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### **Chapter 39 Ecology and Evolution of Fungal-Bacterial Interactions**



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This is the author's final version of the contribution published as:

Olsson, Stefan; Bonfante, Paola; Pawlowska, Teresa E.. Chapter 39 Ecology and Evolution of Fungal-Bacterial Interactions. CRC Press Taylor & Francis Group. 2017. pp: 563-584.

in

The Fungal Community Its Organization and Role in the Ecosystem,Fourth Edition

The publisher's version is available at: http://www.crcnetbase.com/doi/pdf/10.1201/9781315119496-40

When citing, please refer to the published version.

Link to this full text: http://hdl.handle.net/

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# **Ecology and Evolution of Fungal-Bacterial Interactions**

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#### 32 **I. General Introduction**



### 49 **II. Definitions and Concepts**

50 We use the term **symbiosis** in the de Bary's sense of "the living together of unlike organisms", 51 without implications whether this relationship has positive or negative fitness consequences for 52 any of the interacting partners (Martin and Schwab 2012). Thus in terms of fitness outcomes, the 53 symbiosis can assume the forms of a **mutualism** (+/+), **commensalism** (+/0), and **antagonism**,











**Eliminato:** Staněk 1984



181 *Pseudomonas aeruginosa* attracted a lot of attention, which, in turn, yielded important insights 182 into the molecular mechanisms that underlie the coexistence of these two organisms in the 183 context of human disease (Peleg, Hogan, and Mylonakis 2010). *C. albicans* is a commensal 184 yeast found in the normal microbial flora of human oral, digestive, or vaginal mucosa (McManus 185 and Coleman 2014). It is acquired at birth or during physical contact. Factors affecting the 186 mucosal microbiome, such as the use of antibiotics, hormonal imbalance, or diet, can induce 187 non-life threatening *C. albicans* infections of mucosal surfaces, candidiasis (Scully, el-Kabir, and 188 Samaranayake 1994). In severely ill and immunocompromised individuals, *C. albicans* can 189 spread into the blood stream causing invasive and often fatal candidaemia (Eggimann, Garbino, 190 and Pittet 2003). *C. albicans* invasions of host tissues are associated with a morphogenic switch 191 from yeast-like to filamentous growth, which can be induced by changes in environmental

192 conditions, such as shifts in temperature and pH (Berman and Sudbery 2002).

193 *C. albicans* history is intimately linked with the history of humans. Phylogenetic data 194 suggest that its diversification occurred  $\sim$ 3 to 16 MYA and coincided with the evolution of early 195 hominids (Lott et al. 2005). Moreover, it is believed that humans are the main environmental 196 reservoir of *C*. *albicans* (Angebault et al. 2013). In contrast to *C. albicans*, *P. aeruginosa* is a 197 ubiquitous microbe that can be isolated from diverse environments, including humans (Lister, 198 Wolter, and Hanson 2009). However, unlike *C. albicans*, it is rarely a member of the normal 199 microbial flora in humans. Instead, it is a causal agent of community-acquired and, more often, 200 nosocomial infections in individuals who are immunocompromised or suffered a breach in 201 cutaneous or mucosal barriers. The recently observed rise in opportunistic *P. aeruginosa*  202 infections appears to be related to the ability of this microbe to rapidly develop multidrug-203 resistant phenotypes. 204 Mixed infections in which *P. aeruginosa* coexists with *C. albicans* often occur in patients 205 with burn wounds (Gupta et al. 2005) and chronic lung diseases (Hughes and Kim 1973).In 206 such infections the two organisms display an array of antagonistic interactions centered on 207 competition for the host resources and mediated by several mechanisms.For example, *C.*  208 *albicans* responds to the *P. aeruginosa* quorum-sensing signal 3-oxo-C12 homoserine lactone 209 (3OC12HSL) as well as its 12 carbon chain analogs C12HSL and dodecanol with the inhibition 210 of yeast cell filamentation and conversion of previously formed filaments to yeast cells (Hogan, 211 Vik, and Kolter 2004). These are likely defensive responses, as *P. aeruginosa* can attach to the 212 surface of *C. albicans* hyphae and kill them through the action of phospholipase C and 213 phenazines; yeast cells are not susceptible to *P. aeruginosa* attachment (Hogan and Kolter 2002; 214 Gibson, Sood, and Hogan 2009).





261 for mycophagous bacteria to lyse the fungal hyphae in order to parasitize the fungus, proliferate, 262 and inhibit the fungus efficiently. Some bacteria kill the fungus and multiply without penetrating 263 its cell walls, while others proliferate without any negative effects to the fungus (Cuong et al. 264 2011).

265 With the advent of transcriptomics and proteomics, new insights have been gained into 266 these antagonistic of interactions. For example, dual transcriptomic studies of both the fungus 267 and the bacterium challenging each other on agar plates focused on interactions between 268 *Aspergillus niger* and *Collimonas fungivorans* (Mela et al. 2011) as well as *Rhizoctonia solani* 269 and *Serratia plymuthica* (Gkarmiri et al. 2015; Neupane et al. 2015). In these studies, the 270 partners were not allowed to come into physical contact but could exchange metabolites, and in 271 both cases the portion of the fungal colony that was transcriptionally profiled was the one 272 adjacent to the inhibition zone. Both studies found that the fungi reacted by upregulating defense 273 responses (detoxification, efflux pumps), changes to membrane permeability, and increased 274 oxalate production. In contrast, the only response common in bacteria was the upregulation of 275 genes involved in production of secondary metabolites (Mela et al. 2011; Gkarmiri et al. 2015). 276 The two interactions were in many other ways quite different. The *Aspergillus-Collimonas* 277 interaction was mainly characterized by a competition for nitrogen (Mela et al. 2011), while the 278 *Rhizoctonia-Serratia* interaction involved a mutual chemical warfare, as both the fungus and the 279 bacteria upregulated transcription of genes responsible for secondary metabolites/toxins and 280 defenses (Gkarmiri et al. 2015; Neupane et al. 2015). 281 Another example of fungal-bacterial antagonistic interactions comes from *Magnaporthe* 

- 282 *oryzae* transcriptional responses after direct contact with *Lysobacter enzymogenes*, both a wild
- 283 type (WT) strain and a mutant strain deficient in virulence (Mathioni et al. 2013). Four





329 Like with trophic interactions in which bacteria feed on fungi, fungal predation and 330 farming of bacteria are most likely widespread and underappreciated features of terrestrial 331 ecosystems. While some of them can be readily reproduced under laboratory conditions, others 332 need to be studied *in situ* in their natural environments to understand how they connect to more 333 conventional food webs.

334

335 **E.** Highways carrying hyphae-associated bacteria

336 Fungal hyphae expanding in and through unsaturated soil can spread in a soil volume easier than 337 bacteria, as they can bridge over aerial pores and other hydrophobic regions (Kohlmeier et al. 338 2005). The surfaces of the fungus assimilatory hyphae are hydrophilic and thus the fungal 339 hyphae form hydrophilic tracks through soil. These tracks are referred to as **fungal highways** 340 that the bacteria can follow and are generally regarded as beneficial to both the host and the 341 bacterial symbionts (Kohlmeier et al. 2005). The fungal highways have been studied in relation 342 to dissemination of pollutant-degrading bacteria (Kohlmeier et al. 2005; Furuno et al. 2010). In 343 particular, it has been shown that the fungal hyphae might not just help to spread the bacteria but 344 could also function as conduits of pollutants to bacteria (Banitz et al. 2014; Furuno et al. 2010; 345 Wick et al. 2007). In this respect, substrate is channeled from a source along the hyphae to 346 bacteria that are associated with these hyphae. The fungal host seems to nourish the bacterial 347 symbionts inhabiting and spreading on the highways (Bravo et al. 2013; Nazir et al. 2013). The 348 number of bacterial taxa associating and travelling along the fungal highways is probably a 349 combination of selection for the specific prevalent conditions, available substrates, and also by 350 direct activities of the host, *e.g.* a consequence of mutualist recognition or absence of parasite 351 recognition. Bacterial motility by flagella as well as other types of motility have been suggested



376 **F.** Mycorrhiza helper bacteria

377 Mycorrhizal fungi form with the roots of terrestrial plants symbiotic associations of distinct 378 morphologies and functions, collectively referred to as mycorrhizas (Smith and Read 2008). In 379 the most common among them, ecto- and arbuscular mycorrhizas, fungi facilitate plant mineral 380 nutrient uptake from the soil in return for photosynthetic carbon. As a consequence, these 381 symbioses are of great significance in both natural and managed ecosystems, with a particular 382 impact on agriculture and forestry. Current observations indicate that mycorrhizas are, in fact, 383 complex multipartner interactions (Bonfante and Anca 2009), due to the presence of bacteria that 384 can be either loosely or tightly associated with mycorrhizal fungi (Jansa, Bukovská, and 385 Gryndler 2013; Bianciotto et al. 2001; Perotto and Bonfante 1997). Garbaye (1994) pioneered 386 the work on these associations with the now widely accepted term **mycorrhiza helper bacteria,**  387 **MHB**, which defines bacteria that help mycorrhizal establishment. Since the time of MHB 388 discovery and thanks to the advent of the omics era, new knowledge and insights have 389 accumulated, with a particular focus on the microbiota present in the rhizosphere and endosphere 390 of poplar (*Populus*). 391 As a host for both ecto- and arbscular mycorrhizal fungi (AMF), poplar is an excellent 392 model for understanding interactions that govern establishment and functioning of mycorrhizal 393 symbioses, including the role of MHB. For example, the genomes of 21 strains of *Pseudomonas* 394 isolated from the *Populus deltoides* rhizosphere and endosphere have been sequenced (Brown et 395 al. 2012), giving rise to extensive genetic and bioinformatic resources. As a further step, these 396 bacterial isolates were screened for MHB effectiveness expressed as the effects on the *Laccaria*  397 *bicolor* S238N growth rate, mycelial architecture, transcriptional changes and symbiosis with



421 compounds. Moreover, the α- and β-proteobacteria-dominated community of *T. borchii* was able 422 to produce thiophene volatiles from *T. borchii* fruiting body extract, irrespective of their isolation 423 source (truffle or other sources).

424 The complexity of interactions between fungi and both MHB and sporocarp-associated 425 bacteria makes them uniquely difficult to study. However, the tools of symbiotic network 426 construction and testing the applicability of the GMC model to these systems may provide 427 structured approaches to make rapid progress in understanding of these systems.

428

429 **G.** Recognition and assembly of the non-heritable symbionts to form the fungal-bacterial 430 metaorganism

431 Both plant and animal epithelial surfaces coming in contact with bacteria share a similar problem

432 in that they should actively select for beneficial/commensal bacteria and discourage the

433 colonization by antagonists (McFrederick et al. 2012; Artis 2008; Ausubel 2005; Zamioudis and

434 Pieterse 2012). Innate immunity recognition of bacterial cues as MAMPs (microbial associated

435 molecular patterns) plays a key role in this selection in both plants and animals (Artis 2008;

436 Nürnberger et al. 2004). However, the immune reaction is balanced so as not to kill eventual

437 beneficial bacteria, as is done in tissues not normally colonized by bacteria (Artis 2008;

438 Zamioudis and Pieterse 2012). Fungal hyphae growing in most natural environments face a

439 similar need to promote the beneficial and inhibit the antagonistic microbes. Fungal reactions to

440 a bacterial MAMP have been demonstrated (Xu et al. 2008), the existence of innate immunity

441 type recognition has been suggested (Paoletti and Saupe 2009; Paoletti, Saupe, and Clavé 2007),

442 and recently transcriptomic innate immunity type responses have been found in fungi (Ipcho et

443 al. 2016). Fungal innate immunity is thus most likely involved in the active selection for









536 Even though some of the features displayed by the *Rhizopus-Burkholderia* symbiosis are 537 typical for a mutualism, the *Burkholderia* endobacteria appear to be facultative endosymbionts, 538 capable of living both inside and outside eukaryotic cells, a lifestyle similar to that of pathogenic 539 *Legionella*, *Salmonella*, or *Bartonella*. This duality, combined with the ease of experimental 540 manipulation, propelled the *Rhizopus-Burkholderia* symbiosis to become a model for studying 541 the evolution of heritable symbioses. In particular, addressing questions concerning its 542 evolutionary origins, whether it started with the partners interacting as antagonists (Fig. 2), and 543 whether it has already achieved evolutionary stability (Fig. 3) will be a source of rich insights not 544 only into the genetic mechanisms of symbiont vertical transmission but also into other facets of 545 partner coevolution. 546 547 **V. Heritable Symbiotic Interactions** 548 **A.** Introduction

549 As discussed in the preceding sections, symbiont vertical transmission is a principal factor 550 contributing to both the establishment and stability of mutualisms. Importantly, vertical 551 transmission is not exclusive to mutualisms; it can also occur in antagonistic interactions. 552 Vertical transmission can be strict or mixed. In strict vertical transmission symbionts are 553 transferred from a parent exclusively to offspring. In mixed transmission, in addition to being 554 passaged between generations, symbionts move horizontally between members of the same 555 generation. Symbioses with strict vertical transmission are characterized by congruity of partner 556 phylogenetic histories, consistent with partner codiversification (Page 2003). In symbioses with 557 mixed transmission, the extent of horizontal transmission determines the degree of incongruity 558 between partner phylogenies. Interestingly, strict vertical transmission of symbionts tends to be

559 associated with reciprocally obligate partner dependence, whereas mixed transmission is found 560 in associations in which either one or both partners are facultatively dependent on the symbiosis 561 (Fig. 3).

562 Importantly, while in fungi all known heritable associations involve endobacteria that 563 reside inside fungal cells, not all associations formed by fungi with endobacteria are known to be 564 heritable. In heritable symbioses, bacteria are either facultatively or obligately dependent on the 565 fungus. The *Burkholderia* symbionts of *Rhizopus,* discussed in the previous section, as well as 566 *Rhizobium radiobacter* in the root-colonizing *Piriformospora indica* (Sharma et al. 2008) 567 represent facultative heritable endobacteria. In contrast, obligate heritable endosymbionts 568 include two groups of bacteria associated with AMF, *Ca.* Glomeribacter gigasporarum 569 (Bianciotto et al. 2003) and the mycoplasma-related endobacteria, MRE (Naumann, Schüßler, 570 and Bonfante 2010). It is unclear whether the unnamed heritable endosymbiont of *Mortierella*  571 *elongata* (Sato et al. 2010) is a facultative or obligate endobacterium. Remarkably, we are not 572 aware of heritable fungal-bacterial symbioses in which the interacting partners are obligately 573 dependent on each other. Such associations are common in insects, which depend on 574 endobacteria for provision of essential nutrients (McCutcheon and Moran 2012). It remains to 575 be investigated whether this knowledge gap represents a true dearth of reciprocally obligate 576 fungal-bacterial interactions or a detection bias. Recent accumulation of newly discovered 577 associations that involve non-heritable endobacteria suggests that the latter might be the case. 578 Such non-heritable associations include, among others, *Helicobacter pylori* in *Candida albicans*  579 (Siavoshi and Saniee 2014), *Nostoc punctiforme* in *Geosiphon pyriforme* (Schüßler et al. 1994), 580 *Bacillus* spp. in *Ustilago maydis* (Ruiz-Herrera et al. 2015), α-proteobacteria in the 581 ectomycorrhizal fungus *Laccaria bicolor* (Bertaux et al. 2005; Bertaux et al. 2003), and diverse

582 bacteria that inhabit hyphae of phylogenetically diverse fungal endophytes of plants (Hoffman

583 and Arnold 2010). Due to the lack of sufficient data from other systems, our discussion in the

584 following two sections will focus on *Ca.* Glomeribacter gigasporarum and MRE associated with

586

585 AMF.

587 **B.** Heritable facultative mutualisms

588 *Ca.* Glomeribacter gigasporarum, referred hereafter as *Glomeribacter*, is a stable, and 589 structurally integrated endosymbiont found in many representatives of the AMF family 590 Gigasporaceae (Bianciotto, Bandi, et al. 1996; Bianciotto et al. 2003; Mondo et al. 2012). It 591 thrives inside the fungal cells along the different stages of the fungal life cycle, always located 592 inside a compartment structurally resembling a fungal vacuole (Bianciotto, Minerdi, et al. 1996). 593 On the fungal side, the Gigasporaceae, like other AMF, form symbiotic associations with roots 594 of many plants, and may proliferate also in the absence of the endobacteria (Lumini et al. 2007), 595 giving rise to an association that is obligate for the bacterial partner and facultative for the fungal 596 host. A similar disparity is true for all AMF, as they fully depend on their host plants for energy, 597 while plants may complete their life cycle in the absence of AMF. 598 While biodiversity studies have demonstrated that *Glomeribacter* is widespread, they 599 have not identified factors responsible for the evolutionary stability of the Gigasporaceae-600 *Glomeribacter* symbiosis, which dates back to the early Devonian (Mondo et al. 2012). The 601 *Glomeribacter* genome sequencing revealed that this endobacterium has a reduced genome of 602 1.7 Mb (Ghignone et al. 2012), consistent with its uncultivable status (Jargeat et al. 2004). It 603 lacks metabolic pathways leading to important amino acids, but has many amino acid permeases 604 for uptake of nutrients from the fungus, as expected of an endobacterium that depends on its host

605 for nutrients and energy (Fig. 4). Interestingly, the whole operon for biosynthesis of vitamin B12 606 is present in the *Glomeribacter* genome, but it is not clear whether this might represent any 607 benefit for the fungus. In contrast to animals, which use B12-dependent enzymes for methionine 608 synthesis and methylmalonate metabolism, fungi and land plants rely on B12-independent 609 enzymes for these pathways (Young, Comas, and de Carvalho 2015)*.* Consistent with this 610 expectation, the genome of a model AMF, *Rhizophagus irregularis,* encodes B12-independent 611 enzymes (Tisserant et al. 2013). 612 While the significance of *Glomeribacter* to the AMF hosts could not be gleaned from its

613 genomic sequence, the availability of a stable endosymbiont-free AMF *Gigaspora margarita* 614 BEG34 line, designated as  $B(-)$ , allowed for direct comparisons with the line containing the 615 endobacterium, B(+). These comparisons revealed several differences, both phenotypic (Lumini 616 et al. 2007) and transcriptional (Salvioli et al. 2016), that speak to the role of *Glomeribacter* in 617 the AMF host. For example, the  $B(-)$  AMF line was able to colonize its plant host but was 618 impaired in mycelial growth and spore production compared to the  $B(+)$  line (Lumini et al. 619 2007). Moreover, benefits of the endosymbiont presence appeared to extend to the plant host, as 620 the phosphate measurements in *Lotus japonicus* plants revealed a statistically higher phosphate 621 quantity in the symbiosis established by the  $B(+)$  versus the  $B(-)$  AMF line (Salvioli et al. 2016). 622 In turn, the transcriptome analysis showed that the endobacterium had a stronger effect on the 623 pre-symbiotic phase of the fungus, supporting earlier phenotypic observations that 624 *Glomeribacter* promotes germ tube extension in the AMF host (Lumini et al. 2007; Salvioli et al. 625 2016). Coupling of transcriptomics with physiological and cell biology approaches 626 demonstrated that the bacterium increases the AMF sporulation success, raises the AMF 627 bioenergetic capacity, increasing ATP production, and elicits mechanisms to detoxify reactive



651 compensatory evolution/addiction (Aanen and Hoekstra 2007).

652 Despite the remarkable progress made recently in understanding the Gigasporaceae-653 *Glomeribacter* symbiosis, there are many outstanding questions. For example, it remains unclear 654 what factors keep this association from evolving towards reciprocally obligate partner 655 dependence predicted by evolutionary theory (Fig. 3). It could be speculated that the benefits to 656 the AMF host depend on the environmental context and the association may break up when the 657 cost of supporting the endosymbiont becomes prohibitive. This scenario would explain why the 658 endobacteria in the Gigasporaceae-*Glomeribacter* symbiosis appear to retain the potential to 659 transmit horizontally and exchange genes, attributes that may have contributed to their 660 evolutionary longevity (Mondo et al. 2012). 661 662 **C.** Heritable antagonisms 663 The symbiosis between AMF and MRE (mycoplasma related endobacteria) represents an 664 outstanding deviation from the molecular evolution patterns both expected by evolutionary 665 models and detected thus far in heritable endobacteria (McCutcheon and Moran 2012), including 666 *Glomeribacter* (Mondo et al. 2012). In particular, MRE display extraordinary intra-host 667 diversity of their 16S rRNA gene (Naumann, Schüßler, and Bonfante 2010; Desirò et al. 2014; 668 Desirò et al. 2015; Toomer et al. 2015) and genomic sequences (Naito, Morton, and Pawlowska 669 2015; Torres-Cortés et al. 2015). In part, this diversity could be attributed to a high mutation 670 rate, related to the loss of DNA repair machinery from the MRE genomes, combined with the 671 apparent activity of mechanisms contributing to genome plasticity, such as recombination 672 machinery and mobile genetic elements (Naito, Morton, and Pawlowska 2015; Naito and 673 Pawlowska 2016). While the mechanisms responsible for genome plasticity are not expected to



696 vulnerable to genomic degeneration and extinction (McCutcheon and Moran 2012). Yet, co-



708 **A.** Introduction

709 The establishment and outcomes of the fungal bacterial interactions are most probably a result of 710 chemical communication where a compound from one partner elicits a response with another 711 compound from the other partner (Baruch et al. 2014; Piispanen and Hogan 2008; Xu et al. 2008; 712 Badri et al. 2009; Nazir et al. 2010; Schroeckh et al. 2009; Sengupta, Chattopadhyay, and 713 Grossart 2013). This is typical for "ping-pong" type communications, where a communication 714 from one interacting partner draws a response from the other partner (Griffin 2012). The correct 715 order of events in ping-pong communication, rather than unique metabolites, could be selective 716 and instrumental in establishing the relationship (like a combinatorial lock). With the advent of 717 modern omics, these ping-pong events could be studied using transcriptomics (Mela et al. 2011; 718 Gkarmiri et al. 2015; Neupane et al. 2015; Mathioni et al. 2013), proteomics (Moretti et al. 719 2010), and aided with metabolomics, allowing for hourly resolution of events during the



742 **C.** Physiological processes known from other host-symbiont systems

- 743 In this section, we list a few physiological processes known from other host-microbe systems
- 744 that are also likely to be involved in fungal-bacterial interactions.
- 745 **Extracellular vesicle trafficking:** All organisms can produce extracellular vesicles (Deatherage
- 746 and Cookson 2012). In fungal pathogens of humans, these exosomes are important in
- 747 interactions with the host (Rodrigues et al. 2014), whereas in bacteria they play a role in biofilm
- 748 communication between cells (Remis et al. 2014; Kulp and Kuehn 2010) and interaction with
- 749 other bacteria (Kulp and Kuehn 2010; Vasilyeva et al. 2013).
- 750 **Transfer of interfering RNA:** Extracellular vesicles have been shown to sometimes carry small
- 751 RNA (Samuel et al. 2015) or DNA (Kulp and Kuehn 2010), which opens up possibilities for
- 752 interfering with partner organisms (Nicolás and Ruiz-Vázquez 2013).
- 753 **Unconventional secretion:** Fungi, like all eukaryotes, secrete proteins mainly through the ER-
- 754 Golgi pathway using N-terminal signal peptides to guide the proteins into the pathway. Proteins
- 755 without signal peptides can also be secreted through unconventional secretion pathways (Zhang
- 756 and Schekman 2013). These pathways are important during interaction between host and
- 757 microorganisms in both plant and animal systems (Ding, Robinson, and Jiang 2014; Öhman et al.
- 758 2014) and additionally also involved in the production of extracellular vesicles (Zhang and
- 759 Schekman 2013).
- 760 **Priming of responses against pathogens by beneficial organisms:** Beneficial bacteria are
- 761 recognized by similar systems as pathogens and can induce enhanced immune functions against
- 762 later attacks by pathogens, thus priming the defenses. Such priming responses are a hot topic in
- 763 both plant and animal systems (Chu and Mazmanian 2013; Conrath 2009; Aranega-Bou et al.
- 764 2014; Val et al. 2008) and can be expected to be important for both non-heritable and heritable
- 765 fungal bacterial interactions.


## 787 **Acknowledgements**

788 We thank Olga Lastovetsky for comments on the manuscript. This work was supported by the

- 789 National Science Foundation grant IOS-1261004 to TEP and the Torino University 60% grant to
- 790 PB.

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- 1376 Zhang, M., and R. Schekman. 2013. Unconventional secretion, unconventional solutions.
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- 1379 Table 1. Mechanisms shared by diverse eukaryotic hosts to select beneficial organisms
- 1381 colonizing host surfaces involved in nutrient uptake.





1383 **Figure Captions** 

1384 **Figure 1. Metaorganisms comprise fungal hosts and their various bacterial symbionts.** 1385

1386 **Figure 2. Evolutionary theory predictions on the role of vertical transmission in the**  1387 **evolution of mutualisms from antagonisms.** Hosts are depicted as red ovals; host-positive 1388 symbionts are shown as green dots, host-negative symbionts as purple dots. Relative host fitness 1389 is reflected by the size of ovals. 1390 1391 **Figure 3. Hypothetical evolutionary trajectories in heritable mutualisms.** Hosts are 1392 depicted as red ovals; endosymbionts are shown as green dots. Relative host fitness is reflected 1393 by the size of ovals. (A) Evolutionary trajectory leading to obligate reciprocal partner 1394 dependence. (B) Shifting environmental conditions are expected to arrest an association at the 1395 facultative dependence stage. If conditions remain unfavorable for prolonged periods of time, 1396 host populations would be expected to completely lose endosymbionts. Modified from Mondo et 1397 al. (2012). 1398 1399 **Figure 4. Model of plant-fungus-endobacterium interaction** (Courtesy of M. Novero). 1400 Genome-sequencing results for *Candidatus* Glomeribacter gigasporarum indicate that the 1401 bacterium fully depends on the fungal metabolism, including carbon (C), phosphorus (P), and

1402 nitrogen (N) metabolism. In contrast, the fungus depends on its green plant host for C uptake

1403 only.

1404

#### 1405 **Figure 5. Evolutionary theory predictions linking the type of symbiosis with the intra-host**

1406 **relatedness of symbionts and symbiont transmission**.Hosts are shown as red ovals. Relative

- 1407 host fitness is reflected by the size of ovals. Endosymbionts are represented by green and purple
- 1408 dots with different shades depicting different genotypes. Modified from Toomer et al. (2015).

1410 Figure 1



## 1413 Figure 2



## 1416 Figure 3





1419 Figure 4


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1423

Figure 5 1422



 $71\,$