Online supplementary material

Hybridization as a threat in climate relict *Nuphar pumila* (Nymphaeaceae)

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Online resource 1. Primers and PCR amplification conditions. The primer name and sequences, fluorophore, multiplex pool, annealing temperature, PCR temperature ramps, template DNA dilution (applied for limiting the effects of PCR inhibitors), allele size range, number of alleles and publication are detailed for each marker used in the study.

Marker name	Primer forward (5'-3')	Primer reverse (5'-3')	Fluorophore	Multiplex	T. Anneal	T. ramp	DNA dil.	Size range	N.alleles	Reference
NLGA2	CTTTAGGAGGGTCTTTAGCC	CCAATCTCTAGTAGGAGGAGC	ATTO	п	52°C	5°C/s	dil 1/20	93-123	12	Ouborg et al. (2000)
NLGA3	GTTGTAACGTAAATGCCGTCC	CTTGCCGATGAAACCCAT	ATTO	Ι	55°C	5°C/s	dil 1/20	99-183	14	Ouborg et al. (2000)
NLGA5	CCCGCCATATCTGATGAC	AAGTGGAGGGGGACGAAAG	HEX	I	52°C	5°C/s	dil 1/20	70-100	5	Ouborg et al. (2000)
NLGA7	ATTTATTCCCAGCACTTTGG	CTTGACATGATTTCTCTGAACC	HEX	п	52°C	2°C/s	dil 1/5	58-104	12	Ouborg et al. (2000)
NLCA1	CTCAGAAACGAGGCTCTATG	TTTGGTTGGAAGACAAGAAG	FAM	п	52°C	5°C/s	dil 1/20	182-242	15	Ouborg et al. (2000)
NLTG/GA1	AAGCAGCAGCAAAATTTGTA	TGTGCAAGTTACCTGTTTCC	FAM	п	52°C	5°C/s	dil 1/20	117-135	8	Ouborg et al. (2000)
Nsub033	ACACACACACACTCTCTCTCTC	ACTTGCAAAGATCCTCTCAGAT	ATTO	I	57°C	5°C/s	dil 1/10	222-241	6	Yokogawa et al. (2008)
Nsub176	AGAGAGAGAGAGAGACACACACAC	GGCAACAGGTCTATTAATCTCA	FAM	I	57°C	2°C/s	dil 1/5	91-146	12	Yokogawa et al. (2008)

Online resource 2. Morphological and ecological differences between *Nuphar pumila* (specialist) and *N. lutea* (generalist).

Trait/attribute	N. pumila	N. lutea
Morphology		
Stigmatic disc diameter Stigmatic disc form Stigmatic rays Flower diameter Perigon length Fruit length Fruit form Floating leaves Petiole length Petiole form (under the blade) Rhizome diameter	6-8.8 mm deeply lobed 6-13 2-3 cm 1-2 cm 1-3 cm slightly curved, grooved 10 x 12 cm 50-150 (350) cm compressed 1-2 cm	10-20 mm entire 12-25 4-5 cm 2-3 cm 2-4 cm straight, not grooved 30 x 40 cm 50-250 (500) cm trigonous 3-8 cm
Ecology		
Flowering period Waterbodies/habitat Tolerance to water movement Tolerance to wave action Tolerance to salinity Water trophic level Water temperature Water pH Substrates Maximal depth of water Maximum length of peduncle Maximum altitude in Europe Distribution southwards	VI-VIII stagnant no no dystrophic to mesotrophic cool slightly acidic mainly over mud or peat 3.5 m 2 m ca. 1700 m a.s.l. to the Alps	VII-IX stagnant to slowly flowing yes yes moderately eutrophic wide amplitude wide amplitude wide amplitude (also sand and gravel) 6 m 4 m ca. 1000 m a.s.l. to Northern Africa



PCA axis 1 (18.7% variance explained)

Online resource 3. Principal coordinates analysis of individual genotypes. Our sampling includes 194 N. pumila specimens (small pie charts) collected in 13 natural populations, completed with 20 N. lutea specimens (large pie charts) from natural populations (KES - 15 specimens, STI -2 specimens) and botanical gardens (LAU - 3 specimens). Distances among specimens are computed according to their genotype, as characterized by 8 SSR loci. Allele sizes are not accounted in distance calculations. In parallel, we display the admixture levels of specimens, estimated with the "hybrid index" (Buerkle 2005), using pie-charts. Every specimen is assigned either to N. lutea (black) or N. pumila (white) genetic pool using a probabilistic framework; pure breed specimens receive a probability of 0 (N. lutea) or 1 (N. pumila) while first generation hybrids and further admixed genotypes get intermediate probabilities. Note that several hybrid specimens appear as differentiated from both N. pumila and N. lutea genotypes (i.e. appearing with negative coordinates on the PCA1 and PCA2 eigenaxes). Those hybrids actually carry alleles with SSR sizes suggesting a *N. lutea* origin, that were however not directly observed in pure *N. lutea* specimens. This pattern arises due to the limited amount of specimens sampled for N. lutea. It should be noted that Hindex accommodates such data limitations by excluding the alleles being absent from both reference pools (i.e. representing here 28 alleles / 84 in total).



Online resource 4. Model selection. A model selection procedure was used to test for the presence of introgression among N. lutea and N. pumila (i.e. model 1) versus a null hypothesis (model 0) assuming no gene flow among species. The procedure started by estimating the posterior probability of each model [P(model | empirical data)], using a pairwise model comparison where 500'000 simulations were performed for each model (using priors and parameters defined as in the main manuscript) and compared to the empirical data. The 1'000 best simulations, collected from the compared models, were then used to estimate posterior probabilities [i.e. P(model | empirical data), using neural-nets implemented in the "abc" R CRAN package, with 50 iterations]. Cross-validations were then used to evaluate the robustness of this selection procedure. Briefly, 1'000 simulations were randomly picked from each of the compared models and used as pseudo-observations to feed the model selection procedure. This allowed to check i) whether the compared models could be discriminated from each other and ii) check how often posterior probabilities were designating either the correct [i.e. in blue on density plots, P(model X | model X), true positives] or the wrong model [i.e. in red on density plots, P(model X | model Y), false positives]. These counts then allowed computing p-values, indicating the risk of picking the wrong model, at a given posterior probability value. The p-value was estimated as $Pval(model X) = 1 - D_{post.prob.emp}[P(model X |$ model X)] / $(D_{post.prob.emp}[P(model X | model X)] + D_{post.prob.emp}[P(model X | model Y)])$, where D_{post.prob.emp} is the density of cross-validations picking a given model, at the posterior probability obtained with empirical observations.



Online resource 5. Cross-validation of ABC estimates. We used 1,000 simulations as pseudoobservations to assess the robustness and accuracy of our ABC estimations. The results are displayed as scatterplots, where the estimated parameters (i.e. Estimated value) are compared to those that were actually used to set the simulations (i.e. True value). The median and 95% confidence intervals of the obtained estimates as displayed as solid and dashed lines, respectively. These curves outline the level of technical uncertainty and systematic bias (i.e. deviations from the 1:1 gray line) associated to the ABC procedure itself. **a**) - **d**) Note that most of our estimations actually overestimate the actual parameter values (i.e. the median lines are shifted left compared to the 1:1 line), at least when considering the parameter space close to the empirical estimates (i.e. "empir. est.", displayed as a red line). It is therefore likely that our empirical estimations suffer from this same systematic bias. Hence, bias-corrected values (i.e. "correct est." - displayed as the bluedashed line), obtained by intersecting of our empirical estimates with the median line, were considered for further discussion. **e**) – **f**) The cross-validations indicate that our ABC procedure yields essentially random estimations for these two parameters.



Online resource 6. Morphological differences between *N. pumila* and *N. lutea*. Leaves and rhizomes of *N. pumila* (A, C) and *N. lutea* (B, D). Bar: 5 cm. Plants obtained from the Botanic Garden of the University of Fribourg (*N. pumila*) and Botanic Garden of Lausanne (*N. lutea*). Photographs by Hans-Rüdiger Siegel (Natural History Museum Fribourg, Switzerland).