.B., Laugen, A.T., Laurila, A., Primmer, C.R. & Merilä, J. (2003) Latitudinal divergence of common frog (*Rana temporaria*) life history traits by natural selection: evidence from a comparison of molecular and quantitative genetic data. *Molecular Ecology*, **12**, 1963–1978.
- Pannell, J.R., Dorken, M.E. & Eppley, S.M. (2005) Haldane's Sieve in a metapopulation: sifting through plant reproductive polymorphisms. *Trends in Ecology & Evolution*, **20**, 374–379.
- Parr, M.J. (1965) A population study of a colony of imaginal *Ischnura elegans* (Van der Linden)(Odonata: Coenagrionidae) at Dale, Pembrokeshire. *Field Studies*, **2**, 237–282.
- Parr, M.J. (1970) The life histories of *Ischnura elegans* (van der Linden) and *Coenagrion puella* (L.)(Odonata) in south Lancashire. *Proceedings of the Royal Entomological Society of London. Series A, General Entomology*, **45**, 172–181.
- Parr, M.J. (1973) Ecological studies of *Ischnura elegans* (Vander Linden)(Zygoptera: Coenagrionidae). I. Age groups, emergence patterns and numbers. *Odonatologica*, **2**, 139–157.
- Parr, M.J. & Palmer, M. (1971) The sex ratios, mating frequencies and mating expectancies of three coenagrionids (Odonata: Zygoptera) in northern England. *Insect Systematics & Evolution*, **2**, 191–204.
- Parr, M.J. & Palmer, M. (1972) Survival rates, population density and predation in the damselfly, *Ischnura elegans* (Vander Linden)(Zygoptera: Coenagrionidae). *Odonatologica*, **1**, 137–141.
- Parsons, S.M. & Joern, A. (2014) Life history traits associated with body size covary along a latitudinal gradient in a generalist grasshopper. *Oecologia*, **174**, 379–391.
- Phifer-Rixey, M., Heckman, M., Trussell, G.C. & Schmidt, P.S. (2008) Maintenance of clinal variation for shell colour phenotype in the flat periwinkle *Littorina obtusata*. *Journal of Evolutionary Biology*, **21**, 966–978.
- Plummer, M., Best, N., Cowles, K. & Vines, K. (2006) CODA: convergence diagnosis and output analysis for MCMC. *R News*, **6**, 7–11.
- R Core Team (2018) *R: A Language and Environment for Statistical Computing*. R Foundation for Statistical Computing, Vienna, Austria.
- Riener, K., Guralnick, R.P. & White, E.P. (2018) No general relationship between mass and temperature in endothermic species. *eLife*, **7**, e27166.
- Robertson, H.M. (1985) Female dimorphism and mating behaviour in a damselfly, *Ischnura ramburi*: females mimicking males. *Animal Behavior*, **33**, 805–809.
- Roux, C. & Pannell, J.R. (2019) The opposing effects of genetic drift and Haldane's sieve on floral-morph frequencies in tristylous metapopulations. *New Phytologist*, **224**, 1229–1240.
- Salewski, V. & Watt, C. (2017) Bergmann's rule: a biophysiological rule examined in birds. *Oikos*, **126**, 161–172. <https://doi.org/10.1111/oik.03698>.

- Sánchez-Guillén, R.A., Van Gossum, H. & Cordero Rivera, A. (2005) Hybridization and the inheritance of female color polymorphism in two ischnurid damselflies (Odonata: Coenagrionidae). *Biological Journal of the Linnean Society*, **85**, 471–481.
- Sanmartín-Villar, I. & Cordero-Rivera, A. (2016) The inheritance of female colour polymorphism in *Ischnura genei* (Zygoptera: Coenagrionidae), with observations on melanism under laboratory conditions. *PeerJ*, **4**, e2380.
- Schmidt, P.S. & Paaby, A.B. (2008) Reproductive diapause and life-history clines in North American populations of *Drosophila melanogaster*. *Evolution*, **62**, 1204–1215.
- Shelomi, M. (2012) Where are we now? Bergmann's rule sensu lato in insects. *American Naturalist*, **180**, 511–519.
- Slatkin, M. (1987) Gene flow and the geographic structure of natural populations. *Science*, **236**, 787–792.
- Svensson, E.I., Abbott, J. & Härdling, R. (2005) Female polymorphism, frequency dependence, and rapid evolutionary dynamics in natural populations. *American Naturalist*, **165**, 567–576.
- Svensson, E.I., Abbott, J.K., Gosden, T.P. & Coreau, A. (2009) Female polymorphisms, sexual conflict and limits to speciation processes in animals. *Evolutionary Ecology*, **23**, 93–108.
- Svensson, E.I., Willink, B., Duryea, M.C. & Lancaster, L.T. (2020) Temperature drives pre-reproductive selection and shapes the biogeography of a female polymorphism. *Ecology Letters*, **23**, 149–159.
- Takahashi, Y., Morita, S., Yoshimura, J. & Watanabe, M. (2011) A geographic cline induced by negative frequency-dependent selection. *BMC Evolutionary Biology*, **11**, 256.
- Takahashi, Y., Kagawa, K., Svensson, E.I. & Kawata, M. (2014) Evolution of increased phenotypic diversity enhances population performance by reducing sexual harassment in damselflies. *Nature Communications*, **5**, 4468.
- Takahashi, M., Takahashi, Y. & Kawata, M. (2019) Candidate genes associated with color morphs of female-limited polymorphisms of the damselfly *Ischnura senegalensis*. *Heredity*, **122**, 81–92.
- Takahashi, M., Okude, G., Futahashi, R., Takahashi, Y., Kawata, M. (2020) Involvement of the *doublesex* gene in body color masculinization of the blue-tailed damselfly, *Ischnura senegalensis*. *BioRxiv*: 036715.
- Van Gossum, H., Stoks, R., Matthysen, E., Valck, F. & De Bruyn, L. (1999) Male choice for female colour morphs in *Ischnura elegans* (Odonata, Coenagrionidae): testing the hypotheses. *Animal Behaviour*, **57**, 1229–1232.
- Van Gossum, H., Stoks, R. & De Bruyn, L. (2001) Frequency-dependent male mate harassment and intra-specific variation in its avoidance by females of the damselfly *Ischnura elegans*. *Behavioral Ecology and Sociobiology*, **51**, 69–75.
- Van Gossum, H., Bots, J., Van Heusden, J., Hammers, M., Huyghe, K. & Morehouse, N.I. (2011) Reflectance spectra and mating patterns support intraspecific mimicry in the colour polymorphic damselfly *Ischnura elegans*. *Evolutionary Ecology*, **25**, 139–154.
- Van Noordwijk, M. (1978) A mark-recapture study of coexisting zygopteran populations. *Odonatologica*, **7**, 353–374.
- Waller, J.T. & Svensson, E.I. (2017) Body size evolution in an old insect order: no evidence for Cope's Rule in spite of fitness benefits of large size. *Evolution*, **71**, 2178–2193.
- Willink, B. & Svensson, E.I. (2017) Intra- and intersexual differences in parasite resistance and female fitness tolerance in a polymorphic insect. *Proceedings of the Royal Society of London B: Biological Sciences*, **284**, 20162407.
- Willink, B., Duryea, M.C. & Svensson, E.I. (2019) Macroevolutionary origin and adaptive function of a polymorphic female signal involved in sexual conflict. *American Naturalist*, **194**, 707–724.
- Willink, B., Duryea, M.C., Wheat, C. & Svensson, E.I. (2020) Gene expression changes during female reproductive development in a colour polymorphic insect. *Evolution*, **74**, 1063–1081.
- Wonglersak, R., Fenberg, P.B., Langdon, P.G., Brooks, S.J. & Price, B.W. (2020) Temperature-body size responses in insects: a case study of British Odonata. *Ecological Entomology*, **45**, 795–805. <https://doi.org/10.1111/een.12853>.
- Zeileis, A. & Grothendieck, G. (2005) zoo: S3 infrastructure for regular and irregular time series. *Journal of Statistical Software*, **14**, 1–27.
- Zuk, M. (2016) Temperate assumptions: how where we work influences how we think. *American Naturalist*, **188**, S1–S7.

Accepted 22 November 2020

First published online 4 January 2021

Associate Editor: Christopher Hassall