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Geosmithia – a widespread, abundant, but long-time ignored symbionts of subcortical insects

by

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HABILITATION THESIS

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1.2 OUTLINE OF THE THESIS AND SUMMARY OF THE RESULTS

The work is a continuation of my PhD thesis, submitted in 2006, which dealt with the study of the host spectrum, geography and taxonomy of the genus *Geosmithia*. At that time, the association of *Geosmithia* with subcortical insects (mostly bark beetles), was very little known and mostly considered incidental, not specific, similar to many of the moulds we find in bark beetle galleries. At the time of PhD thesis origin, only the species *G. putterillii* and *G. lavendula* were recognized, isolated from various non-specific substrates such as soil or cereals. This, together with the absence of slime spores, a typical entomochoric adaptation, made the mycologists sceptical to the importance of this association. The thesis gave an overview of the composition of the communities of the genus *Geosmithia* on bark beetles in temperate Europe and showed that these fungi make communities that are host-specific and considerably stable over

large geographical distances. This pattern indicates entomochory and long-term association. The work was published in 2008 (Kolařík et al., 2008) and became a baseline study for the symbiosis of *Geosmithia* with bark beetles. The PhD thesis also published a taxonomic revision, which led to the revision of old taxa (*Penicillium pallidum*, *G. putterillii*) and the description of four new species (Kolařík et al., 2005; Kolařík et al., 2004). These results raised a number of additional questions that constitute the Aims of the habilitation thesis.

1.2.1 Aim 1. To assess Geosmithia host range and community pattern at the global scale

The aim was to expand sampling in terms of number of vectors and geography.

Outputs. Geosmithia communities were studied on diverse insect hosts in Central Europe (Jankowiak and Kolarik, 2010; Jankowiak et al., 2014; Kolařík and Jankowiak, 2013; Pepori et al., 2015; Strzałka et al., 2021), Mediterranean basin (Kolařík et al., 2007), Western (Kolařík et al., 2017; Pietsch et al., 2022) and Eastern USA (Huang et al., 2017; Huang et al., 2019), Costa Rica (Kolařík and Kirkendall, 2010) and China (Zhang et al., 2022). The survey showed that Geosmithia is worldwide distributed on numerous bark beetles and other phloem- and wood boring insects. Surprisingly, Geosmithia were found also as primary ambrosia fungi living in obligate symbiosis with ambrosia beetles (Kolařík and Kirkendall, 2010). The vectors differed in the frequency of association with Geosmithia and can be roughly separated to regular and moderate vectors, and to insect species that have very accidental or do not have an association with Geosmithia. The degree of association with Geosmithia across vectors is determined by the combination of organ preference and host tree species evidently primarily shaped by the substrate quality type, mainly correlated with host tree mass and bark thickness (see chapter Host range and strength of the association with bark beetles). Based on the host range, Geosmithia can be divided into generalist species that are very common across many vectors, and to specialist species that occur on vectors inhabiting? host plants of the same plant family. Consequently, the host preference influences the Geosmithia community composition of the particular beetle species. It is obviously shaped by the degree of spatial isolation of given bark beetle species in a given area. If there are only host-specific species on a given host plant, more sharply delimited communities are formed. If there are more polyphagous vectors, the differences are erased. See chapters Vector specificity, community composition and Biogeography.

1.2.2 Aim 2. To evaluate diversity and describe newly found *Geosmithia* species

The first study in Central Europe (Kolařík et al., 2008), showed that most of the presented *Geosmithia* strains cannot be ascribed to the known species. Such a large proportion of unknown diversity called for its study and for a formal description of the species found, which is a necessary step for their next study.

Outputs. The global survey (Aim 1) revealed presence of more than 69 phylogenetic *Geosmithia* species. Seven of them belonged to the known taxa (*G. putterillii*, *G. pallida*, *G. flava*, *G. obscura*, *G. lavendula*, *G. fassatiae*, *G. langdonii*) and for the others, a serial number designation was introduced. In my study, I focused on the description of the most important species such as ambrosia species (*Geosmithia*

eupagioceri, G. microcorthyli, G. rufescens, C. cnesini) (Kolařík et al., 2015; Kolařík and Kirkendall, 2010), phytopathogenic G. morbida (Kolařík et al., 2011), G. ulmacea and G. omnicola, both species frequently associated with Dutch elm disease (Pepori et al., 2015), G. proliferans and G. brunea (Huang et al., 2017), G. fagi, G. longistipitata and G. pazoutovae (Strzałka et al., 2021). I collaborated on the description of nine other species, some of which belong to species isolated also from my collections, i.e. Geosmithia sp. 2 - G. pumila, Geosmithia sp. 3 - G. pulverea, Geosmithia sp. 20 - G. granulata (Zhang et al., 2022), Geosmithia sp. 5 - G. funiculosa (Crous et al., 2022), and some were newly described based on new collections of the colleagues from China - G. luteobrunnea, G. radiata, G. brevistipitata, G. bombycina, G. subfulva and G. fusca (Zhang et al., 2022) (see chapter Taxonomy and Diversity). Some species common in particular areas (e.g. G. sp. 41 in North America, G. funiculosa in Europe), are absent in others, showing clear biogeographical pattern. Thus, study of unexplored areas and vectors has a great chance to substantially increase our knowledge about Geosmithia diversity. See chapter Vector specificity, community composition and Biogeography.

1.2.3 Aim 3. To evaluate the biotechnological potential

Fungi from the order Hypocreales are known producers of various secondary metabolites, but nothing was known in case of *Geosmithia* at the beginning of my research. *Geosmithia spp.* are easily cultivable with a rapid growth, which makes them attractive also for biotechnological purposes. Another advantage is that these fungi are regularly in contact with bark beetles, and thus should have minimal cytotoxicity to animal cells, but still have to compete with co-occurring mites, nematodes, fungi and bacteria. This makes them good target for bioprospecting related with new drug and pesticide discovery. In addition to the biotechnological significance, information on the biological activity of extrolytes is interesting from an ecological point of view, helping us to understand interactions between the fungus and other members of the bark beetle holobiont.

Outputs. Our study on crude extracts shows the huge potential of antibacterial and antifungal activity across the whole genus (Veselská et al., 2019). From the ecological point of view, *Geosmithia* species in the bark beetle galleries have ability to compete with various moulds and bacteria, including insect-associated fungi. Prominent yellow, orange, and red pigments produced by *G. lavendula* and other species were identified as set of anthraquinones, often novel to science, several of them with antibacterial or anti-inflammatory activity (Stodůlková et al., 2009; Stodůlková et al., 2010) and with a potential as highly persistent textile dyes or mordants (Flieger et al., 2009). We also developed a UPLC-MS based method for the separation of *Geosmithia* secondary metabolites (Tylová et al. 2011). See chapter Secondary metabolite production and biotechnological potential.

During the study of the violet-coloured G. *lavendula*, I came across another so far neglected bark beetle symbiont, *Quambalaria cyanescens*. This fungus belongs to a very little explored fungal lineage of Basidiomycota (Exobasidiomycetes, Microstromatales), again with missing knowledge about its secondary

metabolite production. A subsequent study showed that it is a producer of various naphthoquinones, together with newly described quambalarine A-D, which have antimicrobial potential and negligible cytotoxicity to healthy human cells (Prochazkova et al., 2020; Stodůlková et al., 2015; Stodulkova et al., 2008). In addition, they have antiviral activity and selective toxicity to the human carcinoma cells which stimulated a subsequent exploration of its mode of action. That outputs are not part of the Thesis and are summarized in several studies (Grobárová et al., 2016; Matoušková, 2020; Vališ et al., 2017; Zima et al., 2020).

1.2.4 Aim 4. To understand *Geosmithia* ecology. How do they interact with bark beetles or host trees?

There are several types of interactions known between bark beetles and their associated fungi. It can therefore be expected that some of these interactions will also occur in the genus *Geosmithia*.

Outputs. Bark beetle associated fungi are known to have diverse symbiotic (i.e. mutualistic, neutral or antagonistic) interactions with its environment. The most straightforward is the ability to invade healthy plant tissues, exploit these protected nutrient sources by the fungus, which enables to increase it's the beetle's fitness. Studied strains typically show no signs of phytopathogenicity (Jankowiak and Kolarik, 2010; Strzałka et al., 2021). Mild, but significant lesions were found in case of two strains by Li et al. (2022). The only undisputed case where the fungus makes significant necrosis, is the pathogenic complex of walnut twig beetle, *Pityophthorus juglandis/ G. morbida* responsible for the Thousand Cancers Disease of black walnut, *Juglans nigra* (Kolařík et al., 2011). The massive dieback of black walnut has promoted *Geosmithia* research worldwide, and is behind the wider recognition of *Geosmithia* as entomochoric and symbiotic fungi. The *G. mobida* related research, to which I furher contributed, involved study of its genetic variability (Hadziabdic et al., 2014a). Walnut twig beetle transmits other *Geosmithia* species (Kolařík et al., 2017), and as can be seen from preliminary results with infection experiments with *G. obscura* (Pietsch et al., 2022), it needs to be studied whether these species can contribute to necrosis formation. See chapter Phytopathogenic potential and TCD.

1.2.5 Aim 5. To understand *Geosmithia* evolution and biology

As we accumulated knowledge about the host range and pathogenicity, it became obvious that the genus comprised species with variable life strategies, host specificity, degree of affinity to the host beetle and pathogenicity to the host plant. We thus decided to study evolution of the genus by incorporating information about phylogeny, genome size and various phenotype traits (morphology, enzymatic capacity, antibiosis etc.).

Outputs. The study showed the limits of rDNA for robust phylogenetic hypotheses testing, for genus *Geosmithia*. The observed incongruence between rDNA and protein coding genes was attributable to GC content and heterotachy-based artifacts (Kolařík et al., 2017; Veselská et al., 2019). That seems to be a consequence of mechanisms such as the fluctuations in the effective population change, bottlenecks,

usually related to the life history changes, especially those related to the switch between free living style to host-associated life strategy (Kolařík and Vohník, 2017; Kolařík et al., 2021) (papers not included into the Thesis). The study of the evolution of the genus showed that the ancestral species were generalists and later specialized in several lineages to their host vectors. All that shifts were accompanied by loss of metabolic capacity and genome size inflation. We identified three independent origins of primary ambrosia fungi (Kolařík et al., 2015; Kolařík and Kirkendall, 2010), which was accompanied by the cell and genome size inflation and production of particular fatty acids (Veselská and Kolařík, 2015; Veselská et al., 2019). One lineage, G. morbida, became plant pathogen, with the unique feature to digest all components of lignocellulose, what feature can be supposed as G. morbida virulence factor (Veselská and Kolařík, 2015; Veselská et al., 2019). The genome size and DNA content of the cells were measured by flow cytometry. This method has so far been little used in fungi, mainly due to their very small genomes. Therefore, the method needed to be optimized and suitable standards for determining genome size were sought. This led to the first ever methodological work on flow cytometry in mycology (Veselská et al., 2014). The study of genus evolution and the use of flow cytometry was the subject of Tereza Veselská's master and doctoral thesis (see Student's theses related to the topic of the habilitation thesis). My study further contributed to knowledge about hydrophobins, which showed that ability to adhesion is important in the evolution of the genus (Frascella et al., 2014). Beside of them, there is a good evidence that Geosmithia obtained hyrophobin, cerato-ulmin, by the horizontal transfer from *Ophiostoma novo-ulmi* (Bettini et al., 2014). See chapter Evolution and biology.

1.3 PAPERS INCLUDED IN THE THESIS

(the most important studies are underlined)

- <u>Kolařík M, Kostovčík, M Pažoutová S (2007)</u> Host range and diversity of the genus *Geosmithia* (: Hypocreales) living in association with bark beetles in the Mediterranean area. *Mycological Research* 101: 1298-1310.
- 2. Stodůlková E, **Kolařík M,** Křesinová Z, Kuzma M, Šulc M, Man P, Novák P, Maršík, P, Landa, P, Olšovská, J, Chudíčková M, Pažoutová S, Černý J, Bella J, Flieger M (2009) Hydroxylated anthraquinones produced by *Geosmithia* species. *Folia Microbiologica* 54: 179-187.
- 3. Jankowiak R, **Kolařík M** (2010) Fungi associated with the fir bark beetle *Cryphalus piceae* in Poland. *Forest Pathology* 40: 133-144.
- 4. **Kolařík M,** Kirkendall LR (2010) Evidence for a new lineage of primary ambrosia fungi in *Geosmithia* Pitt (Ascomycota: Hypocreales). *Fungal Biology* 114: 676-689.
- 5. Stodůlková E, Man P, **Kolařík M**, Flieger M (2010) High-performance liquid chromatography-off line mass spectrometry analysis of anthraquinones produced by *Geosmithia lavendula*. *Journal of Chromatography A* 1217: 6296-6302.

- 6. **Kolařík M,** Freeland E, Utley C, Tisserat N (2011) Geosmithia morbida sp. nov. a new phytopathogenic species living in symbiosis with the walnut twig beetle (*Pityophthorus juglandis*) on *Juglans* in the USA. *Mycologia* 103: 325-32.
- 7. Tylová T, **Kolařík M**, Olšovská J (2011) The UHPLC-DAD fingerprinting method for analysis of extracellular metabolites of fungi of the genus *Geosmithia* (Acomycota: Hypocreales). *Analytical and Bioanalytical Chemistry* 400: 2943-2952.
- 8. <u>Kolařík M, Jankowiak R (2013)</u> Vector affinity and diversity of *Geosmithia* fungi living on subcortical insects inhabiting Pinaceae species in Central and Northeastern Europe. *Microbial Ecology* 66: 682-700.
- 9. Hadziabdic D, Vito L, Windham M, Pscheidt J, Trigiano R, **Kolařík M** (2014) Genetic differentiation and spatial structure of *Geosmithia morbida*, the causal agent of thousand cankers disease in black walnut (*Juglans nigra*). *Curr Genet* 60: 75–87.
- 10. Bettini PP, Frascella A, Kolařík M, Comparini C, Pepori AL, Santini A, Scala F, Scala A (2014) Widespread horizontal transfer of the cerato-ulmin gene between *Ophiostoma novo-ulmi* and *Geosmithia* species. *Fungal Biology* 118: 663-674.
- 11. <u>Jankowiak R, **Kolařík M**, Bilańskic P</u> (2014) Association of *Geosmithia* fungi (Ascomycota: Hypocreales) with pine- and spruce-infesting bark beetles in Poland. *Fungal Ecology* 11: 71–79.
- 12. Veselská T, Svoboda J, Růžičková Z, **Kolařík M** (2014) Application of flow cytometry for genome size determination in *Geosmithia* fungi: A comparison of methods. *Cytometry Part A*. 85: 4–861.
- 13. Frascella A, Bettini PP, **Kolařík M**, Comparini C, Pazzagli L, Luti S, Scala F, Scala A (2014) Interspecific variability of class II hydrophobin GEO1 in the genus *Geosmithia*. *Fungal Biology* 118: 862-871.
- 14. Veselská T, Kolařík M (2015) Application of flow cytometry for exploring the evolution of Geosmithia fungi living in association with bark beetles: the role of conidial DNA content. Fungal Ecology 13: 83-92.
- 15. Stodůlková E, Císařová I, **Kolařík M**, Chudíčková M, Novák P, et al. (2015) Biologically active metabolites produced by the basidiomycete *Quambalaria cyanescens*. *PLoS ONE* 10(2): e0118913.
- 16. Pepori AL, **Kolařík M**, Bettini PP, Vettraino AM, Santini A (2015) Morphological and molecular characterisation of *Geosmithia* species on European elms. *Fungal Biology* 119:1063-1074.
- 17. **Kolařík M**, Hulcr J, Kirkendall LR. (2015) New species of *Geosmithia* and *Graphium* associated with ambrosia beetles in Costa Rica. *Czech Mycology* 67: 29-35.

- 18. Kolařík M, Hulcr J, Tisserat N, De Beer W, Kostovčík M, Kolaříková Z, Seybold SJ, Rizzo DM (2017) *Geosmithia* associated with bark beetles and woodborers in the western USA: taxonomic diversity and vector specificity. *Mycologia* 109:185-199
- 19. Huang Y-T, **Kolařík M**, Kasson M, Hulcr J (2018) Two new *Geosmithia* species in *G. pallida* species complex from bark beetles in eastern USA. *Mycologia* 109: 790-803.
- 20. <u>Huang Y.-T, Skelton J, Johnson AJ, **Kolařík M**, Hulcr J (2019) *Geosmithia* species in southeastern USA and their affinity to beetle vectors and tree hosts. *Fungal Ecology* 39: 168-183.</u>
- Veselská T, Skelton J, Kostovčík M, Hulcr J, Baldrian P, Chudíčková M, Cajthaml T, Vojtová T, Garcia-Fraile P, Kolařík M (2019) Adaptive traits of bark and ambrosia beetle-associated fungi. Fungal Ecology 41:165-176.
- 22. Procházková E, Kucherak O, Stodůlková E, Tošner Z, Císařová I, Flieger M, **Kolařík M,** Baszczyňski O (2021) NMR structure elucidation of naphthoquinones from *Quambalaria* cyanescens. Journal of Natural Products 84: 46–55.
- 23. <u>Strzałka B, **Kolařík M**, Jankowiak R (2021) *Geosmithia* associated with hardwood-infesting bark and ambrosia beetles, with the description of three new species from Poland. *Antonie van Leeuwenhoek* 114: 169-194.</u>
- 24. Li Y, Bateman C, Skelton J, Wang B, Black A, Huang Y-T, Gonzalez A, Jusino MA, Nolen ZJ, Freeman S, Mendel Z, Kolařík, M., Knížek, M, Park J-H, Sittichaya W, Pham T-H, Ito S, Torii M, Gao L, Johnson AJ, Lu M, Sun J, Zhang Z, Adams DC, Hulcr J (2022) Preinvasion assessment of exotic bark beetle-vectored fungi to detect tree-killing pathogens. *Phytopathology* 112: 261-270.
- 25. Zhang X, Li Y, Dai M, Si H, Zhao G, **Kolařík M**, Hulcr J, Jiang X, Chang R (2022) *Geosmithia* species associated with bark beetles from Southern China, with the description of four new Species. *Frontiers in Microbiology* 13: 10.3389/fmicb.2022.820402
- 26. Pietsch GM, Gazis R, Klingeman WE, Huff ML, Staton ME, Kolařík M, Hadziabdic D (2022) Characterization and microsatellite marker development for a common bark and ambrosia beetle associate, *Geosmithia obscura*. *MicrobiologyOpen* 11:e1286
- 27. Crous PW, Boers J, Holdom D, Steinrucken T, Tan Y, Vitelli J, Shivas R, Barrett M, Boxshall A-G, Broadbridge J et al. (2022) Fungal Planet description sheets: 1383–1435. Persoonia-Molecular Phylogeny and Evolution of Fungi 48: 261-371

1.4 STUDENT'S THESES RELATED TO THE TOPIC

In the course of the *Geosmithia* research, six students had the opportunity to work on attractive and unexplored topics what resulted in ten Theses defended at three departments of Faculty or Sciences at

Charles University and University of Chemistry and Technology, Prague. The list of theses, elaborated with my contribution (supervisor, official or non-official consultant), is provided below.

- Veselská T (2022) Comparative ecophysiology as a tool for the study of adaptive traits of fungal symbionts and pathogens. Doctoral thesis (PhD. degree). Department of botany, Faculty of Sciences, Charles University. Supervisor: M. Kolařík
- 2. Lovás D (2022) Identifikace bioaktivních metabolitů hub asociovaných s kůrovci. Bachelor Thesis (Bc. Degree). University of Chemistry and Technology, Prague, Department of Biotechnology. Bachelor thesis (Bc. degree). Supervisor prof. Ing. Jan Masák, CSc.
- 3. Fabryová A (2016) Study of culturable anaerobic bacterial communities living in symbiosis with bark beetles; its isolation, taxonomy and biotechnical potential. Diploma thesis (MSc. degree). Department of genetics and microbiology, Faculty of Sciences, Charles University. Supervisor: MSc. Paula García Fraile, Ph.D.
- 4. Veselská T (2013) Evoluční ekologie rodu *Geosmithia*. Diploma thesis (MSc. degree). Department of botany, Faculty of Sciences, Charles University. Supervisor: M. Kolařík
- 5. Musil K, (2013) Studium biologicky aktivních sekundárních metabolitů produkovaných vybraným kmenem hub rodu *Geosmithia* metodou UPLC-DAD-TOF-MS. Bachelor thesis (Bc. degree). Department of analytical chemistry and microbiology, Faculty of Sciences, Charles University. Supervisor: RNDr. Mgr. Jana Olšovská, Ph.D.
- 6. Tylová T (2013) Metody kapalinové chromatografie pro analýzu biologicky aktivních mikrobiálních sekundárních metabolitů. Doctoral thesis (Ph.D. degree). Department of analytical chemistry and microbiology, Faculty of Sciences, Charles University. Supervisor: RNDr. Mgr. Jana Olšovská, Ph.D.
- Veselská T (2010) Genetika hub, evoluce genomu a vyuziti prutokoveho cytometru pri studiu DNA. Bachelor thesis (Bc. degree). Department of botany, Faculty of Sciences, Charles University. Supervisor: M. Kolařík
- 8. Křesinová Z (2007) Studium sekundárních metabolitů houby *Geosmithia lavendula*. Doctoral thesis (RNDr. degree). Department of analytical chemistry and microbiology, Faculty of Sciences, Charles University.
- 9. Křesinová Z (2007). Studium sekundárních metabolitů houby *Geosmithia lavendula*. Diploma thesis (MSc. degree). Department of analytical chemistry and microbiology, Faculty of Sciences, Charles University. Supervisor: prof. RNDr. Zuzana Bosáková, CSc.

10. Kostovčík M (2006) Molekulárne genetická a morfologická analýza komplexu *Geosmithia lavendula*. Diploma thesis (MSc. degree). Department of genetics and microbiology, Faculty of Sciences, Charles University. Supervisor: RNDr. Sylva Pažoutová, CSc.

1.5 EXTENDED REVIEW OF THE GEOSMITHIA TAXONOMY, GEOGRAPHY, DIVERSITY, ECOLOGY AND BIOTECHNOLOGICAL POTENTIAL

1.5.1 Introduction

Bark and ambrosia beetles (Coleoptera: Curculionidae: Scolytinae, Platypodinae) are associated with a diverse set of ecto- and endosymbionts, classified among the prokaryotes, filamentous fungi, yeasts, and microinvertebrates. Fugal symbionts are the most studied and their dependency on the insect vector ranges from obligatory, in strictly entomochoric fungi, to incidental, acquired from the environment. Fungal symbionts interact with the host insect and tree, forming mutualistic, commensal or antagonistic interactions (Beaver, 1989; Hofstetter et al., 2015; Six, 2013). The best studied fungal symbionts of bark beetles belong to ophiostomatoid fungi (Ascomycota: Ophiostomatales, Microascales). However, beetle galleries harbour many other fungal families equally frequently, but many of the non-ophiostomatalean have been historically ignored (Jankowiak and Kolarik, 2010; Kirschner, 2001; Kirschner et al., 2001; Kolařík et al., 2006). Filamentous fungi placed into the genus Geosmithia (Ascomycota: Hypocreales, Bionectriaceae) used to be sporadically reported as plant or soil saprobes (Pitt, 1979; Pitt and Hocking, 2009). The very first record of Geosmithia from the bark beetle niche, and a suggestion of its phytopathogenicity, was from the fir bark beetles in the USA by Wright (1938), but the fungus was misidentified as Spicaria anomala (Kolarik et al. 2017). The regular association of Geosmithia fungi with bark beetles was simultaneously discovered in Germany (Kirschner, 1998, 2001) and Czechia (Kubátová et al., 2004; Kubátová et al., 1999). During the first decade of the new millennium, the question of the tightness of the association of Geosmithia with bark beetles was not yet settled. The reasons for these doubts were numerous and relevant. The identified species, such as G. putterilllii have been known from various non-specific substrates such as soil or cereals (Kolařík et al., 2004; Pitt and Hocking, 2009). In addition, the generic concept of Geosmithia before 2012 included species of Hypocreales (Geosmithia in the current definition) but also Eurotiales, which have no connection to insects (Houbraken et al., 2012). Further, Geosmithia strongly resembles Penicillium, Paecilomyces or Mariannaea, which are common and widely ignored contaminants of bark beetle galleries. Geosmithia produces masses of dry spores, a typical feature of airborne fungi, but do not form slimy spores, a typical entomochoric adaptation. In addition, Geosmithia is typically found on hardwoods and conifers of the cypress family, associated with littlestudied secondary bark beetles of minor economic importance. Finally, Geosmithia is highly sensitive to cycloheximide, an antifungal agent often used in the study of ophiostomatalean fungi which are resistant to it. Thus, the Geosmithia presence has been frequently missed and there was a scepticism about the significance of the association.

In the following years, however, many independent studies confirmed *Geosmithia* as a stable, and often dominant symbiont of many bark beetles worldwide, forming fungal communities specific to the host trees frequented by the vector beetles. The subsequent discovery of a phytopathogenic species *G. morbida* (Kolařík et al., 2011), and also species living as primary ambrosia fungi (Kolařík and Kirkendall, 2010),

resulted in the recognition of *Geosmithia* as a genus containing regular bark beetle symbionts with apparent long-term coevolution. Here we summarize and interpret *Geosmithia* biology based on a review more than 140 publications (Fig. 1).

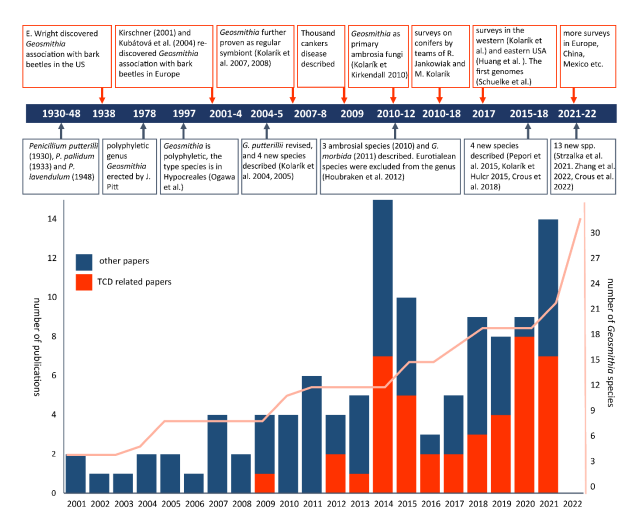


Fig. 1 Upper part. Chronology of important events related to *Geosmithia* taxonomy (below) and ecology, host range and biogeography (above). Lower part. Overview of publications on Hypocrealean *Geosmithia* species over the last 20 years (2001-2021), with a breakdown of papers focusing on Thousand Cankers Disease (orange bar) and on other aspects (blue bar). The graph is based on articles excerpted by the Scopus database and few other important papers. The chart does not include the numerous papers that focus primarily on the biology of the walnut twig beetle, a TCD vector. The graph also presents the increase of described species within the genus (pink line).

1.5.2 History of the genus definition and the main differentiation features

Like in other morpho-genera of anamorphic fungi, the *Geosmithia* generic concept has undergone a dramatic changes (Fig. 1). In the current concept, its characteristics include the following: absence of sexual state, the presence of many colony color but not green color (which diagnoses it as distinct from *Penicillium*), presence of cylindrical shape of phialides without prominent neck and with roughened walls, elliptical to cylindrical conidia produced in chains and presence of specific cellular initials and the conidiophore basis (Kolařík et al., 2004). The discovery of the morphologically unique ambrosia fungi

expands this morphological concept to also include solitary and globose conidia (Kolařík and Kirkendall, 2010). The colony color ranges from white to cream, to various shades of yellow, brown, rusty or red. *Geosmithia* produce the *Penicillium*-like type conidiophore, or conidiophores can be much more complex, irregularly and repeatedly branched. Besides macronematous conidiophores with enteroblastic phialides, microcronematous conidiophores can also be formed on aerial or substrate mycelium (Kolařík et al., 2004). Whereas *Penicillium*-like conidiophores produce columns of dry conidia, microcronematous conidiophores form holoblastic, solitary conidia in slimy droplets. This conidial type, referred to as substrate conidia, is another feature found in related genera such as *Gliocladium* and *Nalanthamala* (Schroers et al., 2005). Other typical *Geosmithia* feature is conidiophore basis, making so-called "peg foot" with smooth cell wall and curved shapes (Kolařík et al., 2004) (Fig. 2).

The type species, G. putterillii, was first described as a Penicillium putterillii and the history of Geosmithia is linked to the morphologically similar Penicillium genus. Species of Geosmithia was first aggregated into the series P. pallidum in Asymmetrica-Funiculosa section that was established for P. pallidum, P. putterillii, P. lavendulum (now in Geosmithia). P. namyslowskii (now in Penicillium, Eurotiales) (Raper and Thom, 1949). John Pitt (1979) proposed a new genus Geosmithia, named in hour of George Smith, to include species from P. pallidum series and some species nowadays classified in Eurotiales. Although at first the concept was not accepted by some authors (Ramirez, 1982; Stolk and Samson, 1986) it was soon solidified in taxonomic lists (Pitt and Samson, 1993; Pitt et al., 2000) and other authors begun to use the name Geosmithia for newly discovered species of similar morphology (Pitt and Hocking, 1985; Yaguchi et al., 1993; Yaguchi et al., 1994; Yaguchi et al., 2005). The first studies utilizing molecular data showed that, while some of the species originally placed in the genus, including the type species G. putterilli, belonged to Hypocreales, others were in fact within the Eurotiales (Iwamoto et al., 2002; Ogawa and Sugiyama, 2000; Ogawa et al., 1997; Peterson, 2000). An eventual revision resulted in the creation of the monophyletic Geosmithia within Hypocreales, and placed other species into the genera of Penicillium, Rasamsonia and Talaromyces within Eurotiales (Houbraken et al., 2012). These changes also affect the classification of Rasamsonia argillacea, a fungus of clinical importance (Giraud et al., 2013),

which is still sometimes incorrectly identified by the old name *Geosmithia argillacea* (Giordano et al., 2021).



Fig. 2 Morphological features of *Geosmithia*. **A** Colony morphology on MEA can range from brown (*G. funiculosa*), lilac (*G. carolii*), white to cream (*G. putterilli*) and yellowish (*Geosmithia* sp. 11). **B, C** Yeast-like stage is presented in some species during the initial growth phase. *Geosmithia carolii* on MEA, 1 day, 24 °C. **D** Oblong and catenate conidia of *G. carolii*. **E** Globose and multinucleate conidia of *G. eupagioceri* stained by propidium iodide and observed under confocal microscope. **F** Long conidial chains of *Geosmithia* sp. 8 CCF4528. **G** Solitary produced conidia of *G. microcorthyli*. **H** Substrate conidia of *G. carolii*. **I** Penicillate conidiophore in *G. putterillii*. **J** Complexly branched conidiophore of *G. eupagioceri*. **K** Simple conidiophore in *Geosmithia* sp. 31. Scale bars **B** 500 μm, **C-E**, **H, K** 10 μm, **I, J** 20 μm

1.5.3 Taxonomy and Diversity

The genus possess relatively high phylogenetic diversity, with over 67 phylogenetic species, from which 32 were formally described (Fig. 1, Table 1). Most of the remaining species have been studied to a degree that allows diagnosis to the species level, but they have not been described formally. These species are

informally identified by numbers. This numbering system originated in Kolařík et al. (2007; 2008) and species thus labeled are frequently used in literature (Table 1).

Before the year 2004, only two species, G. lavendula and G. putterillii (incl. its synonym, P. pallidum), were formally accepted. Later, G. putterillii was found to be a complex of three species, G. putterillii, G. pallida (it itself consisting of five phylogenetic species) and G. flava (Kolařík et al. 2004). Three other species, G. fassatiae, G. langdoni and G. obscura were described from bark beetles in Europe (Kolařík et al., 2005). A large survey of Geosmithia in Europe and Mediterranean basin recognised other 23 undescribed species marked as Geosmithia sp. 1-5, 8-13, 16, 19-31 (Kolařík and Jankowiak, 2013; Kolařík et al., 2007; Kolařík et al., 2008). Subsequent surveys in the USA revealed other 20 species, classified as Geosmithia sp. 32-48 (Huang et al., 2019; Kolařík et al., 2017) or described as G. morbida (Kolařík et al., 2011), G. proliferans and G. brunea (Huang et al., 2017). Recently, several numbered species were formally described: G. ulmacea (sp. 13) and G. omnicola (sp. 10) (Pepori et al., 2015), G. xerotolerans (sp. 21), G. carolliae (sp. 19) (Crous et al., 2018) and G. longistipitata (sp. 28) (Strzałka et al., 2021). Some of the previously recognised taxa (sp. 2 - G. pumila, sp. 3, 23 - G. pulverea, sp. 20 - G. granulata), and others newly found (G. luteobrunnea, G. radiata, G. brevistipitata, G. bombycina, G. subfulva and G. fusca) were described from China (Zhang et al., 2022) and Europe (G. cupressina, G. fagi and G. pazoutovae) (Meshram et al., 2022; Strzałka et al., 2021)). Four species, G. eupagioceri, G. microcorthyli, G. rufescens and C. cnesini were described from ambrosia beetles in Costa Rica (Kolařík et al., 2015; Kolařík and Kirkendall, 2010). Other five tentative and undescribed species were recognized during the surveys on bark beetles in South Africa and Israel (Dori-Bachash et al., 2015; Machingambi et al., 2014) or on other substrates (Deka and Jha, 2018; Sun et al., 2018) (Table 1). The species G. tibetensis (Wu et al., 2013), described from the soil in Tibet, may not be a true *Geosmithia*; no molecular data were provided and its morphology fits that of Eurotiales.

The methods used to characterize *Geosmithia* species follow those used in studies on the genus *Penicillium* and *Aspergillus*. The most commonly used culture substrates are two nutrient-rich media, Malt extract agar (MEA) and Czapek Yeast Autolysate Agar (CYA), and the basal medium Czapek Dox Agar (CZD), the combination of which provides good resolution between most species. Regarding the cultivation temperature, studying growth at 24-25 °C, optimal temperature for perhaps all species, and 37 °C, tolerated by few species only (e.g. *G. lavendula* and *G. morbida*) is used.

The ITS rDNA marker, commonly used to delimit species across fungi, is used to characterize *Geosmithia* species, but it has its limits, especially among closely related species. Therefore, alternative markers are needed for better resolution in some species complexes. Other commonly used markers include RNA polymerase II second largest subunit (RPB2, region defined by the primers fRPB2-5F/fRPB2-7R), β-tubulin gene (TUB2, primers T10/Bt2b) and translation elongation factor 1-α gene (TEF-1α) including the large exon part (primers EF1- 983F/EF1-2218R) and the intron area (EF1-728F/EF1-986R). The latter

shows by far the greatest variability among *Geosmithia* species (Strzałka et al., 2021). The discriminatory power of the alternative markers can be assessed by studying groups of species that are clearly distinguishable morphologically and ecologically, yet have identical ITS sequences, such as *G. microcorthyli* (Kolařík and Kirkendall, 2010), *G. longistipitata* (Strzałka et al., 2021), *Geosmithia* sp. 24 (Dori-Bachash et al., 2015), *Geosmithia* sp. 16 (Kolařík and Jankowiak, 2013) and *G. langdoni* species complexes (Kolařík et al., 2017).

1.5.4 Host range and strength of the association with bark beetles

Geosmithia species are most commonly isolated from the subcortical niche created by bark beetles. The materials which yield most colony forming units are the internal surfaces of galleries, particularly the pupal chambers, but also the surface of eggs, larvae, and adults, and the gallery detritus. Adults captured outside of galleries, prior to the gallery initiation or after emergence from pupation, also frequently yield Geosmithia cultures. Geosmithia are usually isolated from all the gallery throughout its life cycle and can be visually conspicuous, particularly in pupal chambers and detritus in larval passages (Fig. 3). They are best isolated from active gallery systems, but also found in abandoned galleries for some time, as are ophiostomatoid fungi.

Each of the above substrates requires a different approach for optimal *Geosmithia* recovery. Spores attached to surfaces of beetle adults and larvae are cultured using a wash on standard agar media MEA and PDA, and spore load is quantified by serial dilution. Fungi from gallery detritus or walls can be cultured by directly spreading this material onto agar plates. This method readily yields *Geosmithia*, but it is not quantitative. To reduce contaminating fast-growing molds from adults trapped outside of galleries, a rinse in a modified White solution can be used (Kolařík et al., 2008). *Geosmithina* communities can be documented without culturing by using DNA metabarcoding with the standard ITS rDNA primers (Morales-Rodríguez et al., 2021).

Since the pioneering work of Wright (1938), 153 species of subcortical insects (Curculionidae, Scolytinae, Platypodinae: 140; other Curculionidae: 5; Cerambycidae: 2; Bostrichidae: 6) have been studied for the presence of *Geosmithia*; this fungus was found on 119 of them (Table 2). Within scolytine beetles, it was common on phloem-feeding species (111 out of 140 species) but also on ambrosia beetles (10 species out of 14). It also has been found on seed-feeding *Coccotrypes* (Scolytinae). *Geosmithia* vectors from other beetle groups include the Bostrichidae (6 out of 6 studied species) and Cerambycidae (2 of 2 studied species). It was absent in conifer-associated weevils of the genera *Hylobius* and *Pissodes* but it was isolated from another subcortical weevil, *Magdalis armigera* from elm. Surveys focused specifically on *Geosmithia*, or comprehensively documenting fungal communities of subcortical beetles, have been carried out mainly in Europe, the Mediterranean basin, and North America, with fewer studies from the rest of the world, such as from South America, South Africa and China (Fig. 4, Table 1, 2).

The degree of *Geosmithia* association with tree hosts of with beetle vectors can be determined by various quantitative approaches. Unfortunately, different approaches have been used by different authors, including nonstandard definitions of a sample and of sample size, making it difficult to compare between studies. We recommend using basic measures such as the proportion of gallery systems (e.g. (Kolařík et al., 2017), insect bodies (adults, larvae), or gallery segments (eg. (Dori-Bachash et al., 2015; Jankowiak et al., 2014), with *Geosmithia* out of all sampled. A more quantitative estimate of prevalence is the percentage of CFU counts belonging to *Geosmithia* within the whole sampled fungus community (Skelton et al., 2018).

Already Roland Krischner (2001) noted that bark beetles differ in their degree of association with Geosmithia. He also noted that beetles frequently transmitting Geosmithia tend to carry lower frequency and diversity of ophiostomatoid fungi, and called Geosmithia an ecological replacement for ophiostomatoids. Subsequent studies confirmed this pattern. Subcortical insects (mostly bark beetles) can be divided into those with whom Geosmithia is associated strongly, moderately, or not at all (Table 2, Fig. 5). Several beetle-tree networks are regular Geosmithia vectors: 1) broad leaved shrubs and trees, except of Betula and Alnus, and beetle species preferring trunk bases, 2) trees in the family Cupressaceae, except for *Calocedrus*, 3) trees in the family *Pinaceae*, mostly on beetles that colonize parts with the thinner bark. Geosmithia beetle vectors associated with hardwoods include subcortical Curculionidae (several subfamilies: Cossoninae, Scolytinae, and Mesoptiliinae), and Bostrichidae. Wood borers which occur under bark only as larvae but not as adults (Cerambycidae and Buprestidae) do not serve as reliable vectors, and therefore are not typically associated with Geosmithia. Geosmithia are rare or absent on insects colonizing large limbs and trunks of Pinaceae and Betula. Within Pinaceae, Geosmithia abundance and diversity is negatively correlated with thickness of the wood substrata preferred by the insects (Jankowiak and Bilanski, 2018; Kolařík and Jankowiak, 2013). Similarly on Betula, the bark beetle Scolytus ratzeburgi feeds in very moist substrate, under the impermeable bark, and hosts an abundance of ophiostomatoid fungi, but no Geosmithia (Linnakoski et al., 2008). On most other hardwoods, such as Fraxinus, Ulmus and woody plants from the Rosaceae family, vectors specific to trunk bases has much less frequent association. Most conifers within Cupressaceae support diverse communities of Geosmithia, with the exception of Calocedrus. Isolations from the beetle Phloesinus fulgens, specific to Calocedrus, typical yield low abundance of Geosmithia, and mostly Pinaceae-specific species. This may reflect the larger size of the tree and more humid environment than in most other Cupressaceae (Table 2).

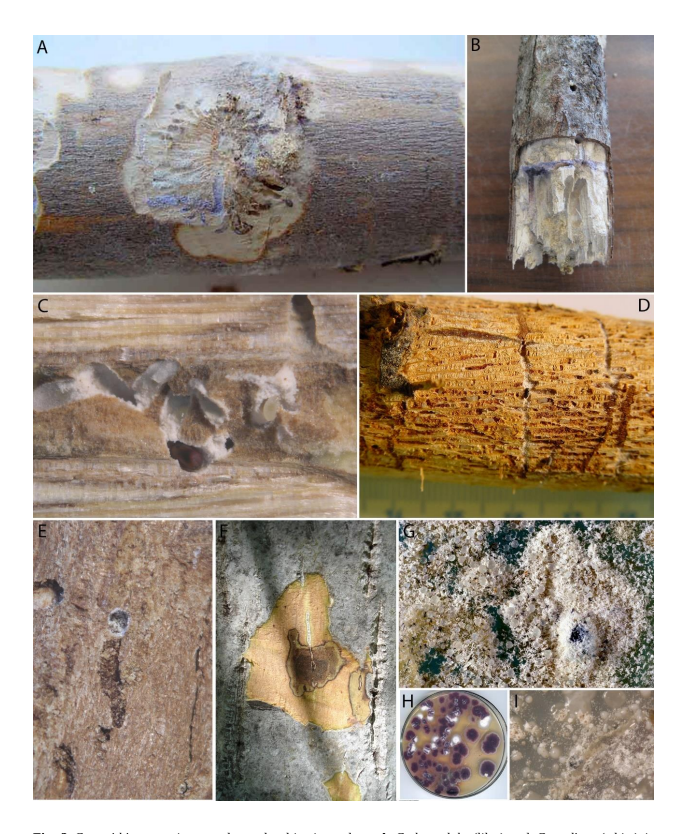


Fig. 3 Geosmithia on native samples and cultivations plates **A** G. lavendula (lilac) and G. radiata (white) in Hypoporus ficus galleries (Ficus, Croatia). **B** G. lavendula in bostrichid gallery (Toxicodendron, California). **C** G. microcorthili in galleries of ambrosia beetle Microcorhyllus sp. (Costa Rica). **D** G. flava in galleries of Ernoporus tiliae (Tilia, Czechia). **E** G. flava in pupal chamber of Cryphalus piceae (Abies, Czechia). **F** Necrosis caused by G. morbida in the phloem of Juglans (testing hole of Pityophthorus juglandis, U.S). **G** G. flava and Ophiostoma novo-

ulmi (white droplets) growing on agar plated with *Scolytus multistiatus* adults (*Ulmus*, Czechia). **H** Agar plate with *Geosmithia* colonies obtained from *H. ficus* galleries (*Ficus*, Croatia). **I.** *Pityophthorus pityographus* adult and detritus from the gallery overgrown by yeasts and *Geosmithia* sp. 24 (*Pinus*, Czechia).

1.5.5 Vector specificity, community composition and Biogeography

The recent twenty years of research on Geosmithia worldwide has finally enabled the first attempt at a synthesis of the ecology and distribution of these fungi (Table 1, 2, Fig. 4). Geosmithia can be divided into generalist species that are common across vectors worldwide, and can also be found outside of the subcortical habitat, such as in decaying wood, soil, cereals and foodstuffs (Kolařík et al., 2004; Labuda and Tancinová, 2006; Pitt and Hocking, 2009), sea sediments (Ameen et al., 2014; Sun et al., 2018), cave environment (Bastian et al., 2009; Crous et al., 2018), or as plant endophyte (Deka and Jha, 2018; McPherson et al., 2013; Sakalidis et al., 2011) (Table 1). This is typical of species in the G. pallida complex (G. pumilla, G. pulverea), then G. fassatiae, G. flava, G. granulata, G. langdonii, G. obscura, G. omnicola, G. putterilli, G. xerotolerans and Geosmithia sp. 1. In contrast, specialists species occur on vectors sharing the host plants of the same plant family. These include species that are restricted in occurrence to *Pinaceae* hosts, then G. morbida (Juglans, Europe, North America), G. ulmacea (Ulmus, Europe, North America). Sometimes the host preference is maintained in the particular geographical area with occurrence on other hosts in different areas (e.g. Geosmithia sp. 12 - Fraxinus, G. sp. 32 - Cupressaceae, G. sp. 11 - Olea, G. carolliae - Ficus). Some species common in some areas (e.g G. sp. 41 in North America, G. funiculosa in Europe), are absent in others, suggesting biogeographical patterns independent of the tree vector/host distribution (Fig. 4, Table 1). We do not have enough data for ambrosial Geosmithia species, but a high vector specificity can be expected. This appears to be the case with G. eupagioceri, which has so far only been found on the beetle Eupagiocerus dentipes in two separate collections in Central America (Kolařík and Kirkendall, 2010, J. Hulcr, unpublished).

Geosmithia communities in any given locality is structured by the influence of the local host tree availability, biogeographical limits, and the presence of suitable vector beetles. The strongest predictors is the host plant. Increasingly it is becoming evident that the beetle vectors are passive (with exception of primary ambrosia species), not actively involved with Geosmithia, and that these apparent fungus-beetle associations are derived from the underlying patterns of the tree host use by Geosmithia. As Geosmithia depend on host trees for development, and on beetles frequenting those trees for transmission, their ecological specialization is best understood on the level of tree-beetle networks. Consequently, insect vector species who regularly co-occur in the same tree part consequently have similar communities of fungi without being able to actively select them. For example, Pinus trees support the same community across different vectors beetles, and the community is different from those associated with Picea (Jankowiak et al., 2014; Kolařík and Jankowiak, 2013) and Abies (Jankowiak and Bilanski, 2018; Jankowiak and Kolarik,

2010), and even further distant from those specific to angiosperms and *Cupressaceae* (except of *Calocedrus*). As an example, polyphagous vectors such as *Pityogenes chalcographus* and *Pityophthorus pityographus* carry fungi specific to *Pinus* or *Picea*, depending on the substrate from which were collected (Jankowiak et al., 2014). This same pattern of *Geosmithia* community structure has been observed in angiosperms in temperate Europe and USA (Huang et al., 2019; Kolařík et al., 2008; Strzałka et al., 2021). Community composition is also shaped by the degree of specificity of the bark beetle vectors available in a given area. If there are only host-specific beetles on a given host plant, more sharply delimited *Geosmithia* communities are formed. Conversely, polyphagous beetle vectors create more diffuse fungus communities (Kolařík et al., 2017; Kolařík et al., 2007). In turn, this regional species pool and its dynamics also influences the richness of *Geosmithia* in individual beetle galleries: in mixed forest, *Geosmithia* communities are more diverse and then in conifer monocultures (Jankowiak et al., 2014).

Some *Geosmithia* species may be primarily endophytic, and only secondarily associated with bark beetle galleries. In California, *G. langdonii* was isolated from both bark beetle galleries as well as an end, whereas two other only isolated from bark beetle galleries (McPherson et al., 2013).

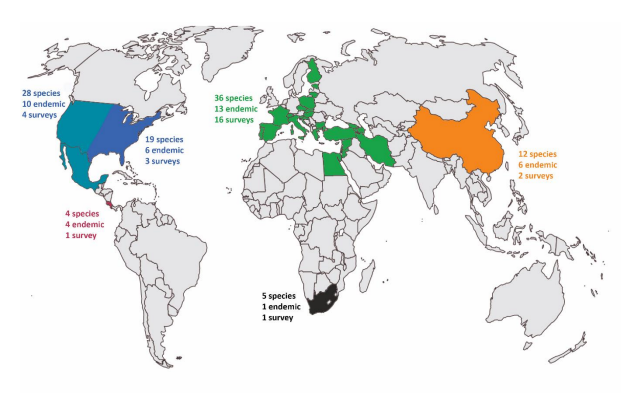


Fig. 4 Map showing the locations where *Geosmithia* species spectrum and diversity has been studied and indicates the total number of species found in each area and the number of species not yet found outside that area ("endemic" species). The map is based on Table 2.

1.5.6 Geosmithia interactions with host insect and plant

It is not completely clear how adult bark beetles transport *Geosmithia*. The majority of known and reliable vectors lack mycangia or any other organs adapted to fungus dispersal, and propagules appear to be transmitted passively in the gut or opportunistically attached to crevices and punctures of the exoskeleton. Several reports show *Geosmithia* presence in mycangia (Belhoucine et al., 2011; Kolařík et al., 2017; Six et al., 2009). Phoretic mites are also able to vector *Geosmithia* fungi (Machingambi et al., 2014).

Bark beetle associated fungi are known to have diverse symbiotic (i.e. mutualistic, neutral or antagonistic) interactions with their beetle vectors. The most straightforward is commensalism or byproduct mutualism, in which the fungus benefits from the beetle's ability to invade fresh plant tissues, which enables the fungus to exploit these nutrient sources, but the fungus does not necessarily benefit the beetle vector (Six, 2020; Six and Wingfield, 2011). Ambrosial *Geosmithia* species are mutualistic, as they provide nutrition to the beetle hosts, but it remains unknown whether the non-ambrosia *Geosmithia* also provide any benefit. Most species are good degraders of hemicellulose, and some are able to also degrade cellulose and lignin; which may benefit the beetle directly or indirectly. Some are able to utilize uric acid as nitrogen sources (Veselská et al., 2019), and thus recycling of nitrogen from the beetle waste product may be a benefit to their hosts that has not been tested. *Geosmithia* can further interact with the insect via volatile chemicals. Volatiles of *G. morbida* attract its insect vector and may synergize beetle aggregation (Blood et al., 2018).

Geosmithia also interacts with other fungi in the beetle galleries. For example, mycoparasitism by Geosmithia was observed on Ophiostoma novo-ulmi, the fungus responsible for the Dutch elm disease (Pepori et al., 2018). Geosmithia produces variety of biologically active compounds, through which they can interact with the ambient microbial community. Antibiosis towards fungi and bacteria has been found in many Geosmithia species (Veselská et al., 2019), and tested most extensively in G. lavendula (Hadj Taieb et al., 2019; Malak et al., 2013a; Stodůlková et al., 2009) and G. pallida KU693285 (Deka and Jha, 2018). Machingambi et al. (2014) have suggested that mites (bark beetle parasites) were unable to feed or reproduce in the presence of Geosmithia associates; the miticidal potential of Geosmithia should be studied in detail. G. lavendula and other species produce hydroxylated anthraquinones (hAQs) with many bioactive properties (Ganapaty et al., 2004; Hilker and Köpf, 1994; Poche, 1998; Stodůlková et al., 2009); the role of hAQs in the bark beetle ecosystem has not been evaluated.

1.5.7 Evolution and biology

The reconstruction of phylogenetic relationships among *Geosmithia* has been conducted primarily using protein-coding genes, as the ribosomal DNA markers, typically used in other fungi, genes have several limitations in *Geosmithia*. Specifically, *Geosmithia* sp. 26 is a species complex that has very different rDNA sequences from others and a very low GC content, preventing a quality alignment (Kolařík et al.,

2017). Subsequently, phylogenies inferred from rDNA and from protein coding genes are in conflict (Veselská et al., 2019). The rapid rDNA sequence evolution and GC content deviation in *Geosmithia* sp. 26 may be a consequence of fluctuations in the effective population change and bottlenecks, possibly related to the switch between free living to host associated life strategy (Kolařík and Vohník, 2017; Kolařík et al., 2021).

Geosmithia species feature many life history traits distributed across the phylogeny, making the genus an ideal model for studying the evolution of individual life styles and associated phenotypic traits (Veselská et al., 2019). Basal Geosmithia lineages are generalists, with broad host ranges across Angiosperms and Gymnosperms and sometimes found also outside of bark beetle habitat. At least six lineages convergently evolved specificity to the *Pinaceae* family (Strzałka et al., 2021; Veselská et al., 2019; Zhang et al., 2022). The shifts were accompanied by losses of metabolic capacity and by genome size inflation. In vitro, this is apparent by the inability to growth on basal CZD agar, which lacks important nutrient supplements such as vitamins (in particular the B group). Three other derived lineages converged on the ambrosia strategy, providing nutrition to specific beetle vectors. This was accompanied by the cell and genome size inflation and the production of large amounts of oleic fatty acid, likely associated with the nutritive function (Veselská and Kolařík, 2015; Veselská et al., 2019). In terms of morphology, ambrosia species produce large conidia, a phenotype seen in other ambrosia fungi (Kolařík et al., 2015; Kolařík and Kirkendall, 2010). One lineage, G. morbida, became plant pathogen, with the unique ability to digest all components of lignocellulose, what can serve as its virulence factor (Veselská et al., 2019), similarly as in other plant pathogenic fungi (Doehlemann et al., 2017; Jagadeeswaran et al., 2021). In general, specialists, such as those on *Pinaceae* and the *Junglans*-specific G. morbida, have a reduced metabolic breadth in comparison to generalists. The genome size in Geosmithia, correlates with cell size (e.g., conidia), as in most eukaryotes, and is related to the ecology of the species. Specifically, species specialized to a narrow host range (including G. morbida) have relatively large genomes, compared to generalists. The largest genomes are present in the ambrosial species (Veselská and Kolařík, 2015).

Relatively little research has been done on the genetics and mating behaviour of these fungi. As with other filamentous ascomycetes, there is a system of vegetative incompatibility that leads to some isolates making mycelial fusions with each other but not with others. In practice, this is manifested by the presence of non-coalescent lesions in the case of *G. mrobida* (Montecchio et al., 2015). The sexual stage has never been observed, and the only population genetics study in the genus suggested absence of recombination in *G. morbida* (Zerillo et al., 2014). Both mating gene idiomorphs (MAT1-1, MAT1-2) are present across the genus (M.K. unpublished) and targeted crossing experiments should be carried out to induce the sexual stage, as has been done in other moulds where the sexual stage was unknown (O'Gorman et al., 2008). In *Geosmithia*, a cleistothecial type of sexual state can be expected, as is the case with related fungi such as *Nigrosabulum*, *Mycoarachis* or *Hapsidospora* (Plishka et al., 2009; Rossman et al., 1999). The genome size, determined by flow cytometry, is 20.5 to at least 54 Mb. The largest genomes are those

of ambrosial species, probably due to the ancient polyplodisation (Veselská and Kolařík, 2015). The genome size values measured by flow cytomertry in *G. morbida*, *G. flava* and *G. putterilli* (24.4-24.7, 25.5-25.8, 26-26.3 Mbp) agree with those measured by whole genome sequencing (26.5, 29.6, 30.0 Mbs) (Schuelke et al., 2017). The number of genes is around 6,000 and only 73-146 were found to be species-specific. Between 300-400 (349-403) protein-coding genes belong to secreted proteins. There are few genes involved in secondary metabolism compared to related taxa such as *Acremonium chrysogenum* and *Stanjemonium grisellum*. In *G. morbida*, 26 genes have homologs with known involvement in interactions with the plant host and thus a potential role in pathogenesis (Schuelke et al., 2017).

Geosmithia, like other Dikarya, have hyphae coated with hydrophobins, that are small proteins, forming and hydrophic membrane and have a crucial role in interactions with hydrophobic substrates such as plant or insect cuticle. Geosmithia have class II hydrophobins, called GEO1 (Bettini et al., 2012; Frascella et al., 2014). They possess intragenic tandem repeat sequence implicated in the rapid generation of variation and subsequent adaptation. GEO1 is also under strong selection pressure, suggesting that the capacity for adhesion is important in the evolution of the genus. The cluster is evolving either via multiple horizontal transfer events and/or birth-and-death evolution. There is also good evidence that at least six Geosmithia obtained another hyrophobin, cerato-ulmin, by a horizontal transfer from Ophiostoma novo-ulmi. Cerato-ulmin is involved in the virulence of O. novo-ulmi, a causal agent in Dutch elm disease of elms, and is presented only in Geosmithia strains from elms, but not in those from other tree hosts (Bettini et al., 2014).

1.5.8 Phytopathogenic potential and TCD

While most *Geosmithia* appear to be saprophytes, the pathogenicity capabilities of some species deserves closer scrutiny. Already the first study on *Geosmithia* (Wright, 1938) studied the infectious potential of *Geosmithia* from *Scolytus praeceps* and *S. subscaber*. When inoculated into a live plant host, these strains were able to cause significant necrosis in the cambium of *Abies concolor* trunk. Based on the morphology, the strain used in the study probably belongs to *Geosmithia* sp. 34 or sp. 34 (Kolařík et al., 2017), and the pathogenicity observations, while convincing, require further verification.

Until now, the pathogenicity was studied by inoculating the phloem of seedlings or adult tree branches in more than 20 *Geosmithia* species and mostly showed no evidence of pathogenicity. In particular, no pathogenic effect was found in *Geosmithia* sp. 16 on *Abies alba* (Jankowiak and Kolarik, 2010), two species from *Geosmithia* sp. 24 species complex on *Pinus* spp. in Israel (Dori-Bachash et al., 2015), four species (*G. cupressina*, *G. langdonii*, *G. omnicola*, *G.* sp. 708) on *Cupressus* (Meshram et al., 2022), five species (*G. flava*, *G. ominicola*, *G. pumila* (=*G.* sp. 2), *G.* sp. 8, *G.* sp. A) on *Virgilia* (Machingambi et al., 2014), six *Geosmithia* spp. (*G. fagi*, *G. flava*, *G. langdonii*, *G. ulmacea*, *G. pulverea* (= sp. 3) and *G. funiculosa* (= sp. 5)) on *Acer*, *Fagus*, *Quercus*, *Tilia* and *Ulmus* (Crous et al., 2022; Strzałka

et al., 2021), *G. luteobrunnea* on *Liquidambar* (Gao et al., 2021; Zhang et al., 2022) and 11 *Geosmithia* strains originating from Czechia, Korea, Vietnam, China, Papua New Guinea, Taiwan on *Quercus shumardii*, and *Q. virginiana* (Li et al., 2022).

A few *Geosmitia* do induce phloem necroses, however, and several are involved in plant diseases. Pathogenicity assays performed using the excised shoot method showed ability of tissue lesion formation in *G. granulata* (= sp. 20), *G. lavendula*, *G. omnicola* and *G. pallida* on *Pistacia vera* (Hadj Taieb et al., 2019). However, testing pathogenicity on detached shoots is questionable, as the results cannot be extrapolated to natural field conditions. Mild, but significant, lession were created by *Geosmithia* sp. 12568 (*Cryphalus piceus*, South Korea) on *Pinus* spp. (Li et al., 2022) and by *Geosmithia* sp. on an artificially inoculated *Olea europea* trunk (van Dyk et al., 2021). Čížková et al. (2005) have shown that *G. pumila* (= sp. 2) *G. langdonii* inhibited the growth of garden cress *Lepidium sativum*.

Several tree diseases are caused by bark beetles which carry *Geosmithia* species, and the fungi may form discolored areas around the beetle galleries, but are not pathogenic themselves. In the so-called Foamy Bark Canker of *Quercus agrifolia* in California (USA), the disease appears to be caused by infestation by the bark beetle *Pseudopityophthorus pubipennis*. The beetle vectors *Geosmithia* sp. 41 and other species ((Kolařík et al., 2017)). This fungus produces significant lesions on artificially inoculated excised oak shoots (Lynch et al., 2014), but both the disease and its causal agent needs further study. Large mortality of American sweetgum (*Liquidambar styraciflua*) planted in China caused by the bark beetle *Acanthotomicus suncei* also involves several species of *Geosmithia*, most commonly *G. luteobrunnea*, around the beetle galleries, but again, the fungus is not pathogenic on its own (Gao et al., 2021; Zhang et al., 2022). Similarly puzzling is the presence of the Dutch elm disease pathogenicity factor cerato-ulmin in *Geosmithia* spp. (Bettini et al., 2014; Scala et al., 2007) while the no active role of the fungus in the disease has been demonstrated yet.

The only case where the fungus makes significant necrosis and together with its vector kills the host plant, is *Geosmithia morbida*. Together with its vector, the walnut twig beetle *Pityophthorus juglandis*, the two organisms contributed to the phenomenon of the Thousand Cancers Disease (TCD) of black walnut, *Juglans nigra* (Kolařík et al., 2011; Tisserat et al., 2009). While *G. morida* and its vector beetle *P. juglandis* are to West of the Norther America, and recently dispersed to other parts of the continent, as well as to Europe (reviewed in Daniels et al. (2016). For several years following this expansion and a drought, there was a notable dieback of black walnut across the U.S. The dieback has recently subsided, with the exceptions of locations where black walnut is planted outside of its typical growing conditions (i.e., California), suggesting that the disease has been largely a symptom of drought. While temporary, the impact of TCD spurred research on *Geosmithia* and its symbiosis with bark beetles (Fig. 1). Research on *G. mobida* involved its genetic variability (Hadziabdic et al., 2014a; Zerillo et al., 2014), host tree range a virulence (Hefty et al., 2018; Sitz et al., 2017; Sitz et al., 2021), vectors (Chahal et al., 2019), migration

(Hadziabdic et al., 2014b; Marchioro and Faccoli, 2022; Montecchio and Faccoli, 2014; Moricca et al., 2020), detection (Stackhouse et al., 2021), eradication (Dal Maso et al., 2019; Juzwik et al., 2021; Seabright et al., 2019) and competition with co-occurring fungi (Gazis et al., 2018). Walnut twig beetle transmits other *Geosmithia* species (Kolařík et al., 2017), and as can be seen from preliminary results with infection experiments with *G. obscura* (Pietsch et al., 2022).

A cross-phylogeny comparison of pathogenic and non-pathogenic species at the genome (Schuelke et al., 2017)) and phenotype level has shown that *G. morbida* is unique among *Geosmithia* species in producing an enzyme that breaks down both cellulose and lignin (Veselská et al., 2019). This capacity can be considered one of the virulence factors responsible for the ability to necrotize the phloem of walnut (Veselská et al., 2019). An interesting avenue of research is the study of the presence of viruses in *G. morbida* that may modulate virulence (Montecchio et al., 2015).

1.5.9 Secondary metabolite production and biotechnological potential

The order Hypocreales is known for the ability to produce a variety of secondary compounds, including toxins. Even crude extracts from *Geosmithia* shows the potential for antibacterial and antifungal activity across the genus (Deka and Jha, 2018; Veselská et al., 2019). Aside from common fungal metabolites, 48 secondary metabolites were found uniquely in *Geosmithia* (Table 3). Prominent yellow, orange, and red pigments produced by *G. lavendula* represent more than twenty different hydroxylated anthraquinones, often novel to science, several of them with antibacterial or anti-inflammatory activity (Malak et al., 2013c; Stodůlková et al., 2009; Stodůlková et al., 2010) and with the potential as persistent textile dyes or mordants (Flieger et al., 2009). *Geosmithia pallida* complex strain FS140 (Table 2) produced 12 different thiodiketopiperazines, including three previously unknown ones (Sun et al., 2018). A single strain identified morphologically as *G. langdonii* yielded 14 metabolites, including four new ones (Malak et al., 2013b; Malak et al., 2018). Their biological activities include antimicrobial, cytotoxic, angiotensin-converting enzyme inhibitory, antileishmanial or nemacitidal (Table 3).

While these first studied on secondary metabolites in *Geosmithia* yielded a large proportion of novel compounds and broad biological activity, the chemical arsenal is rather limited in terms of biosynthetic pathways, yielding mostly low molecular weight, structurally simple metabolites. The three species studied - *G. morbida*, *G. putterilli* and *G. flava* - produce only 14 to 19 secondary metabolite gene clusters only, which contrasts with related filamentous fungi having four-time greater number of similar genes clusters (Schuelke et al., 2017). However, the genetics of secondary metabolite production was explored in these three species only, all belonging to a single phylogenetic lineage, and the novelty of these products bids for further bioprospecting.

1.5.10 Conclusion and future research

Geosmithia has been in the spotlight only for the last decade, and so it is not surprising that many questions, long studied in ecologically similar taxa, are still unanswered. The broad evolutionary direction towards long-term and stable adaptation to beetle vectors observed in Geosmithia is the same as that observed in ophiostomatoid fungi. In both groups, it culminated in the evolution of ambrosia lineages from phloem inhabiting ancestors, and a coevolutionary response from the beetles. One of the most important paradigms that has emerged from the surge of studies on Geosmithia is that Geosmithia are an ecological complement to the ophiostomatoid fungi (Kirschner, 2001). We suggest here the terms Geosmithia-type and ophiostomatoid fungi-type association. Both fungal groups are dependent on bark beetle vectors for their dispersal and reproduction. Geosmithia, however is almost exclusively found in phloem that is drier, and typically more advanced in decay, and as a result are associated with bark beetle communities utilizing upper and thinner parts of trees. Ophiostomatoid fungi, in turn, dominate phloem which retains moisture longer, and therefore are associated with bark beetles on the trunk and roots. This patterns is replicated all around the world, but the factors responsible for it remain unclear. One of these factors could be the relatively greater tolerance to desiccation and osmotolerance in Geosmithia, as is known in G. xerotolerans (Crous et al., 2018), and greater competitivenes under drought conditions, as found in G. morbida (Williams and Ginzel, 2021). Other abiotic variables such as oxygen level and resin concentration have been identified as distinguishing the growth of Edoconidiophora polonica, living in the fresh phloem of the tree trunks, from *Ophiostoma* species living in the dead phloem and in thinner tree parts (Solheim, 1991). Their effect on the growth of *Geosmithia* should be tested (Fig. 5). In terms of pathogenicity, many bark beetle-associates cause discoloration of the phloem around the beetle gallery, but bona fide pathogenicity in the absence of the beetle is rare, truly present only in G. morbida. Several species, such as those on fir in North America, are good candidates for verification of possible weak pathogenicity (Wright, 1938).

The other major lineages of fungi associated with bark beetles – the *Ophiostomatales*, *Microascales*, and several groups within *Fusarium* – also include a range of specificity, from plant pathogens, to soil saprobes, to obligate ambrosia fungi. Sometimes closely related species display dramatically different ecology. *Geosmithia* shows similar patterns, and is an excellent model for the study of adaptive traits related to species interactions. The evolutions of these traits in *Geosmithia* has been documented at the phenotypic level (Veselská et al., 2019), and the next step needs to include a deeper, genomic level.

The main lesson learned from the recent surge of interest in the study of *Geosmithia* is that these fungi are woefully undersampled geographically. A few species are cosmopolitan generalists, but many show considerable specificity to hosts and locations (Fig. 4). Continued studies on this genus needs to emulate the methods from a better studied taxa such as *Penicillium*, *Aspergillus* or *Fusarium*. More variable DNA markers are needed to answer taxonomic, evolutionary and molecular biology questions. Similarly,

and broader array of differentiated media (DG18, G25N, MY70S, CREA) are needed for morphological and metabolic characters. *Geosmithia* also still lacks sufficient genomic data, as only three genomes have been published to date.

Given the many new chemicals isolated from *Geosmithia*, these fungi deserve research also for their biotechnological potential. These fungi do not appear to produce structurally complex substances, and also the diversity of secondary metabolites and biosynthetic pathways is modest. However, the known substances show no or very little cytotoxicity to animal cells, and at the same time they have a number of biological activities. The bioactivity is highly selective, i.e., the fungi do not harm insects, while showing antibacterial antibiosis. Their potential to interact with organisms that are pathogens of bark beetles, such as nematodes and mites, should be tested. Ambrosia species potentially an interesting target for fungal food research, since they provide a complete nutrition to their animal vectors concentrated in enlarged conidia rich in proteins and oils, while being entirely non-toxic and non-melanized (Veselská et al., 2019), M. K. unpublished).

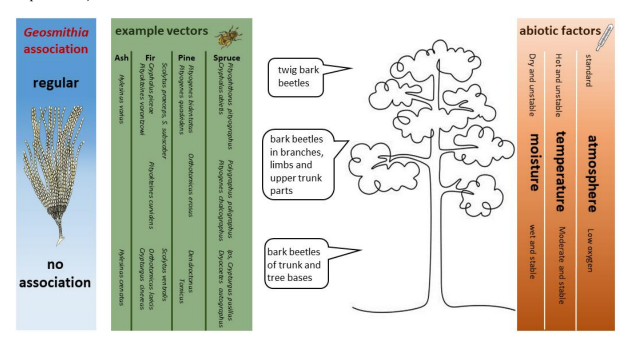


Fig 5 Schematic presentation of the *Geosmithia* association across bark beetles of different organ preference. The abiotic factors of different beetle substrates are shown.

Table 1 List of recognised Geosmithia species with geographical distribution and substrate origin. The host spectrum is expressed as list of host plant families from which the insect vector was collected

Species ¹	Central and North Eastern Europe (~Temperate Europe)	Mediterranean Basin and Black and Caspian Sea region	Western U.S. and Northern Mexico	South and Eastern U.S.	China	Other substrates or locations	References
G. bombycina	_	_	_	_	Rosaceae	_	(Zhang et al., 2022)
G. brevistipitata	_	_	_	_	Cupressaceae	_	(Zhang et al., 2022)
G. brunnea	_	_	_	Altingiaceae, Fagaceae	_	_	(Huang et al., 2017; Huang et al., 2019)
G. carolliae (G. sp. 19)	_	Moraceae	_	_	_	wing of the bat, Brazil	(Crous et al., 2018; Kolařík et al., 2007)
G. cnesini	_	_	_	_	_	ambrosia beetle Cnesinus lecontei, Costa Rica	(Kolařík et al., 2015; Kolařík and Kirkendall, 2010)
G. cupressina	_	Cupressaceae	_	_	_	_	(Meshram et al., 2022)
G. eupagioceri	_	_	_			ambrosia beetle Eupagiocerus dentipes, Costa Rica	(Kolařík and Kirkendall, 2010),
G. fagi	Rosaceae, Fagaceae	_	_	_	_	_	(Strzałka et al., 2021)

¹ The numbering for species no. 1—31 follows Kolařík et al. (2007, 2008,2013), 33-44 (Kolařík et al., 2017), 45-48 (Huang et al., 2019). *Geosmithia pulverea* and *Geosmithia* sp. 23 may represent a same species (Zhang et al., 2022).

G. fassatiae	Fagaceae, Rosaceae	_	Pinaceae, Fagaceae, Salicaceae, Lauraceae	_		_	(Kolařík et al., 2017; Kolařík et al., 2008; McPherson et al., 2013)
G. flava	Araliaceae, Betulaceae, Cupressaceae, Fagaceae, Oleaceae, Pinaceae, Rosaceae, Tiliaceae, Ulmaceae	Anacardiaceae, Lauraceae, Moraceae, Rosaceae	Anacardiaceae, Cupressaceae, Fagaceae, Juglandaceae, Pinaceae, Salicaceae	_	_	scolytids from Virgilia spp., South Africa; Ulmus glabra, Hordeum sp. Grain, England	(Kolařík et al., 2017; Kolařík et al., 2007; Kolařík et al., 2008; Kolařík et al., 2004; Machingambi et al., 2014; Pepori et al., 2015; Strzałka et al., 2021),
G. funiculosa (G. sp. 5)	Fagaceae, Oleaceae, Pinaceae, Rosaceae, Tiliaceae, Ulmaceae	Anacardiaceae	_	_	_	Scolytus beetle, UK	(Crous et al., 2022; Kolařík et al., 2007; Kolařík et al., 2008; Kolařík et al., 2004; Pepori et al., 2015; Strzałka et al., 2021)
G. fusca	_	_	_	_	Fabaceae, Phyllanthaceae, Malvaceae	_	(Zhang et al., 2022)
G. granulata (= G. sp. 20)	_	Asteraceae, Fabaceae, Moraceae, Oleaceae, Ulmaceae, Anacardiaceae	Cupressaceae, Ulmaceae	_	Fabaceae, Malvaceae		(Hadj Taieb et al., 2019; Kolařík et al., 2017; Kolařík et al., 2007; Zhang et al., 2022)

G. langdonii	Cupressaceae, Betulaceae, Fabaceae, Fagaceae, Tiliaceae, Rosaceae, Ulmaceae	Anacardiaceae, Euphorbiaceae, Fagaceae, Lauraceae	Asteraceae, Cupressaceae, Fagaceae, Lauraceae, Pinaceae	_	_		(Benvenuti et al., 2021; Hanzi et al., 2016; Juan Alfredo et al., 2020; McPherson et al., 2013; Meshram et al., 2022; Strzałka et al., 2021; Vitale et al., 2021)
G. lavendula		Anacardiaceae, Fabaceae, Moraceae, Ulmaceae	Anacardiaceae, Cupressaceae, Fagaceae, Juglandaceae, Pinaceae, Rosaceae	Fagaceae, Vitaceae, unknown		laboratory contaminant, U.S; Carya wood, Israel, soi, Venezuela	(Hadj Taieb et al., 2019; Huang et al., 2017; Huang et al., 2017; Kolařík et al., 2017; Kolařík et al., 2007; Morales-Rodríguez et al., 2021; Pitt, 1979; Six et al., 2009)
G. longistipitata (G. sp. 28)	Pinaceae	_	_	_		_	(Jankowiak et al., 2014; Kolařík and Jankowiak, 2013; Strzałka et al., 2021)
G. luteobrunnea	_	_	_	_	Altingiaceae, Ulmaceae		(Zhang et al., 2022)
G. microcorthyli	_	_	_	_	_	ambrosia beetle Microcorthylus sp., Costa Rica	(Kolařík and Kirkendall, 2010)
G. morbida		Juglandaceae	Juglandaceae	Juglandaceae	_	_	(Hadziabdic et al., 2014b; Kolařík et al.,

							2017; Montecchio et al., 2015) and others
G. obscura	Betulaceae, Fagaceae		_	Cupressaceae, Juglandaceae, Vitaceae	_	_	(Huang et al., 2017; Huang et al., 2019; Kolařík et al., 2008; Six et al., 2009)
G. omnicola (G. sp. 10)	Araliaceae, Betulaceae Cupressaceae, Fagaceae, Oleaceae, Rosaceae, Salicaceae, Tiliaceae, Ulmaceae	Anacardiaceae, Cupressaceae, Fabaceae, Lauraceae, Moraceae, Rosaceae,	_	Fagaceae, unknown	_	scolytids from Virgilia spp., South Africa; air, Israel	(Huang et al., 2017; Huang et al., 2019; Kolařík et al., 2017; Kolařík et al., 2007; Kolařík et al., 2008; Machingambi et al., 2014; Meshram et al., 2022; Pepori et al., 2015)
G. pallida s. s.	_	_	_	Juglandaceae, Fagaceae, unknown, Arecaceae	Fabaceae	cotton yarn, England; soil, Nigeria	(Huang et al., 2017; Huang et al., 2019; Kolařík et al., 2017; Pitt, 1979; Zhang et al., 2022)
G. pazoutovae	Fagaceae			_	_	_	(Strzałka et al., 2021)
G. proliferans	_	_	_	Sapindaceae	_	_	(Huang et al., 2017; Huang et al., 2019)
G. pulverea (G. sp. 3)	Betulaceae, Fagaceae, Rosaceae	_	_	_	Gnetaceae Altingiaceae, Fabaceae, Rosaceae, Anacardiaceae, Ulmaceae	roots of <i>Quercus robur</i> , soil, Czechia	(Kolařík et al., 2008; Kolařík et al., 2004; Strzałka et al., 2021; Zhang et al., 2022)

G. pumila (G. sp. 2)	Fagaceae, Oleaceae, Rosaceae, Ulmaceae	Fagaceae, Lauraceae, Ulmaceae	Rosaceae	Cupressaceae, Fagaceae, Juglandaceae, Oleaceae, Ulmaceae, unknown	Ulmaceae	scolytids from Virgilia, South Africa; apple tree, Cyprus, Cucumis melo, Peru	(Hanzi et al., 2016; Huang et al., 2017; Huang et al., 2019; Kolařík et al., 2017; Kolařík et al., 2008; Kolařík et al., 2008; Kolařík et al., 2004; Machingambi et al., 2014; Morales- Rodríguez et al., 2021; Pepori et al., 2015; Strzałka et al., 2021; Zhang et al., 2022)
G. putterillii	Rosaceae	Lauraceae	Cupressaceae, Ericaceae, Fagaceae, Juglandaceae, Lauraceae, Pinaceae, Salicaceae	_	Lauraceae	Beilschmiedia tawa wood, New Zealand	(Kolařík et al., 2017; Kolařík et al., 2007; Kolařík et al., 2008; Kolařík et al., 2004; Zhang et al., 2022)
G. radiata	_	_	_	_	Altingiaceae, Ulmaceae	_	(Zhang et al., 2022)
G. rufescens	_	_	_			two ambrosia beetle species, Costa Rica	(Kolařík and Kirkendall, 2010)
G. subfulva	_	_	_	_	Rosaceae, Anacardiaceae	_	(Zhang et al., 2022)
G. xerotolerans (= G. sp. 21)	_	Fabaceae, Moraceae, Oleaceae	Cupressaceae, Fagaceae, Pinaceae, Rosaceae	Cupressaceae, Fagaceae	Cupressaceae	house wall, Spain	(Crous et al., 2018; Huang et al., 2017; Huang et al., 2019;

							Juan Alfredo et al., 2020; Kolařík et al., 2017)
Geosmithia sp. 1	Cupressaceae, Ranunculaceae, Ulmaceae	Fabaceae, Moraceae	_	_	_	_	(Hanzi et al., 2016; Kolařík et al., 2007; Kolařík et al., 2008)
Geosmithia sp. 4	Ulmaceae	_	_	_	_	_	(Kolařík et al., 2008)
Geosmithia sp. 8	Fagaceae	_	_	unknown	_	_	(Huang et al., 2017; Huang et al., 2019; Kolařík et al., 2008)
Geosmithia sp. 9	Pinaceae	_	_	_	_		(Jankowiak and Bilanski, 2018; Jankowiak et al., 2014; Kolařík and Jankowiak, 2013; Kolařík et al., 2008)
Geosmithia sp. 11	Fagaceae	Oleaceae	_	Fagaceae	_	endophyte of Adansonia gregorii, Australia, based on sequence similarity (99%, GU19942)	(Huang et al., 2017; Huang et al., 2019; Kolařík et al., 2007; Kolařík et al., 2008; Sakalidis et al., 2011)
Geosmithia sp. 12	Fagaceae, Oleaceae	_	Oleaceae	Juglandaceae, Fagaceae, Oleaceae	_		(Huang et al., 2017; Huang et al., 2019; Kolařík et al., 2017; Kolařík et al., 2008; Strzałka et al., 2021)

Geosmithia ulmacea (G. sp. 13)	Ulmaceae	_	Ulmaceae	_	_	_	(Kolařík et al., 2017; Kolařík et al., 2008;
1							Pepori et al., 2015)
Geosmithia sp. 16	Pinaceae	_	_	_	_		(Jankowiak and Bilanski, 2018; Jankowiak and Kolarik, 2010; Jankowiak et al., 2014; Kolařík and Jankowiak, 2013; Kolařík et al., 2008; McPherson et al., 2013)
Geosmithia sp. 22	_	Fagaceae, Moraceae, Oleaceae, Rosaceae	_	_	_	_	(Kolařík et al., 2007)
Geosmithia sp. 23	Betulaceae	Moraceae, Rosaceae	Ulmaceae	Ulmaceae, unknown	_	scolytid on <i>Persea</i> gratissima, Seychelles; Malus pumila branches, Cyprus	(Huang et al., 2019; Kolařík et al., 2017; Kolařík et al., 2007; Kolařík et al., 2008)
Geosmithia sp. 24	Pinaceae	Pinaceae	_		_	_	(Dori-Bachash et al., 2015)
Geosmithia sp. 25	Pinaceae	_	_	_	_	_	(Kolařík and Jankowiak, 2013)

Geosmithia sp. 26	Pinaceae	_	Pinaceae	_	_		(Jankowiak et al., 2014; Kolařík et al., 2017; Kolařík and Jankowiak, 2013)
Geosmithia sp. 27	Pinaceae	_	Pinaceae	_		_	(Jankowiak et al., 2014; Kolařík et al., 2017; Kolařík and Jankowiak, 2013)
Geosmithia sp. 29	Pinaceae	_	_	_	_	_	(Kolařík and Jankowiak, 2013)
Geosmithia sp. 30	Pinaceae	_	_	_	_	_	(Jankowiak et al., 2014; Kolařík and Jankowiak, 2013)
Geosmithia sp. 31	Pinaceae	_	Pinaceae	_	_	_	(Jankowiak et al., 2014; Kolařík and Jankowiak, 2013)
Geosmithia sp. 32	Cupressaceae	Oleaceae	Cupressaceae			_	(Juan Alfredo et al., 2020; Kolařík et al., 2017; Kolařík et al., 2008)
Geosmithia sp. 33	_	_	Pinaceae	_	_	_	(Kolařík et al., 2017)
Geosmithia sp. 34	_	-	Cupressaceae — Calocedrus, Pinaceae	_	_	_	(Kolařík et al., 2017)

Geosmithia sp. 35	_	_	Pinaceae	_	_	_	(Kolařík et al., 2017)
Geosmithia sp. 36	_	_	Pinaceae	_	_	_	(Kolařík et al., 2017)
Geosmithia sp. 37	_	_	Pinaceae	_	_	_	(Kolařík et al., 2017)
Geosmithia sp. 38	_	_	Fagaceae	_	_	_	(Kolařík et al., 2017)
Geosmithia sp. 40	_	_	Pinaceae	_	_	_	(Kolařík et al., 2017)
Geosmithia sp. 41	_	_	Anacardiaceae, Asteraceae, Fagaceae, Lauraceae, Pinaceae, Rosaceae	Juglandaceae, Fagaceae, unknown	_		(Huang et al., 2017; Huang et al., 2019; Kolařík et al., 2017)
Geosmithia sp. 42	_	_	Cupressaceae - Calocedrus, Pinaceae, Rosaceae	_	_	_	(Kolařík et al., 2017)
Geosmithia sp. 43	_	_	Pinaceae	_	_	_	(Kolařík et al., 2017)
Geosmithia sp. 44	_	_	Cupressaceae, Pinaceae	Pinaceae	_	_	(Huang et al., 2017; Huang et al., 2019; Kolařík et al., 2017)
Geosmithia sp. 45	_	_	_	Pinaceae	_	_	(Huang et al., 2017; Huang et al., 2019)
Geosmithia sp. 39	_	_	Juglandaceae	_	_	_	(Kolařík et al., 2017)
Geosmithia sp. 46	_	_	_	Juglandaceae, Fagaceae	_	_	(Huang et al., 2017; Huang et al., 2019)

Geosmithia sp. 47	_	_	_	Juglandaceae, Fagaceae	_	_	(Huang et al., 2017; Huang et al., 2019)
Geosmithia sp. 48	_	_	_	Cupressaceae	_	_	(Huang et al., 2017; Huang et al., 2019)
Geosmithia sp. (G. pallida complex)	_	_	_	_	_	Scolytoplatypus fasciatus from Virgilia, South Africa	(Machingambi et al., 2014)
Geosmithia sp. (G. pallida complex)		_	_	_	_	endophyte of <i>Brucea</i> mollis, India, unique lineage based on KU693285	(Deka and Jha, 2018).
Geosmithia sp. (G. pallida complex)	-	_	-	-	-	sea sediment, China, unique lineage based on MK047400	(Sun et al., 2018).
Geosmithia sp. (G. sp. 24 complex)	_	_	_	_	_	scolytids from <i>Pinus</i> , Israel, sister to <i>Geosmithia</i> sp. 24	(Dori-Bachash et al., 2015)
Geosmithia sp. (G. sp. 8 complex)	_	_	_	_	_	Phloeosinus spp. from Cupresssus, Izrael	(Meshram et al., 2022)

Table 2 Summary of the insect vectors studied for Geosmithia presence and the strength of Geosmithia-vector association

Region	Host plant	Insect vector	Tree part and ecology ¹	Relative <i>Geosmithia</i> abundance ²	Total numbe r of Geosm ithia spp.	References
	Cassia	Microcorthylus sp.	T,B/AMB	100%	1	(Kolařík and Kirkendall, 2010)
Costa Rica	Croton	Cnesinus lecontei	T,B/AMB	100%	2	(Kolařík and Kirkendall, 2010)
	Paulinia	Eupagiocerus dentipes	T,B/AMB	100%	2	(Kolařík and Kirkendall, 2010)
China	Pinus	Dendroctonus armandi	T/PHL	0%	0	(Hu et al., 2015)

¹ The organ preference and feeding habit was classified in the following categories: T − small twigs, B − branches, L − limbs and top of the trunk, thin barked parts, small diameter trunk, T − large diameter trunk, stumps, trunk bases, V − any part, PHL − phloem and bark, AMB − ambrosia beetle, XYL − sapwood; and is based on Postner (1974) and Foit (2010) and Kula et al. (2000) for European, Wood (1982), Bright et Stark (1973) and Smith et al. (2014) for American and Machingambi et al. (2014) for South African species.

² Relative frequency is given, depending on the study, as the number of independent gallery systems, or beetle individuals (adults, or larvae) where *Geosmithia* have been found. If *Geosmithia* were found but not quantified, a + symbol is given.

	various host	Acanthotomicus suncei, Crossotarsus emancipates, Cryphalus eriobotryae, C. kyotoensis, Dinoderus sp., Ernoporus japonicus, Hypothenemus sp., Microperus sp., Phloeosinus cf. hopehi, Phloeosinus sp., Scolytus jiulianshanensis, S. semenovi, Sinoxylon cf. cucumella (Bostrichidae), Xylocis tortilicornis (Bostrichidae)	-	+	3	(Zhang et al., 2022)
	Abies	Cryphalus piceae	T, B,L/PHL	37-82%	6	(Jankowiak and Bilanski, 2018; Jankowiak and Kolarik, 2010; Kirschner, 2001; Kolařík et al., 2008)
		Orthotomicus laricis	T/PHL	0%	0	(Jankowiak and Bilanski, 2018)
		Pisodes piceae (Curculionidae, Molytinae)	L, T/PHL,XYL	0%	0	(Jankowiak and Bilanski, 2018)
		Pityokteines curvidens	L,T/PHL	24%	1	(Jankowiak and Bilanski, 2018)
		Pityokteines vorontzowi	B,L/PHL	70%	1	(Jankowiak and Bilanski, 2018)
Europe,		Pityophthorus pityographus	T, B/PHL	80%	2	(Jankowiak and Bilanski, 2018)
Mediterra nean		Trypodendron lineatum	T/PHL	0%	0	(Jankowiak and Bilanski, 2018)
		Xyleborinus saxesenii	T/PHL	0%	0	(Jankowiak and Bilanski, 2018)
	Alnus	Dryocoetes alni	B,L/PHL	5%	1	(Strzałka et al., 2021)
	Betula	Scolytus ratzeburgi	L,T/PHL	0%	0	(Kolařík et al., 2008; Linnakoski et al., 2008; Strzałka et al., 2021)
	Carpinus	Scolytus carpini	B,T/PHL	100%	5	(Kolařík et al., 2008)
	Clematis	Xylocleptes bispinus	liana stem/PHL	20%	1	(Kolařík et al., 2008)
	Cupressace ae	Phloeosinus armatu, P. bicolor	T, B/PHL	100%	4	(Meshram et al., 2022)

Cupressace ae	Phloeosinus henschi, P. thujae	T, B/PHL	100%	4	(Kolařík et al., 2007; Kolařík et al., 2008)
Cytisus	Phloeotribus rhododactylus, Phloeophthorus cristatus	B,L/PHL	50-100%	1	(Kolařík et al., 2007; Kolařík et al., 2008)
Euphorbia	Aphanarthrum sp.	stem/PHL	100%	1	(Kolařík et al., 2007)
Fagus	Ernoporicus fagi, Taphrorychus bicolor	T,B inner bark of trunk/PHL	27-29%	4	(Kolařík et al., 2008; Strzałka et al., 2021)
Ficus	Hypoborus ficus	B,L,T/PHL	98%	9	(Kolařík et al., 2007)
	Hylesinus crenatus	T/PHL	24%	1	(Strzałka et al., 2021)
Fraxinus	Hylesinus varius, H. orni, H. toranio	B,L/PHL	90-100%	5	(Kolařík et al., 2008; Strzałka et al., 2021)
Hedera	Kissophagus hederae	liana stem/PHL	33%	2	(Kolařík et al., 2008)
	Ips cembrae	L,T/PHL	0%	0	(Jankowiak and Rossa, 2007)
Larix	Orthotomicus laricis	T/PHL	0%	0	(Kirschner, 2001)
	Trypodendron lineatum	T/PHL	0%	0	(Jankowiak and Bilanski, 2018; Kirschner, 2001)
Laurus	Liparthrum colchicum	B,L/PHL	100%	3	(Benvenuti et al., 2021; Kolařík et al., 2007; Vitale et al., 2021)
Olea	Phleotribus scarabeiodes	B,L/PHL	100%	4	(Kolařík et al., 2007)
	Cryphalus abietis	B,L/PHL	100%	3	(Kolařík and Jankowiak, 2013)
Picea	Crypturgus cinereus	L,T/PHL	0%	0	(Kirschner, 2001)
	Crypturgus pusillus	B/PHL	0%	0	(Kirschner, 2001)

	Dendroctonus micans	L,T/PHL	0%	0	(Dohet et al., 2016; Kolařík and Jankowiak, 2013)
	Dryocoetes autographus	T/PHL	0-6%	2	(Jankowiak et al., 2014; Kirschner, 2001)
	Hylurgops palliatus	T/PHL	0-2%	1	(Jankowiak, 2006b; Jankowiak et al., 2014; Kirschner, 2001)
	Ips amitinus	L/PHL	2%	1	(Jankowiak et al., 2014)
	Ips duplicatus	L,T/PHL	0%	0	(Kolařík and Jankowiak, 2013)
	Ips typograhpus	T/PHL	0-1%	2	(Jankowiak and Hilszczański, 2005; Kirschner, 2001; Persson et al., 2009)
	Pityogenes chalcographus	L/PHL	0-24%	6	(Jankowiak et al., 2014; Kirschner, 2001)
	Pityophthorus pityographus	B,L/PHL	58%	7	(Jankowiak et al., 2014)
	Polygraphus poligraphus	L/PHL	24%	3	(Jankowiak et al., 2014; Kirschner, 2001)
	Trypodendron lineatum	T/PHL	0%	0	(Jankowiak and Bilanski, 2018; Kirschner, 2001)
	Acanthocinus aedilis (Cerambycidae)	L,T/PHL,XYL	0%	0	(Jankowiak and Rossa, 2007)
	Crypturgus cinereus	L,T/PHL	0%	0	(Kirschner, 2001)
Pinus	Hylobius abietis (Curculionidae, Molytinae)	T/PHL,XYL	0%	0	(Kolařík and Jankowiak, 2013)
	Hylurgus ligniperda	T/PHL	0%	0	(Davydenko et al., 2014)
	Ips acuminatus	L/PHL	0%	0	(Davydenko et al., 2017)

Ips sexdentatus	T/PHL	0%	0	(Davydenko et al., 2021; Kirschner, 2001; Kolařík and Jankowiak, 2013)
Monochamus galloprovincialis (Cerambycidae)	L/PHL,XYL	3.30%	1 (?)	(Jankowiak and Rossa, 2007)
Orthotomicus erosus	L/PHL	23%	1	(Dori-Bachash et al., 2015)
Orthotomicus laricis	T/PHL	0%	0	(Jankowiak and Bilanski, 2018), (Kirschner, 2001)
Pissodes castaneus, P. piniphilus (Curculionidae, Molytinae)	T/PHL,XYL	0%	0	(Kolařík and Jankowiak, 2013)
Pityogenes bidentatus	B,L/PHL	41-82%	6	(Jankowiak et al., 2014; Jankowiak and Rossa, 2008)
Pityogenes calcaratus	T, B,L/PHL	84%	1	(Dori-Bachash et al., 2015)
Pityogenes chalcographus	L/PHL	24%	4	(Jankowiak et al., 2014)
Pityogenes quadridens	B,L/PHL	86%	2	(Kolařík and Jankowiak, 2013)
Pityophthorus pityographus	B,L/PHL	69%	7	(Jankowiak et al., 2014)
Tomicus destruens	L/PHL	0%	0	(Dori-Bachash et al., 2015; Muñoz- Adalia et al., 2017)
Tomicus minor	L,T/PHL	0%	0	(Jankowiak, 2008)
Tomicus piniperda	T/PHL	0%	0	(Jankowiak, 2006a; Jankowiak and Bilański, 2007; Muñoz-Adalia et al., 2017; Silva et al., 2015)
Trypodendron lineatum	T/PHL	0%	0	(Jankowiak and Bilanski, 2018; Kirschner, 2001)

Pistatia	Chaetoptelius perrisi, Ch. vestitus	B,L/PHL	20-100%		(Hadj Taieb et al., 2019; Kolařík et al., 2007)
Populus	Trypophloeus spp.	T,B/PHL	33%	1	(Kolařík et al., 2008)
	Scolytus mali	L,T/PHL	42%	3	(Strzałka et al., 2021)
Rosaceae	Scolytus rugulosus	B,L/PHL	75-100%	8	(Kolařík et al., 2007; Kolařík et al., 2008; Strzałka et al., 2021)
	Dryocoetes villosus	thick bark of B,L/PHL	0%	0	(Strzałka et al., 2021)
Quercus	Scobicia pustulata (Bostrichidae)	B,T/XYL	100%	1	(Kolařík et al., 2007)
Quercus	Scolytus intricatus	T, B/PHL	15-100%	11	(Kolařík et al., 2008; Kubátová et al., 2004; Strzałka et al., 2021)
	Platypus cylindrus	T, B/AMB	Anisandrus dispar	1	(Belhoucine et al., 2011)
Spartium	Liparthrum genistae, Phloeotribus rhododactylus, Phloeophthorus cristatus	B,L/PHL	50-100%	1	(Kolařík et al., 2007; Kolařík et al., 2008)
Tilia	Ernoporus tiliae	B,T/PHL	73-100%	4	(Kolařík et al., 2008; Strzałka et al., 2021)
	Magdalis armigera (Curculionidae, Magdalinae)	B,L/PHL,XYL	+	1	(Kolařík et al., 2008)
Ulmus	Scolytus multistriatus, S. pygmaeus, S. kirsch, Pteleobius vittatus	B,L/PHL	47-100%	7	(Kolařík et al., 2007; Kolařík et al., 2008; Strzałka et al., 2021)
	Scolytus scolytus	L,T/PHL	22%	3	(Strzałka et al., 2021)
various	Anisandrus dispar	V/AMB	4%	1	(Strzałka et al., 2021)
hardwoods	Trypodendron domesticum	L/AMB	0%	0	(Strzałka et al., 2021)

		Xyleborinus saxeseni	B,L/AMB	0%	0	(Strzałka et al., 2021)
		Xyleborus monographus	B/L/AMB	0%	0	(Strzałka et al., 2021)
		Scolytus subscaber	B/PHL	72%	1 (?)	(Wright, 1938)
	Abies	Scolytus ventralis	T/PHL	0%	0	(Kolařík et al., 2017; Wright, 1938)
		Scolytus praeceps	B,L/PHL	87-88%	4	(Kolařík et al., 2017; Wright, 1938)
	Acer	Phloeotribus frontalis	V, inner bark/PHL	33%	1	(Huang et al., 2017)
	Calocedrus	Phloeosinus fulgens	B,L/PHL	20%	4	(Kolařík et al., 2017)
	Carya	Hypothenemus dissimilis	T,B/PHL	50-100%	2	(Huang et al., 2017; Huang et al., 2019)
		Hypothenemus rotundicollis	T,B/PHL	100%	2	(Huang et al., 2019)
North		Xylobiops basilaris (Bostrichidae)	V/XYL	0-100%	3	(Huang et al., 2019)
America	Celtis	Chramesus chapuisii	T,B/PHL	10-11%	2	(Huang et al., 2017; Huang et al., 2019)
	Cents	Phloeotribus texanus	B,L/PHL	17%	1	(Huang et al., 2017; Huang et al., 2019)
	Cinnamom um	Cnesinus strigicollis	T,B,L/PHL	0%	0	(Huang et al., 2017)
		Phloeosinus cupressi, P. sequoia, P. canadensis, P. punctatus	V/PHL	80-100%	6	(Kolařík et al., 2017)
	Cupressace	Phloeosinus serratus, P. deleoni	V/PHL	100%	2	(Juan Alfredo et al., 2020)
	ае	Phloeosinus dentatus	V/PHL	50-60%	4	(Huang et al., 2017; Huang et al., 2019)

	Hylesinus aculeatus	L/PHL	17%	2	(Huang et al., 2017)
Fraxinus	Hylesinus oregonus	L, T/PHL	80%	1	(Kolařík et al., 2017)
Juglans	Pityophthorus juglandis	T,B/PHL	90-100%	5	(Kolařík et al., 2017), various others studies
Juniperus	Ambrosiodmus lecontei	V/AMB	0%	0	(Huang et al., 2017; Huang et al., 2019)
Notholithoc arpus	Pseudopityophthorus pubipennis	T, B,L/PHL	100%	7	(Kolařík et al., 2017)
Phoenix	Coccotrypes dactyliperda	seeds	75%	1	(Huang et al., 2017)
	Dendroctonus frontalis	T/PHL	0%	0	(Dighton et al., 2021)
	Dendroctonus ponderosae	T/PHL	0%	0	(Lee et al., 2006; Lim et al., 2005)
	Dendroctonus punctatus, D. valens	T/PHL	0%	0	(Dohet et al., 2016)
	Dendroctonus rhizophagus	T, seedlings/PHL	0%	0	(Gonzalez-Escobedo et al., 2019)
Pinus	Ips avulsus	L,T/PHL	0%	0	(Huang et al., 2017; Huang et al., 2019)
	Ips pini	T,L/PHL	0%	0	(Lim et al., 2005)
	Ips plastographus	T/PHL	100%	2	(Kolařík et al., 2017)
	Orthotomicus latidens	L/PHL	45%	2	(Kolařík et al., 2017)
	Orthotomicus spinifer	T/PHL	100%	1	(Kolařík et al., 2017)
	Pityogenes knechteli	B,L/PHL	30%	4	(Kolařík et al., 2017)

	Pityophthorus confusus	little known, B,L/PHL	29-50%	1	(Huang et al., 2017; Huang et al., 2019)	
	Pityophthorus pulicarius	T/PHL	0-12.5%	1	(Huang et al., 2017; Huang et al., 2019)	
	Pseudips mexicanus	T/PHL	50%	1	(Kolařík et al., 2017)	
	Pityophthorus annectens	little known, B/PHL	10%	1	(Huang et al., 2019)	
Pistatia	Scobicia sp. (Bostrichidae)	V/XYL	100%	2	(Kolařík et al., 2017)	
Prunus	Phloeotribus liminaris	L/PHL	25%	1	(Huang et al., 2017)	
	Carphoborus vandykei	unknown/PHL	15%	1	(Kolařík et al., 2017)	
Pseudotsug a	Cryphalus pubescens	L, seedlings/PHL	40%	3	(Kolařík et al., 2017)	
	Scolytus oregoni	L/PHL	100%	5	(Kolařík et al., 2017)	
Rosaceae	Scolytus rugulosus	B,L/PHL	100%	5	(Kolařík et al., 2017)	
Quercus	Micracisella nanula	T,PHL	50%	1	(Huang et al., 2017; Huang et al., 2019)	
	Pseudopityophthorus minutissimus	B,L/PHL	28-56%	1	(Huang et al., 2017; Huang et al., 2019)	
Quercus	Pseudopityophthorus pubipennis	T, B,L/PHL	100%	7	(Kolařík et al., 2017; McPherson et al., 2013)	
Toxicodend ron	Cactopinus rhois	liana stem/PHL	67%	2	(Kolařík et al., 2017)	
Ulmus	Scolytus multistriatus, S. schevyrewi	B,L/PHL	100%	3	(Kolařík et al., 2017)	

	Umbellular ia	Scobicia declivis (Bostrichidae)	V/XYL	72%	4	(Kolařík et al., 2017)
South Africa	various hardwoods or unknown	Chaetophloeus sp.	T,B/PHL	100%	1	(Huang et al., 2017)
		Cryptocarenus seriatus	T/PHL	100%	1	(Huang et al., 2017)
		Xyleborus celsus	T/AMB	100%	1	(Huang et al., 2017)
		Hylocurus hirtellus	T,B/PHL	80%	3	(Kolařík et al., 2017)
		Hypothenemus eruditus	V, mostly T, B/PHL	0-25%	1	(Huang et al., 2017; Huang et al., 2019)
		Xylosandrus compactus	T/AMB	0-50%	1	(Huang et al., 2017; Huang et al., 2019)
		Xylosandrus crassiusculus	V/AMB	0-50%	1	(Huang et al., 2017; Huang et al., 2019)
		Xylosandrus crassiusculus, Xylosandrus compactus	V, mostly T,B/AMB	0%	0	(Huang et al., 2017; Huang et al., 2019)
		Xylosandrus mutilatus	V/AMB	+	2	(Six et al., 2009)
		Scolytoplatypus fasciatus	V/AMB	100%	1	(Machingambi et al., 2014)
		Cryphalini sp. 1, Hapalogenius fuscipennis, Liparthrum sp. 1	B/PHL	100%	5	(Machingambi et al., 2014)

Table 3 List of secondary metabolites reported from Geosmithia

Compound	Chemical class	Activity	Occurrence	Reference
Rheoemodin (1,3,6,8- tetrahydroxyanthraquinone)	anthraquinone	inflammatory activity (10 μg/mL), Acinetobacter baumannii (MIC 12.5 μg/mL)	G. lavendula	(Stodůlková et al., 2009), (Wang et al., 2019)
Rhodolamprometrin (1-acetyl-2,4,5,7- tetrahydroxyanthraquinone)	anthraquinone	Staphylococcus aureus, Bacillus subtilis (MIC 64– 512 μg/mL, affects morphology of mammalian cells	G. lavendula	(Stodůlková et al., 2009), (Malak et al., 2013c)
1-acetyl-2,4,5,7,8- pentahydroxyanthraquinone	anthraquinone	S. aureus, Bacillus subtilis (MIC 64–512 μg/mL), inflammatory activity (10 μg/mL), affects cell cycle mammalian cell	G. lavendula	(Stodůlková et al., 2009), (Malak et al., 2013c)
2,4,5,7-tetrahydroxy AQ-1-carboxylic acid methyl ester	anthraquinone	n.a.	G. lavendula	(Stodulkova et al., 2010)
1,x-diacetyl-2,4,5,7-tetrahydroxy AQ				
1,x-diacetyl-2,4,5,7,8-pentahydroxy AQ				
1-acetyl-2,4,5,7,8-pentahydroxy AQ				
1-acetyl-2,4,5,7-tetrahydroxy AQ				
1,x-diacetyl-monomethoxy-trihydroxy AQ				
1,3,6,8-Tetrahydroxy AQ				
1,3,5,6,8-pentahydroxy AQ				
1-acetyl-dimethoxy-dihydroxy AQ				
1,x-diacetyl-dimethoxy-dihydroxy AQ				
1-acetyl-monomethoxy-tetrahydroxy AQ				
1,x-diacetyl-monomethoxy-tetrahydroxy AQ				
1-acetyl-monomethyl-trihydroxy AQ				
1-acetyl-monomethoxy-trihydroxy AQ				
1,x-diacetyl-trimethoxy-hydroxy AQ				

1-acetyl-2,4,6,8-tetrahydroxy-9,10-anthraquinone	anthraquinone	anti methicillin resistant Staphylococcus aureus (IC50 16.1 µg/mL)	G. lavendula	(Malak et al., 2013c)
2-acetyl-1,4,5,7-tetrahydroxy-9,10-anthraquinone	anthraquinone	n.a.	G. lavendula	(Malak et al., 2013c)
1-acetyl-2,4,5,6,7-pentahydrxy-9,10-anthraquinone	anthraquinone	n.a.	G. lavendula	(Malak et al., 2013c)
4-(2',4'-dihydroxy-6'- (hydroxymethyl)benzyl)benzene-1,2-diol	benzhydryl	n.a.	G. langdonii	(Malak et al., 2014)
p-hydroxybenzyl alcohol	benzyl alcohol	broad application in human medicine, mostly neuroactive (Zhu et al., 2018)	G. lavendula	(Malak et al., 2013c)
4-hydroxybenzyl alcohol	benzyl alcohol	antioxidant and anti- inflammatory activities (Kumar et al., 2017)	G. langdonii	(Malak et al., 2014)
(1S,2R,3R,4R,5R)-2,3,4-trihydroxy-5-methylcyclohexyl-2',5'-dihydroxybenzoate	carbasugar	antileishmanial (IC50 100 $\mu M)$	G. langdonii	(Malak et al., 2018)
1S,2S,3S,4R,5R)-4-[(2',5'-dihydroxybenzyl)oxy]-5-methylcyclohexane-1,2,3-triol	carbasugar	antileishmanial (IC50 57 $$ $\mu M)$	G. langdonii	(Malak et al., 2018)
3,4-dihydroxytoluene (4-methylcatechol)	catechol	n.a.	G. langdonii	(Malak et al., 2014)
bisdethiobis (methylthio)gliotoxin	cyclic dipeptide	antibacterial (Ratnaweera et al., 2016)	G. pallida MK047400	(Sun et al., 2018)
6-acetylbis(methylthio)gliotoxin	cyclic dipeptide	no antibacterial or cyotoxic (Liang et al., 2014)	G. pallida MK047400	(Sun et al., 2018)
6-deoxy-5a,6-didehydrogliotoxin	cyclic dipeptide	cytotoxic (Sun et al., 2012)	G. pallida MK047400	(Sun et al., 2018)
5a,6-didehydrogliotoxin	cyclic dipeptide	n.a.	G. pallida MK047400	(Sun et al., 2018)
6-(phenylmethyl)-(3R,6R)-2,5- piperazinedione	cyclic dipeptide	n.a.	G. pallida MK047400	(Sun et al., 2018)
3-(hydroxymethyl)-3,6-bis(methylthio)-6-(phenylmethyl)-(3R,6R)-2,5-piperazinedione	cyclic dipeptide	n.a.	G. pallida MK047400	(Sun et al., 2018)

3-(hydroxymethyl)-6-(methoxyl)-6- (phenylmethyl)-(3R,6R)-2,5- piperazinedione	cyclic dipeptide	n.a.	G. pallida MK047400	(Sun et al., 2018)
5a,6- anhydrobisdethiobis(methylthio)gliotoxin	cyclic dipeptide	n.a.	G. pallida MK047400	(Sun et al., 2018)
Geospallin A	cyclic dipeptide	angiotensin-converting enzyme	G. pallida MK047400	(Sun et al., 2018)
Geospallin B	cyclic dipeptide	angiotensin-converting enzyme	G. pallida MK047400	(Sun et al., 2018)
Geospallin C	cyclic dipeptide	angiotensin-converting enzyme	G. pallida MK047400	(Sun et al., 2018)
(+)-epiepoformin	cyclohexane epoxide	antifungal, zootoxic and phytotoxic (Cala et al., 2018)	G. langdonii	(Malak et al., 2014)
(-)-dihydroepiepoformin	cyclohexane epoxide	n.a.	G. langdonii	(Malak et al., 2014)
(4S,5S)-4,5-dihydroxy-2-methylcyclohex-2-enone	cyclohexene and cyclohexenone	potato microtuber induction (Salvatore et al., 2020)	G. langdonii	(Malak et al., 2014)
(4R,5R,6R)-4,5-dihydroxy-6-(6'-methylsalicyloxy)-2-methyl-2-cyclohexen-1-one	cyclohexene and cyclohexenone	n.a,	G. langdonii	(Malak et al., 2014)
didodecyl thiodipropionate	dicarboxylic acid	antioxidant	G. lavendula	(Malak et al., 2013c)
6-methylsalicylic acid	hydroxybenzoate	n.a.	G. langdonii	(Malak et al., 2014)
3-hydroxytoluene (m-Cresol)	phenol derivate	n.a.	G. langdonii	(Malak et al., 2014)
2,5-dihydroxybenzaldehyde	phenolic aldehyde	nematicidal, cytotoxic (Kim et al., 2021)	G. langdonii	(Malak et al., 2014)
gentisylquinone	quinone	antibiotic, herbicide (Buckingham, 1996)	G. langdonii	(Malak et al., 2014)
(22E)-ergosta-6,22-diene-3 β ,5 α ,8 α -triol	sterol	n.a.	G. lavendula	(Malak et al., 2013c)

1.5.11. References

Ameen, F., Moslem, M.A., Hadi, S., Al-Sabri, A., 2014. Biodegradation of cellulosic materials by marine fungi isolated from South Corniche of Jeddah, Saudi Arabia. J. Pure Appl. Microbiol 8, 3617-3626.

Bastian, F., Alabouvette, C., Saiz-Jimenez, C., 2009. The impact of arthropods on fungal community structure in Lascaux Cave. J. Appl. Microbiol. 106, 1456-1462.

Beaver, R.A., 1989. Insect-fungus relationships in the bark and ambrosia beetles, in: Wilding, N., Collins, N.M., Hammond, P.M., Webber, J.F. (Eds.), Insect-Fungus Interactions. Academic Press, London, pp. 121-143.

Belhoucine, L., Bouhraoua, R.T., Meijer, M., Houbraken, J., Harrak, M.J., Samson, R.A., Equihua-Martinez, A., Pujade-Villar, J., 2011. Mycobiota associated with *Platypus cylindrus* (Coleoptera: Curculionidae, Platypodidae) in cork oak stands of North West Algeria, Africa. African Journal of Microbiology Research 5, 4411-4423.

Benvenuti, C., Strangi, A., Iovinella, I., Barzanti, G.P., Simoni, S., Vitale, S., Luongo, L., Francardi, V., Roversi, P., 2021. *Xylosandrus compactus* and *Liparthrum colchicum* (Coleoptera Scolytinae) in Tuscany: a preliminary screening of associated Fungi. J. Zool. 104, 139-146.

Bettini, P.P., Frascella, A., Comparini, C., Carresi, L., Pepori, A.L., Pazzagli, L., Cappugi, G., Scala, F., Scala, A., 2012. Identification and characterization of GEO1, a new class II hydrophobin from *Geosmithia* spp. Canadian Journal of Microbiology 58, 965-972.

Bettini, P.P., Frascella, A., Kolařík, M., Comparini, C., Pepori, A.L., Santini, A., Scala, F., Scala, A., 2014. Widespread horizontal transfer of the cerato-ulmin gene between *Ophiostoma novo-ulmi* and *Geosmithia* species. Fungal Biol 118, 663-674.

Blood, B., Klingeman, W., Paschen, M., Hadžiabdić, Đ., Couture, J., Ginzel, M., 2018. Behavioral responses of *Pityophthorus juglandis* (Coleoptera: Curculionidae: Scolytinae) to volatiles of black walnut and *Geosmithia morbida* (Ascomycota: Hypocreales: Bionectriaceae), the causal agent of thousand cankers disease. Environ. Entomol. 47, 412-421.

Bright, D.E., Stark, R.W., 1973. The bark and ambrosia beetles of California (Coleoptera: Scolytidae and Platypodidae). Bulletin of the California Insect Survey 16.

Buckingham, J., 1996. Dictionary of organic compounds. CRC Press.

Cala, A., Masi, M., Cimmino, A., Molinillo, J.M., Macias, F.A., Evidente, A., 2018. (+)-epi-epoformin, a phytotoxic fungal cyclohexenepoxide: structure activity relationships. Molecules 23, 1529.

Chahal, K., Gazis, R., Klingeman, W., Hadziabdic, D., Lambdin, P., Grant, J., Windham, M., 2019. Assessment of alternative candidate subcortical insect vectors from walnut crowns in habitats quarantined for thousand cankers disease. Environ. Entomol. 48, 882-893.

Čížková, D., Šrůtka, P., Kolařík, M., Kubátová, A., Pažoutová, S., 2005. Assessing the pathogenic effect of *Fusarium*, *Geosmithia* and *Ophiostoma* fungi from broad-leaved trees. Folia Microbiol. 50, 59-62.

Crous, P., Luangsa-Ard, J., Wingfield, M., Carnegie, A., Hernández-Restrepo, M., Lombard, L., Roux, J., Barreto, R., Baseia, I., Cano-Lira, J., 2018. Fungal Planet description sheets: 785–867. Persoonia: Molecular Phylogeny and Evolution of Fungi 41, 238.

Crous, P.W., Boers, J., Holdom, D., Steinrucken, T., Tan, Y., Vitelli, J., Shivas, R., Barrett, M., Boxshall, A.-G., Broadbridge, J., 2022. Fungal Planet description sheets: 1383–1435. Persoonia 48, 261-371.

Dal Maso, E., Linaldeddu, B.T., Fanchin, G., Faccoli, M., Montecchio, L., 2019. The potential for pesticide trunk injections for control of thousand cankers disease of walnut. Phytopathologia Mediterranea 58, 73-79.

Daniels, D.A., Nix, K.A., Wadl, P.A., Vito, L.M., Wiggins, G.J., Windham, M.T., Ownley, B.H., Lambdin, P.L., Grant, J.F., Merten, P., 2016. Thousand cankers disease complex: a forest health issue that threatens *Juglans* species across the US. Forests 7, 260.

Davydenko, K., Vasaitis, R., Elfstrand, M., Baturkin, D., Meshkova, V., Menkis, A., 2021. Fungal communities vectored by *Ips sexdentatus* in declining *Pinus sylvestris* in Ukraine: focus on occurrence and pathogenicity of ophiostomatoid species. Insects 12, 1119.

Davydenko, K., Vasaitis, R., Menkis, A., 2017. Fungi associated with *Ips acuminatus* (Coleoptera: Curculionidae) in Ukraine with a special emphasis on pathogenicity of ophiostomatoid species. Eur. J. Entomol. 114, 77-85.

Davydenko, K., Vasaitis, R., Meshokova, V., Menkis, A., 2014. Fungi associated with the redhaired bark beetle, *Hylurgus ligniperda* (Coleoptera: Curculionidae) in the forest-steppe zone in eastern Ukraine. Eur J Entomol 111, 561-565.

Deka, D., Jha, D.K., 2018. Optimization of culture parameters for improved production of bioactive metabolite by endophytic *Geosmithia pallida* (KU693285) isolated from *Brucea mollis* Wall ex. Kurz, an endangered medicinal plant. Kurz, an endangered medicinal plant. J Pure Appl Microbiol 12, 1205-1213.

Dighton, J., Walsh, E., Groben, G., Zhang, N., 2021. Influence of southern pine beetle on fungal communities of wood and bark decomposition of coarse woody debris in the New Jersey pine barrens. Forestry Research 1, 17.

Doehlemann, G., Ökmen, B., Zhu, W., Sharon, A., 2017. Plant pathogenic fungi. Microbiology spectrum 5, 5.1. 14.

Dohet, L., Grégoire, J.-C., Berasategui, A., Kaltenpoth, M., Biedermann, P.H.W., 2016. Bacterial and fungal symbionts of parasitic *Dendroctonus* bark beetles. FEMS Microbiol. Ecol. 92, fiw129.

Dori-Bachash, M., Avrahami-Moyal, L., Protasov, A., Mendel, Z., Freeman, S., 2015. The occurrence and pathogenicity of *Geosmithia* spp. and common blue-stain fungi associated with pine bark beetles in planted forests in Israel. Eur. J. Plant Pathol. 143, 627-639.

Flieger, M., Stodulková, E., Kolařík, M., Šulc, M., Kuzma, M., 2009. Submerzní kmeny *Geosmithia* sp. MK385B (CCM 8366) a *Geosmithia* sp. MK583B (CCM 8365), směs antrachinonových barviv jimi produkovaná a způsob její produkce, in: Institute of Microbiology, A. (Ed.), Czech Republic.

Foit, J., 2010. Distribution of early-arriving saproxylic beetles on standing dead Scots pine trees. Agric. For. Entomol. 12, 133-141.

Frascella, A., Bettini, P.P., Kolařík, M., Comparini, C., Pazzagli, L., Luti, S., Scala, F., Scala, A., 2014. Interspecific variability of class II hydrophobin GEO1 in the genus *Geosmithia*. Fungal Biol 118, 862-871.

Ganapaty, S., Thomas, P.S., Fotso, S., Laatsch, H., 2004. Antitermitic quinones from *Diospyros sylvatica*. Phytochemistry 65, 1265-1271.

Gao, L., Li, Y., Wang, Z.X., Zhao, J., Hulcr, J., Wang, J.G., Li, Y.Z., Ju, R.T., 2021. Biology and associated fungi of an emerging bark beetle pest, the sweetgum inscriber *Acanthotomicus suncei* (Coleoptera: Curculionidae). J. Appl. Entomol. 145, 508-517.

Gazis, R., Poplawski, L., Klingeman, W., Boggess, S.L., Trigiano, R.N., Graves, A.D., Seybold, S.J., Hadziabdic, D., 2018. Mycobiota associated with insect galleries in walnut with thousand cankers disease reveals a potential natural enemy against Geosmithia morbida. Fungal Biol 122, 241-253.

Giordano, A., Di Landro, F., De Carolis, E., Criscuolo, M., Dragonetti, G., Fianchi, L., Pagano, L., 2021. Disseminated *Geosmithia argillacea* infection in a patient with Ph-positive acute lymphoblastic leukemia. Case report and literature review. Journal of Fungi 7, 778.

Giraud, S., Favennec, L., Bougnoux, M.-E., Bouchara, J.-P., 2013. Rasamsonia argillacea species complex: taxonomy, pathogenesis and clinical relevance. Future Microbiology 8, 967-978. Gonzalez-Escobedo, R., Briones-Roblero, C.I., López, M.F., Rivera-Orduña, F.N., Zúñiga, G., 2019. Changes in the microbial community of *Pinus arizonica* saplings after being colonized by the bark beetle *Dendroctonus rhizophagus* (Curculionidae: Scolytinae). Microb. Ecol. 78, 102-112. Grobárová, V., Vališ, K., Talacko, P., Pavlů, B., Hernychová, L., Nováková, J., Stodůlková, E., Flieger, M., Novák, P., Černý, J., 2016. Quambalarine B, a Secondary Metabolite from Quambalaria cyanescens with Potential Anticancer Properties. J. Nat. Prod. 79, 2304-2314.

Hadj Taieb, K., Gharsallah, H., Ksentini, I., Schuster, C., Fernández-Bravo, M., Garrido-Jurado,

I., Quesada-Moraga, E., Leclerque, A., Triki, M., Ksantini, M., 2019. Phytopathogenic and antagonistic potentialities of fungi associated with pistachio bark beetle, *Chaetoptelius vestitus* (Coleoptera, Curculionidae), infesting pistachio (*Pistacia vera*) in Tunisia. J. Appl. Microbiol. 126, 1821-1834.

Hadziabdic, D., Vito, L.M., Windham, M.T., Pscheidt, J.W., Trigiano, R.N., Kolarik, M., 2014a. Genetic differentiation and spatial structure of *Geosmithia morbida*, the causal agent of thousand cankers disease in black walnut (*Juglans nigra*). Curr. Genet. 60, 75-87.

Hadziabdic, D., Windham, M., Baird, R.E., Vito, L., Cheng, Q., Grant, J., Lambdin, P., Wiggins, G., Windham, A., Merten, P., 2014b. First report of *Geosmithia morbida* in North Carolina: the pathogen involved in thousand cankers disease of black walnut. Plant Dis. 98, 992.

Hanzi, M., Cochard, B., Chablais, R., Crovadore, J., Lefort, F., 2016. First report of *Geosmithia langdonii* and *Geosmithia* spp. isolated from a decaying elm (*Ulmus minor*) in Geneva, Switzerland. Folia Forestalia Polonica. Series A. Forestry 58, 96-102.

Hefty, A.R., Aukema, B.H., Venette, R.C., Coggeshall, M.V., McKenna, J.R., Seybold, S.J., 2018. Reproduction and potential range expansion of walnut twig beetle across the Juglandaceae. Biol. Invasions 20, 2141-2155.

Hilker, M., Köpf, A., 1994. Evaluation of the palatability of chrysomelid larvae containing anthraquinones to birds. Oecologia 100, 421-429.

Hofstetter, R.W., Dinkins-Bookwalter, J., Davis, T.S., Klepzig, K.D., 2015. Symbiotic associations of bark beetles, Bark Beetles. Elsevier, pp. 209-245.

Houbraken, J., Spierenburg, H., Frisvad, J.C., 2012. *Rasamsonia*, a new genus comprising thermotolerant and thermophilic *Talaromyces* and *Geosmithia* species. Antonie Van Leeuwenhoek 101, 403–421.

Hu, X., Li, M., Chen, H., 2015. Community structure of gut fungi during different developmental stages of the Chinese white pine beetle (*Dendroctonus armandi*). Scientific Reports 5, 8411.

Huang, Y.-T., Kolarik, M., Kasson, M., Hulcr, J., 2017. Two new *Geosmithia* species in *G. pallida* species complex from bark beetles in eastern USA. Mycologia 109, 790-803.

Huang, Y.T., Skelton, J., Johnson, A.J., Kolařík, M., Hulcr, J., 2019. *Geosmithia* species in southeastern USA and their affinity to beetle vectors and tree hosts. Fungal Ecol 39, 168-183.

Iwamoto, S., Tokumasu, S., Suyama, Y., Kakishima, M., 2002. Molecular phylogeny of four selected species of the strictly anamorphic genus *Thysanophora* using nuclear ribosomal DNA sequences. Mycoscience 43, 169-180.

Jagadeeswaran, G., Veale, L., Mort, A.J., 2021. Do lytic polysaccharide monooxygenases aid in plant pathogenesis and herbivory? Trends Plant Sci. 26, 142-155.

Jankowiak, R., 2006a. Fungi associated with *Tomicus piniperda* in Poland and assessment of their virulence using Scots pine seedlings. Annals of Forest Science 63, 801-808.

Jankowiak, R., 2006b. Mycobiota associated with *Hylurgops palliatus* (Gyll.) on *Pinus sylvestris* L. in Poland. Acta Societatis Botanicorum Poloniae 75, 333-338.

Jankowiak, R., 2008. Fungi associated with *Tomicus minor* on *Pinus sylvestris* in Poland and their succession into the sapwood of beetle-infested windblown trees. Canadian Journal of Forest Research 38, 2579-2588.

Jankowiak, R., Bilanski, P., 2018. *Geosmithia* species associated with fir-infesting beetles in Poland. Acta Mycol. 53, 1115.

Jankowiak, R., Bilański, P., 2007. Fungal flora associated with *Tomicus piniperda* L. in an area close to a timber yard in southern Poland. J. Appl. Entomol. 131, 579-584.

Jankowiak, R., Hilszczański, J., 2005. Ophiostomatoid fungi associated with *Ips typographus* (L.) on *Picea abies* [(L.) H. Karst.] and *Pinus sylvestris* L. in north-eastern Poland. Acta Societatis Botanicorum Poloniae 74, 345-350.

Jankowiak, R., Kolarik, M., 2010. Fungi associated with the fir bark beetle *Cryphalus piceae* in Poland. For. Pathol. 40, 133-144.

Jankowiak, R., Kolarik, M., Bilanski, P., 2014. Association of *Geosmithia* fungi (Ascomycota: Hypocreales) with pine- and spruce-infesting bark beetles in Poland. Fungal Ecol 11, 71-79.

Jankowiak, R., Rossa, R., 2007. Filamentous fungi associated with *Monochamus galloprovincialis* and *Acanthocinus aedilis* (Coleoptera: Cerambycidae) in Scots pine. Polish Botanical Journal 52, 140-143.

Jankowiak, R., Rossa, R., 2008. Associations between *Pityogenes bidentatus* and fungi in young managed Scots pine stands in Poland. For. Pathol. 38, 169-177.

Juan Alfredo, H.-G., Gerardo, C.-R., Nallely Guadalupe, A.-O., Lourdes, V.-T., César, H.-R., Francisco, A.-T., 2020. Phylogenetic Position of Geosmithia spp.(Hypocreales) Living in Juniperus spp. Forests (Cupressaceae) with Bark Beetles of Phloeosinus spp.(Scolytinae) from the Northeast of Mexico. Forests 11, 1142.

Juzwik, J., Yang, A., Heller, S., Moore, M., Chen, Z., White, M., Wantuch, H., Ginzel, M., Mack, R., 2021. Vacuum steam treatment effectiveness for eradication of the Thousand Cankers Disease vector and pathogen in logs from diseased walnut trees. J. Econ. Entomol. 114, 100-111.

Kim, Y.J., Duraisamy, K., Jeong, M.-H., Park, S.-Y., Kim, S., Lee, Y., Nguyen, V.T., Yu, N.H., Park, A.R., Kim, J.-C., 2021. Nematicidal activity of grammicin biosynthesis pathway intermediates in *Xylaria grammica* KCTC 13121BP against *Meloidogyne incognita*. Molecules 26, 4675.

Kirschner, R., 1998. Diversität mit Borkenkäfern assoziierter filamentöser Mikropilze, Fakultät für Biologie. Eberhard-Karls Universität, Tübingen, p. 573.

Kirschner, R., 2001. Diversity of filamentous fungi in bark beetle galleries in Central Europe, in: Misra, J.K., Horn, B.W. (Eds.), In *Trichomycetes* and other fungal groups: Professor Robert W. Lichtwardt commemoration volume Science Publishers, Enfield (NH), USA, pp. 175-196.

Kirschner, R., Sampaio, J.P., Gadanho, M., Weiss, M., Oberwinkler, F., 2001. *Cuniculitrema polymorpha* (Tremellales, gen. nov. and sp. nov.), a heterobasidiomycete vectored by bark beetles, which is the teleomorph of *Sterigmatosporidium polymorphum*. Antonie Van Leeuwenhoek 80, 149-161.

Kolařík, M., Freeland, E., Utley, C., Tisserat, N., 2011. *Geosmithia morbida* sp. nov., a new phytopathogenic species living in symbiosis with the walnut twig beetle (*Pityophthorus juglandis*) on *Juglans* in USA. Mycologia 103, 325-332.

Kolařík, M., Hulcr, J., Kirkendall, L.R., 2015. New species of *Geosmithia* and *Graphium* associated with ambrosia beetles in Costa Rica. Czech Mycol 67, 29-35.

Kolařík, M., Hulcr, J., Tisserat, N., De Beer, W., Kostovčík, M., Kolaříková, Z., Seybold, S.J., Rizzo, D.M., 2017. *Geosmithia* associated with bark beetles and woodborers in the western USA: taxonomic diversity and vector specificity. Mycologia 109, 185-199.

Kolařík, M., Jankowiak, R., 2013. Vector affinity and diversity of *Geosmithia* fungi living on subcortical insects inhabiting Pinaceae species in central and Northeastern Europe. Microb. Ecol. 66, 682-700.

Kolařík, M., Kirkendall, L.R., 2010. Evidence for a new lineage of primary ambrosia fungi in *Geosmithia* Pitt (Ascomycota: Hypocreales). Fungal Biol 114, 676-689.

Kolařík, M., Kostovčík, M., Pažoutová, S., 2007. Host range and diversity of the genus *Geosmithia* (Ascomycota: Hypocreales) living in association with bark beetles in the Mediterranean area. Mycol. Res. 111, 1298-1310.

Kolařík, M., Kubátová, A., Čepička, I., Pažoutová, S., Šrůtka, P., 2005. A complex of three new white-spored, sympatric, and host range limited *Geosmithia* species. Mycol. Res. 109, 1323-1336.

Kolařík, M., Kubátová, A., Hulcr, J., Pažoutová, S., 2008. *Geosmithia* fungi are highly diverse and consistent bark beetle associates: evidence from their community structure in temperate Europe. Microb. Ecol. 56, 198-199.

Kolařík, M., Kubátová, A., Pažoutová, S., Šrůtka, P., 2004. Morphological and molecular characterisation of *Geosmithia putterillii*, *G. pallida* comb. nov. and *G. flava* sp. nov., associated with subcorticolous insects. Mycol. Res. 108, 1053-1069.

Kolařík, M., Sláviková, E., Pažoutová, S., 2006. The taxonomic and ecological characterization of the clinically important heterobasiodiomycete *Fugomyces cyanescens* and its association with bark beetles. Czech Mycol 58, 81-98.

Kolařík, M., Vohník, M., 2017. When the ribosomal DNA does not tell the truth: The case of the taxonomic position of Pekloderma argillaceum, an ericoid mycorrhizal fungus residing among hymenochaetales. Fungal Biol.

Kolařík, M., Wei, I., Hsieh, S.-Y., Piepenbring, M., Kirschner, R., 2021. Nucleotide composition bias of rDNA sequences as a source of phylogenetic artifacts in *Basidiomycota*—a case of a new lineage of a uredinicolous *Ramularia*-like anamorph with affinities to *Ustilaginomycotina*. Mycological Progress 20, 1553-1571.

Kubátová, A., Kolařík, M., Prášil, K., Novotný, D., 2004. Bark beetles and their galleries: Well-known niche for little known fungi, case of *Geosmithia*. Czech Mycol 55, 1–18.

Kubátová, A., Novotný, D., Prášil, K., 1999. Bělokaz dubový jako přenašeč mikroskopických hub [Oak bark beetle as vector of microscopic fungi], in: Jankovský, L., Krejčíř, R., Antonín, V. (Eds.), Houby a les [Fungi and forest], Brno, pp. 235-236.

Kula, E., Zabecki, W., 2000. Vliv některých růstových a porostních faktorů na obsazení smrku kambioxylofágy (Effect of selected growth and vegetations factors on distribution of cambioxylophages on spruce). Lesnická práce 6, 257-260.

Kumar, C.G., Mongolla, P., Pombala, S., Bandi, S., Babu, K., Ramakrishna, K., 2017. Biological evaluation of 3-hydroxybenzyl alcohol, an extrolite produced by *Aspergillus nidulans* strain KZR-132. J. Appl. Microbiol. 122, 1518-1528.

Labuda, R., Tancinová, D., 2006. Fungi recovered from Slovakian poultry feed mixtures and their toxinogenity. Ann. Agric. Environ. Med. 13, 193-200.

Lee, S., Kim, J.J., Breuil, C., 2006. Diversity of fungi associated with the mountain pine beetle, *Dendroctonus ponderosae* and infested lodgepole pines in British Columbia. Fungal Diversity 22, 91-105.

Li, Y., Bateman, C., Skelton, J., Wang, B., Black, A., Huang, Y.-T., Gonzalez, A., Jusino, M.A., Nolen, Z.J., Freeman, S., Mendel, Z., Kolařík, M., Knížek, M., Park, J.-H., Sittichaya, W., Pham, T.-H., Ito, S.-i., Torii, M., Gao, L., Johnson, A.J., Lu, M., Sun, J., Zhang, Z., Adams, D.C., Hulcr, J., 2022. Preinvasion assessment of exotic bark beetle-vectored fungi to detect tree-killing pathogens. Phytopathology 112, 261-270.

Liang, W.-L., Le, X., Li, H.-J., Yang, X.-L., Chen, J.-X., Xu, J., Liu, H.-L., Wang, L.-Y., Wang, K.-T., Hu, K.-C., 2014. Exploring the chemodiversity and biological activities of the secondary metabolites from the marine fungus *Neosartorya pseudofischeri*. Mar. Drugs 12, 5657-5676.

Lim, Y.W., Kim, J.J., Lu, M., Breuil, C., 2005. Determining fungal diversity on *Dendroctonus ponderosae* and *Ips pini* affecting lodgepole pine using cultural and molecular methods. Fungal Diversity 19, 79-94.

Linnakoski, R., de Beer, Z.W., Rousi, M., Niemelä, P., Pappinen, A., Wingfield, M.J., 2008. Fungi, including *Ophiostoma karelicum* sp. nov., associated with *Scolytus ratzeburgi* infesting birch in Finland and Russia. Mycol. Res. 112, 1475-1488.

Lynch, S.C., Wang, D.H., Mayorquin, J.S., Rugman-Jones, P., Stouthamer, R., Eskalen, A., 2014. First report of *Geosmithia pallida* causing foamy bark canker, a new disease on coast live

oak (*Quercus agrifolia*) in association with *Pseudopityophthorus pubipennis* (western oak bark beetle, Coleoptera: Curculionidae: Scolytinae) in California. Plant Dis. 98, 1276-1276.

Machingambi, N.M., Roux, J., Dreyer, L.L., Roets, F., 2014. Bark beetles (Curculionidae: Scolytinae), their phoretic mites (Acari) and associated *Geosmithia* species (Ascomycota: Hypocreales) from *Virgilia* trees in South Africa. Fungal Biol 118, 472-483.

Malak, L., Bishay, D., Abdel-Baky, A., Moharram, A., Cutler, S.J., Ross, S.A., 2013a. New secondary metabolites from *Geosmithia lavendula*. Planta Medica 79, P51.

Malak, L.G., Bishay, D.W., Abdel-baky, A.M., Moharram, A.M., Cutler, S.J., Ross, S.A., 2013b. Bioactive secondary metabolites from *Geosmithia langdonii*. Planta Medica 79, PL7.

Malak, L.G., Bishay, D.W., Abdel-Baky, A.M., Moharram, A.M., Cutler, S.J., Ross, S.A., 2013c. New anthraquinone derivatives from *Geosmithia lavendula*. Natural product communications 8, 1934578X1300800215.

Malak, L.G., Ibrahim, M.A., Bishay, D.W., Abdel-Baky, A.M., Moharram, A.M., Tekwani, B., Cutler, S.J., Ross, S.A., 2014. Antileishmanial metabolites from *Geosmithia langdonii*. J. Nat. Prod. 77, 1987-1991.

Malak, L.G., Ibrahim, M.A., Moharram, A.M., Pandey, P., Tekwani, B., Doerksen, R.J., Ferreira, D., Ross, S.A., 2018. Antileishmanial carbasugars from *Geosmithia langdonii*. J. Nat. Prod. 81, 2222-2227.

Marchioro, M., Faccoli, M., 2022. Dispersal and colonization risk of the Walnut Twig Beetle, *Pityophthorus juglandis*, in southern Europe. J. Pest Sci. 95, 303-313.

Matoušková, Z., 2020. ledování vlivu quambalarinu B na aminokyselinový metabolismus leukemických buněčných linií, Přírodovědecká fakulta, Katedra biochemie. Univerzita Karlova, Praha.

McPherson, B.A., Erbilgin, N., Bonello, P., Wood, D.L., 2013. Fungal species assemblages associated with *Phytophthora ramorum*-infected coast live oaks following bark and ambrosia beetle colonization in northern California. For. Ecol. Manage. 291, 30-42.

Meshram, V., Sharma, G., Maymon, M., Protasov, A., Mendel, Z., Freeman, S., 2022. Symbiosis and pathogenicity of *Geosmithia* and *Talaromyces* spp. associated with the cypress bark beetles *Phloeosinus* spp. and their parasitoids. Environ. Microbiol. 24, 3369–3389.

Montecchio, L., Faccoli, M., 2014. First record of thousand cankers disease *Geosmithia morbida* and walnut twig beetle *Pityophthorus juglandis* on *Juglans nigra* in Europe. Plant Dis. 98, 696-696.

Montecchio, L., Fanchin, G., Berton, V., Scattolin, L., 2015. Vegetative incompatibility and potential involvement of a mycovirus in the Italian population of *Geosmithia morbida*. Phytopathologia Mediterranea 54, 465-476.

Morales-Rodríguez, C., Sferrazza, I., Aleandri, M.P., Dalla Valle, M., Speranza, S., Contarini, M., Vannini, A., 2021. The fungal community associated with the ambrosia beetle *Xylosandrus compactus* invading the Mediterranean maquis in central Italy reveals high biodiversity and suggests environmental acquisitions. Fungal Biol 125, 12-24.

Moricca, S., Bracalini, M., Benigno, A., Ghelardini, L., Furtado, E.L., Marino, C.L., Panzavolta, T., 2020. Observations on the non-native Thousand Cankers Disease of walnut in Europe's southernmost outbreak. Global Ecology and Conservation, e01159.

Muñoz-Adalia, E.J., Sanz-Ros, A.V., Flores-Pacheco, J.A., Hantula, J., Diez, J.J., Vainio, E.J., Fernández, M., 2017. *Sydowia polyspora* dominates fungal communities carried by two *Tomicus Species* in pne plantations threatened by *Fusarium circinatum*. Forests 8, 127.

O'Gorman, C.M., Fuller, H.T., Dyer, P.S., 2008. Discovery of a sexual cycle in the opportunistic fungal pathogen *Aspergillus fumigatus*. Nature 457, 471-474.

Ogawa, H., Sugiyama, J., 2000. Evolutionary relationships of the cleistothecial genera with *Penicillium, Geosmithia, Merimbla* and *Sarophorum* anamorphs as inferred from 18S rDNA sequence divergence, in: Samson, R.A., Pitt, J.I. (Eds.), Integration of modern taxonomic methods for *Penicillium* and *Aspergillus* classification. Harwood, Amsterdam, pp. 149-161.

Ogawa, H., Yoshimura, A., Sugiyama, J., 1997. Polyphyletic origins of species of the anamorphic genus *Geosmithia* and the relationships of the cleistothecial genera: evidence from 18S, 5S and 28S rDNA sequence analyses. Mycologia 89, 756-771.

Pepori, A.L., Bettini, P.P., Comparini, C., Sarrocco, S., Bonini, A., Frascella, A., Ghelardini, L., Scala, A., Vannacci, G., Santini, A., 2018. *Geosmithia-Ophiostoma*: a new fungus-fungus association. Microb. Ecol. 75, 632-646.

Pepori, A.L., Kolařík, M., Bettini, P.P., Vettraino, A.M., Santini, A., 2015. Morphological and molecular characterisation of *Geosmithia* species on European elms. Fungal Biol 119, 1063-1074.

Persson, Y., Vasaitis, R., Långström, B., Öhrn, P., Ihrmark, K., Stenlid, J., 2009. Fungi vectored by the bark beetle *Ips typographus* following hibernation under the bark of standing trees and in the forest litter. Microb. Ecol. 58, 651-659.

Peterson, S.W., 2000. Phylogenetic analysis of *Penicillium* species based on ITS and LSU-rDNA nucleotide sequences, in: Samson, R.A., Pitt, J.I. (Eds.), Integration of modern taxonomic

methods for *Penicillium* and *Aspergillus* classification. Harwood Academic Publishers, Reading, pp. 163-178.

Pietsch, G.M., Gazis, R., Klingeman, W.E., Huff, M.L., Staton, M.E., Kolařík, M., Hadziabdic, D., 2022. Characterization and microsatellite marker development for a common bark and ambrosia beetle associate, *Geosmithia obscura*. MicrobiologyOpen 11, e1286.

Pitt, J.I., 1979. The genus *Penicillium* and its teleomorphic states *Eupenicillium* and *Talaromyces*. Academic Press, London.

Pitt, J.I., Hocking, A.D., 1985. Interfaces among genera related to *Aspergillus* and *Penicillium*. Mycologia 77, 810-824.

Pitt, J.I., Hocking, A.D., 2009. Fungi and food spoilage, 3rd ed. Springer, Dordrecht, The Netherlands.

Pitt, J.I., Samson, R.A., 1993. Species names in current use in the *Trichocomaceae* (Fungi: Eurotiales). Regnum vegetabile 128, 13-57.

Pitt, J.I., Samson, R.A., Frisvad, J.C., 2000. List of accepted species and their synonyms in the family *Trichocomaceae*., in: Samson, R.A., Pitt, J.I. (Eds.), Integration of modern taxonomic methods for *Penicillium* and *Aspergillus* classification. Harwood Academic Publishers, pp. 9-49.

Plishka, M.J.R., Tsuneda, A., Currah, R.S., 2009. Morphology and development of *Nigrosabulum globosum*, a cleistothecial coprophile in the Bionectriaceae (Hypocreales). Mycol. Res. 113, 815-821.

Poche, R.M., 1998. Development of a new bird repellent, Flight Control, Proceedings of the Vertebrate Pest Conference, pp. 337-334.

Postner, M., 1974. Scolytidae (= Ipidae), Borkenkaäfer, in: Schwenke, W. (Ed.), Die Forstschaädlinge Europas, II Käfer. Verlag Paul Parey, Hamburg, Berlin, pp. 334-482.

Prochazkova, E., Kucherak, O., Stodůlková, E., Tošner, Z.k., Cisarova, I., Flieger, M., Kolarik, M., Baszczynski, O., 2020. NMR Structure Elucidation of Naphthoquinones from Quambalaria cyanescens. J. Nat. Prod. 84, 46-55.

Ramirez, C., 1982. Manual and Atlas of the Penicillia. Elsevier Biomedical Press, Amsterdam etc.

Raper, K.B., Thom, V.C., 1949. A manual of the Penicillia. The Williams & Wilkins Comp., Baltimore.

Ratnaweera, P., de Silva, E.D., Wijesundera, R.L., Andersen, R.J., 2016. Antimicrobial constituents of *Hypocrea virens*, an endophyte of the mangrove-associate plant *Premna serratifolia* L. J. Natl. Sci. Found. Sri Lanka 44, 43–51.

Rossman, A.Y., Samuels, G.J., Rogerson, C.T., Lowen, R., 1999. Genera of *Bionectriaceae*, *Hypocreaceae* and *Nectriaceae* (Hypocreales, Ascomycetes). Studies in Mycology 42, 1-248.

Sakalidis, M.L., Hardy, G.E.S., Burgess, T.I., 2011. Endophytes as potential pathogens of the baobab species *Adansonia gregorii*: a focus on the Botryosphaeriaceae. Fungal Ecol 4, 1-14.

Salvatore, M.M., Alves, A., Andolfi, A., 2020. Secondary metabolites of *Lasiodiplodia theobromae*: distribution, chemical diversity, bioactivity, and implications of their occurrence. Toxins 12, 457.

Scala, A., Comparini, C., Tegli, S., Scala, F., 2007. A non-Ophiostoma fungus expresses the gene encoding the hydrophobin cerato-ulmin. J. Plant Pathol. 89, 233-240.

Schroers, H.J., Geldenhuis, M.M., Wingfield, M.J., Schoeman, M.H., Yen, Y.F., Shen, W.C., Wingfield, B.D., 2005. Classification of the guava wilt fungus *Myxosporium psidii*, the palm pathogen *Gliocladium vermoesenii* and the persimmon wilt fungus *Acremonium diospyri* in *Nalanthamala*. Mycologia 97, 375-395.

Schuelke, T.A., Wu, G., Westbrook, A., Woeste, K., Plachetzki, D.C., Broders, K., MacManes, M.D., 2017. Comparative genomics of pathogenic and nonpathogenic beetle-vectored fungi in the genus *Geosmithia*. Genome Biol Evol 9, 3312-3327.

Seabright, K.W., Myers, S.W., Fraedrich, S.W., Mayfield III, A.E., Warden, M.L., Taylor, A., 2019. Methyl bromide fumigation to eliminate thousand cankers disease causal agents from black walnut. For. Sci. 65, 452-459.

Silva, X., Terhonen, E., Sun, H., Kasanen, R., Heliövaara, K., Jalkanen, R., Asiegbu, F.O., 2015. Comparative analyses of fungal biota carried by the pine shoot beetle (*Tomicus piniperda* L.) in northern and southern Finland. Scand. J. For. Res. 30, 497-506.

Sitz, R.A., Luna, E.K., Caballero, J.I., Tisserat, N.A., Cranshaw, W.S., Stewart, J.E., 2017. Virulence of genetically distinct *Geosmithia morbida* isolates to black walnut and their response to coinoculation with *Fusarium solani*. Plant Dis. 101, 116-120.

Sitz, R.A., Luna, E.K., Ibarra Caballero, J., Tisserat, N.A., Cranshaw, W.S., McKenna, J.R., Stolz, J., Stewart, J.E., 2021. Eastern black walnut (*Juglans nigra* L.) originating from native range varies in their response to inoculation with *Geosmithia morbida*. Front For Glob Change 4, 12.

Six, D.L., 2013. The bark beetle holobiont: why microbes matter. J. Chem. Ecol. 39, 989-1002.

Six, D.L., 2020. Niche construction theory can link bark beetle-fungus symbiosis type and colonization behavior to large scale causal chain-effects. Current Opinion in Insect Science 39, 27-34.

Six, D.L., Doug, S.W., de Beer, Z.W., Woolfolk, S.W., 2009. *Ambrosiella beaveri*, sp. nov., associated with an exotic ambrosia beetle, *Xylosandrus mutilatus* (Coleoptera: Curculionidae, Scolytinae), in Mississippi, USA. Antonie Van Leeuwenhoek 96, 17-29.

Six, D.L., Wingfield, M.J., 2011. The role of phytopathogenicity in bark beetle-fungus symbioses: a challenge to the classic paradigm. Annu. Rev. Entomol. 56, 255-272.

Skelton, J., Jusino, M.A., Li, Y., Bateman, C., Thai, P.H., Wu, C., Lindner, D.L., Hulcr, J., 2018. Detecting symbioses in complex communities: the fungal symbionts of bark and ambrosia beetles within Asian pines. Microb. Ecol. 76, 839-850.

Smith, S.M., Cognato, A.I., 2014. A taxonomic monograph of Nearctic *Scolytus* Geoffroy (Coleoptera, Curculionidae, Scolytinae). ZooKeys 450, 1-182.

Solheim, H., 1991. Oxygen deficiency and spruce resin inhibition of growth of blue stain fungi associated with *Ips typographus*. Mycol. Res. 95, 1387-1392.

Stackhouse, T., Boggess, S.L., Hadziabdic, D., Trigiano, R.N., Ginzel, M.D., Klingeman, W.E., 2021. Conventional gel electrophoresis and TaqMan probes enable rapid confirmation of Thousand Cankers Disease from diagnostic samples. Plant Dis. 105, 3171-3180.

Stodůlková, E., Císařová, I., Kolařík, M., Chudiččková, M., Novák, P., Man, P., Kuzma, M., Pavlu, B., Černý, J., Flieger, M., 2015. Biologically active metabolites produced by the Basidiomycete Quambalaria cyanescens. PLoS ONE 10.

Stodůlková, E., Kolařík, M., Křesinová, Z., Kuzma, M., Šulc, M., Man, P., Novák, P., Maršík, P., Landa, P., Olšovská, J., Chudíčková, M., Pažoutová, S., Černý, J., Bella, J., Flieger, M., 2009. Hydroxylated anthraquinones produced by *Geosmithia* species. Folia Microbiol. 54, 179–187.

Stodulkova, E., Man, P., Kolarik, M., Flieger, M., 2010. High-performance liquid chromatography-off line mass spectrometry analysis of anthraquinones produced by Geosmithia lavendula. J. Chromatogr. 1217, 6296-6302.

Stodůlková, E., Man, P., Kolařík, M., Flieger, M., 2010. High-performance liquid chromatography - off line mass spectrometry analysis of anthraquinones produced by *Geosmithia lavendula*. J. Chromatogr. 1217, 6296-6302.

Stodulkova, E., Sulc, M., Cisarova, I., Novak, P., Kolarik, M., Flieger, M., 2008. Production of (+)-globulol needle crystals on the surface mycelium of Quambalaria cyanescens. Folia Microbiol. 53, 15-22.

Stolk, A.C., Samson, R.A., 1986. A new taxonomic scheme for *Penicillium* anamorphs, Advances in *Penicillium* and *Aspergillus* systematics. Springer, pp. 163-192.

Strzałka, B., Kolařík, M., Jankowiak, R., 2021. *Geosmithia* associated with hardwood-infesting bark and ambrosia beetles, with the description of three new species from Poland. Antonie Van Leeuwenhoek 114, 169-194.

Sun, Y., Takada, K., Takemoto, Y., Yoshida, M., Nogi, Y., Okada, S., Matsunaga, S., 2012. Gliotoxin analogues from a marine-derived fungus, *Penicillium* sp., and their cytotoxic and histone methyltransferase inhibitory activities. J. Nat. Prod. 75, 111-114.

Sun, Z.-H., Gu, J., Ye, W., Wen, L.-X., Lin, Q.-B., Li, S.-N., Chen, Y.-C., Li, H.-H., Zhang, W.-M., 2018. Geospallins A–C: New thiodiketopiperazines with inhibitory activity against angiotensin-converting enzyme from a deep-Sea-derived Fungus *Geosmithia pallida* FS140. Mar. Drugs 16, 464.

Tisserat, N., Cranshaw, W., Leatherman, D., Utley, C., Alexander, K., 2009. Black walnut mortality in Colorado caused by the walnut twig beetle and Thousand Cankers Disease. Plant Health Progress 10, 1-10.

Vališ, K., Grobárová, V., Hernychová, L., Bugáňová, M., Kavan, D., Kalous, M., Černý, J., Stodůlková, E., Kuzma, M., Flieger, M., 2017. Reprogramming of leukemic cell metabolism through the naphthoquinonic compound Quambalarine B. Oncotarget 8, 103137.

van Dyk, M., Spies, C.F.J., Mostert, L., van der Rijst, M., du Plessis, I.L., Moyo, P., van Jaarsveld, W.J., Halleen, F., 2021. Pathogenicity testing of fungal isolates associated with olive trunk diseases in South Africa. Plant Dis. 105, 4060-4073.

Veselská, T., Kolařík, M., 2015. Application of flow cytometry for exploring the evolution of *Geosmithia* fungi living in association with bark beetles: the role of conidial DNA content. Fungal Ecol 13, 83-92.

Veselská, T., Skelton, J., Kostovčík, M., Hulcr, J., Baldrian, P., Chudíčková, M., Cajthaml, T., Vojtová, T., Garcia-Fraile, P., Kolařík, M., 2019. Adaptive traits of bark and ambrosia beetle-associated fungi. Fungal Ecol 41, 165-176.

Veselská, T., Svoboda, J., Růžičková, Z., Kolařík, M., 2014. Application of flow cytometry for genome size determination in Geosmithia fungi: A comparison of methods. Cytometry Part A.

- Vitale, S., Luongo, L., Barzanti, G.P., Binazzi, F., Petrucci, M., Galli, M., Pennacchio, F., Francardi, V., 2021. First report of *Geosmithia pallida* and *G. langdonii* associated with *Liparthrum colchicum* in Central Italy. J. Zool. 104, 167-170.
- Wang, M., Kornsakulkarn, J., Srichomthong, K., Feng, T., Liu, J.-K., Isaka, M., Thongpanchang, C., 2019. Antimicrobial anthraquinones from cultures of the ant pathogenic fungus Cordyceps morakotii BCC 56811. The Journal of Antibiotics 72, 141-147.
- Williams, G.M., Ginzel, M.D., 2021. Competitive advantage of *Geosmithia morbida* in low-moisture wood may explain historical outbreaks of Thousand Cankers Disease and predict the future fate of *Juglans nigra* within its native range. Front For Glob Change 4, 725066.
- Wood, S.L., 1982. The bark and ambrosia beetles of North and Central America (Coleoptera: Scolytidae), a taxonomic monograph. Great Basin Naturalist Memoirs 6, 1-1359.
- Wright, E., 1938. Further investigations of brown-staining fungi associated with engraver beetles (*Scolytus*) in white fir. J. Agric. Res. 57, 759-773.
- Wu, Y.-M., Xu, J.-J., Wang, H.-F., Zhang, T.-Y., 2013. *Geosmithia tibetensi*s sp. nov. and new *Gibellulopsis* and *Scopulariopsis* records from Qinghai-Tibet. Mycotaxon 125, 59-64.
- Yaguchi, T., Miyadoh, S., Udagawa, S., 1993. *Chromocleista*, a new cleistothecial genus with a *Geosmithia* anamorph. Trans Mycol Soc Japan 34, 101-108.
- Yaguchi, T., Someya, A., Udagawa, S., 1994. Two new species of *Talaromyces* from Taiwan and Japan. Mycoscience 35, 249-255.
- Yaguchi, T., Udagawa, S., Nishimura, K., 2005. *Geosmithia argillacea* is the anamorph of *Talaromyces eburneus* as a heat resistant fungus. Cryptogamie: Mycol. 26, 133-141.
- Zerillo, M.M., Caballero, J.I., Woeste, K., Graves, A.D., Hartel, C., Pscheidt, J.W., Tonos, J., Broders, K., Cranshaw, W., Seybold, S.J., 2014. Population structure of *Geosmithia morbida*, the causal agent of thousand cankers disease of walnut trees in the United States. PLoS ONE 9, e112847.
- Zhang, X., Li, Y., Si, H., Zhao, G., Kolařík, M., Hulcr, J., Jiang, X., Dai, M., Chang, R., 2022. *Geosmithia* species associated with bark beetles from China, with the description of nine new species. Frontiers in Microbiology 13, 820402.
- Zhu, H.-y., Zhang, D., Zhang, Q., Zhao, Y., He, Z.-m., Gao, Y.-g., Zhang, L.-x., 2018. 4-Hydroxybenzyl alcohol derivatives and their sedative—hypnotic activities. RSC advances 8, 19539-19550.

Zima, V., Radilová, K., Kožíšek, M., Albiñana, C.B., Karlukova, E., Brynda, J., Fanfrlík, J., Flieger, M., Hodek, J., Weber, J., 2020. Unraveling the anti-influenza effect of flavonoids: Experimental validation of luteolin and its congeners as potent influenza endonuclease inhibitors. European Journal of Medicinal Chemistry 208, 112754.

1.6 APPENDIX – SCIENTIFIC PAPER INCLUDED INTO THE HABILITATION THESIS

