

Supporting Information for: Ecological drivers of flower-leaf sequences: aridity and pollination success select for flowering-first in the American Plums

Tables

species	n FLS	n petal length	n pdsi
alleghaniensis	17	39	114
americana	95	271	200
angustifolia	77	238	200
gracilis	85	289	200
hortulana	106	254	200
maritima	75	255	200
mexicana	64	284	200
munsoniana	117	279	200
nigra	118	230	200
rivularis	111	225	200
subcordata	46	71	30
texana	19	38	39
umbellata	70	284	200

Table S1: Sample sizes of each for each species used in this study

species	index BBCH 0-11	index BBCH 0-11 w/o day of season	index BBCH 0-09
mexicana	0.85	0.90	0.69
umbellata	0.82	0.83	0.58
angustifolia	0.76	0.77	0.51
maritima	0.68	0.76	0.42
gracilis	0.64	0.68	0.36
americana	0.62	0.55	0.38
munsoniana	0.60	0.67	0.38
alleghaniensis	0.59	0.65	0.32
nigra	0.55	0.62	0.30
hortulana	0.51	0.52	0.28
texana	0.51	0.54	0.25
rivularis	0.44	0.53	0.22
subcordata	0.16	0.18	0.05

Table S2: Hysteranthly index score for 13 species in the American plums based on model predictions defining flowering during BBCH stages 0-11 as hysteranthis for models with an without day of season of observation as a co-variate, and with hysteranthis defined as flowering during BBCH 0-09 only.

	Estimate	Est.Error	Q5.5	Q25	Q75	Q94.5
Intercept	0.34	0.23	-0.02	0.20	0.48	0.70
phi_Intercept	1.92	0.42	1.22	1.65	2.21	2.55
pdsi.z	-0.47	0.30	-0.96	-0.66	-0.28	0.01
petal.z	-0.14	0.24	-0.54	-0.29	0.01	0.23
pdsi.z:petal.z	-0.14	0.49	-0.91	-0.46	0.16	0.65

	Estimate	Est.Error	Q5.5	Q25	Q75	Q94.5
Intercept	0.49	0.25	0.09	0.33	0.65	0.88
phi_Intercept	1.77	0.41	1.09	1.50	2.06	2.39
pdsi.z	-0.43	0.32	-0.92	-0.63	-0.22	0.07
petal.z	-0.14	0.27	-0.56	-0.30	0.03	0.27
pdsi.z:petal.z	-0.16	0.54	-1.01	-0.50	0.17	0.69

Table S3: Model parameter estimates of the relationship between environmental/trait predictors and hys-
teranthly index based on hys-teranthly index derived from a model included day of season of observation (top
table) and one that excluded this predictor from the index derivation (bottom table)

Figures

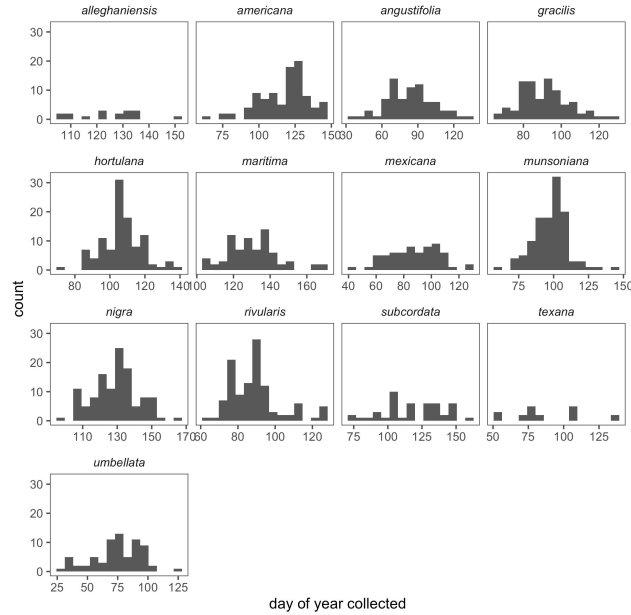


Figure S1: Histograms of collect day of year for each of the American plum species used in these analyses

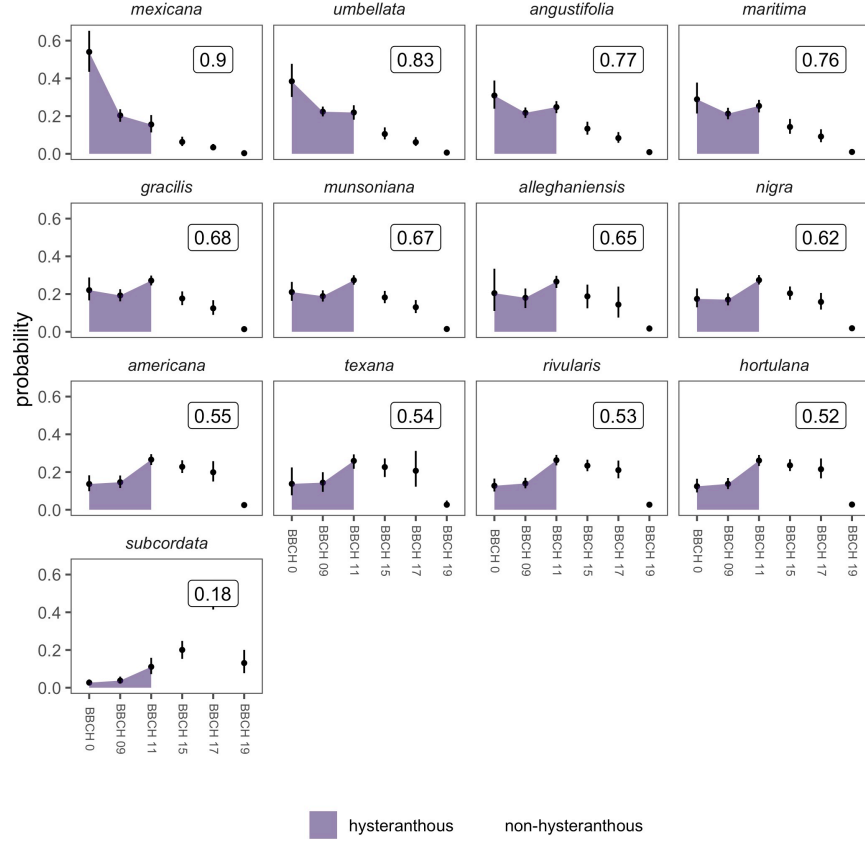


Figure S2: Predicted likelihood that a species would be in flower during each vegetative BBCH phase for each species in the American plums. Points are the mean likelihood while bars represent 89% uncertainty intervals. Hysteresis index values in each box were derived from the summed likelihood species would be found at BBCH stages 0-11 (purple fill).

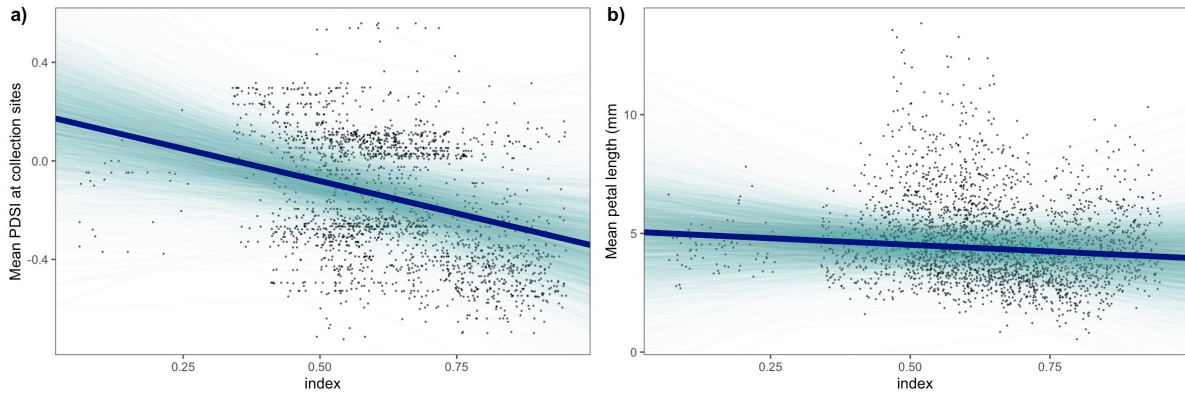


Figure S3: Relationships between the hysteresis index scores and environmental and biological traits based on Bayesian phylogenetic mixed models where the relationship between hysteresis score and PDSI a) and mean petal length b) were each modeled separately. Solid lines indicate the mean posterior estimate and shaded areas 4000 draws from the posterior distribution as a display of uncertainty. The points are jittered along the x-axis only for visibility. This plots complements the model presented in Fig. 3

Extended Methods:

To complement our main analyses in which we fit models based on species-level means of PDSI and petal length as predictors, we also implemented additional models where we assessed the relationship between PDSI and hysternathy, and the relationship between petal length and hysternathy separately. With this model structure our analyses accounted for the intra-specific variation in both PDSI and petal length as the phylogenetic structure of these variables.

In these models, we modeled species and phylogeny as above.

The model structure is:

$$y_{trait} = \alpha + \alpha_{sp} + \alpha_{phylo} + \beta_{hyst.index} * X_{hyst.index} + \epsilon$$

$$\epsilon \sim N(0, \sigma_y^2)$$

where y_{trait} is observed trait values (PDSI or petal length), and the slope $\beta_{hyst.index}$ describes the relationship between extended hysternathy (higher hysternathy index value) and the trait of interest. α describes a grand intercept, and α_{sp} and α_{phylo} describe the species and phylogenetic effects respectively.