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Life-history traits and climatic responsiveness in noctuid moths 0

Zsolt Végvári, Edit Juhász, János Pál Tóth, Zoltán Barta, Sándor Boldogh, Sándor Szabó and 5 Zoltán Varga

Z. Végvári (vegvari@hnp.hu), Dept of Conservation Zoology, Hortobágy National Park Directorate, Univ. of Debrecen, HU-4024 Debrecen Sumen u.2, Hungary. – E. Juhász, S. Szabó and Z. Varga, Dept of Evolutionary Zoology and Human Biology, Univ. of Debrecen, HU-4032 Debrecen Egyetem tér 1., Hungary. – J. Pál Tóth, Res. Inst. for Viticulture and Oenology, Tarcal, Könyves Kálmán Street 54., Tarcal, HU-3915, Hungary. – Z. Barta, MTA-DE "Lendület" Behavioural Ecology Research Group, Dept of Evolutionary Zoology, Univ. of Debrecen, Hu-4032 Debrecen Egyetem tér 1., Hungary. – S. Boldogh, Aggtelek National Park Directorate, HU-3758 Jósvafő, Tengerszem oldal. 1., Hungary.

Emergence phenology has been shown to advance considerably in the past decades in many lepidopterans. Noctuid moths 15 (Noctuidae) constitute a species-rich family of lepidopterans with a large diversity of life history traits presumably driving climatic responsiveness. In our study we aim to assess the role of life-history and ecological traits in climatic responsiveness of noctuid moths, whilst controlling for phylogenetic dependence. We used a long-term dataset of European noctuid moths collected from a light-trap in northeastern Hungary. As the study site is located at the intersection of several biogeographical zones harbouring a large number of noctuid moth species, our dataset provides a unique possibility to investigate 20 the moths' climatic sensitivity. To estimate the role of life-history traits and ecological factors in driving lepidopterans' response to climatic trends, we employed three proxies related to the species' ecology (habitat affinity, food plant specialization and food type) and two robust types of life-history traits (migration strategy and hibernation form). The degree of temporal shifts of various measures of emergence phenology was related to hibernation stage, food type and migration strategy. Large-scale phylogenetic relatedness exerted little constraint in all models fitted on each measure of phenology. Our results imply that noctuid moths overwintering as adults exhibited greater degrees of phenological shifts than species 25 hibernating as larvae or pupae. It implies that moths hibernating as adults are forced to suspend activity in our climate and the prolongation of autumn activity might be the result of increased plasticity in flight periods.

- 30 During the past decades an increasing number of studies have reported on the effects of global climatic change on ecosystem structure and functioning (Parry et al. 2007, Walther 2010, Burrows et al. 2011). Climatic processes influence phenology, population dynamics and life-history traits
- 35 in a broad range of organisms, including a large number of endangered taxa (Parmesan 2006, Forrest and Miller-Rushing 2010, Jenouvrier and Visser 2011). Specifically, evidence is accumulating that temporal shifts in insect phenology as a response to climatic alteration has accelerated over
- 40 the past few decades (Singer and Parmesan 2010, Hodgson et al. 2011). For instance, emergence phenology has been shown to advance considerably in the past decades in many
- lepidopterans (Stefanescu et al. 2003, Hekkinen et al. 2010, [AQ3] Diamond et al. 2011). Besides, studies have recently detected
 - 45 that the annual number of generations of night-active moths has changed over the last decades (Altermatt 2010a, Kocsis and Hufnagel 2011, Gimesi et al. 2012). Global warming is assumed to be one of the most probable drivers of the increasing number of generations presumably by allowing
 - 50 more time for activity (Altermatt 2010a). Nocturnal moths have also been found to respond strongly to global warming
 - 52 by advancing spring flight periods but an immense number

of species respond in a varied way to climatic changes in this taxa (Hodgson et al. 2011, Valtonen et al. 2011).

85 European noctuid moths constitute a representative taxonomic group for studying the effects of climatic trends on insect phenology, as they represent one of the most species-rich insect families subdivided into a large number of subfamilies and tribes characterised by a high diversity of life 90 history, possibly related to the observed variation in climatic responsiveness (Karsholt and Razowski 1996, references in Supplementary material Appendix 1). A specific explanation for this large variation in climatic responsiveness may be that the ability of lepidopterans to track food plant phenology is 95 assumed to be constrained by a number of ecological and life history factors (van Asch and Visser 2007, Altermatt 2010b, Breed et al. 2013). As a consequence, declining population sizes in butterflies (and also in moths) are often attributed to spatial mismatches between their distribution area and 100 changing climatic envelopes, leading to regional or total extinctions of entire subspecies (Devictor et al. 2012).

The developmental stage in which individuals hibernate has already been shown to affect climatic responsiveness. For instance, species overwintering as adults were shown to exhibit higher degrees of phenological flexibility than

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0 lepidopterans hibernating as larvae or pupae, as these taxa can emerge immediately after hibernation (Diamond et al. 2011).

Recent studies have detected that climatic warming seems to enhance the range of migratory movements of some moth

species, as migrant populations might track climatic trends faster owing to their large mobility (Sparks et al. 2007).

Evidence is accumulating that climatic responsiveness shows contrasting regional patterns across Europe, both in the frequency of supplementary generations and temporal

10 patterns of emergence phenology of lepidopterans (Tobin et al. 2008, Alternatt 2010a, Pöyry et al. 2011, Kharouba et al. 2014).

In our study we aim to assess the role of life-history and ecological traits in climatic responsiveness in a species-rich

15 lepidopteran family (Noctuidae) with a control for phylogenetic dependence. To accomplish this objective, we used a long-term dataset of European noctuid moths collected by a light-trap in Jósvafő (Aggtelek karst, northeastern Hungary, 48.5°N, 20.51°E, 287 m a.s.l), eastern Europe, where such

20 investigations are still scarce.

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Although a recent study has shown that many life-history traits might have important roles in climatic responsiveness in lepidopterans (Radchuk et al. 2013), we aim to assess only those which emerged as substantial predictors of climatic

- 25 responsiveness in lepidopterans. Accordingly, we employed three proxies related to the species' ecology (habitat affinity, characterised as habitat specialist or generalist; food plant specialization and food type, defined as herbaceous or woody) and two types of life-history traits (migration strategy, and hibernation form) already shown to affect the climatic
- sensitivity of lepidopterans (Forrest and Miller-Rushing 2010, Diamond et al. 2011).

Specifically, our hypotheses are as follows: 1) species with broader habitat range would be less responsive to climatic

- 35 change, as these noctuids are less constrained by climateinduced changes in habitat phenology. 2) Food-generalist noctuid moths would exhibit smaller degrees of advancement in emergence phenology (as estimated by capture dates) than specialist species as they are less dependent on the
- 40 phenology of one or a few specific food plants. 3) Migrant moths would track climatic variability to a greater degree than resident ones, as they have more developed sensitivity to weather conditions promoting flexible onsets of northward migration (Sparks et al. 2007, Alerstam et al. 2011).
- 45 4) Finally, noctuids which hibernate as adults might track climatic trends faster than species hibernating as larvae or pupae as these moths can emerge immediately after finishing hibernation (Nyman et al. 2012).
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Material and methods

Study area

- 55 The moth fauna of the Aggtelek karst area has been intensively surveyed over the last decades of the last century (Varga 1999, Szabó et al. 2007). We collected data during a long term light trap survey. The Jermy-type light trap was established in 1988 on the northeast-facing slope of the Tohonya Valley in Aggtelek National Park, northeast
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61 Hungary (48.1°N, 20.51°E, 287 m). The trap equipped with a 125 W mercury bulb was activated in 1988 and operated in each year for a 20-year period until 2007 between March and November, spanning 4788 days in total. The study area 65 provides a unique opportunity to investigate the moths' climatic sensitivity, as this part of the Carpathian basin is located in the intersection of several biogeographical zones harbouring 407 noctuid moth species, an unusually high number in central Europe (Varga 1999, 2012). Furthermore, 70 the study area is surrounded by a broad variety of habitats hosting a set of noctuid taxa highly representative for the region. The light trap was located in an abandoned orchard, presently used as a semi-dry haymeadow, surrounded by scattered rests of karstic oak-hornbeam Waldsteinio-Carpinetum 75 and xerothermic oak forests (Corno-Quercetum pubescentis) and hazel shrub Coryletum avellanae communities (Szabó et al. 2007).

Climate data

Weather data were obtained from the nearest meteorological station at Miskolc (48°1N, 20°4E), 60 km from the light-trap. Out of a numerous set of climatic variables, daily mean temperature and daily precipitation were available for the studied period. To assess temporal patterns in climatic 85 predictors relevant for lepidopteran biology, we calculated monthly and seasonal (March to May for spring, June to August for summer, September to November for autumn and December to February for winter) means and variances of daily average temperature and monthly as well as seasonal 90 precipitation sums and variances. Further, we computed temporal patterns in cumulative temperature, a biologically more relevant predictor of climatic responsiveness than temperature itself (Pöyry et al. 2011). The computation went as follows: we set a temperature limit of 5.0°C, and summed 95 daily average temperature values above this limit for the following periods in each year,: 1) 1 January to 31 March, 2) 1 January to 31 April, 3) 1 January 1 to 31 May, 4) 1 March to 31 May. To detect temporal trends in weather proxies, we fitted linear regressions on these variables as a function 100 of years.

Noctuid data

In each year the light trap was operated from the first days of 105 March until the first days of November (with a small, 1–2 day variation per year due to accidentally unfavourable weather conditions), totalling 4788 days (mean \pm SD = 239.4 \pm 12 days per year). Captured specimens were collected on a daily basis and were stored in cotton layers or pinned until 110 identification. Species-level identification was carried out by SSz and ZV and the data were entered into a database by recording date, taxon and number of individuals.

During the study period 128 045 individuals of 417 noctuid taxa were collected. To acquire robust measures on climatic responsiveness, we used only the data of species with records from at least ten years spanning for at least 15 years, yielding 200 species in total. In the following step we retained only species with more than twenty individuals caught in total in every year, to be able to obtain reliable estimates of the annual population development. As a result, we used the 121

- 0 records of 70 species in subsequent analyses (filtered species hereafter; Supplementary material Appendix 2). To test the robustness of the above filtering conditions, we removed noctuids with less than ten distinct capture dates per year from the set of 200 species and repeated all analyses.
- 5 To test the effects of first and final operation dates of the light trap, we calculated the distribution of capture dates of filtered species. This shows that the percentage of first and last capture dates within seven days from first and last operation dates, respectively, were less than 0.1%. Therefore it is
- 10 unlikely that between-year variation in first and final operation dates can influence phenological patterns.

To derive standard measures of climatic responsiveness for each species, we calculated slopes of linear regressions fitted on years using the following estimates of emergence phenol-

- 15 ogy as response variables: 1) first, 2) median and 3) last julian dates (day-of-year number) of capture and 4) activity length (defined as the number of days between first and last capture dates), calculated using corresponding capture dates. We refer to these response variables as climatic responsiveness
- 20 of first, median and last julian days as well as activity length. High slopes from these regressions indicate that a given species can respond faster to climatic changes. The linear regression slopes were then applied in phylogenetic least squares models as response variables to investigate which
- 25 ecological and life history traits influence climatic responsiveness. To test the robustness of the emergence predictors, we applied the approach proposed by van Strien et al. (2008) to our dataset. Accordingly, we calculated 1) the date of the 25th individual captured (FA25) and 2) the date of the first
- 30 25% of all observed individuals in the entire flight period (FP25) for each species in the filtered dataset. In the next step we calculated Pearson's correlation coefficients medium capture dates and only FP25 which revealed significant relationships (r = 0.73, p < 0.03) between medium capture
- 35 dates and those proposed by van Strien et al. (2008).

Ecological and life-history predictors

We obtained data on ecological variables and life-history 40 traits from the series of Noctuidae Europae (Ronkay et al. 2001, Hacker et al. 2002, Fibiger et al. 2009, 2010). Habitat affinity was classified as specialist or generalist. Feeding strategy was characterized by diet specialization (categorised as food specialist or food generalist) and food 45 type (classified according to the food type of caterpillars:

- 45 type (classified according to the food type of caterpinals, herbaceous or woody). Migration strategy was categorised as resident and migratory. Hibernation type was classified as imago, pupal and larval stages (Varga et al. 2005).
- All used data on the species' life history and ecology are 50 presented in the Supplementary material Appendix 2.

Phylogenetic and statistical analyses

In the first step we constructed a composite tree (Supplementary material Appendix 3) using the data of the most recent molecular survey applying gradual branch lengths (Mitchell et al. 2006, Zahiri et al. 2011). To account for the lack of knowledge of species-level tree topology, we included only subfamily-level information provided by the most recent molecular phylogecontinuous (Mitchell et al. 2007, Zahiri et al. 2011)

60 netic analyses (Mitchell et al. 2006, Zahiri et al. 2011).

To evaluate the degree of association between climatic 61 responsiveness measures and ecological and life-history predictors, we fitted phylogenetic generalized least squares (PGLS) models controlling for the non-independence among taxa by incorporating a variance-covariance matrix 65 that represents their phylogenetic relationships (Martins and Hansen 1999, Pagel 1997, 1999). The degree of phylogenetic dependence (λ) was set to the most appropriate value obtained for each model by likelihood ratio statistics (Freckleton et al. 2002). In the following step we calcu-70 lated the strength of phylogenetic relatedness in each trait separately by quantifying the maximum-likelihood value of λ with its statistical significance estimated by likelihood ratio tests (i.e. by comparing the log-likelihood of the model 75 in which λ is set to the most appropriate value to the loglikelihood of a model in which $\lambda = 0$). As recent studies on the phylogeny of Noctuidae have suggested a number of major changes in the structure of the phylogenetic tree (references provided in the Supplementary material Appen-80 dix 4), we repeated assessing phylogenetic signal in response and predictor variables by formulating linear mixed models that include only genus nested in subfamily as a random term, and compared model performance by ANOVA tests.

After model fitting, the relative importance of life-history traits and ecological variables was calculated using informa-85 tion-theoretic model-comparison (Burnham and Anderson 2002). First we obtained the values of Akaike's information criterion corrected for small sample size (AIC_c) and the corresponding Akaike weight of each model (ω). Second, we selected models with substantial support ($\Delta_i = AIC_i$) 90 - AIC_{min} < 2.0) and calculated model-averaged parameter estimates (β) and unconditional standard errors (SE_u; Burnham and Anderson 2002) of each variable by the sums of their Akaike weights across all models with substantial 95 support containing the given predictor. Akaike differences in the range 0-2 indicate substantial level of empirical support of a given model, whereas $\Delta_i > 7$ implies very weak support (Burnham and Anderson 2002).

For all analyses we used the R statistical computing environment including its 'ape' (Paradis et al. 2011), 'geiger' 100 (Harmon et al. 2009), 'MuMIn' (Barton 2011) and 'nlme' (Pinheiro et al. 2009) packages.

Results

Climatic patterns

During the 20 years of data collection monthly mean temperature has significantly increased only in June ($b = 0.140^{\circ}C$ year⁻¹, $F_{1,17} = 7.292$, p = 0.015, Fig. 1). Seasonal mean and cumulative temperature has not shown significant trends. Further, we detected no clear temporal patterns in variance of monthly temperature and all precipitation metrics (results not shown). 115

Temporal trends in capture data

Noctuids showed a considerable amount of variation in slopes of various estimates of emergence phenology as a function of years. Out of the 70 taxa in our study, 38 shifted 121

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Figure 1. June mean temperature as a function of years during the study.

25 their first capture to earlier dates with four of them exhibiting significant advance. Median capture dates, indicative of the climatic responsiveness of the bulk of the population, showed similar patterns with 34 noctuids emerging earlier but the trend was only significant for a single species. Last julian dates of annual records have advanced in 27 taxa,

four of which were significant (Supplementary material Appendix 2, Fig. 2).

In contrast, first capture of 32 noctuids shifted to later dates with only a single species showing significant patterns. Median capture date was detected to change to later dates in

35 Whethan capture date was detected to change to later dates in
 36 noctuids including three species with significant trends.
 Last capture dates of 43 noctuids have shifted later, five of



Figure 2. Number of species shifting emergence phenology to earlier and later dates.

Table 1. λ -statistics for phylogenetic signal in the investigated 61 response variables and predictors. $\lambda = 0$ indicates no phylogenetic signal.

Variable	λ	р	
Slope of first capture date	0.000	0.000	0.999
Slope of median capture date	0.000	0.000	0.999
Slope of last capture date	0.000	0.000	0.999
Slope of activity length	0.000	0.000	0.997
Migration type	0.000	0.000	0.999
Hibernation type	0.000	0.000	0.422
Food specialisation	0.543	0.898	0.343
Food type	0.811	2.249	0.134
Habitat affinity	0	0.001	0.98

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which exhibited significant temporal patterns. Although activity length shortened in 28 species with significant trends in three species, it became longer in 42 noctuids including significant relationships for nine taxa (Supplementary material Appendix 2).

On average, noctuids shifted their first capture dates by 4.705 (SE = 0.902) days to later dates after 1998 than before this year across all species considered. Similarly, median capture occurred 4.525 (SE = 2.210) days later in the second period than in the first one. In contrast, overall last emergence shifted by 7.547 (SE = 1.325) days to later dates, representing a delay almost twice as long as the delay at the beginning of the flight season. As a result, activity length increased by 2.842 (SE = 1.151) days calculated over the 90 filtered set of species.

Ecological and life history correlates of phenological change

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Phylogenetic signal (λ) was zero for all measures of all response and proxy variables (Table 1). Quantitatively similar results were obtained by linear mixed models which showed no significant effect of genus nested within subfamily (p = 0.067).

Responsiveness of first capture dates was significantly affected by food type and migration strategy – parameter estimates of which had confident intervals not overlapping with zero. Noctuids with larvae feeding primarily on woods are shifting first emergence to later dates while species feeding on herbaceous plants advance first capture dates ($\Sigma = 0.826$; Table 2, 3; Supplementary material Appendix 5).

Our results show that migratory strategy has emerged as the second most important proxy of climatic responsiveness measured as first capture dates based on its Akaike sums in PGLS-models fitted on first capture dates ($\Sigma = 0.749$; Table 2, 3; Supplementary material Appendix 5): resident noctuids advanced climatic responsiveness measured as first capture dates to a greater degree than migratory species. Parameter estimates for all other variables have confidence intervals overlapping with zero, therefore we consider their effects as non-significant.

Considering last emergence phenology hibernation stage has emerged with the highest Akaike scores calculated over supported models fitted on the slope of last capture dates ($\Sigma = 1.0$, Table 2). Hibernation stage was the only predictor 121

0	Table 2. Akaike sums, model-averaged parameter estimates (β), standard errors (SEu) and 95% confidence intervals for predictors across best	61
	supported models ($\Delta_i < 2.0$).	

	Slope of first emergence date	Slope of median emergence date	Slope of last emergence date	Slope of activity length	
5	Food type $\Sigma = 0.826$ $\beta(SE_u) = -0.395 (0.145)$	Migration strategy $\Sigma = 0.483$ $\beta(SE_u) = -0.497 (0.575)$	Hibernation stage $\Sigma = 1.0$ $\beta(SE_u) = -2.674 (0.543)$	Hibernation stage $\Sigma = 1.0$ $\beta(SE_u) = -3.018 (0.608)$	65
10	95% Cl = $(-0.678; -0.111)$ Migration strategy $\Sigma = 0.749$ $\beta(SE_{u}) = -0.547 (0.243)$ 95% Cl = $(-1.023; -0.07)$ Habitat affinity $\Sigma = 0.296$	95% CI = $(-1.625; 0.631)$ Food type $\Sigma = 0.404$ $\beta(SE_u) = -0.376 (0.365)$ 95% CI = $(-1.091; 0.339)$ Habitat affinity $\Sigma = 0.299$	95% CI = $(-3.737; -1.61)$ Migration strategy $\Sigma = 0.246$ $\beta(SE_u) = -0.227 (0.406)$ 95% CI = $(-1.023; 0.57)$ Food type $\Sigma = 0.209$	95% CI = $(-4.21; -1.826)$ Migration strategy $\Sigma = 0.282$ $\beta(SE_u) = 0.265 (0.467)$ 95% CI = $(-0.651; 1.18)$ Habitat affinity $\Sigma = 0.213$	70
15	$\beta(SE_u) = 0.252 (0.15)$ 95% CI = (-0.099; 0.548) Food specialisation $\Sigma = 0.212$ $\beta(SE_u) = 0.224 (0.165)$ 95% CI = (-0.099; 0.548)	$\beta(SE_u) = 0.206 (0.36) 95\% CI = (-0.499; 0.912) Food specialisation \Sigma = 0.293 \beta(SE_u) = 0.052 (0.404) 95\% CI = (-0.74; 0.843)$	$\beta(SE_u) = -0.217 (0.301)$ 95% CI = (-0.807; 0.373) Food specialisation $\Sigma = 0.183$ $\beta(SE_u) = 0.151 (0.293)$ 95% CI = (-0.424; 0.725)	$\beta(SE_{u}) = -0.212 (0.287)$ 95% CI = (-0.774; 0.35) Food type $\Sigma = 0.195$ $\beta(SE_{u}) = 0.099 (0.339)$ 95% CI = (-0.564; 0.763)	75
20	Hibernation stage $\Sigma = 0.151$ $\beta(SE_u) = 0.386 (0.303)$ 95% CI = (-0.208; 0.981)	Hibernation stage $\Sigma = 0.278$ $\beta(SE_u) = 0.097 (0.755)$ 95% CI = (-1.384; 1.577)	Habitat affinity $\Sigma = 0.142$ $\beta(SE_u) = -0.018 (0.252)$ 95% CI = (-0.512; 0.475)	Food specialisation $\Sigma = 0.186$ $\beta(SE_u) = -0.06 (0.331)$ 95% CI = (-0.708; 0.588)	80

of last capture dates and activity length in the best supported models: species hibernating as imagos shifted last capture to

²⁵ later dates and prolonged activity length to a greater degree than noctuids hibernating as larvae or pupae (Table 2, Fig. 3, 4; Supplementary material Appendix 7).

We obtained qualitatively similar results for species with more than ten capture dates per year (Supplementary material Appendix 6).

Discussion

- ³⁵ In this study we aimed to 1) describe the extent of phenological responses presumably caused by ongoing climate change in a species-rich group of noctuid moths and 2) investigate the possible ecological and life history correlates of these responses. We believe that our results are based on
- 40 a unique combination of methodological elements. First, our dataset was collected over 20 years applying highly standardised collection methodology. Second, the single collection site allows high-level replication of species-specific climatic responsiveness of large number of noctuid taxa

allowing multispecies analyses of 70 species after rigorous filtering. Third, we calculated multiple phenological estimates in order to avoid collection bias primarily affecting first emergence phenology.

Considering noctuid phenology, we detected no clear patterns in changes of emergence phenology, which might be 90 related to the diversity in a number of life history traits of the studied species, as shown by recent investigations (Altermatt [AQ2] 2010a or b????). Additionally, complex trends of emergence phenology can also be a result of our finding that significant trends in climatic patterns were only found for June tem-95 perature, which is in line with a number of recent studies showing no clear temperature trends after 1990 (reviewed by Stocker et al. 2013). Although Hungary has experienced considerable increase of annual mean temperature over the past century (Molnár V. et al. 2012), our study covered only 100 two decades and was located in an area where the foothills of the Carpathian Mountains might have had a buffering effect.

Our findings show that the degree of temporal shifts of various measures of emerging phenology was primarily related to hibernation stage, food type and migration strategy.

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Table 3. Supported models ($\Delta_i\!<\!2.0)$ fitted on all types of response v	ariables.
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	Response	Model formula	DF	AICc	Δ_{i}	W	
	Slope of first emergence date	food type $+$ migration strategy	4	123.811	0	0.279	11(
50		food type + habitat affinity + migration strategy	5	124.904	1.092	0.161	11
50		food type	3	125.321	1.509	0.131	
	Slope of median emergence date	migration strategy	3	246.815	0	0.123	
	1 0	food type	3	247.018	0.203	0.111	
		food type $+$ migration strategy	4	247.695	0.88	0.079	
		hibernation stage	4	247.858	1.043	0.073	11°_{11}
55		food specialisation	3	247.881	1.067	0.072	
		habitat affinity	3	247.996	1.181	0.068	
		habitat affinity + migration strategy	4	248.586	1.771	0.051	
	Slope of last emergence date	hibernation stage	4	203.804	0	0.395	
	Slope of activity length	hibernation stage	4	219.033	0	0.362	
60		hibernation stage + migration strategy	5	220.947	1.914	0.139	12



Figure 3. Shift of last capture date as a function of hibernation stage in noctuid moths.

- 25 Specifically, hibernation stage is the strongest predictor of climatic responsiveness measured as last capture dates and activity length in noctuid moths, similarly to the results of Diamond et al. (2011). This might indicate that noctuid moths overwintering as adults exhibited greater degrees 30 of phenological shifts than species hibernating as larvae or pupae. This finding extends the results of a recent study
- reporting that species with more advanced overwintering stages have experienced relatively greater advances in their date of first appearance (Diamond et al. 2011). However, the difference between the reactions of the bulk of the popula-
- tion and last active individuals to climatic trends seems to be



Figure 4. Shift of activity length as a function of hibernation stage 60 in noctuid moths.

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different in our case. Since noctuids hibernating as adults in 61 our database showed no shifts in median emergence to earlier dates, the last individuals of these moths have prolonged capture dates over the study period in comparison to species hibernating in larval or pupal form. Prolonged autumn 65 activity may allow longer time for replenishing fat reserves before hibernation which can increase survival chances and improve reproductive success. Moreover, all of these species are winter-active moths in the southern part of their 70 distribution areas implying that noctuid taxa hibernating as imagos are forced to suspend activity in our climate and the prolonged autumn activities of the last emerging individuals might be the result of enhanced temporal plasticity of the timing of autumn emergence (László et al. 2000).

75 Food type was included in best supported models for climatic responsiveness measured as first capture dates, similarly to previous investigations: noctuids feeding on herbaceous plants advanced first capture dates to a greater degree than species connected to woody materials (Diamond et al. 2011, Altermatt 2010a or b????). Although a previous study showed that herbaceous plants were more likely than [AQ2] woody plants to advance phenology (Walther et al. 2002), our results imply that moths feeding on herbaceous plants are able to track plant phenology faster than noctuids feed-85 ing on woody plants which evokes the following hypotheses. First, as herbaceous plants are shorter-lived on average than woody taxa, their herbivores are forced to synchronise phenology to a greater degree than moth larvae feeding on woody plants. Second, woody plants are on average larger 90 than herbaceous species implying higher levels of phenological variance of plant parts consumed by herbivores in individual trees which allows for a wider range of phenological synchronisation. However, the complexity of climatic responses driven by trophic interactions is enhanced by the fact that the highly diverse family of Noctuidae is linked to 95 a great variety of larval food plants (references in Supplementary material Appendix 1). For instance, the majority of these species is connected to forested habitats and majority of spring- and autumn-active moths are linked to woody food plants (details shown in Supplementary material Appendix 100 2; references in Supplementary material Appendix 1). A large group of genera (Supplementary material Appendix 2) use a wide variety of food plants which are most often herbaceous (grasses for mostly monovoltine Apamea, and often Urticaceae for the phenologically more plastic Plusiinae), or 105 partly herbaceous and woody (Acronictinae) (Supplementary material Appendix 2).

Migration strategy has emerged as a supported predictor of climatic responsiveness. Migrant noctuids have shifted first captures to later dates than resident moths which implies that migratory species arriving from southern regions may detect warming spring weather later than locally hibernating taxa (Alerstam et al. 2011). Indeed, migrant moths need time to reach central Europe and thus are outcompeted in advancements of climatic responsiveness measured as first capture dates by locally wintering moths hibernating as imagos.

Throughout our analyses we detected no phylogenetic relatedness on the subfamily level in climatic responsiveness of noctuid moths. As the existing politomies are largely unresolved, this implies that the absence of phylogenetic signal in climatic responsiveness of noctuid moths 121

- 0 can be considered as the result of poor resolution of clades rather than missing phylogenetic relationships. Since the majority of noctuid taxa in our dataset belongs to the tribe Hadenini previously merged into the subfamily Noctuinae by Lafontaine and Schimdt (2011), for which recent
- 5 investigations on evolutionary relatedness do not provide better phylogenetic resolutions, we were not able to detect phylogenetic signal on finer scales (Zahiri et al. 2013). This warrants further research on phylogenetic relatedness among noctuid moths to be able to detect if climatic
- 10 responsiveness is independent of phylogenetic relatedness in noctuid species as shown by a number of investigations showing that phenological shifts are not driven by common descent in a broad range of taxa (Végvári et al. 2010, Diamond et al. 2011). However, the lack of phylogenetic
- 15 signal in emergence phenology of the species in our dataset can also be an outcome of phenotypic plasticity which drives phenological patterns as a response to climatic variability, suggested by the findings of recent studies across a
- [AQ4] number of taxonomic levels (Hoffmann and Sgrò 2011).
 20 The similar results of Diamond et al. (2011) showing that phylogenetic relatedness among species predicted very little variation in butterflies' phenological responses imply that climatic responsiveness in lepidopterans can be treated as a species-specific reaction rather than driven by evolutionary relatedness (Altermatt 2010a).

Our study implies recommendations for lepidopteran conservation, as a reflection to current declining trends in a number of insect species on a global scale. We found that noctuid moths hibernating as adults are able to track

30 climatic trends faster than species overwintering in less developed stages which suggest higher declining probabilities in these taxa, considering current climatic predictions (Stocker et al. 2013). Our result that resident moths are responding to ongoing climatic patterns to a greater degree

35 than migratory taxa, calls for more intensive surveys of migrating lepidopterans populations.

In summary, we have demonstrated in a diverse set of noctuid moth species that temporal trends in emergence phenology (which is associated with a measure of tempera-

40 ture increase in the study area) is primarily related to hibernation stage, diet specialisation and migration strategy.

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⁵⁰ **References**

- Alerstam, T. et al. 2011. Convergent patterns of long-distance nocturnal migration in noctuid moths and passerine birds. – Proc. R. Soc. B 278: 3074–3080.
- 55 Altermatt, F. 2010a. Climatic warming increases voltinism in European butterflies and moths. – Proc. R. Soc. B 277: 1281–1287.
 - Altermatt, F. 2010b. Tell me what you eat and I'll tell you when you fly: diet can predict phenological changes in response to climate change. – Ecol. Lett. 13: 1475–1484.

Barton, K. 2011. MuMIn: Multi-model inference. – R package ver. 61 1.0. 0.

Breed, G. A. et al. 2013. Climate-driven changes in northeastern US butterfly communities. – Nature Climate Change 3: 142–145.

Burnham, K. P. and Anderson, D. R. 2002. Model selection and multi-model inference: a practical information-theoretic approach. – Springer.

- Burrows, M. T. et al. 2011. The pace of shifting climate in marine and terrestrial ecosystems. Science 334: 652–655.
- Devictor, V. et al. 2012. Differences in the climatic debts of birds and butterflies at a continental scale. – Nature Climate Change 2: 121–124. 70
- Diamond, S. E. et al. 2011. Species' traits predict phenological responses to climate change in butterflies. Ecology 92: 1005–1012.
- Fibiger, M. et al. 2009. Noctuidae Europaeae, Pantheinae– Bryophilinae. – Entomological Press, Sorø.
- Fibiger, M. et al. 2010. Noctuidae Europaeae. Rivulinae– Phytometrinae, and Micronoctuidae, including Supplement to Noctuidae Europaeae. – Entomological Press Sorø.

Forrest, J. and Miller-Rushing, A. J. 2010. Toward a synthetic understanding of the role of phenology in ecology and evolution. – Phil. Trans. R. Soc. B 365: 3101–3112.

Freckleton, R. P. et al. 2002. Phylogenetic analysis and comparative data: a test and review of evidence. – Am. Nat. 160: 712–726.

Gimesi, L. et al. 2012. The effect of climate change on the phenology of moth abundance and diversity. – Appl. Ecol. Environ. Res. 10: 349–363.

- Hacker, H. et al. 2002. Noctuidae Europaeae, Hadeninae I. – Entomological Press, Sorø, pp. 419.
- Harmon, L. et al. 2009. geiger: analysis of evolutionary diversification. – R package ver. 1.3-1.
- Heikkinen, R. K. et al. 2010. Assessing the vulnerability of [AQ3]
 European butterflies to climate change using multiple criteria.
 Biodivers. Conserv. 19: 695–723.
- Hodgson, J. A. et al. 2011. Predicting insect phenology across space and time. – Global Change Biol. 17: 1289–1300.
- Hoffmann, A. A. and Sgró, C. M. 2011. Climate change and [AQ4] evolutionary adaptation. – Nature 470: 479–485. 95
- Jenouvrier, S. and Visser, M. E. 2011. Climate change, phenological shifts, eco-evolutionary responses and population viability: toward a unifying predictive approach. – Int. J. Biometeorol. 5: 905–919.
- Karsholt, O. and Razowski, J. 1996. The Lepidoptera of Europe: a distributional checklist. – Apollo Books.
- Kharouba, H. M. et. al. 2014. Predicting the sensitivity of butterfly phenology to temperature over the past century. – Global Change Biol. 20: 504–514.
- Lafontaine, J. D. and Schmidt, B. C. 2011. Additions and corrections to the check list of the Noctuoidea (Insecta, Lepidoptera) of North America north of Mexico. – ZooKeys 149: 145.
- Kocsis, M. and Hufnagel, L. 2011. Impacts of climate change on Lepidoptera species and communities. – Appl. Ecol. Environ. Res. 9: 43–72.
- László, G. M. et. al. 2000. On the genesis of the Himalayan-Sino Pacific Thyatiridae (Lepidoptera) fauna, with special reference to Taiwan. – In: Biodiversity across the Taiwan Strait. Natl Mus. Nat. Sci., Taichung, poster volume, 65.
- Martins, E. P. and Hansen, T. F. 1999. Phylogenies and the comparative method: a general approach to incorporating phylogenetic information into the analysis of interspecific data – Am. Nat. 153: 448–448.
- Mitchell, A. et al. 2006. Systematics and evolution of the cutworm moths (Lepidoptera : Noctuidae): evidence from two proteincoding nuclear genes. – Syst. Entomol. 31: 21–46.
- Molnár, A. et al. 2012. Pollination mode predicts phenological response to climate change in terrestrial orchids: a case study from central Europe. J. Ecol. 100: 1141–1152.

121

115

75

85

100

105

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Nyman, T. et al. 2012. Climate-driven diversity dynamics in plants 0 and plant-feeding insects. - Ecol. Lett. 15: 889-898.

Pagel, M. 1997. Inferring evolutionary processes from phylogenies. – Zool. Scripta 26: 331–348.

- Pagel, M. 1999. Inferring the historical patterns of biological evolution. - Nature 401: 877-884.
- 5 Paradis, E. et al. 2011. Package 'ape': analysis of phylogenetics and evolution. Publisher????
 - Parmesan, C. 2006. Ecological and evolutionary responses to recent climate change. - Annu. Rev. Ecol. Evol. Syst. 37: 637-669.
- Parry, M. L. et al. 2007. Climate change 2007: impacts, adaptation and vulnerability. Intergovernmental Panel on Climate Change. 10 - Cambridge Univ. Press.
 - Pinheiro, J. et al. 2009. Nlme package. Publisher????
 - Pöyry, J. et al. 2011. Climate-induced increase of moth multivoltinism in boreal regions. - Global Ecol. Biogeogr. 20: 289 - 298.
- Radchuk, V. et al. 2013. Each life stage matters: the importance of 15 assessing the response to climate change over the complete life cycle in butterflies. - J. Anim. Ecol. 82: 275-285.
 - Ronkay, L. et al. 2001. Noctuidae Europaeae. Hadeninae II. - Entomological Press, Sorø, pp. 452.
- Singer, M.C. and Parmesan, C. 2010. Phenological asynchrony 20 between herbivorous insects and their hosts: signal of climate change or pre-existing adaptive strategy? - Phil. Trans. R. Soc. B 365: 3161-3176.
 - Sparks, T. H. et al. 2007. Increased migration of Lepidoptera linked to climate change. - Eur. J. Entomol. 104: 139-143.
- Stefanescu, C. et al. 2003. Effects of climatic change on the phenol-25 ogy of butterflies in the northwest Mediterranean Basin. Global Change Biol. 9: 1494–1506.
 - Stocker, T. F. et al. 2013. Climate change 2013: the physical science basis. Working Grp I Contrib. to the 5th Assess. Rep. of the Intergovernmental Panel on Climate Change. Summary for Policymakers (IPCC, 2013). Publisher???
 - Supplementary material (available online as Appendix oik.01655 at <www.oikosjournal.org/readers/appendix>).
- 35 Appendix 1-7.

- Szabó, S. et al. 2007. Long-term light trap study on the macro-61 moth (Lepidoptera : Macroheterocera) fauna of the Aggtelek National Park. - Acta Zool. Acad. Sci. Hung. 53: 257-269.
- Tobin, P. C. et al. 2008. Historical and projected interactions between climate change and insect voltinism in a multivoltine 65 species. - Global Change Biol. 14: 951-957.
- Valtonen, A. et al. 2011. Environmental controls on the phenology of moths: predicting plasticity and constraint under climate change. - Oecologia 165: 237-248.
- van Asch, M. and Visser, M. E. 2007. Phenology of forest caterpillars and their host trees: the importance of synchrony. 70 Annu. Rev. Entomol. 52: 37–55.
- van Strien, A. J. et al. 2008. Bias in phenology assessments based on first appearance data of butterflies. - Öecologia 156: 227-235.
- Varga, Z. 1999. The Lepidoptera of the Aggtelek National Park. 75 – In: Mahunka S. (ed.), The fauna of the Aggtelek National Park II. Hung. Nat. Hist. Mus., Budapest, pp. 443-504.
- Varga, Z. 2012. Magyarország nagylepkéi Macrolepidoptera of Hungary. - Heterocera Press, Budapest.
- Varga, Z. et al. 2005. Checklist of the fauna of Hungary. Macrolepidoptera. – Hung. Nat. Hist. Mus., Budapest
- Végvári, Z. et al. 2010. Life history predicts advancement of avian spring migration in response to climate change. - Global Change Biol. 16: 1-11.

Walther, G. R. 2010. Community and ecosystem responses to recent climate change. - Phil. Trans. R. Soc. B 365: 2019-2024 [AQ1]

- Walther et al. 2002
- Zahiri, R. et al. 2011. A new molecular phylogeny offers hope for a stable family level classification of the Noctuoidea (Lepidoptera). - Zool. Scripta 40: 158-173.
- Zahiri, R. et al. 2013. Relationships among the basal lineages of 90 Noctuidae (Lepidoptera, Noctuoidea) based on eight gene regions - Zool. Scripta 42: 488-507.

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