



Life history evolution, species differences, and phenotypic plasticity in hemiparasitic eyebrights (*Euphrasia*)

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PREMISE: Species delimitation in parasitic organisms is challenging because traits used to identify species are often plastic and vary depending on the host. Here, we use species from a recent radiation of generalist hemiparasitic *Euphrasia* to investigate trait variation and trait plasticity. We tested whether *Euphrasia* species show reliable trait differences, investigated whether these differences correspond to life history trade-offs between growth and reproduction, and quantified plasticity in response to host species.

METHODS: Common garden experiments were used to evaluate trait differences between 11 *Euphrasia* taxa grown on a common host, document phenotypic plasticity when a single *Euphrasia* species is grown on eight different hosts, and relate observations to trait differences recorded in the wild.

RESULTS: *Euphrasia* exhibited variation in life history strategies; some individuals transitioned rapidly to flowering at the expense of early season growth, while others invested in vegetative growth and delayed flowering. Life history differences were present between some species, though many related taxa lacked clear trait differences. Species differences were further blurred by phenotypic plasticity—many traits were plastic and changed with host type or between environments.

CONCLUSIONS: Phenotypic plasticity in response to host and environment confounds species delimitation in *Euphrasia*. When grown in a common garden environment, some morphologically distinct taxa can be identified, though others represent morphologically similar shallow segregates. Trait differences present between some species and populations demonstrate the rapid evolution of distinct life history strategies in response to local ecological conditions.

KEY WORDS flowering time; host range; life history evolution; parasitic plants; phenotypic plasticity; Orobanchaceae.

Parasitism is a ubiquitous feature of the natural world, with parasitic organisms found living in every ecosystem and exploiting all free-living organisms (Price, 1980; Windsor, 1998). Parasitic plants are a group of ca. 4500 species of 12 separate evolutionary origins that have evolved a modified feeding organ, the haustorium, which allows them to attach to a host plant and extract nutrients and other compounds (Westwood et al., 2010; Nickrent and Musselman, 2017; Twyford, 2018). Parasitic plants are morphologically diverse and present a broad range of life history strategies and host interactions (Schneeweiss, 2006; Těšitel et al., 2010b). Hemiparasitic plants, i.e., those that are parasitic but also photosynthesize, can often attach to a broad range of hosts; the well-studied grassland parasite *Rhinanthus* has been found to attach to over 50 co-occurring grass and herbaceous species (Cameron et al., 2006). All hemiparasitic

plants are exoparasites; leaves, stems, roots, and flowers grow outside the host, and only the haustorium invades and grows within the host (Twyford, 2017).

Research to date has largely focused on three aspects of life history variation in parasitic plants. First, a body of work has looked to understand variation for specific traits between populations and related species. For example, work on natural populations of the hemiparasite *Pedicularis* has shown how investment in male reproductive organs depends on extrinsic environmental conditions (Guo et al., 2010a), while seed mass is primarily determined by intrinsic factors such as plant size rather than extrinsic factors such as elevation (Guo et al., 2010b). Second, researchers have investigated how parasite life history traits are affected by interactions with their host. In the widespread and weedy

obligate holoparasite *Phelipanche ramosa*, the duration of the life cycle differs depending on the host (Gibot-Leclerc et al., 2013), with a more rapid life cycle on local rather than non-local hosts. In hemiparasitic Rhinanthus minor, biomass depends on the host species and the number of haustorial connections (Rowntree et al., 2014). Finally, a number of studies have looked at life history variation between species studied in a phylogenetic context (Schneeweiss, 2006; Těšitel et al., 2010a). For example, broadscale analyses of the Rhinantheae clade in the Orobanchaceae has shown a shift from a perennial ancestor to annuality, with correlated shifts to a reduced seed size (Těšitel et al., 2010a). Despite the diversity of this research, there are still considerable gaps in our knowledge as to how life history trait variation is maintained (e.g., how common are trade-offs between life history traits), how much of this variation is genetic and how much is plastic, and which traits are the targets of natural selection.

In this study, we explore trait variation in generalist hemiparasitic eyebrights (Euphrasia, Orobanchaceae). Euphrasia is one of the largest genera of parasitic plants and is characterized by recent transoceanic dispersal and rapid species radiations (Gussarova et al., 2008). In the United Kingdom, there are 21 Euphrasia species, which are mostly indistinguishable at DNA barcoding loci (Wang et al., 2018), show complex morphological variation (Yeo, 1968; Metherell and Rumsey, 2018), and readily hybridize (Liebst, 2008; Stace et al., 2015). Despite shallow species differences due to postglacial divergence, Euphrasia species demonstrate substantial ecological divergence, with many taxa restricted to specific habitats such as coastal turf, mountain scree, heathland, or open grassland. Habitat differences would be expected to exert strong selection on life history traits, and this may include selection on growth to match seasonal water availability and to exploit local hosts, or selection on flowering time in response to local competition from surrounding plants, or in response to mowing or grazing (Hellström et al., 2004).

Our research builds on a large body of experimental work, with Euphrasia used in common garden studies for over 125 years (Koch, 1891). The first experimental work on Euphrasia revealed that phenotypic differences between two related species, E. rostkoviana and E. montana, are maintained in a common garden environment (Wettstein, 1895). Experimental work in the 1960s showed the growth of various Euphrasia species differs depending on the host species (Wilkins, 1963; Yeo, 1964). More recent experiments using large sample sizes in common gardens (Matthies, 1998; Zopfi, 1998; Lammi et al., 1999; Svensson and Carlsson, 2004) or in experimental field sites (Seel and Press, 1993; Hellström et al., 2004) have shown the effect of commonly encountered hosts such as grasses and legumes on hemiparasite biomass, mineral accumulation, plant architecture and reproductive output. Despite this extensive experimental work, studies in Euphrasia have yet to compare life history strategies of different species and the extent of phenotypic plasticity in life history traits. This work is critical for improving our knowledge of hemiparasite evolution and for understanding the nature of species differences in a taxonomically complex group. It is also unclear whether Euphrasia are restricted to growing on hosts such as grasses and herbaceous species or can parasitize a broad range of taxa including novel hosts rarely encountered in the wild. To address these questions requires simultaneously investigating the growth of multiple Euphrasia species and multiple host species with sufficient replication to enable suitable statistical comparisons.

Here, we used a series of common garden experiments, in conjunction with field observations, to understand life history trait

evolution, species differences, and phenotypic plasticity in hemiparasitic Euphrasia. Our first experiment assessed the morphological distinctiveness among several Euphrasia species and their hybrids when grown on a single host species in standardized common garden conditions. This experiment also addressed whether there is life history trait divergence among recently diverged hemiparasite species and whether these trait differences correspond to life history trade-offs. We then inspected the plasticity of a single focal Euphrasia population grown on many different hosts. This experiment quantified the magnitude of trait change when Euphrasia are grown on different hosts. It also tested whether they are truly generalist parasites by observing their growth on a wide range of hosts and without a host. Finally, we related our trait observations in a common garden to records of herbarium specimens collected in the wild. This comparison will help us understand whether life history traits and species' morphological differences are consistent between the common garden and the wild. Overall, our joint observations of phenotypic variation between closely related taxa and the extent of host-induced plasticity within a species, in an experiment and in the wild, provide new insights into variation in life history strategies in these hemiparasitic plants.

MATERIALS AND METHODS

Experimental design and plant cultivation

We performed two common garden experiments to investigate phenotypic variation in *Euphrasia*. Both common garden experiments took place in parallel in 2016. The experiments used wild-collected, openpollinated *Euphrasia* seeds that were pooled across individuals in a population. Seeds were contributed by plant recorders as part of the Eye for Eyebrights (E4E) public engagement project and as such included a scattered geographic sample across Great Britain (Appendix S1). All *Euphrasia* species were identified from the herbarium specimens of field collections, and from living material grown in the glasshouse, by *Euphrasia* referee Chris Metherell. Host seeds were sourced from commercial suppliers and from field collections (Appendix S2).

Species differences experiment—We observed trait differences of 24 populations from five *Euphrasia* species and six natural *Euphrasia* hybrids when grown on clover (*Trifolium repens*). This experiment included sampling multiple populations of three widespread and closely related grassland species, *E. arctica*, *E. confusa*, and *E. nemorosa*, and sparse population sampling of the moorland specialist *E. micrantha* (one population) and calcareous grassland specialist *E. pseudokerneri* (two populations). We chose clover as a host because it usually supports vigorous hemiparasitic growth and confers high survival (Zopfi, 1998).

Phenotypic plasticity experiment—We measured traits of a focal *Euphrasia* taxon, *E. arctica*, when grown with eight potential hosts (*Arabidopsis thaliana*, *Equisetum arvense*, *Festuca rubra*, *Holcus lanatus*, *Marchantia polymorpha*, *Pinus sylvestris*, *Plantago lanceolata*, and *Trifolium repens*) and without a host. These hosts were chosen to include a broad representation of functional groups and phylogenetic diversity, with species encountered in the wild and with novel hosts (full details in Appendix S2). The novel hosts were included to see the limits to which parasitic *Euphrasia* can associate, namely with a tree (*Pinus*), a pteridophyte that produces adventitious roots (*Equisetum*), and a liverwort that produces rhizoids (*Marchantia*).

Cultivation protocol—Reliable cultivation of *Euphrasia* can be challenging due to low seed germination, variation in time to establishment, the requirement of seed stratification, and high seedling mortality when transplanted (Yeo, 1961; Zopfi, 1998). We developed cultivation protocols that combine winter germination cues that improve germination and mimic nature, but also used highly standardized and replicated pot conditions that avoid transplanting Euphrasia and thus maximize survivorship. We filled 9-cm plastic pots with Melcourt Sylvamix Special growing medium (Tetbury, Gloucestershire, UK) in December, placed one Euphrasia seed per pot, and left pots outside over winter at the Royal Botanic Garden Edinburgh (RBGE) for seeds to experience natural seed stratification. Hosts were planted in seed trays in April. Euphrasia plants were moved to an unheated and well-ventilated greenhouse in the spring once the cotyledons were fully expanded, and a single seedling from each host (or a 1-cm² clump of Marchantia) was transplanted into the pot containing Euphrasia. Hosts that died within 10 days of planting were replaced. Twenty or more replicates were grown for each host-parasite combination. Plants were subsequently grown to flowering with regular watering, the locations of pots randomized at weekly intervals, and foreign weed seedlings removed.

Common garden trait measurements and statistical analyses

We measured seven morphological traits at first flowering related to life history variation, indicators of plant vigor, or characters used in taxonomy. In addition to date of first flowering, we recorded corolla length, the ratio of cauline leaf length to internode length below the measured leaf ("internode ratio"), number of leaf teeth on the lower floral leaf (bract), number of nodes to flower, number of branches, and plant height. All lengths were measured to the nearest millimeter as done by Metherell and Rumsey (2018). For the phenotypic plasticity experiment, we also recorded early season growth (height 6 weeks after transplantation of potential host) and height at the end of season after senescence. We did not directly observe host attachment, as preliminary investigations revealed a fine root structure where haustoria were difficult to observe. Instead, we inferred that attachment is likely to have taken place based on observations of height according to Yeo (1964). In his study, Euphrasia that attached to a "good" host tended to grow tall with elongated internodes, while Euphrasia that did not attach or attached to a "bad" host were much smaller (see discussion for more details).

We used a combination of fixed effect and mixed models to gain insights into the differences in means and the magnitude of variability in our data. In all models, response variables were analyzed as either Gaussian (and log-transformed if necessary) or Poisson. If the response variable was analyzed as Poisson, the model was checked for overdispersion and if it was overdispersed, an observation-level random effect was fitted. All correlations between variables were Pearson's correlations. Multiple correlation comparisons were corrected using Holm's correction method. Phenotypic clustering was inspected using principal component analysis (PCA). All analyses were done in R version 3.4.3, with the packages lme4 (Bates et al., 2015) and MCMCglmm (Hadfield, 2010) for generalized linear mixed effects models, base R for linear models, RcmdrMisc for correlations (Fox, 2018) and ggplot2 for data visualisation (Wickham, 2016). MCMCglmm models were run for a minimum of 70,000 iterations using either inverse Wishart or parameter-expanded priors with a minimum burn-in period of 30,000 iterations. Model convergence was assessed visually by plotting the posterior distributions and Markov chains.

In the species differences experiment, species of Euphrasia was fitted as a fixed effect, and population of Euphrasia was treated as a random effect. We excluded hybrids from these analyses because we were interested in testing differences between species. In the case of height and cauline to internode ratio, the traits were log transformed. Likelihood ratio tests calculated the overall significance of species, where this was not possible, deviance information criteria were used to test better model fit. We calculated proportion variance explained by population of Euphrasia (after accounting for fixed effects) by dividing the population random effect variance by the total variance in the model. Tukey post hoc tests were performed on each pairwise comparison of *Euphrasia* species and adjusted p-values calculated, using Tukey honestly significant difference (HSD) test and correcting for family-wise error rate in the emmeans R package (Lenth, 2019). For the phenotypic plasticity experiment, host species was fitted as a fixed effect. The models were re-levelled so that "no host" was the baseline. Analysis of variance was used to determine overall significance of host species. Tukey post hoc tests were then performed on each pairwise comparison of host species, with adjusted p-values calculated in base R and the multcomp package (Hothorn et al., 2008).

Trait variation in the wild

We tested how phenotypes in the experiments related to those in nature by comparing results from the species differences experiment to phenotypic measurements of herbarium specimens of the same population sampled in the wild. Three individuals were measured from each collection sheet for a given population for each trait. Pressed plants submitted by collectors varied in quality, and therefore, we were unable to measure the height of these plants, nor was it possible to infer date of first flowering. We analyzed the data using generalized linear mixed effect models with where individuals were grown (i.e., common garden or wild-collected) as a fixed effect, with each of five traits as the response variable. We treated species and population of Euphrasia as random effects to understand the relative contributions of each to the overall variability in a given trait. Response variables that were considered count data were analyzed with a Poisson distribution, in all other cases a Gaussian distribution was used. R-values were calculated using Pearson's correlations of the population level means between the common garden and the wild samples.

RESULTS

Species differences

Our species differences experiment revealed extensive morphological trait variation across Euphrasia species when compared at first flowering. From the 222 Euphrasia individuals that survived to flower on their clover host, the greatest variation was seen in number of branches (9-fold difference between species), internode ratio (2.7-fold) and height (2.5-fold), while traits such as node to flower (1.6-fold) and corolla length (1.6-fold) proved less variable (Fig. 1A–D; Appendix S3). A large degree of this variation was separated by species and by population (Table 1). The species with the most distinct life history strategy was E. micrantha, which flowered from a low node on the plant

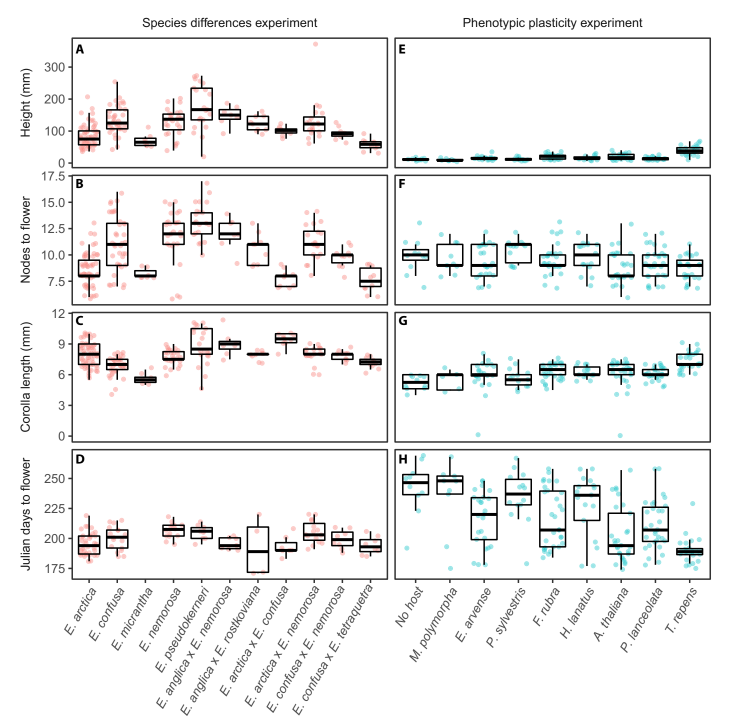


FIGURE 1. Trait variation in a common garden experiment of (A–D) diverse *Euphrasia* species and hybrids grown on clover and (E–H) *Euphrasia arctica* grown on many different hosts. The edges of the box plots show the first and third quartiles, the solid lines the median, the whiskers the highest and lowest values within 1.5-fold of the inter-quartile range, and the jittered dots each individual measurement.

 $(8.3\pm0.2~{\rm nodes})$ while it was short $(70\pm8~{\rm mm};{\rm Appendix~S3})$. It also formed a partly distinct cluster in the PCA (Appendix S4). *Euphrasia pseudokerneri* was relatively distinct, flowered once it had grown tall $(176\pm16~{\rm mm})$ and from a high node on the plant $(13.2\pm0.4~{\rm nodes})$, but showed little separation in the PCA. The morphologically similar *E. arctica, E. confusa*, and *E. nemorosa* differed for some traits, with *E. nemorosa* initiating flowering 14 days later and from 3.3 nodes

higher than *E. arctica*, but overlapped in many other traits and in overall multi-trait phenotype (Appendices S4, S5). Despite species being a significant factor in the models, and some notable differences in specific traits, there were few significant pairwise Tukey comparisons due to substantial within-species variation (Appendix S6). Of the seven significant pairwise trait differences, three were for node to flower and three for number of leaf teeth, with four of the seven

TABLE 1. Summary of generalized linear mixed effects models for Euphrasia trait values measured in a common garden environment. Model outputs are summarized for five Euphrasia species grown with clover in the species differences experiment and for E. arctica grown with eight hosts and without a host in the phenotypic plasticity experiment. For the phenotypic plasticity experiment, we report model outputs with all potential hosts, as well as models excluding Pinus and Marchantia when there was no evidence of attachment or interactions (reported in square brackets). The percentage variance explained by random effects are reported in parentheses along with the 95% credibility interval. ***p < 0.001, **p < 0.05.

	Species	Phenotypic plasticity		
rait	Species	Population	Host	
ranches	DIC _{full} = 676.08; DIC _{-spp} = 679.48	DIC _{full} = 676.08; DIC _{pop} = 714.87 (25.9%, 13.4–57.4%)	NA	
orolla length	$\chi^2(4) = 11.91*$	$\chi^2(1) = 41.38^{***} (54.1\%, 24.4-69.2\%)$	$(F_{8,173} = 9.85)^{***}$ $[(F_{6,157} = 11.38)^{***}]$	
leight	$\chi^2(4) = 11.67^*$	$\chi^2(1) = 57.13*** (61.2\%, 35.0-79.7\%)$	$(F_{8,185} = 23.14)^{***}$ $[(F_{6,164} = 24.39)^{***}]$	
nternode ratio	$\chi^2(4) = 13.00*$	$\chi^2(1) = 34.38^{***}$	$(F_{8,184} = 3.36)^{**}$ $[(F_{6,163} = 4.11)^{***}]$	
ulian days to flower	$\chi^2(3) = 2.26$	$\chi^2(1) = 1.42 \text{ E-}14$ (58.7%, 28.6–80.9%)	$\chi^{2}(8) = 192.39***$ [$\chi^{2}(6) = 141.67***$]	
lodes to flower	$\chi^2(4) = 15.42^{**}$	$\chi^2(1) = 2.87$ (14.1%, 1.0–33.5%)	$\chi^2(8) = 5.02$ [$\chi^2(6) = 3.04$]	
Number of leaf teeth	$\chi^2(3) = 12.45**$	$\chi^2(1) = 0.0059$ (0.12%, 2.8–23%)	$\chi^2(8) = 26.79^{***}$ [$\chi^2(6) = 17.04^{**}$]	

Note: Models for number of branches were implemented with a different statistical approach in MCMCglmm, with models compared using the deviance information criterion (see methods)

significant comparisons involving E. micrantha. In most cases, hybrids combined morphological characters of their parental progenitors. For example, hybrids involving E. nemorosa flowered later in the season and initiated flowering from a higher node than E. arctica hybrids (Fig. 1A-D).

Correlation analyses across species revealed clear suites of traits that are related. Significant correlations were found between 12 of the 21 pairwise comparisons after correcting for multiple tests, with five of these correlations with R > 0.6 (Table 2A). Plants flowering at a late node are more likely to be tall, more highly branched, and have many teeth on the lower floral leaf. The relationship of traits is also supported in the PCA, with many traits contributing to multiple principal components (Appendix S5). Traits related to height and flowering node were largely uncorrelated with internode ratio and corolla length.

Phenotypic plasticity

Our phenotypic plasticity experiment showed substantial morphological variation across 194 E. arctica plants grown with eight different potential host species and the 22 plants grown without a host. Plants growing on clover transitioned to flower quickly (189.8 \pm 2.0 Julian days), grew tall by the time of first flowering (39 \pm 3 mm), and produced large flowers (7.4 \pm 0.2 mm; Fig. 1E– H, Appendix S7). These results contrast with Euphrasia with no host, which flowered on average 52 days later (241.3 \pm 7.9 Julian days), were extremely short at first flowering (11 \pm 1 mm), and produced small flowers (5.3 \pm 0.2 mm). Euphrasia arctica grown on Arabidopsis, Equisetum, Festuca, Holcus, or Plantago were all statistically significantly different from no host for at least one trait (Tukey comparisons, p < 0.05), while *E. arctica* on *Marchantia* or Pinus was not significantly different from no host for any trait (p > 0.05; Appendix S8). While the overall effect of host was significant for all traits except nodes to flower (Appendices S9–S11), three traits showed relatively little plasticity, with few statistically

significant pairwise Tukey comparisons for nodes to flower (0 significant comparisons), number of leaf teeth (3), and internode length (4), while the other three traits showed many pairwise differences (days to flower, 21 significant comparisons; height, 16; corolla length, 12; Appendix S8). Our comparison of growth traits across host treatments measured through the year showed that height at the end of the season was weakly predicted from height 6 weeks after introducing a host (R = 0.47), but strongly correlated with height at first flowering (R = 0.82; Appendix S12). Plants that flowered early were more likely to grow larger by the end of season (R = -0.55) and become more highly branched (R = -0.57;Appendix S12).

Across host treatments, there was a significant negative correlation between Julian days to flower and most other traits (Table 2B). We find that late flowering individuals are likely to be smaller at first flowering and have fewer branches, leaves with fewer teeth, and smaller flowers. While these traits were strongly correlated, there were substantial differences in the magnitude of response. For example, days to flower differed considerably depending on host, with a 3.8-fold greater difference than seen between means for different Euphrasia species grown on the same host (Fig. 1D, H). In contrast, corolla length and node to flower proved less variable depending on host, with a 1.4-fold and 1.2-fold change between means, respectively.

Variation in the wild

The comparison between the species differences common garden experiment and wild-collected herbarium specimens revealed population means of a single trait, nodes to flower, are strongly correlated (R = 0.79), and trait values are not significantly different ($p_{MCMC} = 0.71$) between environments (Fig. 2; Appendices S13, S14). All other traits did differ significantly between environments ($p_{MCMC} < 0.05$), with *Euphrasia* plants in the common garden having corollas on average 1.4 mm longer, with

TABLE 2. Pearson's correlation coefficients for seven phenotypic traits measured in a common garden experiment for (A) five *Euphrasia* species and six hybrids, (B) *Euphrasia arctica* grown with eight hosts and without a host. ***p < 0.001, **p < 0.05. Asymptotic p-values are reported from the Hmisc package in R using the rcorr() function.

(A) Species differences experiment									
Trait	Corolla length (mm)	Height (mm)	Internode ratio	Julian days to flower	Leaf teeth	Nodes to flower			
Branches	0.260**	0.609***	-0.116	0.057	0.658***	0.775***			
Corolla length (mm)		0.319***	-0.161	-0.127	0.197*	0.049			
Height (mm)			0.246**	0.292*	0.563***	0.628***			
Internode ratio				0.204	-0.120	0.076			
Julian days to flower					0.053	0.249*			
Leaf teeth						0.651***			

(B) Phenotypic plasticity experiment									
Trait	Corolla length (mm)	Height (mm)	Internode ratio	Julian days to flower	Leaf teeth	Nodes to flower			
Branches Corolla length (mm) Height (mm) Internode ratio Julian days to flower Leaf teeth	0.524***	0.834*** 0.503***	-0.299*** 0.098 0.477***	-0.572*** -0.406*** -0.481*** -0.034	0.694*** 0.536*** 0.692*** 0.168 -0.691***	-0.572** -0.166 -0.186 -0.009 0.530*** -0.239**			

0.2 more teeth on the lower floral leaves, an increase in internode ratio of 1.0 mm, and 4 more pairs of branches. Despite these differences, there were correlations between the common garden and the wild-collected specimens for corolla length (R=0.93, $p_{\rm MCMC}<0.001$), internode ratio (R=0.65, $p_{\rm MCMC}<0.001$) and number of branches (R=0.29, $p_{\rm MCMC}<0.001$), but not for number of leaf teeth (R=0.07, $p_{\rm MCMC}=0.034$).

DISCUSSION

Our study sheds light on species differences, life history evolution and phenotypic plasticity of the generalist parasitic plant *Euphrasia*. We found different life history strategies between recently diverged species, with some species rapidly transitioning to flower at the expense of growth-related traits, while others delay flowering and invest in early-season vegetative growth. However, many traits are phenotypically plastic and change in response to the host. While plants in benign common garden conditions grew vigorously, the correlation between life history traits in a common garden and in the wild suggests our experimental observations are indicative of patterns observed in nature. Morphological differences between species in the common garden also suggest that the currently delimited Euphrasia species are, at least in part, distinct. Overall, our study highlights the value of integrating trait data from multiple common garden experiments and field collections to study life history strategies in parasitic plants and demonstrates the rapid evolution of life history differences in a postglacial radiation of hemiparasites.

Life history variation in a generalist hemiparasitic plant

We found evidence for different life history strategies in British Euphrasia. Euphrasia arctica, E. micrantha, and hybrids such as E. $arctica \times E$. confusa, transition rapidly to flower, flower while they are short, and produce their first flower from a low node on the plant. These rapid flowering species contrast with E. pseu-dokerneri, E. nemorosa and hybrids involving E. nemorosa that

delay flowering until later in the season, grow tall before flowering, and produce their first flower from a late node on the main axis. These different life history strategies correspond to the known ecology of these species, with E. nemorosa flowering late in tall mixed grassland, while E. micrantha flowers early in patchy heathland (Metherell and Rumsey, 2018). While species show some general differences in life history strategies, there is also significant variation between populations within species. A relationship between internode number and habitat has previously been observed within Euphrasia species, with populations of E. rostkoviana in Sweden flowering at a lower node in a common garden if they have been collected from intensely grazed pasture (Zopfi, 1998). Overall, these observations within and between species are consistent with the classic life history trade-off between growth and reproduction (Stearns, 1992; Roff, 2002). For Euphrasia growing in the wild, early reproduction allows the plants to reliably complete their life cycle before summer competition, herbivory, mowing, summer drought, and other seasonal abiotic and biotic stresses. However, early flowering involves reproducing at the expense of early-season growth and at a time when the resource budget may be constrained by relatively few haustorial connections. These trait trade-offs pose an interesting comparison to the well-studied Mimulus guttatus (syn. Erythranthe guttata), a nonparasitic relative in the Lamiales that shares the same basic plant architecture. In M. guttatus, multiple traits related to growth and reproduction are correlated, both within and between populations, due to genetic trade-offs between time to flower and fecundity (Mojica et al., 2012; Friedman et al., 2015). In Euphrasia, the genetics underpinning this life history trade-off have yet to be characterized and may be a consequence of multiple independent loci or trade-offs at individual loci (Hall et al., 2010).

While much life history variation is captured by differences in time to flower and growth-related traits, we also see evidence for flower size representing a separate axis of variation across *Euphrasia* species. In our common garden, *E. micrantha* has small corollas, while *E. arctica* and *E. nemorosa* have larger corollas, and corolla size is not strongly correlated with other traits.

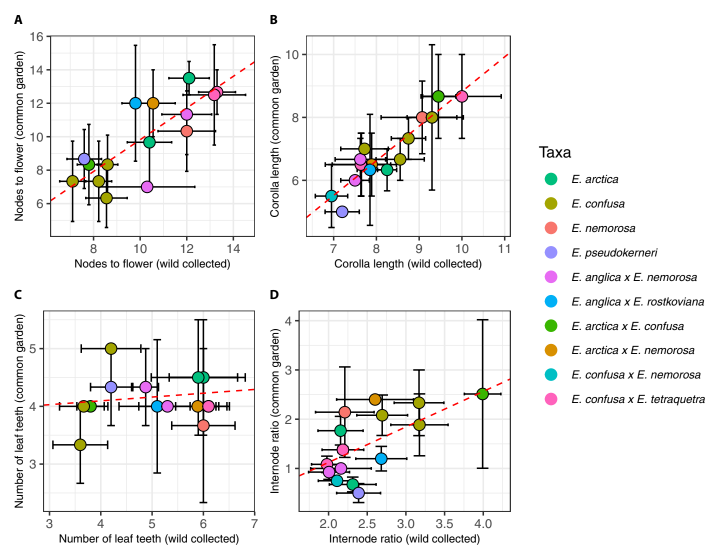


FIGURE 2. Relationship between morphological trait measurements made in the common garden and on wild-collected herbarium specimens for diverse Euphrasia species. Points are Euphrasia population means; bars represent the standard error of measurements. The line of best fit was calculated using coefficients from linear regression models on the means of each Euphrasia population. Length measurements are reported in millimeters. For an alternative representation of pairwise comparisons, see Appendix S13.

Euphrasia species are well known to have flower size variation, with a continuum between small-flowered species that are highly selfing (e.g., E. micrantha, corolla size = 4.5-6.5 mm, inbreeding coefficient $F_{1s} > 0.88$; Stone, 2013) and large-flowered species that are highly outcrossing (e.g., E. rostkoviana, flower size 8–12 mm, $F_{\rm IS} = 0.17 - 0.25$; French et al., 2005). Such wide variation in outcrossing rate has been documented in species of Datura (Motten and Stone, 2000), Mimulus (Karron et al., 1997), and Nicotiana (Breese, 1959). Small flowers have shorter anther-stigma separation and thus increased potential for autogamous selfing (Karron et al., 1997), while also having reduced attractiveness to pollinators and thus receiving less outcross pollen (Mitchell et al., 2004). In addition to differences in corolla size between Euphrasia species, corolla size also shows a change of up to 2 mm in response to host species. This change in flower size is of a magnitude that may potentially affect the mating system (Luo and Widmer, 2013) and suggests host species represents a previously unaccounted factor affecting the mating system of parasitic plants.

Our comparisons of Euphrasia species in a common garden also shed light on the distinctiveness of these recently diverged species and can be used to refine the suite of traits that are reliable in telling Euphrasia species apart. Euphrasia is a taxonomically complex plant genus, with the 21 currently described British species presenting complex and often overlapping morphological variation (French et al., 2008; Metherell and Rumsey, 2018; Wang et al., 2018). Our study suggests varying degrees of morphological distinctiveness of Euphrasia species. We see E. micrantha is morphologically distinct in the common garden and E. pseudokerneri somewhat distinct, while the closely related species E. arctica, E. confusa, and E. nemorosa differ in life history traits such as nodes to flower, but overlap in many other traits and are not clearly separated in the PCA. The morphological trait differences between species observed under standardized conditions are correlated with values from field-collected herbarium specimens where plants have associated with diverse hosts, been exposed to different ecological conditions, and were collected at

different life-stages. These correlations suggest that our common garden results generalize to observations in nature. However, our study is likely to overestimate the distinctiveness of taxa by only including a subset of UK species and by choosing populations that could be identified to species level in the field. We suspect adaptive divergence between closely related *E. arctica*, *E. confusa*, and E. nemorosa is a consequence of differential natural selection for local ecological conditions such as soil water availability or mowing. Selection appears to be operating at a fine spatial scale, with significant life history trait differences evident between populations within species. Euphrasia taxa may be genetically cohesive, either showing genome-wide divergence or divergence in genomic regions underlying life history differences (Twyford and Friedman, 2015), or alternatively these taxa may be polytopic and not genetically cohesive (Hollingsworth et al., 2017). Genomic sequencing of natural populations will help resolve the nature of species differences in Euphrasia.

Phenotypic plasticity in response to host

Our phenotypic plasticity experiment shows Euphrasia are affected by growing with a range of different hosts. Specifically, E. arctica with a host such as clover rapidly transitions to flowering. At the other extreme, Euphrasia grown without a host are small and flower late. These differences in growth are established early in the season, and early-flowering plants go on to grow the tallest and are more highly branched. Most other hosts result in a continuum of Euphrasia phenotypes between these extremes. Two surprising results were that E. arctica parasitizing Arabidopsis grew relatively tall despite the host senescing early in the growth season and that growth of Euphrasia with Equisetum was similar to growth on the commonly encountered grass Holcus lanatus. This result suggests that it attached to Equisetum, which would need to be confirmed by excavating root systems and observing haustoria, or it indirectly benefits without attachment through association with Equisetum fungal symbionts (Bouwmeester et al., 2007). Less surprising was the poor growth of E. arctica with Pinus. However, an association between Melampyrum pratense and Pinus sylvestris suggests at least some hemiparasitic Orobanchaceae benefit from attachment to woody host species or from interactions with their associated ectomycorrhizal fungi (Salonen et al., 2000).

The diverse effects of host on parasite growth are complex, but the variation we saw in our experiments may be attributed to host root architecture, germination time, and resource availability, as well as the presence of mechanisms to defend against parasite attack, such as cell wall thickening, localized host dieback, and chemical defence (Cameron et al., 2006; Twyford, 2018). While *Euphrasia* is generally thought to have low reliance on host resources, deriving only ~30% of carbon heterotrophically (Těšitel et al., 2010b), at least under our experimental conditions *Euphrasia* only produced multiple flowers on certain hosts. Overall, our results point to *E. arctica* being a true generalist hemiparasite, but one where vigorous growth is only observed with a subset of potential hosts.

In terms of specific traits, only three pairs of trait correlations show consistent correlation coefficients in both *Euphrasia* common garden experiments (between height, number of branches, and leaf teeth), with other correlations between species breaking down when *Euphrasia* are grown on different hosts. The most notable plasticity is seen in flowering time, with plants on clover

rapidly transitioning to flower within ~100 days of germination, while plants with a more typical host (e.g., *Holcus lanatus*) flower a month later. Phenotypic plasticity in flowering time in response to resource availability is well documented in many plant groups, particularly *Arabidopsis* (e.g., Zhang and Lechowicz, 1994), but has received less attention in studies of parasitic plants, which are more likely to look at growth-related traits such as biomass (Ahonen et al., 2005; Matthies, 2017). However, date of first flowering has been shown to differ by up to 10 weeks in populations of *Rhinanthus glacialis* across Switzerland (Zopfi, 1995). Overall, we expect date of first flowering to be critical for the lifetime reproductive success of parasitic plants in the wild.

In contrast to seeing traits with extensive plasticity, we also saw evidence of developmental constraint in number of nodes to flower. For E. arctica, this trait showed the least plasticity with different hosts, is consistent between populations within species, and between the common garden and the field. Thus, the developmental event of transitioning to flower may be genetically determined, with changes in flowering time altered by plasticity in internode length and not nodes to flower. This developmental constraint may explain why nodes to flower is such an important diagnostic trait for species identification in Euphrasia and related species in the Rhinantheae (Jonstrup et al., 2016). Despite nodes to flower changing little in response to host species, our overall impression is that *Euphrasia* show considerable plasticity and little developmental constraint in many aspects of growth. In particular, differences between individuals on a given host also suggests other sources of variation, such as genetic background in host and parasite, as well as the timing of attachment, may be crucial in determining performance.

CONCLUSIONS

Despite over a century of experimental studies in parasitic plants, our understanding of the evolution of life history strategies in these diverse organisms is extremely limited. Our results with *Euphrasia* provide strong support for the rapid evolution of distinct life history strategies in response to local ecological conditions, with phenotypic plasticity further altering plant growth in response to host availability. We anticipate that future studies that test lifetime reproductive success of many parasitic plant species grown on many different host species will give further insight into the complex nature of host–parasite interactions.

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AUTHOR CONTRIBUTIONS

A.D.T conceived and designed the research. A.D.T., N.F., and E. L. Y.W. carried out the experiments. C.M. identified the plants. A.D.T. and M.B. analyzed the data. A.D.T. and M.B. wrote the manuscript. All authors read and approved the manuscript.

DATA AVAILABILITY

Phenotypic data from both common garden experiments and from herbarium collections, as well as the R scripts used for data analysis, are deposited in Dryad (doi.org/10.5061/dryad.cfxpnvx2d). Source code and data are also deposited on Github: https://github.com/ Euphrasiologist/phenotypic_plasticity_euphrasia.

SUPPORTING INFORMATION

Additional Supporting Information may be found online in the supporting information tab for this article.

APPENDIX S1. Host species used in the common garden experiment.

APPENDIX S2. Collection details for *Euphrasia* species used in the common garden experiment.

APPENDIX S3. Summary of trait values for many Euphrasia species and hybrids grown on a clover host.

APPENDIX S4. Principal component analysis of morphological variation of Euphrasia in a common garden.

APPENDIX S5. Factor loadings for the principal component analyses of (A) five species and six hybrids, (B) five species, and (C) Euphrasia arctica with eight hosts and no host.

APPENDIX S6. Pairwise differences in trait value of Euphrasia species grown with clover in a common garden experiment.

APPENDIX S7. Summary of trait values for Euphrasia arctica grown on many different hosts.

APPENDIX S8. Comparison of Euphrasia arctica traits in the phenotypic plasticity common garden experiment.

APPENDIX S9. Analysis of deviance for each trait in the phenotypic plasticity experiment with Euphrasia arctica grown with many different hosts.

APPENDIX \$10. ANOVAs for traits measured in the phenotypic plasticity experiment with Euphrasia arctica grown with many dif-

APPENDIX S11. Summary of generalized linear models for the phenotypic plasticity experiment with Euphrasia arctica grown on many hosts in a common garden.

APPENDIX S12. Relationship between growth-related traits and end of season height for Euphrasia arctica grown with eight hosts and no host.

APPENDIX S13. Comparison between trait values for wildcollected herbarium specimens and common garden plants of diverse Euphrasia species.

APPENDIX S14. Model output from MCMCglmm comparing traits for the wild collected Euphrasia specimens to the baseline of the common garden data.

LITERATURE CITED

- Ahonen, R., S. Puustinen, and P. Mutikainen. 2005. Host use of a hemiparasitic plant: no trade-offs in performance on different hosts. Journal of Evolutionary Biology 19: 513-521.
- Bates, D., M. Maechler, B. Bolker, and S. Walker. 2015. Fitting linear mixed-effects models using lme4. Journal of Statistical Software 67: 1-48.
- Bouwmeester, H. J., C. Roux, J. A. Lopez-Raez, and G. Becard. 2007. Rhizosphere communication of plants, parasitic plants and AM fungi. Trends in Plant Science 12: 224-230.
- Breese, E. 1959. Selection for differing degrees of out-breeding in Nicotiana rustica. Annals of Botany 23: 331-344.
- Cameron, D. D., A. M. Coats, and W. E. Seel. 2006. Differential resistance among host and non-host species underlies the variable success of the hemiparasitic plant Rhinanthus minor. Annals of Botany 98: 1289-1299.
- Fox, J. 2018. RcmdrMisc: R commander miscellaneous functions. R package version 2.5-1. Website: https://CRAN.R-project.org/package=RcmdrMisc.
- French, G., R. Ennos, A. Silverside, and P. Hollingsworth. 2005. The relationship between flower size, inbreeding coefficient and inferred selfing rate in British Euphrasia species. Heredity 94: 44.
- French, G., P. Hollingsworth, A. Silverside, and R. Ennos. 2008. Genetics, taxonomy and the conservation of British Euphrasia. Conservation Genetics 9: 1547-1562.
- Friedman, J., A. D. Twyford, J. H. Willis, and B. K. Blackman. 2015. The extent and genetic basis of phenotypic divergence in life history traits in Mimulus guttatus. Molecular Ecology 24: 111–122.
- Gibot-Leclerc, S., F. Dessaint, C. Reibel, and V. Le Corre. 2013. Phelipanche ramosa (L.) Pomel populations differ in life-history and infection response to hosts. Flora - Morphology, Distribution, Functional Ecology of Plants 208:
- Guo, H., S. J. Mazer, and G. Du. 2010a. Geographic variation in primary sex allocation per flower within and among 12 species of *Pedicularis* (Orobanchaceae): Proportional male investment increases with elevation. American Journal of Botany 97: 1334-1341.
- Guo, H., S. J. Mazer, and G. Du. 2010b. Geographic variation in seed mass within and among nine species of Pedicularis (Orobanchaceae): effects of elevation, plant size and seed number per fruit. Journal of Ecology 98: 1232-1242.
- Gussarova, G., M. Popp, E. Vitek, and C. Brochmann. 2008. Molecular phylogeny and biogeography of the bipolar Euphrasia (Orobanchaceae): recent radiations in an old genus. Molecular Phylogenetics and Evolution 48: 444-460.
- Hadfield, J. D. 2010. MCMC methods for multi-response generalized linear mixed models: the MCMCglmm R package. Journal of Statistical Software 33: 1-22.
- Hall, M. C., D. B. Lowry, and J. H. Willis. 2010. Is local adaptation in Mimulus guttatus caused by trade-offs at individual loci? Molecular Ecology 19: 2739-2753.
- Hellström, K., P. Rautio, A.-P. Huhta, and J. Tuomi. 2004. Tolerance of an annual hemiparasite, Euphrasia stricta agg., to simulated grazing in relation to the host environment. Flora - Morphology, Distribution, Functional Ecology of Plants 199: 247-255.
- Hollingsworth, P. M., L. E. Neaves, and A. D. Twyford. 2017. Using DNA sequence data to enhance understanding and conservation of plant diversity at the species level. In S. Oldfield and S. Blackmore [eds.], Plant conservation science and practice: the role of Botanic Gardens, 23-48. Cambridge University Press, Cambridge, UK.
- Hothorn, T., F. Bretz, and P. Westfall. 2008. Simultaneous inference in general parametric models. Biometrical Journal 50: 346-363.
- Jonstrup, A., M. Hedrén, and S. Andersson. 2016. Host environment and local genetic adaptation determine phenotype in parasitic Rhinanthus angustifolius. Botanical Journal of the Linnean Society 180: 89–103.

- Karron, J. D., R. T. Jackson, N. N. Thumser, and S. L. Schlicht. 1997. Outcrossing rates of individual Mimulus ringens genets are correlated with anther-stigma separation. Heredity 79: 365.
- Koch, L. 1891. Zur Entwicklungsgeschichte der Rhinanthaceen (II Euphrasia officinalis L.). Jahrbücher für Wissencshafltiche Botanik 22: 1-34.
- Lammi, A., P. Siikamäki, and V. Salonen. 1999. The role of local adaptation in the relationship between an endangered root hemiparasite Euphrasia rostkoviana, and its host, Agrostis capillaris. Ecography 22: 145-152.
- Lenth, R. 2019. emmeans: estimated marginal means, aka least-squares means. R package version 1.3.5.1. Website: https://CRAN.R-project.org/package=
- Liebst, B. 2008. Do they really hybridize? A field study in artificially established mixed populations of Euphrasia minima and E. salisburgensis (Orobanchaceae) in the Swiss Alps. Plant Systematics and Evolution 273: 179-189.
- Luo, Y., and A. Widmer. 2013. Herkogamy and its effects on mating patterns in Arabidopsis thaliana. PLoS ONE 8: e57902.
- Matthies, D. 1998. Influence of the host on growth and biomass allocation in the two facultative root hemiparasites Odontites vulgaris and Euphrasia minima.
- Matthies, D. 2017. Interactions between a root hemiparasite and 27 different hosts: growth, biomass allocation and plant architecture. Perspectives in Plant Ecology, Evolution and Systematics 24: 118-137.
- Metherell, C., and F. J. Rumsey. 2018. Eyebrights (Euphrasia) of the UK and Ireland. Botanical Society of Britain and Ireland, Bristol, UK.
- Mitchell, R., J. Karron, K. Holmquist, and J. Bell. 2004. The influence of Mimulus ringens floral display size on pollinator visitation patterns. Functional Ecology 18: 116-124.
- Mojica, J. P., Y. W. Lee, J. H. Willis, and J. K. Kelly. 2012. Spatially and temporally varying selection on intrapopulation quantitative trait loci for a life history trade-off in Mimulus guttatus. Molecular Ecology 21: 3718-3728.
- Motten, A. F., and J. L. Stone. 2000. Heritability of stigma position and the effect of stigma-anther separation on outcrossing in a predominantly selffertilizing weed, Datura stramonium (Solanaceae). American Journal of Botany 87: 339-347.
- Nickrent, D. L., and L. J. Musselman. 2017. Parasitic plants. In B. H. Ownley and R. N. Trigiano [eds.], Plant pathology: concepts and laboratory exercises, 277-288. CRC Press, Boca Raton, LA, USA.
- Price, P. W. 1980. Evolutionary biology of parasites. Princeton University Press, Princeton, NJ, USA.
- Roff, D. A. 2002. Life history evolution. Sinauer, Sunderland, MA, USA.
- Rowntree, J. K., D. Fisher Barham, A. J. Stewart, and S. E. Hartley. 2014. The effect of multiple host species on a keystone parasitic plant and its aphid herbivores. Functional Ecology 28: 829-836.
- Salonen, V., H. Setälä, and S. Puustinen. 2000. The interplay between Pinus sylvestris, its root hemiparasite, Melampyrum pratense, and ectomycorrhizal fungi: influences on plant growth and reproduction. Écoscience 7: 195-200.
- Schneeweiss, G. M. 2006. Correlated evolution of life history and host range in the nonphotosynthetic parasitic flowering plants Orobanche and Phelipanche (Orobanchaceae). Journal of Evolutionary Biology 20: 471-478.
- Seel, W., and M. Press. 1993. Influence of the host on three sub-Arctic annual facultative root hemiparasites. New Phytologist 125: 131-138.

- Stace, C. A., C. D. Preston, and D. A. Pearman. 2015. Hybrid flora of the British Isles. Botanical Society of Britain and Ireland, Bristol, UK.
- Stearns, S. C. 1992. The evolution of life histories, vol. 575. Oxford University Press, Oxford, UK.
- Stone, H. 2013. Evolution and conservation of tetraploid Euphrasia L. in Britain. PhD thesis, University of Edinburgh, Edinburgh, UK.
- Svensson, B. M., and B. Å. Carlsson. 2004. Significance of time of attachment, host type, and neighbouring hemiparasites in determining fitness in two endangered grassland hemiparasites. Annales Botanici Fennici 41: 63 - 75.
- Těšitel, J., L. Plavcová, and D. D. Cameron. 2010a. Heterotrophic carbon gain by the root hemiparasites, Rhinanthus minor and Euphrasia rostkoviana (Orobanchaceae). Planta 231: 1137-1144.
- Těšitel, J., P. Říha, Š. Svobodová, T. Malinová, and M. Štech. 2010b. Phylogeny, life history evolution and biogeography of the rhinanthoid Orobanchaceae. Folia Geobotanica 45: 347-367.
- Twyford, A. D. 2017. New insights into the population biology of endoparasitic Rafflesiaceae. American Journal of Botany 104: 1433-1436.
- Twyford, A. D. 2018. Parasitic plants. Current Biology 28: R857-R859.
- Twyford, A. D., and J. Friedman. 2015. Adaptive divergence in the monkey flower Mimulus guttatus is maintained by a chromosomal inversion. Evolution 69: 1476-1486.
- Wang, X., G. Gussarova, M. Ruhsam, N. de Vere, C. Metherell, P. M. Hollingsworth, and A. D. Twyford. 2018. DNA barcoding a taxonomically complex hemiparasitic genus reveals deep divergence between ploidy levels but lack of species-level resolution. AoB PLANTS 10: ply026-ply026.
- Westwood, J. H., J. I. Yoder, and M. P. Timko. 2010. The evolution of parasitism in plants. Trends in Plant Science 15: 227-235.
- Wettstein, R. 1895. Der Saison-Dimorphismus als Ausgangspunkt für die Bildung neuer Arten im Pflanzenreiche. Berichte der Deutschen Botanishchen Gesellschaft 33: 303-313.
- Wickham, H. 2016. ggplot2: Elegant graphics for data analysis. Springer-Verlag, NY, NY, USA.
- Wilkins, D. 1963. Plasticity and establishment in Euphrasia. Annals of Botany 27: 533-552.
- Windsor, D. A. 1998. Controversies in parasitology, Most of the species on Earth are parasites. International Journal for Parasitology 28: 1939-1941.
- Yeo, P. 1961. Germination, seedlings, and the formation of haustoria in Euphrasia. Watsonia 5: 1-22.
- Yeo, P. 1964. The growth of Euphrasia in cultivation. Watsonia 6: 1-24.
- Yeo, P. 1968. The evolutionary significance of the speciation of Euphrasia in Europe. Evolution 22: 736-747.
- Zhang, J., and M. J. Lechowicz. 1994. Correlation between time of flowering and phenotypic plasticity in Arabidopsis thaliana (Brassicaceae). American Journal of Botany 81: 1336-1342.
- Zopfi, H.-J. 1995. Life history variation and infraspecific heterochrony in Rhinanthus glacialis (Scrophulariaceae). Plant Systematics and Evolution 198: 209-233.
- Zopfi, H. J. 1998. The genetic basis of ecotypic variants of Euphrasia rostkoviana Hayne (Scrophulariaceae) in relation to grassland management. Flora 193: 41 - 58.