Ergosterol Content of Fungi Associated with *Dendroctonus ponderosae* and *Dendroctonus rufipennis* (Coleoptera: Curculionidae, Scolytinae)

BARBARA J. BENTZ¹ AND DIANA L. SIX²

Ann. Entomol. Soc. Am. 99(2): 189–194 (2006)

ABSTRACT  Insects require sterols for normal growth, metamorphosis, and reproduction, yet they are unable to synthesize these organic compounds and are therefore dependent upon a dietary source. For phloephagous species, such as *Dendroctonus* bark beetles, whose food does not necessarily contain appropriate types or adequate quantities of sterols, fungal symbionts may provide an alternative source in the form of ergosterol. We determined and compared the relative amounts of ergosterol in the primary fungal associates of *Dendroctonus ponderosae* Hopkins and *Dendroctonus rufipennis* Kirby. Ergosterol content of host tree phloem naturally infested with larvae (and their fungal symbionts) of both species also was compared with ergosterol contents in uninfested phloem tissue. Mycelia of *Ophiostoma montium* (Rumford) von Arx and *Ophiostoma clavigerum* (Robinson-Jeffrey & Davidson) Harrington isolated from *D. ponderosae* mycangia, and *Leptographium abietinum* (Peck) Wingfield isolated from the exoskeleton of *D. rufipennis* contained relatively large quantities of ergosterol, although no significant differences in content were found among these fungal species. Phloem colonized by larvae of both species contained significantly more ergosterol than did uninfested host phloem tissue. Our results suggest that larval life stages of *D. ponderosae* and *D. rufipennis* may obtain vital nutrients not only from the host tree phloem but also from fungal symbionts, in the form of ergosterol, while mining larval galleries.

KEY WORDS  bark beetle, symbiosis, mycangial fungi, mycophagy

SEVERAL BARK BEETLE SPECIES within the genus *Dendroctonus* (Coleoptera: Curculionidae, Scolytinae) can exhibit dramatic population eruptions at periodic intervals, resulting in tree mortality across vast forested areas. Although the mechanisms are not always clear, both density-dependent and -independent processes can contribute to these population fluctuations, including interactions with other organisms (Bridges 1983). Many *Dendroctonus* are symbiotically associated with at least one filamentous fungal species that is mutually beneficial to the beetle host, whereas relationships with other fungal associates may be antagonistic or commensalistic (Six 2003b). The fungi benefit from association with the beetle host by consistent dissemination to an ephemeral and relatively rare resource (a freshly killed tree) upon which they are dependent for growth and reproduction. In some associations, growth and reproductive success of beetle brood is increased in the presence of fungi, often because of a nutritional benefit (Whitney 1971, Bridges 1983, Goldhammer et al. 1990, Coppedge et al. 1995, Six and Paine 1998, Ayres et al. 2000). Although mycophagy may not be obligatory for all bark beetle species, fungal feeding may provide vital nutrients not found in woody plant tissues or only present in inadequate concentrations (Harrington 2005). For example, most plant tissues have relatively low levels of nitrogen, and a fungal mutualist of *D. frontalis* Zimmermann was found to benefit larval growth by concentrating nitrogen in phloem tissue where feeding occurs (Ayres et al. 2000).

Insects also require sterols for normal growth, metamorphosis, and reproduction, yet are unable to synthesize these organic compounds and are therefore dependent upon a dietary source (Clayton 1964, Richmond and Thomas 1975, Svoboda et al. 1978). Sterols in woody plant tissues are typically present only in low concentrations (Kramer and Kozlowski 1960) or in forms not usable by insects (Clayton 1964). For insects that feed in woody tissues, fungi may provide an alternate source of sterols in the form of ergosterol (*24*β-methylcholesta-5,7,trans-22-trien-3β-ol) (Norris et al. 1969, Kok et al. 1970, Norris 1972, Maurer et al. 1992, Morales-Ramos et al. 2000). This major fungal sterol is produced by most fungi (Weete 1973, 1989) and is highly usable by many insects for the production of hormones and cell membranes (Clayton 1964). Although ubiquitous in fungi, ergosterol varies in its concentration in mycelia by species, age of the cul-

This paper was written and prepared by a U.S. Government employee on official time and therefore is in the public domain and not subject to copyright.

¹ Corresponding author, address: USDA Forest Service, Rocky Mountain Research Station, Logan, UT 84321 (e-mail: bbentz@fs.fed.us).

² Department of Ecosystem and Conservation Sciences, The University of Montana, Missoula, MT 59812.
ture, developmental stage, and growth conditions (Pasanen et al. 1999).

In several insect–fungus symbioses, the insect associate has been found to be dependent upon sterols provided by the fungal associate (Norris et al. 1969, Kok et al. 1970, Norris 1972, Maurer et al. 1992, Morales-Ramos et al. 2000, Nasir and Noda 2003). These include ambrosia beetles (Scolytinae) that feed solely on fungi they garden in galleries in the sapwood of trees (Norris 1972, Kok 1979, Mueller et al. 2005). Ambrosia beetles are dependent upon ergosterol produced by their fungal symbionts for successful oocyte pupation (Norris and Baker 1967, Norris et al. 1969, French and Roepert 1975, Kok 1979). This relationship holds for another scolytine, the coffee berry borer, *Hypothenemus hampei* (Ferrari). This beetle bores in, and feeds on, coffee berries but cannot molt or reproduce without ergosterol from its symbiotic fungus *Fusarium solani* Martius (Morales-Ramos et al. 2000).

Although almost all fungi produce ergosterol, the varying levels of ergosterol present in the mycelia of different species of fungi may account for the relative differences in benefits to host beetles exhibited by some fungal associates. For example, studies by Kok et al. (1970) and Kok and Norris (1973) found that the most beneficial species of fungi associated with *Xyleborus* ambrosia beetles were those that possessed the greatest concentrations of ergosterol.

*Dendroctonus rufipennis* Kirby and *Dendroctonus ponderosae* Hopkins, two economically important bark beetle species in western North American forests, are closely associated with fungi. *Leptographium abietinum* (Peck) Wingfield is commonly found in specialized pits on adult *D. rufipennis* pronota and elytra (Solheim 1995, Six and Bentz 2003). Although *Ceratocystis rufipenni* (Wingfield) Harrington & Solheim has been isolated from wood colonized by *D. rufipennis* in some populations (Reynolds 1992, Solheim 1995), Six and Bentz (2003) did not isolate this fungus from any *D. rufipennis* adults collected in Alaska, Utah, Colorado, and Minnesota, and they consider it to be a minor or incidental associate. Little research has been conducted on these fungi, and it is unclear whether they play a mutualistic or antagonistic role in *D. rufipennis* population dynamics.

*D. ponderosae* is closely associated with two fungi, *Ophiostoma montium* (Rumford) von Arx and *Ophiostoma clavigerum* (Robinson-Jeffrey & Davidson) Harrington. Both species can be found on the exoskeleton of adult *D. ponderosae* and in specialized structures of the maxillae termed sac mycangia that function in fungal transport (Six 2003a,b). Sac mycangia are relatively rare in bark beetles, and those that do possess them are also mycophagous, suggesting that mycangia evolved to maintain tight associations with specific fungi that provide sources of nutrition for the beetle host (Harrington 2005). Effects of the two fungi on their host beetle are not well understood. Only two studies have been conducted investigating potential nutritional effects of the fungi, and the results of these two studies seem to conflict (Six and Paine 1998; K. Bleiker and D.L.S., unpublished data). In one study, *D. ponderosae* were reared in logs colonized with parental beetles associated with either *O. clavigerum*, *O. montium*, or without fungi (Six and Paine 1998). Resulting brood that developed with *O. clavigerum* were more numerous and emerged significantly earlier than brood that developed with *O. montium*. Beetles introduced without fungi did not produce brood. These results indicate that both fungi support brood production but that *O. clavigerum* may be a superior associate in conferring nutritional benefit. However, in a recent study conducted in phloem sandwiches (K. Bleiker and D.L.S., unpublished data), beetles developing with *O. montium* consumed significantly less phloem than insects developing with *O. clavigerum* or no fungi, suggesting that *O. montium* provides superior nutritional benefits. Beetles developing with *O. clavigerum* or no fungus had to consume more phloem to meet their dietary requirements.

As has been found with other mycophagous scolytine beetles, we hypothesized that *D. ponderosae* and *D. rufipennis* obtain sterols, in particular ergosterol, from their close fungal associates. Because *D. rufipennis* does not have sac mycangia, as has been found for *D. ponderosae*, and thus may not have as highly an evolved association with its fungus or as strong a dependence on fungi for nutrients, we predicted that the fungal associate of *D. rufipennis* may provide less of a beneficial nutrient resource in the form of ergosterol compared with one or both of the fungi associated with *D. ponderosae*. Our objective was to determine and compare the relative amounts of ergosterol contained in the primary fungal associates of *D. ponderosae* and *D. rufipennis*. Furthermore, the ergosterol content of phloem from host trees of both beetle species containing feeding larvae (and symbiotic fungi) also was compared with that of uninfested phloem.

**Materials and Methods**

**Fungal Identification and Growth for Extraction.** Fungal isolates used in this study were taken from the culture collection of D.L.S. and were initially identified by using morphological characteristics as well as keys and descriptions contained in Upadhyay (1981), Grylls and Seifert (1993), and Jacobs and Wingfield (2003). The identity of select isolates was further confirmed using DNA sequencing and morphological comparisons with known strains (Lee et al. 2003) (J. Kim and C. Breuil, personal communication) (Table 1).

Three replicates of each isolate of each of the three fungal species were analyzed for ergosterol content. Each replicate was grown on 2% malt extract agar for 10 d. Then, a small plug of mycelium of each isolate was placed into a 123-ml Erlenmeyer flask containing 2% malt yeast extract broth and allowed to grow for an additional 10 d. Fungal mats were removed from the liquid broth, placed on a filter, and rinsed with distilled water while a vacuum was applied to remove culture broth residue. Mycelial mats were then freeze-dried in individual vials. After lyophilization, vials were topped...
with nitrogen and held in the dark at 2°C until ergosterol extraction.

**Phloem Sampling.** To compare concentrations of ergosterol estimated from known fungal isolates in culture with actual ergosterol concentrations consumed by larvae feeding in trees colonized by fungal associates, we collected and extracted ergosterol from phloem from trees infested by the two host beetles. Samples of phloem were collected on 16 April 2003 from four *D. rufipennis*-infested *Picea englemanni* Parry and four uninfested *P. englemanni* at three sites on the Fishlake National Forest, Utah. Phloem samples from one *D. ponderosa*-infested *Pinus contorta* Douglas and four uninfested *P. contorta* were collected 28 January 2003 on the Wasatch Cache National Forest, Utah. Additional phloem samples were collected 23 April 2003 from four *D. ponderosa*-infested *P. contorta* and two uninfested *P. contorta* on the Sawtooth National Forest, Idaho. Phloem was collected by using a sterile knife to remove a 100-cm² section of phloem from live and beetle-infested tree boles 1.37 m above the ground. Samples removed from trees infested with *D. rufipennis* and *D. ponderosa* were taken from areas where larvae were mining. Phloem samples were transported to the laboratory on ice, oven-dried for 24 h, and stored at −30°C until lyophilization.

**Ergosterol Determination.** Before transfer to extraction vials, the dry weight of each mycelial and phloem sample was recorded to the nearest 0.1 mg. Freeze-dried fungal samples used in the extraction process weighed ~0.03 g and phloem samples ~1.0 g. Ergosterol content of the fungal mats and phloem was analyzed using previously published methods (Newell et al. 1988, Padgett and Posey 1993). High-performance liquid chromatography (HPLC)-grade methanol (20 ml) was added to a round-bottomed flask containing the dried mycelium or phloem and refluxed for 2 h at 80°C. Five milliliters of a 4% solution of KOH in 95% ethanol was added to the neutral extracts obtained from the mehanolic pathway and refluxed for 30 min at 80°C to hydrolyze the esters. Sterols were removed from the alcoholic base by partitioning into pentane after cooling to 25°C and adding 10 ml of distilled water. A series of three petri dish additions (10, 5, and 5 ml) were mixed by repeatedly inverting tightly sealed flasks. After each addition and mixing, the solution was placed in a separatory funnel. The pentane layer was withdrawn, and pentane fractions from each sample were pooled. The pentane samples were evaporated in open glass flasks at 30°C under a stream of nitrogen. Each sample was redissolved in 2 ml of methanol, and the vials were tightly sealed and sonicated for 5 min to enhance the rate of dissolution. Redissolved samples were filtered through 0.22-µm polytetrafluoroethylene membranes and stored in the dark at 2°C for 1–7 d until HPLC analysis.

Samples were analyzed by isocratic liquid chromatography on an Agilent 1100 (Agilent Technologies, Palo Alto, CA) by using a 250- by 4.6 mm 5-µm Supelco LiChrospher RP18 column (Supelco, Bellefonte, PA) held at 45°C. Samples (5 µl) were injected by a thermostatted autosampler cooled to 5°C by using methanol (2 ml/min) as a mobile phase. Ergosterol was detected by an Agilent diode array detector at 282 nm and quantified using an ergosterol four point external standard calibration curve (Sigma, St. Louis, MO). Three replicates of each extracted fungal and phloem isolate were analyzed. Final ergosterol concentration of each sample was expressed as milligrams of ergosterol per gram of fungal or phloem dry mass.

**Statistical Analysis.** Differences in ergosterol concentration among the fungal species were analyzed using mixed model analysis within SAS (Littell et al. 1996). Fungal species was included as a fixed effect, and replicate and isolate within species were included as random effects. Because the phloem data violated normality assumptions, and our main objective was to test for presence or absence of ergosterol in uninfested and infested phloem, we transformed the data to a binomial format. Samples with an ergosterol quantity >0 were coded as 1 and as 0 otherwise. GLIMMIX, a SAS procedure for fitting generalized linear mixed models (Littell et al. 1996), was used to accommodate the random effects of replicate and isolate within species, with host tree species included as a fixed effect and a binomial error distribution.
Results

Mycelia of the three fungal species contained relatively large levels of ergosterol, as determined after 20 d growth at 20°C. However, no significant differences were found among the fungal species in total ergosterol content (Fig. 1). The probability of containing ergosterol was significantly greater in *P. engelmannii* and *P. contorta* phloem infested with *Dendroctonus* larvae (and their fungal symbionts) than uninfested phloem samples which contained either no detectable ergosterol or only trace amounts (*F* = 8.92; *df* = 1, 21.56; *P* = 0.0069) (Fig. 2). We found no significant differences in ergosterol concentrations between the two host tree species or among sample dates.

Discussion

We found no differences in the ergosterol content of mycelia of *O. montium* and *O. clavigerum* isolated from *D. ponderosa* mycangia and *L. abietinum* isolated from the exoskeleton of *D. rufipennis*. Mean percentage of ergosterol (0.88–1.06% dry weight) of these fungi was similar to levels found in filamentous fungi associated with decaying building materials (0.26–1.40%) (Pasanen et al. 1999), but it was much greater than levels quantified from fungal associates of *Xyleborus* ambrosia beetles (0.12–0.24%) (Kok and Norris 1973). *Xyleborus* feed almost entirely on concentrated spore layers of their associated fungi, and the ergosterol obtained from these layers is vital to successful development and reproduction (Baker and Norris 1968). Because *Dendroctonus* larvae feed on phloem colonized by fungal hyphae (A.S. Adams and D.L.S., unpublished data), they consume fungal biomass in much lower concentrations than do ambrosia beetles. Consequently, the ergosterol in the mycelium of the fungi would in effect be diluted by the phloem within which the fungi grow. This may necessitate a greater amount of ergosterol per dry weight of mycelium to fulfill the basic sterol requirements of the host beetles.

As expected, phloem of uninfested host trees contained zero to only trace levels of ergosterol. Conversely, phloem taken from trees colonized by *D. ponderosa* and *D. rufipennis* and their fungal symbionts contained significantly more ergosterol than did phloem taken from uninfested trees. Samples were taken from infested trees when both *D. ponderosa* and *D. rufipennis* were in larval life stages. Larvae of *Dendroctonus* feed upon both fungi and phloem as they mine (A.S. Adams and D.L.S., unpublished data). Consequently, they potentially gain nutrients from both the host tree and the fungal symbionts. Our results suggest that larval life stages of *D. ponderosa* and *D. rufipennis* may obtain sterols, in the form of ergosterol, while mining larval galleries.

Only one nonambrosia scolytine, the coffee berry borer, has been assessed for dependence upon ergosterol provided by its symbiotic fungus (Morales-Ramos et al. 2000). In laboratory experiments, it was determined that a diet containing an ergosterol concentration between 0.015 and 0.020% (dry weight) was required to achieve normal reproduction by the beetle. Although uninfested coffee berries contained no ergosterol, coffee berries colonized by the fungal symbiont contained levels of ≈0.011%. Although this level is slightly below that required by the beetle, the authors attributed this to the fact that whole berry extractions were used in their analyses rather than extractions of berry tissues directly adjacent to larvae where colonization of the berry by the fungus, and consequently ergosterol concentrations, are likely to be the greatest. Similarly, our infested phloem samples, which contained less ergosterol than did the fungal-colonized coffee berries (0.006%), were
taken from areas of host trees where larvae were mining, but most likely also included phloem uncolol-
ized by fungi. Because of the presence of sac mycangia and known mycophagy in *D. pondersoae*, we hypothesized that its associated fungus would contain more ergosterol and thereby potentially confer a greater benefit to its host than would *L. abietinum* associated with *D. rufipennis*, which possesses only rudimentary pit mycangia. Sac mycangia are specialized structures thought to have evolved specifically to carry fungi, thereby suggesting a more obligate relationship between beetle host and fungal associate (Six 2003a). We also hypothesized that *O. clavigerum* would provide greater quantities of ergosterol to *D. pondersoae* than would *O. montium*. *O. clavigerum* has a long shared evolutionary history with *D. pondersoae* (Six and Paine 1999), whereas *O. montium* is considered a more recent associate (Six and Paine 1999). However, although we found no difference in the amount of ergosterol among any of the primary fungi, our results suggest that these associates of *D. pondersoae* and *D. rufipennis* do make relatively large quantities of ergosterol available to their beetle hosts. Future studies should investigate the degree of mycophagy and the use and degree of dependence of *Dendroctonus* beetles on ergosterol provided by their fungal symbionts.

**Acknowledgments**

We thank Jackie Redmer and Chris Heck for technical support in ergosterol extraction and analysis and the Utah State University Biotechnology Center for equipment use. Joyce Knoblett (USDA-ARS Bee Lab, Logan, UT) conducted HPLC analyses of all samples. We thank Jae Kim and Colette Breuil for assistance with fungal identifications. Matt Hansen, Jim Vandygriff, and Rebecca Gerhardt assisted with field collections. Aaron Adams, Kathy Bleiker, and Colette Breuil provided valuable comments on an earlier version of this manuscript. Partial funding for this study was provided by USDA-CSREES National Research Initiative Grant 2001-35302-10985.

**References Cited**

Ayres, M. P., R. T. Wilkens, J. J. Ruel, M. J. Lombardero, and E. Vallery. 2000. Nitrogen budgets of phloem-feeding bark beetles with and without symbiotic fungi (Coleoptera: Scolytidae). Ecology 81: 2198–2210.

Baker, J. M., and D. M. Norris. 1965. A complex of fungi mutualistically involved in the ambrosia beetle *Xyleborus ferrugineus*. J. Invertebr. Pathol. 11: 246–250.

Bridges, J. R. 1983. Mycangial fungi of *Dendroctonus frontalis* (Coleoptera: Scolytidae) and their relationship to beetle population trends. Environ. Entomol. 12: 858–861.

Clayton, R. B. 1964. The utilization of sterols by insects. J. Lipid Res. 5: 3–19.

Coppedge, B. R., F. M. Stephen, and G. W. Felton. 1995. Variation in female southern pine beetle size and lipid content in relation to fungal associates. Can. Entomol. 127: 145–154.

French, J.R.J., and R. A. Roepers. 1975. Studies on the biology of the ambrosia beetle, *Xyleborus dispar* (F.) (Coleoptera: Scolytidae). Z. Ang. Entomol. 78: 241–247.

Goldhammer, D. S., F. M. Stephen, and T. D. Paine. 1990. The effect of the fungi *Ceratocystis minor* (Hedgecock) Hunt, *Ceratocystis minor* (Hedgecock) Hunt var. *Barrasii* Taylor, and SJ.B 122 on reproduction of the southern pine beetle, *Dendroctonus frontalis* Zimmermann (Coleoptera: Scolytidae). Can. Entomol. 122: 407–418.

Grylls, B. T., and K. A. Seifert. 1993. A synoptic key to species of *Ophiostoma*, *Ceratocystis*, and *Ceratoxyctis*. pp. 261–268. In M. J. Wingfield, K. A. Seifert, and J. F. Webber [eds.]. *Ceratocystis and Ophiostoma*: taxonomy, ecology, and pathogenicity. APS Press, St. Paul, MN.

Harrington, T. C. 2005. Ecology and evolution of mycophagous bark beetles and their fungal partners, pp. 257–291. In F. E. Vega and M. Blackwell [eds.]. Insect-fungal associations: ecology and evolution. Oxford University Press, New York.

Jacobs, K., and M. J. Wingfield. 2003. *Leptographium* species: tree pathogens, insect associates, and agents of blue stain. APS Press, St. Paul, MN.

Kok, L. T. 1979. Lipids of ambrosia fungi and the life of mutualistic beetles, pp. 33–52. In L. R. Batra [ed.], *Insect fungus symbioses: nutrition, mutualisms and commensals*. Wiley, New York.

Kok, L. T., and D. M. Norris. 1973. Comparative sterol compositions of adult female *Xyleborus ferrugineus* and its mutualistic fungal ectosymbionts. Comp. Biochem. Physiol. 44: 499–505.

Kok, L. T., D. M. Norris, and H. M. Chu. 1970. Sterol metabolism as a basis for a mutualistic symbiosis. Nature (Lond.) 225: 661–662.

Kramer, P. J., and T. T. Kozlowski. 1960. Physiology of trees. McGraw-Hill, New York.

Littell, R. R., G. A. Milliken, W. W. Stroup, and R. D. Wolfinger. 1996. SAS system for mixed models. SAS Institute, Cary, NC.

Lee, S., J. J. Kim, S. Fung, and C. Breuil. 2003. A PCR-RFLP marker distinguishing *Ophiostoma clavigerum* from morphologically similar *Leptographium* species associated with bark beetles. Can. J. Bot. 81: 1104–1112.

Maurer, P., D. Debieu, C. Malosse, P. Leroux, and G. Ribé. 1992. Sterols and symbiosis in the leaf-cutter ant *Acro-
myrmex octospinosus* (Reich) (Hymenoptera, Formi-
cideae: Attini). Arch. Insect Biochem. Physiol. 20: 13–21.

Morales-Ramos, J. A., M. G. Rojas, H. Sittertz-bhatkar, and G. Saldaña. 2000. Symbiotic relationship between *Hy-
pothenus hempei* (Coleoptera: Scolytidae) and *Fusar-
ium solani* (Moniliales: Tuberculariaceae). Ann. Entomol. Soc. Am. 93: 541–547.

Mueller, U., N. Gerardo, D. Aanen, D. L. Six, and T. Schultz. 2005. The evolution of agriculture in insects. Annu. Rev. Ecol. Evol. Syst. 36: 563–595.

Newell, S. Y., T. L. Arsuffi, and R. D. Fallon. 1988. Fundamental procedures for determining ergosterol content of decaying plant material by liquid chromatography. Appl. Environ. Microbiol. 54: 1857–1879.

Norris, D. M. 1972. Dependence of fertility and progeny development of *Xyleborus ferrugineus* upon chemicals from its symbionts, pp. 299–310. In D.J.L. Harding and R. A. Stuttard [eds.], *Insect and mite nutrition*. North-Holland, Amsterdam, The Netherlands.

Norris, D. M., and J. K. Baker. 1967. Symbiosis: effects of a mutualistic fungus upon the growth and reproduction of *Xyleborus ferrugineus*. Science (Wash. DC) 156: 1120–1122.

Norris, D. M., J. K. Baker, and H. M. Chu. 1969. Symbiotic interrelationships between microbes and ambrosia beetles. III. Ergosterol as the source of sterol to the insect. Ann. Entomol. Soc. Am. 62: 413–414.
Nasir, H., and H. Noda. 2003. Yeast-like symbiotes as a sterol source in anobiid beetles (Coleoptera, Anobiidae): possible metabolic pathways from fungal sterols of 7-dehydrocholesterol. Arch. Insect Biochem. Physiol. 52: 175–182.

Padgett, D. E., and M. H. Posey. 1993. An evaluation of the efficiencies of several ergosterol extraction techniques. Mycol. Res. 97: 1476–1480.

Pasanen, A., K. Yki-Pietila, P. Pasanen, P. Kalliokoski, and J. Tarhanen. 1999. Ergosterol content in various fungal species and biocontaminated building materials. Appl. Environ. Microbiol. 65: 138–142.

Reynolds, K. M. 1992. Relations between activity of Dendroctonus rufipennis Kirby on Lutz spruce and blue stain associate with Leptographium abietinum (Peck) Wingfield. For. Ecol. Manage. 47: 71–86.

Richmond, J. A., and H. A. Thomas 1975. Hylobius pales: effect of dietary sterols on development and on sterol content of somatic tissue. Ann. Entomol. Soc. Am. 68: 329–332.

Six, D. L. 2003a. A comparison of mycangial and phoretic fungi of individual mountain pine beetles. Can. J. For. Res. 33: 1331–1334.

Six, D. L. 2003b. Bark beetle-fungus symbioses. pp. 99–116. In K. Bourtizs and T. A. Miller [eds.], Insect symbiosis. CRC, Boca Raton, FL.

Six, D. L., and B. J. Bentz. 2003. Fungi associated with the North American spruce beetle, Dendroctonus rufipennis. Can. J. For. Res. 33: 1815–1820.

Six, D. L., and T. D. Paine. 1998. Effects of mycangial fungi and host tree species on progeny survival and emergence of Dendroctonus jeffreyi and D. ponderosae. Environ. Entomol. 27: 1393–1401.

Six, D. L., and T. D. Paine. 1999. Phylogenetic comparison of ascomycete mycangial fungi and Dendroctonus bark beetles (Coleoptera: Scolytidae). Ann. Entomol. Soc. Am. 92: 159–166.

Solheim, H. 1995. A comparison of blue-stain fungi associated with the North American spruce beetle Dendroctonus rufipennis and the Eurasian spruce bark beetle Ips typographus. pp. 61–67. In D. Amlid [ed.], Proceedings, Forest Pathology Research in the Nordic Countries. SNS Conference, 9–12 August 1994, Biri, Norway. Norsk Institutt for Skogforskning, As, Norway.

Svoboda, J. A., M. J. Thompson, W. E. Robbins, and J. N. Kaplanis. 1978. Insect Sterol Metab. Lipids 13: 742–753.

Upadhyay, H. P. 1981. A monograph of Ceratocystis and Ceratocystiopsis. University of Georgia Press, Athens, GA.

Weete, J. D. 1973. Sterols of the fungi: distribution and biosynthesis. Phytochemistry 12: 1843–1864.

Weete, J. D. 1989. Structure and function of sterols in fungi. Adv. Lipid Res. 23: 115–167.

Whitney, H. S. 1971. Association of Dendroctonus ponderosae (Coleoptera: Scolytidae) with blue-stain fungi and yeasts during brood development in lodgepole pine. Can. Entomol. 103: 1495–1503.

Received 14 July 2005; accepted 6 December 2005.