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TROPHIC RELATIONSHIPS IN ORCHID MYCORRHIZA – DIVERSITY AND IMPLICATIONS FOR CONSERVATION

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Introduction

Orchid species are perennial, and though demographic data suggest that the family includes r- as well as K-strategists (Whigham & Willems 2003), most species are potentially long-lived. Individual plants may be kept in living plant collections or in nature reserves for practically unlimited periods of time. There are several reports on natural populations suspected of little or no seedling recruitment, "senile populations" (Tamm 1991, Rasmussen 1995), especially among rare orchids under critical surveyance. Such populations may function as a seed source to neighbouring areas but are likely to eventually disappear from the site.

Sustainable conservation thus requires the preservation of conditions that enable the species to carry through its entire life cycle. The conservational concern should also involve species of other organisms that are associated during a critical life stage, such as a pollinator during flowering, or a symbiotic fungus during seed germination (Zettler *et al.* 2003). Not only that, but the requirements of these organisms must be considered, such as appropriate substrates for the fungi. Clearly, "orchids require an ecosystem approach to their conservation" (Roberts 2003).

In light of recent research, the orchid-fungus relationship has proved particularly complex, as it may be subject to trophic changes during the lifetime of the orchid. The degree of specificity, and the paths of biological energy are major concerns in these relationships. All of this may need to be assessed in cases of severely endangered orchid species.

Orchid mycorrhiza is still considered a unilateral relationship

Transport of carbohydrates from fungi to seedlings of orchids has been amply demonstrated, beginning with Smith's experiments (1966, 1967). There is no other feasible explanation for the long-term deficiency in the photoassimilating apparatus known from orchid seedlings generally and adult stages widespread in the family (e.g., Girlanda et al. 2006). Recent stable isotope analyses support the fungal origin of a significant part of the C and N found in aboveground structures of orchids (Gebauer & Meyer 2003, Julou et al. 2005). Hyphal coils within the orchid tissues become degraded by an enzymatic process and transfer is assumed to occur entirely or predominantly over a dead fungal interface, as ultrastructural studies suggest (Peterson et al. 1996). This further adds to the evidence of an asymmetric relationship, with the orchid as the receiving and depen-

A recent report stating mutualism in orchid-fungus relationships (Cameron et al. 2006) was based on a set of special experimental circumstances: Surface sterilized plantlets were planted on an inert agar, and internal hyphae from within the rhizome were allowed to colonize the agar. When ¹⁴CO₂ was subsequently supplied to the leaves, about 2% of the photoassimilated labeled carbon could later be traced to the mycelium. Physiologically interesting as this may be, it is important to note that the result was obtained under an extreme starvation of the mycelium. Such conditions would hardly ever occur under field conditions where complex carbon

sources abound. Furthermore, we do not know whether the transfer occurred via an intact plant interface surrounding hyphal pelotons. Field studies suggest that low substrate carbon supply may increase the virulence of the fungi and turn the situation into parasitism of the fungus on the orchid (Beyrle *et al.* 1995).

Orchids are never "fully autotrophic"

Seedlings of Neuwiedia veratrifolia, belonging to the subfamily Apostasioideae, usually considered the most basal in orchid phylogeny, were found for the first time by Kristiansen et al. (2001). They develop typical protocorms with pelotons, and the fungi associated with them in the wild proved to belong to Tulasnella and Thanathephorus, two genera that are known to develop Rhizoctonia-stages (Kristiansen et al., 2004). VAM is the only type of mycorrhiza found in monocotyledons outside of Orchidaceae, and it seems a plausible scenario that ancestors of the orchid family developed a seedling mycotrophy, based on invasive saprotrophic rhizoctonioid mycelia in conjunction with the evolution of micro-seeds. From the beginning the whole range of this rather mixed assembly of imperfect mycelia of Basidiomycetes (i.e., Rhizoctonia s.l., Table 1, below) appears to have been employed.

Orchid seed evolution seems to have run towards further reduction in size, the epiphytic orchid groups tending to produce smaller seeds than terrestrials (Rasmussen, 1995). Assuming an evolutionary reduction of seed nutrient reserves within the orchid family clade, a secondary loss of seedling mycotrophy appears unlikely, and is not supported by any observations so far. In other words: seedling mycotrophy seems to be a uniquely derived and omnipresent orchid character.

Plant seedlings generally begin life by utilizing seed reserves that consist of stored photoassimilates from their autotrophic mother plant. In contrast, the orchid seedling relies not only on reserves from the mother plant but also on carbohydrates from mycotrophy. Otherwise the seedlings will not develop in the field. Thus, if the whole life history is considered, orchids are never fully autotrophic. When this description is sometimes used about

orchids (e.g., "the fully autotrophic *Listera ovata*", Girlanda *et al.*, 2006), this either refers to the adult stage only, or must be considered a slip of the pen.

In terms of preserving an orchid species, this means that fungi employed during germination cannot be disregarded. All orchids are to some degree mycoheterotrophic, although this designation has somewhat misleadingly been restricted to species with obviously chlorophyll-deficient adult stages. The sequential or simultaneous combination of mycotrophy and phototrophy, that is charactistic of orchids, may be described as mixotrophy. The only exception from mixotrophy would be the entirely mycotrophic orchids. Cephalanthera damasonium is an example of a species that segregates into holomycotrophic and mixotrophic individuals: the albinos showed no trace of photoassimilation as adult plants, whereas the adult green individuals were found to be mixotrophic with about fifty-fifty contribution of carbon from either system (Julou et al. 2005). Other studies, also based on the distribution of stable carbon and nitrogen isotopes, indicate that green leaved forms may aquire a significant fraction of their C and N through fungi, but that species differ considerably in this respect (Gebauer & Meyer, 2003). Thus orchids are arranged in a continuum from holomycotrophy to various degrees of mixotrophy.

Useful terminology from the animal kingdom

Taylor (2004) put it aptly: "let's be clear – we are talking about plants that consume fungi." Much confusion may arise from inadequte or misleading designations. The phytobiont (orchid) has colloquially been referred to as the 'host', notwithstanding the fact that the mycobiont is providing the meal! Even worse are anthropomorphic expressions that seem to imply voluntary and mutualistic associations ('marriage', 'fidelity', 'promiscuity'), or deception ('cheater') which suggests a previous mutualism or presupposes a "normal" behavior deviated from. Even the idea of 'specificity' implies a degree of mutual selection. Such expressions should be avoided as they are inconsistent with our observations and present knowledge.

Table 1. Above: Examples of orchid-fungus-substrate relationships. Below: Fungus genera mentioned above, listed with taxonomic position according to Kirk *et al.* 2001.

Orchid species	Trophic stage	Provider/Prey	Ultimate food source
Neuwiedia veratrifolia: inc	dividually monophagous? (Kristiansen et al	. 2004)	
	seedling mycotrophy	Tulasnella Thanathephorus	leaf litter (Kristiansen et al., 2001) leaf litter
	adult mycotrophy adult phototrophy	Tulasnella sp., Thanathephorus sp.	
Cypripedium, several spec	ies: mono-oligophagous (Shefferton et al. 2	005)	
	seedling mycotrophy	?	
	adult mycotrophy adult phototrophy	Tulasnellaceae	organic debris
Goodyera pubescens and I	Liparis lilifolia: mono(-oligo)phagous (McC	Cormick et al. 2004)	
	seedling mycotrophy	Tulasnella cf. bifrons	organic debris
	adult mycotrophy	Tulasnella cf. bifrons	organic debris
	adult phototrophy		
Epipactis microphylla: oli	gophagous (Selosse et al. 2004)		
	seedling mycotrophy	Tuber?	
	adult mycotrophy	mainly Tuber cf. excavatum	live trees (ECM)
	phototrophy (in green individuals) and not (in albinos)		
Neottia nidus-avis: oligopl	hagous, locally monophagous? (McKendric	k et al. 2002. Selosse et al. 2002)	
veolita muus-uvis. ongopi		,	Line to an (ECM)
	seedling mycotrophy adult mycotrophy	Sebacina Sebacina	live trees (ECM) live trees (ECM)
	no phototrophy	Sebacina	live trees (ECIVI)
	(sources cited in Mckendrick <i>et al.</i> 2002)		
Limodorum abortivum: ol	igophagous, obligate fungal switch? (Girlan	nda 2006)	
	seedling mycotrophy	Ceratobasidium?	organic debris
	adult mycotrophy	Russula spp.	live trees (ECM)
	very little phototrophy		
Tipularia discolor: switch	from germination fungus, polyphagous as a	dult (McCormick et al. 2004)	
	seedling mycotrophy	Tomentella sp.	large woody debris
	adult mycotrophy	4 groups of tulasnelloids	mussen & Whigham 1998 organic debris
	y y y	+ some persistance of <i>Tomentella</i>	3
	adult phototrophy	•	
Corallorhiza trifida: mono	ophagous (McKendrick et al. 2000a+b)		
	seedling mycotrophy	Tomentella	Salix and Betula ECM
	adult mycotrophy	Tomentella	Salix and Betula ECM
	no phototrophy		
Epidendrum rigidum: mor	nophagous (Pereira et al. 2005)		
	seedling mycotrophy	Epulorhiza	? saprophyte
	adult mycotrophy	<i>Epulorhiza</i>	? saprophyte
	adult phototrophy		
Hexalectris spicata: oligop	phagous (Taylor et al. 2003)		
	seedling mycotrophy	?	?
	adult mycotrophy	Sebacinaceae+ Thanathephorus	live trees (ECM)?
	chlorophyll deficient		

TABLE 1. Continue.

Psathyrella

Russula

Sebacina

Tomentella

Tulasnella

Tuber

Thanathephorus

Orchid species	Trophic stage	Provider/Prey		Ultimate food source
Gastrodia elata: serial	monophagy, obligate switch of fungus (Xu	ı & Guo, 2000)		
	seedling mycotrophy adult mycotrophy no phototrophy	Mycena osmundicola Armillaria mellea s.l.		leaf litter live and dead wood
Epipogium roseum: ol	ligophagy? Yamato et al. 2005			
	seedling mycotrophy	? Coprinus + Psathyrella		dung, dead wood
	adult mycotrophy			
	no phototrophy?			
Teleomorph	Anamorph	Family	Order and class	
Armillaria mellea		Marasmiaceae	Aga	ricales Basidiomycetes
Ceratobasidium	Ceratorhiza (Rhizoctonia s.1.)	Ceratobasidiaceae	Ceratobasidiales Basidiomycetes	
Coprinus		Coprinaceae	Agaricales Basidiomycetes	
Mycena		Tricholomataceae	Aga	ricales Basidiomycetes

Coprinaceae

Russulaceae

Exidiaceae

Ceratobasidiaceae

Thelephoraceae

Tuberaceae

Tulasnellaceae

It seems about time to acknowledge that orchids are mycophagous and that the orchid-fungus association is more like a predator-prey-relationship. A set of concepts and terminology from the zoological vocabulary comes to mind. Recent research has revealed a trophic diversity in orchids so great that we need these concepts to encompass the whole range. Thus, we have examples of orchids with a broad food selection (i.e. polyphagous), the diet spanning several fungal families (Tipularia discolor, Table 1) as well as examples of orchids that are oligophagous, utilizing a minor group of related fungi. Verification of strict monophagy requires the analysis of the plant species through much of its geographic and ecological range. Normally monophagy would be an orchid species-to-fungal species relationship but it might also exist on the individual level, as shown in Goodyera pubescens (McCormick et al. 2006). In this species germina-

Epulorhiza

Epulorhiza

(Rhizoctonia s.l.)

(Rhizoctonia s.l.)

Rhizoctonia s.str.

tion could be carried out with a range of *Rhizoctonia* spp., but the first strain to infect an individual protocorm seemed to be subsequently preferred. Young plants of *Goodyera pubescens* only rarely switched from their initial fungus, which shows a surprising ability of seedlings to discriminate hyphae. When a switch was induced experimentally, it carried a considerable risk of mortality.

Agaricales Basidiomycetes

Russulales Basidiomycetes

Tremellales Basidiomycetes

Ceratobasidiales Basidiomycetes

Thelephorales Basidiomycetes

Pezizales Ascomycetes

Tulasnellales Basidiomycetes

Nevertheless, an obligate switch of fungus at some point during adolescence is well documented in *Gastrodia elata*, that is known to germinate on *Mycena osmundicola* and switch to *Armillaria mellea* later (i.e. serial monophagy). There are no reports of other food sources for *G. elata* and the switch appears to be necessary for life cycle progression (Xu & Guo 2000). The same applies to *Tipularia discolor* (McCormick *et al.* 2004), along with a successional change in the growing environment (Rasmussen & Whigham 1998).

Fungal switch may be a more wide-spread phenomenon, however. The sporadic occurrence of various *Rhizoctonia* mycelia in adult orchids otherwise feeding on ectomycorrhizal fungi as noted by Taylor *et al.* (2003), Selosse *et al.* (2004) and Girlanda *et al.* (2006) might be traces of persisting germination fungi. This parallels the situation in which *Rhizoctonia* pelotons are sporadically found in orchid species that go almost entirely phototrophic soon after germination (Bayman *et al.* 2002).

Conservational implications

The identification of the fungi carries a great deal of information about the natural requirements of the orchid species, since the ultimate food sources may be identified, be it leaf litter, woody debris or certain live host trees (Table 1). In a conservation context that would enable the detection of recruitment sites or encouragement of new ones (Batty *et al.* 2001).

A broad food selection may render an orchid species comparatively robust to environmental changes. On the other hand, the generalist strategy is considered costly in terms of defence mechanisms to keep the fungi from becoming virulent. The mono- or oligophagous orchid can be optimally adapted to a narrow food selection but is more likely to experience food limitation that might prevent sexual reproduction and threaten individual survival, if photosynthesis is not a sufficient option. It would also be more dependent on the quality of this narrow food base.

Assessing the relative importance of phototrophic assimilation is also important, because this identifies the light requirements of the orchid species in question. The epilithic *Lepanthes rupestris* appears to be an example of fungal dependency ending soon after germination (Bayman *et al.* 2002), pelotons being extremely rare in the roots of young and more mature plants at two sites studied (but no leafless seedlings were seen). The same seems to apply to several species of *Cypripedium* and many epiphytic species, the canopy environment probably offering opportunities for a largely phototrophic existence. The holomycotrophic species, of course, represent the other extreme, being able to survive in deep

shade or even as entirely subterraneans (Rhizanthella).

Over and above specific inherited trophic traits there is, of course, in many orchids a phenotypic plasticity in respect to mycotrophic persistence, which is influenced by the growing conditions offered at each site and time. For instance, a lack of mycorrhizal infection in adult plants needs not be interpreted as an inherently low dependence on mycotrophy. The plants in question could simply be optimizing their individual survival in an environment with much light and exhausted fungal food sources.

Evolutionary considerations

Rhizoctonia-based seedling mycotrophy was probably the first step in the evolution of orchid mycorrhiza, possibly from an arbuscular mycorrhiza-dependent ancestor, and hence is a plesiomorphic condition within Orchidaceae. The adult orchid would be expected to be at first predominantly phototrophic, as in non-mycorrhizal or arbuscular-dependent ancestors. However, an obvious adaptation to a rich fungal food supply and/or limiting light would be a paedomorphic extension of the

Stage 4: Germination, seedling and adult mycotrophy

using non-rhizoctonioid mycelia. High mycotrophic efficacy without risky change of host. innovation: ability to germinate utilizing non-rhizoctonioid mycelia Stage 3: Germination and seedling mycotrophy using rhizoctonioid mycelia, change to non-rhizoctoniods mycelia for adult mycotrophy. Higher mycotrophic efficacy on a potentially more stable fungal host, utilizing a broader range of substrates. Fungal switch may accommodate successional changes. innovation: ability to utilize non-rhizoctonioid mycelia Stage 2: Germination, seedling and adult mycotrophy using rhizoctonioid mycelia. Mixed phototrophic/mycotrophic strategy in adults with an option for holomycotrophy. innovation: paedomorphic retention of seedling mycotrophy. Stage 1: Germination and seedling mycotrophy using rhizoctonioid mycelia. Fast growth of young plants from very small seeds. innovation: ability to control rhizoctonioid mycelia

FIGURE 1. Hypothetical steps in the evolution of orchid mycorrhiza from stage 0, the non-orchid ancestor. See text for actual examples of species demonstrating the stages. Obligate ontogenetic switch of fungal host is known to occur in stage 3 orchids. It is likely that stages 1-4 have evolved several times, but it is unknown to what extent reversal may happen.

Stage 0: VAM root symbiosis

Rhizoctonia-dependency into adult life history (Fig. 1, stage 1 to 2).

Retention of this seedling mycotrophy combined with alternative fungal food sources in adult plants could be the next step in optimization of mycophagy. This evolution in orchids might be accelerated by ample available biomass of fungal species that for some reason are unable to trigger orchid seed germination. The challenges would consist of inducing initial invasion and peloton formation from mycobionts whose biology does not predispose them for entering living plant tissue, and futhermore developing novel defence mechanisms taylored to keep that infection under control (Fig. 1, stage 3).

The ultimate adaptation to such alternative food sources would be evolution of compatibility of the fungus to orchid seeds and the germination process (Fig. 1, stage 4). So far, germination by non-Rhizoctonia (in the broadest sense) has only been documented within a few, advanced orchid groups: Tipularia and Corallorhiza, Gastrodia (Table 1) and possibly Cyrtosia (Galeola) septentrionalis (discussed in Rasmussen, 1995).

We do not know if there is any impact on fungal fitness and evolution by this symbiosis. One might speculate that orchid predation is too slight to impact on fungal life strategies. As for the fitness, low fruiting body production has been reported in mycelia that support orchids as compared to mycelia of related fungal species (Jones & Smith, 2004, Taylor & Bruns, 1999).

Conclusions

- Orchid mycorrhiza is a non-mutualistic symbiosis and it is practical to think of it as a predator-prey or parasite-provider relationship, with the orchid as the beneficiary. Terms implying mutualism or defection from a presupposed mutualism are mislegding.
- The entire life history is important in conservation of orchid species. Fungi that assist in germination are essential.
- Fungi involved in the various life phases need to be identified and their contribution to growth of the orchid assessed, be it brief or lasting, high or low.

 The ultimate food source in mycotrophy, i.e., the substrate for fungal preys needs to be rated as a maintaining factor for the orchid population in question.

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