Detection and quantification of Leptographium wageneri, the cause of black-stain root disease, from bark beetles (Coleoptera: Scolytidae) in Northern California using regular and real-time PCR

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Abstract: Black-stain root disease is a threat to conifer forests in western North America. The disease is caused by the ophiostomatoid fungus Leptographium wageneri (W.B. Kendr.) M.J. Wingf., which is associated with a number of bark beetle (Coleoptera: Scolytidae) and weevil species (Coleoptera: Curculionidae). We developed a polymerase chain reaction test to identify and quantify fungal DNA directly from insects. Leptographium wageneri DNA was detected on 142 of 384 bark beetle samples (37%) collected in Lassen National Forest, in northeastern California, during the years 2001 and 2002. Hylastes macer (LeConte) was the bark beetle species from which Leptographium DNA was amplified most regularly (2001: 63.4%, 2002: 75.0% of samples). Lower insect–fungus association rates were found for Hylurgops porosus (LeConte), Hylurgops subcostatus (Mannerheim), Hylastes gracilis (LeConte), Hylastes longicollis (Swaine), Dendroctonus valens (LeConte), and Ips pini (Say). The spore load per beetle ranged from 0 to over 1 × 10^3 spores, with only a few beetles carrying more than 1 × 10^3 spores. The technique permits the processing of a large number of samples synchronously, as required for epidemiological studies, to study infection rates in bark beetle populations and to identify potential insect vectors.

Résumé : Le noircissement des racines est une maladie qui menace les forêts de l’ouest de l’Amérique du Nord. La maladie est causée par Leptographium wageneri (W.B. Kendr.) M.J. Wingf., un champignon de la famille des Ophiostomatidae qui est associé à plusieurs espèces de scolytes (Coléoptères : Scolytidae) et de charançons (Coléoptères : Curculionidae). Nous avons mis au point un test de réaction en chaîne de la polymérase pour identifier et quantifier l’ADN fongique directement à partir des insectes. L’ADN de L. wageneri a été détecté sur 142 des 384 (37 %) échantillons de scolytes collectés dans la forêt nationale de Lassen, dans le nord-est de la Californie, au cours des années 2001 et 2002. Hylastes macer (LeConte) était l’espèce de scolyte à partir duquel l’ADN de Leptographium a été le plus régulièrement amplifié (2001 : 63.4 %, 2002 : 75.0 % des échantillons). Des taux d’association plus faibles ont été observés pour Hylurgops porosus (LeConte), Hylurgops subcostatus (Mannerheim), Hylastes gracilis (LeConte), Hylastes longicollis (Swaine), Dendroctonus valens (LeConte), et Ips pini (Say). La charge de spores par insecte variait de 0 à plus de 1 × 103 spores, avec seulement quelques insectes transportant plus de 1 × 10^3 spores. La technique permet de traiter simultanément un grand nombre d’échantillons, comme l’exigent les études épidémiologiques pour étudier le taux d’infection dans les populations de scolytes et identifier les insectes vecteurs potentiels.

Introduction

Black-stain root disease (BSRD) causes considerable damage to conifer forests in western North America. Symptoms associated with BSRD include reduced growth, chlorosis, dark staining of the tracheids extending from the roots to the lower bole, and ultimately tree death (Jacobs and Wingfield 2001). The causal agent of BSRD is Leptographium wageneri (W.B. Kendr.) M.J. Wingf. (syn. Verticiladiella wageneri W.B. Kendr.), which occurs in three host-specific varieties: L. wageneri var. ponderosum infects ponderosa pine (Pinus ponderosa Dougl. ex P. & C. Laws.), Jeffrey pine (Pinus jeffreyi Grev. & Balf.), and lodgepole pine (Pinus contorta Dougl. ex Loud.); L. wageneri var. pseudotsugae infects Douglas-fir (Pseudotsuga menziesii (Mirbel) Franco); and L. wageneri var. wageneri infects mainly pinyon pine (Pinus monophylla Torr. & Frem., Pinus edulis Engelm.), although a number of other coniferous hosts have been reported (Jacobs and
Wingfield 2001). The teleomorph stage of *L. wageneri* var. *ponderosum*, *Ophiostoma wageneri* (Goheen & F.W. Cobb) T.C. Harr. (Harrington 1987), has been isolated only once from ponderosa pine (Goheen and Cobb 1978), and its ecological role remains unknown. No sexual stages of the varieties *pseudotsugae* and *wageneri* have been reported.

There appear to be significant differences in the epidemiology and biology of the three host-specific varieties, and not all three are equally understood. For instance, in the Douglas-fir variety, disease severity is related to forest silvicultural practice; for example, young, dense Douglas-fir plantations are much more vulnerable than old-growth stands (Hansen 1997).

In ponderosa pine, short-distance spread of BSRD occurs by root-to-root transmission of the pathogen (Wood et al. 2003). Insect vectors are assumed to play a crucial role in overland spread of fungal inoculum, but experimental evidence is lacking for both the ponderosa and the pinyon varieties of the pathogen. Conidia of *Leptographium* and related ophiostomatoid fungi are produced in sticky masses (conidial droplets) at the apex of stalked conidiophores, an adaptation for dispersal by insect vectors. *Leptographium wageneri* is associated with bark beetles (Coleoptera: Scolytidae) such as *Dendroctonus* spp. (Wagener and Mielke 1961), *Hylastes* spp. (Goheen 1976), *Hylurgops* spp. (Wagner 1977), and *Ips* spp. (Morrison and Hunt 1988). Weevils (Coleoptera: Curculionidae) of the genera *Pissodes* (Witcosky 1981) and *Steremnius* (Witcosky 1981) have also been implicated as vectors in Douglas-fir.

Detection of black-stain fungi from infested wood and insect vectors has been carried out traditionally using selective media containing cycloheximide (Harrington 1992; Witcosky 1985). The isolates obtained from the environmental samples have to be purified and then identified morphologically. The procedure is time-consuming and successful isolations of *L. wageneri* from insects are rarely obtained. Isolation of *L. wageneri* var. *ponderosum* from insects attacking ponderosa pine has never been reported to our knowledge. These problems have severely limited ecological and pathological studies of this insect–disease complex. Recently, molecular detection methods based on genus- and (or) species-specific primers have been developed for a variety of ophiostomatoid strains (Kim et al. 1999), but such tests have not been developed yet for studies on BSRD.

The purpose of this study is (i) to establish a rapid molecular method to detect *L. wageneri* DNA from insects, (ii) to determine the occurrence of *L. wageneri* var. *ponderosum* on different scolytid bark beetle species in a heavily infected ponderosa pine forest in Northern California, and (iii) to quantify the individual spore load on a given beetle using real-time polymerase chain reaction (PCR).

## Materials and methods

### Study site

The study area (40°39'N, 121°12'W, 1800 m elevation), located on the Lassen National Forest, Lassen County, California, has active black-stain root disease centers resulting in current tree mortality. Ponderosa and Jeffrey pines are the predominant tree species with white fir (*Abies concolor* (Gord. & Glend.) Lindl. ex. Hildebr.) representing less than 15% of the basal area; thus infection can be assumed to be exclusively *L. wageneri* var. *ponderosum*. The approximate age of the pines is 80–100 years, and the average diameter at breast height is 45 cm. Sixteen permanent 3.0-ha plots, previously established on this location to study effects of silvicultural treatments on disease spread, were utilized in this study to obtain insect samples. Existing black-stain root disease center sizes in these plots ranged from individual infected trees to 0.5-ha mortality gaps.

### Insect sampling

Bark beetles were collected weekly during the flight period from mid-April to mid-June in 2001 and 2002. Insects were trapped using 68 Lindgren funnel traps (4-funnels, Phero Tech Inc., Delta, British Columbia) baited with the standard ethanal lure (Phero Tech Inc.) and UHR alpha pinene lure (Phero Tech Inc.) (Lindgren 1983). Trapped insects were stored at −20 °C before processing. Insect species were identified based on morphology. To determine phoresy of *L. wageneri* by individual beetle species, one to five specimens belonging to the same species and collected in the same trap were pooled together and screened by PCR as an individual sample. In the case of *Hylurgops porosus*, the most abundant species, up to 20 specimens from the same trap were pooled together. Length and diameter measurements were taken from 25 specimens of *Hylurgops porosus* (LeConte), *Hylurgops subcostatus* (Mannerheim), *Hylastes macer* (LeConte), *Dendroctonus valens* (LeConte), and *Ips pini* (Say), and from three specimens of *Hylastes gracilis* (LeConte). Surface area and volume of the beetles were calculated by assuming a cylindrical body shape.

### Fungal growth and spore harvest

*Leptographium* strains (Table 1) were grown for 8 weeks at 18 °C on semi-selective medium, which consists of the following per litre: 15.0 g Bacto™ malt extract (Becton Dickinson, Sparks, Maryland), 1.5 g Bacto™ yeast extract (Becton Dickinson), 15 g agar (Fisher Scientific, Pittsburgh, Pennsylvania), 0.3 g streptomycin sulfate (Sigma, St. Louis, Missouri), and 0.2 g cycloheximide (Sigma). Conidiophores growing on 1.0-cm² agar slices (five replicates) were counted at 15× magnification using a stereomicroscope (Nikon SMZ 1000). Each agar slice was added to a microtube containing 1 mL H₂O and vortexed for 2 min to dislodge spores. The harvested spores were counted using a hematocytometer (Hauser Scientific, Horsham, Pennsylvania) with a light microscope (Nikon Eclipse E400) and diluted to final concentrations of 1 × 10⁵ – 1 × 10⁶ spores/mL.

### Primer design

The *L. wageneri* specific primer combination LEPTO1 (CAAAGACGCGACGACGAGTCCTC) and LEPTO2 (GTTG-CCAGGGAACCTCGGAAG) were developed using the primer3 software (www-genome.wi.mit.edu/cgi-bin/primer/primer3 www.cgi). Primer binding sites were chosen based on the alignment using ClustalX (Thompson et al. 1997) of about 640 bp in the ITS2 and 28S rRNA gene in the nuclear ribosomal region of 44 *Leptographium* and *Ophiostoma* strains (Jacobs et al. 2001) (Fig. 1). Because DNA sequences at the primer binding sites for the ITS2 and 28S loci are identical across all three varieties of *L. wageneri*, the
primers are specific to the species complex rather than one variety. However, because the pathogen varieties are host specific and do not co-occur in our study area, this is unlikely to be problematic in our or other study sites. The amplicon size was 230 bp. Primers were synthesized by Qiagen–Operon Inc., Alameda, California. Specificity of the primers was tested by running PCR reactions on 16 isolates of L. wageneri from across the pathogen’s range as well as on 15 other Leptographium and forest pathogens (Table 1).

DNA extraction and PCR amplification

DNA from fungal colonies grown on selective medium was extracted using a “hyphal tipping” method: mycelium was scraped from the Petri dish and suspended in 100 µL of distilled water, frozen on dry ice for 3 min, thawed at 75 °C, and vortexed for 2 min. Freezing and thawing was repeated three times, with the last thaw extended to 15 min. Samples were spun down for 5 min at 15 700g, and 6.25 µL of the supernatant was used for PCR reactions. Suspensions of varying concentrations of fungal spores were employed to develop reference standards for DNA-based quantification of spore loads found on insects. Fungal spore suspensions and bark beetles were added to a microtube and mechanically disrupted by shaking with 6.35 mm diameter glass beads for 30 s using an amalgamator (BIO 101, Carlsbad, California). After adding 0.3 mL of 2% CTAB (cetyltrimethylammonium bromide), the samples were subjected to three freeze–thaw cycles (2 min on dry ice, 2 min at 75 °C, repeated three times with the last thaw extended to 30 min). DNA was extracted using 0.35 mL phenol–chloroform–isoamyl alcohol followed by purification using the Geneclean kit (BIO 101), following instructions by the manufacturer.

The PCR mix consisted of 8.0 µL H₂O, 2.5 µL PCR buffer (Promega, Madison, Michigan), 5.0 µL MgCl₂ (25 mmol/L, Promega), 2.5 µL dNTP (2 mmol/L, Promega), 0.25 µL primer LEPTO1 (50 mmol/L), 0.25 µL primer LEPTO2 (50 mmol/L), 0.25 µL Taq polymerase (Promega), and 6.25 µL DNA template. The cycling profile was denaturation at 94.0 °C for

<table>
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</table>

aATCC, American Type Culture Collection, Rockville, Md.; UC Berkeley, University of California, Berkeley. T.C. Harrington, Department of Natural Resource Ecology and Management, Iowa State University.
180 s, followed by 45 cycles of 94 °C for 35 s, 65 °C for 55 s, 72 °C for 50 s, then final extension at 72 °C for 10 min. PCR products were checked on a 1.5% agarose gel stained with ethidium bromide. Negative PCR reactions were checked for failed DNA extraction using the universal primers ITS1 and ITS4 (White et al. 1990).

Real-time PCR amplification for the quantification of fungal spores

Spore suspensions from *L. wageneri* var. *ponderosum* (ATCC 58574) diluted in H$_2$O to final concentrations of 1 × 10$^1$ – 1 × 10$^6$ spores/mL were used to establish standard curves. Real-time PCR using SYBR$^\text{®}$ Green I (Applied Biosystems, Foster City, California) was performed on an iCycler (Bio-Rad, Hercules, California). To quantify the starting amount of template DNA, the threshold cycle (Ct) for each sample was calculated. The Ct value, which is the cycle number when a significant increase of SYBR Green fluorescence is first recorded against the background fluorescence level, was calculated using baseline cycles 2 to 10. The Ct value is proportional to the logarithm of the initial DNA concentration. The data analysis window was set at 10.00% of a cycle and centered at the end of the cycle. Reaction conditions for real-time PCR were 7.5 µL H$_2$O, 2.5 µL PCR buffer, 5.0 µL MgCl$_2$ (25 mmol/L), 2.5 µL dNTP, 0.25 µL primer LEPTO1 (50 mmol/L), 0.25 µL primer LEPTO2 (50 mmol/L), 1.25 µL SYBR Green, 1.25 µL fluorescein (Bio-Rad, Hercules, California), 0.25 µL Taq polymerase, and 6.25 µL DNA template. Cycling parameters were 94.0 °C for 180 s, followed by 45 cycles of 94 °C for 35 s, 62 °C for 55 s, 72 °C for 50 s, and a final extension at 72 °C for 10 min. Melting curves were measured for 110 cycles of 10 s each, starting at 62 °C and increasing by 0.3 °C each cycle. Melt-curve peaks can be displayed graphically as d(relative fluorescence units)/d temperature. The temperature at which dsDNA fragments are completely denatured into single strands is dependent not only on the size of the fragments, but also on their guanine–cytosine content. Melt-curve analysis is thus more powerful than gel electrophoresis and can be used to verify specificity of PCR amplicons (Hayden et al. 2004).

To test the possible inhibitory effects of beetle DNA on quantification of *L. wageneri* spore, loads spore standards were spiked with extracted DNA from PCR negative beetles. These were run alongside clean reactions containing the same spore standard to compare threshold cycles for signs of inhibition. Reactions were carried out using identical real-time PCR protocols (see above) except that 6.25 µL of beetle DNA extract was added to the spiked standard in lieu of an equal volume of H$_2$O in the reaction. Multiple replicates were performed for all spore concentrations for each of the four
beetle species that were collected in traps both years (D. valens, 
Hylurgops subcostulatus, Hylastes macer, Hylurgops porosus).

The quantification parameters described herein were used to perform quantitative PCR reactions on individual beetles that had resulted positive for *L. wageneri* using the standard PCR assay. Quantification of fungal spores was not attempted for samples consisting of pooled insects.  

**Sequence analyses**

Regular PCR products were cleaned via QiaQuick PCR Purification kit (Qiagen Sciences, Valencia, California), as per kit instructions, except that cleaned products were eluted in 30 μL salt-free water. Cleaned products were cycle sequenced with 4 μL BigDye Terminator version 3.0 (Applied Biosystems (ABI)), 2.4 pmol salt-free primer, and 5–20 ng DNA (template concentration determined by gel). Cycle sequencing was performed on an iCycler (Bio-Rad) according to ABI-recommended protocol: 26 cycles of 96 °C for 10 s, 55 °C for 5 s, 60 °C for 4 min, followed by a hold at 4 °C. Samples were desalted in ethanol as per ABI instructions (80 °C for 5 s, 60 °C for 4 min, followed by a hold at 4 °C). Samples were brought up in 10 μL salt-free water. Cleaned products were cycle sequenced for samples consisting of pooled insects.

**Results**

**Insect sampling**

Bark beetles were collected weekly from all traps during both sampling seasons (Table 2). Overall insect captures were considerably higher in 2001 (250 samples) than in 2002 (134 samples). Four species were collected in both years: *Hylurgops porosus* was the most prevalent species in both years, followed by *Hylurgops subcostulatus, Hylastes macer,* and *D. valens* (Table 2). Low numbers of *I. pini, Hylastes gracilis,* and *Hylurgops reticulatus* (Wood) were collected only in 2001. *Hylastes tenuis* (Eichhoff) was collected from two traps only in 2002.

** Primer specificity**

The primers LEPTO1 and LEPTO2 amplified a single PCR product of 230 bp from genomic DNA in all 16 isolates of *L. wageneri* tested, including all three varieties of the pathogen (Table 1). The primers did not amplify most other *Leptographium* and *Ophiostoma* strains tested, with the exception of *Leptographium neomexicanum, Leptographium reconditum,* and *Leptographium serpens.*

**Detection of Leptographium DNA from naturally infected bark beetles using regular PCR**

A total of 384 insect samples (for a total of 908 insect specimens) were analyzed for the presence of *Leptographium* DNA using the primers LEPTO1 and LEPTO2 (Table 2). An
initial set of experiments was carried out using *Hylurgops porosus*, the most abundant bark beetle species, to test for the lowest number of insects from which fungal DNA could be detected. The 230-bp *L. wageneri* DNA fragment was amplified regularly from tubes were DNA extractions were carried out on 20, 10, 5, and even single individual insects (Fig. 2). For the other experiments, one to five insects were pooled and processed in a single tube. A PCR product was obtained from 91 of 250 samples (36.4%) collected in 2001. In 2002, 134 samples were collected, with 51 showing a positive PCR amplification (38.1%). PCR positives were not found among bark beetles collected from seven traps (10.3%). All PCR negatives using LEPTO1 and LEPTO2 were successfully amplified using the universal primers ITS1 and ITS4.

**Identity of the PCR fragments**

Specificity of all products was verified by analyses of melt curves. Representative PCR fragments obtained from each insect species (two per species) and from each tester *L. wageneri* var. *ponderosum* isolate (Table 1) were sequenced and compared with the GenBank using BLAST search. All samples showed a 100% identity with the published sequence of *L. wageneri* var. *ponderosum*.

**Number of fungal spores and conidiophores on selective medium**

The number of conidiophores and spores produced by reference strains of *L. wageneri* var. *ponderosum* (Table 1) was counted after 8 weeks of growth on selective medium. On average, 143–202 conidiophores were counted per square centimetre (mean = 169/cm²). The number of spores ranged between 1.13 × 10⁷ and 2.40 × 10⁷ conidiospores/cm² (mean = 1.85 × 10⁷). This would correspond to approximately. 0.70 × 10⁵ – 1.38 × 10⁵ conidiospores per conidiophore (mean = 1.10 × 10⁵ spores per conidiophore). The diameter of a single conidial droplet was approximately 100 μm.

**Quantification of fungal spores using real-time PCR**

The calibration curve for DNA extracted from *L. wageneri* spores is shown in Fig. 3 (three replicates). Threshold cycles (Ct) were negatively correlated with the number of spores in the range from 1 × 10² to 1 × 10⁶ spores/mL. When a spore concentration of 1 × 10⁵ spores/mL was used, a PCR fragment was still amplified and detected on the agarose gel, but the threshold cycles became inconsistent (data not shown). The 230-bp PCR fragment showed a sharp melt peak at 89.5 °C (±1.0 °C), which was clearly distinguishable from the flattened peaks produced by primer dimers at about 82.0 °C (data not shown).

At spore concentrations >1 × 10³, there was no significant difference between clean and spiked standards (one-tailed paired sample t test, t = 1.21, 34 df, p = 0.12). At spore concentrations ≤1 × 10³ *L. wageneri* var. *ponderosum* DNA could be amplified in spiked samples, but its quantification was unreliable. In 89% of 44 reactions, in fact, PCR resulted in amplicons characterized by the exact expected size and melt-curve profile for *L. wageneri*, but threshold cycles were often too low or unscorable for quantitative purposes. The effect of beetle species on standard threshold cycles was not significant in a one-way ANOVA ($F_{3,42} = 0.8767$, p = 0.46).

Only PCR-positive insect samples consisting of a single specimen were used for the quantitative analysis with real-time PCR. DNA quantification was thus performed for a total of 44 samples from 2001 and 27 samples from 2002. The number of *Leptographium* spores carried by single bark beetles was inferred from the comparative analysis between DNA amounts found in suspensions of known spore concentration and DNA amounts in the tested insect samples (Fig. 4).

To be conservative in accounting for effects of beetle DNA on quantification we grouped samples with ≤1 × 10³ spores in one category. Spore loads within a given species varied across five orders of magnitude. All species contained a relatively high number of specimens without detectable spores. The lowest detectable spore load was estimated to be 10–1000 spores per insect. The highest estimates of the number

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**Fig. 2.** Polymerase chain reaction amplification of a 230-bp DNA fragment using the primers LEPTO1 and LEPTO2 from fungal colonies (lanes 3–9) and bark beetles (lanes 13–23). Lane 1, 100-bp DNA ladder (Promega); 2, water negative control; 3, *L. reconditum*; 4, *L. abietinum*; 5, *O. piceaparum*; 6, *L. lundbergii*; 7, *L. procerum*; 8, *L. terebrantis*; 9, water negative control; 10, 11, DNA ladder; 12, water negative control; 13, DNA extracted from *Hylurgops porosus* (20 beetles pooled); 14, DNA extracted from *Hylurgops porosus* (one beetle); 15, DNA extracted from *Hylurgops porosus* (five beetles pooled); 16, DNA extracted from *Hylurgops porosus* (one beetle); 17, DNA extracted from *Hylurgops porosus* (20 beetles pooled); 18, DNA extracted from *Hylurgops subcostatus* (one beetle); 19, DNA extracted from *Dendroctonus valens* (one beetle); 20, DNA extracted from *Hylurgops porosus* (one beetle); 21, DNA extracted from *Hylurgops porosus* (one beetle); 22, DNA extracted from *Hylurgops subcostatus* (one beetle); 23, DNA extracted from *D. valens* (one beetle); 24, water negative control; 25, DNA ladder.
Fig. 3. Standard curves for quantifying L. wageneri DNA using real-time polymerase chain reaction with SYBR Green® as the fluorescent dye. Threshold cycles (Ct), corresponding to the increase of template DNA above background level, were plotted against the log of genomic DNA extracted from spore suspensions in the range from $1 \times 10^2$ to $1 \times 10^6$ spores/100 µL.

of spores were in the range of $1 \times 10^5$ to $1 \times 10^6$ spores per individual beetle. However, this high association rate was found only for two specimens of Hylastes macer (2001 and 2002) and for one specimen of each Hylurgops subcostulatus (2002) and Hylastes gracilis (2001). In most cases, the number of spores per individual beetle was in the range of $1 \times 10^1$ to $1 \times 10^4$. In both years total spore load was highest for Hylastes macer.

Correlation between detected amounts of L. wageneri DNA and size of beetles

The surface area of individual adult bark beetles varied greatly among the different species, with D. valens being the largest, followed by the medium-sized species Hylastes macer, Hylurgops porosus, and Hylurgops subcostulatus, and the smallest species I. pini and Hylastes gracilis (Table 3). The relative size of these three beetle groups corresponded roughly to 7:2:1. The bark beetle most often associated with fungal DNA, Hylastes macer, was the second largest of these species. The two other species of similar size, Hylurgops porosus and Hylurgops subcostulatus, showed an intermediate association with L. wageneri DNA. Dendroctonus valens, on the other hand, was the species among the four captured in both 2001 and 2002 with the lowest association rate with fungal DNA. The association rate with fungal DNA for Hylastes gracilis was very similar to that for D. valens. Ips pini, the smallest of all bark beetles, had the lowest association rate.

Discussion

Previous evidence regarding the role played by insects in the epidemiology of BSRD

Vectoring of fungal inoculum by bark beetles is thought to play a crucial role in the epidemiology of BSRD (Hansen 1997; Wood et al. 2003). Detection and quantification of L. wageneri by incubating dead bark beetles on artificial media is complicated by its slow growth, causing inhibition and overgrowth by faster growing filamentous fungi, yeasts, and other ophiostomatoid species (Zhou et al. 2001), even when cycloheximide is added to the medium (W. Schweigkofler, unpublished data). Witcosky (1985) presented low association rates of Verticiladiella wageneri (= L. wageneri) with beetles in Douglas-fir stands in Oregon where the fungus was isolated from less than 5% of 668 Hylastes nigrinus, Stereumius carinatus, and Pissodes fasicatus beetles. Even given the limited success in isolating this fungus from Douglas-fir-associated bark beetles, processing large numbers of insect samples can be very time-consuming and laborious. Rapid detection of ophiostomatoid fungi from infected wood using PCR was reported previously (Kim et al. 1999), but to our knowledge molecular techniques have not been used yet for the detection of Leptographium from associated insects in ponderosa pine.

Species detected by method described and potential for cross-reactivity

The primers LEPTO1 and LEPTO2 amplified a fragment from L. wageneri and L. reconditum as was expected based on published sequence information (Jacobs et al. 2001). No sequences were available previously for L. neomexicanum and L. serpens, which have an identical primer binding site based on our results and newly available sequence data (Fig. 1). However, geographic distribution and (or) host range of these species differ widely from those of L. wageneri. Leptographium reconditum was isolated exclusively from the rhizosphere of monocots (Triticum spp. and Zea mays) in South Africa, and no association with insects is known (Jooste 1978). Also L. neomexicanum is not known to be vectored by bark beetles and has only been reported from New Mexico (Jacobs and Wingfield 2001). Leptographium serpens occurs mainly in Europe and South Africa, with very few observations from North America (Jacobs and Wingfield 2001). On the other hand, primers LEPTO1 and LEPTO2 do not amplify DNA of Leptographium terebrantis, a pathogenic species that is widespread in western North America, attacks a wide range of conifers (e.g., P. ponderosa), and is associated with several bark beetle species (Jacobs and Wingfield 2001). To further confirm the identity of the PCR fragments, we sequenced representative samples from every bark beetle species and found them to be identical with published L. wageneri sequences.

Leptographium wageneri occurs in three varieties that are distinguishable mainly by host specificity. In addition, isozyme analysis (Oetrosina and Cobb 1987; Zambino and Harrington 1992) and DNA fingerprinting using random amplified polymorphic DNA – PCR (Withuhn et al. 1997) showed differences among the varieties. Minor morphological differences are also present but difficult to observe for nonspecialists. Although our study focused exclusively on the single variety known to infect P. ponderosa, the ribosomal gene region we used for developing the primers LEPTO1 and LEPTO2 is identical for the three varieties (Jacobs et al. 2001). We conclude therefore that the molecular marker used in the present study is well suited for ecological studies on the association
between all three varieties of the black stain pathogen *L. wageneri* and associated bark beetles.

**Insects trapped in our study: differences and similarities amongst them**

Of the nine bark beetle species we collected, only four species (*Hylurgops porosus*, *Hylurgops subcostulatus*, *Hylastes macer*, and *D. valens*) were sampled in both 2001 and 2002. Extreme fluctuations of bark beetle densities are well known and can reach up to four orders of magnitude within 5 years (Turchin et al. 1991; Reeve et al. 1995). The bark beetles *Hylastes gracilis*, *Hylastes longicollis*, and *Hylastes macer* infest a range of pine species in the western United States, among them ponderosa pine, sugar pine (*Pinus lambertiana* Dougl.), Jeffrey pine, and lodgepole pine (Wood et al. 2003). The broods develop under the bark of the root crown and in the roots of dying and dead trees, in stumps and in the underside of logs, and in slash in contact with the ground (Furniss and Carolin 1977; Wood 1982). *Hylurgops* spp. also attack the phloem of the root collar, the main roots of

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**Fig. 4.** Quantification of *Leptographium* spores from bark beetles using real-time polymerase chain reaction. *Hylastes macer* (*a*), *Hylurgops porosus* (*b*), *Hylurgops subcostulatus* (*c*), and *Dendroctonus valens* (*d*) were collected both in 2001 (black bars) and 2002 (grey bars). *Ipis pini*, *Hylastes gracilis*, *Hylastes longicollis*, and *Hylastes reticulatus* (three beetles) were collected only in 2001.
dying or dead trees, or freshly cut stumps (Furniss and Carolin 1977; Wood 1982). In western North America, they can be found on most pine species and some fir species.

*Dendroctonus valens* is the largest and most widely distributed bark beetle in North America and has been reported from more than 40 conifer species. Vertical galleries are excavated in the phloem of the lower trunk and to a lesser extent in larger roots (Wood et al. 2003). This species usually attacks trees of reduced vigor or those infested with other bark beetles, but it also can attack apparently healthy trees (Smith 1971).

*Ips pini* attacks many conifers, including ponderosa, Jeffrey, and lodgepole pines. This beetle species infests mainly saplings and pole-sized trees, logging slash, windthrown trees, or trees broken by wind or snow (Kegley et al. 1997). Females construct tunnels or “egg galleries” in the phloem layer, slightly broken by wind or snow (Kegley et al. 1997). Reports on the ecology of *Hylastes tenuis* and *Hylurgops reticulatus* are scant (Hanula et al. 1998; Kelsey and Joseph 2003; Miller et al. 1986).

The bark beetle *Hylastes nigrinus* (Mannerheim) and the weevils *Pissodes fasciatus* (LeConte) and *Stererninus carinatus* (Boheman) (Coleoptera: Curculionidae) were shown to carry *L. wageneri* inoculum in Douglas-fir stands in the Pacific Northwest (Witcosky and Hansen 1985) and to transmit *L. wageneri* to seedlings under laboratory conditions (Witcosky 1985) but were not collected in our study.

**Mitotic spores as the main inoculum for BSRD**

Sexual recombination and the development of ascospores are known to play a crucial role in the epidemiology of other ophiostomatoid fungi, e.g., *Ophiostoma novo-ulmi*, the causal agent of Dutch elm disease (Brasier 2001). In the case of *L. wageneri* var. *ponderosum*, a sexual stage (= *O. wageneri*) was detected only once from ponderosa pines (Goheen and Cobb 1978). Sexual recombination may be a rare event in this pathogen (Otrosina and Cobb 1987), and we assume that the inoculum carried by the bark beetle consists predominantly of asexual conidia.

Contamination of insects with fungal spores could occur when different species of bark beetles are captured in the same trap. This could theoretically be the case for *D. valens*, *I. pini*, *Hylastes gracilis*, and *Hylastes longicollis*, which were found to be less often associated with *L. wageneri* DNA than were *Hylurgops porosus*, *Hylurgops subcostulatus*, and *Hylastes macer*. However, at least one sample of each of the former four species was found to carry *L. wageneri* DNA when no other PCR-positive insect sample was present in the same trap. This indicates that cross-contamination in the trap from other insects is unlikely.

**Table 3. Size of bark beetle species used for the detection of Leptographium DNA.**

<table>
<thead>
<tr>
<th>Species</th>
<th>Length (mm)</th>
<th>Diameter (mm)</th>
<th>Volume (mm³)</th>
<th>Surface (mm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Dendroctonus valens</em></td>
<td>7.68±1.38</td>
<td>3.1±0.9</td>
<td>52.1±14.3</td>
<td>84.5±16.7</td>
</tr>
<tr>
<td><em>Hylastes gracilis</em></td>
<td>4.0±0.0</td>
<td>1.0</td>
<td>3.1±0.0</td>
<td>14.1±0.0</td>
</tr>
<tr>
<td><em>Hylastes macer</em></td>
<td>5.78±0.72</td>
<td>1.5</td>
<td>10.2±1.4</td>
<td>30.8±3.7</td>
</tr>
<tr>
<td><em>Hylurgops porosus</em></td>
<td>4.98±1.02</td>
<td>1.5</td>
<td>8.8±1.8</td>
<td>27.0±4.8</td>
</tr>
<tr>
<td><em>Hylurgops subcostulatus</em></td>
<td>4.28±0.72</td>
<td>1.5</td>
<td>7.6±1.2</td>
<td>23.7±3.4</td>
</tr>
<tr>
<td><em>Ips pini</em></td>
<td>2.88±0.38</td>
<td>1.0</td>
<td>2.3±0.7</td>
<td>10.6±2.7</td>
</tr>
</tbody>
</table>

**Note:** Values are means ± standard deviations.

1 Twenty-five specimens were measured for each species (*Hylastes gracilis*: three specimens).

2 Standard deviations were not determined for the diameters of the very small and uniform bark beetle species.

Inoculum phoresy determined first by ecology of insect species and second by size of insect

The presumed physical attachment of *L. wageneri* spores to the exoskeleton of bark beetles could suggest a correlation between the size of the beetles and the number of spores they carry. Our study shows that some insect genera are more effective carriers than others (e.g., *Hylastes > Hylurgops > Dendroctonus > Ips*), independently of insect size; for example, *D. valens*, the largest species, showed only a low association with *L. wageneri* DNA. The insect species from which *L. wageneri* DNA was found most consistently (*Hylastes macer*, *Hylurgops porosus*, and *Hylurgops subcostulatus*) are of intermediate size. In the Dutch elm disease pathosystem, low inoculum numbers of *O. novo-ulmi* were isolated from the two smaller bark beetle species *Scolytus kirschi* and *Scolytus multistratus*, whereas the larger beetle *Scolytus scolytus* was associated with a higher number of fungal spores (Webber et al. 1987; Webber 1990, 2000).

It is not surprising that different insect species may vary in their ability to carry inoculum. Differential niche colonization and reproductive strategies may be one of the factors explaining such variations among groups of insects. For instance, the location of the pupal chamber is of crucial importance for the amount of fungal inoculum per bark beetle (Webber 2000). Because of the total removal of surface tissue during insect metamorphosis, conidia that were attached to the larvae will not be present at the surface of the imago. Consequently, all fungal spores carried by the vector must attach to the imago during emergence from the pupal chamber to the bark surface. Therefore, beetles pupating in inner bark chambers of the lower trunk or in roots (e.g., *Hylastes macer*) are more likely to carry a high spore load than beetles pupating in the outer bark.

Quantification of spore loads and putative classification of insects as carriers based on spore load

In the Dutch elm disease pathosystem, Webber (1990) estimated the spore load for *S. scolytus* to be in the range of 1 to 350 000 spores per beetle, for *S. multistratus* 1 to 30 000 spores, and *S. kirschi* had less than 300 spores per beetle. *Scolytus scolytus* is similar in size to *Hylastes macer* (mean length: 5.4 ± 0.3 mm; Webber 1990); the size of *S. kirschi* (2.5 ± 0.3 mm) corresponds roughly to that of *I. pini*. Our
quantification of L. wageneri spores using real-time PCR resulted in similar estimates of the number of spores (Fig. 4). For example, we detected between 0 to several hundred thousand spores on Hylastes macer, Hylurgops subcostatus, and Hylastes gracilis, but only a few specimens yielded more than 10,000 spores. On average, the spore load on single beetles was highest for Hylastes macer in both years. Hylurgops porosus was associated with only a moderate number of spores, but could still contribute significantly to L. wageneri inoculum dispersal because of high insect numbers in both years. The number of spores on D. valens and L. pini was, with the exception of a single D. valens individual, below 1000. This is consistent with their known ecological niche, making them unlikely to vector L. wageneri var. ponderosum.

In the case of O. novo-ulmi, a minimum of 1 × 10^5 conidia is required for infection of Ulmus procera. Infection was not always assured even with an inoculum load of 5 × 10^6 conidia (Webber 1987; Webber and Gibbs 1989). If a similar inoculum threshold would be necessary for the infection of ponderosa pines with L. wageneri, then only a small percentage of the insects in our study would act as effective vectors. A single conidiophore of L. wageneri contains, on average, ca. 1 × 10^9 conidia in a slimy droplet. In pure culture up to 200/cm^2 conidial droplets were produced. Therefore it is highly probable that a beetle may touch more than one droplet when emerging from its pupal chamber to the tree surface. However, not all conidia will stick to the exoskeleton, and some might be lost during the flight of the beetle to new hosts (e.g., because of UV inactivation).

Our results substantiate the hypothesis that Hylastes macer might be the most important vector for the long-distance dispersal of L. wageneri, although other root feeding bark beetle species such as Hylurgops porosus might be involved in the disease cycle. The DNA techniques we report provide a new tool to study the epidemiology of black-stain root disease. Understanding the interactions among inoculum potential, beetle ecology, host tree physiology, and site edaphic factors as they relate to disease susceptibility and severity is crucial to minimizing disease impact.

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