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Thirteen New Plastid Genomes from Mixotrophic and Autotrophic Species Provide Insights into Heterotrophy Evolution in Neottieae Orchids

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Abstract

Mixotrophic species use both organic and mineral carbon sources. Some mixotrophic plants combine photosynthesis and a nutrition called mycoheterotrophy, where carbon is obtained from fungi forming mycorrhizal symbiosis with their roots. These species can lose photosynthetic abilities and evolve full mycoheterotrophy. Besides morphological changes, the latter transition is associated with a deep alteration of the plastid genome. Photosynthesis-related genes are lost first, followed by housekeeping genes, eventually resulting in a highly reduced genome. Whether relaxation of selective constraints already occurs for the plastid genome of mixotrophic species, which remain photosynthetic, is unclear. This is partly due to the difficulty of comparing plastid genomes of autotrophic, mixotrophic, and mycoheterotrophic species in a narrow phylogenetic framework. We address this question in the orchid tribe Neottieae, where this large assortment of nutrition types occurs. We sequenced 13 new plastid genomes, including 9 mixotrophic species and covering all 6 Neottieae genera. We investigated selective pressure on plastid genes in each nutrition type and conducted a phylogenetic inference of the group. Surprisingly, photosynthesis-related genes did not experience selection relaxation in mixotrophic species compared with autotrophic relatives. Conversely, we observed evidence for selection intensification for some plastid genes. Photosynthesis is thus still under purifying selection, maybe because of its role in fruit formation and thus reproductive success. Phylogenetic analysis resolved most relationships, but short branches at the base of the tree suggest an evolutionary radiation at the beginning of Neottieae history, which, we hypothesize, may be linked to mixotrophy emergence.

Key words: Mycorrhiza, mycoheterotrophy, mixotrophy, phylogeny, Neottieae, plastome.

Introduction

Mixotrophic organisms grow by using both inorganic and organic carbon resources, either simultaneously or successively,

and therefore represent intermediates between autotrophy and heterotrophy (Selosse et al. 2017; Těšitel et al. 2018). Land plants widely associate with fungi through mycorrhizal

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Table 1
Length, GC Content, and Genes Lost of the 13 Neottieae Plastomes Sequenced

Species ^a	Accession	Length				GC Content				Gene Loss ^b
		Total	LSC	SSC	IR	Total	LSC	SSC	IR	<i>Pseudogene/Undetected</i>
<i>P. pabstii</i>	MH590357	163,909	90,710	18,823	27,188	37.3	35	31	43.3	
<i>Ce. damasonium</i>	MH590345	161,699	88,720	19,085	26,947	37.3	35.1	30.7	43.2	
<i>Ce. longibracteata</i>	MH590346	161,986	88,888	19,138	26,980	37.2	35	30.6	43.1	
<i>Ce. rubra</i>	MH590347	162,277	88,814	19,199	27,132	37.2	35	30.6	43.1	
<i>E. albensis</i>	MH590348	159,763	87,237	18,782	26,872	37.3	35.1	30.7	43.2	
<i>E. atrorubens</i>	MH590349	159,790	87,237	18,803	26,875	37.3	35.1	30.7	43.3	
<i>E. gigantea</i>	MH590350	158,977	87,101	18,664	26,606	37.3	35.2	30.8	43.2	
<i>E. helleborine</i>	MH590351	159,822	87,313	18,785	26,862	37.3	35.1	30.7	43.2	
<i>E. microphylla</i>	MH590352	159,236	86,706	18,784	26,873	37.4	35.2	30.7	43.3	<i>ndhD, ndhE, ndhF, ndhH</i>
<i>E. palustris</i>	MH590353	159,134	87,114	18,702	26,659	37.4	35.2	30.8	43.2	
<i>E. purpurata</i>	MH590354	159,864	87,246	18,786	26,916	37.3	35.1	30.7	43.2	
<i>L. abortivum</i>	MH590355	128,822	85,544	15,099	27,102	36.3	35.1	30.4	43.1	<i>ndhA, ndhB, ndhC, ndhD, ndhE, ndhF, ndhG, ndhH, ndhI, ndhJ, ndhK, cemaA</i>
<i>N. cordata</i>	MH590356	147,034	82,416	12812	25903	37.5	35.1	29.3	43.4	<i>ndhA, ndhB, ndhC, ndhD, ndhE, ndhF, ndhG, ndhH, ndhI, ndhJ, ndhK</i>

^aComplete genera names are *Palmorchis*, *Cephalanthera*, *Epipactis*, *Limodorum*, and *Neottia*.

^bA gene was presumed pseudogenized if a frameshift indel resulted in a reduction in protein length higher than 25% of its original length.

symbiosis and usually exchange their photosynthates for mineral nutrients (Smith and Read 2008). Repeatedly, plants evolved the ability to gain organic compounds also from their mycorrhizal fungi, a mixotrophic nutrition also called partial mycoheterotrophy (Selosse and Roy 2009; Hynson et al. 2013; Těšitel et al. 2018). We hereafter use the word mixotrophy for these plants that combine photosynthetic and mycorrhizal carbon sources. In many mixotrophic lineages from different families, species eventually lost photosynthesis and became fully mycoheterotrophic (Leake 1994; Merckx, Freudenstein, et al. 2013). Traits of mycoheterotrophy such as chlorophyll loss, reduction in leaf surface area, development of short clumpy roots, or production of reserve-less “dust” seeds have been first studied (Merckx, Mennes, et al. 2013). An increasing amount of molecular data now clarifies the genetic traits linked with the evolution to mycoheterotrophy.

The plastid genomes (or plastomes) of 24 fully mycoheterotrophic species have been sequenced to date (Graham et al. 2017; Petersen et al. 2018). Their comparative analysis has resulted in a model of sequential plastid genome degradation associated with mycoheterotrophy, which parallels what has been observed for heterotrophic parasitic plants (Funk et al. 2007; Barrett and Davis 2012; Barrett et al. 2014; Wicke et al. 2016; Graham et al. 2017). Genes encoding subunits of the nicotinamide adenine dinucleotide H dehydrogenase (NDH)-like complex, which regulates excessive electron flow in the plastidial electron transfer chain, are the first to be lost, followed by most photosynthesis-related genes. Rubisco large subunit (*rbcl*), adenosine triphosphate (ATP)-synthase, and plastid-encoded RNA polymerase (PEP) genes are often

retained in the early stage of photosynthesis loss. Eventually, degradation encompasses housekeeping genes involved in plastid translation and other specific functions. This model summarizes well how the plastid genome evolves in mycoheterotrophic and parasitic plant as they become more and more dependent on organic carbon uptake (Wicke et al. 2016; reviewed in Graham et al. [2017] and Wicke and Naumann [2018]).

Due to the few data available, still little is known about what occurs in the evolutionary steps prior to mycoheterotrophy, that is, in mixotrophic species that retain photosynthetic abilities. Whether these species already experience relaxed selective pressure on photosynthesis remains unclear in the general case. This is a challenging question because of the scarcity of lineages that have retained together autotrophic, mixotrophic, and mycoheterotrophic species. Known examples include the genus *Burmannia* L. (Burmanniaceae; Bolin et al. 2017), the tribe Pyroleae (Ericaceae; Lallemand et al. 2016), and the tribe Neottieae (Orchidaceae; Selosse and Roy 2009; Gonneau et al. 2014). In the leafless genus *Corallorhiza* Gagnebin (Orchidaceae), where no full autotroph occurs, comparison of plastomes from mixotrophic (green) and mycoheterotrophic (nongreen) species showed very limited gene loss in the former and relaxed selective constraints on photosynthesis and ATP-synthase genes in the latter (Barrett et al. 2014). Consistently, the selective regime does not differ between mixotrophic *Corallorhiza* and autotrophic outgroups (Barrett et al. 2014). The absence of autotrophic *Corallorhiza* as reference yet prevents firm conclusions regarding selective regime in mixotrophs. In the orchid genus *Cymbidium* Sw., *C. macrorhizon* is leafless but still holds

chlorophyll in the stem and fruits and the importance of its photosynthesis for fruiting has recently been demonstrated (Suetsugu et al. 2018). The plastome of this mixotrophic species shows no gene loss except for a few NDH genes and is mostly under purifying selection (Kim et al. 2018). The two genera *Corallorhiza* and *Cymbidium* where photosynthesis already undergoes relaxed selection in mixotrophic species (Barrett et al. 2014) call for extensive analyses in other lineages.

We sequenced plastomes of mixotrophs in Neottieae, an orchid tribe mostly found in northern hemisphere forests. Neottieae include 6 genera and about 200 species (WCSP 2018) with various nutrition types: *Palmorchis* Barb. Rodr. (all putative autotrophs), *Neottia* Guett. (many mycoheterotrophs and some probable mixotrophs; Těšitelová et al. 2012; Yagame et al. 2016; Schiebold et al. 2018), *Epipactis* Zinn (numerous mixotrophs and some autotrophs; Lallemand et al. 2018), *Cephalanthera* Rich. (some mycoheterotrophs and many mixotrophs), *Limodorum* Boehm. (all mixotrophs; Girlanda et al. 2006), and *Aphyllorchis* Blume (all mycoheterotrophs; Roy et al. 2009). Early studies on mycoheterotrophic and mixotrophic Neottieae contributed to our understanding of the ecophysiology and evolution of plants feeding on fungal carbon (Selosse et al. 2002; Gebauer and Meyer 2003; Julou et al. 2005). The transition to mixotrophy in this group is consistently linked to a shift in type of mycorrhizal fungi (supplementary table S1, Supplementary Material online). Autotrophic orchids associate with a polyphyletic group of saprotrophic and endophytic fungi, the rhizoctonias (Dearnaley et al. 2012), whereas mixotrophic Neottieae use ectomycorrhizal fungi from different genera, which simultaneously form mycorrhizas with surrounding trees (Selosse and Martos 2014). For the purpose of this study, we had to assign a given nutrition type to each species investigated. Even if there is a continuum between autotrophy and mixotrophy (Jacquelyn et al. 2017), we decided to consider a species as mixotrophic when it dominantly associated with ectomycorrhizal fungi and showed an enrichment in ^{13}C compared with surrounding autotrophs, a usual marker of fungal carbon gain (Hynson et al. 2013; see supplementary table S1, Supplementary Material online, for references regarding species nutrition).

Plastome sequencing of some Neottieae species by Feng et al. (2016) has yielded important results: 1) new hypotheses regarding the controversial phylogenetic relationships between Neottieae genera (Bateman et al. 2005; Roy et al. 2009; Górniak et al. 2010; Xiang et al. 2012), 2) patterns of gene loss in mycoheterotrophs matching the model described above, and 3) no relationship between the chlorophyll level (presence or absence based on visual estimation) and the plastome degradation in the leafless *Neottia* clade. Their sampling, however, included only four out of six Neottieae genera and one mixotrophic species (*Cephalanthera longifolia* (L.) Fritsch), limiting the phylogenetic resolution and the analysis

of plastome evolution during the emergence of mixotrophy and mycoheterotrophy. Here, we sequenced 13 additional Neottieae plastomes, including representatives of the 2 remaining genera, as well as 9 mixotrophic species (table 1). We investigated how gene content and selective pressure are related to nutrition type and carried out a tentative plastome-based phylogenetic reconstruction of Neottieae.

Plastome Structure, Gene Content, and Selective Regime

None of the plastomes of the 13 photosynthetic species sequenced displayed loss of large chromosome regions (>5 kb) or changes in synteny compared with related autotrophic Neottieae (Feng et al. 2016). Their sizes and GC contents were similar, with the exception of *Neottia cordata* and *Limodorum abortivum* which have smaller genomes as a consequence of NDH gene loss (table 1). A given gene was presumed pseudogenized if a frameshift indel resulted in a reduction in protein length higher than 25% of its original length. It was considered physically lost when it failed to pass the 77% similarity threshold (compared with closest available reference species) used for gene annotation and was thus undetected. Gene loss (pseudogenization or undetected) was restricted to the NDH complex in a few species (table 1): *N. cordata* (10 genes lost, *ndhI* with 148 amino acids vs. 182 in autotrophic *Neottia* species may also be nonfunctional), *L. abortivum* (all genes) and *Epipactis microphylla* (4 genes). An exception was *cemA* (chloroplast envelope membrane protein), which is lost in *L. abortivum* similarly to some mixotrophic *Corallorhiza* (Barrett et al. 2014), which may indicate a nonessential photosynthetic function. A 9-amino-acid deletion and a 28-amino-acid insertion occurred respectively in *rpoC1* for *Cephalanthera rubra* and *accD* for *N. cordata*. However, we did not find any stop codons and/or frameshift indels in their sequence, and *accD* is known to be highly variable in length (Kim and Lee 2004; Gurdon and Maliga 2014; Wicke and Naumann 2018), suggesting that both proteins may still be functional in these species.

The selective regime was analyzed for five main gene subsets (as in Graham et al. [2017]): photosynthesis-related, NDH, ATP, PEP, and housekeeping (table 2). We used two different methods, both based on ω , the ratio of nonsynonymous substitutions per nonsynonymous site (d_N) to synonymous substitutions per synonymous site (d_S). ω provides insights into selection intensity but is also sensitive to population size and other parameters such as time of divergence (Ohta and Ina 1995). The first analysis was done with the PAML program, which allows to compute different ω values for different groups according to their nutrition type. Although this method provides useful information, it is not sufficient to confidently conclude about intensification or relaxation of selective pressure. A higher ω value can especially be a consequence of either increased positive selection or

Table 2

Analysis of Selective Pressure for Different Sets of Genes Conserved among Autotrophic and Mixotrophic Neottieae

Genes and Models	PAML Analysis				RELAX Analysis	
	np ^a	ln L ^b	Λ ^c	P Value	k ^d	P Value
Photosynthesis^e						
M0: ω ₀ = 0.11	43	-33,266.07				
M1: ω ₀ = 0.12, ω _N = 0.11	44	-33,265.49	1.17	0.28	1.03	0.67
M2: ω ₀ = 0.12, ω _p = 0.15, ω _{N*} = 0.10	45	-33,263.31	4.35	0.04	1.13	0.44
M3: ω ₀ = 0.12, ω _p = 0.15, ω _{Na} = 0.11, ω _{Nx} = 0.09	46	-33,263	0.62	0.43	0.91	1
rbcl						
M0: ω ₀ = 0.12	43	-2,687.57				
M1: ω ₀ = 0.21, ω _N = 0.09	44	-2,685.94	3.26	0.07	0.57	0.03
M2: ω ₀ = 0.20, ω _p = 0.52, ω _{N*} = 0.04	45	-2,676.02	19.85	10 ⁻⁵	5.24	10 ⁻⁴
M3: ω ₀ = 0.20, ω _p = 0.52, ω _{Na} = 0.05, ω _{Nx} = 0.01	46	-2,675.11	1.82	0.18	10.97	0.04
NDH^f						
M0: ω ₀ = 0.21	29	-18,772.43				
M1: ω ₀ = 0.18, ω _N = 0.22	30	-18,771.82	1.22	0.27	1.63	0.22
M2: ω ₀ = 0.18, ω _p = 0.16, ω _{N*} = 0.24	31	-18,770.10	3.44	0.06	1.61	0.10
M3: ω ₀ = 0.18, ω _p = 0.16, ω _{Na} = 0.24, ω _{Nx} = 0.23	32	-18,770.09	0.02	0.90	0.57	0.77
ATP^g						
M0: ω ₀ = 0.13	43	-10,149.71				
M1: ω ₀ = 0.15, ω _N = 0.13	44	-10,149.43	0.56	0.45	1.08	0.49
M2: ω ₀ = 0.15, ω _p = 0.12, ω _{N*} = 0.13	45	-10,149.4	0.05	0.82	0.72	0.16
M3: ω ₀ = 0.15, ω _p = 0.12, ω _{Na} = 0.13, ω _{Nx} = 0.12	46	-10,149.36	0.09	0.76	1.01	0.93
PEP^h						
M0: ω ₀ = 0.26	43	-22,109.79				
M1: ω ₀ = 0.23, ω _N = 0.26	44	-22,109.33	0.92	0.34	0.93	0.83
M2: ω ₀ = 0.23, ω _p = 0.19, ω _{N*} = 0.28	45	-22,107.93	2.8	0.09	1.98	0.04
M3: ω ₀ = 0.23, ω _p = 0.19, ω _{Na} = 0.30, ω _{Nx} = 0.25	46	-22,107.14	1.59	0.21	0.69	0.39
Housekeepingⁱ						
M0: ω ₀ = 0.26	43	-25,635.43				
M1: ω ₀ = 0.23, ω _N = 0.27	44	-25,634.46	1.94	0.16	1.16	0.56
M2: ω ₀ = 0.22, ω _p = 0.30, ω _{N*} = 0.26	45	-25,634.21	0.49	0.48	1.1	0.75
M3: ω ₀ = 0.22, ω _p = 0.30, ω _{Na} = 0.27, ω _{Nx} = 0.25	46	-25,633.98	0.46	0.5	1.18	0.02
matK						
M0: ω ₀ = 0.35	43	-4,121.08				
M1: ω ₀ = 0.39, ω _N = 0.34	44	-4,120.92	0.33	0.56	1.11	0.7
M2: ω ₀ = 0.39, ω _p = 0.37, ω _{N*} = 0.34	45	-4,120.9	0.05	0.83	1.34	1
M3: ω ₀ = 0.39, ω _p = 0.37, ω _{Na} = 0.45, ω _{Nx} = 0.23	46	-4,118.08	5.63	0.02	0.48	0.52
ycf1 + ycf2						
M0: ω ₀ = 0.68	43	-27,092.79				
M1: ω ₀ = 0.78, ω _N = 0.65	44	-27,091.67	2.23	0.14	1.12	0.46
M2: ω ₀ = 0.78, ω _p = 0.57, ω _{N*} = 0.66	45	-27,091.38	0.59	0.44	1.1	1
M3: ω ₀ = 0.78, ω _p = 0.57, ω _{Na} = 0.75, ω _{Nx} = 0.55	46	-27,088.8	5.16	0.02	1.13	0.13

^aNumber of parameters for the model.

^bLog-likelihood of the data for the model.

^cLog-likelihood ratio test statistic used to compute P value.

^dSelection intensity parameter.

^eccsA, petA, petB, petD, petG, petL, petN, psaA, psbA, psbC, psal, psaj, psbA, psbB, psbC, psbD, psbE, psbF, psbH, psbl, psbj, psbK, psbl, psbM, psbN, psbT, psbZ, rbcl, ycf3, and ycf4.

^fndhA, ndhB, ndhC, ndhD, ndhE, ndhF, ndhG, ndhH, ndhI, and ndhK; species analyzed were *Cephalanthera damasonium*, *Ce. longibracteata*, *Ce. longifolia*, *Ce. rubra*, *Epipactis albensis*, *E. atrorubens*, *E. gigantea*, *E. helleborine*, *E. palustris*, *E. purpurata*, *Neottia fugongensis*, *N. ovata*, *Palmeria pabstii*, *Sobralia callosa*, and *Calanthe triplicata*.

^gatpA, atpB, atpE, atpF, atpH, and atpI.

^hrpoA, rpoB, rpoC1, and rpoC2.

ⁱaccD, clpP, infA, matK, rpl2, rpl14, rpl16, rpl20, rpl22, rpl23, rpl32, rpl33, rpl36, rps2, rps3, rps4, rps7, rps8, rps11, rps12, rps14, rps15, rps16, rps18, and rps19.

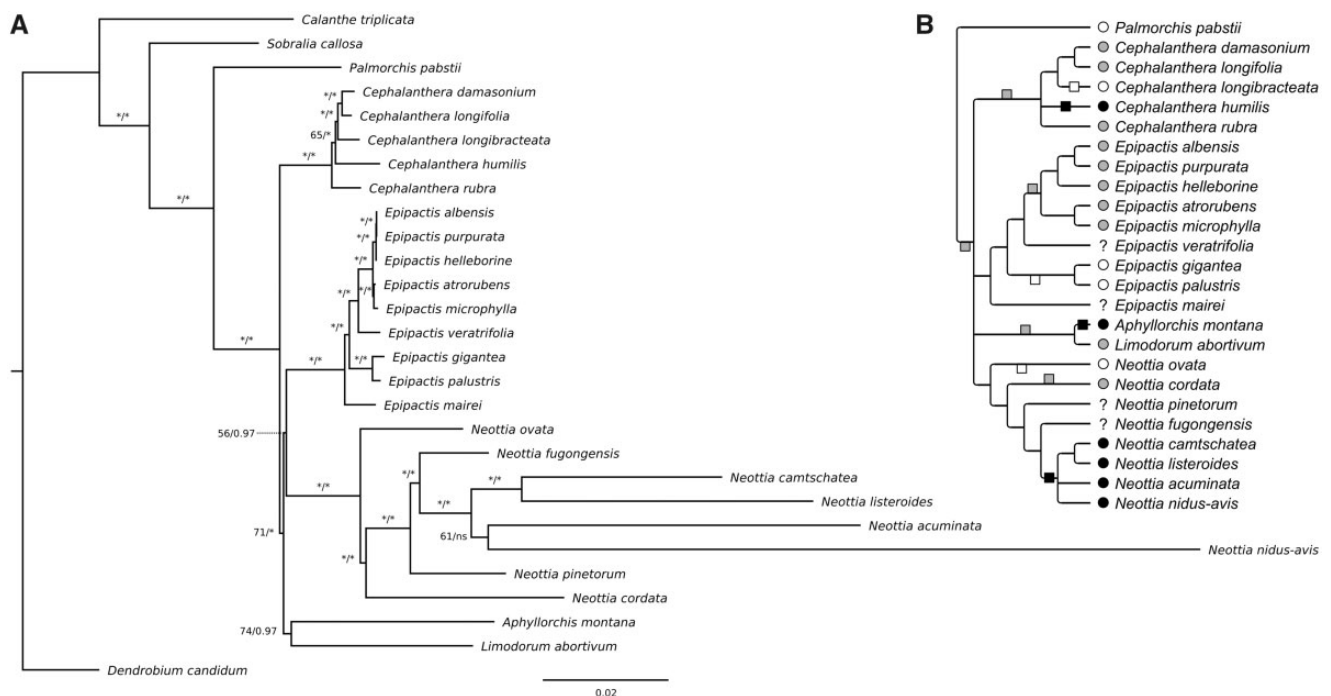


Fig. 1.—(A) Phylogeny of Neottieae based on whole plastome analysis. Numbers above or to the left of branches represent bootstrap values (1,000 replicates) from ML analysis (left, stars indicate numbers above 95) and posterior probabilities from Bayesian inference (right, stars indicate probabilities above 0.98), ns means that the branch was not recovered in the Bayesian analysis. Scale bar: number of substitutions per site. (B) Two scenarios of mixotrophy and mycoheterotrophy evolution among Neottieae, that is, autotrophy (changes above the line) versus mixotrophy (changes below the line) of the Neottieae (excl. *Palmorchis*) ancestor. The tree is a consensus of the three analyses carried out on different sets of species and plastid DNA data (see [supplementary fig. S1, Supplementary Material](#) online, for other analyses). Nutrition type evolution is mapped considering the most parsimonious scenario in each case. Circles in front of species names indicate autotrophy (white), mixotrophy (gray), or mycoheterotrophy (black) and squares on the branches indicate shifts to these nutrition types in each scenario. Nutrition type evolution is mapped considering the most parsimonious scenario in each case. Boxes on the branches show changes occurring in both scenarios.

relaxation (Wertheim et al. 2015). We used the program RELAX as a second method to refine our analysis. RELAX allows more subtle estimation of ω variations by considering different categories of sequence sites that can evolve under different selective constraints (Wertheim et al. 2015).

For PAML analysis, four nested models allowing the calculation of different ω values for some subsets of branches were defined: M0, the null model with one ω for the whole tree; M1, which allowed ω to differ between outgroups (ω_0) and Neottieae (ω_N); M2, same as M1 but allowing a different ω for *Palmorchis* (ω_p) and the remaining Neottieae (ω_{N^*} ; below noted as Neottieae* = Neottieae without *Palmorchis*); M3, same as M2 but allowing ω to differ between autotrophic (ω_{Na}) and mixotrophic Neottieae* (ω_{Nx} , see [fig. 1B](#) for species considered as mixotrophic). Models were compared by likelihood ratio tests following this scheme: M1 versus M0, M2 versus M1, and M3 versus M2. For all subsets, successive comparisons of the four models of ω variation among branches did not show a better fit for one model against another one, except for the photosynthesis genes for M2 versus M1 ([table 2](#)). M2 allowed different ω for outgroups ($\omega_0 = 0.12$), *Palmorchis pabstii* ($\omega_p = 0.15$), and the

remaining Neottieae ($\omega_{N^*} = 0.10$; [table 2](#)). These results were not supported by the RELAX analysis, which indicated instead intensification of selective constraints for PEP genes comparing Neottieae* and *Palmorchis* ($k = 1.98$; $P = 0.04$) and for housekeeping genes comparing mixotrophic and autotrophic Neottieae* ($k = 1.18$; $P = 0.02$; [table 2](#)). But one should consider that these results are not adjusted for multiple comparisons.

The same analysis carried out for individual photosynthesis-related genes showed a variation in selective regime only for *rbcl* ([table 2](#) for *rbcl*, other genes shown in [supplementary table S2, Supplementary Material](#) online). *rbcl* data fitted models M2 much better than M1. In M2, ω was lower for Neottieae* ($\omega_{N^*} = 0.04$) compared with outgroups ($\omega_0 = 0.20$) and *P. pabstii* ($\omega_p = 0.52$; [table 2](#)). This was supported by the RELAX analysis, which showed an intensification of selective constraints in Neottieae* compared with *Palmorchis* on the one hand ($k = 5.24$; $P = 10^{-4}$), and in mixotrophic compared with autotrophic Neottieae* on the other hand ($k = 10.97$; $P = 0.04$). Thus, *rbcl* experiences stronger purifying selection in Neottieae*, and more particularly in mixotrophic species. For individual nonphotosynthesis genes,

PAML analysis showed that both *matK* and *ycf1 + ycf2* fitted M3 better than M2, with higher ω for autotrophic ($\omega_{Na} = 0.45$, resp. 0.75) compared with mixotrophic Neottieae* ($\omega_{Nx} = 0.23$, resp. 0.55; [table 2](#)). This was, however, not supported by the RELAX analysis ([table 2](#)).

To summarize, we did not find evidence for relaxed selection in plastomes of mixotrophic species. By contrast, both PAML and RELAX analyses showed stronger selective pressure in mixotrophic Neottieae* compared with *Palmorchis* for *rbcl*.

Plastome Evolution in Mixotrophic Neottieae

The observed loss of NDH genes in three plastomes confirms that this gene complex is sensitive to evolutionary decay in Neottieae (Feng et al. 2016) and more generally in orchids (Barrett et al. 2014; Luo et al. 2014; Kim et al. 2015; Lin et al. 2017; Kim et al. 2018) and some other photosynthetic lineages (Wicke and Naumann 2018). Interestingly, it does not correlate with species nutrition. For example, the 11 genes were lost in *L. abortivum* and *N. cordata* (if *ndhI* is indeed nonfunctional). The first is mixotrophic and highly dependent on fungal carbon because its photosynthesis does not even compensate for its respiration (Girlanda et al. 2006); the second displays a more versatile use of fungal resources (Schiebold et al. 2018), which may even be optional (Těšitelová et al. 2015). Similarly, *E. microphylla* is the only species of the mixotrophic *Epipactis* section (*E. albensis*, *E. atrorubens*, *E. helleborine*, and *E. purpurata*) (Jin et al. 2014) for which NDH gene loss occurred, whereas some losses have been reported in *E. veratrifolia* and *E. mairei* (Feng et al. 2016) that are supposed autotrophic (pending more investigations). This suggests a species-specific decay of NDH genes in orchids, independently of their autotrophic or mixotrophic habit. In addition, we did not observe any evidence for intensification or relaxation of selective pressure on these genes ([table 2](#)). We suppose that relaxation on NDH genes occurs in the whole orchid clade, making any difference among orchid species hardly detectable. Changes in nutrition type do not seem to alter this pattern.

Photosynthesis-related genes neither are lost nor experience relaxed selective constraints in mixotrophic species but, in Neottieae*, they showed a slightly lower ω ; *rbcl* was even under stronger purifying selection compared with *P. pabstii* ([table 2](#)). Some parts of the plastomes unrelated to photosynthesis also experienced higher selective constraints in mixotrophs, as shown by 1) lower ω for *matK* and *ycf1 + ycf2* in mixotrophic than in autotrophic Neottieae* and 2) the RELAX analyses of PEP and housekeeping genes (displaying $k > 1$). These results are concordant with what has been reported for mixotrophic *Corallorhiza*, which 1) retain all photosynthesis-related genes except *cemA* and *psbM*, 2) display apparently unimpaired photosynthesis, and 3) show no difference in plastome selective regime compared with autotrophic references

(Barrett et al. 2014). This also agrees with the retention of purifying selection on plastidial genes observed in the mixotrophic *C. macrorhizon* (Kim et al. 2018). However, our results contrast with other kinds of mixotrophy where relaxed purifying selection has been observed for photosynthesis, PEP and ATP-synthase genes, namely 1) in carnivorous plants (*psa*, *atp*, and *rpo* genes in Wicke et al. [2014]) and 2) in obligate hemiparasites (*psa*, *psb*, *pet*, *atp*, and *rbcl* in Petersen et al. [2015]; photosynthesis and *atp* genes in Wicke et al. [2016]). For this latter trophic strategy, retention of selective constraints on photosynthesis genes has nevertheless been reported for photosynthesis genes in *Cuscuta* (McNeal et al. 2007) and also for some housekeeping genes in *Viscum* (*accD*, *cemA*, *clpP*, and *ycf2*, Petersen et al. 2015). However, these comparisons between studies have to be taken with caution given the differences in scales among them or in a given study (individual genes vs. gene subsets).

In Neottieae, depending on the plastid genes considered, mixotrophic species thus feature either no change or a lowly supported increase in selective pressure compared with other mixotrophic groups such as *Corallorhiza*, carnivorous or parasitic plants (Barrett et al. 2014; Wicke et al. 2014, 2016; Petersen et al. 2015). Photosynthesis therefore seems to remain advantageous in mixotrophic Neottieae, and this is congruent with physiological evidence for the importance of photosynthetic carbon in fruit development and reproduction in these species (Roy et al. 2013; Bellino et al. 2014; Gonneau et al. 2014; Lallemand et al. 2019). We hypothesize that the retention of genes and purifying selection in the plastome of *C. macrorhizon* (Kim et al. 2018) indicates a similar crucial role of photosynthesis in reproduction, as illustrated by Suetsugu et al. (2018).

Neottieae Phylogeny and Mycoheterotrophy Evolution

Phylogenetic analysis of the 28 available plastomes confirmed Neottieae monophyly with *Palmorchis* in sister position to the remaining species ([fig. 1A](#)). *Neottia*, *Epipactis*, and *Cephalanthera* genera were each monophyletic and most intrageneric relationships were well defined. *Aphyllorchis* and *Limodorum* were sister genera but with mild support from maximum likelihood (ML) analysis (bootstrap = 74, [fig. 1A](#)). However, very short branches with low support prevented strong conclusions about intergeneric relationships ([fig. 1A](#)). Because mycoheterotrophic *Neottia* species display accelerated evolutionary rates (see long branches in [fig. 1A](#)), we removed them to increase alignment quality: We then obtained a congruent tree with better support for most but not all clades ([supplementary fig. S1A, Supplementary Material](#) online). In particular, *Neottia*, *Epipactis*, *Aphyllorchis*, and *Limodorum* clustered together in a monophyletic group sister to *Cephalanthera* with better support from ML analysis (bootstrap = 87 vs. 71). When restricting

the analysis to the plastidial coding sequences (CDS) of non-mycoheterotrophic species, that is, excluding noncoding regions to allow better alignment, *Limodorum* was sister to *Epipactis* but *Neottia* no longer clustered with these genera in a monophyletic group (supplementary fig. S1B, Supplementary Material online). Combining these results suggests that at least *Cephalanthera* is sister clade to the rest of Neottieae. Analysis based on ITS (internal transcribed spacer of nuclear ribosomal DNA, including ITS1-5.8S rRNA-ITS2) confirmed *Aphyllorchis* and *Limodorum* as sister genera but failed to clarify further intergeneric relationships (supplementary fig. S1C, Supplementary Material online).

Phylogenetic relationships between the Neottieae genera have long been controversial and studies using different species and marker combinations yielded contradictory results, without any clear intergeneric relationships (ITS and the plastid *trnL* intron in Bateman et al. [2005]; ITS, *trnS-G* spacer, and *rbcl* in Roy et al. [2009]; ITS, *Xdh*, *rbcl*, *matK*, *psaB*, and *trnL-F* spacer in Xiang et al. [2012]; ITS, *rbcl*, and *matK* in Zhou and Jin [2018]). More confusingly, the phylogeny of Feng et al. (2016), based on a subset of the plastomes we analyzed, disagrees with our results, displaying *Cephalanthera* and *Aphyllorchis* as sister genera, and *Epipactis* and *Neottia* as successive sisters. This inconsistency may result from the difference in the number and diversity of species included in the analysis. We added 13 species and 2 new genera (*Palmorchis* and *Limodorum*) but failed to reproduce their results when considering only their subset of species, suggesting that this is rather due to different phylogenetic methods and/or to evolutionary signals conveyed by the plastome regions used. Feng et al. (2016) selected conserved plastome regions, whereas we made a dual analysis of whole plastome alignment and CDS-restricted alignment. Moreover, we chose not to filter or manually edit alignments following Tan et al. (2015) for accurate phylogenetic inferences (see also Hallas et al. 2017).

Neottieae intergeneric relationships thus remain difficult to estimate confidently. Our plastome-based analysis showed that the branches separating these genera are very short ($<10^{-3}$ substitutions per site). Thus, the common ancestor of Neottieae* (i.e., Neottieae with exclusion of *Palmorchis*) likely experienced rapid speciation, with little time for informative mutations to accumulate (Glor 2010). Such a fast evolution of ancestral lineage makes it prone to hybridization and incomplete lineage sorting (Glor 2010; Wang et al. 2014), so that independent loci (e.g., nuclear and plastidial markers) may encapsulate different phylogenetic stories and make any combined analysis unreliable. Another possibility is heterotachy (Fitch and Markowitz 1970), a variation (here a decrease) in evolutionary rate in the ancestral lineage of Neottieae*. Arbitrating between fast speciation and heterotachy would be particularly challenging given the very short branch lengths. Confident reconstruction of early speciation in Neottieae will be complicated, but different gene trees

could provide interesting insights into early Neottieae evolution and possible hybridization events.

We also wanted to compare two possible evolution of nutrition in Neottieae*. Based on our plastid phylogeny, we display two hypotheses, namely 1) an autotrophic ancestor and 2) a mixotrophic ancestor for Neottieae* (see Materials and Methods for reconstruction of intermediate characters). Assuming an autotrophic common ancestor, mixotrophy evolved at least four times (fig. 1B, gray boxes above the branches). Conversely, assuming a mixotrophic common ancestor would give three reversions to autotrophy (fig. 1B; white boxes below the branches). Each scenario implies two emergences of mycoheterotrophy, but unfortunately, the lack of information about the trophic status of some taxa (such as *E. mairei* or *Neottia fugongensis*) and the absence of several mycoheterotrophic clades limit our analysis. For example, green *Neottia* other than *N. cordata* or *Epipactis* species outside the *Epipactis* section may actually show mixotrophic nutrition. Assuming a mixotrophic common ancestor implies one change less than an autotrophic one but includes reversion to autotrophy. Although unlikely at first glance, reversion is allowed because mixotrophy is flexible and displays a continuum from autotrophy to mycoheterotrophy (Jacquemyn et al. 2017). Most importantly, the selective pressure maintaining photosynthesis genes reported here allows a reversion from mixotrophy to autotrophy, because potential for autotrophy remains intact in mixotrophic Neottieae. The question whether mixotrophy is a plesiomorphy of Neottieae thus remains open, pending for analyses of more species.

A final speculation comes at this point: If mixotrophy turns out to be indeed the ancestral state of Neottieae, could it have been a key factor in the fast early diversification speculated above? Mixotrophy allows plants to grow in light-limited forest understory and so to occupy a broad new ecological niche (Těšitel et al. 2018). Orchids show important dispersion abilities and most mixotrophic Neottieae do not seem very selective regarding the fungi they associate with (Těšitelová et al. 2012). One can therefore speculate that an early mixotrophic Neottieae lineage could have rapidly colonized distant and heterogeneous forest patches, with ability to keep or lose photosynthetic abilities, thus favoring radiation.

Materials and Methods

DNA Extraction and Sequencing

Total DNA was extracted from the leaves or fruits of the following Neottieae species (table 1 and supplementary table S1, Supplementary Material online): *Palmorchis pabstii* Veyret, *N. cordata* (L.) Rich., *E. albensis* Nováková & Rydlo, *E. atrorubens* (Hoffm.) Besser, *E. gigantea* Douglas ex Hook., *E. helleborine* (L.) Crantz, *E. microphylla* (Ehrh.) Sw., *E. palustris* (L.) Crantz, *E. purpurata* Sm., *Cephalanthera damasonium* (Mill.) Druce, *Ce. longibracteata* Blume, *Ce. rubra* (L.)

Rich., and *L. abortivum* (L.) Sw. The following extraction kits and protocols were used (supplementary table S3, Supplementary Material online): DNeasy Plant Mini Kit (Qiagen) with final elution in distilled water, NucleoSpin Plant II kit (Macherey-Nagel), CTAB extraction as in Porebski et al. (1997).

Given low yield due to DNA degradation in some old samples, the Accel-NGS 1S Plus DNA Library Kit for Illumina with the 1S Plus Indexing kit (Swift Biosciences Inc., Ann Arbor, MI) was used for preparation of libraries, following the manufacturer's instructions. We did so for the set of samples handled in EPGV laboratory (supplementary table S3, Supplementary Material online). For other samples, handled in Moscow laboratory, the TruSeq DNA sample preparation kit (Illumina, USA) and the NEBNext Ultra DNA kit (New England Biolabs, USA) were used for preparation of libraries (supplementary table S3, Supplementary Material online). Paired-end sequencing on multiplexed libraries was performed either on Illumina HiSeq 2000, Illumina HiSeq 2500 using the rapid run mode, and Illumina HiSeq 4000 or Illumina MiSeq, depending on the samples (different material available in the different teams involved in the study; supplementary table S3, Supplementary Material online). Raw sequences for *Ce. longibracteata*, *Ce. rubra*, *E. albensis*, *E. atrorubens*, *E. gigantea*, *E. microphylla*, *E. palustris*, *E. purpurata*, and *P. pabstii* were submitted to the NCBI database: BioProject PRJNA484137, SRA accession number ongoing.

Read Cleaning, Plastome Assembly, and Annotation

Redundant reads were removed with a homemade C++ script. The redundancies were sought by pairs read1–read2, the redundancy with the best Phred score was kept. Trimming of the low-quality 3' end was done using a homemade C++ script, one base after the other until reaching a base with a Phred score > 30 or until the mean Phred score of the read was above 30. Pairs with at least one read shorter than 30 bp after this step or with N were removed. Read number before and after cleaning are indicated in supplementary table S4, Supplementary Material online.

For samples that were prepared with the Accel-NGS 1S Plus DNA Library Kit, adapter trimming was conducted using BBDuk Trimmer plugin version 37.64 in Geneious version 11.1.3 (<http://www.geneious.com>; last accessed September 2018; Kearsse et al. 2012) with the following parameters: adapters = Illumina Truseq DNA adapters, ktrim = r, k = 27, hdist = 1, edist = 0, mink = 6, minlength = 10, trimbyoverlap = t, and minoverlap = 6. Addition of a low complexity tail to the 3' end of fragments was part of the library preparation kit and had to be removed from the beginning of reads 2. Following the manufacturer's recommendations, trimming an additional 10 bp at the 5' end was done for all reads.

De novo assembly was done differently depending on the species (supplementary table S3, Supplementary Material online). 1) A subset of 25% of the reads was assembled using

the Geneious algorithm with medium-low sensitivity parameters. Resulting contigs were mapped on the plastome of *Neottia ovata* (L.) Bluff & Fingerh. (KU551271), *Cephalanthera longifolia* (L.) Fritsch (KU551263), or *E. veratrifolia* Boiss. & Hohen (KU551267), depending on the species. Contigs corresponding to the plastome were then dissolved and reassembled with medium sensitivity parameters to increase assembly quality. The resulting contigs were again mapped to the reference genome and large single copy (LSC), inverted repeat B (IRB), and small single copy (SSC) regions were isolated. IRB was duplicated and reverse complemented to obtain inverted repeat A (IRA). Concatenation of LSC, IRB, SSC, and IRA led to the final plastome sequence. 2) The CLC Genomics Workbench de novo assembly procedure was used. In this latter case, reads were first trimmed with the following settings: quality limit = 0.01 (corresponds to $Q \geq 20$), remove adapters = yes (Illumina Truseq adapters), discard short reads = yes (limit 50 nt for MiSeq data, 25 nt for HiSeq data). Then de novo assembly was performed with the following settings: automatic bubble size = yes, minimum contig length = 1,000 bp, automatic word size = yes, perform scaffolding = yes, and auto-detect paired distances = yes. Contigs of plastid origin were selected based on the results of BLAST search with *Phalaenopsis aphrodite* plastome used as query. Contigs corresponding to LSC, IR, and SSC were concatenated accordingly and resulting sequences were verified and corrected by performing the gap closure method using back mapping of the reads. IRs were duplicated and reverse complemented to obtain the final parts (IRA) of the plastomes.

Coverage values of the contigs used for assembly are indicated in supplementary table S4, Supplementary Material online. Depending on the species, the phylogenetically closest reference plastome was used for annotations transfer in Geneious with a 77% similarity threshold that turned out, after preliminary attempts, to be low enough to detect similar gene sequences and sufficiently high to avoid false positives. Manual correction and addition of missing information was then carried out for each plastome. Annotated plastome sequences were submitted to GenBank under the accession numbers MH590345–MH590357 (table 1).

Phylogenetic Inferences

Including the 12 already published sequences, we ended up with a set of 25 Neottieae plastomes and added 3 outgroup species from the same subfamily Epidendroideae (supplementary table S1, Supplementary Material online). The four mycoheterotrophic *Neottia* were either kept or removed to test whether their rapidly evolving plastomes (much higher than the two other mycoheterotrophs; fig. 1A) and the increased length and quality of the alignment obtained without them alter phylogeny reconstruction. The second copy of the inverted repeat region (IRA) was removed from all plastomes

and the resulting sequences were aligned a first time using the Mauve progressive algorithm, as implemented in Geneious (parameters set to automatically calculate seed weight and minimum locally collinear block (LCB) score, compute LCBs, full alignment), to detect potential rearrangements (Darling et al. 2004). The LCBs obtained were individually aligned a second time using the MAFFT online service (Kato et al. 2017) with automatic selection of alignment strategy, gap opening penalty set to 3.0, offset value set to 0.1 and other parameters left as default. The resulting alignments were then concatenated. Model selection was done with the ModelFinder algorithm (Kalyaanamoorthy et al. 2017) implemented in the IQ-TREE program (Nguyen et al. 2015). The model with lowest Bayesian information criterion score was used for ML analysis with IQ-TREE (Nguyen et al. 2015). The first model with the lowest Bayesian information criterion compatible with MrBayes v3.2 parameters (Ronquist et al. 2012) was chosen for Bayesian inference with this program. A complementary analysis using the nuclear marker ITS, including ITS1-5.8S rRNA-ITS2, was done using available Neottieae sequences (supplementary table S1, Supplementary Material online).

On the plastid phylogeny (fig. 1B), we displayed two possible evolution of nutrition based on the hypotheses of 1) an autotrophic ancestor or 2) a mixotrophic ancestor for Neottieae*. Intermediate characters for other nodes were proposed in order to minimize the number of transitions between trophic types (autotrophic, mixotrophic, or mycoheterotrophic), that is, the most parsimonious scenario under each hypothesis, considering the nutritional status of extant species (supplementary table S1, Supplementary Material online). A given species was considered as mixotrophic when it dominantly associated with ectomycorrhizal fungi and showed an enrichment in ^{13}C compared with surrounding autotrophs, a usual marker of fungal carbon gain (Hynson et al. 2013; see supplementary table S1, Supplementary Material online, for references regarding species nutrition).

Selective Regime Analyses

All CDS were extracted from the same set of plastomes used for phylogeny reconstruction, except the six mycoheterotrophic species. Except for NDH complex, genes that were lost or pseudogenized in at least one of the remaining species were discarded from subsequent analyses (see table 2 for the remaining genes). For NDH, because some species had lost all the genes, we rather only kept for subsequent analysis only the 15 species, which retained the 11 NDH genes (table 2). CDS were aligned based on their amino acid sequences with subsequent backtranslation using the AlignTranslation function (Wright 2015) of the DECIPHER package (Wright 2016) in the R environment for statistical computing (R Development Core Team 2007). Functional gene groups were built by concatenating CDS for photosynthesis, NDH, ATP-synthase,

PEP, and housekeeping genes (table 2). *ycf1* and *ycf2* were combined and analyzed independently because of their specific evolutionary rates (Barrett et al. 2014). In addition, some genes were analyzed individually when long enough for confident parameter estimation: *ccsA*, *psaA*, *psaB*, *psbA*, *psbB*, *psbC*, *psbD*, *rbcl*, *accD*, and *matK* (see supplementary table S2, Supplementary Material online). ω , the ratio of nonsynonymous substitutions per nonsynonymous site (d_N) to synonymous substitutions per synonymous site (d_S), was used to estimate selective pressure for different sets of genes. The codeml program implemented in PAML version 4.9 (Yang 2007) was used to compute ω values, with the branch model and the tree shown in figure 1B. The codon frequency parameter was set to F3X4. Four nested models allowing the calculation of different ω values for some subsets of branches were defined: M0, the null model with a unique ω for the whole tree; M1, which allowed ω to differ between outgroups (ω_O) and Neottieae (ω_N); M2, same as M1 but allowing a different ω for *Palmorchis* (ω_P) and the remaining Neottieae (ω_{N^*}); M3, same as M2 but allowing ω to differ between autotrophic (ω_{Na}) and mixotrophic Neottieae* (ω_{Nx} , see fig. 1B for species considered as mixotrophic). Models were compared by likelihood ratio tests following this scheme: M1 versus M0, M2 versus M1, and M3 versus M2. The statistic of the test (Λ) was calculated as two times the difference in model likelihood and compared with a chi-squared distribution with degrees of freedom equaling the difference in parameters number between the two models. *P* values were obtained using the function pchisq of the R stats package.

We completed the PAML analysis with a specific test dedicated to the detection of changes in selective pressure (relaxation or intensification). This was carried out with the program RELAX, available at the Datamonkey webserver <http://datamonkey.org/relax>; last accessed March 6, 2019. RELAX allows more subtle estimation of ω variations by considering different categories of sequence sites that can evolve under different selective constraints (Wertheim et al. 2015). ω distributions are estimated for a test (T) and a reference (R) subset of branches and a selection intensity parameter k such that $\omega_T = \omega_R^k$ is introduced. The program then compares the goodness of fit of the data to a branch-site evolutionary model with $k = 1$ (null model) or k being a free parameter (alternative model). When the alternative model shows a better fit following a likelihood ratio test, $k < 1$ (resp. $k > 1$) indicates a relaxation (resp. an intensification) of selective constraints in the test group. Three different tests were done with three different designs of test versus reference groups, copying what has been done with the PAML nested analysis (see supplementary fig. S2, Supplementary Material online, for group designs).

Supplementary Material

Supplementary data are available at *Genome Biology and Evolution* online.

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Author Contributions

Conceptualization: F.L. and M.-A.S.; formal analysis: F.L.; investigation: F.L., M.L., M.M., I.L.C., A.B., and M.-C.L.P.; resources: M.L., I.L.C., A.B., E.Z., M.M., and M.-C.L.P.; data curation: F.L., M.L., M.M., M.J., and A.B.; writing—original draft: F.L.; writing—review and editing: F.L., M.L., E.Z., E.D., M.J., M.-C.L.P., and M.-A.S.; supervision: M.-C.L.P. and M.-A.S.; funding acquisition: M.-A.S.

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