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Disease-Causing Fungi in Homes and Yards in the Midwestern United States

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Abstract

A number of fungal pathogens that may result in a variety of human diseases are found in residential homes and yards. The growth of these microscopic fungi is often favored by particular characteristics of the dwelling and nearby outdoor environment. Evolved virulence factors or increased ability of specific fungi to grow in diverse, and sometimes harsh, microenvironments presented by the domestic environment may promote growth and pathogenesis. Infection may occur by inhalation or direct inoculation and include endemic fungi in addition to opportunistic or emerging species. Systemic or locally aggressive fungal infections are particularly likely and may be life-threatening in those with compromised immune systems. Allergic disease may include sinusitis, pneumonitis and immediate hypersensitivity. Controversial topics include mycotoxins, volatile organic compounds and sick building syndrome. This narrative review describes the usual presentations, domestic environmental sources, prevention techniques and risk of acquiring these diseases in the Midwestern United States. (J Patient Cent Res Rev. 2016;3:99-110.)

Keywords

mycoses; fungi; histoplasmosis; blastomycosis; sporotrichosis; Aspergillus

A number of potential fungal pathogens are found in residential homes and yards. These microscopic fungi may result in a variety of human disease. This includes potentially fatal systemic fungal infections, traumatic wound infections, allergic manifestations, exacerbations of asthma, and sometimes vague, troubling, nonspecific signs and symptoms. Fortunately, these instances are relatively uncommon; however, domestically acquired fungal-related disease in humans may be seen more frequently as the proportion of immunocompromised patients living at home increases. Except for dermatophytes, most such fungal infections were apparently uncommon until the use of immunosuppressive agents and drugs in the second half of the 20th century. No doubt, significant improvements in the clinical and laboratory diagnosis of fungal infections in the last several decades has increased recognition of these diseases.

Allergic, toxic and irritant diseases from inhaled mold spores have been known since antiquity. The purpose of this narrative review is to describe disease that may result from fungal infection or exposure in one’s home or yard, and to increase clinician awareness that such illnesses may occur without significant occupational, recreational or exotic outdoor exposure. Included are potentially pathogenic fungi found in microenvironments immediately outside the home as well as on indoor or outdoor household surfaces, or that are associated with large appliances or other home furnishings. This article will not include dermatophyte or fungal infections acquired through contaminated or colonized food substances or personal care items, nor will it provide diagnostic or treatment detail for the diseases discussed. Rather, it is an overview of the genesis and clinical manifestations of the more common (e.g. blastomycosis) or emerging (e.g. black yeasts) types of fungal-related diseases and recent societal concerns (e.g. sick building syndrome) one might encounter at home in the Midwestern United States.

Home and Yard Microbial Ecology

Residential Soil and Building Materials: Yards contain some combination of plants, rocks/concrete/stone, decaying small animal carcasses and waste, processed
natural materials such as decorative bark, fencing and, the principal component, soil. Soil is a mixture of mineral and organic (humus) constituents. Soil-particle surfaces, pore spaces and nutrient-rich zones such as the rhizosphere (soil surrounding a plant root) form particular habitat for microorganisms, often in biofilms. The quantity and types of fungi in a specific soil microenvironment are determined by a complex interaction of physical attributes (nutrients, moisture, sunlight, temperature, pH level and redox potential) and competing organisms (other fungi, bacteria, archaea, viruses and larger organisms such as amoebae or worms) (Figure 1).

Factors contributing to the sometimes extreme environment of residential soil include applied herbicides, pesticides and other chemicals, temperature differentials due to the nearby heated dwelling and radiation of heat from sunlit patios, tannins from new decorative bark, etc. Concrete or stone are even more extreme environments for microorganisms. The ability of fungal pathogens to grow in such diverse environmental stress may explain their ability to “opportunistically” inhabit a plethora of challenging niches, including the human body. Pathogenic fungi may be indigenous to the native soil or enter the yard through animal deposits, manure or purchased materials.

Home exterior and interior building materials, surfaces and appliances can be even more extreme microenvironments for fungi than the surrounding yard. Kitchens, bathrooms, metals, hard polymers, silicones, antiseptics and cleaning products cause oxidative and other stress on microbial cells. Additionally, indoor aromatic pollutants may be metabolized by certain black yeasts. As with residential soil, successful indoor colonization by a fungus is determined by its ability to tolerate the particular physiochemical properties of the building materials or surfaces and derive sufficient nutrients from them. Despite the significant differences in appearance, indoor surface habitats are often similar to extreme natural outdoor environments. The high-moisture, high-temperature, detergent-laden, alkaline environment of a household dishwasher is one example. The ability to develop so-called “extremotolerance” may favor genetically and physiologically adaptable “generalist” fungi such as black yeast and Aspergillus spp. (both perhaps ancestrally rock surface inhabitants) as well as Blastomyces spp.

**Figure 1.** Microbial soil ecology. Pictorial representation of factors determining distribution of bacterial and fungal microorganisms in soil. Side panels represent climatic, physical and geobiochemical factors. Central panel includes a magnification (upper circle) representing soil particles, pore spaces and bacteria in biofilm, and another (lower circle) representing the rhizosphere and associated bacteria and fungi (and amoebae and nematodes). (Reproduced from Baumgardner DJ. Soil-related bacterial and fungal infections. J Am Board Fam Med. 2012;25:734-44, by permission of the American Board of Family Medicine.)
Climate Conducive to Fungi: Most environmental fungi cannot grow at human body temperatures. An in vitro study of 4,802 fungal strains representing 144 genera revealed that each 1°C temperature increase between 30° and 40° restricted the growth of approximately 6% additional fungal strains. In addition, our complex, redundant immune system inhibits a majority of potentially pathogenic fungal species. Again, the ability of a particular organism to survive our warm-blooded, slightly alkaline internal environment may be the result of harsh selective factors in domestic outdoor and indoor environments including thermal, physical, chemical, oxidative and competitive stresses. In addition to thermal tolerance and biofilms (common among black yeasts and Candida spp.), virulence factors for human invasion may have originated to counterattack bacteria, amoeba, nematodes and other organisms in the natural environment. As Gostinčar and colleagues suggested, “We have made our indoor (surface) environments prohibitive for the growth of the majority of microbes. However, the most resistant ones (and as such possibly the most dangerous) have succeeded in surviving and adapting.”

Mold growth and mold problems are common in homes. Visible mold may be identified in rooms whether a problem is suspected or not. It is frequently found on windowsills, refrigerator seals, spaces underneath kitchen sinks and air vents. Establishing normal levels for indoor fungal load has been difficult and controversial. A North Carolina study suggested that heating, ventilation and air conditioning (HVAC) systems and associated ductwork may transmit mold spores from the crawl space to the home interior. A recent university housing study from California examined fungi on residential surfaces including kitchens and bathroom drains, windowsills and the skin of human inhabitants. This housing had no reported mold problems and did not allow pets. The surfaces yielded a fungal profile similar to that expected in the outdoor air of that locale, and differed by surface type. Fungi typical of indoor air, such as Cladosporium and Cryptococcus, were found on windowsills. Drains yielded thermotolerant fungi including Candida, Exophiala and Fusarium. Human forehead skin contained a variety of indoor- and outdoor-associated fungi. In a separate study, fungi, especially Aspergillus, were identified on home pillows.

Other Environmental Factors: Pathogenic fungi may be introduced into the respiratory tract via bioaerosols (e.g. wind- or fan-borne spores, dusts and airborne soil or construction particles). Concentrations of fungal spores (up to and exceeding 100,000 spores/m³) vary widely by geographic factors like temperature, moisture, wind, photoperiod and time of day. Common outdoor spores include those from Cladosporium, Alternaria, Penicillium, Aspergillus and others. Indoor spore levels are generally 40–80% of outdoor levels and reflective of outdoor species. Humidity, inadequate ventilation and increased ambient temperature can increase the fungal burden of a home. A number of indoor sources, such as plants, pets and mold-transporting footwear and clothing, also may contribute mold. Fungal pathogens can enter humans by traumatic inoculation into wounds. Simultaneously introduced dirt-related minerals may promote infection by suppressing local host defenses. Much less is known about the contribution to human disease of fungal toxins and volatile organic compounds released by fungi.

On occasion, fungal infections may be acquired in one’s home or yard from a source external to one’s property, as in the case of an earthquake or dust storm. In the Midwest, both histoplasmosis and blastomycosis apparently have been acquired at home via airborne spores from a more remote source. Cases of mucormycosis caused by Apophysomyces trapeziformis were reported following a 2011 tornado in Joplin, Missouri.

Fungal Infections Primarily Acquired Outdoors

Histoplasmosis: The thermally dimorphic systemic fungus, Histoplasma capsulatum, is endemic to the Midwest, where a number of histoplasmosis outbreaks have occurred. Evidence of exposure is found in a majority of residents in this region. Histoplasmosis, a primary pulmonary disease, is often asymptomatic. The intensity of the acute pneumonia likely correlates with the burden of inhaled conidia. Pulmonary or mediastinal masses, nodules, granulomas, fibrosis or pericarditis may result. Arthralgias or erythema nodosum may be seen. Acute heavy-inoculum pneumonia may be severe and require mechanical ventilation. A bilateral, reticulonodular pattern may be seen on chest X-ray or computed tomography. The fungus may disseminate in any pulmonary infection to a variety of organ systems.
(liver, spleen, bone, lymph nodes, central nervous system, etc.). Patients with underlying lung disease may experience chronic, progressive (cavitary) pulmonary histoplasmosis.

*H. capsulatum* can cause opportunistic infections. Histoplasmosis-associated eye disease, a choroiditis often associated with serotype HLA-B7, is relatively unique to the Midwest. Patients are often asymptomatic unless scarring involves the optic nerve or fovea. A precise etiologic link to *H. capsulatum* as the causative agent is still lacking. Pediatric disease is generally asymptomatic or mild (fever, cough, chest pain and malaise) and limited to 2 weeks in duration, although fatigue may linger. Localized skin disease following direct inoculation is rare.

Growth of *H. capsulatum* is associated with 67–87% relative humidity, 90–127 cm of annual precipitation and mean soil temperatures of 22–29°C. The latter may explain the relative paucity of infections in the most upper portions of the Midwest. *H. capsulatum* lives in soil that is generally moist, acidic and high in nitrogen, carbohydrate and cationic salt content, particularly (but not exclusively) in association with bat or bird guano. The ability of *Histoplasma* to assimilate uric acid and related compounds may explain the latter association. Conidia (spores) are inhaled following nearby soil disturbance as a result of construction or excavation or around potted plants. The problem of birds nesting in large numbers on, in or near buildings is a particular problem. The large amounts of guano and nesting materials may remain for years and pose a health hazard during removal or even fairly minimal disturbance or dust formation.

While *H. capsulatum* spores inhaled from one’s yard are often generated by a more remote source (often part of a large urban outbreak), there have been specific instances of exposure from sources on the property. Examples include infection of homeowners and tenants during demolition of the exterior walls of an old brick house that included bird or bat guano, infection of visiting relatives at a family reunion in Iowa exposed to an old barn, and 10 sporadic cases from the same duplex housing complex on an Air Force base in central Illinois. In the latter cases, several soil sites about the complex were culture-positive for *H. capsulatum*. In the former two examples, environmental testing was apparently not attempted.

**Blastomycosis:** Blastomycosis is a potentially life-threatening systemic and cutaneous fungal infection. Pulmonary disease is the most common manifestation, and it has a broad differential diagnosis. The pneumonia may be asymptomatic, chronic or subacute, or sometimes rapidly progressive with adult respiratory distress syndrome and death. Acute or chronic dissemination of the fungus to skin, brain, genitourinary system, bones or virtually any organ system may result.

Blastomycosis is caused by *Blastomyces dermatitidis* or the newly discovered cryptic species *Blastomyces gilchristii*, both are dimorphic fungi endemic to the Midwest. Annual incidence ranges from <1–100 per 100,000 persons. *Blastomyces* occurs in and near soil microenvironments. Soil isolations and sites of likely exposure, including home sites, are often associated with sand soils, nearby waterways and animal excreta. It also appears to be a survivor of harsh or changing environmental conditions and may utilize a wide variety of substrates. It is proposed that the ability to grow and sporulate in carbon-poor, high-ammonia environments may be a key to its competitive success. Preceding precipitation and environmental temperatures may impact the occurrence of *Blastomyces*. No animal reservoir has been identified. Except for rare inoculation disease (usually localized to the skin), fungal spores (conidia) enter the body upon inhalation from the environment into the lungs. Sources of conidia include wind- or excavation-generated dusts, disturbance of soil or other microenvironments and likely moisture-induced natural spore release from the local environment.

While we feel there is good evidence that blastomycosis may be acquired on one’s own property, and others have concurred, some have disputed this hypothesis. Evidence includes case studies and repeated cases of blastomycosis at the same home site with different families, several studies that failed to implicate specific outdoor or occupational activities with acquisition of blastomycosis in most cases, and the isolation of *B. dermatitidis* from an abandoned house and 20 m from a human home in northern Wisconsin. Evidence, therefore, has been largely circumstantial. In-home cultures have been otherwise negative, but
B. dermatitidis is exceedingly difficult to culture from the environment.47,57

Sporotrichosis: Sporotrichosis is an uncommon subacute or chronic mycosis. In normal hosts, it is primarily a local lymphocutaneous infection resulting from direct environmental inoculation (thorns, splinters, cuts, etc.). Days or weeks later, a papule forms at the site of inoculation, becomes nodular and may ulcerate. Further nodules typically appear along the lymphatic distribution of that body site. This ascending lymphangitis picture also may be caused by nocardiosis, atypical mycobacteria, leishmaniasis and cutaneous anthrax.63 Osteoarticular disease may occur by direct inoculation or disease extension. Extensive cutaneous and systemic dissemination often occurs in untreated patients with underlying disease (alcoholism, acquired immune deficiency syndrome, diabetes or chronic obstructive pulmonary disease). Subacute or chronic pulmonary disease may follow, usually in the immunocompromised, through dissemination from the lymphatics or direct inhalation of conidia.63

The etiologic dimorphic fungus Sporothrix schenckii (and related strains) is found worldwide in soil, vegetation, sphagnum moss, potting soil, decaying wood, cornstalks, hay and several warm-blooded animals. Exposure to infected cats also may result in disease through scratches, bites or contact with a lesion on the cat.63-65 Unlike human lesions where the number of S. schenckii organisms may be low, cat lesions may have a very high fungal burden that allows transmission through lesion contact or splash without penetrating injury.63 Sporotrichosis may be acquired in the yard or home. A large multistate outbreak took place in 1988 related to sphagnum moss harvested in Wisconsin.66 A number of victims were patients who had received seedlings for their home sites. S. schenckii was cultured from five samples of unopened moss bales from one implicated nursery. Another outbreak was epidemiologically associated with bales of hay from a Halloween haunted house (there was no residual hay to culture, and cultures from the implicated fields 5 months after exposure were negative for S. schenckii).67 Such bales are often purchased for yard decorations. The author is personally acquainted with a poorly controlled diabetic who died of disseminated sporotrichosis acquired via a thorn puncture from her home garden in Illinois. In our own health system’s patient database (representing several hundred thousand individuals), there were 15 cases of sporotrichosis diagnosed on clinical grounds from 2000 to 2009. All but one had upper extremity disease. Of these 15 suspected cases, 5 had confirmed thorn punctures and 3 had exposure to other outdoor activity risks.68

Other Mycoses: A variety of nondimorphic, soil-related yeasts and molds found in residential areas can cause significant local or systemic human infection, particularly in the immunocompromised.63,69 Scedosporium apiospermum is an illustrative example of such a fungus.70 S. apiospermum may cause local, usually deep, trauma-related infections in normal hosts, colonization or infection of the respiratory tract in persons with predisposing conditions, systemic invasive disease in immunocompromised hosts and pneumonia or brain abscess in near-drowning or trauma victims.69 Strains of this fungus and the closely related Pseudallescheria boydii are thermotolerant and have been associated with agricultural, garden, potting or feces-enriched soil, sewers, polluted fresh water and salt-water habitats.70-73 S. apiospermum has been isolated preferentially from urban parks, playgrounds and other developed areas compared to natural areas,74 including from old decorative bark beneath a home dryer vent in suburban Milwaukee County, Wisconsin.75 Other examples include Rhizopus and Mucor spp. (sinus, pulmonary, gastrointestinal and wound infections). Transmissions of these fungi typically occur via sporagiospores in outdoor or indoor air that is inhaled or alighted on open wounds. Victims are usually diabetics or immunocompromised.76,77

Fungal Disorders Primarily Acquired Indoors

The problem of indoor household mold has been an area of active research and controversy the past three decades. Some of this has been fueled by lawsuits and media reports on “toxic mold” and “sick building syndrome.” It is difficult to prove cause and effect for these health conditions. Summaries of initial literature on this subject suggest some of these concerns were founded and others not. There is no general dose-response relationship between fungal concentrations and specific human disease.17,21,24 Furthermore, fungal species detection and enumeration is dependent on the type of sampling and
analytic technique used (quantitative polymerase chain reaction may be the most useful to identify associations between particular fungi and disease).\textsuperscript{78}

**Respiratory Tract Illnesses:** Relatively noncontroversial diseases caused by indoor molds include allergic bronchopulmonary aspergillosis (ABPA), allergic sinusitis, and hypersensitivity pneumonitis. Clinical allergic disease due to molds occurs in about 5\% of the total population, and these fungal allergies usually occur in patients with a history of atopy (about 40\% of the U.S. population). The most common allergenic indoor molds are *Penicillium* and *Aspergillus*. *Cladosporium* and *Alternaria* are predominately outdoor molds commonly found in indoor air and may provoke allergy.\textsuperscript{17,21,22} Allergic symptoms are often more prevalent in people living in damp residences.\textsuperscript{17} Humidity and warmer indoor air promote the growth of both mold (Figure 2) and house dust mites (a common allergen). Molds also serve as food for *Dermatophagoides* dust mites, further complicating the assignment of allergic etiologic factors.\textsuperscript{21}

The role of the fungal cell wall components, chitin and β-glucans in allergy is unclear. Human studies are limited.\textsuperscript{17,79} Smaller chitin fragments appear to be protective of allergic responses in animal models, however, some studies in human cells and nasal passages are contradictory. β-glucans have been shown to be both proinflammatory and anti-inflammatory.\textsuperscript{80}

Fungal hypersensitivity pneumonitis results from an exaggerated normal immunoglobulin G immune response to inhaled fungal proteins in conjunction with inhalation of a very large quantity of such fungal proteins. The interaction between the inhaled proteins and the fungus-directed, cell-mediated antibody reaction leads to an intense local immune response manifested as chronic pneumonitis of the peripheral airways and interstitium. Hypersensitivity pneumonitis has been linked to humidifiers and HVAC systems (most cases are occupational in origin). Humidifier exposures are more often due to filamentous bacteria rather than fungi.\textsuperscript{21,22}

Previous literature has associated indoor fungal exposure with more respiratory tract illnesses in infants and children\textsuperscript{17} and with increased asthma occurrence and severity (as well as subsequent improvement after household mold remediation).\textsuperscript{81} More recent studies applying the environmental relative moldiness index (ERMI) to homes have shown an association between higher ERMI score and severe asthma versus no asthma (but not versus moderate asthma) in household children in Detroit,\textsuperscript{82} and between the ERMI and later respiratory illnesses in atopic infants born in the Cincinnati area.\textsuperscript{83} The summation of levels of three molds (two *Aspergillus* species and one *Penicillium*) was significantly associated with the development of asthma at age 7 years in infants from this same area who were studied, along with their homes, beginning at age 1 year.\textsuperscript{84} This same authorship group subsequently noted that the home characteristics of air conditioning (negatively), carpeting (negatively), age of home (positively) and dust mite allergen (positively) were associated with the ERMI. Several variables were confounded, and multivariate analysis suggested that lower overall socioeconomic status was associated with higher ERMI.\textsuperscript{85} Similar associations of certain indoor molds with asthma and allergic symptoms (particularly when mold odor was present) were found in single studies and meta-analyses from other Midwestern and worldwide locations.\textsuperscript{85-92}

**Toxic and Irritant Effects:** Controversial concerns regarding indoor molds are especially focused around toxic effects. The major concerns are: 1) mycotoxins,

![Figure 2. Unidentified mold growing on a household windowsill in Milwaukee County, Wisconsin, in the month of February. This is a common occurrence in winter in the upper Midwest due to condensation at the window interface between warmer, moist indoor air and very cold outdoor air.](image-url)
which are relatively large, generally nonvolatile molecules (i.e. require inhalation of aerosols of fungal fragments, spores or large molecules) that are not required for fungus growth or survival but often assist in microbial competition; and 2) volatile organic compounds (VOCs) released by molds. The latter is responsible for the characteristic moldy odor. In the United States, acute mold poisonings are uncommon and mostly result from eating mold-contaminated food or by acute inhalations of high concentrations of mold particles or mycotoxins as part of agricultural or other occupational exposure. Indoor residential mold toxicity appears to be much less common and poorly documented.\textsuperscript{17,21-23} Mycotoxins may persist long after fungal particles are nonviable and may only become a problem when dusts are formed. In addition, there are numerous toxins and a poor understanding of their genetic control and potential effects. Thus, it is difficult to relate most toxins to specific symptoms.\textsuperscript{22}

There has been particular interest in the trichothecene mycotoxins of \textit{Stachybotrys chartarum} and its related species since a mid-1990s outbreak of acute pulmonary hemorrhage in Cleveland, Ohio, infants was initially associated with this fungus.\textsuperscript{21,93} While \textit{Stachybotrys} spp. have been isolated from other substrates, including gypsum (drywall), fiberglass wallpaper and aluminum foil, it has an affinity for water-damaged cellulose. However, this latter substance also promotes the growth of \textit{Cladosporium, Penicillium} and \textit{Aspergillus}. In nutrient-poor, damp-cellulose environments, \textit{Stachybotrys} is a tertiary colonizer following the primary (\textit{Penicillium} and \textit{Aspergillus versicolor}) and secondary (\textit{Cladosporium}) fungal inhabitants. This fact alone makes association of \textit{Stachybotrys} with human disease difficult,\textsuperscript{24} and many other molds are capable of producing mycotoxins.\textsuperscript{21} The presence of toxigenic molds in a household does not prove the presence of mycotoxins or that the occupants were exposed to a toxic dose.\textsuperscript{22} Recently it has been proposed that molds from water-damaged buildings may reside internally only in biofilms, likely in the sinuses. These molds may continue to release and produce mycotoxins that contribute to ongoing illness.\textsuperscript{94}

Within 10 years of the Cleveland outbreak, considerable doubt had been placed on the likelihood of \textit{Stachybotrys} toxicity due to serious methodological flaws, difficulty assessing mycotoxin levels in homes and lack of other supporting evidence,\textsuperscript{21,22,24} although some investigators remain more favorable toward this epidemiologic association.\textsuperscript{93}

Fungal VOCs may cause direct surface irritation that manifests as a burning sensation of the skin, eyes and upper airways. These compounds also have been suggested to cause symptoms such as lethargy and headache, but these effects have not been confirmed by controlled studies. It is more difficult to explain other symptoms such as dizziness, memory loss, gastrointestinal symptoms and muscle cramps. The contribution to these symptoms of potential psychological responses to VOCs notwithstanding, there is little scientific evidence for VOC causality. As with many fungal diseases, individual differences in susceptibility may lead to discordant symptom histories from equally exposed persons.\textsuperscript{17}

Sick building syndrome is a controversial concept defined as nonspecific symptoms of the eyes, skin and upper airways, headaches and fatigue of uncertain cause, all of which the patient associates with the indoor environment (though not necessarily fungi). It is more common among women and those with self-reported allergy or with certain personality traits such as somatization and neuroticism.\textsuperscript{79} There is substantial overlap of this concept and other controversial fungus-related concerns discussed earlier.

**Fungal Disease Acquired in all Residential Settings**

**Aspergillosis:** Conditions caused by \textit{Aspergillus} may occur from indoor or outdoor exposures, be acute or chronic and have allergic or toxin-mediated manifestations. Most infections occur in immunocompromised patients, usually due to \textit{Aspergillus fumigatus}, and include invasive pulmonary aspergillosis (cough, dyspnea with or without fever, chest pain, hemoptysis, wheezing — symptoms often confused with other entities), pulmonary or sinus fungus balls, chronic pulmonary aspergillosis, allergic fungal sinusitis, invasive sinusitis, posttraumatic keratitis, otomycosis (benign growth on external auditory canal debris), endocarditis and other local, superficial or disseminated forms.\textsuperscript{95,96} Which manifestation of \textit{Aspergillus}-related disease occurs, if any, following...
exposure depends upon the quantity and virulence of the encountered strain and the host response.\textsuperscript{97}

Forms of chronic pulmonary aspergillosis (e.g. simple aspergilloma, chronic cavitary pulmonary aspergillosis, chronic necrotizing pulmonary aspergillosis) are uncommon and nearly always occur in those with underlying chronic obstructive pulmonary disease, diabetes or other immunosuppressive states. Patients usually present with chronic relapsing cough, dyspnea and weight loss. Chest pain, fatigue, fever and acute hemoptysis also may occur.\textsuperscript{98}

Allergic bronchopulmonary aspergillosis is a pulmonary manifestation of allergy to \textit{A. fumigatus} in rural and urban asthmatics that may start in childhood and remain unrecognized for years (some advocate screening all asthmatics for ABPA). The process begins with sensitization in an asthmatic who remains in good control but may progress to steroid-dependent asthma, then to end-stage fibrotic pulmonary disease and “honeycomb lung.” Some patients experience a prolonged remission. \textit{Aspergillus} allergic disease also complicates cystic fibrosis, but laboratory and clinical features are distinct.\textsuperscript{96,97}

Most infections occur following inhalation of conidia. \textit{Aspergillus} is ubiquitous in air, and sources of spores include soil, decaying vegetation and dust. The relative importance of soil as a source of infection compared to plants, flowers, building materials, water and other environments is unclear.\textsuperscript{11,99} \textit{Aspergillus} fungi were commonly isolated from refrigerator surfaces and foods, pillows,\textsuperscript{20} laundered socks,\textsuperscript{100} composted wood chips and yard waste.\textsuperscript{96} Sputum isolations of \textit{A. fumigatis} have been associated with higher levels of the fungus in the air inside patient homes (although correlation with \textit{A. fumigates} sensitization was not demonstrated).\textsuperscript{101}

\textbf{Fusarium:} \textit{Fusarium} spp. may cause many of the nonallergic manifestations seen with \textit{Aspergillus}, frequently in immunocompromised persons. \textit{Fusarium} is found worldwide in soil, water, decaying organic debris and on plants and is spread by air or rain. Infection, and perhaps hypersensitivity pneumonitis, may be acquired at home.\textsuperscript{102,103}

\textbf{Black Fungi:} Phaeohyphomycosis is a term applied to clinical syndromes caused by fungi containing high levels of melanin, exclusive of eumycetoma and chromoblastomycosis. Diseases that characterize phaeohyphomycosis include allergic fungal sinusitis and allergic bronchopulmonary mycosis similar to that of \textit{Aspergillus}, superficial infections (often starting as a single inflammatory cyst on an extremity), deep local infections (often related to trauma or immunosuppression), disseminated infections (usually in the immunocompromised), pulmonary disease (usually in immunocompromised patients or those with underlying lung disease) and central nervous system infection (often in normal hosts). Melanin appears to be a virulence factor in complex, multifactorial and incompletely understood ways.\textsuperscript{104-106}

A variety of genera may be agents of phaeohyphomycosis. Three examples are given here. \textit{Alternaria} and \textit{Cladosporium} are ubiquitous fungi commonly found in outdoor and indoor air. Both may cause chronic allergic sinusitis in normal hosts. The former has caused ulcerated cutaneous infections, visceral infections, osteomyelitis, keratitis, onychomycosis and invasive sinusitis in immunocompromised individuals, especially those with bone marrow transplants. The latter rarely causes similar disease and pulmonary infections.\textsuperscript{104,105} \textit{Exophiala} (black yeasts) may cause a variety of deep and cutaneous infections, sinusitis and rare other entities in normal and immunocompromised hosts.\textsuperscript{104,105,107} Infections by these fungi may be underdiagnosed in the United States.\textsuperscript{107} They may be found in water, soil, plants and decaying woody material. Indoors, \textit{Exophiala} may be identified in dishwashers,\textsuperscript{13} sink drains, bathwater,\textsuperscript{12} and other bathroom and shower surfaces (along with \textit{Cladosporium}).\textsuperscript{108} Infection by \textit{Exophiala} occurs by traumatic inoculation and rarely by inhalation.\textsuperscript{105}

\textbf{Prevention}

A recent commentary from the Centers for Disease Control and Prevention pointed out that “fungal infections remain serious and underappreciated causes of illness and death.”\textsuperscript{109,109} Prevention of sequelae from these infections includes clinician and public awareness.

Immunocompromised patients should be especially warned regarding risks for fungal infection and ways to prevent fungal exposure. In the domestic environment, this would include maintaining proper personal skin
integrity and hand hygiene before eating or handling food, after going outdoors and after touching plants, pets or soil. Household measures include keeping water-exposed surfaces and areas of dust accumulation cleaned and disinfected where appropriate, avoiding standing water, using sterile water in nebulizers and humidifiers, and removing water-damaged materials. It is best to avoid home (or yard) renovation, attics, crawl spaces, excavation, caves and musty areas during immunosuppression. There is moderate evidence that remediation of moldy houses reduces asthma symptoms and respiratory infections, but controlled trials regarding allergic rhinitis have been few and nonconfirmatory. Mold spores are present in all outdoor and indoor environments and cannot be eliminated.

General measures for all persons to reduce the risk of domestically acquired fungal disease include maintaining relative humidity below 60% and limiting shower and pot-boiling times, the number of plants to be watered and the use of exhaust fans. Cold surfaces, such as windowsills in winter, promote mold growth. Visible mold in the home should be removed, with particular attention to the underlying cause of excessive moisture. In particular, water leaks should be remediated immediately, and proper HVAC system maintenance ensured. Health departments are often helpful in guiding home remediation. Commercial mold remediation companies may be used if qualified and ethical.

Summary
A variety of fungal microorganisms are capable of departing a soil environment and causing serious focal or systemic infection in humans. Specific evolved virulence factors for the ability to grow in diverse, sometimes harsh, domestic microenvironments may promote infection acquired on one’s personal property. Although many of the fungal diseases listed in this article are uncommon, they can occur in individuals largely restricted to a domestic environment in the Midwest. Delay in diagnosis from dismissing a disease such as blastomycosis or failing to consider a fungal etiology for a deep wound infection due to the apparent lack of traditional outdoor recreational or occupational risk factors may lead to patient morbidity or mortality. If a home- or yard-related fungal infection is discovered, source investigation may be an important public health measure. Similarly, in patients who present with self-described mold-related illnesses, an appropriate differential diagnosis should be followed and other causes of the particular complaints ruled out.

Patient-Friendly Recap
- Myriad fungal spores live in Midwestern homes and yards and can occasionally cause serious infections in residents.
- The author reviewed more than 100 sources to identify the most notable disease-causing domestic fungi (excluding contaminated food or toiletries) and related symptoms.
- Delay in diagnosis can be avoided if clinicians learn to spot the signs of these often challenging fungal infections.

Conflicts of Interest
None.

References
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