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Laboulbeniales (Ascomycota) of the Boston Harbor Islands I: Species Parasitizing Coccinellidae and Staphylinidae, with Comments on Typification

Danny Haelewaters¹,* , Serena Y. Zhao¹, André De Kesel², Rebecca E. Handlin¹, Isabel R. Royer³, Brian D. Farrell⁴, and Donald H. Pfister¹

Abstract - This paper, based on a recent comprehensive sampling of insects, is the first report of Laboulbeniales from the New England region since the 1930s. We present 7 new records of laboulbenialean parasites on Staphylinidae (rove beetles) and Coccinellidae (lady beetles) from the Boston Harbor Islands National Recreation Area. These are Clonophoro-mycetes nipponicus Terada & I.I. Tav., Hesperomyces virescens Thaxt., Ilyomyces cf. mairei F. Picard, Laboulbenia philonthi Thaxt., Peyritschiella protea Thaxt., Stichomyces conosomatis Thaxt., and Teratomyces actobii Thaxt. One of these parasite species, C. nipponicus, has not been found previously outside of its type locality in Japan. Examination of Roland Thaxter’s 1891–1932 slides led to the designation of lectotypes for L. philonthi, P. protea, S. conosomatis, and T. actobii. The following synonymy is established: Teratomyces brevicaulis Thaxt. = T. actobii. In addition, we discovered new localities for H. virescens (from Canada, Cuba, Guatemala, and Japan) and L. philonthi (from Canada, Grenada, Panama, Trinidad, and Venezuela).

Introduction

Existing museum collections provide host specimens from a wide range of collection locations and dates for the study of arthropod ectoparasites. Collections made in the context of an all taxa biodiversity inventory (ATBI) provide sample populations from host collections that should exhibit minimal bias in parasite distribution and diversity. Parasite density for obligate ectoparasites can be determined because they are fairly easily observed on the exoskeleton of a given host; these parasites persist on dead individuals, and thus, historical collections of infected hosts serve as records of species relationships and population dynamics through time. They also provide a measure of parasite abundances and host usage across a single habitat or system. In this paper, we present fungal parasites of 2 host families collected in a survey of all arthropods in an island system.

The Order Laboulbeniales

The Laboulbeniales form an order of fungi (phylum Ascomycota, class Laboulbeniomycetes) that obligately parasitize arthropod hosts. They are

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ectoparasites, that is, they occur externally on various parts of the insect. Phylogenetic analyses of DNA sequences indicate that the class Laboulbeniomycetes, which also includes the order Pyxidiophorales, is sister to the Sordariomycetes—a clade that includes saprotrophs and pathogens of plants, animals, and fungi, and that generally produce perithecia (Blackwell and Malloch 1989, Schoch et al. 2009, Weir and Blackwell 2001).

Roland Thaxter, a professor at Harvard University (Cambridge, MA) from 1891 to 1932, undertook the first systematic study of the Laboulbeniales, and his 5 monographic volumes (1896, 1908, 1924, 1926, 1931) and many non-illustrated papers are still among the most-often cited papers in Laboulbeniales research. Despite Thaxter’s intense local collecting activities, there is no evidence that he explored the sites now included in the Boston Harbor Islands (BHI) National Recreation Area.

The Laboulbeniales lack hyphae and, unique among related fungi, they exhibit determinate growth resulting in a well-defined thallus, which can be interpreted as a reduced hyphal system (Santamaria 1998, Tavares 1985). Thalli are often less than 300 µm in length, and consist of a receptacle with a perithecium or multiple perithecia and appendages bearing antheridia, which produce spermatia (some species in the genera *Aporomyces*, *Laboulbenia*, and *Rickia*, however, seem to lack antheridia [Rossi and Santamaria 2008, 2015; Thaxter 1926]). We currently do not know how these fungi reproduce. The thallus attaches to the host integument at the foot cell, and at least some species form haustoria—simple or branched rhizoidal structures that penetrate the host’s integument to provide additional stability and surface area for nutrient uptake (Benjamin 1971; Gäumann and Dodge 1928; Thaxter 1896, 1908). Presumably, haustoria penetrate the host’s integument to make contact with the cavity (haemocoel) and draw nutrient material from it. Although only observed in some genera, it is thought that all Laboulbeniales produce haustoria whether simple and minute or well developed (Benjamin 1971).

Laboulbeniales species can be monoecious (with both male and female sexual organs on the same thallus) or dioecious (with separate female and male thalli), and they rely on host contact to disperse their sticky, 2-celled ascospores (De Kesel 1993, 1995). Most Laboulbeniales exhibit a high degree of host specificity, with a host spectrum ranging from a single species to several related or congeneric species (De Kesel 1996, Majewski 1994, Scheloske 1969, Tavares 1985, Thaxter 1896). Phylogenetically unrelated insects living together in the same microhabitat (e.g., in ant nests or saltmarshes; Blum 1924, De Kesel and Haelewaters 2014) may become infected by the same species of Laboulbeniales. This finding implies that the parasite’s success is dependent on host characteristics as well as environmental features (De Kesel 1996).

Although some studies have suggested that insects infected with Laboulbeniales do not experience adverse effects on their fitness (Benjamin 1971, Scheloske 1969, Whisler 1968), negative effects on hosts have been reported (Báthori et al. 2015; Kamburov et al. 1967; Riddick 2006, 2010; Strandberg and Tucker 1974). Laboulbeniales can alter reproductive behaviors of infected hosts, such as oviposition patterns (Strandberg and Tucker 1974), and cause injury to the host’s appendages.
and abdominal tissues (Gemeno et al. 2004, Thaxter 1908). Some researchers have suggested that heavily infected hosts with high numbers of thalli on the head, eyes, antennae, mouthparts, legs, and/or elytra may no longer be able to detect food efficiently, mate, or behave as predators (Nalepa and Weir 2007, Scheloske 1969). Laboulbeniales have also been implicated in playing an influential role in the population dynamics of their hosts (Kamburov et al. 1967).

Laboulbeniales infect hosts selectively across the phylum Arthropoda, including representatives of the subphyla Cheliceriformes (order Acari; mites), Myriapoda (class Diplopoda; millipedes), and Hexapoda (class Insecta; true insects) (Haelewaters et al. 2012, Weir and Hammond 1997). The majority of known Laboulbeniales parasitize 9 orders among the Insecta: Blattodea (cockroaches and termites), Coleoptera (beetles), Dermaptera (earwigs), Diptera (true flies), Hemiptera (true bugs), Hymenoptera (bees, wasps, and ants), Mallophaga (bird lice), Orthoptera (crickets and allies), and Thysanoptera (thrips). Note that termites, previously ranked in the order Isoptera, were recently included in the order Blattodea, based on phylogenetic data (Beccaloni and Eggleton 2013). Although undeniably diverse, about 80% of the known fungal-parasite species are found on beetles (Weir and Hammond 1997). Parasite prevalence, often used as a measure for population dynamics of Laboulbeniales, is reportedly highest in species that overwinter as adults, have overlapping generations, form dense or continuous stable populations, and live in moist, damp habitats (De Kesel et al. 2011, Huldén 1983; recent examples in De Kesel 2011 and Wang et al., in press).

The family Coccinellidae includes species with aggregating behavior and overwintering patterns that are conducive to infection by Laboulbeniales (De Kesel 2011), and the family Staphylinidae includes the most commonly infected taxa in both the tropics and temperate regions (Weir and Hammond 1997).

**Methods**

The Harvard Museum of Comparative Zoology houses a collection of the Coccinellidae and Staphylinidae from the BHI that includes 1431 individuals representing more than 85 species. These specimens were collected for the ATBI from 13 islands using a variety of methods: litter sampling; pitfall, malaise, light, and bowl traps; and hand collecting (for details, see Rykken and Farrell 2013).

We examined pinned insects under dissecting microscopes at 10–50x magnification to record the diversity of the Laboulbeniales on their hosts. When we found infected insects, we removed individual fungal thalli from the host using Minuten Pins (BioQuip #1208SA, Rancho Dominguez, CA) inserted onto wooden rods. We removed thalli or groups of thalli at the foot and then mounted them in eosin dye in glycerol (1 g eosin powder per 2 ml 25% glycerol), PVA Mounting Medium (BioQuip #6371A), or Amann solution (Benjamin 1971). We placed a tiny amount of Hoyer’s medium on each slide with the tip of a Minuten pin, positioned thalli in the Hoyer’s, allowed the specimen to dry briefly, and placed a drop of the mounting medium on the cover slip, which was then dropped sideways onto the Hoyer’s medium. In this way, the briefly fixed, dried thalli remained in place.
when we added the cover slip, which was ringed with transparent or white nail varnish. We viewed mounted specimens at 400–1000x magnification for identification using numerous relevant systematic and taxonomic sources (listed in the Literature Cited). Slides are deposited at the Farlow Herbarium (FH; Harvard University, Cambridge, MA).

**Study site: Boston Harbor Islands**

BHI consists of 34 islands and peninsulas in waters near Boston, MA. The islands range in size from 3 to 150 ha and lie as far as 20 km from shore. The islands support a variety of habitats, including marine and estuarine intertidal wetlands and freshwater marshes (Tiner et al. 2003). The vegetation of the islands reflects succession after varying levels of human disturbance, and individual island floras are heavily influenced by usage history (Elliman 2005). Although fungi have not been specifically targeted in surveys of the islands, the invertebrates of the BHI are well documented (Rykken and Farrell 2013).

An ATBI, initiated by the National Park Service in 2005, sought to document all species inhabiting the islands, with particular attention to Arthropoda. The documented diversity of the BHI (Rykken and Farrell 2013) likely captures the true diversity of the islands. We expected lower diversity relative to the mainland, as is commonly found in island ecosystems (Gillespie and Roderick 2002) because of the islands’ geographic isolation, small size, and level of human disturbance (Davidson et al. 2011). Island areas and distances from the mainland are available at the National Park Service website (http://www.nps.gov/boha/).

**Results**

We screened 253 Coccinellidae, 27 (10.7%) of which were infected with Laboulbeniales, and 1178 Staphylinidae, 20 (1.7%) of which were infected. For an overview of the BHI sites with Laboulbeniales records, see Figure 1.

Screening records by host are included in the Supplementary Materials (see Supplemental File 1, available online at http://www.eaglehill.us/NENAonline/suppl-files/n22-3-N1361-Haelewaters-s1, and for BioOne subscribers, at http://dx.doi.org/10.1656/N1361.s1).

In the course of the study we also found new localities for *Hesperomyces virescens* Thaxt. (from Canada, Cuba, Guatemala, and Japan) and *Laboulbenia philonthi* Thaxt. (from Canada, Grenada, Panama, Trinidad, and Venezuela).

**Taxonomy**

*Clonophoromyces nipponicus* Terada & I.I. Tav. (Fig. 2), Transactions of the Mycological Society of Japan 34:357 (1993)

*Distribution and hosts.* Previously known only from Japan, where it was found on *Bryoporus gracilis* (Sharpe, 1888) (Staphylinidae, Tachyporinae) (Terada and Tavares 1993).

*New records from the BHI.* Suffolk County, THOMPSON ISLAND, 42°18′47″N, 71°0′33.6″W, 3–13 July 2007, J. Rykken, on *Bryoporus testaceus* LeConte,
1863, MCZ-ENT00601078, slides FH 00313408 (6 thalli from left metatarsus), FH 00313409 (2 thalli from sternites), FH 00313410 (2 thalli from tergites), FH 00313411 (1 thallus from left tarsus III), and FH 00313412 (4 thalli from tergites); same data, MCZ-ENT00601079, slide FH 00313511 (1 adult thallus from left elytron); Suffolk County, THOMPSON ISLAND, 42°18’47"N, 71°0’33.6"W, 6–13 June 2007, J. Rykken, on *Bryoporus testaceus*, MCZ-ENT00601092, slides FH 00313413 (3 thalli from tergites), FH 00313414 (1 thallus from mouthparts), and FH 00313413 (5 thalli from legs).

Figure 1. Overview of the Boston Harbor. Islands in grey and black are part of the Boston Harbor Islands National Recreation Area. Those in black have been reported in this paper: B = Bumpkin Island, C = Calf Island, G = Grape Island, GB = Great Brewster Island, L = Langlee Island, S = Snake Island, T = Thompson Island, and W = World’s End peninsula.
Remarks. Species of *Clonophoromyces* Thaxt. are rarely collected. To date, 2 species have been described, both of which are known from their type locality only (Terada and Tavares 1993, Thaxter 1931). The genus is characterized by cell II-carrying secondary axes, each of which produces a perithecium and accessory axes on the second cell. *Clonophoromyces nipponicus* differs from *C. grenadinus* Thaxt. in many characters (summarized in Terada and Tavares 1993), the most striking of which is that each secondary axis in *C. nipponicus* produces 2 accessory axes and a perithecium rather than the 1 accessory axis and a perithecium produced by *C. grenadinus*.

Figure 2. *Clonophoromyces nipponicus* from *Bryoporus testaceus* (Coleoptera, Staphylinidae). A: Adult thallus with broken primary axis, only basal part of cell III is left (slide FH 00313511). B: Young thallus with several developing secondary axes on cell II (slide FH 00313415). C: Young thallus with broken foot and intact primary axis (slide FH 00313409). Scale bar = 50 µm.
The material from the BHI is nearly identical to the original description of *C. nipponicus*, except for cell I. In North American thalli, cell I is widely suffused with dark brown color, whereas in the Japanese thalli, cell I is only colored strictly at the base (Terada and Tavares 1993). Color variation, however, is a seldom-used character for distinguishing species.

**Hesperomyces virescens** Thaxt., Proceedings of the American Academy of Arts and Sciences 25:264 (1891)
≡ *Stigmatomyces virescens* (Thaxt.) Thaxt., Proceedings of the American Academy of Arts and Sciences 29:106 (1894)
≡ *Hesperomyces hyperaspidis* Thaxt., Memoirs of the American Academy of Arts and Sciences 16:111 (1931)

*Distribution and hosts.* Described on *Chilocorus stigma* (Say, 1835) [as *C. bivulnerus*] (Coccinellidae, Coccinellinae) from California, US. This is a widespread species known from all continents except Antarctica, hosted by several genera of Coccinellidae (reviewed in Ceryngier et al. 2012, Santamaria et al. 1991).

**New records from the BHI.** Plymouth County, WORLD’S END PENINSULA, 42°15′51.9″N, 70°52′37.8″W, 16 August 2006, J. Rykken, on *Harmonia axyridis* (Pallas, 1773), MCZ-ENT00602410, slide FH 00313516 (8 adult thalli from left elytron); Plymouth County, LANGLEE ISLAND, 42°15′36.6″N, 70°53′13.3″W, 7–8 September 2005, J. Rykken, on *H. axyridis*, MCZ-ENT00602433, slide FH 00313517 (12 thalli from right elytron); Plymouth County, LANGLEE ISLAND, 42°15′37.2″N, 70°53′11.4″W, 21 September 2005, J. Rykken, on *H. axyridis*, MCZ-ENT00602435, slide FH 00313518 (11 thalli from right elytron); Suffolk County, CALF ISLAND, 42°20′28″N, 70°53′46″W, 19 June 2007, J. Rykken, on *H. axyridis*, MCZ-ENT00602436, slide FH 00313519 (6 thalli from right elytron); Plymouth County, LANGLEE ISLAND, 42°15′37.2″N, 70°53′11.4″W, 7–8 September 2005, J. Rykken, on *H. axyridis*, MCZ-ENT00602441, slide FH 00313520 (3 thalli from tip of right elytron); Plymouth County, BUMPKIN ISLAND, 42°16′51″N, 70°53′58″W, 10–19 July 2006, M. Wheat, on *H. axyridis*, MCZ-ENT00602442, slide FH 00313521 (9 adult thalli from right elytron); Suffolk County, THOMPSON ISLAND, 42°19′2″N, 71°0′31″W, 9 October 2006, B. Farrell & OEB10, on *H. axyridis*, MCZ-ENT00602484, slide FH 00313522 (6 adult thalli from tip of left elytron); same data, MCZ-ENT00602487, slide FH 00313523 (22 thalli from tip of left elytron); Plymouth County, WORLD’S END PENINSULA, 42°15′39.7″N, 70°52′14.5″W, 6–20 July 2006, J. Rykken, on *Psyllobora vigintimaculata* (Say, 1824), BHI-010084, slide FH 00313107 (2 thalli).

**Additional new records (non-BHI).** CANADA, Québec, Berthierville, 4 August 1917, G. Maheux, on *Hippodamia tredicimpunctata tibialis* (Say, 1824), D. Haelew. 526, in Collection d’insectes du Québec (CIQ), slide FH 00313524 (9 adult thalli from elytral tips); CUBA, Santiago de Cuba Province, Parque Nacional de Baconao, Gran Piedra, no date, no collector, on *Cycloneda sanguinea* (L., 1763), D. Haelew. 614, in American Museum of Natural History collection, slide FH 00313529 (3 thalli from left elytron); GUATEMALA, Huehuetenango Department, La Laguna, 15°45′48.1″N, 91°50′56.9″W, 9 May 2013, R.S. Zack, on *Cycloneda*

**sanguinea sanguinea** (L., 1763), D. Haelew. 167, in James Entomology Collection of Washington State University, slides FH 00313509 (9 thalli from left elytron) and FH 00313510 (17 thalli from left elytron); JAPAN, Hokkaido Prefecture, Hokuto, Murayama, 1918, no collector, on *Propylea* sp., D. Haelew. 555, in Collection d’insectes du Québec (CIQ), slides FH 00313527 (5 thalli from tip of right elytron), and FH 00313528 (7 adult thalli from tip of right elytron).

**Remarks.** On the Boston Harbor Islands, *H. virescens* has only been found on the native *Psyllobora vigintimaculata* and the invasive *Harmonia axyridis* (Multi-colored Asian Lady Beetle). Infection rates are 3.2% (*n* = 34) and 20.6% (*n* = 126), respectively.

Other lady beetle species at the BHI were found uninfected: *Anatis labiculata* (Say, 1824) (*n* = 1); *Brachiacantha ursina* (Fabricius, 1787) (*n* = 2); *Coccinella septempunctata* L., 1758 (*n* = 34); *Coleomegilla maculata* (DeGeer, 1775) (*n* = 10); *Cycloneda munda* (Say 1835) (*n* = 8); *Hippodamia convergens* Guérin-Menéville, 1842 (*n* = 2); *H. variegata* (Goeze, 1777) (*n* = 14); *Hyperaspis binotata* (Say, 1826) (*n* = 4); and *Propylea quatuordecimpunctata* (L., 1758) (*n* = 18). Of these, *C. septempunctata*, *C. munda*, and *H. convergens* have been previously observed with *H. virescens* in the US (Harwood et al. 2006, Thaxter 1931). The genera *Brachiacantha* Dejean, 1837; *Hyperaspis* Redtenbacher, 1844; and *Propylea* Mulsant, 1846 are known to host *H. virescens* (Ceryngier et al. 2012). The native *C. maculata* has been found with *Hesperomyces coleomegillae* W. Rossi & A. Weir and *Hesperomyces palustris* W. Rossi & A. Weir in Costa Rica, Cuba, Ecuador, and the US (Goldmann et al. 2013; D. Haelewaters, unpubl. data).

On 2 slides (FH 00313516 and FH 00313517) taken from *H. axyridis*, we found thalli with long median outgrowths of the perithecial tip together with thalli having short median outgrowths. This observation has also been made on *Adalia bipunctata* (L., 1758) from French Guyana (Bernardi et al. 2014) and *Cycloneda sanguinea sanguinea* (L., 1763) from Guatemala (this study).

**Ilyomyces cf. mairei F. Picard**, Bulletin Scientifique de la France et de la Belgique 50:446 (1917)

**Distribution and hosts.** Described from France on *Stenus elegans* Rosenhauer, 1856 (Staphylinidae, Steninae) and subsequently only reported on *Stenus aceris* Stephens, 1833 from France (Balazuc’s material, originally labelled as *I. lavagneii* (F. Picard) F. Picard; Santamaría 2003), on *Stenus* sp. from Spain (Santamaría, 1992), and on *Stenus clavicornis* (Scopoli, 1763) from the US (Haelewaters 2013).

**Record from the BHI.** Suffolk County, THOMPSON ISLAND, 42°18'59.6"N, 71°0'39.9"W, 3 July 2007, J. Rykken, on *Stenus clavicornis* (Scopoli, 1763), MCZ-ENT00601500, slides FH 00313250 (7 juvenile thalli from elytra) and FH 00313251 (1 juvenile thallus from elytra).

**Remarks.** The first and only record for North America, 8 thalli of *I. cf. mairei* were found on 1 specimen of *S. clavicornis* from Thompson Island (MCZ-ENT00601500), collected in 2007 (Haelewaters 2013). *Stenus* rove beetles are common in riparian habitats but only very rarely are reported with Laboulbeniales (Santamaría 2003). Prior studies reported the genus *Ilyomyces* on representatives
of the subfamily Steninae (Coleoptera, Staphylinidae) from France (Picard 1917, Santamaria 2003), Spain (Santamaria 1992, 2003), and Indonesia (Weir 1995), with *I. mairei* previously only known in Europe (Picard 1917; Santamaria 1992, 2003).

*Ilyomyces* cf. *mairei* in North America is reported on *Stenus clavicornis*, a congener of its European hosts, *S. aceris* and *S. elegans*. *Stenus clavicornis* is native to Europe and has been known in North America since 1968 (Majka and Klimaszewski 2008). *Ilyomyces* species have been reported on several *Stenus* species in Europe, and this incidence at the BHI likely represents either unintended co-colonization of North American habitats by the staphylinid host and its fungal parasite; host pursuit, in which host-range expansions are followed by the parasite from the same origin populations; or host-shift events, although this would imply that the parasite

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**Figure 3. Laboulbenia philonthi** from *Philonthus carbonarius* (Coleoptera, Staphylinidae). A: Adult thallus, showing scar of the trichogyne (arrowhead) (slide FH 00313515). B, C: Juvenile thalli (slide FH 00313515). Scale bar = 100 µm.
was already present in the US on another (ecologically) similar host, which as yet has not been proven (De Kesel and Haelewaters 2014, Haelewaters 2015, Nicholls et al. 2010, Roy et al. 2011).

**Laboulbenia philonthi** Thaxt. (Fig. 3), Proceedings of the American Academy of Arts and Sciences 28:174 (1893)

*Lectotype, designated here.* USA, Massachusetts, Belmont/Waltham, Waverley neighborhood, “April 23, 1813” [sic], no collector, on *Philonthus* sp., slide FH 00313494 (5 adult thalli). We recorded the collection date as written by Thaxter, but infer the collecting year to be 1893 based on surrounding materials.

**Distribution and hosts.** So far reported in Europe from: Austria, Czech Republic, France, Germany, Great Britain, Greece, Italy, Latvia, Lithuania, the Netherlands, Poland, and Spain; as well as from Korea, Turkey, Argentina, Guatemala, Mexico, and the US (Haelewaters et al. 2014, Majewski 2008, Santamaría et al. 1991). It parasitizes representatives of the genus *Philonthus* Stephens, 1829, and related genera (Staphylinidae, Staphylininae).

**New records from the BHI.** Suffolk County, CALF ISLAND, 42°20'25.5"N, 70°53'48.9"W, 28 August–4 September 2007, J. Rykken, on *Philonthus lomatius* Erichson, 1840, MCZ-ENT00601709, slide FH 00313392 (3 juvenile and 3 adult thalli from tergites); Norfolk County, GRAPE ISLAND, 42°16'7.4"N, 70°55'14.7"W, 2–10 July 2008, S.W. Cho, on *P. lomatius*, MCZ-ENT00601705, slides FH 00313512 (1 juvenile thallus from tergite) and FH 00313513 (4 thalli from tergites); Plymouth County, BUMPKIN ISLAND, 42°16'52.4"N, 70°54'8.1"W, 23–30 August 2006, M. Wheat, on *Philonthus carbonarius* (Gravenhorst, 1802), MCZ-ENT00601692, slide FH 00313520 (2 adult thalli from tergite); Suffolk County, GREAT BREWSTER ISLAND, 42°19'58.2"N, 70°53'48.3"W, 24 July–2 August 2006, R. Becker, on *P. carbonarius*, MCZ-ENT00601693, slide FH 00313515 (11 thalli from tergites).

**Additional new records (non-BHI).** CANADA, Québec, Granby, 4 July 1939, P.E. Mercier, on *Philonthus aurulentus* Horn, 1884, D. Haelew. 535, in Collection d’insectes du Québec (CIQ), slides FH 00313525 (1 juvenile and 4 adult thalli from tergites) and FH 00313526 (1 juvenile and 5 adult thalli from right metatibia); GRENADE, Saint Andrew Parish, Grand Etang, no date, no collector, on *Philonthus* sp., Thaxter 2923, slide FH 00313495 (1 juvenile and 10 adult thalli from “tip abdomen”); TRINIDAD, Port of Spain, no date, no collector, on *Philonthus* sp., Thaxter 2865, slide FH 00313496 (7 juvenile thalli); PANAMA, termite nest, no date, no collector, on *Philonthus* sp., Thaxter 3197, slide FH 00313497 (2 adult thalli); VENEZUELA, Los Chorros, no date, L.R. Reynolds (communicated by F.J. Psota to R. Thaxter), on *Philonthus* sp., Thaxter 3394, slide FH 00313498 (13 thalli); same data, on *Oligotergus fasciatus* (Nordmann, 1837) [as *Philonthus segmentarius*], Thaxter 3396, slide FH 00313499 (13 adult or submature thalli); same data, on *Oligotergus fasciatus* (Nordmann, 1837) [as *Philonthus anceps*], Thaxter 3395, slide FH 00313500 (27 thalli).

**Remarks.** Thaxter (1896, 1908) collected this species only on the American continent, with records from the New England region, California, and Florida in
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the US; and from Mexico and Guatemala. Hence, the suggestion was made that *L. philonthi* “appears to be strictly American” (Thaxter 1908), which is not the case as exemplified by the more recent numerous records across 3 climate zones in Europe: subboreal (De Kesel and Krastina 2006), temperate (Haelewaters et al. 2014, Majewski 2009, Santamaría et al. 1991), and mediterranean to subtropical (Santamaría et al. 1991). Between the time of Thaxter (1908) and this study, only 1 American collection has been made, in Ecuador (Proaño Castro and Rossi 2008).

Because Thaxter (1893, 1896, 1908) designated no type specimen, we decided to re-examine Thaxter’s slides of *L. philonthi*, which are deposited at FH. This led to our selection of a slide to serve as lectotype.

Study of Thaxter’s material also revealed new, unpublished records of this species for Grenada, Trinidad, Panama, and Venezuela, and extends the total thallus length recognized for this species. In his original description, Thaxter (1893) stated that the total length from foot to perithecial tip was 290–360 µm. Thalli from Poland and the Iberian Peninsula measure up to about 550 µm (Majewski 1994) and 558 µm (Santamaría 1998). The Venezuelan material, however, provided us with the longest thalli observed so far, up to 830 µm in length (slide FH 00313498). Thaxter (1908) mentioned that *L. philonthi* is the common species on *Philonthus* spp. in temperate South America, which may be supported by the current new records.

**Peyritschiella protea** Thaxt. (Fig. 4), Proceedings of the American Academy of Arts and Sciences 35:427 (1900)

*Lectotype, designated here.* GERMANY, Freistaat Thüringen, no date, no collector, on *Bledius tricornis* (Herbst, 1784), Thaxter 392, in British Museum collection, slide FH 00313501 (9 thalli from legs).

*Distribution and hosts.* Recorded on the following staphylinid genera: *Anotylus* Thomson, 1859; *Bledius* Samouelle, 1819; *Manda* Blackwelder, 1952; *Oxytelus* Gravenhorst, 1802; *Planeustomus* Jacquelin du Val, 1857; *Styloxys* Gozis, 1886 (subfamily Oxytelinae). The genus *Philonthus* Stephens, 1829 (subfamily Staphylininae) is mentioned as a host only twice, from Poland and Tibet (Lee 2006, Majewski 1994). *Peyritschiella protea* is widespread in Europe (Santamaría et al. 1991), but seldom reported from other continents: Asia (Tibet [Lee et al. 2006]), Africa (Algeria [Maire 1920]), and North America (Cambridge, MA, US [Thaxter 1908]).

*New records from the BHI.* Suffolk County, GREAT BREWSTER ISLAND, 42°20’1.7”N, 70°53’48.1”W, 14–21 June 2006, J. Rykken, on *Anotylus insecatus* (Gravenhorst, 1806), MCZ-ENT00601478, slides FH 00313424 (2 thalli from tergites) and FH 00313425 (4 thalli from last tergite); Suffolk County, GREAT BREWSTER ISLAND, 42°19’58.2”N, 70°53’48.3”W, 24 July–2 August 2006, R. Becker, on *A. insecatus*, MCZ-ENT00601477, slides FH 00313426 (10 thalli from tergites) and FH 00313427 (3 thalli from head); Suffolk County, GREAT BREWSTER ISLAND, 42°20’1.7”N, 70°53’48.1”W, 14–21 June 2006, J. Rykken, on *A. insecatus*, MCZ-ENT00601479, slides FH 00313428 (1 thallus from right elytron) and FH 00313429 (1 thallus from left elytron); Suffolk County, GREAT BREWSTER ISLAND, 42°20’1.7”N, 70°53’48.1”W, 21–28 June 2006, S. Madden,
on *A. insecatus*, MCZ-ENT00601472, slide FH 00313430 (1 thallus from pygidium).

**Remarks.** This is the second report of this species in the US. The first collection was made at the Fresh Pond (Cambridge, MA) and consists of a single (!) thallus (slide FH 00313502).

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Figure 4. *Peyritschiella protea* from *Anotylus insecatus* (Coleoptera, Staphylinidae). A: Adult thallus with 1 perithecium arising from the fourth layer of the receptacle (slide FH 00313424). B: Adult thallus with 1 perithecium produced at the third layer of the receptacle, another perithecium at the fourth layer (slide FH 00313424). Scale bar = 50 μm.
**Stichomyces conosomatis** Thaxt., Proceedings of the American Academy of Arts and Sciences 37:38 (1901)

*Lectotype, designated here.* USA, Massachusetts, Belmont/Waltham, Waverley neighborhood, October 1901, R. Thaxter, on *Sopedophilus littoreus* (L., 1758), slide FH 00313505 (11 thalli; 2 fully adult, others in different stages of development).

**Distribution and hosts.** On species of the genus *Sopedophilus* Gistel, 1856 (Staphylinidae, Tachyporinae) in various European countries: Belgium, Great Britain, the Netherlands, Poland, and Spain; as well as in North America (US), South America (Ecuador), Africa (Algeria), and Asia (Japan) (Bernardi et al. 2014, Haelewaters et al. 2012, Majewski 2008).

**New records from the BHI.** Suffolk County, GREAT BREWSTER ISLAND, 42º19'59.3"N, 70º53'46.1"W, 23–30 August 2006, J. Rykken, on *Sopedophilus testaceus* (Fabricius, 1792), MCZ-ENT00601267, slide FH 00313423 (3 thalli from left-hand side of pronotum); Norfolk County, GRAPE ISLAND, 42º16'15.3"N, 70º55'2.7"W, 2–10 July 2008, S.W. Cho, on *S. testaceus*, MCZ-ENT00601268, slide FH 00313432 (1 adult thallus from right elytron).

**Remarks.** Among Thaxter’s material, there is a series of 9 slides of *S. conosomatis*, labeled as having been collected in Belmont and Waverley, both in close proximity in eastern Massachusetts. Collections in this series were made in 1900 and 1901, and we assume that Thaxter had all 9 slides available when he made the description of *S. conosomatis* (Thaxter 1901). Because the slide that Thaxter indicated as holotype (FH 00313506, unpublished) is in unsatisfactory condition, we decided to designate a lectotype (= FH 00313505).

This species was commonly found on *Sopedophilus littoreus* (L., 1758) [as *Conosoma pubescens*; see Herman 2001 for the complex taxonomic history of this species] in the New England states (Thaxter 1901, 1931). This is the first published record from the US since Thaxter’s contributions.

Most thalli of *S. conosomatis* have only 1 perithecium and 0, 1, or 2 perithecial primordia. Both Tavares (1985) and Majewski (1994) described the development of secondary perithecia upon the cell above cell II. Since cell III never bears perithecia, this cell should be considered cell II'. Of the examined thalli on slide FH 00313423, 2 thalli have a perithecial primordium in addition to a normal (but broken) perithecium, both borne on cell II; in the other thallus, the third cell of the receptacle axis (cell II') gives rise to the single perithecium.

**Teratomyces actobii** Thaxt., Proceedings of the American Academy of Arts and Sciences 29:98 (1894)


= *Teratomyces brevicaulis* Thaxt., Proceedings of the American Academy of Arts and Sciences 29:99 (1894), syn. nov.

*Lectotype, designated here.* USA, Maine, Kittery Point, 18 August 1893, [R. Thaxter], on *Erichsonius nanus* (Horn, 1884) [as *Actobius*], slide FH 00313507 (1 thallus).
Distribution and hosts. Described from the US on *Erichsonius nanus* (Horn, 1884) [as *Actobius nanus*] (Staphylinidae, Staphylininae). Also reported from various European countries (Huggert and Eriksson 2010, Santamaria et al. 1991), Algeria (Maire 1920), and Sri Lanka (Thaxter 1931) on different species of *Erichsonius* Fauvel, 1874 and *Parerichsonius* Coiffait 1963.

**New records from the BHI.** Suffolk County, SNAKE ISLAND, 42°22′0.4″N, 70°58′44.3″W, 25 September 2007, J. Rykken, on *Erichsonius brachycephalus* Frank, 1975, MCZ-ENT00601563, slides FH 00313356 (3 juvenile thalli from tergite), FH 00313357 (1 juvenile thallus from last tergite), FH 00313358 (1 juvenile thallus from penultimate tergite).

Remarks. Of several available slides of collections from Kittery Point, ME, in agreement with the original description (Thaxter 1894), slide FH 00313507 is in good condition and contains a thallus with diagnostic characters. This specimen is designated above as the lectotype.

The genus *Teratomyces* Thaxt. consists of 10 species, 9 described by Thaxter (1893, 1894, 1896, 1900, 1901, 1931) and 1 by Rossi (2010). Additionally, 2 collections of *Teratomyces* have been made that are undescribed (New Zealand [Hughes et al. 2004], Bolivia [Weir and Rossi 2001]). Identification of species is difficult in this genus. Species delimitation is based on characters such as receptacle structure and color, length of cell VI, number of perithecia, and length and color of appendage; these characters can vary among thalli from a single host specimen (Hughes et al. 2004). The extent of morphological variability is often subject to debate in its use in defining species limits, especially when “morpho-species” occur on a single host specimen.

Although our material from the BHI is immature, the receptacle structure and especially the typical variability in blackening make us believe that it belongs to *T. actobii* (Thaxter 1896). Thaxter (1931) stated that *T. actobii* was a common species in the New England area, where he collected it several times in Maine (Kittery Point) and Massachusetts (Arlington). Other species of *Teratomyces* present in New England are *T. mirificus* on *Acylophorus pronus* Erichson, 1840; *T. quedianus* on *Hemiquedius ferox* (LeConte, 1878) [as *Quedius*]; and *T. brevicaulis* on *Erichsonius nanus* [as *Actobius*]. In *T. quedianus*, cell II is elongate, tapering downwards; its blackening typically extends to the basal part of cell III. These characteristics separate it clearly from *T. actobii*. The distinction between the latter species and *T. brevicaulis* is based on the size of cell VI (stalk cell of perithecium) and the larger clavate appendage cells in *T. brevicaulis*. Confusion arises because both species are sometimes found on a single host specimen (Thaxter 1896). In all of the Thaxter slides we examined, the distinction between *T. actobii* and *T. brevicaulis* as described by Thaxter is clear except for 1 record on *Erichsonius basalalis* Motschulsky, 1858 [as *Actobius*] from Sri Lanka (slide FH 00313508; see Thaxter 1931); these specimens combine features of *T. actobii* (the receptacle varies from hyaline to black) and of *T. brevicaulis* (having a short cell VI and slender perithecium of *T. brevicaulis*). For this reason, we consider *T. actobii* and *T. brevicaulis* to be synonyms thereby accepting considerable within-species morphological variability.
Both names were introduced in the same publication (Thaxter 1894). We choose to use the *T. actobii* for this species over *T. brevicaulis* because this name has been more widely used. We were unable to find any position-relatedness for either of the two forms; both were removed from legs and abdomen.

Generating DNA-sequence data will be the final key to unlock the phylogenetic relationship among morpho-species in *Teratomyces*.

**Discussion**

These observations are the first to document the Laboulbeniales of the Boston Harbor Islands, and the first reports of Laboulbeniales on Coccinellidae and Staphylinidae in the New England area since the time of Roland Thaxter (1858–1932).

The study of Laboulbeniales can bring new insight to the community dynamics of host–parasite interactions. Insects disperse with their parasites; thus, there is potential for infection of native hosts by infected introduced insect species. Although more research is needed, we hypothesize that host pursuit (Nicholls et al. 2010) followed by host shift is the mechanism at play in (the distribution of) *Ilyomyces cf. mairei* on the BHI.

Helmus et al. (2014, references therein) demonstrated that the acceleration of global shipping has increased establishment rates of exotic species, including beetles. The rapid and far-reaching transport of these beetles provides increased opportunities for dispersal together with their associated fungi. In addition, Humans often actively transport and spread insects used or studied for biological control. The study of obligate and non-motile ectoparasites of insects, such as the Laboulbeniales, provides an avenue to document effects of human-mediated host dispersal on parasite dynamics and distributions. However, further research is needed to fully characterize the diversity of the Laboulbeniales. Advances in molecular methods for the Laboulbeniales will no doubt accelerate determinations of parasite identities and host relationships.

In this study, we documented new records and revised species ranges from a relatively well-sampled region. With more thorough documentation of species diversity, future studies should characterize the mechanisms governing the distribution of these fascinating fungal parasites. With respect to Laboulbeniales research, the BHI collection at the Harvard Museum of Comparative Zoology is still largely unexplored. Of the Laboulbeniales discussed in the present paper, some species were previously known from the US (*H. virescens*) or specifically from New England (*L. philonthi, P. protea, S. conosomatis, T. actobii*). Others had not been reported from North America (*I. cf. mairei, C. nipponicus*) and are new records for the continent. We found the Laboulbeniales on the predicted host genera. *Bryoporus testaceus, Hippodamia tredicimpunctata tibialis*, and *Stenus clavicornis* are new host species for *C. nipponicus, H. virescens*, and *I. cf. mairei*, respectively.

Future work will include screening the Carabidae and other families in the BHI collection, as well as continued sampling on the Boston Harbor Islands.
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