Appendices

Appendix 1: Chapter 2

Figure 1: The distribution of ploidy levels across the British and Irish angiosperms in the four families with the highest number of species. Shown are Rosaceae, Poaceae, Asteraceae and Fabaceae. Each family has distinct distributions of ploidy levels.

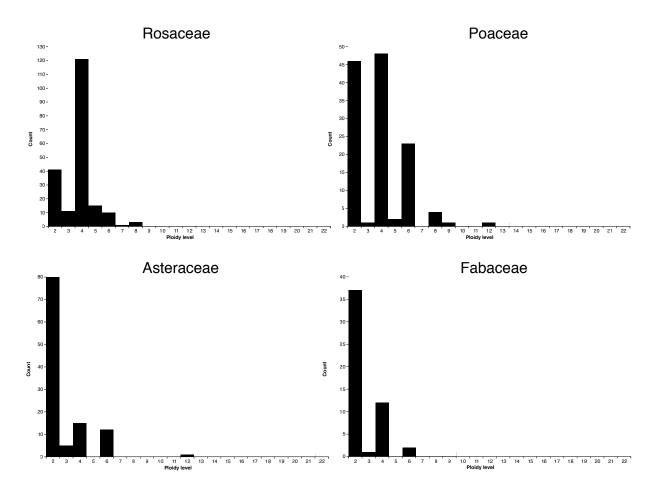


Table 1: Search strings for Google Scholar searches used to generate the list of examples of cross-ploidy hybrids in Chapter 2. Note that other examples were added if they were deemed to be important and/or well known.

Journal	Search string
Molecular Ecology	Ploidy hybrid genetic introgression diploid OR tetraploid OR hexaploidy OR octoploid source: "Molecular Ecology"
Evolution	Ploidy hybrid genetic introgression diploid OR tetraploid OR hexaploidy OR octoploid site:onlinelibrary.wiley.com source:"Evolution" -source:"and Evolution" -source:"Organic Evolution"
Heredity	Ploidy hybrid genetic introgression diploid OR tetraploid OR hexaploidy OR octoploid source: "Heredity"
Annals of Botany	Ploidy hybrid genetic introgression diploid OR tetraploid OR hexaploidy OR octoploid source: "Annals of Botany"
American Journal of Botany	Ploidy hybrid genetic introgression diploid OR tetraploid OR hexaploidy OR octoploid source:" American Journal of Botany"
New Phytologist	Ploidy hybrid genetic introgression diploid OR tetraploid OR hexaploidy OR octoploid source:" New Phytologist"
PNAS	Ploidy hybrid genetic introgression diploid OR tetraploid OR hexaploidy OR octoploid source:" PNAS"
Biological Journal of the Linnean Society	Ploidy hybrid genetic introgression diploid OR tetraploid OR hexaploidy OR octoploid source:" Biological Journal of the Linnean Society"
Botanical Journal of the Linnean Society	Ploidy hybrid genetic introgression diploid OR tetraploid OR hexaploidy OR octoploid source:" Botanical Journal of the Linnean Society"
Journal of Evolutionary Biology	Ploidy hybrid genetic introgression diploid OR tetraploid OR hexaploidy OR octoploid source:" Journal of Evolutionary Biology"
PLoS One	Ploidy hybrid genetic introgression diploid OR tetraploid OR hexaploidy OR octoploid source:" PLoS One"

Appendix 2: Chapter 3

Figure 2: Trees with root nodes containing the highest and lowest posterior mean probability of hybridisation from Model 1 (BLUP's of nodes in the phylogeny). A is the top tree (subset of Orchidaceae) whilst B is the tree with lowest probability of hybridisation (Fabaceae and Polygalaceae).

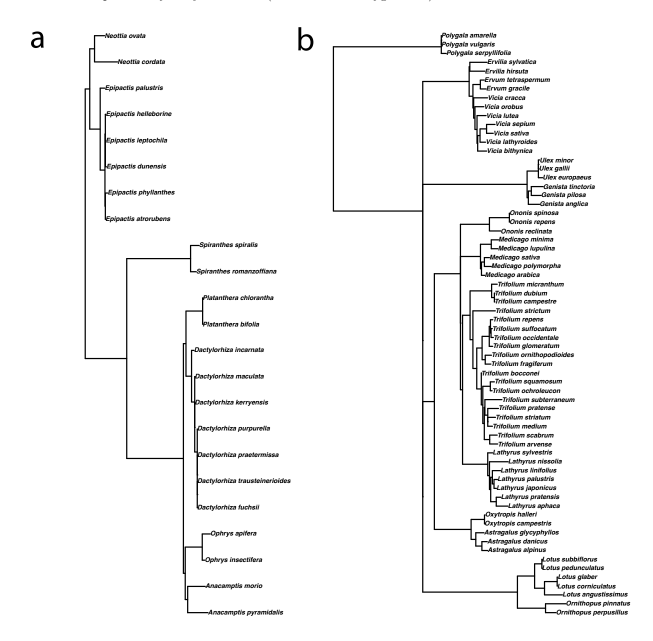


Figure 3: The joint probability of hybridisation between two parental species give both branch length between species (tree based genetic distance) and geographical overlap between parental species (measured as overlap in occupancy of 10x10 km grid squares in the UK). The degree of shading in the scale bar and tiles represent the posterior probability of hybridisation from Model 1 given parameter values for each variable. Estimates are visualised at mean genus size, for annual-perennial parental combinations and accounting for phylogenetic relationships between species.

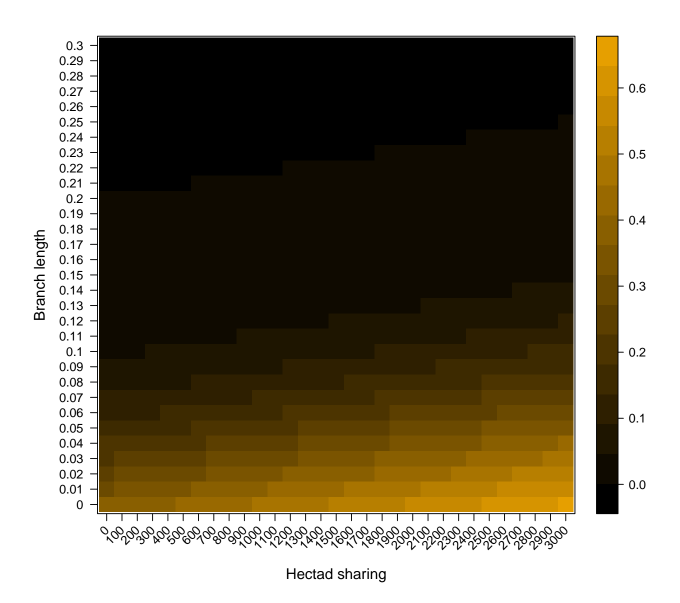


Figure 4: Predicted fit of probability of hybridisation given hectad sharing and ploidy difference of parental species from Model 2. Dashed lines indicate the 95% Credible Intervals, and the bold lines represent the posterior mode of the coefficients of congeneric pairs of species hybridising as a function of pairwise overlap in distribution, conditional on parental ploidy status. The effect is visualised at mean genetic distance for annual-perennial parent combinations and accounting for phylogenetic effects. The bold red dashed line indicates mean pairwise overlap in distribution (10x10km²).

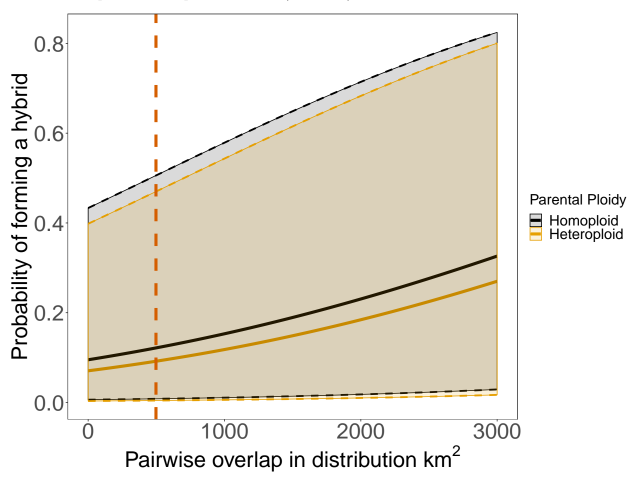


Figure 5: Predicted fit of probability of hybridisation given branch length between parental species and ploidy difference of parental species from Model 2. Homoploid indicates parental species of the same ploidy level, and heteroploidy indicates parental species of different ploidy levels. Dashed lines indicate the 95% Credible Intervals, and the bold lines represent the posterior mode of the coefficients of congeneric pairs of species hybridising as a function of pairwise branch length, conditional on parental ploidy status. The effect is visualised at mean hectad sharing for annual-perennial parent combinations and accounting for phylogenetic effects. The red dashed line indicates mean pairwise branch length between all pairs of species.

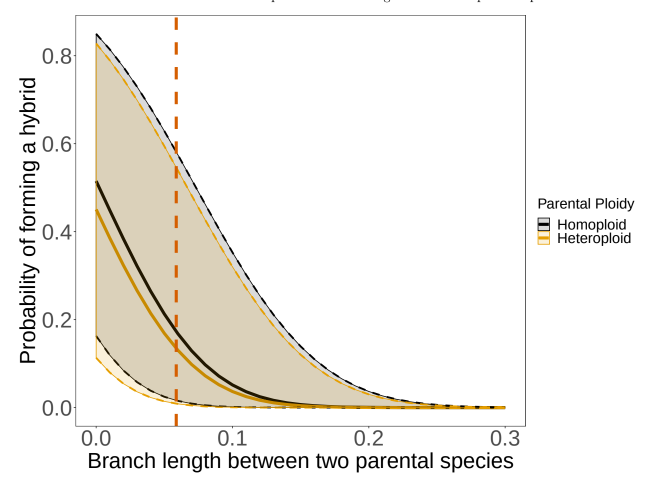


Figure 6: Predicted fit of probability of hybridisation given branch length between parental species from Model 1. Black dashed lines are the 95% Credible Intervals, bold line is the posterior mean of the coefficient for the probability of congeneric pairs of species hybridising as a function of branch length. This effect is visualised at mean hectad sharing, for annual-perennial parent combinations and accounting for phylogenetic effects. The bold red dashed line indicates mean genus level genetic distance between pairs of species.

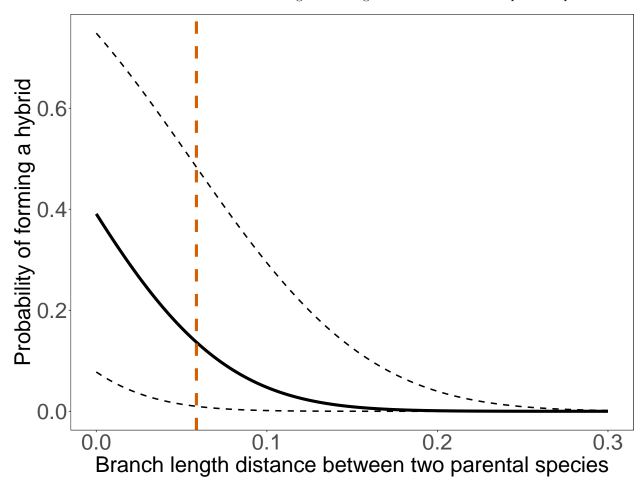


Table 2: Model 1: Probability of hybridisation on the probit scale with genetic distance, hectads shared and life history of parental species as fixed covariates. The posterior mean of the distribution of each coefficient is given, along with lower and upper 95% Credible Intervals. The p-value (pMCMC) is also reported and given in bold where significant. Annual-perennial and perennial-perennial levels are jointly tested using a Wald test in the main text.

Covariate	Posterior mean	l-95% CI	u-95% CI	Effective sample size	pMCMC
(Intercept)	-1.31	-3.76	0.60	1000	0.22
Branch length between species pairs	-59.75	-66.69	-51.98	185.41	0.0010
Hectads shared between species pairs	0.001	0.0007	0.0012	1000	0.0010
Annual-perennial parent pair	-0.12	-0.97	0.66	1000	0.76
Perennial-perennial parent pair	0.64	-0.25	1.58	1000	0.16
Genus size	-0.0014	-0.041	0.031	1107	0.92

Table 3: Phylogenetic signal of probability of hybridisation and the species variance independent of phylogenetic effects on the probit scale. 95% Credible Intervals of the variances are also presented. See Methods in Chapter 3 for calculation.

Variance Component	Posterior Mode	Lower Credible Interval	Upper Credible Interval
Model 1 Phylogenetic Variance	0.62	0.32	0.77
Model 1 Species Variance	0.33	0.18	0.58
Model 2 Phylogenetic Variance	0.61	0.30	0.82
Model 2 Species Variance	0.34	0.084	0.44

Table 4: Model 2: Probability of hybridisation on the probit scale with ploidy, genetic distance, hectads shared and life history of parental species as covariates. The posterior mean of the distribution of each coefficient is given, along with lower and upper 95% Credible Intervals. The p-value (pMCMC) is also reported and given in bold where significant.

Covariate	Posterior mean	l-95% CI	u-95% CI	Effective sample size	pMCMC
(Intercept)	-0.11	-2.41	2.12	813	0.93
Branch length between species pairs	-74.93	-88.38	-63.46	319	0.0010
Crosss ploidy effect	-0.73	-1.02	-0.40	1000	0.0010
Hectads shared between species pairs	0.0013	0.0009	0.0016	883	0.0010
Annual-perennial parent pair	0.093	-1.15	1.23	836	0.89
Perennial-perennial parent pair	0.82	-0.40	1.96	836	0.16
Genus size	-0.029	-0.084	0.031	621	0.32

Appendix 3: Chapter 4

Figure 7: Distributions of parameters fitted to the models with constant gene flow (blue), secondary contact (green), and without gene flow (grey) in the demographic simulation software, $\delta a \delta I$. The model with historic gene flow (orange) had poor AIC values, and so is omitted here.

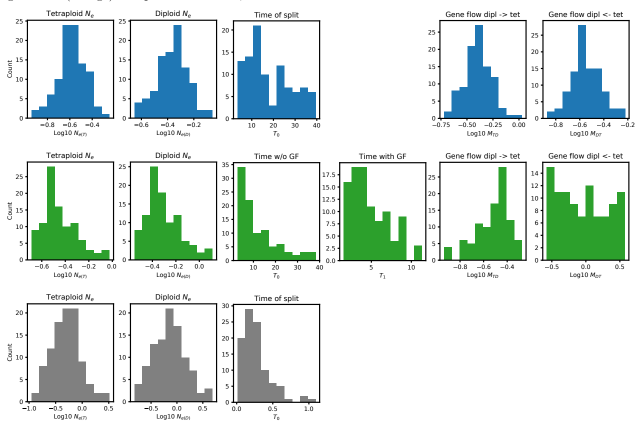


Table 5: Primers and PCR conditions used to amplify the rpL32- $trnL_{\rm UAG}$ plastid marker in Euphrasia species.

Primer	Orientation	Sequence (5'-3')	Reagents (1 reaction)	PCR conditions	References
rpL32-F	Forward	CAGTTCCAAAAAAACGTACTTC	$12.5\mu\mathrm{M}$ Taq $2\mathrm{X}$ Master Mix, $0.5\mu\mathrm{L}$ Bovine Serum Albumen, $0.5\mu\mathrm{L}$ forward and reverse primers at $10\mu\mathrm{M}$, $10.5\mu\mathrm{L}$ water, $1\mu\mathrm{L}$ sample DNA	5 min at 94°C, 35× (30 s at 94°C, 45 s at 50°C, 40 s at 72°C), 5 min at 72°C	(Wang et al., 2018)
$trnL_{\mathrm{UAG}}$	Reverse	CTGCTTCCTAAGAGCAGCGT			

Table 6: Primers and PCR conditions used to amplify the ITS1 nuclear marker in Euphrasia species.

Primer	Orientation	Sequence (5'-3')	Reagents (1 reaction)	PCR conditions	References
ITS4	Forward	TCCTCCGCTTATTGATATGC	$12.5\mu\mathrm{M}$ Taq $2\mathrm{X}$ Master Mix, $0.5\mu\mathrm{L}$ Bovine Serum Albumen, $0.5\mu\mathrm{L}$ forward and reverse primers at $10\mu\mathrm{M}$, $10.5\mu\mathrm{L}$ water, $1\mu\mathrm{L}$ sample DNA	5min at 94°C, 30 x (30s at 94°C, 30s at 54°C, 2min at 72°C), 10 min at 72°C.	(Wang et al., 2018)
ITS5	Reverse	${\tt GGAAGTAAAAGTCGTAACAAGG}$	•		

Appendix 4: Chapter 6

Table 7: Host species used in the common garden experiment in Chapter 6. The species along with the taxonomic family they belong to, their ecological functional group and the source of the seeds are also given. Commercial seed stocks list the original collection where known.

Common name	Species name	Family	Functional group (informal)	Seed source
	Species name	· · · · · · · · · · · · · · · · · · ·		
Thale cress	$Arabidopsis\ thaliana$	Brassicaceae	Herb	Laboratory stock
Field horsetail	$Equisetum\ arvense$	Equisetaceae	Fern	Wild collected in
				Edinburgh (GPS
				coordinates: 55.9679,
D 16		To.		-3.2129)
Red fescue	Festuca rubra	Poaceae	Grass	Commerical:
				Emorsgate seeds
T. 1 1	TT 1 1 .	To the state of th		(Yorkshire + Dorset)
Yorkshire fog	$Holcus\ lanatus$	Poaceae	Grass	Commerical:
		3.5		Emorsgate seeds
Common liverwort	$Marchantia\ polymorpha$	Marchantiaceae	Bryophyte	Wild collected in
				Edinburgh (GPS
				coordinates: 55.9679,
D.1		T01	TT 1	-3.2129)
Ribwort plantain	$Plantago\ lanceolata$	Plantaginaceae	Herb	Commerical:
				Emorsgate seeds
~ .		.	_	(Somerset + Wiltshire)
Scots pine	$Pinus\ sylvestris$	Pinaceae	Tree	Commerical: Scotia
			-	Seeds
White clover	$Trifolium\ repens$	Fabaceae	Herb	Commerical:
				Emorsgate seeds
				(Yorkshire + Wiltshire)

Table 8: Collection details for Euphrasia species used in the common garden experiment. *Population also used in the multiple host phenotypic plasticity experiment.

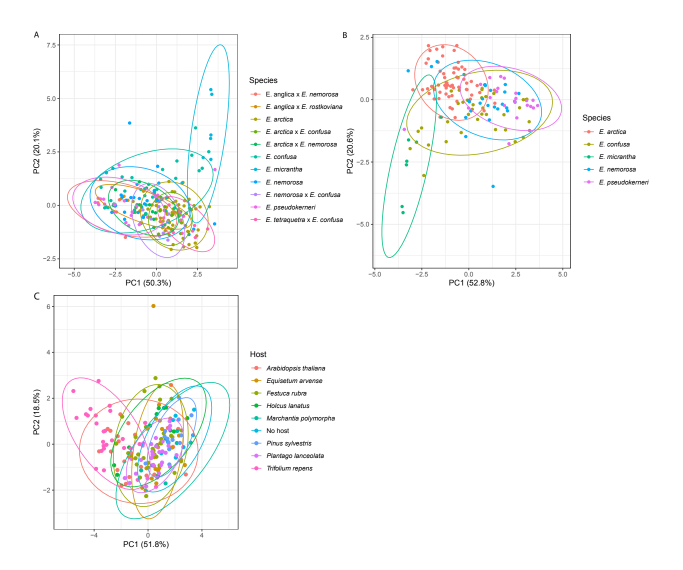
Collection number	Taxon	Locality	Latitude	Longitude	Collector
E4E0138	E. arctica	Fintallick, Glen Ledock, Comrie, Perthshire	56.41318	-4.03085	Dot Hall
E4E0144	E. arctica	Balachuirn, Isle of Raasay	57.38996	-6.06877	S.J. Bungard
E4E0032	E. arctica	South Links, Burray, Orkney	58.85275	-2.88701	John Crossley
E4E0139	E. arctica	Dalreoch Farm, Enochdhu	56.74199	-3.53350	Martin Robinson
E4E0049	E. arctica	Ouaisne, Jersey	49.17707	-2.18293	Anne Haden
E4E0247	E. arctica	Elsdon. Newcastle upon Tyne	55.22770	-2.10234	Stephanie Miles
NBer001*	E. arctica	North Berwick Glenn, East Lothian	56.05696	-2.70456	Alex Twyford
E4E0038	E. confusa	Oldbury, near Hartshill, Warwickshire	52.55285	-1.53980	John and Monika Walton
E4E0114	E. confusa	Trethew Mill, Bodmin, Cornwall 50.39585	-4.709558	Rosemary Parslow	
E4E0095	E. confusa	North Anston Grassland, South Yorkshire	53.34738	-1.20803	Graeme Coles
E4E0009	$E.\ confusa$	Devil's Hole Blowout, Ravenmeols Local Nature Reserve, Merseyside	53.54062	-3.09041	Philip H. Smith
E4E0188	E. micrantha	Meall a Bathaich, Glen Garry, East Perthshire	56.82082	-4.182812	Alistair Godfrey
E4E0064	E. nemorosa	Castle Hill Local Nature Reserve, East Sussex	50.7842	0.052719	David Harris

Collection number	Taxon	Locality	Latitude	Longitude	Collector
E4E0069	E. nemorosa	Meridian Business Park, Leicester	52.60857	-1.19809	Geoffrey Hall
E4E0123	E. nemorosa	Bloody Oaks Triangle, Tickercote, Rutland	52.68950	-0.56263	Geoffrey Hall
E4E0029	E. pseudokerneri	Levin Down, Sussex	50.91346	-0.74150	Elizabeth Sturt
E4E0112	E. pseudokerneri	Beeston Common, Norfolk	52.93442	1.220071	Francis Farrow
E4E0027	$E. \ anglica \times E. \ nemorosa$	West Dean Woods, Sussex	50.93212	-0.79735	Elizabeth Sturt
E4E0016	$E.\ anglica \times E.\ rostkoviana$	Straduff Rathcabbin, Co. Tipperary	53.11902	-8.02454	David Nash
E4E0033	$E. \ arctica \times E. \ confusa$	Nr Quoyorally, South Ronaldsay, Orkney	58.75897	-2.93473	John Crossley
E4E0145	$E. \ arctica \times E. \ nemorosa$	Kylfakin, Wof, Skye	57.26685	-5.76042	S.J. Bungard
E4E0021	$E. \ arctica \times E. \ nemorosa$	Dunamase, Co. Laois	53.03153	-7.21015	David Nash
E4E0031	$E.\ nemorosa \times E.\ confusa$	Dolebury Fort, Somerset	51.32605	-2.79432	C.W. Hurfurt
E4E0143	$E. \ tetraquetra \times E. \ confusa$	Ballyteige Burrow, Co Wexford, Ireland	52.20268	-6.64325	Jim Hurley

Table 9: Summary of trait values for many Euphrasia species and hybrids grown on a clover host (i.e. the species differences experiment). Values are means ± 1 SE. Length measurements are in millimeters. Note: Date of first flower not recorded.

Taxon	Corolla length	Height	Internode ratio	Julian days to flower	Lower floral leaf teeth	Nodes to flower	Number of branches
E. arctica	8.0 ± 0.2	82.9 ± 4.4	1.1 ± 0.1	195.2 ± 1.5	4.4 ± 0.1	8.6 ± 0.2	$*4.56 \pm 0.2$
E. confusa	6.9 ± 0.2	134.4 ± 7.2	1.6 ± 0.1	200.2 ± 2.4	5.3 ± 0.2	11.1 ± 0.4	7.26 ± 0.5
E. micrantha	5.6 ± 0.2	70.6 ± 8.1	3.0 ± 0.4	_	2.4 ± 0.3	8.3 ± 0.2	0.57 ± 0.4
$E.\ nemorosa$	7.7 ± 0.1	127.4 ± 8.1	1.4 ± 0.1	206.6 ± 1.7	5.1 ± 0.2	11.9 ± 0.5	7.67 ± 0.5
E. pseudok-	8.8 ± 0.4	176.4 ± 15.6	1.4 ± 0.1	205.1 ± 2.0	5.5 ± 0.2	13.2 ± 0.4	8.67 ± 0.6
erneri							
$E. \ anglica \ \mathbf{x}$	9.1 ± 0.5	148.1 ± 11.8	1.4 ± 0.1	195.7 ± 1.9	6.0 ± 0.3	12.0 ± 0.6	10.00 ± 1.0
E. nemorosa							
$E. \ anglica \ \mathbf{x}$	7.9 ± 0.2	122.6 ± 8.3	1.3 ± 0.1	192.3 ± 12.3	5.9 ± 0.3	10.6 ± 0.5	7.44 ± 0.7
E.							
rostkoviana							
$E. \ arctica \ \mathbf{x}$	9.5 ± 0.2	100.3 ± 4.3	1.4 ± 0.1	193.4 ± 3.2	3.8 ± 0.1	7.8 ± 0.3	5.70 ± 0.4
$E.\ confusa$							
$E. \ arctica \ \mathbf{x}$	8.0 ± 0.2	132.2 ± 14.5	1.3 ± 0.1	205.3 ± 2.4	6.0 ± 0.3	11.3 ± 0.4	6.50 ± 0.4
$E.\ nemorosa$							
$E. \ arctica \ \mathbf{x}$	7.9 ± 0.2	92.5 ± 5.9	1.0 ± 0.1	199.3 ± 2.8	5.1 ± 0.2	9.8 ± 0.3	7.00 ± 0.5
$E.\ nemorosa$							
$E.\ confusa$ x	7.2 ± 0.2	57.4 ± 5.8	0.7 ± 0.1	194.1 ± 2.7	4.2 ± 0.2	7.6 ± 0.4	4.00 ± 0.3
E. tetraquetra							

Figure 8: Principal component analysis of morphological variation of *Euphrasia* in a common garden. Panels show (A) five species and six hybrids grown with a single clover host, (B) five species grown with a clover host omitting hybrids, and (C) *E. arctica* with nine host treatments. Points represent individuals, and ellipses represent the standard error of the (weighted) average of scores.



Table(s) 10: The first five principal components extracted from the principal component analysis, with the contribution of variance of each trait to each principal component. The last two rows of each table show the standard deviation and the proportion of variance explained by the principal component.

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Species differences (including hybrids)	PC1	PC2	PC3	PC4	PC5
Branches	0.229	0.053	0.071	0.252	0.094
Corolla length	0.089	0.262	0.369	0.032	0.136
Height	0.211	0.115	0.149	0.047	0.379
Internode ratio	0.005	0.441	0.186	0.030	0.190
Leaf teeth	0.213	0.056	0.097	0.428	0.128
Nodes to flower	0.224	0.093	0.126	0.181	0.081
Standard deviation	1.738	1.099	0.964	0.616	0.533
Proportion of variance	0.503	0.201	0.155	0.063	0.047

Species differences (excluding hybrids)	PC1	PC2	PC3	PC4	PC5
Branches	0.226	0.024	0.096	0.233	0.017
Corolla length	0.100	0.269	0.361	0.082	0.141
Height	0.214	0.128	0.151	0.063	0.367
Internode ratio	0.029	0.434	0.202	0.000	0.171
Leaf teeth	0.214	0.032	0.064	0.424	0.159
Nodes to flower	0.217	0.113	0.125	0.198	0.145
Standard deviation	1.780	1.111	0.932	0.612	0.433
Proportion of variance	0.528	0.206	0.145	0.062	0.031

Phenotypic plasticity	PC1	PC2	PC3	PC4	PC5
Branches	0.183	0.065	0.032	0.098	0.220
Corolla length	0.139	0.001	0.252	0.340	0.030
Height	0.179	0.150	0.016	0.065	0.128
Internode ratio	0.070	0.301	0.274	0.119	0.146
Julian days to flower	0.158	0.198	0.056	0.077	0.191
Leaf teeth	0.178	0.024	0.090	0.153	0.166
Nodes to flower	0.093	0.262	0.280	0.147	0.119
Standard deviation	1.904	1.137	0.924	0.725	0.586
Proportion of variance	0.518	0.185	0.122	0.075	0.049

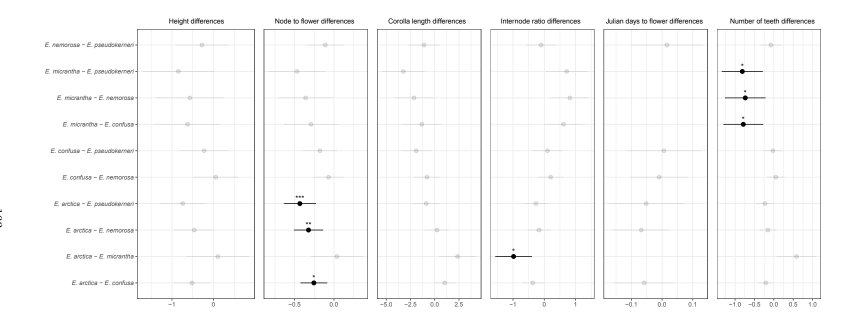


Table 11: Summary of trait values for $Euphrasia\ arctica$ grown on many different hosts. Values are mean +/- one standard error. Length and height measurements are in millimeters.

Early season	At first flowering							End of season	
Host	Height	Corolla length	Height	Internode ratio	Julian days to flower	Lower floral leaf teeth	Nodes to flower	Number of branches	Height
A. thaliana	12.8 ± 1.1	6.1 ± 0.3	19.2 ± 1.6	2.4 ± 0.1	201.6 ± 4.3	3.2 ± 0.1	8.8 ± 0.3	2.1 ± 0.4	30.0 ± 3.2
E. arvense	6.1 ± 0.4	5.9 ± 0.3	15.1 ± 1.1	2.6 ± 0.2	215.3 ± 4.6	2.4 ± 0.1	9.3 ± 0.3	0.4 ± 0.1	35.6 ± 4.8
F. rubra	6.7 ± 0.4	6.3 ± 0.1	19.5 ± 1.4	2.6 ± 0.2	$\begin{array}{c} 216.5\ \pm \\ 4.4 \end{array}$	2.8 ± 0.2	9.6 ± 0.3	0.8 ± 0.3	39.6 ± 4.1
H. lanatus	7.1 ± 1.3	6.3 ± 0.1	16.0 ± 1.6	2.4 ± 0.2	224.5 ± 7.0	2.5 ± 0.2	9.8 ± 0.4	0.8 ± 0.4	33.8 ± 6.8
M. polymorpha	6.3 ± 0.7	5.5 ± 0.4	9.6 ± 1.3	2.9 ± 0.4	222.6 ± 17.0	1.7 ± 0.3	9.7 ± 0.5	0	11.3 ± 2.5
No host	5.9 ± 0.3	5.3 ± 0.2	11.2 ± 1.1	2.8 ± 0.2	241.3 ± 7.9	1.9 ± 0.3	9.9 ± 0.5	0	9.7 ± 2.0
P. lanceolata	7.5 ± 0.5	6.1 ± 0.1	14.1 ± 0.8	2.8 ± 0.1	211.2 ± 3.7	2.9 ± 0.1	10.4 ± 0.3	0.4 ± 0.1	28.3 ± 3.4
P. sylvestris	6.2 ± 0.6	5.7 ± 0.3	12.2 ± 1.3	2.9 ± 0.2	233.8 ± 6.1	1.9 ± 0.2	9.2 ± 0.3	0	17.2 ± 2.6
T. repens	12.9 ± 1.4	7.4 ± 0.2	39.4 ± 2.6	2.1 ± 0.2	189.8 ± 2.0	3.9 ± 0.1	8.7 ± 0.3	4.7 ± 0.4	143.2 ± 8.6

Table 12: Comparison of *E. arctica* traits in the phenotypic plasticity common garden experiment. Tukey comparisons are presented between *E. arctica* traits with two different host treatments. Point estimates are the mean difference of the comparison, calculated from the phenotypic plasticity model using the emmeans R package. *** p < 0.001, ** p < 0.01, * p < 0.05.

Host 1	Host 2	Corolla length	Height	Internode ratio	Julian days to flower	Nodes to flower	Number of leaf teeth
Arabidopsis thaliana	No host	1.065*	0.49*	0.102	-0.178***	-0.116	0.508
Equisetum arvense	No host	0.946*	0.49° 0.304	0.102 0.041	-0.112***	-0.110 -0.066	0.508 0.212
Festuca rubra	No host	1.04*	0.504	0.041 0.073	-0.112***	-0.000 -0.034	0.212 0.397
Holcus lanatus	No host	1.04*	0.329	$0.075 \\ 0.077$	-0.112	-0.054 -0.009	0.397 0.258
Marchantia		0.25		0.077	-0.005 -0.031		-0.136
polymorpha	No host	0.25	-0.181			-0.025	
Pinus sylvestris	No host	0.481	0.067	0.015	-0.03	0.051	0.01
$Plantago\ lanceolata$	No host	0.879	0.246	0.016	-0.137***	-0.071	0.419
Trifolium repens	No host	2.102***	1.241***	0.180*	-0.244***	-0.134	0.711
$Equisetum\ arvense$	$Arabidopsis\ thaliana$	-0.119	-0.186	-0.061	0.066*	0.05	-0.296
$Festuca\ rubra$	$Arabidopsis\ thaliana$	-0.024	0.039	-0.029	0.065**	0.082	-0.111
$Holcus\ lanatus$	$Arabidopsis\ thaliana$	-0.015	-0.158	-0.025	0.114***	0.107	-0.25
Marchantia	$Arabidopsis\ thaliana$	-0.815	-0.671***	-0.032	0.147***	0.091	-0.644
polymorpha							
Pinus sylvestris	$Arabidopsis\ thaliana$	-0.584	-0.423*	-0.087	0.148***	0.167	-0.498
$Plantago\ lanceolata$	$Arabidopsis\ thaliana$	-0.186	-0.244	-0.086	0.041	0.044	-0.089
Trifolium repens	$Arabidopsis\ thaliana$	1.037***	0.751***	0.077	-0.066	-0.018	0.204
$Festuca\ rubra$	$Equisetum\ arvense$	0.095	0.225	0.031	0	0.032	0.185
$Holcus\ lanatus$	$Equisetum\ arvense$	0.104	0.028	0.035	0.049	0.057	0.046
Marchantia polymorpha	$Equisetum\ arvense$	-0.696	-0.486*	0.029	0.081	0.041	-0.348
Pinus sylvestris	Equiportum amuma	-0.465	-0.237	-0.025	0.082**	0.117	-0.202
· ·	Equisetum arvense Equisetum arvense	-0.465 -0.067	-0.257 -0.059	-0.025	-0.025	-0.006	0.202
Plantago lanceolata Trifolium repens	Equisetum arvense Equisetum arvense	1.156***	-0.059 0.937***	0.138*	-0.025 -0.132***	-0.068	0.207
Holcus lanatus	Festuca rubra	0.01	-0.197	0.138 0.003	0.049	0.025	-0.139
Marchantia	Festuca rubra Festuca rubra	-0.79	-0.197 -0.71***	-0.003	0.049 0.081	0.025 0.009	-0.139 -0.533
polymorpha	restuca ruota	-0.79	-0.71	-0.002	0.081	0.009	-0.000
Pinus sylvestris	$Festuca\ rubra$	-0.56	-0.462**	-0.057	0.083**	0.085	-0.387
Plantago lanceolata	$Festuca\ rubra$	-0.161	-0.283	-0.056	-0.025	-0.038	0.022
Trifolium repens	$Festuca\ rubra$	1.062***	0.712***	0.106	-0.132***	-0.1	0.315
$Marchantia \ polymorpha$	Holcus lanatus	-0.8	-0.513*	-0.006	0.033	-0.016	-0.394

Host 1	Host 2	Corolla length	Height	Internode ratio	Julian days to flower	Nodes to flower	Number of leaf teeth
Pinus sylvestris	Holcus lanatus	-0.569	-0.265	-0.061	0.034	0.06	-0.248
Plantago lanceolata	$Holcus\ lanatus$	-0.171	-0.086	-0.06	-0.074**	-0.063	0.161
Trifolium repens	$Holcus\ lanatus$	1.052**	0.909***	0.102	-0.18***	-0.125	0.454
Pinus sylvestris	$Marchantia \ polymorpha$	0.231	0.248	-0.055	0.001	0.076	0.146
Plantago lanceolata	$Marchantia \ polymorpha$	0.629	0.427	-0.054	-0.106***	-0.047	0.555
Trifolium repens	$Marchantia \ polymorpha$	1.852***	1.423***	0.109	-0.213***	-0.109	0.847*
Plantago lanceolata	Pinus sylvestris	0.398	0.178	0.001	-0.107***	-0.123	0.409
Trifolium repens	Pinus sylvestris	1.621***	1.174***	0.164*	-0.214***	-0.185	0.701*
Trifolium repens	Plantago lanceolata	1.223***	0.996***	0.163*	-0.107***	-0.063	0.292

Table 13: Analysis of deviance for each trait in the phenotypic plasticity experiment with E. arctica grown with many different hosts, assuming a Poisson distribution. For each model, we report the change in degrees of freedom (df), deviance, residual degrees of freedom, residual deviance, and p-value generated from the χ^2 distribution. Factor host, where the model includes all host species, is compared to the intercept model where no hosts are fitted.

Trait	Factor	df	Deviance	Resid. df	Resid. Dev	$\Pr(>\chi^2)$
Julian days to flower	Host (Intercept)	8	192.390 192	184 611.5053	419.1153	2.56E-37
Nodes to flower	Host (Intercept)	8	5.020 193	185 43.49272	38.47252	0.755416
Number of leaf teeth	Host (Intercept)	8	26.793 193	185 68.17096	41.37748	0.000767

Table 14: ANOVAs for traits measured in the phenotypic plasticity experiment with *E. arctica* grown with many different hosts, assuming Gaussian distributed residuals. For each model, we report the degrees of freedom (df), sums of squares (SS), mean squares (MS), F-statistic, and p-value.

Trait	Factor	df	SS	MS	F	p
Corolla length	Host Residuals	8 173	49.469 108.555	6.184 0.6275	9.854565	3.00E-11
Height	Host Residuals	8 185	27.021 27.009	$3.378 \\ 0.146$	23.139	2.52E-24
Internode ratio	Host Residuals	8 184	$0.562 \\ 3.845$	$0.070 \\ 0.0209$	3.362213	0.001275

Table 15: Summary of generalised linear models for the phenotypic plasticity experiment with *Euphrasia arctica* grown on many hosts in a common garden. All models compare *E. arctica* grown with a particular host to the intercept of no host. Generalised linear models assuming Poisson residuals with log link function were used in Julian days to flower, nodes to flower and number of leaf teeth, while all others assumed Gaussian residuals. The model coefficient is reported with standard error in brackets. *** p < 0.001, ** p < 0.05.

				Julian days to		Number of leaf
Term	Corolla length	Height (log)	Internode ratio	flower	Nodes to flower	teeth
(Intercept)	5.250 (0.250)***	2.363 (0.115)***	0.353 (0.043)***	5.489 (0.02)***	2.293 (0.095)***	0.646 (0.218)**
Arabidopsis thaliana	1.064 (0.293)***	0.489 (0.135)***	0.102 (0.051)*	-0.177 (0.024)***	-0.115 (0.114)	0.507 (0.241)*
$Equisetum\\ arvense$	0.945 (0.300)**	0.304 (0.138)*	$0.041\ (0.052)$	-0.111 (0.024)***	-0.065 (0.116)	$0.212 \ (0.254)$
Festuca rubra	1.040 (0.288)***	0.529 (0.134)***	$0.073\ (0.050)$	-0.112 (0.023)***	-0.033 (0.111)	$0.396 \ (0.242)$
$Holcus\ lanatus$	1.050 (0.323)**	0.331 (0.147)*	0.077(0.055)	-0.063 (0.025)*	-0.008 (0.123)	0.257 (0.267)
Marchantia polymorpha	0.250 (0.433)	-0.181 (0.171)	0.070 (0.064)	-0.03 (0.029)	-0.024 (0.143)	-0.135 (0.338)
Pinus sylvestris	$0.480 \ (0.333)$	0.067 (0.153)	0.015 (0.058)	-0.029 (0.026)	$0.051 \ (0.126)$	$0.010 \ (0.290)$
Plantago lanceolata	0.879 (0.288)**	0.245 (0.134)	$0.016 \ (0.05)$	-0.136 (0.023)***	-0.071 (0.112)	0.419 (0.242)
Trifolium repens	2.101 (0.293)***	1.241 (0.136)***	0.180 (0.051)***	-0.243 (0.024)***	-0.133 (0.115)	0.711 (0.239)**

Figure 10: Relationship between growth-related traits and end of season height for *E. arctica* grown with eight hosts and no host. (A) height at first flowering, (B) height 6-weeks after germination, (C) Julian days to flower, (D) number of branches. Length measurements are reported in mm.

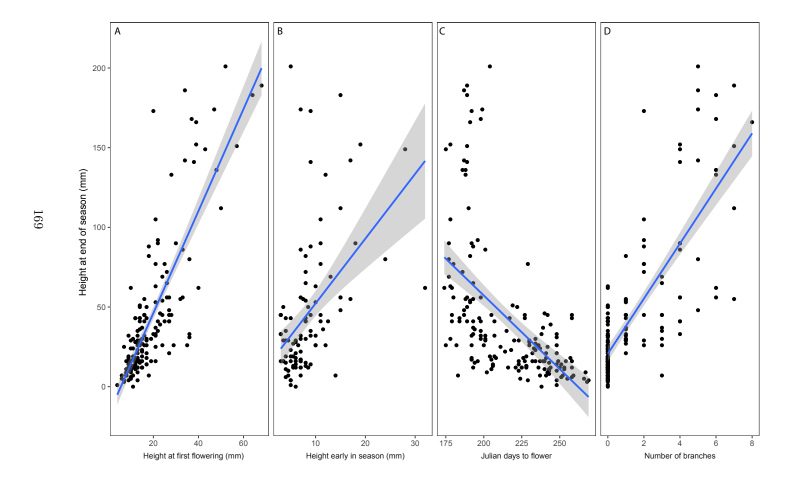


Figure 11: Comparison between trait values for wild-collected herbarium specimens and common garden plants of diverse *Euphrasia* species for (A) nodes to flower, (B) corolla length (mm), (C) number of leaf teeth, (D) internode ratio. Points are for *Euphrasia* population means, with bars representing the standard error of measurements.

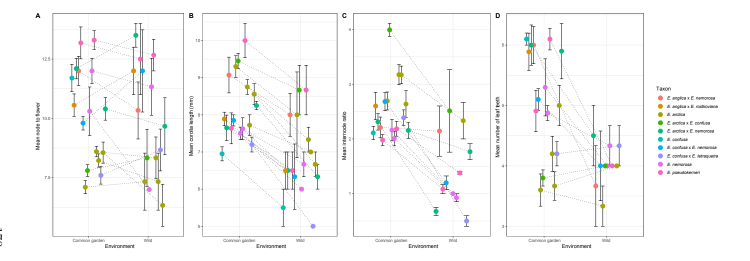


Table 16: Model output from MCMCglmm comparing traits for the wild collected *Euphrasia* specimens to the baseline of the common garden data (Intercept). The posterior means are reported along with the lower and upper 95% credible intervals, as well as the p-value (pMCMC) for the effect.

Trait	Factor	Posterior mean	Lower credible interval	Upper credible interval	$_{ m pMCMC}$
Branches	(Intercept) Wild	1.863 -0.457	1.682 -0.619	2.086 -0.290	0.001 0.001
	collected				
Internode ratio	(Intercept)	2.533	2.118	2.920	0.001
	Wild collected	-1.008	-1.206	-0.823	0.001
Corolla	(Intercept)	8.182	7.477	8.756	0.001
	Wild collected	-1.363	-1.650	-1.032	0.001
Nodes	(Intercept)	2.322	2.189	2.465	0.001
	Wild collected	-0.016	-0.135	0.086	0.800
Teeth	(Intercept)	1.616	1.485	1.722	0.001
	Wild collected	-0.187	-0.369	-0.004	0.050

Appendix 5: Chapter 7

Figure 12: Euphrasia reproductive output over time showing differences in reproductive trajectories, data from Experiment 1. Values represent mean reproductive nodes at a particular time point \pm one standard error. Eleven species of host are shown, along with the average host where points are the mean of all hosts in the experiment.

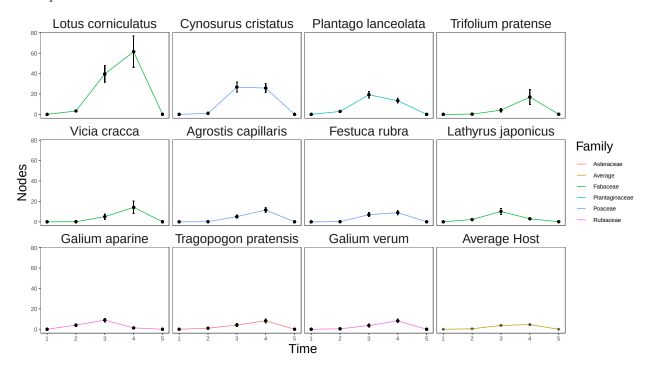


Figure 13: Posterior distributions of the phylogenetic signal for the models from Experiment 1, where 45 different host species were grown with *Euphrasia arctica*. The distributions of phylogenetic signal are shown for three *Euphrasia* traits: survival, total reproductive output at the end of the season, and days to flower. Total reproductive output shows both the highest and least variable estimate of phylogenetic signal, however all are significant as the distributions are not overlapping zero.

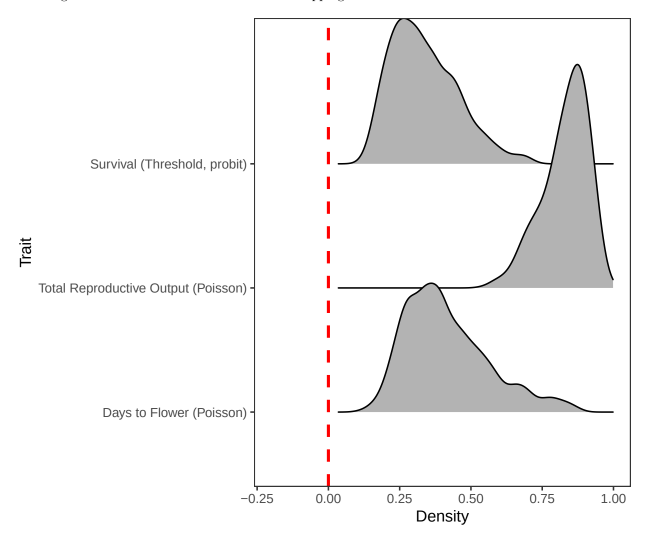


Figure 14. Posterior distribution of the variance for random effects in the model fitted for Experiment 2, where four species of *Euphrasia* were grown on thirteen different species of host. The random effects are the *Euphrasia*-host interaction, the sole effect of host species, and the residual variance. Although the residual variance is the explaining most variation, both the host-parasite interaction and hosts themselves are estimated to be significantly way from zero.

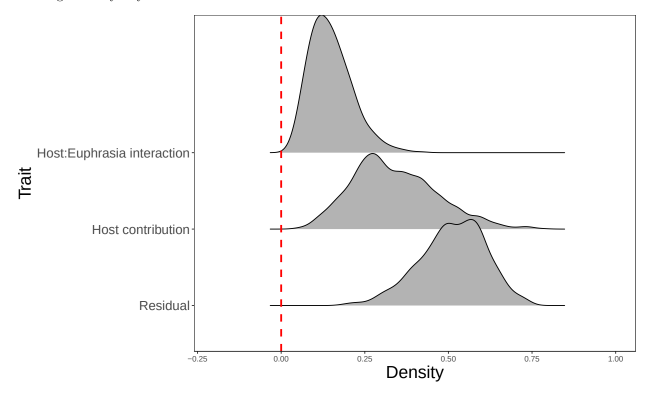


Figure 15. Performance of four species of Euphrasia on thirteen different species of host plants measured as cumulative reproductive nodes. Each panel represents a unique Euphrasia population (a = A1766, b = T1761, c = V1761, d = M1767, e = M1768, f = M1769), coloured by species. Two populations, (e) and (f) co-occur. Host species are ranked by average performance conferred to a Euphrasia species, where HPU = $Hypericum\ pulchrum$, CVU = $Calluna\ vulgaris$, HLA = $Holcus\ lanatus$, OVU = $Origanum\ vulgare$, UGA = $Ulex\ gallii$, PMA = $Plantago\ maritima$, PLA = $Plantago\ lanceolata$, VCH = $Veronica\ chamaedrys$, FOV = $Veronica\ chamaedrys$, FOV

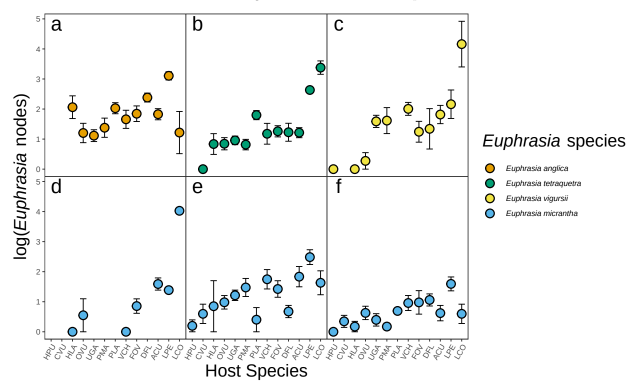


Table 17: Plant names, attributes and collection sources for host species used in Experiment 1.

Host species	Authority	Functional group	Life History	Seed source
No host	-	-	-	-
Agrostis capillaris	L.	Grass	Perennial	Emorsgate
Allium ursinum	L.	Forb	Perennial	RBGE
Anthriscus	(L.) Hoffm.	Forb	Perennial	Emorsgate
sylvestris				
Arabidopsis	(L.) Heynh.	Forb	Annual	Inbred lines
thaliana				University of
				Edinburgh
Centaurea nigra	L.	Forb	Perennial	Emorsgate
Centranthus ruber	(L.) DC.	Forb	Perennial	Chiltern Seeds
Chenopodium	L.	Forb	Annual	Author collections
album				
Chenopodium	L.	Forb	Perennial	Surplus seed
bonus-henricus				$\overline{\mathrm{RBGE}}$
Cynosurus	L.	Grass	Perennial	Emorsgate
cristatus				
Cystopteris	R. Sim	Fern	Perennial	RBGE
dickeniana				
Dactylorhiza	(T.Stephenson &	Forb	Perennial	RBGE
purpurella	T.A.Stephenson)			
• •	Soó			
Equisetum arvense	L.	Fern	Perennial	RBGE
Erica tetralix	L.	Woody	Perennial	RBGE
$Festuca\ rubra$	L.	Grass	Perennial	Emorsgate
Fragaria vesca	L.	Forb	Perennial	Scotia seeds
Galanthus nivalis	L.	Forb	Perennial	RBGE
Galium aparine	L.	Forb	Annual	Author collection,
				Upper Halliford,
				Surrey, Engalnd,
				11/16
$Galium\ verum$	L.	Forb	Perennial	Emorsgate
Helianthemum	(L.) Mill.	Forb	Perennial	Scotia seeds
nummularium	,			
Holcus lanatus	L.	Grass	Perennial	Emorsgate
Hordeum vulgare	L.	Grass	Annual	Wiggly Wigglers
Hyac in tho ides	(L.) Chouard ex	Forb	Perennial	RBGE
non- $scripta$	Rothm.			
Lagurus ovatus	L.	Grass	Annual	www.wildflowershop.co.uk
Lathyrus japonicus	Willd.	Legume	Perennial	RBGE
Leucanthemum	(Vaill.) Lam.	Forb	Perennial	Emorsgate
vulgare	, ,			~
Lotus corniculatus	L.	Legume	Perennial	Emorsgate
Meum	Jacq.	Forb		$\overline{\mathrm{RBGE}}$
at ham anticum	_			
$Mimulus\ guttatus$	DC.	Forb	Perennial	Author collections
· ·				

Host species	Authority	Functional group	Life History	Seed source
Ononis spinosa	L.	Legume	Perennial	Emorsgate & Wild Flower Shop
Papaver rhoeas	L.	Forb	Annual	Emorsgate
Phleum pratense	L.	Grass	Perennial	Wild Flower Shop
Pinus sylvestris	L.	Woody	Perennial	Scotia seeds
Plantago lanceolata	L.	Forb	Perennial	Emorsgate
Pteridium aquilinum	L. (Kuhn)	Fern	Perennial	British Pteridological Society spore exchange
$Rumex\ acetosella$	L.	Forb	Perennial	Scotia seeds
Senecio vulgaris	L.	Forb	Annual	RBGE
Silene dioica	(L.) Clairv.	Forb	Perennial	D. Charlseworth, Univ. Edinburgh
Silene latifolia	Poir.	Forb	Perennial	D. Charlseworth, Univ. Edinburgh
Thymus polytrichus	A.Kern. ex Borbás	Woody	Perennial	Emorsgate
Sorbus aucuparia	L.	Woody	Perennial	RBGE
Tragopogon pratensis	L.	Forb	Perennial	Scotia seeds
Trifolium pratense	L.	Legume	Perennial	Chiltern Seeds & Wild Flower Shop
Ulex europaeus	L.	Legume/Woody	Perennial	Tree Seed Online Ltd
Vicia cracca	L.	Legume	Perennial	Emorsgate
$Zea\ mays$	L.	Grass	Annual	Chiltern Seeds

Table 18: Plant names, attributes and collection sources for host species used in Experiment 2.

Host species	Authority	Source/Location	Plant status
Agrostis curtisii	Kerguélen	Millenium Seed Bank, Kew Gardens	Seed
Calluna vulgaris	(L.) Hull	RBGE	Seed, but small plants from cuttings
Deschampsia (Avenella) flexuosa	(L.) Trin.	Chiltern Seeds	Seed
Festuca ovina	Ĺ.	Emorsgate	Seed
Holcus lanatus	L.	Emorsgate	Seed
Hypericum pulchrum	L.	Scotia Seeds	Seed
Lotus corniculatus	L.	Emorsgate	Seed
Lolium perenne	L.	Emorsgate	Seed
Origanum vulgare	L.	Emorsgate	Seed
Plantago lanceolata	L.	Emorsgate	Seed
Plantago maritima	L.	Scotia Seeds	Seed
Ulex gallii	Planch.	Millenium Seed Bank, Kew Gardens	Seed
Veronica chamaedrys	L.	Scotia Seeds	Seed

Table 19: Euphrasia species collections across both experiments in Chapter 7.

Experiment	Euphrasia species	Location	Grid Reference
1	E. arctica	Inverkeithing, Scotland	NT 1389 82312
2	E. anglica	(A1766)	Cheddar, Somerset
2	E. vigursii	(V1761)	St Agnes Head, Cornwall
2	$E.\ tetraquetra$	(T1761)	St Agnes Head, Cornwall
2	$E.\ micrantha$	(M1767)	Borrowdale, Cumbria
2	$E.\ micrantha$	(M1768)	Alness, Scotland
2	E. micrantha	(M1769)	Orkney, Scotland

Table 20: Model output from MCMCglmm for the event history analysis (survival) model in Experiment 1. The intercept represents the latent probit estimate of mean *Euphrasia* survival on a perennial grass transplanted at the earliest date, measured at the first time point. The posterior means are reported along with the lower and upper 95% credible intervals as well as the effective sample size and p-value for the effect (pMCMC).

Covariates	Posterior mean	l-95% CI	u-95% CI	Effective sample size	pMCMC
(Intercept)	3.0348	1.8630	4.1519	1000	< 0.001
Time	-1.0533	-1.1164	-0.9912	1000	< 0.001
AnnPerAnn	0.1390	-0.2489	0.6076	1000	0.5300
Normalised transplant date	-0.0164	-0.0213	-0.0117	1000	< 0.001
Functional group fern	-0.2583	-1.5117	1.0171	1000	0.6520
Functional group forb	-0.3076	-0.9687	0.3844	1000	0.3700
Functional group legume	-0.0828	-1.0457	0.7646	1000	0.8500
Functional group woody	-0.6675	-1.4986	0.1819	1000	0.0980

Table 21: Model output from MCMCglmm for the days to flower model in Experiment 1. The intercept represents the log of the mean days to flower since germination of *Euphrasia* on a perennial grass transplanted at the earliest date. The posterior means are reported along with the lower and upper 95% credible intervals as well as the effective sample size and p-value for the effect (pMCMC).

Covariates	Posterior mean	l-95% CI	u-95% CI	Effective sample size	pMCMC
(Intercept)	4.6197	4.1765	5.0536	1000	< 0.001
AnnPerAnn	-0.1380	-0.2703	0.0043	1188	0.0560
Functional group fern	-0.1127	-0.5410	0.3556	1000	0.6000
Functional group forb	-0.0879	-0.3087	0.1793	1106	0.3780
Functional group legume	-0.0650	-0.3307	0.3032	860.9	0.6160
Functional group woody	0.0991	-0.2964	0.4466	1000	0.5520
Normalised transplant date	0.0034	0.0008	0.0060	1000	0.0160

Table 22: Model output from MCMCglmm for the number of reproductive nodes over time model in Experiment 1. The intercept represents log of the mean number of reproductive nodes of *Euphrasia* on a perennial grass transplanted at the earliest date, measured at the first time point. The posterior means are reported along with the lower and upper 95% credible intervals as well as the effective sample size and p-value for the effect (pMCMC).

Covariates	Posterior mean	l-95% CI	u-95% CI	Effective sample size	pMCMC
(Intercept)	-4.1298	-17.0773	5.4805	550	0.3420
Time3	2.3713	1.5862	3.2031	773.2	< 0.001
Time4	3.0630	2.1378	3.9166	1000	< 0.001
AnnPerAnn	0.7872	-1.2385	2.8500	1000	0.4460
Functional group fern	-4.3612	-16.8977	6.6709	789.8	0.3960
Functional group forb	-2.3178	-9.4309	3.7584	793.8	0.4420
Functional group legume	-2.3657	-10.7235	5.1473	756.9	0.5760
Functional group woody	-7.6673	-15.5032	-1.0839	549.4	0.0180
Normalised transplant date	-0.0760	-0.0919	-0.0625	1000	< 0.001
Time3:AnnPerAnn	-0.9448	-2.0965	0.1002	1000	0.0920
Time4:AnnPerAnn	-2.3383	-3.6057	-0.8897	1000	0.0040

Table 23: Model output from MCMCglmm for the cumulative reproductive nodes at the end of the season model in Experiment 1. The intercept represents the log of the mean cumulative reproductive nodes at the end of the season of *Euphrasia* on a perennial grass transplanted at the earliest date. The posterior means are reported along with the lower and upper 95% credible intervals as well as the effective sample size and p-value for the effect (pMCMC).

Covariates	Posterior mean	l-95% CI	u-95% CI	Effective sample size	pMCMC
(Intercept)	-0.4637	-9.8823	9.4058	1093	0.9240
AnnPerAnn	-0.3610	-2.9028	2.1730	886.5	0.7720
Functional group fern	-3.6600	-15.1134	6.8501	1000	0.4660
Functional group forb	-2.9965	-8.8016	2.1653	1097	0.2340
Functional group legume	-2.0488	-9.1675	4.6899	1000	0.5500
Functional group woody	-7.5786	-14.1020	-1.0165	633.3	0.0100
Normalised transplant date	-0.0762	-0.0945	-0.0570	1000	< 0.001

Table 24: Model output from MCMCglmm for the number of cumulative reproductive nodes of *Euphrasia* individuals at the end of the season from Experiment 2. The intercept represents log of the mean cumulative number of reproductive nodes of *Euphrasia anglica*, population A1766, on a host that was transplanted at the earliest date. The posterior means are reported along with the lower and upper 95% credible intervals as well as the effective sample size and p-value for the effect (pMCMC).

Covariates	Posterior mean	l-95% CI	u-95% CI	Effective sample size	pMCMC
(Intercept)	1.7842	1.2210	2.2714	787.7	0.0010
$Euphrasia\ micrantha$	-1.2795	-1.7479	-0.8284	1000	0.0010
$Euphrasia\ tetraquetra$	-0.3702	-0.8160	-0.0076	873.2	0.0620
$Euphrasia\ vigursii$	-0.2457	-0.7758	0.2138	1000	0.3340
Population: M1767	0.3269	-0.2098	0.9299	846.7	0.2760
Population: M1768	0.7931	0.4788	1.0699	1000	0.0010
Normalised transplant date	0.0059	-0.0084	0.0237	1208	0.4820