CONTROLLING ZOONOTIC DISEASE VECTORS FROM INSECTS AND ARTHROPODS USING PRECONIDIAL MYCELIA AND EXTRACTS OF PRECONIDIAL MYCELIA FROM ENTOMOPATHOGENIC FUNGI

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ABSTRACT

The present invention utilizes extracts of the pre-sporulation (preconidial) mycelial stage of entomopathogenic fungi as insect and arthropod attractants and/or pathogens and can be employed to limit the zoonotic diseases they transmit. The fungus can be cultivated on grain, wood, agricultural wastes or other cellulosic material and extracts can be made thereof. More than one fungus and substrate can be used in combination with one or more antimicrobial, antiprotozoal, antiviral, and genetically modified agents that result in reduced spread of contagions and lessen the damage they inflict on animals, and plants.
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BACKGROUND OF THE INVENTION

1. Field of the Invention
2. Description of the Related Art
3. Diseases emanating from ecologically distressed and polluted environments increasingly threaten animals and plants. With deforestation, habitat destruction, decline in water quality, decreases in biodiversity, all of which are exacerbated by global climate change and human impacts, zoonotic diseases are increasingly a threat to healthy environments and their inhabitants, especially animal populations, including humans and their livestock. Many of these disease-causing organisms are carried by or bred within insects or other arthropods. Insects are any of the large class (Insecta) of small arthropod animals characterized, in the adult state, by division of the body into head, thorax, and abdomen, three pairs of legs on the thorax, and, usually, two pairs of membranous wings; arthropods are any of the largest phylum (Arthropoda) of invertebrate animals with jointed legs, a segmented body, and an exoskeleton, including herein insects, arachnids such as spiders, mites and ticks, and myriapods. Since many of these bite humans and livestock, as well as damage plants, they transmit a wide variety of diseases, many of which result in billions of dollars worth of damage to economies worldwide.

4. Insects are among the most diverse and numerous life forms on earth. While the majority of the one million named species of insects are considered beneficial, somewhere from 1% to 5% are considered to be pests. Some of these insect pests not only cause tremendous losses in terms of direct destruction of crops, livestock, and human dwellings, they are also vectors for pathogens including protozoa, round worms, bacteria, and viruses that cause devastating human health problems. As climates change, with an overall tendency to warming, tropical and subtropical diseases are spreading into temperate regions, once devoid of these threats. The negative physical, mental, economic, social, and ecological implications of disease carrying pest insects and arthropods are difficult to quantify since their effects are wide-ranging and multidimensional. As ecosystems in which humans dwell are harmed, water is polluted, sanitation huddles mount, toxins are accumulated, and food scarcity increases, animals (including humans) become much more susceptible to infection from pathogen-carrying insects and arthropods as their innate immune systems are weakened. Chemical pesticides, antibiotics, and vaccinations are notoriously ineffective against long-term exposure to populations of rapidly evolving organisms. Additionally, resistance to pesticides and antimicrobials can result in “super-bugs” which often develop in both insects and the microbes they transmit. As diseases ebb and flow, we need a more sophisticated way of out-smarting the vectors that carry them. If the vector can be stopped, the disease can be stopped. By using attractants from entomopathogenic fungi, this new approach allows the unusual flexibility of being able to switch or combine attractant extracts and mycelium sourced by tapping into the vast and continually evolving genome of naturally occurring wild or human-improved strains.

5. Many insects and arthropods are vectors for contagions. Some in particular are common carriers of pathogens and contagions. Many of these contagions are spread by simple contact, some are spread from bites or proboscis punctures, while others can be transmitted to animals when they consume these disease-laden insects.

6. Zoonotic disease is defined as any disease that is spread from animals to people. Any subsequent insect controlling technology can be enhanced since the insects and arthropods become concentrated as a result of the attractive properties of the preconidial mycelium or extract of selected entomopathogenic fungi. The further novelty of this invention is that it allows other technologies that limit disease to work more effectively by concentrating and localizing the disease-spreading organism to a more centralized locus, reducing expenses while enhancing efficacies. In essence, disease vectors by insects and arthropods can be better controlled.

7. Ants can carry diverse populations of pathogenic bacteria. For instance, Pharaoh ants (Monomorium pharaonis and related species) are known as vectors to more than dozen pathogenic bacteria, including Salmonella spp., Staphylococcus spp., and Streptococcus spp., and are especially dangerous to burn victims recovering in hospital environments. See Beaton S. H., “Pharaoh ants as pathogen vectors in hospit-
Although we have identified many diseases mosquitoes carry, we are unlikely to have identified them all. More mosquito-pathogen vectors are likely to be discovered as insects (and arthropods) evolve and species populations remodel. We know that mosquitoes can be the vector for viruses, using their proboscis as a form of a syringe capable for injecting many viruses, specifically West Nile virus, encephalitis viruses (Western equine encephalitis, St. Louis encephalitis, La Crosse encephalitis, Japanese encephalitis, Eastern equine encephalitis), Yellow Fever, and Dengue Fever. How many other viruses carried by mosquitoes, yet unknown or not yet evolved, will be discovered? Surely, there will be more.

Mosquitoes also inject protozoa into humans, including malaria (Plasmodium falciparum), which still results in millions of deaths per year worldwide. Control measures have included the use of chemical pesticides such as DDT™ and Deltamethrin™; however, their recurrent and prolonged use stimulates resistance. It seems Nature always finds a way around chemical “solutions.” To resolve complex problems in Nature, complex solutions are needed. This invention speaks directly to this issue.

Even the use of pesticide impregnated mosquito nets, which have been initially effective at reducing malaria infection, are not a long-term solution. Pandemonium in 2007 was initially found in Dakar, found that malaria infection rates in certain segments of the population rose to levels higher than before the introduction of bed nets. The researchers collected specimens of Anopheles gambiae, the mosquito species responsible for transmitting malaria to humans in Africa. Between 2007 and 2010 the proportion of the insects with a genetic resistance to one type of pesticide rose from 8% to 48%. By 2010, the proportion of mosquitoes resistant to Deltamethrin, the chemical recommended by the World Health Organization for bed nets, was 39%. In the last four months of the study, the researchers found that the incidence of malaria attacks returned to high levels. Among older children and adults the rate was even higher than before the introduction of the nets. The researchers argue that the initial effectiveness of the bed nets reduced the amount of immunity that people acquire through exposure to mosquito bites. Combined with resurgence in resistant insects, there was a rapid rebound in infection rates. The authors are worried that their study has implications beyond Senegal, writing “these findings are a great concern since they support the idea that insecticide resistance might not permit a substantial decrease in malaria morbidity in many parts of Africa.” Trape, J-F. et al., “Malaria morbidity and pyrethroid resistance after the introduction of insecticide-treated bednets and artemisinin-based combination therapies: a longitudinal study.” The Lancet Infectious Diseases, early online publication, doi: 10.1016/S1473-3099 (11)70194-3 (2011).

Below is a short summary of insects and arthropods with some of the zoonotic pathogens they transmit.

Insects and Arthropods Vectoring Zoonotic Pathogens
Ants: Bacteria (Salmonella spp., Staphylococcus spp., Streptococcus spp., etc.) Example: Fire ants spread several bacterial diseases in hospitals, including Staphylococcus, Salmonella and Clostridium.
Mosquitoes: Malaria protozoa (Plasmodium falciparum) carried by 30-40 species, including Anopheles gambiae. Viruses: West Nile (carried by more than 42 species), encephalitis, Yellow Fever and Dengue Fever (carried by several species of Aedes, including A. aegypti).
Flies: Bacteria, protozoa (ex. Tsetse fly carries the protozoan Trypanosoma causing often-fatal "sleeping sickness"). Flies also spread viruses, including influenza strains H5N2 & H5N1 (bird flu) and H11N1 (swine flu), which can also be carried by Blow Flies (Calliphoridae, Calliphora vicina and related species) and the common house fly (Musca domestica and related species). Houseflies can also transmit typhoid (Salmonella typhi) and dysentery (a disease complex caused by viruses, bacteria, protozoa and parasitic worms). White flies can transmit begomoviruses (family Geminiviridae), criniviruses, ipomoviruses, tomodviruses, and some carlaviruses.
Bed Bugs: MRSA (meticillin resistant Staphylococcus aureus bacteria) carried by Cimex species. Other bacteria can be transmitted by bed bugs.
Lice and ticks: Bacteria: Rickettsia spp. causing Rocky Mountain Spotted Fever; Bartonella vinsonii & B. henselae causing intramuscular infectious; and Borrelia burgdorferi causing Lyme disease.
Fleas: Bacteria, including Yersinia pestis causing bubonic plague.
Midges: Viruses (Blue tongue virus to cattle, epizootic hemorrhagic disease).
Leafhoppers: Tomato/Tobacco Mosaic viruses, wheat striate mosaic virus, maize fine streak virus, chickpea chlorotic dwarf virus, green petal virus, and others.
Virtually all biting insects and arthropods can result in bacterial or viral infections, either directly from a contagion reservoir within them or from wound exposure to the open environment. This is true with regard to both animal and plant diseases.
The present invention affords yet another new option for disease control: to attract but not necessarily kill mosquitoes, whilst reducing or eliminating their pathogen payloads. This option is important especially in areas where the insect populations are helpful in maintaining biological diversity of other animals that are dependent upon them for food. Removing all the insects from an ecosystem would likely result in unforeseen consequences, beyond that which is readily obvious. The food web is interconnected, and while most experts will agree that reducing disease vectors is prudent; destroying a native insect population is not.
Moreover, since Metarhizium species are natural parasites of mosquitoes, the natural genome of this and other entomopathogenic fungi offer sources of ever-evolving libraries of new strains, making resistance much more unlikely compared to chemical pesticides. An additional advantage of using preconidial entomopathogenic fungi such as Metarhizium anisopliae is that native strains of this fungus can be isolated wherever mosquitoes live, meaning that the constant co-evolution of this fungus to overcome resistance factors of the mosquitoes provides us with a unique partnership with nature to constantly adapt native, new strains of this
fusus for implementation in controlling mosquitoes. Moreover, if new strains of *Metarhizium anisopliae* are blended with any antimicrobial agent, the insects and the diseases they spread can be further controlled. Should the disease organism being carried by, for instance, a mosquito, develop resistance to an antimicrobial or antiviral drug, then a mixture of more than one drug or remedy can be employed to overcome resistance. Thus, this invention allows for a platform for continually out-smarting resistance by blending technologies and combining antimicrobials—out-racing the ability of insects and pathogens to adapt to either the entomopathogenic fungus or the antimicrobial method employed at the points of contact. Such synergism can have many derivative improvements and are expected by this inventor.

[0026] As an example, Artemesinif from *Artemisia* plants, has been found to be effective against malaria. Either pure or less expensive crude, extracts containing Artemesin can be blended with the preconidial extracts and/or mycelium of *Metarhizium anisopliae*. This combination would both attract mosquitoes and upon ingestion of the blended extract reduce the malarial loads they carry. Similarly, other combinations could include any or a plurality of antimalarial drugs or the crude precursors from which they are derived, including but not limited to: Quinine and related agents, Chloroquine, Amodiaquine, Pyrimethamine, Proguanil, Sulfonamides, Melloquine, Atovaquone, Primaquine, Halofantrine, Doxy-cycline, and Clindamycin. Moreover, the water/ethanol extracts of some polypore mushrooms, particularly *Polyporus umbellatus* has shown strong antimalarial activity, although the active ingredients have not yet been identified. Lovy, A., B. Knowles, R. Labbe & L. Nolan, “Activity of edible mushrooms against the growth of human T4 leukemia cancer cells, and *Plasmodium falciparum*,” *Journal of Herbs, Spices & Medicinal Plants* vol. 6(4): 49-57 (1999). Additionally, other polypore mushrooms, and Basidiomycetes, are likely to produce antimalarial compounds.


[0028] This same principle could also be used to enhance more traditional insect control devices. For example, blends of extracts and preconidial mycelium of entomopathogenic fungi can be used to enhance the performance of UV light based insect traps such as BASF’s “Vector™” or CO₂ emitting suction traps. In essence, any current or future method might well result in greater performance for controlling insects, whether these be mosquitoes, flies or others, by employing extracts and mycelium of preconidial entomopathogenic fungi.

[0029] Using preconidial entomopathogenic fungi to develop new or enhance existing insect control measures may also be used to help mitigate diseases spread by flies. Flies such as the blood sucking Tsetse fly carry the protozoan *Trypanosoma* that causes an often fatal “sleeping sickness” in Africa. Blow flies, aka ‘blue bottle flies’ (*Calliphora nigraria* and *Aldrichina grahami*) and house flies (*Musca domestica*) have both been found by multiple researchers to harbor and carry bird flu viruses, meaning that poultry farms and slaughter houses represent nexus distribution points for this contagion. See http://www.flutrackers.com/forum/showthread.php?t=29335. According to the researchers, “more than one-third of the adult *Musca domestica* sampled contained *AI* [avian influenza] virus particles.” Blow flies swarm and breed upon carcasses, including birds, as well as broken eggs and bird feaces, and can acquire bird flu viruses. The ever-so-common housefly can carry bird flu viruses, and potentially re-infect chickens and other poultry that eat flies regularly. What has not been reported yet is whether or viruses such as bird flu can be transmitted to humans from infected flies. Given the huge swarms of flies that congregate around dead and diseased animals, this vector seems likely. According to the researchers, “more than one-third of the adult *Musca domestica* sampled contained *AI* virus particles” (http://www.flutrackers.com/forum/showthread.php?t=29640).

[0030] As symptoms of bird flu infection may not be evident for a few days, and yet the animals can be infectious, factory farms, and in particular slaughter houses (where blow flies feed on cadavers and also make contact with living animals) can be a serious, although largely unpublicized threat to public health. Flies infected from contacting poultry infected from bird flu, for example, can be eaten by non-infected birds, thus increasing the probably of disease transmission. Thus the need to attract virus-vectoring flies, and to reduce their pathogen payload is dually important. Note that even if the flies are not caught, but seek out, make contact with, and/or ingest the sweet extracts having antiviral or antimicrobial properties, the benefits incurred are that these insects are then less infectious due to reduced levels of contagions.

[0031] Because the purification of antimicrobial and antiviral drugs is typically more expensive than their crude, or semi-pure precursors, this invention anticipates that less-than-pharmaceutical grade antiviral, antimicrobial, and antiprotozoa medicines can be employed in combination with extracts and the mycelium of preconidial entomopathogenic fungi to create a successful treatment in the prevention, mitigation, or curing of contagions transmitted by insects and arthropods. Moreover, the inventor’s prior research on the use of polypore mushroom derivatives to combat viruses, which employ a similar method of extraction to the methods described herein for the creation of attractant preconidial entomopathogenic extracts, is yet another application of this novel way of limiting zoonotic contagions.

[0032] Other insect arthropods such as lice and ticks can carry *Rickettsia* bacteria causing Rocky Mountain Spotted fever. Flies can transmit bubonic plague (*Yersinia pestis* bacteria) and lice can carry *lyme disease* (*Borrelia bacteria*) to humans, deer, and other animals. ‘Bed bugs’ (Cimex species from the Cimicidae) have also recently been found to carry...
drug-resistant staph bacteria (MRSA—methylis resistance *Staphylococcus aureus*), compound the challenge faced by hospitals, hotels, dormitories, army barracks, prisons, and other densely populated areas. Denser populations of humans and animals—especially denser populations of immunocompromised humans and animals—increase the probably of infection and re-transmission. Whether the initial infection being transmitted from a biting insect or arthropod is from a bacterium or a virus, co-occurrence of non-insect borne diseases may more readily ensue. The now-lowered immunity of the infected animal population at large may, for instance, make the spread of Ebola, Hanta, bird flu viruses, diphtheria, dysentery, and any contagion more readily spreadable. The resultant consequences of a population’s lowered immunity can also degrade the overall population’s immunological defenses against cancers. Conversely, those already suffering from cancer, or have compromised immune systems due to other diseases, are more susceptible to infection.

Moreover, insects spread viruses into plants. For instance, caterpillars and grasshoppers spread the Tomato-Tobacco Mosaic Virus. For farmers, there are dual advantages for controlling plant eating insects and the crop destroying diseases they spread. By combining extracts from the polypore mushroom, *Fomes fomentarius*, a source of antiviral agents active against the Tobacco Mosaic Virus with extracts of preconidial mycelium of *Cordyceps* species (well known for infecting caterpillars and grasshoppers), farmers could benefit by both limiting these crop damaging insects and lessening the threat of viruses they spread. This is but one of many examples that will become obvious and are expected manifestations of this over-arching invention.

Hence this inventor sees a two-fold need: to control movement of insects, and to control the population bio-burden of insects and arthropods that transmit diseases to people, animals, and plants. Combining methods and compositions discussed herein to create discrete ways to attract disease-carrying insects and subsequently killing them and/or reducing their pathogenic payloads will be important for protecting environmental health. In the age of technologies creating genetically modified organisms, potentiating pathogen carrying insects as biological weapons is possible and protection from such threats is sorely needed. Hence, this invention could be important for defense against bioterrorism in its many elaborations.

**SUMMARY OF THE INVENTION**

In view of the absence of using the preconidial mycelium of entomopathogenic fungal mycelium to attract insects and arthropods that carry contagions and disease, the present invention provides improved insect bio control agents, and methods and compositions of using such agents.

The present invention offers a unique approach to zoonotic disease control by attracting insects or arthropods that contact or ingest “preconidial” mycelium of entomopathogenic fungi (that is, mycelium in a developmental state prior to conidia or spore formation) which is also combined with any pest or disease controlling mechanism, another drug, plant derived medicine, pharmaceutical, hormone disrupter, attenuation gene, bacteriophage, or fungus or fungi possessing antimicrobial or anti-viral properties that result in arresting movements by such insects or arthropod while limiting the populations and pathogenicity of their carrier diseases.

**0037** Preconidial mycelium is defined as mycelium lacking spores but existing in a state prior to or without spore formation. The preconidial state and preconidial mycelium may include sclerotia or microsclerotia, compact masses of hyphae that are formed by certain fungi and give rise to new fungal growth or spore-producing structures. Commercial conidial formations of *Metarhizium anisopliae* strive to achieve at least 1,000,000 conidia per gram, and optimally 10,000,000,000 conidia per gram. Preconidial mycelium is defined in ranges as preferably having less than 10,000 conidia per gram of myceliated substrate, more preferably less than 1,000, and most preferably less than 100 conidia per gram. The preconidial mycelium is optimally without spores. Preconidial mycelium can be created by selectively cultivating non-sporeulating sectors from entomopathogenic fungi or by chemical agents that temporarily suppress conidia (spore) formation. See U.S. Pat. No. 7,951,389 and other patents by the present inventor. Either way, conidia formation can be re-activated, either naturally or artificially. Using preconidial preparations, mycelium and extracts in a variety of forms—living, frozen, dried, freeze dried, extracted—offers advantages by attracting insects or other arthropods and concentrating them in a more centralized location. Once concentrated, a variety of technologies can be deployed to trap or kill the insects and other arthropods, and reduce the pathogen payload they harbor.

**0038** Such preconidial mycelium of entomopathogenic fungi may be used solely as an attractant (either as an attractant for pest insects or as an attractant for beneficial insects) or as an attractant and pathogen where the preconidial mycelium is both the attractant and the pathogenic agent. Additionally, whence the insects or arthropods make contact with the preconidial entomopathogenic mycelium there is the added advantage of improving the restricting of disease transmission by having another control technology in the same locale.

Where attractant mycopesticidal strains are utilized with insects, the infected insects carrying the fungal hyphae become a vector back into population, further dispersing the antimicrobial mycelium. The preconidial mycopesticidal mycelium can grow within or upon an insect, can be carried to another insect when they touch, or can grow upon organic debris allowing subsequent insect infestation from simple contact. Moreover, some insects will become immunocompromised from contact with *Metarhizium* based products, and the resultant lowered immunity allows for other pathogenic fungi to infect the now weakened insect. This secondary infectious suite of organisms can be more virulent than the *Metarhizium* itself. All these modes of action result in lowering the bio-burden and the pathogenic payloads that these zoonotic disease-bearing insects harbor. Multiple avenues of growth and infection are provided and could be further enhanced if the addition of conidia from entomopathogenic fungi were deployed, as part of the composition of insect control.

**0040** The preconidial mycelium of mycopesticial fungi is grown in pure culture using standard techniques for in vitro propagation. Once inoculated onto a substrate such as grain or wood, the mycelia matures to a state prior to conidia formation. The window of utility extends from post-spore germination through all stages of mycelial growth prior to sporulation. The preconidial mycelium may be utilized as is or may be arrested in its development through means such as flash chilling, freeze-drying, air-drying, extract in acetone, water dehydrtation, cryogenics, refrigeration, gaseous cooling, gas affix-
ation (nitrogen, carbon dioxide, ethylene) and packaged in spoilage-proof or sealed packages. Even with post-conidial cultures of entomopathogenic fungi, methods can be employed which will ‘turn off’ conidial formation and ‘turn on’ non-conidial mycelial growth, resulting in attractancy, phagostimulation, and in some cases trailing following or swarming behavior.

[0041] The end-user facilitates opening the package and placing the exposed mycelia contents in the vicinity of recent pest activity. For use as an attractant, extracts of the preconidial mycelium may also be utilized. It is envisioned that the fungal attractants and/or pesticides may be used in conjunction with any type of appropriate trap or attractant disseminator or delivery system as is known to the art.

[0042] By combining an extract of mycelium from a fungus having antimicrobial and/or antiviral properties with an extract from the preconidial mycelium of an entomopathogenic fungus, the unique mixture can serve as a unique combination for mitigating disease transmission. A novel agent or treatment that kills the contagion but also severely harms the human host, for instance, is neither medically practicable nor commercially attractive. However, a novel agent that neutralizes the bacterium, protozoa or virus being carried by an insect is both medically and commercially significant. Moreover, if the preconidial entomopathogenic fungi attracts and simultaneously carries an infectious agent that controls the insect while also reducing internal pathogens harmful to animals and crops, disease transmission vectors can be limited, arrested, or re-directed using these unique combinations.

[0043] The present invention thus provides improved products and methods wherein the fungal mycelium acts as food and attractant and/or as an ingested or contact insecticide, palatable enough that insects will readily consume it even in the presence of competing food sources, or otherwise repellent materials, with high recruitment of other insects among insects that exhibit such behavior. This results in multiple visits to a highly attractive (and potentially virulent) food, thereby providing numerous individual insect and/or colony vectors of inoculation.

[0044] The present invention further provides these and other advantages with improved control of insect pests using fungal compositions (mycopesticides and mycocontractants) having strong attractant properties and placing these attractant preconidial fungi in or around an object or area to be protected. The present invention also provides insecticidal foods and baits that utilize, as a toxicant, relatively innocuous and naturally occurring materials as the active agent, so as to control insects carrying zoonotic diseases without undue effect on the ecology. Alternatively, the present invention provides attractants that can be utilized with bio-control agents, environmentally benign biopesticides, chemical control agents including insect toxicants and pesticides, human modified organisms, viruses and bacteriophages, physical control agents such as mechanical and electrical devices and combinations thereof. It is to be expected that the number of sub-inventions and applications obvious to those skilled in the relevant arts is limited only by imagination and time, and any such derivative inventions and applications should be considered to be part of the invention disclosed herein. New zoonotic diseases and new disease controlling technologies will emerge and the inventions described herein are likely to enhance many future technologies.

[0045] Still further objects and advantages of the present invention will become more apparent from the following detailed description and appended claims.

[0046] Before explaining the disclosed embodiments of the present invention in detail, it is to be understood that the invention is not limited in its application to the details of the particular products and methods illustrated, since the invention is capable of other embodiments, including those embodiments that have not yet been reduced to practice and tested. In addition, the terminology used herein is for the purpose of description and not of limitation.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

[0047] The concepts of “pathogens” and “pathogenic” (and the related “entomopathogens” and “entomopathogenic”) have implications that extend well beyond the standard dictionary definition of “capable of causing disease or mortality.” Some entomopathogenic fungi are widespread and cause no known affects whatsoever in their insect hosts; Myrmecinaeospomum durum is illustrative of entomopathogenic fungi that cause few symptoms and are consequently hard to detect in the first place. Schmid-Hempel, P., Parasites in Social Insects, Princeton University Press, p. 83 (1998). Entomopathogenic fungi as used herein are those capable of infecting and parasitizing insects, regardless of their actual effect on the host. “Virulence” and “virulent strains” similarly have meanings extending beyond the dictionary definition of extremely infectious, malignant or poisonous. Parasite virulence and host resistance determine how host and parasite interact in ecological time and how they co-evolve. Virulence is often defined as an increase in the host mortality rate as a result of the parasite’s presence. But reduced host fecundity, parasite replication rate within the host, and several other measures have also been used. Virulence should in principle also include instances where the behavior of the host is manipulated by the parasite to increase the probability of its successful transmission and where it places the individual host at greater risk. See Schmid-Hempel, supra, pp. 237-238. Here the terms virulent and virulence are used in a broad sense that encompasses all of these meanings. It will refer to processes which are caused by entomopathogenic fungi and which lead to a reduction in some component of the host’s fitness or an increase in mortality. Virulence and resistance are therefore properties that emerge as a result of host-parasite interaction in a given environment. Expression of virulence is as diverse as the lifestyles and characteristics of the insect hosts and the entomopathogenic fungi themselves.

[0048] The present invention provides improved mycocontractants and mycopesticides (fungal mycelia utilized as insect attractants or baits and/or insect biopesticides, after mycology, the study of fungi) to control zoonotic diseases harbored by and vectored by insects and non-insect arthropods.

[0049] Laboratory procedures for testing entomopathogenic fungi often involve procedures inapplicable in the field, such as “dusting” of many or all of the insects with spores or forced contact with conidia in petri dishes (itself a form a stress). Insects infected with mycopathogenic spores are often rejected or isolated from the general population, thus limiting the further spread of the fungal disease. Wilson, E. O., The Insect Societies, The Belknap Press of Harvard University Press, pp. 103-119 (1974). For these and other reasons,
conidia of entomopathogenic fungi have often been much more effective under laboratory conditions than in the field.

[0050] It was found that the “fragrance signature” of the mycoperistomial mycelium is a strong attractant to insects prior to conidia formation. The genesis for these findings was the initial observation that the odor of the cultured mycelium was similarly pleasing to humans when preconidial and repellent after conidia formation; smell and the fragrance signatures of mycelium are utilized by the present inventor as indicators of the health of the mycelium in large scale production of gourmet and medicinal mushrooms, whereas “petri dish mycologists” and entomologists studying pathogenic fungi are typically trained not to sniff or inhale from the cultures. In fact, most mycologists are trained not to do so, as standard laboratory protocol, because such actions could be a threat to their health. It was noted such fragrance signatures are lost when mycelium is grown via liquid fermentation—this may be due to such fragrance signatures being “washed away” or due to the greatly reduced nutritional base available to the mycelium in liquid fermentation as compared to solid substrates such as grain or wood. “Outgassing” of CO₂ and attractant molecules by the mycelium is believed by the present inventor to be responsible for at least some portion of the attractant value. It was also noted that liquid fermentation utilizing a typical fermenter with bubbled air mixing will promote conidia formation, with such conidia production being even further promoted by the common commercial practice of utilizing bubbled or chemically generated oxygen.

[0051] In addition to the attractant properties and phago-stimulatory (feeding stimulating) properties of preconidial mycoperistostesists, it was further found that pathogenic fungal control agents are much more effective when preconidial (pre-sporulation) mycoperistomial mycelium is ingested and/or contacted by the targeted insect as compared to conidia or post-sporulation mycelium/conidia offered to targeted insects for the purpose of infection by contact. The preconidial mycoperistomial mycelium is thought to be an effective attractant and/or pathogen, at least in part, because it is a preferred food, particularly for social insects and other fungi-feeding insects.

[0052] The preconidial mycelium has been observed to be a preferred food source that stimulates “grazing” of the fungi on wood and/or grain, scattering of the fungus, and caking of the fungus by social insects including termites, carpenter ants, and fire ants. Novel behaviors observed in the social insects include that of formosan termites (Coptotermes formosanus) ignoring available wood while preferring to set up “housekeeping” in the mycelium, and fire ants and carpenter ants moving the preconidial fungi around the feeding arena and/or into nest chambers. Social insect colonies have been described as “factory forresses.” See Wilson, supra, (1974); Oster, G. F. and E. O. Wilson, Caste and Ecology in the Social Insects, Princeton University Press (1978); Schmid-Hempel, supra, (1998). While it may be difficult for a parasite to “break into the fortress” and gain access to a colony, once inside, the opportunities abound (Schmid-Hempel, supra, p. 77 (1998)). Similarly, once the social insect defenses have been penetrated via the attractiveness of preconidial mycoperistomial mycelium, the opportunities abound for further inoculation and spread of the preconidial mycelium both orally and dermally, as well as optional introduction of other bio-control agents. Chemical toxicants. Novel and unique features of the invention include the use of mycoperistomial mycelium or extract as an attractant, the use of a mycoperistomial vector of parasitization that relies directly on hyphal fragments to infect both insects and/or social insect housing structures, the use of high levels of carbon dioxide to grow and maintain preconidial mycelium, the use of late sporulating strains to prolong the attractive preconidial state, the use of various methods to arrest development at the preconidial stage and/or to facilitate growth, packaging, shipping, and convenient application by an end user, and various improvements in methods of attracting, controlling, preventing, eradicating, and limiting the spread of disease vectoring insects and arthropods.

[0053] Preconidial mycelium has proven to be highly effective by ingestion or contact, with the exudate-excreting mycelial hyphae already being in a state of active growth when ingested or contacted. The preconidial mycelium is thought by the present inventor to function both as a “fungal food of infection” and as a contact insecticide. Efficacy as a contact insecticide is believed to be aided by the somewhat “sticky” nature of mycelium. While not wishing to be bound by any theories or hypotheses, the present inventor believes various possible vectors for further spread and growth of the preconidial mycelium include: incidental contact and adhesion; feeding and “sloppy eating” which may spread hyphae to insect cuticles, food caching, individual and social grooming; aerial transmission of hyphal fragments (as dry hyphal fragments are much less dense than spores, they easily become airborne and spread); inhalation; incidental contact; trophalaxis (exchange of liquid food); protodea thorahalaxis (exchange of anal excrement by termites and others); cannibalism; mating; contact with cadaver; inoculation of housing structures; etc. Mycoperistomial species are thought by the present inventor to employ various pathogenic modes when transmitted via ingestion or contact with mycelial hyphae, including: infection via the cuticle, the tracheal openings, the alimentary canal, or wounds with resultant growth upon the insect and resultant depletion of host resources and/or damage or destruction of host tissue; production of antibiotics, antibacterial, and antiprotistoxins with the resultant death of microflora within the gut; production of ant-fungal compounds affecting symbiotic and associated fungi; production of toxic substances by the entomopathogens; suppression or disruption of the immune system response, etc.

[0054] Since mites are non-insect arthropods and mites have long been observed as a pest to mushroom crops, both at the mycelial stage and when mushrooms subsequently form (Stamets, P. and Chilton, J., The Mushroom Cultivator, Agarikon Press, 1983), and since mites can be parasitized by entomopathogenic fungi, the use of preconidial mycelium of entomopathogenic fungi to attract and control mites, and the bacterial “biotext” they inflict to mushroom crops is an important new strategy for limiting losses in mushroom farms, or wherever mites inflict damage and cause bacterial diseases. The same methods described herein can be readily adapted for limiting mites and the diseases they spread to plants, thus protecting crops.

[0055] In utilizing wood and other cellulose containing materials, one preferred method is to grow the pre-sporulation mycoperistomial mycelium on wooden or other cellulose materials “bait blocks” or “bait traps.” Bait chips, blocks, or traps (or optionally other forms such as pellets, extruded pellets, mats, fabrics, ropes, etc.), optionally soaked with a bait solution, hormone, or other sugar and/or nutrient solution, are infused and/or inoculated with preconidial mycoperistomial mycelia which then spread the infection to the targeted
insect pests via any of the mycelium vectors described herein. Biodegradable bait traps may be made of, or have components made of various cellulose, lignin, cellulolignin, carbohydrate, and fiber materials including but not limited to: paper products and cardboard; wood and sawdust; corn cobs and cornstalks; chip board; fibers such as jute, flax, sisal, reeds, grasses, bamboo, paper, and coconut fibers; nut casings such as peanuts, almonds, walnuts, sunflower, pecans, etc.; seed hulls such as cottonseed hulls; agricultural products and byproducts such as hemp, cereal straws, sugar cane bagasse, soybean roughage, coffee wastes, tea wastes, cactus wastes, banana fronds, and palm leaves; industrial byproducts such as fiberized rag stock; combinations thereof, and numerous other forest agricultural, and industrial products and byproducts which will host mycelium and are degradable by mycopesistical fungi. Where rapid biodegradability of the traps is desired, materials such as cardboard or paper may be utilized. For insects including carpenter ants or termites, cockroaches, etc., the bait blocks preferably contain channels, tunnels, grooves, ridges, holes, or perforations specifically sized to allow entry by the targeted species and or its brood, pupae and/or larvae. Inoculation may, for example, be accomplished via grain in the channels and the blocks may optionally be layered or “wafered” together. A composite, layered or intertwined matrix of materials may be utilized, with one set of materials infused with the attractant extract of an entomopathogenic species and the other containing active or metabolically arrested preconidial mycelium. A multiplicity of such bait blocks or traps or barriers may be utilized to protect structures, agricultural locations, hospitals, dormitories, etc. A fungal matrix with a plurality of pre-sporeulating mycopesistical fungal species and/or extracts that are highly attractive to the targeted pest insect, combined with antimicrobial, antiprotease, and anti-viral ingredients, may be created so that the targeted pest is drawn close to a locus where the insect pest becomes infected and is harmed or killed by the selected fungi or via other means.

[0056] The wooden, cardboard, or lignin-cellulose baits and bait traps may optionally be frozen, dried or freeze-dried, or gaseously treated to arrest growth until activated by moisture and air exposure. Either the myceliated bait may be presented to the insect, with rehydration and recovery taking place, for example, within the central nests of social insects, or placed in the migration corridors of traveling insects. The bait block may be rehydrated prior to or during use or presented fresh.

[0057] The highly active nature of preconidial mycopesistidal mycelium indicates that essences extracted from preconidial mycelium of mycopesistical fungi can be expected to be highly attractive in and of themselves, and in conjunction, associated compounds may possess innate antimicrobial or antiviral properties, and thereby similarly useful alone or in conjunction with biological, chemical, mechanical and/or electronic insect control agents, useful as masking agents for otherwise repellent toxicants for insect pests, and useful as “distractors” in diverting insects away from sites that need protection. Such essences include extracts, concentrates, fragrances, derivatives, active constituents, etc. and may be prepared by methods known to the art including extraction with water, alcohols, organic solvents and supercritical fluids such as CO2, etc. Extracts may also be prepared via steam distillation of volatile components, similar to the preparation of “essential oils” from flowers and herbs. Suitable alcohols include those containing from 1 to 10 carbon atoms, such as, for example, methanol, ethanol, isopropanol, n-propanol, n-butanol, 2-butanol, 2-methyl-1-propanol (t-butanol), ethylene glycol, glycerol, etc. Suitable organic solvents include: unsubstituted organic solvents containing from 1 to 16 carbon atoms such as alkanes containing from 1 to 16 carbon atoms; alkenes containing from 2 to 16 carbon atoms; alkynes containing from 2 to 16 carbon atoms; and aromatic compounds containing from 5 to 14 carbon atoms, for example, benzene, cyclohexane, cyclopentane, methylcyclohexane, pentanes, hexanes, heptanes, 2,2,4-trimethylpentane, toluene, xylene, etc.; ketones containing from 3 to 13 carbon atoms such as, for example, acetone, 2-butanone, 3-pentanone, 4-methyl-2-pentanone, etc.; ethers containing from 2 to 15 carbon atoms such as such as 2-butyl methyl ether, 1,4-dioxane, diethyl ether, tetrahydrofuran, etc.; esters containing from 2 to 18 carbon atoms such as, for example, methyl formate, ethyl acetate and butyl acetate; nitriles containing from 2 to 12 carbon atoms such as, for example acetonitrile, propionitrile, benzonitrile, etc.; amides containing from 1 to 15 carbon atoms such as, for example, formamide, N,N-dimethylformamide, N,N-dimethylacetamide; amides and nitrogen-containing heterocycles containing from 1 to 10 carbon atoms such as pyrrolidine, 1-methyl-2-pyrrolidinone, pyridine, etc.; halogen substituted organic solvents containing from 1 to 14 carbon atoms such as, for example, bromotrichloromethane, carbon tetrachloride, chlorobenzene, chloroform, 1,2-dichloroethane, dichloromethane, 1-chlorobutane, trichloroethylene, 1,2-dichloroethane, 1,2,4-trichlorobenzene, 1,1,2-trichloroethane, etc.; aldehydes, aryls, arylalkyl, aryl, alkaryl and aralkyl substituted organic solvents containing from 3 to 13 carbon atoms such as, for example, 2-butoxy ethanol, 2-ethoxy ethanol, ethylene glycol dimethyl ether, 2-methoxy ethanol, 2-methoxethyl ether, 2-ethoxyl ether, etc.; acids containing from 3 to 10 carbon atoms such as acetic acid, trifluoroacetic acid, etc.; carbon disulfide, methyl sulfide, nitromethane and combinations thereof. Extracts may also be prepared via sequential extraction with any combination of the above solvents. The extracts may optionally be combined with fixatives, enhancing agents, oils, alcohols, solvents, glycerin, water and other substances that aid in distributing the attractant and/or enhancing its fragrance value. Essences extracted from preconidial mycelium of mycopesistical fungi can be used as a protectant or distractors, luring insects away from a locus and preventing insect damage to a locus, habitat, structure, crop, animal, human, etc. Such attractant essences and extracts may be utilized with wicking agents, sprayers, etc. to enhance their effectiveness. Preliminary indications are that such attractant molecules are polar and thus best extracted with polar and/or hydrophilic solvents. The present invention in conjunction with the principles of chemical ecology and evolutionary biology raise the possibility that the entomopathogenic fungal species produce attractant molecules (or more likely, groups of attractant molecules) that have co-evolved over evolutionary time with species of insects or groups of insects. Such attractant molecules, optimized for one species of insect, may well show attractant properties to larger groups of insects. Since all these fungi produce fatty acids, particularly linoleic acids, these and other sterols, all have within them the preparation of these attractant molecules. It will be apparent to these skilled in the art that numerous such molecules or groups of attractant molecules may be isolated.
and/or characterized from the preconidial fungi of the present invention and as such should be considered part of the present invention.

[0058] The preconidial mycelium or extracts thereof may be utilized solely as an attractant for various purposes. For example, preconidial mycelium may be utilized to affect insect choice of geographical location, destructive and zoonotic disease bearing pests being attracted and distracted away from structures, agricultural plots, hospitals, army barracks, theaters, convention centers, schools, etc. Fungal species and strains particularly attractive to beneficial insects may be utilized to attract desired insect species, the fungi acting as a biological catalyst to steer the course of the insect community evolution. Alternatively, varying insects may simply be attracted to occupy the environment and thus forestall pest invasions. It is known that virulence of entomopathogenic strains varies widely in the laboratory when tested via typical conidial based assays, with mortalities from 0% to 100% being recorded dependent upon such factors as number of conidia applied per insect and the insect species and the entomopathogenic species and strain being tested. Similar results may be expected for preconidial formulations, although a greater effectiveness in general may be expected since lack of virulence in the typical bioassay is often related to a failure of conidia to adhere to the insect and/or failure of the conidia to germinate as discussed above. Thus strains of “pathogenic” or “entomopathogenic” fungal species may be selected which actually vary in virulence from non-pathogenic to relatively weakly virulent to strongly virulent. Non-virulent preconidial mycelium may be used to attract beneficial predator and parasitic insects. Alternatively, non-virulent strains may be utilized as a distractors, for example attracting Coccinellidae, the lady beetles, away from areas where they may be a pest (such as office buildings) and into “ladybug motels.” Alternatively, virulent strains may be utilized as an olfactory attractant but made inaccessible with devices such as screens or slots.

[0059] The mycoinsecticides and/or mycopesticides disclosed herein may also be optionally enhanced by the use of other baits, foods, attractants, arrestants, feeding stimuliants, sex pheromones, aggregating pheromones, trail pheromones, etc. For example, a bait box overgrown with preconidial mycopesticide mycelium might contain other attractants and contact pesticides, and contain antinmicrobial, antiprotozoa, and antiviral ingredients.

[0060] Attractant preconidial or pre-sporation mycelium (virulent, weakly virulent and/or non-virulent) or extracts may also be used in conjunction with other biological organisms, chemical pesticides and physical control agents as part of integrated pest management (IPM) systems that incorporate multiple pest control tools and seek to minimize pesticide inputs. The use of attractant fungi in combination with other insect control agents affords the advantage of attracting the targeted pest to a locus, which, by other treatments, results in sterility and/or death of the targeted insect.

[0061] The weakened immune systems of pest insects exposed to pathogenic or virulent mycopestidial organisms allows other beneficial parasitic and predator species to flourish. Such beneficial biological control agents include microbial pathogens, predator insects (entomopathogenic insects which eat other insects) and parasitic insects (those which reproduce by laying eggs in or on any stage of the host insect, from egg to adult), as well as non-insect predators such as birds and beneficial nematodes, spiders, and mites. Examples of biological control agents include: entomopathogenic fungal species and their spores; Bacillus thuringiensis, B. popilliae, B. subtilis, and Pseudomonas; fire ant parasites (such as Phorid flies); fly parasites including wasps such as Muscidae raptor, Spalangia cerneri; hister beetles such as Carcinops parvus; dung beetles including Otnerophagus spp.; parasitic nematodes such as Steinernema feltiae; cockroach parasites such as Anastatus teniipes, Aprostocetus hagenowi, Compsa merceti and nematodes; lacewings; ladybugs; big-eyed bugs; damsel bugs; praying mantises; Trichogramma wasps; beneficial mites; ant parasites; flea parasites; lygus bug parasites; mealybug; aphid and whitefly pests and predators; caterpillar parasites; spider mite predators; looper parasites; diamondback and moth parasites; scale parasites and predators; mites; parasitoids and predators; etc. Strains may be selected, utilizing those methods known to the art, for virulence against the targeted pest insects and/or non-virulence or weak virulence against predator insect species as well as such qualities as resistance to pesticides, etc. If desired, resistant predator or parasitic species may be selected for, bred and released to further control the targeted pest species. Blends of beneficial insect attractant plants and habitat plants may also be utilized in combination with antimicrobial, antiprotozoa and antiviral agents. This multiphase approach is not limited to just one pairing of fungus, one beneficial organism and one anti-disease component, but as many permutations as can be implemented for the purpose of creating an environmental equilibrium affording long-term protection of the inhabitants from other insects, animals, and plants. Other fungal attractants may also be optionally utilized. Thus, a combination of the preconidial mycelium of mycopesticidal species and Oyster mushrooms (Pleurotus and Hymenogastrum species, the mycelium and mushrooms of which are very attractive to Phorid flies) might be utilized to attract phorid flies in the genus Pseudacteon that parasitize fire ants and leaf-cutter ants.

[0062] The preconidial mycopesticides (both virulent and non-virulent strains) and extracts may also be utilized as “masking agents” as well as attractants in conjunction with insect chemical control agents, toxicants and/or pesticides, thereby preventing aversion to other effective compounds that may otherwise repel the insect. Chemical control agents include insect toxicants, poisons, regulators and pesticides as well as the chemicals (semiochemicals) which mediate interactions between individuals of a insect species (pheromones) or between co-evolved species (allelochemicals, such as kaurones and allatones). Residual (persistent), non-residual (nonpersistent), and soil liquid, aerosol or fog control chemical agents include, by way of example but not of limitation: stomach poisons such as sulfochlor; pyrethrum extracts; natural and synthetic pyrethroids; parpyrethroids (non-ester pyrethroids) such as silfluofen, etofenprox and cyfluthrin; pyrethroid analogs such as fenvalerate, permethrin, phenopropthrin, flualinate, flucythrinate, fenpropathrin, cypermethrin, deltamethrin, tralomethrin, cycloprop, esfenvalerate and zeta-cypermethrin; allethrin; lethanes; nicotinyl compounds such as imidacloprid; phenylpyrazoles such as fipronil; amidonilhydrazones such as hydramethylnon (a respiratory poison); abamectin (a mixture of avermectins, insecticidal or antihelminthic compounds derived from the soil bacteria Streptomyces avermectilis); Spinosad (spinosyns, the active ingredients produced by S. spinosus); artemisinin from Artemisia plants; nitromethanes; carbamates such as propoxur and fenoxycarb; organophosphates
such as acephate and chlorpyrifos; pyriproxyfen; insect growth regulators; synthesis inhibitors; chitin synthesis inhibitors such as hexafluorunuron and diflubenzuron; mineral acids such as boric acid; alcohols and organic solvents; elements such as sulfur; and combinations thereof. Such chemical control agents may optionally be combined with synergistic compounds that increase the toxicity and/or enhance the biological activity of another, for example by inhibiting the enzymatic detoxification of insecticides by microbial oxidases or hydrolytic enzymes such as esterases. Examples of synergists include: methylenedioxyphenyl (MDP) compounds such as piperonyl butoxide, piperonal bis-(2,2-butoxyethoxy)-ethyl acetate, 1,2-methylenedioxyphenethylamine, tropial (polyalkoxy acetyl of piperonaldehyde) and sesamex; trisubstituted aliphatic and aromatic phosphates such as TOCP (tri-o-cresyl phosphate); a number of non-insecticidal carbamates; EPN (O-ethyl-O-p-nitrophenyl phenolphosphonothionate); sulfoxide; propoxyn ethers; p-nitrobenzothio cyanate; 2-((4-hydroxy-2-(2-hydroxyethyl)-oxy) triethylamine; 2-(diethylamino)ethyl 2,2-diphenyl pentanolate; 2-propynyl 4-chloro-2-nitrophenyl ether; N-octyl bicycloheptene dicarboximide; and n-propyl isoume. Use of attractant or attractant/pesticidal preconidial mycelium or extracts, in combination with antibiotics and antivirals, enables the use of extremely small amounts of toxicant or pesticide to effectively control insect populations and the diseases they transmit. Alternatively, sublethal doses of pesticides or toxicants may be included to enhance the activity and virulence of the mycotoxicological species; or pathogenic and virulent preconidial mycelium may be utilized as a preconditioning treatment, increasing the susceptibility to and/or potentiating the virulence of other agents (such as pesticidal chemicals, other mycotoxicides, or bacteriological, plasmoidal and viral compounds). Lethal or sublethal doses of insect toxicant and antibiotic materials may optionally be encapsulated within an attractant extract or mycelia-impregnated (virulent or non-virulent) sheath, coating, covering, encapsulative material, protective and/or time degrading envelope, or the toxin may surround, cover or encapsulate a mycelial substance or extract of strong attractive and/or mycotoxicidical properties, or such may be simply mixed.

The mycotoxicants and mycotoxicides of the present invention may also be combined with physical control agents. Physical control agents are devices that destroy insects directly or act indirectly as barriers, excluders, or collectors. Physical controls include the use of mechanical and electrical devices, heat, light, electricity, X-rays, lasers, and so on, to kill insects directly, reduce their reproductive capacity, or to attract them to something that will kill them. Various physical means may be employed to act as barriers to insect movement. Sticky materials in which insects become hopelessly entangled may be used in the form of flypaper or coated objects and materials. Traps may be used for control, survey, and surveillance purposes. Control traps may be used in conjunction with mycotoxicants and with some means of killing the insects that enter (e.g., a pesticide or an electrically charged grid). Mosquito or bed nets can be impregnated to attract disease-carrying insects or arthropods whereupon contact, they are trapped. If not trapped, the escaping insects and arthropods, post contact, may have their pathogenic payloads reduced. This approach has many merits—as the insects and arthropods live after making contact, but now represent less of a threat for infection and disease transmission.

The preconidial mycelium on manufactured, compressed pellets or granules, with or without additional liquid(s), can be used for applications in agricultural, forest, industrial and/or domestic settings, wherein the myceliated pellets become loci for attracting the target pests, and thus through contact become infected. Trends in mushroom spawn for gourmet and bioremediation purposes have long been evolving toward pelletized or granular spawn while mycotoxicidical spore technology similarly has evolved toward granulated or spray formulations. Various forms of pelletized spawn, coated compositions, granules and dusts are known, including those formed from nutrients, with or without carriers and binders, such as peat moss, vermiculite, alginate gel, wheat bran, calcium salts, hydrophilic materials such as hydrogel, perlite, diatomaceous earth, mineral wool, clay, polymers, biopolymers and starch, including wettable powders, emulsifiable concentrates, starch and/or biopolymer coatings, etc. Pelletized spawn is specifically designed to accelerate the colonization process subsequent to inoculation. Idealized pelletized spawn seeks a balance between surface area, nutritional content, and gas exchange and enables easy dispersal of mycelium throughout the substrate, quick recovery from the concussion of inoculation, and sustained growth of mycelium sufficient to fully colonize the substrate. See Stamets and Chilton, supra, pp. 141-142 and U.S. Patent No. 4,551,165 (1985) to Warner, U.S. Pat. No. 4,678,512 (1987) to Lewis et al., U.S. Pat. No. 4,724,147 (1988) to Marois, et al., U.S. Pat. No. 4,818,530 (1989) to Marois, et al., U.S. Pat. No. 5,008,105 (1991) to Lewis, et al., U.S. Pat. No. 5,786,488 (1998) to Lamar, et al., and U.S. Pat. No. 6,145,549 (2000) to Lamar, et al. Liquid sprays include the above wettable powders and emulsifiable concentrates, water-dispersible granules, aqueous solutions, emulsions such as oil-in-water and water-in-oil emulsions, dispersions, suspensions, microemulsions, microparticles, etc. Wettable powders are formulations that are typically uniformly dispersible in water and also contain surface-active agents (surfactants) such as wetting agents, emulsifiers and dispersing agents. Emulsifiable concentrates are prepared with organic solvents and/or one or more emulsifiers. Sticking agents such as oils, gelatin, gums, tackifiers and adhesives may be used to improve the adhesion of the spray. Humectants may also be used to decrease the rate of evaporation, including for example glycols having from 3 to 10 carbon atoms and glycerin and solutes such as salts or sugars in water.

For large scale application, fabric or fiber cloths, landscaping cloths, geofabrics, soil blankets and rugs, mats, tarps, bags, gauzes, fiber mats, brick, fiber netting, felts, tattamis, bags, baskets, etc. made of biodegradable materials infused with preconidial mycelia of mycotoxicidical species, combined with antimicrobial and antiviral agents, may be utilized as a mechanism for attracting, preventing, killing or limiting the spread of targeted insects (or attracting beneficial insects) and zoontic diseases. Thus, for example, barriers or “aprons” of mycotoxicidical mycelium grown on straw, coconut fiber, wood, paper, cardboard or the other forestry and agricultural products, wastes and cellulose sources noted above might be placed around Oak trees to protect from beetles and introduced wilt(s) such as Phytophthora and Ceratocystis fagacearum or around pine trees or stands to protect from destructive fungi and diseases carried by bark beetles. Similarly, such mycotoxicidical aprons might be utilized to protect other trees, shrubs, grasslands, rivers and streams, estuaries, riparian zones, agricultural fields, gardens
and crops, structures, communities, habitats and sensitive ecosystems. Such preconidial mycotoxicidal aprons might alternatively be used to attract pest insects to a site whereupon other biological, chemical, mechanical, electrical and/or other insect reducing treatments become more effective. Conversely, creation of buffers utilizing non-virulent strains selected for attractiveness to beneficial insects can be used to attract beneficial species, which naturally parasitize problem insects.

[0066] Alternatively, woodchips, grains, hydromulch and other substrates infused with preconidial mycelium may be utilized in spray hydroseeding or mobile hydroseeding. Agricultural equipment may be utilized to inoculate fields and agricultural wastes. The mycotoxicidal fungi may also optionally be utilized in conjunction with saprophytic fungi and mycorrhizal fungi to enhance soils and agricultural yields (“companion cultivation” of beneficial fungi). Mycotoxicicial species are also useful in the mycoremediation (fungal bioremediation) of various sites. As one example, reclaimed logging roads could become perimeter-barriers which could forestall and/or prevent beetle-plagues from devastating forests by infusing mycomats or hydromulches with species-specific pathogenic fungi (and optionally saprophytic and mycorrhizal fungi), combined with antimineral or antiviral agents, while simultaneously retaining other benefits of mycoremediation. Thus, mycotoxicisic species such as Mortierellum, Beauveria and Cordyceps, mycorrhizal mycotoxicisic fungi such as Laccaria, and mycenothecial saprophytic fungi such as Pleurotus might be combined with ectomycorrhizal and endomycorrhizal species and saprophytic fungi to provide simultaneous insect control, road reclamation and protection of streams from silt runoff and disease control. As Hypoloma capnoides, a premier wood chip decomposer, mycelium has been observed to be repellent to insects, stretches of insect repellant barriers may be combined with attractant mycotoxicidal kill and/or control zones for insects such as wood-boring beetles. Similarly, control of agricultural runoff utilizing saprophytic fungi on agricultural wastes might be combined with the present mycocontactant and/or mycotoxicisic applications, while in combination with antimineral and antiviral agents, to limit the spread of disease.

[0067] In general, preferred mycotoxicisic species as pathogens are somewhat slow-acting (that is, not immediately fatal) so as to avoid bait shyness and to avoid learning effects in social insects before individuals have distributed mycelium to all other members of the colony. To effect control of the Formosan subterranean termites (Coptotermes formosanus) colonies, bait chemicals must kill slowly enough to allow foraging termites to return to the colony and spread the toxin to other colony members. Wright et al., “Growth response of Mortierellum anisopliae to two Formosan subterranean termite nest volatiles, naphthalene and fenochlor,” Mycologia, 92(1): pp. 42-45 (2000) and the references therein. Bait shyness and other colony defense mechanisms such as segregation or avoidance of infected nestmates or necrophoretic behavior by the workers (i.e., removal of dead nestmates) serves as a means of defense against the spread of such pathogens when the targeted insect dies too quickly. For example, in general, queen fire ants will not feed on new foodstuffs until the food is first sampled by foragers or workers in members of expendable classes and deemed safe after a two or three day waiting period. Note, however, this general pattern may not always apply to the highly attractive mycocontactant and mycotoxicisicides disclosed herein. Preconidial mycelium strains may be selected for virulence after an appropriate time period. In many applications it may be preferable to utilize a mixture or matrix of several species or strains of entomopathogenic fungi with different characteristics, maturation and growth rates including strains with delayed sporulation (and thereby prolonged attractant value) while in other applications a single species may be preferred. Similarly, with reference to a single species, a mixture of strains or a single strain may be utilized. A mixture of species and/or strains both allows the targeted insects to choose the species to which they are most attracted and provides for the possibility of simultaneous infection and insect plagues from multiple virulent species and strains. This makes tolerance or resistance of the insect or arthropod much more unlikely compared to just using one strain or anticoenobial agent.

[0068] Those skilled in the art will recognize that numerous entomogenous and entomopathogenic fungal species are known to the art and the above preconidial mycocontactant and mycotoxicisic methods and products may be favorably applied to many or all such species, and it is the intent of the inventor that the invention be understood to cover such. Suitable entomopathogenic fungi include: the Deuteromycetes Mortierellum, Beauveria, Paecilomyces, Hirataella, Verticillium, Calicifomycet, Nomuraea, Aspergillus and other fungi imperfecti; sexually reproducing fungi such as the Ascomycetes Cordycep, Ophiocordycep, Ascocherat, Torulohella, Hypocrella and its Aschersonia anamorph, and the Paecilomycte Laboulbenia hagenii; the Basidiomycetes such as Laccaria, Pleurotus, Fomes, Fomitopsis, Hypyizygos, Piptoporus, Lenzites, Ganozoma, and combinations thereof. The Entomophorataceae including Entomophaga, Massospora, Neozobites, Zoophthora, Pandora and other Phycomycetes are also considered to be within the scope of the invention. Also included are such entomopathogenic species that have been genetically modified to be more virulent (including those modified via mutagenesis, hybridization and recombinant DNA techniques).

[0069] By way of example, but not of limitation, mycotoxicisic species include Metarhizium anisopliae (“green muscarine”), Mortierellum flaviride, Beauveria bassiana (“white muscarine”), Beauveria bronchii, Paecilomyces farinosus, Paecilomyces fumosoroseus, Verticillium lecanii, Hirataella citriniformis, Hirataella thompsoni, Aschersonia aleyrodias, Entomophaga gyri, Entomophaga maimaiga, Entomophaga muscae, Entomophaga pravissil, Entomophthora phutelae, Zoophthora radicans, Neosyzygites floridana, Nomuraea rileyi, Pandora neoaphisid, Tolypocladium cylindrosorum, Calicifomycet clavosporus and Lagendium giganteum, the wide variety of Cordycep (and Ophiocordycep) and its ascomycetous forms including Cordycep variabilis, Cordycep fac, Cordycep (Elaphocordycep) subessilis, Cordycep myrmecophile, Cordycep sphecocephala, Cordycep entomorrhiza, Cordycep gracilis, Cordycep militaris, Cordycep washingtonensis, Cordycep meloanathae, Cordycep ravenelli, Cordycep unilateralis, Cordycep sinensis and Cordycep clavulata, and mycorrhizal species such as Laccaria bicolor. Other mycotoxicisic species will be apparent to those skilled in the art.

[0070] The concepts of “preconidial” and “spores” or “conidia” are complex, containing a number of different forms and specialized structures for reproduction of fungi. Many fungi are pleomorphic, that is, one fungus may produce several sorts of spores, which may or may not be coincident in
time. With regard to the sexually reproducing *Cordyceps, Laccaria* and other “fungi perfecti,” preconidial or pre-sporulation refers to the pre-fruiting stage. The term “preconidial” or “pre-sporulation” has a somewhat different meaning with regard to the sexually reproducing fungi than with most other entomopathogenic fungi, as sexually reproducing fungi are “fungi perfecti” or mushroom fungi, whereas the non-mushroom fungi such as *Beauveria* and *Metarhizium* are the more primitive “fungi imperfecti.” This situation is complicated by the fact that entomophthoralean fungi have complex life cycles involving non-sexual conidia and sexual resting spores. The situation is further complicated by the fact that some or all *Cordyceps* fungi are dimorphic and have a teleomorphic (the sexual perfect) form or morph, e.g. that characterized by sexual spores including ascospores and basidiospores and one or more anamorphs (the ascum perfect form or morph, e.g. characterized by the presence or absence of conidia) with conidial stages within the imperfect fungal genera including *Beauveria, Metarhizium, Paecilomyces, Hirutella, Verticillium, Aspergillus*, *Akanthomyces*, *Desmidioспорa*, *Hymenositoide*, *Mariannaea*, *Nomuraea*, *Parasaria*, *Tolypocladium*, *Spicaria* (= *Isaria*) ; and *Botrytis*. For example, *Cordyceps subsessilis* is the perfect form of *Tolypocladium inflatum*, an anamorph (imperfect) form which produces andAPER. *Cordyceps militaris* (Fr.) L. is also thought to be dimorphic, the conidial stage of which is believed to be a *Cephalosporium*. *Cordyceps unilateralis* seems specific on *Camponotini*, while *Hirutella sporoachialis* is probably an anamorph of *Cordyceps unilateralis* specific on *Polyrhachis*. Schmid-Hempel, supra, p. 43. The situation is further complicated in that conidia, without ascis, have often been observed in *Cordyceps* by the inventor. DNA studies are expected to better elucidate these relationships. As used herein, unless otherwise specified, preconidial or pre-sporulation mycelium of sexually reproducing fungi refers to the pre-sporulation mycelial stage of the mushrooms, including any preconidal imperfect stages and any preconidial sclerotia or microsclerotia.

[0071] It is further expected that the preconidial products and methods may, with no more than routine experimentation, prove useful against preconidial, parasocial, subsocial and non-social insects including semisocial, quasisocial, communal and solitary insect pests such as: cockroaches including American, German, Surinam, brown-banded, smokybrown, and Asian cockroaches; grasshoppers and locusts; crickets including mole cricket; Mormon crickets (actually a long-horned grasshopper); beetles, beetles grubs and beetle larvae, including Colorado potato beetle (*Leptinotarsa decemlineata*) and other potato beetles, Mexican bean beetle, Japanese beetle, cereal leaf beetle, darkling beetle (lesser mealworm); moths including Gypsy moths (*Lymantria dispar*) and Gypsy moth larvae, diamondback moths (*Plutella xylostella*), codling moth (*Laspeyresia pomonella*), Douglas fir tussock moth (*Orgyia pseudotsugata*), western spruce budworm (*Choristoneura occidentalis*), and grape berry moths (*Lobesia botina*); flies and fly larvae; springtails; large centipedes; shield centipedes; millipedes; European corn borers (*Ostrinia nubilalis*); Asiatic corn borers; caterpillars including velvetbean caterpillar (*Anticarsia geminata*), and other caterpillars and larvae of the Lepidoptera; whiteflies (*Dialeurodes* and *Bemisia*); leafhoppers; and silverleaf whiteflies; thrips (*Thrips spp.*); including melon thrips (*Thrips palm*), and western flower thrips (*Frankliniella occidentalis*); aphids including Russian wheat aphid; spider mites (*Tetranychus spp.*); mealybugs including citrus mealybug (*Planococcus citri*) and solanum mealybug (*Pseudococcus solani*); boll weevils, black vine weevils (*Otiorychnum solncatus*), European pear weevils (*Curculio pyraster*); mosqui- toes; wasps; cotton fleahoppers; pasture scarabs such as *Ado- ryphora columbia* and other Scarabaeidae; spittle bug (*Mahanarva posticata*); corn earworm (*Helicoverpa zea*); American bollworm (*Heliothis armigera*); armyworms including *Pseudaelia unipuncta*, fall armyworm (*Spodoptera frugiperda*), southern armyworm (*Spodoptera eridania*), beet armyworm (*Spodoptera exigua*), and yellow-striped armyworm (*Spodoptera ornithogallii*); black cutworm (*Agrotis ipsilon*); tobacco hornworm (*Manduca sexta*); tobacco budworm (*Helicoverpa* syn. *Helicoverpa* vic- senc); sugar cane frog hopper; rice brown planthopper; ear- wigs; loopers including cabbage looper (*Trichoplusia ni*); soy- bean looper (*Pseudoplusia includens*), forgan looper (*Caucumina erechthea*) and celery looper (*Anaglyphus fal- cifera*); cabbageworms including the imported cabbageworm (*Pieris rapae*) and the European cabbageworm (*Pieris brassicae*); tomato pinworm (*Keiferia lycopersicella*); tomato hornworm (*Manduca quinquemaculata*); leafminers (*Liri- onza spp.*); cotton leafworm (*Alabama argillacea*); corn rootworm; garden webworm (*Achryya rautalis*); grape leaf- folder (*Desmia funeralis*); melonworm (*Diaphania hyalina*); pickleworm (*Diaphania nitidalis*); achen worm (*Eumorpha achenia*); sweet potato hornworm (*Agrms cingulata*); whitelined sphinx (*Hyles lineata*); lygus bugs (*Lygus spp.*); chinch bugs including *Blissus leucopterus* and false chinch bugs; sow bugs; pill bugs; citrus rust mite; pill wood lice; wheat cockchafer; white gnats and cockchafer; *Hop- loclehis marginalis* and *Melolontha melolontha*; storage pests such as *Prostephanus truncatus* and *Sitophilus zeamais*; soil insects; and various other insect pests in the orders, Isopoda, Diplopoda, Chilopoda, Symphyta, Thysanura, Collembola, Orthoptera, Dermaptera, Anoplura, Mallophaga, Thysan- optera, Heteroptera, Homoptera, Lepidoptera, Coleoptera, Diptera, Siphonaptera, Thysanoptera, Acarina, Arachnida, etc. and the families Plutellidae, Acrididae, Tettigoniidae, Gryl- lidae, Grylloptilidae, Pyralidae, Sphingidae, Noctuidae, Pyralidae, Xyphopilidae, Scarabaeidae, Scythridae, Platypo- didae, Lymexyloidae, Nitidulidae, Pseudococcidae, Aphididae, Delphacidae, Coccididae, Cercopidae, Aleyrodidae, Cocc- idae, etc. It will be recognized that the insects listed above are representative examples of insects and arthropods which may be attracted and/or controlled according to the present invention, but such listing is not intended as a limitation to certain species as numerous other insect and arthropod specie to which the invention may be applied will be apparent to those skilled in the art.

[0072] It will be noted from the discussion above and examples and results below that attractiveness, pathogenicity and virulence toward the targeted insect are dependent in some degree upon factors including choice of mycopicidal species, host range and specificity, selection of a strain within that species and selection of substrate. Entomopathogenic fungi also vary greatly in host specificity. Some entomopathogenic fungi are highly specific, such as *Pandora neophilidis*, which is restricted to aphids. Other entomopatho- genic fungi have wide host ranges, such as *Beauveria bassiana* and *Bacillus thuringiensis* which is known to affect over 100 species of arthropods. Other species with wide host ranges include *Metarhizium anisopliae*, *Paecilomyces farinosus* and *Zoophthora radi-
cans. However, in the laboratory, isolates of fungi with wide host ranges are generally more virulent to the host from which they were first isolated; certainly their host range is much more restricted than that of the species to which they belong. Goettel et al., “Safety to Non-target Vertebrates of Fungal Biocontrol Agents,” in: Laird et al. (eds.) Safety of Microbial Insecticides, pp. 209-232 (1990). Furthermore, fungi with wide host ranges are frequently even more specific under field conditions. There are reports of fungi attacking only one host even though closely related host species are present. Discrepancies between reports of social insect host specificity may be related to a general difference between tropical vs. temperate habitats rather than to the specific fungi and social insect species involved. Schmid-Hempel, supra at p. 44. Such specificity is thought to be due to the complex biotic and abiotic interactions in the field. This indicates that it should be possible, using no more than routine experimentation and bioassays of mycophagistic strains and of the appropriate orders, families, genera, species and varieties of targeted pest insects, to isolate and use strains and substrates wherein the desired characters are maximized with respect to either a targeted insect or targeted insect group, thereby producing a species-specific, genus-specific, family-specific or order-specific entomopathogenic host specific fungal strain. Such entomopathogenic strain selected for host range and specificity may be similarly selected for minimal or no infection, or virulence towards beneficial insects or non-targeted insects.

EXAMPLE 1
Attracting and Controlling Mosquitoes, Which Can Carry Viruses

[0073] Rice colonized by preconidial mycelium of Metarhizium anisopliae (ATCC #62716, and “F52”) fungus clearly attracted Aedes aegypti females. Using an olfactometer in choice tests, the mycelium grown on rice attracted the female mosquitoes significantly over the controls. By comparison in the olfactometer, response of these host-seeking Aedes aegypti to a hand is about 83% to CO2 (Allan et al. 2006). Combining the preconidial mycelium and the extracts from the same mycelium resulted in attractiveness of mosquitoes to more than 80% equivalency to a human hand, far more so than the mycelium or extract alone. Since the actively growing mycelium is also outgassing carbon dioxide (but the extract does not), the added attractiveness of using an ethanolic/water extract is significant. Aedes mosquitoes spread viruses such as yellow fever, Chikungunya fever, and Dengue fever. Adding antiviral medicines previously proven useful, or yet to be discovered, to the extracts or mycelium of the preconidial entomopathogenic fungus, would abate the spread of disease, whether or not insect mortality occurred.

EXAMPLE 2
Attracting and Controlling Mosquitoes, Which Can Carry Malaria Protozoa

[0074] Prepare mycelium and extracts by the methods described herein. Mix in DDT, chemical pesticides, purified artemesinin or its crude, less expensive precursors, to the extracts and mycelium from preconidial entomopathogenic fungi such as Metarhizium anisopliae to bait and control stations, nets, or into standing water. Place these mixtures in environments where the mosquitoes exist, including Anoph-
es gambiae or any of its 30-40 species relatives, all of which carry Malaria protozoa (Plasmodium falciparum).

EXAMPLE 3
Attracting and Controlling Flies, Which Can Carry Viruses

[0075] Prepare the preconidial mycelium and extracts of the preconidial mycelium Metarhizium anisopliae according the methods described previously and blend with ribavirin, oseltamivir, and other antiviral drugs in pure or crude form to preconidial extracts and/or mycelium of Metarhizium anisopliae to attract house flies or blow (“blue bottle”) flies and upon contact or ingestion, reduce the viral loads of the viruses they carry, thus reducing their contagiousness.

EXAMPLE 4
Attracting and Controlling Flies, Which Can Carry Bacterial and Protozoa Pathogens

[0076] Prepare the preconidial mycelium and extracts of the preconidial mycelium Metarhizium anisopliae according the methods described previously and blend with antimicrobial agents active against bacteria and protozoa. Use this blend to attract Tsetse fly carrying species of the protozoan genus Trypanosoma causing often-fatal “sleeping sickness.” Use this blend to attract house flies (Musca domestica) and Blow Flies (Calliphoridae, Calliphora vicina, and related species), which carry the pathogens Staphylococcus aureus, Streptococcus pyogenes, Bacillus anthracis, Listeria, Salmonella, Clostridium, and Enterococci, which subsequent to contact, result in reduced pathogen payloads and infectivity.

EXAMPLE 5
Attracting and Controlling Ants, Which Can Carry Pathogenic Bacteria

[0077] Prepare the preconidial mycelium and extracts of the preconidial mycelium Metarhizium anisopliae according the methods described previously and blend with antimicrobial agents active against bacteria and protozoa. Use this blend to attract ants, such a Pharaoh ants and Fire Ants carrying pathogenic bacteria (Salmonella, Staphylococcus, Streptococcus, and Clostridium, etc.) resulting in reductions in their pathogens, making them less contagious and less infectious.

EXAMPLE 6
Attracting and Controlling Cimex Species (Bed Bugs), Which Carry Pathogenic Bacteria

[0078] Prepare the preconidial mycelium and extracts of the preconidial mycelium Metarhizium anisopliae according the methods described previously and blend with antimicrobial agents active against Staphylococcus aureus bacteria. Use this blend to attract and control bed bugs resulting in reductions in their levels of Staphylococcus aureus bacteria, making them less contagious, reducing infectivity.

EXAMPLE 7
Attracting and Controlling Lice and Ticks, Which Carry Pathogenic Bacteria

[0079] Prepare the preconidial mycelium and extracts of the preconidial mycelium Metarhizium anisopliae according
the methods described previously and blend with antimicrobial agents active against *Rickettsia* spp. (the cause of Rocky Mountain Spotted fever), *Bartonella vinsonii* and *B. henselae* causing intramuscular infections, *Borreliia burgdorferi* causing Lyme disease. Use this blend to attract and control pathogen-bearing lice and ticks, resulting in reductions in their levels of pathogenic bacteria, making them less contagious, reducing infectivity.

**EXAMPLE 8**

Attracting and Controlling Flies, Which Carry Pathogenic Bacteria

[0080] Prepare the preconidial mycelium and extracts of the preconidial mycelium *Metarhizium anisopliae* according the methods described previously and blend with antimicrobial agents active against the bacteria *Yersinia pestis* causing bubonic plague. Use this blend to attract and control pathogen-bearing flies, resulting in reductions in their levels of pathogenic bacteria, making them less contagious, reducing infectivity.

**EXAMPLE 9**

Attracting and Controlling Midges, Which Carry Pathogenic Viruses

[0081] Prepare the preconidial mycelium and extracts of the preconidial mycelium *Metarhizium anisopliae* according the methods described previously and blend with antiviral agents active against viruses (Blue tongue virus to cattle, epizootic hemorrhagic diseases). Use this blend to attract and control pathogen-bearing midges, resulting in reductions in their levels of pathogenic bacteria, making them less contagious, reducing infectivity.

**EXAMPLE 10**

Attracting and Controlling Flies Carrying Viruses

[0082] Prepare the preconidial mycelium and extracts of the preconidial mycelium *Metarhizium anisopliae* according the methods described previously and blend with extracts of polypropylene mycelium such as *Fomitopsis officinalis*, *Fomitopsis piniola*, *Fomitopsis robustus*, *Piptoporus betulinus*, *Trametes versicolor*, *Trametes elegans*, *Ganoderma lucidum*, *Ganoderma applanatum*, *Ganoderma annulans*, *Ganoderma oregonense*, *Ganoderma resinaceum*, *Ganoderma tsugae*, *Heterobasidion annosum*, *Inonotus obliquus*, *Antrodia camphorata*, *Rigidoporus ulmarius*, *Pensilpinia fuscascens*, *Psilocybe cyanescens*, *Psilocybe azurescens*, *Psilocybe cubensis* and other mushroom-derived antiviral drugs in pure or crude form to preconidial extracts and mycelium of *Metarhizium anisopliae* to attract house flies or Blow (“blue bottle”) flies and upon contact, reduce the viral loads they carry, thus reducing their contagiousness.

**EXAMPLE 11**

Attract and Control Flies to Insect Control Devices

[0083] Add preconidial extracts and/or mycelium of *Metarhizium anisopliae* (prepared according the methods described previously, and blended with antimicrobial and antiviral agents) to insect trapping and killing contraptions used for limiting the spread of zoonotic disease such as “bug zappers” (BASF’s Vector™), forced airflow (fan) trapping systems, CO₂ emitters, laser target-and-kill systems, soap systems, sticky mats, and bug nets, resulting in reducing the threat of the contagions flying insects carry.

**EXAMPLE 12**

Attracting and Controlling Disease-Bearing Insects with Cellulosic Materials

[0084] Add preconidial extracts and/or mycelium of *Metarhizium anisopliae* (prepared according the methods described previously, and blend with antimicrobial and antiviral agents) to fabric clothes, burlap sacks, wood chips, straw, to attract insects and arthropods carrying pathogens that results in a reduced pathogen load within these insects and arthropods subsequent to contact.

**EXAMPLE 13**

Attracting Mosquitoes to Attract Disease Carrying Bats and Birds

[0085] Prepare the preconidial mycelium and extracts of the preconidial mycelium *Metarhizium anisopliae* according the methods described previously and blend with antimicrobial and antiviral agents active against the contagions carried by disease carrying bats and birds. Use this blend to attract mosquitoes and other flying insects, which in turn will attract and control the movement of bats and birds. The ingestion of the insects, now carrying antimicrobial and antiviral agents, can then reduce the pathogen payload of the bats and birds, thereby reducing contagion risk.

**EXAMPLE 14**

Blending Antiviral Drugs with Extracts and Mycelium of Preconidial Entomopathogenic Fungi

[0086] Blend the extracts or mycelia of preconidial entomopathogenic fungi with the less expensive antiviral drug precursors, expired antiviral drugs, or antiviral drugs such as Abacavir, Aciclovir, Acelovir, Adelovir, Amantadine, Amprenavir, Ampiglen, Arbidol, Atazanavir, Atripla, Boceprevir, Cidofovir, Combidar, Darunavir, Delavirdine, Didanosine, Diducanol, Efinavir, Efavirenz, Emtricitabine, Enufavir, Etenovir, Foscarnet, Fosinovir, Fosamiprovir, Foscaritin, Fosfenet, Ganciclovir, Ibbacitabine, Imunovir, Idoxuridine, Imiquimod, Indinavir, Insinovir, Interferon type III, Interferon type II, Interferon type I, Interferon, Laminovir, Lopinavir, Loviride, Maraviroc, Moroxydine, Methisazone, Nelfinavir, Nevirapine, Nexavir, Nucleoside analogues, Oseltamivir (Tamiflu), Peginterferon alfa-2a, Peniclovir, Peramivir, Pleconaril, Podophyllotoxin, Protease inhibitors, Raltegravir, Reverse transcriptase inhibitor, Ribavirin, Rimantadine, Ritonavir, Pyrimidine, Saquinavir, Stavudine, Tea tree oil, Tenofivir, Tenovir dispersible, Tipranavir, Trifluridine, Trizivir, Truvanavir, Valaciclovir (Valtrex®), Valganciclovir, Vincristine, Vidarabine, Viramidine, Zaletabine, Zanamivir (Relenza®) and Zidovudine to attract disease carrying insects and arthropods, and upon contact or ingestion, reduce their pathogenic payload, thus reducing their contagiousness, and limiting disease transmission.

**EXAMPLE 15**

Blending Antibacterial Drugs with Extracts and Mycelium of Preconidial Entomopathogenic Fungi

[0087] Blend the extracts or mycelia of preconidial entomopathogenic fungi with the less expensive antibacterial
drug precursors, expired antibacterial drugs, or antibacterial drugs such as Amoxycillin, Ampicillin, Cipro, Duricef, Erythromycin, Floxin, Levaquin, Roxithromycin, Suprax, and Zithromax to attract disease carrying insects and arthropods, and upon contact or ingestion, reduce their pathogenic payloads, thus reducing their contagiousness, and limiting disease transmission.

EXAMPLE 16

Blending Antiviral Drugs with Extracts and Mycelium of Preconidial Entomopathogenic Fungi to Protect Plants From Viral Diseases

[0088] Blend the extracts or mycelia of preconidial entomopathogenic fungi with antiviral drugs that protect plants to attract disease carrying insects and arthropods, and upon contact or ingestion, reduce their pathogenic payloads, thus reducing their contagiousness, and limiting disease transmission, thus protecting plants.

[0089] Leafhoppers, and white flies, which transmit viruses to plants, can be attracted to the extracts and mycelium of preconidial entomopathogenic fungi and limit viral disease transmission. Moreover, when antiviral drugs or their less pure, crude precursors are employed in combination with the extracts of preconidial entomopathogenic mycelium or with the preconidial mycelium of entomopathogenic fungi, the viral transmission threat from white flies and leaf hoppers is reduced or eliminated, thus saving crops from the damaging effects of viruses. Two exemplary examples are the beet leafhopper, Circulifer tenellus spreads curly top virus, Macrosytes facsiifrons spreads mycoplasma to hundreds of plants, including many vegetables. Additionally, hundreds of species in family Cicadellidae transmit plant diseases, many of which are viruses.

EXAMPLE 17

Blending Extracts and Mycelium of Preconidial Entomopathogenic Fungi with Genetically Modified Gene Sequences

[0090] Blend extracts of preconidial entomopathogenic mycelium or with the preconidial mycelium of entomopathogenic fungi to attract and control insects and arthropods that transmit contagions that harm plants, and which results in making contact with genetically modified gene sequences, further resulting in the protection of plants from viruses and other contagions carried by insects and arthropods.

Example 18

Blending Extracts and Mycelium of Preconidial Entomopathogenic Fungi With Bacteriophages to Limit Disease Transmission

[0091] Blend extracts of preconidial entomopathogenic mycelium or with the preconidial mycelium of entomopathogenic fungi to attract and control insects and arthropods that transmit contagions that harm plants and animals with bacteriophages, thus protecting plants and animals by the effect of the bacteriophages' ability to reduce or fend off transmissible diseases.

[0092] No limitations with respect to the specific embodiments and examples disclosed herein are intended or should be inferred, as the examples and embodiments are representative only. While examples and preferred embodiments of the present invention have been shown and described, it will be apparent to those skilled in the art, or ascertainable using no more than routine experimentation, that many changes and modifications may be made without departing from the invention in its broader aspects. The appended claims are therefore intended to cover all such changes, modifications and equivalents as fall within the true spirit and scope of the invention.

1 claim:

1. A method comprising using a preconidial preparation of an entomopathogenic fungus to attract and control arthropods that carry zoonotic diseases, wherein the preconidial preparation is selected from the group consisting of preconidial mycelium and extract of preconidial mycelium, wherein non-sporulating sectors are selectively cultured to produce the preconidial mycelium, and wherein the arthropods are selected from the group consisting of insects, arachnids and myriapods.

2. The method of claim 1 wherein the insects are selected from the group consisting of malaria-carrying mosquitoes, virus-carrying mosquitoes, contagion-carrying flies, contagion-carrying ants, contagion-carrying bedbugs, carrying-flies and contagion-carrying midges and the arachnid is selected from the group consisting of virus-carrying ticks and contagion-carrying ticks.

3. The method of claim 1 wherein the entomopathogenic fungus is selected from the group consisting of Deuteromycetes Metarhizium, Beauveria, Paecilomyces, Hirsutella, Verticillium, Cucilinomyces, Nomuraea and Aspergillus, the Ascomycetes Cordyceps, Ophiocordyceps, Ascochaeta, Toxobilia, Hypocrella and its Aschersonia annamor, the Pyrenomycete Laboulbenia hagern, the Entomophthoraceae Entomophaga, Massospora, Neozymes, Zoophtora and Pandora, the Phycomyctes and combinations thereof.

4. The method of claim 1 wherein the entomopathogenic fungus is selected from the group consisting of Metarhizium anisopeiae, Metarhizium flavirete, Beauveria bassiana, Beauveria bronniartii, Paecilomyces farinosus, Paecilomyces fumosoroseus, Verticillium lecanii, Hirsutella citriformis, Hirsutella thompsoni, Aschersonia alyroides, Entomophaga grylli, Entomophaga mainaiga, Entomophaga muscae, Entomophaga praxibuli, Entomophthora plutelae, Zoophthora radicans, Neogyzyges floridanus, Nomuraea rileyi, Pandora neoaphisdis, Toxoplocladium cylindrosporum, Cucilinomyces clavosphorus, Lagenidium giganteum, Cordyceps variabilis, Cordyceps fasci, Cordyceps (Ophiocordyceps) subsectis, Cordyceps myrmecophilus, Cordyceps spherocephala, Cordyceps entomorrhiza, Cordyceps granulis, Cordyceps militaris, Cordyceps washingtonensis, Cordyceps melolonthae, Cordyceps raveneliai, Cordyceps unilateralis, Cordyceps sinensis, Cordyceps clavulata, and combinations thereof.

5. The method of claim 1 wherein the entomopathogenic fungus is selected from the group consisting of Cordyceps sinensis, Cordyceps subsectis, Cordyceps militaris, Cordyceps unilateralis, Ophiocordyceps species and their anamorphs including Metarhizium, Beauveria, Paecilomyces, Hirsutella, Beauveria and combinations thereof, wherein the insects are selected from the group consisting of animal biting insects, non-animal biting insects, mosquitoes, flies, bed bugs, fleas and mites and wherein the arthropods are selected from the group consisting of animal-biting arthropods, non-animal biting arthropods, spiders, ticks and mites.

6. The method of claim 1 wherein the insects and the arthropods that carry zoonotic diseases are attracted to a locus.
where disease carrying insects and arthropods and their diseases can be better controlled.

7. The method of claim 1 wherein the preconidial preparation contains both preconidial mycelium and extract of preconidial mycelium.

8. The method of claim 1 wherein the preconidial mycelium contains less than 100 conidia per gram of myceliated substrate.

9. The method of claim 1 wherein a mosquito net is impregnated with the extract of preconidial mycelium and an antimicrobial drug, whereby mosquitoes that make contact with the mosquito net thereby have a subsequent contagion load reduction after contact or consumption.

10. The method of claim 1 whereby bats and birds have their contagion payload reduced by consuming insects that have been previously attracted to and consumed extracts from preconidial entomopathogenic mycelium combined with antiviral and antimicrobial drugs and the insects then have reduced pathogen payloads and contain residues of the antiviral and antimicrobial drugs, thus conferring a disease reducing benefit to previously contagion-infected birds and bats.

11. The method of claim 10 wherein the preconidial preparation is combined with antimicrobial drugs selected from the group consisting of antibacterial drugs, antiviral drugs and antiprotozoan drugs in order to attract and control insects and arthropods that carry zoonotic diseases to a locus wherein their pathogen payload is thereby reduced after contact or ingestion.

12. The method of claim 11 wherein the antimicrobial drugs are selected from the group consisting of below-pharmaceutical grade drugs, crude forms of drugs, drugs derived from natural products, expired drugs, partially purified drugs and combinations thereof.

13. The method of claim 1 further comprising blending the preconidial preparation of an entomopathogenic fungus with derivatives of mushroom preparations and mushroom mycelium preparations selected from the group consisting of Fomitopsis officinalis, Fomitopsis pinicola, Fomitopsis robustus, Piptoporus betulinus, Trametes versicolor, Trametes elegans, Ganoderma lucidum, Ganoderma applanatum, Ganoderma annulans, Ganoderma oregoneum, Ganoderma resinaceum, Ganoderma tsugae, Heterobasidion annosum, Inonotus obliquus, Antrodia camphorate, Rigidosporus ulmarius, Perenoporia fraxinophila, Psilocybe cyanescens, Psilocybe azurescens and Psilocybe cubensis preparations, extracts, derivatives, drugs and combinations thereof.

14. The method of claim 1 wherein the preconidial preparation is further blended with an antimicrobial substance selected from the group consisting of antimalarial drugs, the crude precursors from which antimalarial drugs are derived, Quinine, Chloroquine, Amodiaquine, Pyrimethamine, Proguanil, Sulfonamides, Mefloquine, Atovaquone, Primaquine, Halofantrine, Doxycycline and Claralamycin, whereby the preconidial preparation attracts mosquitoes harboring malaria protozoa and other diseases, and wherein upon contact or ingestion, pathogenic payloads of malaria protozoa and other diseases within the mosquitoes are reduced, thus reducing their contagiousness and limiting disease transmission.

15. The method of claim 1 wherein the preconidial preparation is further combined with an antiviral selected from the group consisting of less expensive antiviral drug precursors, expired antiviral drugs, Aciclovir, Acyclovir, Adefovir, Amantadine, Amprovir, Ampigen, Arbidol, Azasanavir, Atipil, Boceprevir, Cidofovir, Combivir, Darunavir, Entavir, Didaoxin, DDCosanol, Efavirenz, Etravirine, Emtricitabine, Efaviride, Emecavir, Fomivirsen, Fosamprenavir, Foscarnet, Fosapron, Ganciclovir, Ibricitabine, Imonovir, Ixodixuride, Imitrividol, Indinavir, Ionofox, Interferon type III, Interferon type II, Interferon type I, Interferon, Lamivudine, Lopivir, Loviride, Maraviroc, Moroxidine, Methiszone, Nelfinavir, Nevirapine, Nexavar, Nucleoside analogues, Oseltamivir (Tamiflu®), Peginterferon alfa-2a, Peniclovir, Peramivir, Pleconaril, Podophyllotoxin, Protease inhibitor, Raltegravir, Reverse transcriptase inhibitor, Ribavirin, Rimantadine, Ritonavir, Ritonavir, Saquinavir, Stavudine, Tea tree oil, Tenofovir, Tenofovir disoproxil, Tipranavir, Trifluridine, Trizivir, Truvada, Valaciclovir (Valtrex®), Valganciclovir, Vicriviroc, Vidarabine, Viramidine, Zalcitabine, Zanamivir (Relenza®), Zidovudine and combinations thereof, which attract disease carrying insects and arthropods, and upon contact or ingestion, reduce pathogenic payloads of contagions in the disease carrying insects and arthropods, thus reducing their contagiousness, and limiting disease transmission.

16. The method of claim 1 further comprising blending the preconidial preparation of an entomopathogenic fungus with an antibacterial selected from the group consisting of antibacterial drug precursors, expired antibacterial drugs, antibacterial drugs, Amoxicillin, Amoxicillin, Cipro, Duricef, Erythromycin, Floxin, Levaquin, Roxithromycin, Suprax, Zithromax and combinations thereof in order to attract disease carrying insects and arthropods, upon contact or ingestion, reduce their pathogenic payloads, thus reducing their contagiousness, and limiting disease transmission.

17. The method of claim 1 comprising blending a preconidial preparation of an entomopathogenic fungus with bacteriophages that protect plants and animals from disease carrying insects and arthropods in order to attract the insects and arthropods, and upon contact or ingestion, reduce their pathogenic payloads, thus reducing their contagiousness and limiting disease transmission, thus protecting plants and animals.

18. A method comprising blending a preconidial preparation of an entomopathogenic fungus with antimicrobial substances that protect plants from disease carrying insects and arthropods in order to attract the insects and arthropods, and upon contact or ingestion, reduce their pathogenic payloads, thus reducing their contagiousness and limiting disease transmission, thus protecting plants.

19. The method of claim 18 wherein the preconidial preparation of an entomopathogenic fungus is selected from the group consisting of preconidial mycelium, extracts of preconidial mycelium and combinations thereof and the insects and arthropods are selected from the group consisting of Cicadellidae and whiteflies.

20. The method of claim 19 wherein the antimicrobial substances are selected from the group consisting of antiviral drugs, antibacterial drugs, crude precursors of antimicrobial drugs, expired antimicrobial drugs, antimicrobial drugs of less than pharmaceutical grade, natural substances containing antimicrobial drugs and combinations thereof.