THE EVOLUTION OF AGRICULTURE IN INSECTS

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Abstract  Agriculture has evolved independently in three insect orders: once in ants, once in termites, and seven times in ambrosia beetles. Although these insect farmers are in some ways quite different from each other, in many more ways they are remarkably similar, suggesting convergent evolution. All propagate their cultivars as clonal monocultures within their nests and, in most cases, clonally across many farmer generations as well. Long-term clonal monoculture presents special problems for disease control, but insect farmers have evolved a combination of strategies to manage crop diseases: They (a) sequester their gardens from the environment; (b) monitor gardens intensively, controlling pathogens early in disease outbreaks; (c) occasionally access population-level reservoirs of genetically variable cultivars, even while propagating clonal monocultures across many farmer generations; and (d) manage, in addition to the primary cultivars, an array of “auxiliary” microbes providing disease suppression and other services. Rather than growing a single cultivar solely for nutrition, insect farmers appear to cultivate, and possibly “artificially select” for, integrated crop-microbe consortia. Indeed, crop domestication in the context of coevolving and codomesticated microbial consortia may explain the 50-million year old agricultural success of insect farmers.

1. INTRODUCTION

The cultivation of crops for nourishment has evolved only a few times in the animal kingdom. The most prominent and unambiguous examples include the fungus-growing ants, the fungus-growing termites, the ambrosia beetles and,
course, humans. For humans, who started the transition from an ancestral hunter-gatherer existence to farming only about 10,000 years ago (Diamond 1997, Smith 1998), sustainable, high-yield agriculture has become critical for survival in a global economy with projected food shortages, and diverse research programs are currently devoted to the optimization of agricultural productivity in the context of growing environmental challenges (Green et al. 2005). Agricultural progress has been achieved by humans through a combination of insight, creative planning, and a fair share of contingency and luck (Diamond 1997, Schultz et al. 2005, Smith 1998). However, humans have so far not examined nonhuman agricultural systems, such as the fungus-growing insects, for possible insights to improve agricultural strategies.

This lack of an applied interest in insect agriculture probably derives from a general perception that human agricultural systems (based largely on plant cultivation) function in a fundamentally different manner than insect systems (all based on fungus cultivation). However, humans have learned much of practical value through the close examination of adaptive features of other organisms (including insects), and comparable problems such as crop diseases affect all farmers regardless of their phylogenetic positions or those of their crops (plant, fungus, or otherwise). Because of the universality of crop diseases in both human and insect agriculture, it may be fruitful to examine the short-term and long-term solutions that have evolved convergently in insect agriculture for possible application to human agriculture (Denison et al. 2003). Such a synthesis is the goal of this review.

1.1. Behavioral and Nutritional Elements

Defining Agriculture

Insect fungiculture and human farming share the defining features of agriculture (see Table 1): (a) habitual planting (“inoculation”) of sessile (nonmobile) cultivars in particular habitats or on particular substrates, including the seeding of new gardens with crop propagules (seeds, cuttings, or inocula) that are selected by the farmers from mature (“ripe”) gardens and transferred to novel gardens; (b) cultivation aimed at the improvement of growth conditions for the crop (e.g., manuring; regulation of temperature, moisture, or humidity), or protection of the crop against herbivores/fungivores, parasites, or diseases; (c) harvesting of the cultivar for food; and (d) obligate (in insects) or effectively obligate (in humans) nutritional dependency on the crop. Obligate dependencies of the insect farmers can be readily demonstrated by experimental removal of their cultivated crops, resulting in reduced reproductive output, increased mortality, or even the certain death of the cultivar-deprived insect (Francke-Grosmann 1967, Grasse 1959, Norris 1972, Sands 1956, Weber 1972). Our definition of agriculture does not require conscious intent in planting and harvesting. Conscious planning, learning, and teaching have clearly accelerated the development of complex agriculture in humans, but presumably not in insects (Schultz et al. 2005).
TABLE 1  Agricultural behaviors of farming ants, termites, beetles, and humans

<table>
<thead>
<tr>
<th>Agricultural behavior</th>
<th>Attine ants</th>
<th>Macrotermite termites</th>
<th>Xyloborine ambrosia beetles</th>
<th>Humans</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dependency on crop for food</td>
<td>Obligate</td>
<td>Obligate</td>
<td>Obligate</td>
<td>Facultative</td>
</tr>
<tr>
<td>Engineering of optimal growth conditions for crop (e.g., substrate preparation; moisture or humidity regulation)</td>
<td>Present</td>
<td>Present</td>
<td>Present</td>
<td>Present</td>
</tr>
<tr>
<td>Planting of crop on improved substrate</td>
<td>Present</td>
<td>Present</td>
<td>Present</td>
<td>Present</td>
</tr>
<tr>
<td>Intensive, continuous monitoring of growth and disease status of all crops</td>
<td>Present</td>
<td>Present</td>
<td>Present</td>
<td>Absent</td>
</tr>
<tr>
<td>Sustainable harvesting of crop for food</td>
<td>Present</td>
<td>Present</td>
<td>Present</td>
<td>Present</td>
</tr>
<tr>
<td>Protection of crop from diseases and consumers</td>
<td>Present</td>
<td>Present</td>
<td>Present</td>
<td>Present</td>
</tr>
<tr>
<td>Weeding of alien organisms invading the garden</td>
<td>Present</td>
<td>Unknown</td>
<td>Present</td>
<td>Present</td>
</tr>
<tr>
<td>Use of chemical herbicides to combat pests</td>
<td>Present</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Present</td>
</tr>
<tr>
<td>Use of microbial symbionts for nutrient procurement for the crop</td>
<td>Present</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Present</td>
</tr>
<tr>
<td>Use of disease-suppressant microbes for biological pest control</td>
<td>Present</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Absent¹</td>
</tr>
<tr>
<td>Sociality</td>
<td>Strictly eusocial</td>
<td>Strictly eusocial</td>
<td>Subsocial or communal²</td>
<td>Social</td>
</tr>
<tr>
<td>Task partitioning in agricultural processes</td>
<td>Present</td>
<td>Unknown</td>
<td>Present</td>
<td>Present</td>
</tr>
<tr>
<td>Application of artificial selection for crop improvement</td>
<td>Unknown, but ants exert symbiont choice</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Present</td>
</tr>
<tr>
<td>Learning and cultural transmission of agricultural innovations</td>
<td>Absent</td>
<td>Absent</td>
<td>Absent</td>
<td>Present</td>
</tr>
</tbody>
</table>

¹See Section 7 for some recently discovered microbes with potential disease-suppressant properties.
²One ambrosia beetle species is eusocial, all other species appear to be subsocial or communal (see text).
We restrict our review to ant, termite, and beetle fungiculturists. Cases analogous to human animal husbandry, such as the tending by ants of hemipteran insects (e.g., aphids, treehoppers; Hölldobler & Wilson 1990), are beyond the scope of this review. We also exclude cases that fail to meet all four of the requirements of agriculture as defined above, including, e.g., the ant Lasius fuliginosus, which promotes fungal growth in the walls of its nest, because the fungus is apparently not grown for food but instead for strengthening the walls (Maschwitz & Hölldobler 1970) or for antibiotic protection of the walls (Mueller 2002). On the same grounds, we exclude a number of possible cases of incipient agriculture. For example, Littoria snails may “protofarm” fungi by creating plant wounds that become infected with fungal growth that is part of the snails’ diet, but the snails do not actively inoculate the plant wounds or otherwise garden the fungi (Silliman et al. 2003). Many more such protofarming species probably await discovery, particularly among invertebrates, and all of the known insect agriculturists (fungus-growing ants, termites, and beetles) probably originated from comparable protoagricultural ancestors (Mueller et al. 2001, Schultz et al. 2005). Comparison of these protofarming insects with “primitive” human agriculture exceeds the scope of this review.

1.2. A Coevolutionary Approach to Understanding Agriculture

We will analyze agriculture as a type of strong coevolutionary interaction, defined by the nutritional and behavioral criteria summarized above, in which natural selection acts upon both farmers and crops as reciprocally interdependent lineages (Futuyma & Slatkin 1983, Rindos 1984). Our coevolutionary approach to agriculture considers not only the interactions between a specific farming insect and a single cultivated crop, but also its interactions with other pathogenic and mutualistic microbes that have recently been discovered in insect gardens. Like the cultivars, some of these microbes are also managed by the insect farmers for specific purposes (Figure 1). In other words, an insect garden is not a pure monoculture, but a sequestered and engineered ecological community consisting of several interacting microbes, some beneficial and others detrimental to the farmers. To gain a comprehensive understanding of the principles of insect agriculture, it will therefore be necessary to examine the nature of insect-microbe interactions in gardens, the evolutionary origins of these interactions, and the convergent and divergent evolutionary trajectories that culminated in the extant agricultural systems of insects.

2. THE THREE INSECT-AGRICULTURE SYSTEMS

Behaviorally complex systems of insect agriculture are known from only three groups of insects: ant, termites, and beetles.
2.1. Ant Fungiculture

The fungus-growing ants are a monophyletic group of about 220 described and many more undescribed species in the tribe Attini (subfamily Myrmicinae) (Price et al. 2003, Schultz & Meier 1995). Attine ants occur only in the New World (Argentina to the southern United States) and attain their greatest diversity in the wet forests of equatorial South America, the region of their presumed evolutionary origin (Mueller et al. 2001). Attine ants are obligate agriculturists; their cultivated fungi are the sole source of food for the larvae and an important source of food for the adults. Although adults are able to supplement their diets by feeding on plant juices (Bass & Cherrett 1995, Murakami & Higashi 1997), the cultivated fungi are nutritionally sufficient to support the ants even in the absence of additional nutrients (Mueller 2002, Mueller et al. 2001). Garden fungi are transmitted vertically across generations when daughter queens transport small pellets of natal-nest mycelium within their infrabuccal pockets, pouches present in the mouthparts of all ants (Fernández-Marín et al. 2004, Huber 1905, Mueller 2002). In the derived leafcutter ants, the workers are divided into a remarkable range of differently sized morphological castes, each specialized on a different task (Hart et al. 2002, Weber 1972) (Figure 2d).

Different attine ant lineages cultivate their fungi on different substrates. The ancestral gardening substrate, still used by the so-called lower attines, consists of flower parts, arthropod frass, seeds, wood fragments, or other similar plant debris, whereas the leafcutting genera *Atta* and *Acromyrmex* primarily use freshly cut leaves and flowers. Despite these distinct substrate specializations, all attine systems contain at least four symbionts: (a) the fungus-growing ants; (b) their fungal cultivars (basidiomycetes in the mushroom families Lepiotaceae and Pterulaceae; Mueller et al. 1998, Munkacsi et al. 2004); (c) mutualistic antibiotic-producing actinomycete bacteria (family Pseudonocardiaceae; Currie et al. 1999b); and (d) garden parasites in the ascomycete fungal genus *Escovopsis* (Currie et al. 1999a, Currie et al. 2003b). Additional bacteria and yeasts also occur in attine gardens and may function as mutualists, e.g., by secreting digestive enzymes or antibiotics (Carreiro et al. 1997, Craven et al. 1970, Santos et al. 2004).

2.2. Termite Fungiculture

Of the more than 2600 described termite species, about 330 species in the subfamily Macrotermitinae cultivate a specialized fungus, genus *Termitomyces*, for food. Nests are generally founded by a single pair of reproductives, the future queen and king. They seal themselves permanently in a cell of hard clay (the so-called royal chamber) where they rear the first brood of sterile workers. In most termite species, a new colony acquires a fungal strain from wind-dispersed sexual *Termitomyces* spores shortly after nest founding and begins construction of the first gardens (De Fine Licht et al. 2005). These spores come from fruiting bodies (mushrooms) that arise from mature termite colonies. The fruiting of the fungus appears to be roughly synchronized to the period when the first foraging workers emerge from a
new nest, a few months after the nest-founding stage. Termite gardens are grown on dead plant material that is only partially decomposed, such as leaf litter, dead grass, dead wood, or dry leaves.

Termite gardens are built from spore-containing fecal pellets in chambers that the termites construct either inside a mound or dispersed in the soil. Fecal pellets are added continuously to the top of the comb and fungal mycelium rapidly permeates the new substrate (Figure 2e). After a few weeks, the fungus starts to produce vegetative nodules that are consumed by the termites. These nodules are a rich source of nitrogen, sugars, and enzymes. The nodules are also covered with indigestible asexual spores (conidia), so that consumption serves the additional function of inoculating the feces with spores, which pass through the gut unharmed and are then planted in new comb with the deposition of feces (Leuthold et al. 1989). Mature comb is also consumed (Darlington 1994), but it is nutritionally inferior to the nodules.

2.3. Beetle Fungiculture

Ambrosia beetles make up around 3400 of the 7500 species in the weevil subfamily Scolytinae (the bark and ambrosia beetles, including the traditionally separate Platypodinae; Farrell et al. 2001, Harrington 2005, Wood 1982). Most ambrosia beetles construct tunnel systems (galleries; Figure 2f) in woody tissues of trees (typically in weakened or recently dead trees or, more rarely, in vigorous hosts), although some species are specialized to colonize pith, large seeds, fruits, and leaf petioles (Harrington 2005, Wood 1982). The term ambrosia refers to the fungi cultivated by the beetles on gallery walls, upon which they feed as an exclusive, or near exclusive, food source. The beetles are obligately dependent upon the fungi, from which they acquire essential vitamins, amino acids, and sterols (Beaver 1989, Kok et al. 1970).

The most advanced fungiculturists among the ambrosia beetles occur in the Xyleborini, a large monophyletic tribe of about 1300 species (Farrell et al. 2001, Jordal 2002). It is this group of ambrosia beetles that we primarily focus on in this review. Although life histories among the Xyleborini vary considerably, most share a number of fungicultural characteristics. There is a sexual division of labor in the Xyleborini; only females perform gardening tasks, whereas males are short-lived and flightless (Norris 1979). After mating, females disperse to new host substrate, carrying the fungi in specialized pockets termed mycangia. Once within a new host, founding females “plant” the fungi on the walls of the excavated tunnels, lay eggs, and tend the resulting garden and brood (Norris 1979). In ways not fully understood, they are able to control the growth of the fungal crop, as well as, to a degree, the composition of its multiple fungal species (Beaver 1989, French & Roeper 1972, Kingsolver & Norris 1977, Roeper et al. 1980). If the female dies, the garden is quickly overrun by contaminating fungi and bacteria, which ultimately results in the death of the brood (Borden 1988, Norris 1979).

The ambrosia gardens of xyleborine beetles are not pure monocultures as was once believed, but are typically composed of an assemblage of mycelial fungi,
yeasts, and bacteria (Batra 1966, Haanstadt & Norris 1985). These assemblages were termed multi-species complexes by Norris (1965), who suggested that it is a complex as a whole, rather than any one individual microbe, that allows the beetles to exploit nutrient-poor substrates such as wood. However, most subsequent work has revealed that one “primary” fungus always dominates in beetle gardens (Baker 1963, Batra 1966, Gebhardt et al. 2004, Kinuura 1995). Furthermore, the beetles typically carry only the primary fungus in the mycangium (although secondary fungi are sometimes also isolated from mycangia), and the cultivation efforts of female beetles tend to favor the primary fungus, which imparts the greatest nutritional benefit (Francke-Grosmann 1967, Gebhardt et al. 2004, Morelet 1998, Norris 1979). Some auxiliary fungi also support beetle development, but survival on the auxiliary fungi alone is often greatly reduced (Norris 1979). These observations implicate the primary fungus as the intended crop, whereas the secondary fungi, yeasts, and bacteria may be contaminant “weeds” or may play additional auxiliary roles in the gardens, paralleling the hypothesized roles of the auxiliary bacteria and yeasts in attine gardens (see above).

3. EVOLUTIONARY ORIGINS OF INSECT AGRICULTURE

Phylogenetic analyses reveal nine independent origins of insect agriculture (Figure 2; Table 2). In ants, fungal cultivation arose only once, probably 45–65 Mya in the Amazon rainforest (Mueller et al. 2001; Schultz & Meier 1995). In termites, fungiculture likewise had a single origin, approximately 24–34 Mya in the African rainforest (Aanen et al. 2002; D.K. Aanen & P. Eggleton, submitted). In ambrosia beetles, however, agriculture arose independently seven times between 20–60 Mya, six times in various nonxyleborine lineages, and once in the ancestor of the Xyleborini about 30–40 Mya (Farrell et al. 2001). Whereas the common ancestors of the macrotermes and of the xyleborines each domesticated a single, specific primary cultivar clade to which their descendants have adhered throughout subsequent evolution (Figure 2, Table 2), attine ants maintain associations with multiple independently domesticated cultivar lineages (which are for the most part very closely related; Mueller et al. 1998; Munkacsi et al. 2004; Table 2). Interestingly, there are no known cases of reversal from agricultural to nonagricultural life in any of the nine agricultural insect lineages (Figure 2a, 2b, and 2c), suggesting that the transition to fungiculture is a drastic and possibly irreversible change that greatly constrains subsequent evolution.

Two main models have been suggested for the independent evolutionary transitions to agriculture in insects, the “consumption-first” versus the “transmission-first” models (Mueller et al. 2001). In the consumption-first model (the likely model for the termites), an insect lineage initially begins to incorporate fungi into its more generalist diet, then becomes a specialized fungivore, and finally evolves adaptations for cultivating fungi. In the transmission-first model (the likely model for the beetles), the insect lineage begins its association with a fungus by serving
### TABLE 2  Crop ecology and evolution in ant, termite, beetle, and human agriculture

<table>
<thead>
<tr>
<th>Crop ecology &amp; evolution</th>
<th>Attine ants</th>
<th>Macrotermite termites</th>
<th>Xyleborine ambrosia beetles</th>
<th>Humans</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of inferred evolutionary origins of agricultural behavior</td>
<td>Single origin of agriculture in the Attini</td>
<td>Single origin of agriculture in the Macrotermite</td>
<td>Single origin of agriculture in the Xyleborini</td>
<td>Multiple independent origins of agriculture</td>
</tr>
<tr>
<td>Estimated date and region of origin of agricultural behavior</td>
<td>45–65 Mya in Amazonian rainforests</td>
<td>24–34 Mya in African rainforests</td>
<td>30–40 Mya in xyleborine beetles; region of origin unknown</td>
<td>10,000 years ago for the earliest known origins of human agriculture; multiple regions of origin</td>
</tr>
<tr>
<td>Number of crop clades cultivated</td>
<td>Multiple cultivar clades One cultivar clade belongs to the Pterulaceae; at least three additional polyphyletic cultivar clades belong to the Lepiotaecae (tribe Leucocoprineae).</td>
<td>Single cultivar clade All termite cultivars belong to the genus Termitomyces.</td>
<td>Multiple cultivar clades The ambrosia cultivars Ambrosiella and Raffaelleca have a polyphyletic origin within the genera Ophiostoma and Ceratocystis.</td>
<td>Multiple crop clades from diverse lineages of plants and fungi</td>
</tr>
</tbody>
</table>
Crop transmission
Vertical cultivar inheritance from parent to offspring (i.e., cultivar transfer between generations)

Present
Vertical cultivar inheritance is the rule in all attine ant lineages.

Variable
Vertical cultivar inheritance is typical for two independently derived macrotermite lineages; all other macrotermites acquire cultivars horizontally via wind-dispersed spores from other colonies.

Present
Vertical cultivar inheritance is the rule in all xyleborine lineages.

Specialization on crops
Specialization at higher phylogenetic levels

Clade-clade correspondence
Defined clades of ant species are specialized in that they only grow fungi typical for their own ant-specific cultivar clade.

Clade-clade correspondence
Defined clades of termite species are specialized in that they only grow fungi typical for their own termite-specific cultivar clade.

No strict clade-clade correspondence
The fungal lineages associated with xyleborine ambrosia beetles are only distantly related to each other, and all are also associated with some bark beetles.

Not applicable
Not applicable

Specialization at level of farmer species

Present
In all cases studied in detail, single species of ants grow only a single, phylogenetically narrow group of fungi (i.e., a single, phylogenetically defined "species" of fungus).

Present
Single termite species generally associate with multiple fungal lineages, but within the limits of specific cultivar clades (see above); however, some termite species are very specialized on a particular fungus.

Present
Single beetle species generally associate with multiple, distantly related fungal lineages; the primary cultivar is typically a single fungal species, but secondary cultivars vary and often come from different fungal lineages than the primary-cultivar lineage.

Not applicable
Specialization on crops is historically manifested by different human lineages (reflecting separate domestication events), but no single human lineage is nearly as specialized on a single crop as are the insect farmers.

(Continued)
### Table 2  (Continued)

<table>
<thead>
<tr>
<th>Crop ecology &amp; evolution</th>
<th>Attine ants</th>
<th>Macrotermite termites</th>
<th>Xyleborine ambrosia beetles</th>
<th>Humans</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Crop sharing and crop exchange</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lateral exchange of cultivars within the same farmer species (i.e., exchange within same farmer generation)</td>
<td>Present</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Present</td>
</tr>
<tr>
<td>Within a species of ant, cultivars are probably occasionally exchanged between colonies (e.g., after cultivar loss when cultivar-deprived colonies may acquire replacement cultivars from neighboring colonies).</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cultivar exchange between farmer species</td>
<td>Present</td>
<td>Present</td>
<td>Rare</td>
<td>Not applicable</td>
</tr>
<tr>
<td>Horizontal cultivar exchange occurs between sympatric ant species if they are specialized on the same cultivar lineage.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Genetic exchange between domesticated and wild populations (e.g., genetic interbreeding between</td>
<td>Variable</td>
<td>Absent</td>
<td>Absent</td>
<td>Absent</td>
</tr>
<tr>
<td>Primitive attine ants probably import novel cultivar genotypes</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wild populations are unknown for termite cultivars.</td>
<td></td>
<td>Wild populations are unknown for ambrosia beetle cultivars.</td>
<td></td>
<td>Some traditional cultivation regimes (e.g., potato farming in the</td>
</tr>
</tbody>
</table>
domesticated and wild populations; or import of new domesticates from wild populations) regularly from wild populations; cultivars of the derived higher attine ants do not appear to have free-living populations.

<table>
<thead>
<tr>
<th>Crop monoculture</th>
<th>Present</th>
<th>All available data indicate that only a single cultivar is grown within a nest.</th>
<th>Present</th>
<th>All available data indicate that only a single cultivar is grown within a nest.</th>
<th>Present</th>
<th>Most beetle species appear to be associated with a single primary cultivar and one or more secondary cultivars; it is unclear whether secondary cultivars are truly cultivars, weeds, or switch between these roles.</th>
<th>Variable</th>
<th>Monoculture predominates agriculture in many but not all regions.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crop sexuality</td>
<td>Variable, but sexual recombination is rare</td>
<td>Predominant clonal cultivar propagation within nests and between nests is punctuated occasionally by genetic exchanges, outbreeding, or other events generating recombinants.</td>
<td>Variable, but most cultivars are propagated sexually Cultivar sexuality is typical for those termite lineages that acquire cultivars from the environment every generation, but cultivar sexuality appears absent in the two agricultural systems with vertical cultivar transmission.</td>
<td>Variable, but sexual recombination is probably rare Asexuality is typical for both primary and secondary cultivars; some secondary cultivars are sexual.</td>
<td>Variable, but most cultivars are propagated sexually The great majority of human cultivars are sexually propagated, but some cultivars are largely clonally (e.g., potato) or strictly clonally propagated (e.g., banana).</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
as a vector of that fungus, then begins to derive nutrition from it, and finally becomes a fungus cultivator. In a third possibility, an insect-fungus association evolves because the insects originally use fungi as a source of antibiotics, as for example in the lower termite *Reticulotermes speratus* that derives antibiotic protection from fungal sclerotia mixed into egg piles (Matsuura et al. 2000). Lastly, insect-associated fungi may have undergone even more complicated evolutionary histories, originating from the exploitation by one insect lineage (e.g., the ancestor of attine ants) of a preexisting insect-fungus association (the fungi ancestrally associating with beetles) when it encounters these insect-adapted fungi in a shared nest environment (e.g., decaying wood; Sanchez-Peña 2005). This latter hypothesis, however, is not supported by the phylogenetic relationships between beetle and ant fungi and is inconsistent with the estimated dates of origin of these insect-fungal associations (i.e., attine agriculture probably arose well before ambrosia beetle agriculture; see Table 2).

For attine ants, it is unclear whether agriculture arose from a state of ancestral fungivory, antibiotic acquisition, or fungal vectoring (Mueller et al. 2001). Termite agriculture most likely originated via the consumption-first route, because many nonfarming termite species are attracted to and feed on fungus-infested wood, which suggests that the nonfarming ancestors of the farming termites may have fed on fungi as well (Batra & Batra 1979, Rouland-Lefevre 2000). The nonfarming ancestors of the fungus-growing beetles appear to have associated with fungi even before the origin of fungiculture, because many of the more primitive nongardening scolytines act as fungal vectors without apparent dependence on their fungal associates (Harrington 2005, Malloch & Blackwell 1993, Six 2003, Six & Klepzig 2004). This suggests non-nutritional dependencies on fungi that predate the origins of fungiculture in the various ambrosia beetle lineages (Six 2003). However, many nonambrosial scolytines carry fungi in mycangia and feed as larvae on ungardened mycelium that colonizes host plants and feed as new adults on spore layers lining pupal chambers (Ayres et al. 2000, Barras 1973, Six & Klepzig 2004, Six & Paine 1998; also A. Adams & D.L. Six, unpublished data), suggesting a stage of nutritional dependency predating the origin of fungiculture. Thus, some of the seven agricultural origins in beetles appear to have followed the transmission-first route, whereas others followed the consumption-first route.

Insect agriculture is restricted to the cultivation of fungi rather than plants, which predominate in human agriculture. Although it is true that some insects are specialized on host plants that they protect from other herbivores (e.g., *Pseudomyrmex* ants protect acacia trees in exchange for shelter and nutritional benefits; Janzen 1966, Hölldobler & Wilson 1990), none of these insect-plant mutualisms possesses all four of the components of agriculture listed above. One could therefore ask what factors have predisposed insects to evolve fungal rather than plant agriculture. Indeed, there are several advantages of fungal agriculture over plant agriculture, and several characteristics of plants may even preclude their easy cultivation. First, unlike fungi, plants typically have stringent light and space requirements, excluding them from cultivation in the subterranean or otherwise enclosed
nests of insects. Such nesting habits may facilitate fungiculture by shielding fungal crops from unwanted consumers (i.e., other fungivores) and wind-dispersed diseases. Furthermore, unlike plants, which usually require regular pollination for long-term cultivation, fungi can be maintained indefinitely in a nonsexual mycelial state, yielding a more consistent food source. Thus, although seeds and plant material can be readily harvested, fungi are likely more cultivatable, explaining the predominance of fungal rather than plant agriculture among insects.

Ant, termite, and most beetle agriculturists are social. All ants and termites are eusocial (characterized by reproductive division of labor, cooperative brood care, and overlap of generations; Hölldobler & Wilson 1990). Only one ambrosia beetle (*Austroplatypus incompertus*) is known to be eusocial (Kent & Simpson 1992); the remainder are subsocial, in which a single female cares for her brood, or communal, in which several reproductive females cooperate in brood care and gardening (Kirkendall et al. 1997). Sociality may have facilitated the evolution of agriculture because of the inherent advantage to agriculture of division of labor, which enables the partitioning of agricultural tasks and augments agricultural efficiency (Hölldobler & Wilson 1990, Hart et al. 2002). In ant and termite farmers, for example, agricultural tasks are partitioned in a conveyor-belt-like series between different worker castes, each specialized on one main task: foraging; processing and cleaning of substrate before incorporation into the garden; planting of mycelium onto new substrate; monitoring and weeding of the garden; or disposal of diseased or senescent garden (Bot et al. 2001a, Hart et al. 2002, Traniello & Leuthold 2000). Task partitioning has so far not been investigated in the ambrosia beetles because of the logistical difficulties of studying beetle behavior in their concealed tunnels. Task partitioning likely facilitates great efficiency in defense against nest and garden robbers (Adams et al. 2000a,b; LaPolla et al. 2002), in monitoring gardens for diseases, and in modulating optimal environmental conditions for crop growth.

4. AGRICULTURAL EVOLUTION AND ECOLOGY

A series of convergent and divergent features of agricultural evolution emerge from a comparative analysis of ant, termite, and beetle fungiculture (summarized in Tables 1–3).

4.1. Cultivar Transmission Between Farmer Generations

In attine ant and xyleborine beetle agriculture, fungal cultivars are transmitted vertically by trophophoresy from parent to offspring generations (Fernández-Marín et al. 2004, Francke-Grosmann 1967, Haanstad & Norris 1985, Huber 1905). Female reproductive ants and beetles acquire inocula from their natal gardens, carry these inocula with them in specialized pockets during dispersal flights early in life, and use these inocula as starter cultures for their new gardens. Trophophoretic
### TABLE 3  Disease ecology and evolution in ant, termite, beetle, and human agriculture

<table>
<thead>
<tr>
<th>Agriculture in:</th>
<th>Attine ants</th>
<th>Macrotermite termites</th>
<th>Xyloborine ambrosia beetles</th>
<th>Humans</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Disease ecology and evolution</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. Disease and pest types</td>
<td>Crops are attacked by fungal and bacterial pests, including specialized (Escovopsis) and unspecialized (Trichoderma) fungal pathogens. Microbial “weeds” (e.g., Xylaria fungi) and fungivorous arthropods (e.g., mites) can also inhabit gardens.</td>
<td>Specialized crop parasites have not been documented, but some species of Xylaria are specialized as weeds of fungus gardens. Common soil fungi are present in gardens as well.</td>
<td>Most ambrosial gardens consist of a complex of mycelial fungi, yeasts, and bacteria. At least some of these are likely to cause disease or act as pests.</td>
<td>Crops are infected by fungal, bacterial, and viral diseases and are attacked by invertebrate and vertebrate herbivores. Weeds are an additional problem in human agriculture.</td>
</tr>
<tr>
<td>2. Disease prevalence</td>
<td>Infection of gardens by the crop parasite Escovopsis is frequent and sometimes devastating.</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Ubiquitous and diverse diseases cause immense loss in crop harvests.</td>
</tr>
<tr>
<td>3. Defense mechanisms by farmers against crop disease</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3a. Protection from disease</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-Sequestration, sheltering</td>
<td>Present</td>
<td>Present</td>
<td>Present</td>
<td>Absent$^{a}$</td>
</tr>
<tr>
<td>-Substrate sterilization or cleaning</td>
<td>Present</td>
<td>Present</td>
<td>Unknown</td>
<td>Uncommon</td>
</tr>
<tr>
<td>-Guarding or protecting against disease vectors</td>
<td>Present</td>
<td>Present</td>
<td>Present</td>
<td>Uncommon</td>
</tr>
<tr>
<td>-Partitioning of disease-removal tasks to minimize contact between contaminated workers and healthy crop</td>
<td>Present An uncontaminated worker-caste specializes on gardening; a contaminated worker-caste specializes on disease removal.</td>
<td>Present Foraging and gardening are performed by different worker castes in many species.</td>
<td>Unknown</td>
<td>Uncommon</td>
</tr>
<tr>
<td>3b. Eradication of disease</td>
<td>Present</td>
<td>Present</td>
<td>Present</td>
<td>Very rare</td>
</tr>
<tr>
<td>---------------------------</td>
<td>---------</td>
<td>---------</td>
<td>---------</td>
<td>-----------</td>
</tr>
<tr>
<td>-Continuous, intensive monitoring of all crop</td>
<td>Present</td>
<td>Present</td>
<td>Present</td>
<td>Present</td>
</tr>
<tr>
<td>-Physical weeding</td>
<td>Ant gardeners continuously “weed” and “groom” their gardens, excising infected garden fragments.</td>
<td>Termite gardeners groom gardens to eliminate some alien microbes (e.g., Xylichromites stromata).</td>
<td>Female beetles continually tend crop gardens but whether directed weeding is part of this activity is unknown.</td>
<td>Physical weeding is more integral in primitive agricultural systems, but less so in modern monoculture systems.</td>
</tr>
<tr>
<td>-Application of chemical herbicides</td>
<td>Present</td>
<td>Present</td>
<td>Unknown</td>
<td>Present</td>
</tr>
<tr>
<td>Metapleural and mandibular gland secretions have general antibiotic properties; antibiotics derived from actinomycete bacteria have specific antibiotic effects against Escovopsis parasites.</td>
<td>Salivary gland secretions have antibiotic properties; defensive secretions of soldiers have antibiotic properties, but their sanitary function in the garden is unknown.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-Routine use of disease-suppressant microbes</td>
<td>Present</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Absent</td>
</tr>
</tbody>
</table>

4. Defense mechanisms of crop against crop diseases

| 4a. Induced resistance of crop | Unknown | Unknown | Unknown | Present |
| 4b. Constitutive resistance of crop | Present | Unknown | Unknown | Present |

| aGreenhouses provide a sequestered environment for some human crops, but greenhouse farming is costly and contributes only a small fraction (about 0.02%, Paulitz & Bélanger 2001) to the total productivity of human agriculture; the bulk of human agriculture operates in open landscapes that expose crops to environmental stresses (fluctuations in moisture and temperature; wind-borne pathogens; migratory herbivores; etc.).

bSome recently developed uses of disease-suppressant microbes in agricultural pest control are discussed in the text; application of disease-suppressant microbes has yet to become an integral part of mainstream human agriculture (but see Morrissey et al. 2004).
vertical transmission also occurs in two macrotermitine groups, except that in one of these two groups the fungal cultivar is transmitted via the king (the single species *Macrotermes bellicosus*), whereas in the other group (the genus *Microtermes*) the fungus is transmitted via the queen. In the few cases where fungal transmission has been studied in the remaining macrotermitines (Johnson 1981, Johnson et al. 1981, Korb & Aanen 2003, Sieber 1983), these termites rely on horizontal acquisition of fungal crops from the environment in each generation.

4.2. Higher-Level Specialization (Clade-Clade Congruence) Between Farmers and Crops

Vertical transmission of cultivars leads to the expectation of clade-clade correspondences and topological congruence between the phylogenies of insect farmers and those of their cultivars. Indeed, in all insect farming systems, major groups of farmers (large clades or paraphyletic grades, e.g., the lower attine ants) strictly specialize on major groups of corresponding fungal cultivars (Figure 2). The expected farmer-cultivar congruence therefore does occur at higher (i.e., broad) phylogenetic levels, possibly because of ancient evolutionary codependencies (e.g., physiological/nutritional requirements of the farmers, cultivation requirements of the fungi, etc.) that strictly preclude switches by farmers to cultivars outside of their specialized major cultivar groups.

Phylogenetic patterns (Figure 2) indicate, however, that within these strictly constrained major cultivar groups, insect-farmer species occasionally switch between fungal species or strains. This combination of lower-level, within-group switching and higher-level major-group specialization in insect farmers would be analogous, in humans, to defined clades of specialized wheat-farmers, rice-farmers, potato-farmers, bean-farmers, etc., each of which is able to switch between varieties within their area of specialization (e.g., between varieties of wheat and to closely related species such as barley), but which cannot switch across major groups (e.g., from wheat to beans). Among insect farmers, switches to novel major cultivar groups have been exceedingly rare evolutionary events (Villesen et al. 2004).

4.3. Lower-Level Specialization on Cultivars

Though low-level switching between cultivar species and strains within major cultivar groups occurs occasionally over evolutionary time, over ecological spans of time most insect farmer species associate with only a very narrow subgroup of cultivars (species or strains). For example, every attine ant species surveyed to date cultivates only a phylogenetically narrow set of cultivars (e.g., a single species of fungus), implicating species specificity between ants and cultivars at very recent levels of evolutionary diversification (Bot et al. 2001b, Green et al. 2002, Schultz et al. 2002). In ambrosia beetles, like ants, only one primary cultivar is associated with a particular beetle species within a particular geographic region (Gebhardt et al. 2004, Batra 1967). However, although most beetles are associated with a species-specific, primary fungus across their entire geographic
ranges, some beetle species associate with different primary cultivars in different geographic regions (Baker 1963, von Arx & Hennebert 1965). Among macrotermitine species, there exists considerable variation in cultivar specialization: Some species are limited to a single, unique cultivar, whereas other species cultivate a great diversity of fungal cultivars, which they sometimes share with other, usually closely related, macrotermitine species (Katoh et al. 2002; D.K. Aanen & P. Eggleton, submitted). The factors underlying variation in termite specialization are unknown, but different cultivars may serve different primary functions, providing specific, termite-adapted enzymes in some cases (leading to termite-cultivar specialization), while providing generalized food in other cases (permitting exchange between termite species; D.K. Aanen, V.I. Ros, H.H. de Fine Licht, C. Roulant-LeFévre, J. Mitchel, et al., in review).

4.4. Cultivar Sharing and Exchange Between Farmer Species

Even though each attine ant species is specialized on a single cultivar species, a given cultivar species may be cultivated by several sympatric species of ants, and these sympatric ant species may not necessarily be closely related to one another (e.g., they may represent different ant genera) (Bot et al. 2001b, Green et al. 2002, Villesen et al. 2004). Cultivar transfer between ant species may occur via direct or indirect avenues. Direct avenues may include raiding of neighboring colonies (Adams et al. 2000a, Rissing et al. 1989) or, in polygynous species, cofounding of colonies by multiple queens that exchange cultivars or recombine them in the cofounded garden. Indirect avenues may include cultivar escapes from gardens, followed by a free-living (feral) existence and subsequent reincorporation into a symbiosis when a different attine colony imports the free-living strain into its nest (Mueller et al. 1998).

For ambrosia beetles, the available phylogenetic evidence points to cultivar sharing between different sympatric beetle species (Farrell et al. 2001, Gebhardt et al. 2004) but few investigations have addressed this question. Distantly related ambrosia beetle species are sometimes associated with the same cultivar (Gebhardt et al. 2004), implicating fungal exchange, either direct or indirect, as explained above for the ants. Cultivar exchange between and within beetle species may occur when different female beetles colonize the same tree and the fungal associates cross-contaminate adjacent galleries.

In contrast to attine ants and ambrosia beetles that all transmit their cultivars vertically between generations, most macrotermitine species acquire their fungi horizontally each generation. This implies that new termite-cultivar combinations arise each generation, which should facilitate cultivar exchange between species, as well as between lineages of the same species. Cultivar surveys of sympatric macrotermitine communities indeed indicate that cultivars are generally shared between closely related species via interspecific cultivar exchanges (Aanen et al. 2002; but see the exceptions mentioned above in Section 4.3). Intra-specific cultivar exchanges have so far not been investigated in macrotermitines.
4.5. Propagation of Sexual Versus Asexual Cultivars and Links to Free-Living Cultivar Populations

All vertically transmitted insect cultivars, including the cultivars of attine ants, ambrosia beetles, and termites in the genus *Microtermes* and the species *Macrotermes bellicosus*, seem to be asexually propagated by their insect farmers across multiple farmer generations. In contrast, the horizontally transmitted termite cultivars (propagated by all other macrotermitine genera) undergo regular meiosis and sexual recombination (see above).

Evidence for cultivar asexuality in attine ants comes from DNA fingerprinting studies that indicate that all gardens of a single leafcutter colony contain a single cultivar clone (monoculture) (Kweskin 2003, Poulsen & Boomsma 2005; J. Scott & U.G. Mueller, unpublished data); that identical cultivar clones occur in different colonies of the same geographically widespread attine ant species (Bot et al. 2001b, Green et al. 2002, Mueller et al. 1996); and that different sympatric ant species occasionally share genetically identical cultivar clones (see above; Bot et al. 2001b, Green et al. 2002, Mueller et al. 1998). Contrary to previous suggestions (Chapela et al. 1994), however, attine cultivar clones are not ancient. Although attine cultivars are clonally propagated across many ant generations (Mueller 2002), this clonality is punctuated by occasional recombination events, involving either sexual (meiosis, mating) or parasexual (e.g., mitotic recombination, exchange of haploid nuclei) processes. Evidence for occasional recombination includes: (a) fruiting structures (mushrooms), which are known for cultivars of nearly all genera of attine ants, contradicting the expectation of loss of fruiting ability under strict clonality spanning millions of years (Mueller 2002); (b) rates of allele sequence divergence in attine cultivars that are similar to those of closely related, sexually reproducing fungi (Mueller et al. 1998; S.A. Rehner, unpublished data); and (c) cultivars of the lower attine ants that have close genetic links to free-living fungal populations (Green et al. 2002, Mueller et al. 1998), suggesting that these fungi are capable of moving in and out of the symbiosis, that cultivar lineages may regularly interbreed with wild lineages, or both. Taken together, the genetic and natural-history information suggest predominantly asexual cultivar propagation within ant nests and across many ant generations, punctuated by occasional genetic recombination events.

As in attine ants, a single cultivar monoculture is grown in a single termite colony (Aanen et al. 2002, Katoh et al. 2002). Within termite nests, the *Termitomyces* cultivar is propagated asexually by inoculating fresh garden substrate with asexual spores (Leuthold et al. 1989), and probably also by transplanting mycelium from older to younger gardens. Although *Termitomyces* species have no known free-living populations existing entirely independent of the termite farmers, they have nonetheless retained the ancestral (presymbiotic) condition of regular sexual reproduction, and most *Termitomyces* cultivars are spread from one termite nest to another horizontally via sexual spores produced by fruiting bodies (mushrooms) growing on the external surfaces of mature nests. The *Termitomyces* cultivar of the
termite *Macrotermes natalensis*, for example, has an outcrossing mating system (De Fine Licht et al. 2005). Asexual cultivar propagation spanning several termite generations only occurs in those species with vertical uniparental propagation. Phylogenetic patterns implicate occasional horizontal cultivar exchange between nests of the same and different termite species (Aanen et al. 2002), but whether such horizontal exchange is associated with cultivar sexual reproduction remains unknown.

In xyleborine beetles, the primary fungi are strictly asexual (Jones & Blackwell 1998, Rollins et al. 2001), whereas the less specific, auxiliary fungi are often sexual (Francke-Grosman 1967). A preponderance of asexual reproduction in fungal cultivars also occurs in the primary fungi of all other non-xyleborine ambrosia beetles, whereas, again, the more incidental fungi are often sexual (Six 2003, Six & Paine 1999), suggesting that this may have been the ancestral condition at the origin of the xyleborine beetle-fungus symbiosis.

### 4.6. Coevolutionary Modifications

Farmer-cultivar specialization enhances the potential for coadaptation, in which evolutionary modification in one of the partners causes a reciprocal coevolutionary modification in the other partner (Futuyma & Slatkin 1983). It is relatively easy to identify evolutionary modifications in the farmer species, such as specialized morphological structures for the trophophoretic transport of cultivars by females during the dispersal flight (e.g., mycangia in the beetles, infrabuccal pocket in the ants), modifications of mandibles and guts of beetle and ant larvae for fungus-feeding (Browne 1961, Schultz & Meier 1995), or the suite of behavioral, glandular, or physiological modifications that form the basis of insect farming. Examples of evolutionary modifications in the cultivars have been more difficult to identify, however, because the cultivated fungi are inherently more difficult to study.

The clearest examples of cultivar modifications are the hyphal-tip swellings (gongylidia) produced by the cultivars of the higher attines and the analogous nodules produced by macrotermitine cultivars. Both gongylidia and nodules are nutrient-rich structures designed for easy harvesting by the farmers, ingesting, and feeding to the larvae or nymphs. Nutrient-rich structures are not known for beetle gardens, although the ambrosia morphology of the beetle cultivars suggests evolutionary modification designed specifically for efficient consumption and digestion by the beetle larvae. Ambrosial growth consists of tightly packed conidiophores with copious spores and is only formed in the presence of the beetles (French & Roeper 1972). Ambrosia formation has not been reported from nonsymbiotic fungus species. Interestingly, two of the major genera of fungi associated with ambrosia beetles (*Ambrosiella, Raffaelea*) are each polyphyletic, and the multiple lineages within each genus have converged on the same ambrosial morphology (Blackwell & Jones 1997, Jones & Blackwell 1998), suggesting evolutionary convergence due to selection. Other likely coevolutionary modifications that have
yet to be investigated include predominantly asexual reproduction in the insect cultivars while under cultivation and the cultivars’ capacity to survive storage in the dispersal pockets of the beetles and ants, or the passage through the alimentary canal of the termites.

4.7. Symbiont Choice and “Artificial Selection” of Cultivars

From an evolutionary perspective, insect agriculture represents a case of cooperative interaction between farmer and cultivar lineages, each exploiting the other for its own reproductive purposes (Herre et al. 1999, Mueller 2002). Such cooperative interactions are frequently unstable and can erode over evolutionary time, for example, when mutant overexploiters arise (so-called cheater cultivars) and invade a mutualism. A series of additional farmer-cultivar conflicts are predicted that could destabilize the mutualism (Mueller 2002, Aanen & Boomsma 2005, Schultz et al. 2005), but at least two evolutionary mechanisms preserve the cooperative nature of the farmer-cultivar association: First, partner feedback, inherent in vertical cultivar transmission, is an automatic feedback mechanism in which an uncooperative partner reduces the other partner’s fitness to the extent that it reduces its own fitness as well; and, second, partner (symbiont) choice in which farmers favor associations with productive cultivars and discriminate against inferior cultivars in specific choice situations (e.g., a choice between cultivar strains that may coexist in a garden or in proximate gardens, exercised either by workers during the planting of new gardens or by reproductives when choosing a cultivar strain for dispersal). In cases where the evolutionary rates differ between two cooperating partners, partner choice is a particularly important mechanism (Sachs et al. 2004). The slower-evolving partner (e.g., the insect farmer) is expected to exert the choice between variants of the faster evolving partner (e.g., the fungal cultivar), and thus the slower-evolving farmer imposes selection favoring beneficial symbiont variants (e.g., productive cultivars) and prevents the spread of nonbeneficial cultivar mutants (e.g., degenerate or suboptimal cultivars; Mueller 2002, Sachs et al. 2004). Symbiont choice has yet to be investigated for termite and beetle farmers, but ant farmers are able to discern surprisingly fine genotypic differences between cultivars (Mueller et al. 2004; also N.K. Advani & U.G. Mueller, submitted), suggesting that cultivar diversity in ant gardens, arising, for example, through mutation in a garden or through the import of novel strains, may evolve under an analog of “artificial selection.”

5. ROLE OF DISEASE IN INSECT AGRICULTURE

“Weedy” fungi frequently invade the gardens of ants, termites, and beetles, and may coexist at low or manageable levels along with the crop. If the gardening insects are removed or if they abandon their nests, the garden is quickly overrun by these weeds (Batra & Batra 1979, Norris 1979). One such group of weeds, wood-degrading
fungi in the endophytic genus *Xylaria*, is found in most gardens of fungus-growing ants and termites, probably because it is introduced with garden substrate (Fisher et al. 1995; V.I. Ros, A.J. Debets, T. Læssøe, D.K. Aanen, submitted; N.M. Gerardo & U.G. Mueller, personal observation). Though weeds like *Xylaria* do not directly attack the cultivar, they compete with it for nutrients and thus decrease crop yield (V.I. Ros, A.J. Debets, T. Læssøe, D.K. Aanen, submitted). *Escovopsis* species, ascomycete fungi found in colonies of fungus-growing ants, are specialized parasites that subsist directly on the cultivars and reduce the nutrients available to the ants (Currie 2001a, Currie et al. 2003b). Weed fungi and bacteria are also known in termite and beetle agriculture, but have yet to be studied in detail (Six 2003; V.I. Ros, A.J. Debets, T. Læssøe, D.K. Aanen, submitted; D.K. Aanen, unpublished data).

*Escovopsis* infections reduce garden productivity, which in turn reduces ant colony growth and the likelihood of colony survival (Currie 2001a, 2001b, Currie et al. 1999a). *Escovopsis* is geographically widespread and taxonomically diverse. The parasite has been isolated from colonies of every attine genus throughout their geographic ranges, and particular *Escovopsis* lineages are specialized to parasite particular cultivar lineages. This high degree of host specificity suggests a long history of host-parasite coevolution in *Escovopsis*, in which the cultivars, the ants, and their mutualistic bacteria have likely coadapted to defend against *Escovopsis* attack and in which each *Escovopsis* species has become narrowly specialized to overcome the defenses of some hosts but not others (Gerardo et al. 2004).

6. DISEASE AND MICROBIAL MANAGEMENT STRATEGIES

In response to the persistent selection pressure imposed by weeds and pathogens, fungus-farming insects have evolved an arsenal of strategies for preventing and suppressing infection (Table 3).

6.1. Sequestration of Gardens

All insect agriculturists sequester and separate their gardens from the surrounding environment, e.g., by growing their gardens in underground chambers or galleries in wood, or by covering them with a protective mycelial veil in the case of some *Apterostigma* ants (Villesen et al. 2004). Although sequestered nests are the ancestral condition in fungus-growing ants, termites, and beetles, and although sequestered nests may serve other agricultural functions such as the regulation of temperature and humidity, sequestration no doubt also buffers the garden against fungivores, wind-borne pathogens, and arthropod vectors of diseases (e.g., mites, collembolans).
6.2. Maintenance of Cultivar Genetic Variability

Although attine ants clonally propagate their cultivars across generations over short evolutionary time spans, no attine cultivar has been found to be an ancient clone. Instead, the evidence indicates that the lower attines occasionally acquire new cultivars from wild (free-living), sexually reproducing fungal populations and that both lower and higher attines occasionally acquire new cultivars from the nests of other attines. The cultivars of higher attines, which are not known to have free-living populations (Mueller 2002), nonetheless retain the ability to fruit (Mueller 2002) and demonstrate patterns of DNA-sequence diversity that suggest occasional genetic recombination through self-mating (S.A. Rehner, personal communication) or through true intercrossing between different cultivar strains (M. Bacci, personal communication). Thus, whereas the crop employed by any attine ant colony at any given time is a clonally propagated monoculture, the genetic variability and resilience necessary for long-term disease management resides in the fungal population external to the nest. As already noted, sexual reproduction is the norm in the cultivars of those termites that reacquire their cultivars horizontally each generation, and sexual reproduction may also occur in the fungi of those termites that transmit their cultivars vertically across generations. Whether the primary cultivars of the ambrosia beetles occasionally reproduce sexually remains unknown. At least for the termites and ants, then, and possibly for the beetles as well, access to a population-level reservoir of cultivar genetic variability is a consistent feature of insect agriculture that may provide alternative crops for dealing with disease.

6.3. Intensive Monitoring of Gardens, Weeding, and Herbicide Application

All insect agriculturists constantly inspect their gardens, and no part of the garden is left untended for periods of time sufficient to allow the establishment and spread of diseases and fungivores (Batra & Batra 1979, Currie & Stuart 2001). Insect agriculturists, particularly the ants and termites, are able to invest in such intensive monitoring because their societies possess a nonreproducing worker caste, a large portion of which is dedicated to garden care. In the beetles, the relatively small size of the garden allows for intensive monitoring by a single female or by a small family of females. Intensive monitoring ensures that diseases are discovered and eradicated in the early stages of infection before they are able to spread and cause significant crop loss. Early detection is an effective defense against novel disease mutants that might evolve greater virulence if left untreated, because in the early stages of infection these strains can more readily be controlled with standard treatments.

Garden treatment in attine ants includes the use of secretions from their metapleural and mandibular glands to clean substrate as it is brought into the nest, probably removing some or most weeds and pathogens from the surface of the substrate before it is added to the fungus garden (Maschwitz et al. 1970, Ortius-Lechner et al. 2000). Although antibiotic-producing glands have not been studied in fungus-growing termites, some secretions of nonfungus-growing termites have
antibiotic properties (Rosengaus et al. 1998, 2004). Antimicrobial glands in ambrosia beetles remain unknown and unstudied. In addition to glandular secretions, attine ants have another antimicrobial defense. Some or all of their integuments are covered with actinomycete bacteria. These bacteria are known to inhibit Escovopsis growth (Currie et al. 1999b), and experimental reduction of actinomycetes in colonies increases Escovopsis infection (Currie et al. 2003a). Garden bacteria in the genus Burkholderia (Santos et al. 2004) also provide antibiotics that provide protection against the garden parasite Escovopsis and against entomopathogenic diseases of the ants themselves. Termite gardens contain actinomycetes and other bacteria (Batra & Batra 1979), and beetle gardens contain a great diversity of bacterial secondary symbionts; however, the exact roles of these bacterial associates remain unknown.

6.4. Microbial Buffering

Beyond the known antibiotic-producing, disease-suppressing bacteria in attine colonies (Currie 2001a, Currie et al. 1999b, Santos et al. 2004), other secondary bacteria and fungi occur in insect gardens (Carreiro et al. 1997, Craven et al. 1970; C. Wang & U.G. Mueller, unpublished data), but their roles remain largely unknown. Although some of these secondary microbes may be neutral or detrimental to garden health and productivity, others may provide disease-modulating effects through competitive exclusion, antibiotic suppression of disease-causing microbes, resistance induction, or other mechanisms of microbial interaction. Competitive exclusion, disease suppression, and resistance induction have been demonstrated in both experimental and natural microbial systems (Hood 2003, Paulitz & Bélanger 2001, Wille et al. 2001). Some secondary microbes may even facultatively switch between beneficial and detrimental roles, depending on garden growth conditions, seasonal factors, or interactions with the insects or other microbes. For example, although the detrimental effects of Escovopsis are obvious in natural garden outbreaks and in interactions with cultivars in vitro (Currie 2001a,b; Currie et al. 1999a; Gerardo et al. 2004), it remains an untested possibility that Escovopsis may provide beneficial effects when present at low levels in the garden matrix.

Secondary microbes in termite gardens remain uninvestigated, but the possible significance of a secondary microbial flora in beetle gardens has been recognized for some time (Norris 1965). Norris (1965) suggested that it is the microbial complex as a whole (filamentous fungi, yeasts, and bacteria), rather than the dominant ambrosia fungus per se, that allows the beetles to exploit nutrient-poor substrates such as wood. Norris did not speculate on any additional roles for the secondary microbes, such as suppressing diseases, but such auxiliary roles deserve further study.

6.5. Management of Crop-Associated Microbial Consortia

If, as recent evidence indicates, secondary microbes serve ancillary functions in gardens by buffering against disease organisms or by producing antibiotics,
enzymes, and metabolites, it is possible that insect farmers have evolved the
capacity to manage these microbial consortia. Such microbial management strategies
by insect farmers could include the following three methods.

6.5.1. STERILIZATION OF SUBSTRATE BEFORE INCORPORATION INTO THE GARDEN
(ANTS, TERMITES) OR USE OF ESSENTIALLY STERILE SUBSTRATE FROM THE OUTSET
(BEETLES) In termites, the passage of substrate through the gut before incorpo-
rating into the garden probably eliminates many unwanted microbes and may
increase the abundance of desired microbes (Figure 1). In attine ants, the con-
siderable effort spent cleaning substrate surfaces appears to partially sterilize
the substrate (Weber 1972). In the case of the ambrosia beetles, gardening occurs in
a closed system because the beetles do not need to leave the nest to forage and
because galleries are excavated in what is essentially a sterile medium, the
sapwood or heartwood of living or recently killed trees, which are generally free of
endophytic fungi and other microbes. This closed system greatly reduces the po-
tential for accidental introduction of unwanted microbes and likely facilitates the
management of desired microbes in beetle gardens.

6.5.2. SPATIALLY STRUCTURED GARDEN MATRIX Structuring of gardens allows in-
sect farmers to assess properties of particular, localized crop-microbe consortia.
Any unwanted mutant genotypes, arising locally under particular microbe-microbe
competitions, thus can be identified indirectly through the detection of their detri-
mental effects on the properties of the subgarden, and that subgarden piece can
then be excised. Conversely, novel microbial mutants with beneficial effects can
be identified indirectly by their beneficial effects on the subgarden, and preferen-
tially subcultured and propagated across the rest of the garden (Figure 1). Such
“symbiont-community choice” is possible only because of the fixed garden matrix,
enabling farmers to assess properties of local consortia.

6.5.3. CONTINUOUS CO-OCCURRENCE OF GARDENS OF ALL AGES IN CLOSE PROXI-
MITY, RANGING FROM “UNPLANTED” TO MATURE GARDENS Coexistence of gar-
dens at different developmental stages is inherent in the vertical structuring of attine
and termite gardens into younger subgardens at the top and older subgardens at the
bottom. A range of differently-aged gardens allows farmers to efficiently practice
one-way, selective transfer of only beneficial crop-microbe consortia from mature
to younger gardens. Age-structuring also delays the spread of mutant microbes
from older garden material to younger, more sterile gardens and thus prevents
deterioration of the symbiont-community.

6.6. Multipartner Coevolution and Coevolving
Antibiotic Defenses

One hypothetical advantage of secondary mutualistic microbes is that, unlike the
insect farmers, microbes can potentially evolve at the same rate as the coevolving
garden pests, enabling mutualistic insect-microbe systems to respond rapidly to the emergence of novel disease genotypes (Currie 2001a, Mueller & Gerardo 2002). Although such rapid microbial antibiotic defenses would obviously confer clear advantages, their evolutionary maintenance remains unclear. One possibility is that any single farmer society may have access to a diverse array of microbes from which it can select particular, desired types as needed. This scenario raises the question of how the insect farmers could maintain such a diverse array in their colonies in the face of both competition between microbes and the regular bottlenecking of the entire microbial “library” that presumably occurs at the founding of every new insect colony. Alternatively, the associated secondary microbes may be inherently fast-mutating, so that novel beneficial genotypes can rapidly arise to muster an appropriate defensive response. This scenario raises the question of how the most beneficial genotypes are recognized by the insect farmers and chosen for selective “amplification” against particular pathogens. The lack of clear evolutionary mechanisms for maintaining functional associations with coevolving, mutualistic microbes is not trivial, and future research needs to assess not only the diversity of microbial genotypes within single farmer colonies, but also to identify the mechanisms underlying adaptive symbiont-choice selection of beneficial, novel microbial genotypes. Future research also needs to address whether the coevolution of several, mutualistically-aligned partners (i.e., a “multidefense alliance” of ants, cultivar, and auxiliary microbes), each mustering its own defense, provides for a more evolutionarily stable disease-management strategy compared to a strategy in which the insect farmers act alone in a coevolutionary arms race against particular pathogens.

7. PRINCIPLES OF INSECT AGRICULTURE: LESSONS FOR HUMAN AGRICULTURE?

Perhaps the most striking feature of insect agriculture is the long-term cultivation of clonal monocultures. Monoculture increases agricultural efficiency through an economy of scale (Wolfe 1985), and clonality preserves the desirable properties of the crop by eliminating sexual recombination, but these advantages come at two costs: (a) increased vulnerability to the rapid spread of disease mutants (Barrett 1981, Shipton 1977, Mitchell et al. 2002, Mundt 2002, Peacock et al. 2001, Piper et al. 1996, Wolfe 1985), and (b) decreased resistance to fast-evolving diseases due to decreased genetic variability in the crop (Barrett 1981, Gustafson et al. 2003, Hamilton et al. 1990, Jaenike 1978, Zhu et al. 2000). These economic trade-offs (i.e., monoculture/clonality efficiency versus disease vulnerability) apply to both human and insect farmers.

The insect farmers’ solution to the monoculture-disease problem appears to be not a single, “magic bullet” strategy (e.g., sole reliance on pesticides), but rather a combination of several strategies consisting of (a) crop sequestration, (b) intensive monitoring of crops for diseases, (c) access to a population-level reservoir of crop
genetic variability, and (d) management of disease-suppressant microbes associated with the crop (Table 3). Of these strategies, large-scale crop sequestration is the least feasible in human agriculture because human crops need exposure to sunlight and because greenhouse cultivation is costly (Paulitz & Bélanger 2001). Intensive (e.g., daily) monitoring of every single crop plant for diseases may be feasible for some crops (e.g., in greenhouse environments); however, hourly monitoring of the kind implemented in insect agriculture seems cost-prohibitive for human agriculture at large.

A more novel approach is to design human agricultural systems that more efficiently take advantage of the microbial consortia that are known to play beneficial roles in crop nutrient uptake and disease resistance (Morrissey et al. 2004, Paulitz & Bélanger 2001, Wardle et al. 2004). Microbes of the rhizosphere (e.g., nitrogen-fixing bacteria and mycorrhizal fungi) have long been managed as critical associates of certain crops and trees (Finlay 2004, Johansson et al. 2004). More recently, disease-suppressant bacteria have been discovered that live on the root exudates of crops and produce antibiotics that protect the crop against pathogens (Haas & Keel 2003, Mazzola 2004, Morrissey et al. 2004, Weller et al. 2002, Whipps 2001). Disease-suppressant effects on crop plants have also been documented for phyllosphere microbes (Lindow & Brandl 2003) and endophytic microbes (Narisawa et al. 2002, Sturz et al. 2000). Agricultural research on rhizosphere, phyllosphere, and endophyte microbes of human crops is a very new field, however, and many beneficial microbes remain to be discovered and put to use (e.g., inoculation of crops with phyllosphere microbes to deter herbivores or to suppress airborne diseases).

Two problems commonly encountered in human agricultural experiments with beneficial microbial consortia are, first, that the composition of microbial species is difficult to manage and stabilize (Garbeva et al. 2004, Mazzola 2004), and, second, that beneficial microbes can rapidly evolve into detrimental ones (Alves et al. 2003, Morrissey et al. 2002). The farming insects’ solution to these problems appears to consist of (a) selection on spatially limited microbial consortia (i.e., high-resolution, spatial separation of evolutionary processes, preventing the uncontrolled spread of microbes from inferior consortia); (b) propagation of crops with fast generational turnovers, thus minimizing the time for the evolution of any deleterious traits in the microbes; and (c) partial or complete sterilization of the substrate prior to planting, thus minimizing the influx of microbial contaminants into a largely closed agricultural system.

Perhaps it is from strategies such as these that humans have the most to learn from insect farmers, certainly if disease-suppressant microbes are ever to be managed in human agriculture (Morrissey et al. 2004). In developing these strategies, agriculturists would need to keep in mind that, during the domestication process, current human crops were not necessarily selected for capacities to interact with auxiliary microbes, i.e., the alleles in the wild ancestors optimally mediating such interactions may have been lost during the domestication process. Thus, a full evaluation of the potential uses of auxiliary microbes in human agriculture may
require the study of the microbial consortia associated with the wild populations from which human-domesticated crops were originally derived. Such domestication within the context of coevolving and codomesticated microbial consortia may well be the key element explaining the 50-million year old agricultural success of the insect farmers.

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EVOLUTION OF AGRICULTURE IN INSECTS

See legend on next page
**Figure 1** Comparison of agriculture in attine ants, xyleborine beetles, macrotermitine termites, and humans. The time series *(left to right)* highlight the roles of beneficial auxiliary microbes *(blue shading)* that suppress diseases *(black dots)* or aid in buffering against contaminant microbes *(orange shading)*. *(a)* **Ant agriculture.** Ants attempt to clean contaminant microbes from garden substrate (not shown) and remove garden diseases *(black dots)* through active weeding *(top ant)*. The ants *(bottom ant)* then plant a crop-microbe consortium *(crop plus beneficial auxiliary blue microbes)* onto the prepared substrate, spreading beneficial microbes through the garden matrix. *(b)* **Beetle agriculture.** Primary fungus *(crop)* lining the tunnel grows intermixed with secondary microbes *(blue shading)* and occasional contaminant microbes *(orange dots)*. No disease microbes *(black dots)* are indicated because they very rarely occur in young gardens near a tunnel head. The exact roles of the secondary microbes in beetle fungiculture are still unknown. *(c)* **Termite agriculture.** Hypothetical passage of a mixture of crop spores, auxiliary microbes, and substrate *(ingested plant material)* through the gut of a termite, followed by defecation of the substrate-crop-microbe consortium in fecal pellets that the termite adds to new garden. Other *(external)* modes of crop-microbe copropagation may exist in termite farmers, paralleling the planting of crop-microbe consortia in attine ants. Selective passage of microbes through the alimentary canals of attine ants and ambrosia beetles is unknown, but has never been investigated. No disease microbes *(black dots)* are indicated because no specialized pathogens have yet been identified in the fungus-growing termite system. *(d)* **Human agriculture** *(wheat).* A seed, fortuitously planted in soil enriched in antibiotic-secreting rhizosphere bacteria *(blue-shaded soil)*, grows into a vigorous, disease-resilient crop plant *(Weller et al. 2002).* However, crops are often planted in microbially suboptimal soil *(orange-shaded soil)*, leading to higher disease loads *(black dots)* on such plants. Traditional human planting schemes passage crops through a seed stage without copropagating disease-suppressant rhizosphere microbes. Illustrations by Barrett Klein.
Figure 2  Evolutionary histories of insect agriculture. (a–c) Comparison of the patterns of evolutionary diversification in the insect farmers (left cladograms) and their cultivated fungi (right cladograms). In the left cladograms, farmer lineages are black and nonfarmer relatives are gray, whereas in the right cladograms, cultivated fungal lineages (cultivars) are black and noncultivated feral fungal lineages are gray. Independent origins of agricultural behavior are indicated for each farmer clade in the left cladograms, and independently domesticated fungal lineages appear as separate cultivar lineages in the right cladograms. (d) Garden of the fungus-growing ant *Atta texana* (photo by Greg Dimijian). The workers are cleaning and shredding leaf cuttings before expanding new gardens through the addition of leaf material. (e) Garden of the fungus-growing termite *Macrotermes bellicosus* (photo by Karen Machielsen). The fungus is grown on fecal pellets that are stacked into lamellar walls of the fungus garden (comb). (f) Gallery of the ambrosia beetle *Trypodendron lineatum* (photo by Susanne Kühnholz) with ambrosia fungus (black) lining the main gallery and beetle brood developing in niches adjacent to the gallery. Galleries are constantly patrolled by adult beetles (not shown). Figure adapted from Mueller & Gerardo 2002.
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