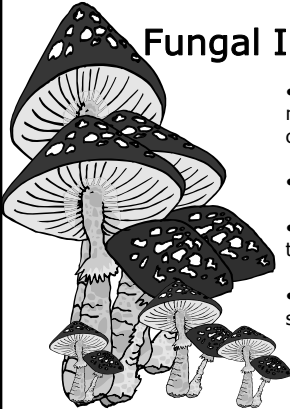
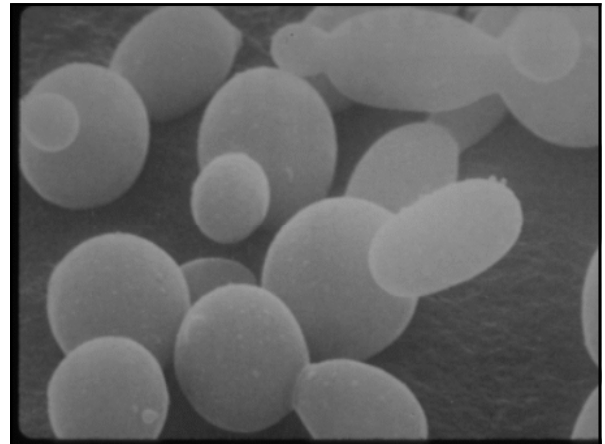


Fungal Infections

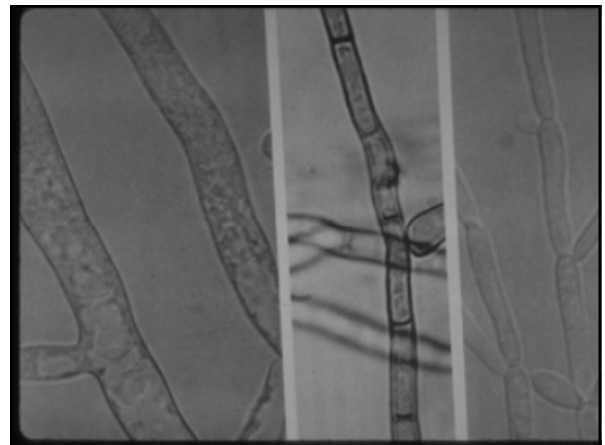


- Once exotic and rare; now increasingly common
- Fungi are not "virulent"
- But they are good at taking advantage
- "Opportunistic" in many senses



Fungal biology

- Eukaryotic (organized nucleus and cell structure)
- Non-motile
- Aerobic
- Saphrophytic or parasitic
- Cell wall contains glucan and chitin
- Cell membrane contains ergosterol



Fungal cell structure

- Yeasts (unicellular, budding)
- Molds (hyphae, mycelia, spores)
- Dimorphs (both)

Pathogenesis

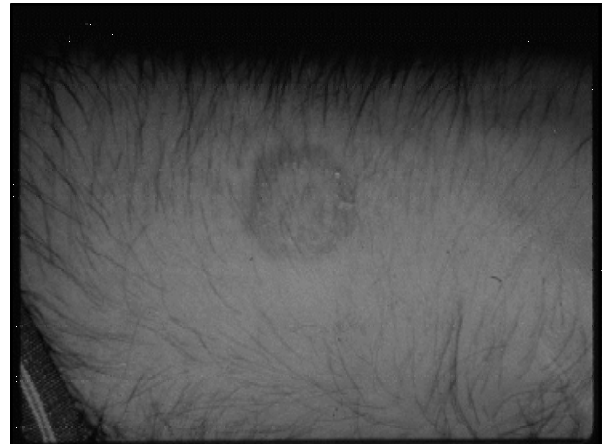
Toxins: produced, but not relevant to human infections

Disease from:

- Bulk of organisms
- Immune response to them or their byproducts

Overview of fungal infections

- Superficial or cutaneous (skin, hair, nails)
- Subcutaneous
- Systemic
 - “true pathogens” may cause disease in normal hosts although worse with immunocompromise
 - “opportunists” cause disease almost exclusively in immunocompromised hosts



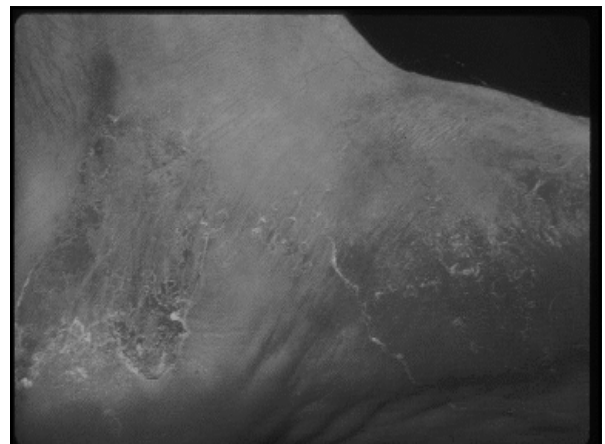
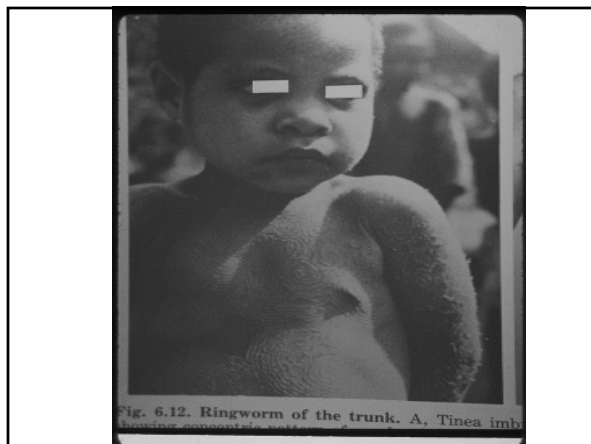
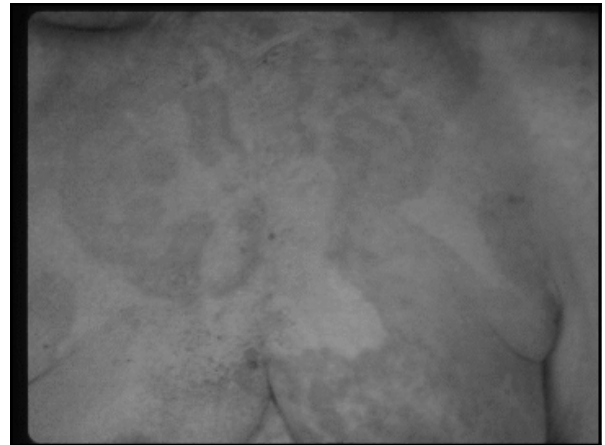
Superficial fungal infections

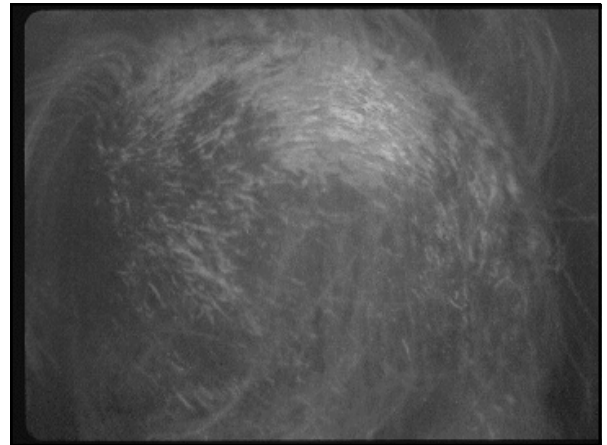
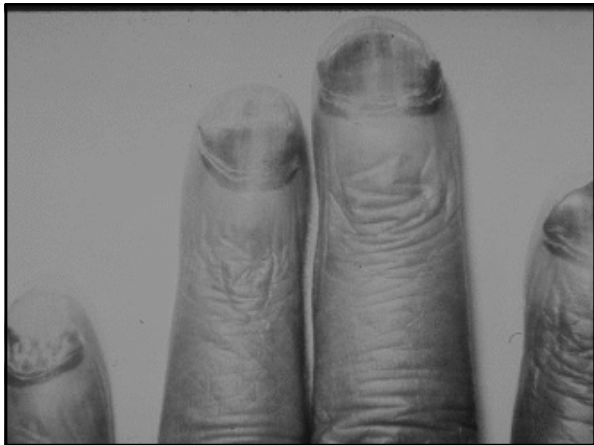
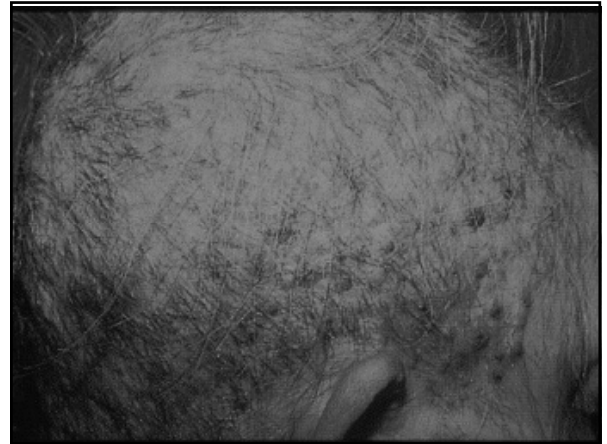
Dermatophytes: molds producing keratinase

Pathogenesis: grow as saprophytes on skin/nails; cause inflammation below

Clinical:

- Tinea corporis Tinea cruris
- Tinea pedis Tinea unguum
- Tinea capitis





Superficial fungal infections

Malassezia furfur: lipophilic yeast (derives nourishment from skin lipids)

Pathogenesis: lives on skin, causes pigment changes and itch underneath

Diseases:

- Tinea versicolor
- Occasionally fungemia with lipid infusion

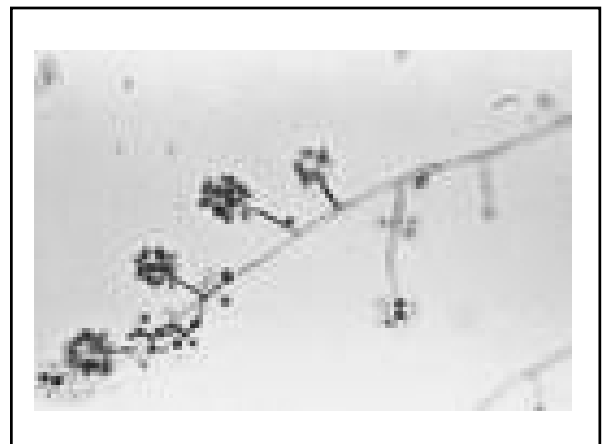


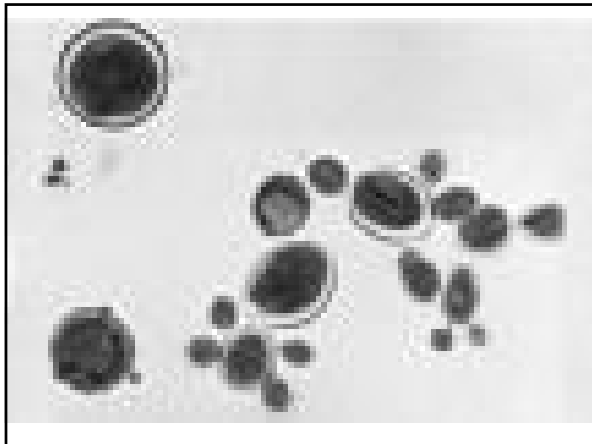
Subcutaneous fungal infection: Sporotrichosis

- Organism: *Sporothrix schenckii*
 - Dimorphic soil fungus (mold in environment, yeast in body)
- Habitat: soil, worldwide
- Pathogenesis: splinters or thorns inoculate organism into subcutaneous tissues

Subcutaneous fungal infections

- Pathogenesis: introduced through skin by foreign body, grow in subcutaneous tissues, spread via lymphatics
- Disease; usual local; may disseminate to adjacent bones, joints.
- Most common in nonindustrialized world (mycetoma of feet)





Sporotrichosis

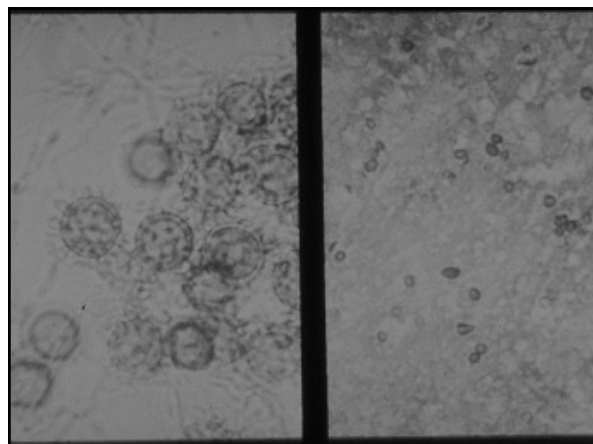
Pathophysiology:

- Spore inoculated by foreign body
- Yeasts travel along lymphatics
- Elicit mixed pyogenic-granulomatous reaction

Clinical:

- Gardeners and outdoorspersons
- Ulcerating nodules along hard cord
- Bone and joint destruction
- Dissemination rare

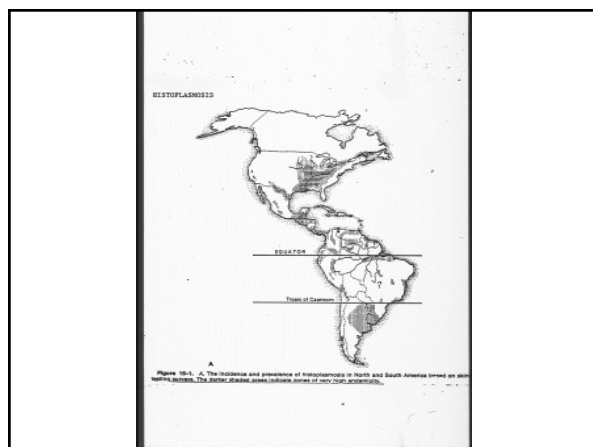




Systemic fungal infections: the “true pathogens”

Histoplasmosis, Coccidioidomycosis,
Blastomycosis

- Dimorphic
- Respiratory acquisition
- Restricted geographic distribution
- Infect normal hosts
- Disease reminiscent of TB



Histoplasmosis

Organism: *Histoplasma capsulatum*

- Soil dimorph (yeast in body, mold in environment)

Habitat: soils with high N content

- Ohio-Mississippi valley; Caribbean; Central and S. America
- Guano of bats, birds, poultry (chicken coops and caves)

Pathogenesis: inhalation of spores

Histoplasmosis

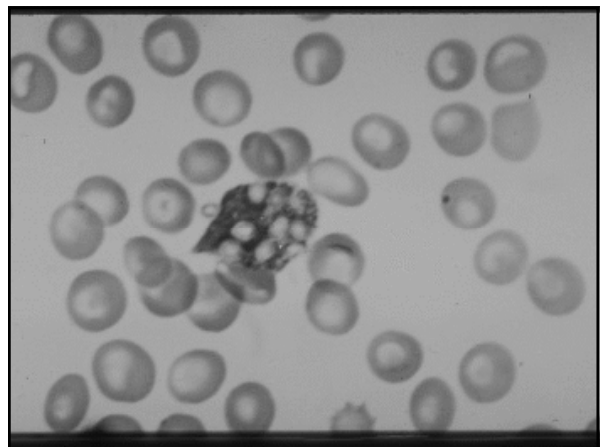
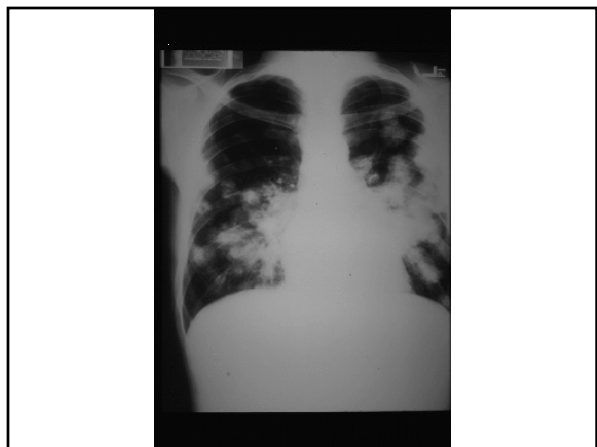
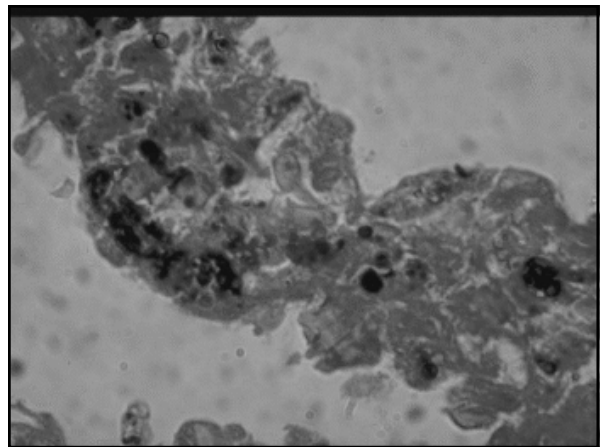
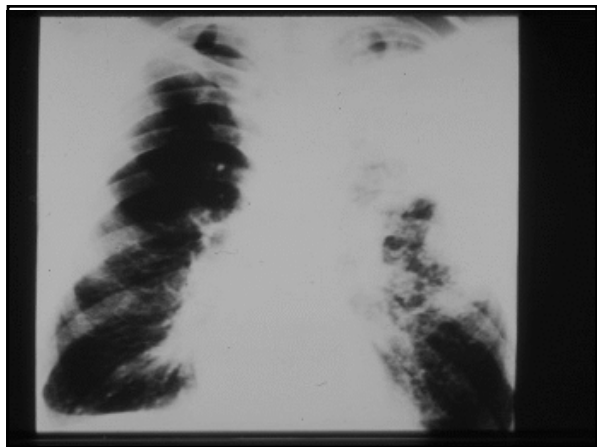
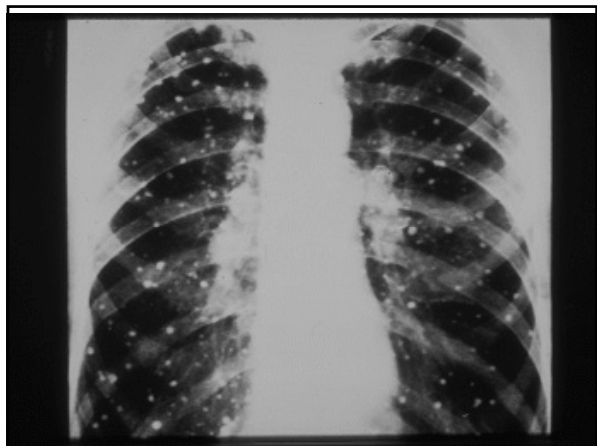
Pathophysiology:

- Mold spores transform into yeast in lung, elicit cellular immunity as per TB
- Hematogenous dissemination
- Skin test reactivity
- Walled off granulomata

Clinical:

Mimics TB. Usually latent disease, but

- may cause acute/chronic cavitary lung disease
- may disseminate after infection (infancy, immunocompromise)
- may reactivate years later



Coccidioidomycosis

Organism: *Coccidioides immitis*

- Dimorph: mold in soil, spherules and endospores in host

Habitat: lower Sonoral life zone (arid):
Southwest US, Mexico, Central and South America

Pathogenesis: inhalation of spores

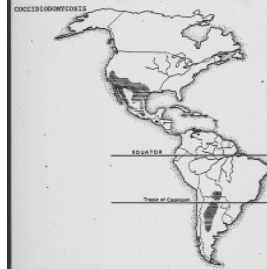
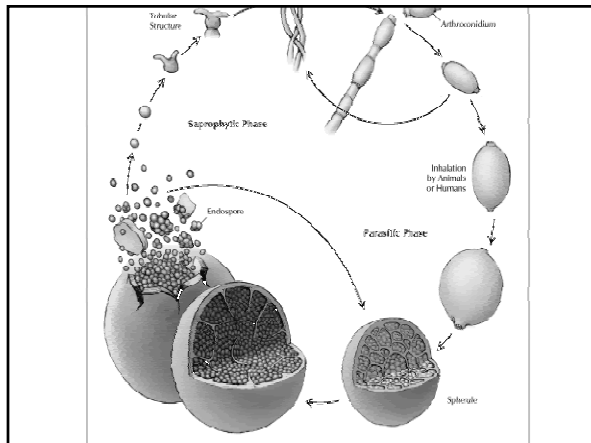


Figure 17-10 Incidence and prevalence of coccidioidomycosis. The areas marked by horizontal lines indicate incidence. The shaded areas indicate regions of high endemicity.



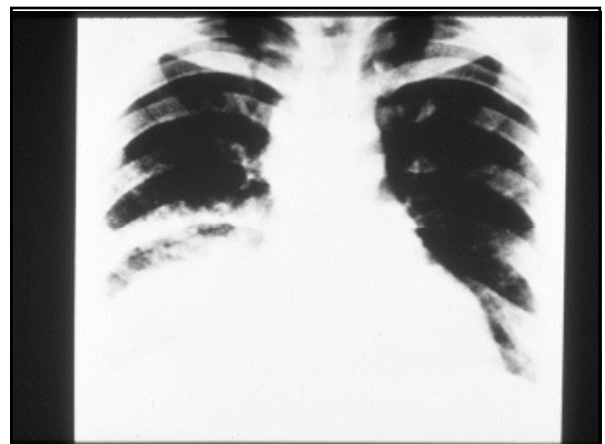
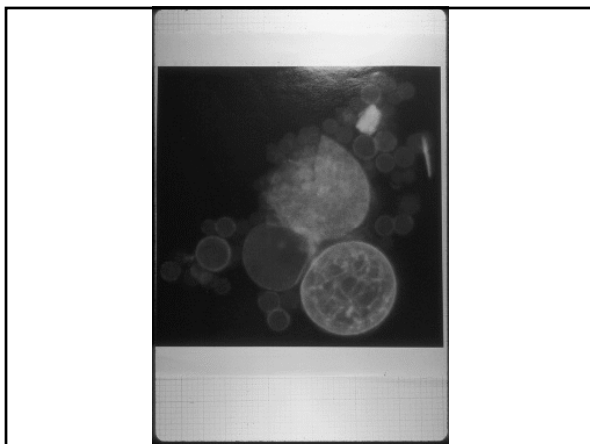
Cocci

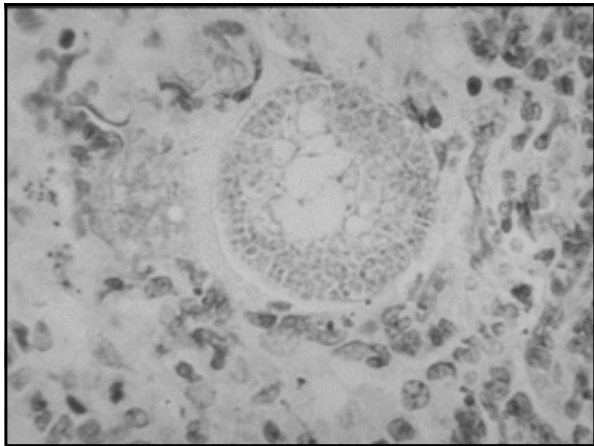
Pathophysiology:

- Spores transform into spherules in lung, elicit cellular immunity as per TB
- Hematogenous dissemination
- Skin test reactivity
- Walled off granulomas

Clinical:

- Acute self-limited flu-like seroconversion syndrome ("Valley fever")
- Acute or chronic lung disease
- Dissemination (pregnancy, dark skin, immunocompromise)
 - Skin
 - Bone
 - CNS





Blastomycosis

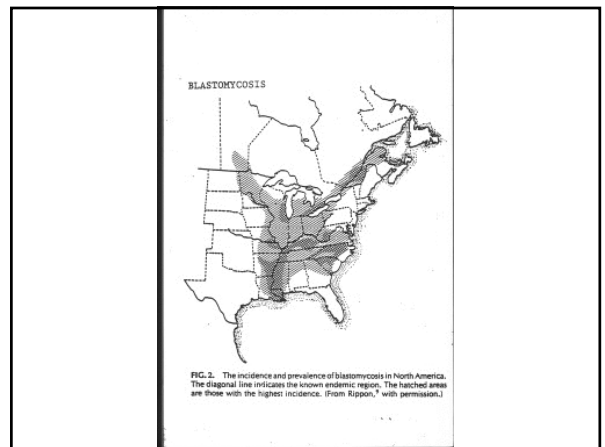
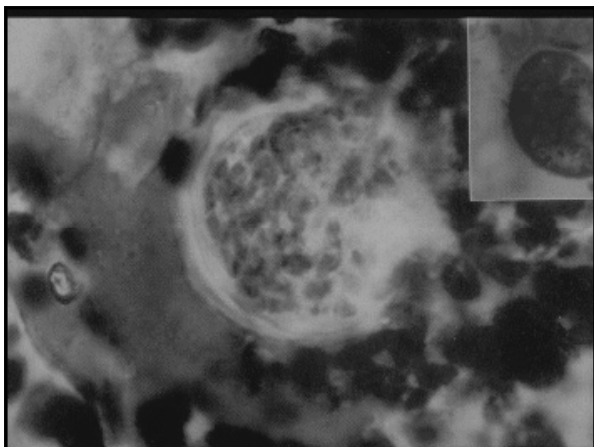
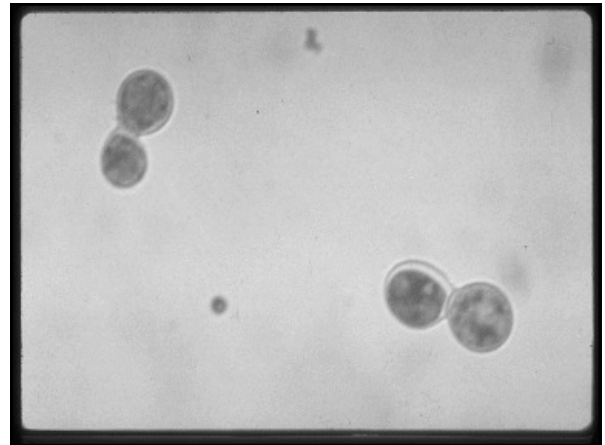
Organism: *Blastomyces dermatitidis*

dimorph: mold to yeast

Habitat: humid woodlands

- MidAtlantic zone
- Beaver dams, peanut farms
- Organic debris rather than soil

Pathogenesis: inhalation of spores



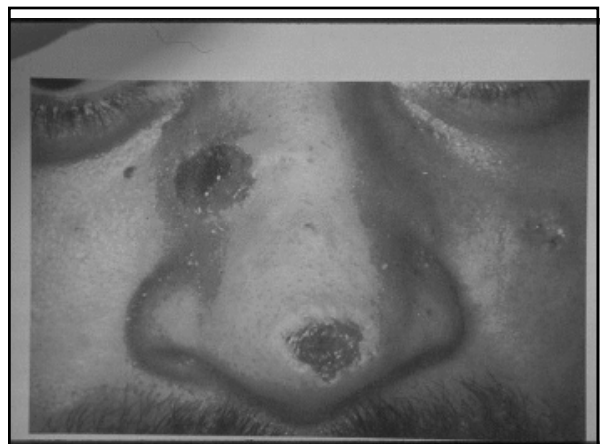
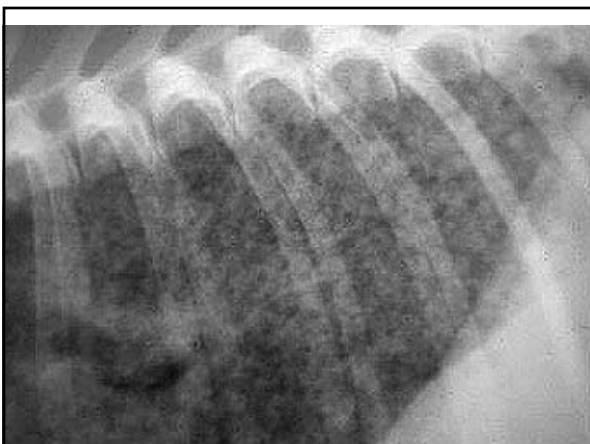
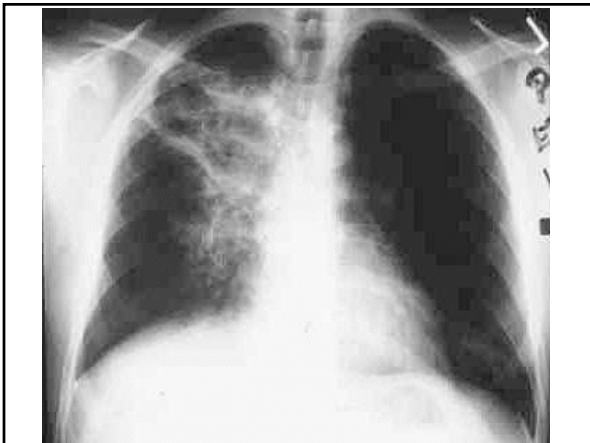
Blastomycosis

Pathophysiology:

- Spores transform into yeast in lung, disseminate
- No good antigen test to define exposed population

Clinical:

- Acute or chronic lung disease (nodular/cavitary)
- Disseminated disease:
 - Skin
 - Bone
 - Urinary tract in men





Systemic fungal infections:

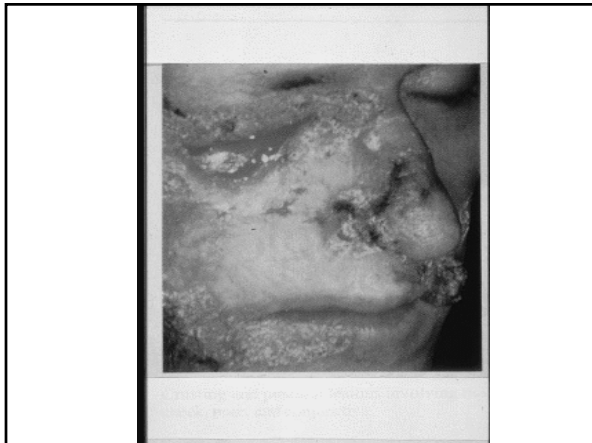
B. "the opportunists"

Histo, Blast, Cocci

- Geographic distribution
- Dimorphic
- Infection by inhalation
- Pyogenic/granulomatous host response
- Similar to TB
- Infection \approx immunity

Opportunists

- Omnipresent
- Yeasts or molds
- Various routes of infection
- Host response varies
- Clinical syndromes vary
- No lasting immunity

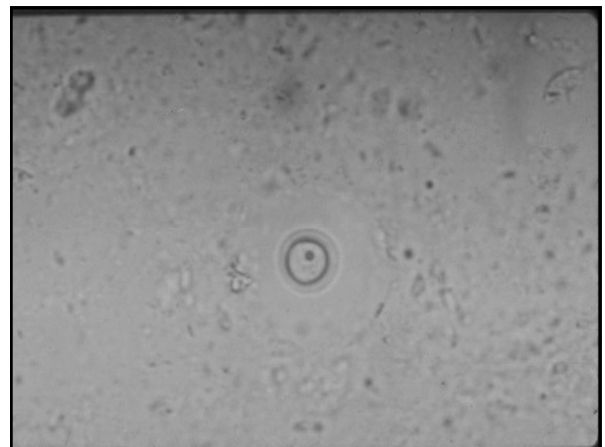


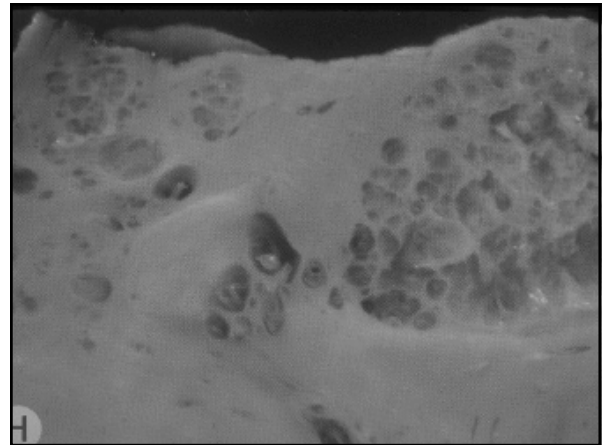
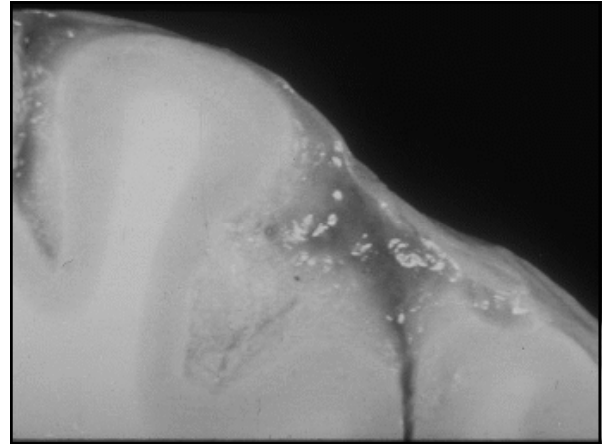
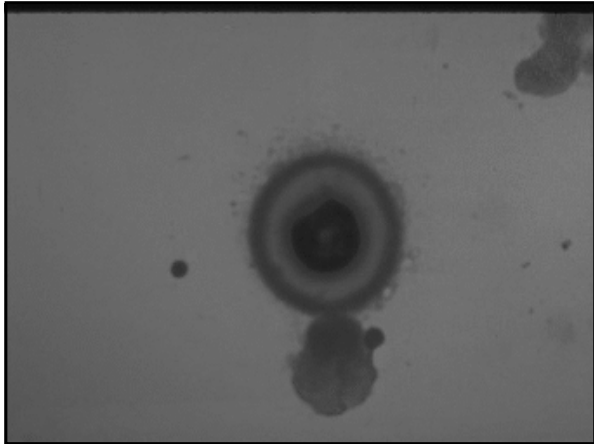
Cryptococcosis

Organism: *Cryptococcus neoformans*
yeast with a thick polysaccharide capsule

Habitat: bioterrorists (of a sort), worldwide

Pathogenesis: inhalation of yeasts





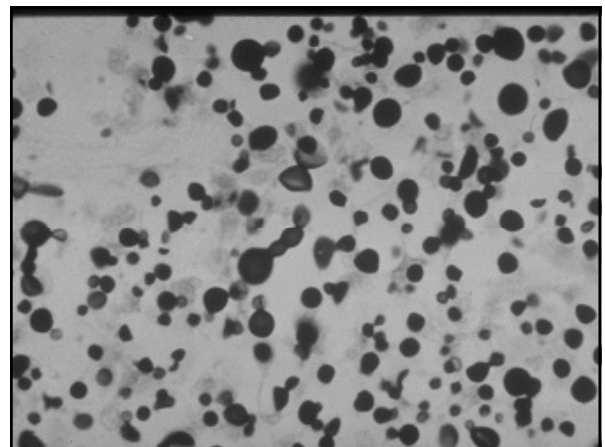
Cryptococcosis

Pathophysiology:

- Inhalation leads to
- Transient colonization OR
- Acute/chronic lung disease OR
- CNS invasion

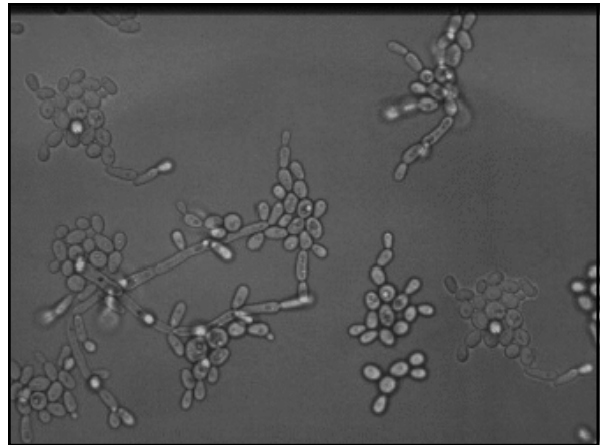
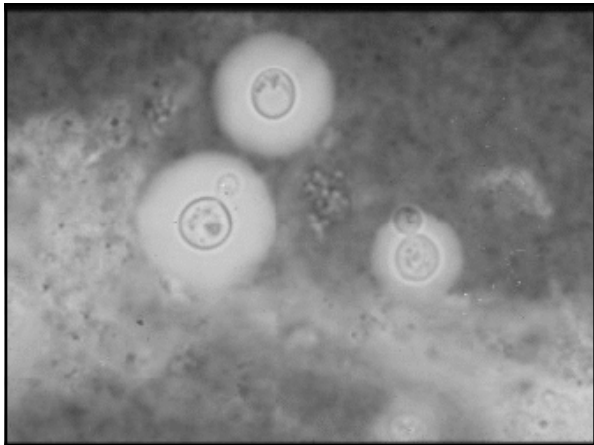
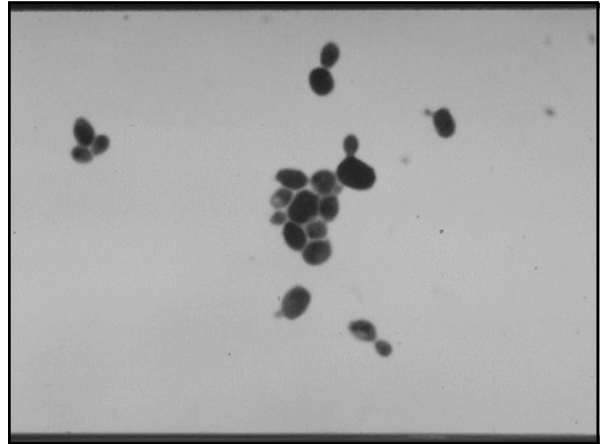
Clinical

- Pneumonia OR
- Meningoencephalitis
- Acute or chronic
- Fever, headache, stiff neck, fever, delirium
- Hydrocephalus



Cryptococcal meningitis

- India ink preparation of CSF may show organisms
- Serum or CSF antigen assay diagnostic in >95% cases of CNS disease



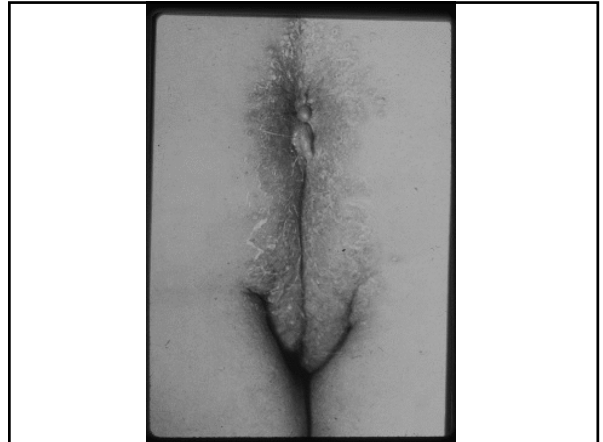
Candidiasis

- Organism: *Candida albicans* et al (yeasts with hyphal forms)
- Habitat: normal human flora
- Pathogenesis:
 - Colonized areas: change in environment leads to overgrowth
 - Noncolonized areas: change in immunity leads to invasion



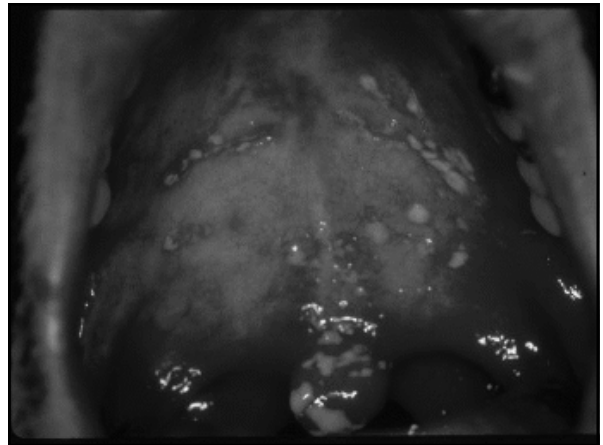
Pathogenesis of Candida infections

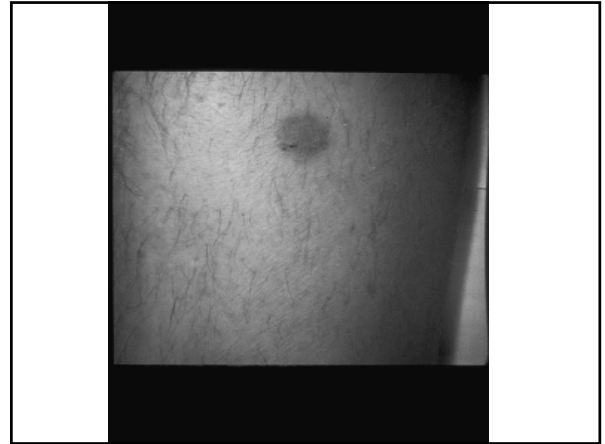
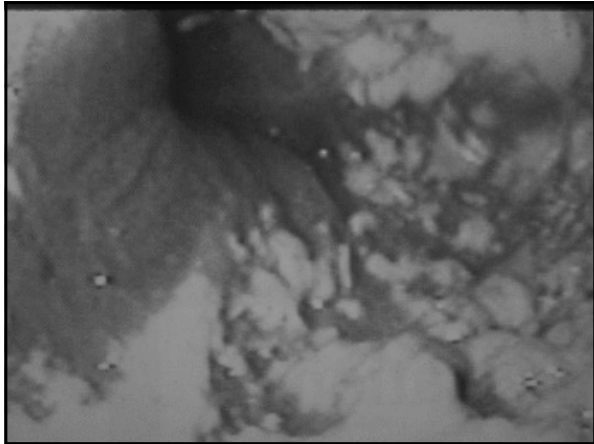
- Primary host defenses:
 - Intact skin
 - Intact mucosa with normal pH and normal flora
 - Functioning lymphocytes
 - Functioning neutrophils



Pathogenesis of local Candida infections

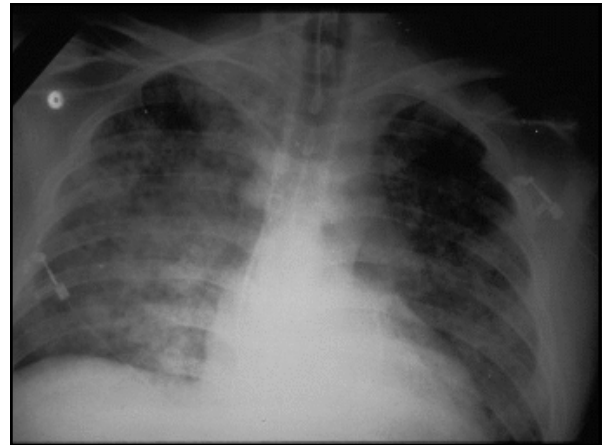
- Environmental changes
 - Wet skin
 - Changes in local flora
 - Hormones, foreign bodies
- Lymphocyte dysfunction
 - Immaturity
 - Destruction (HIV)





Pathogenesis of invasive Candida infections

- Breach in anatomic integrity (often biofilm on catheter)
- Defective PML function (first line of defense)
 - Myeloperoxidase, complement necessary but not sufficient defense
 - Cytokines also essential for recruiting phagocytes in disseminated disease
 - Antibody may or may not be present; may or may not be protective



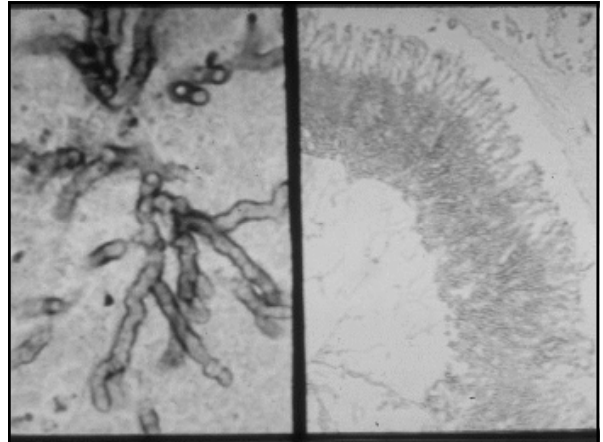
Invasive candidiasis

- Usually in critically ill patients with multiple risks (hospitalized, neutropenic, on antibiotics, many catheters)
- Fever, leukocytosis, organ dysfunction
- Microabscesses in kidney, liver, skin, eye, lung, heart
- Candida endocarditis

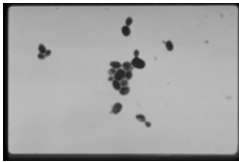


“Virulence” of Candida?

- Inherent “virulence”
 - environmental tolerance
 - Secrete hydrolases, beta proteases, phospholipases
 - Can adhere to plastic
 - Can invade GI, renal epithelium
- Additional hyphal virulence
 - Protects against phagocytosis
 - Knockout strains



Additional comments on candidiasis



- Gram stain may help identify
- Infection and colonization are difficult to distinguish
- Best treatment restores missing defense

Aspergillosis

Pathophysiology

Spores in lung may:

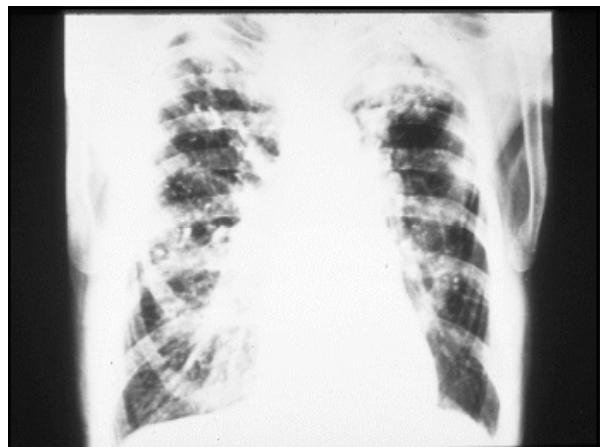
- Elicit allergy
- Grow in preexisting cavity
- Invade vasculature, disseminate with local and distant disease
- Neutrophils prime defenders

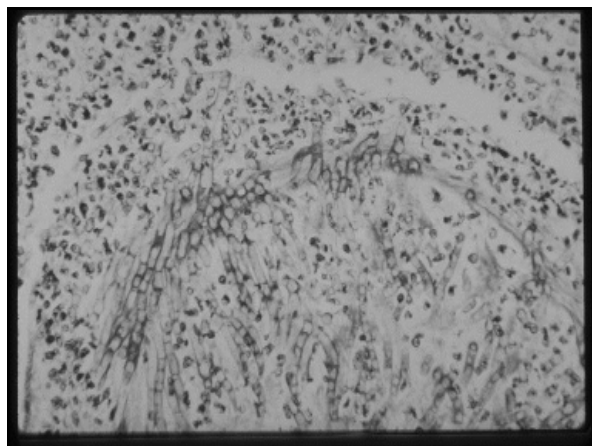
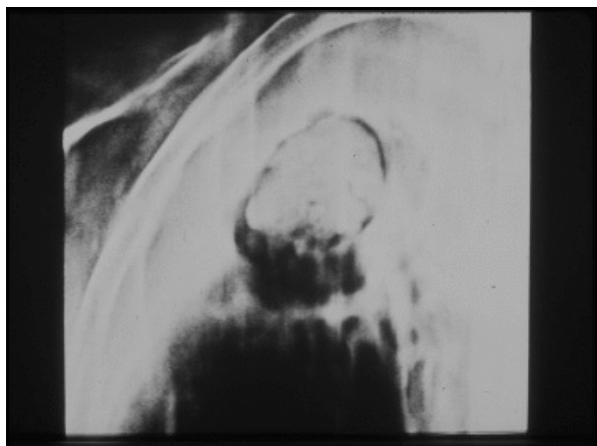
Clinical

- Allergic bronchopulmonary aspergillosis
- Aspergilloma
- Invasive aspergillosis with pneumonia, other end-organ disease

Aspergillosis

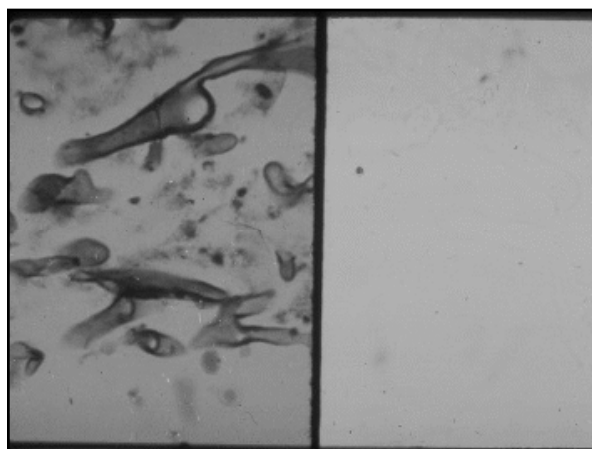
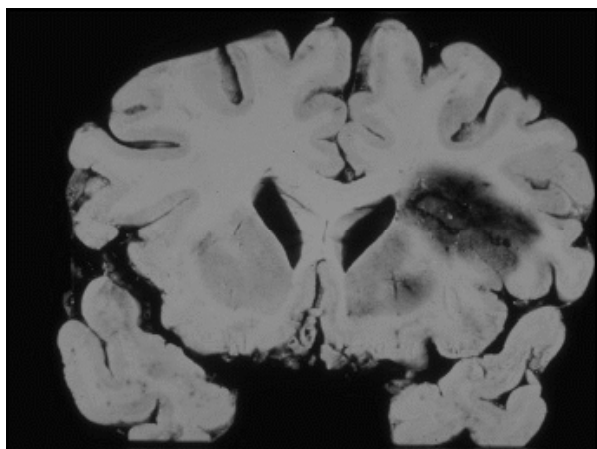
- Organism: *Aspergillus fumigatus* and others
 - Mold without a yeast phase
- Habitat: everywhere, worldwide
- Pathogenesis: inhalation of spores

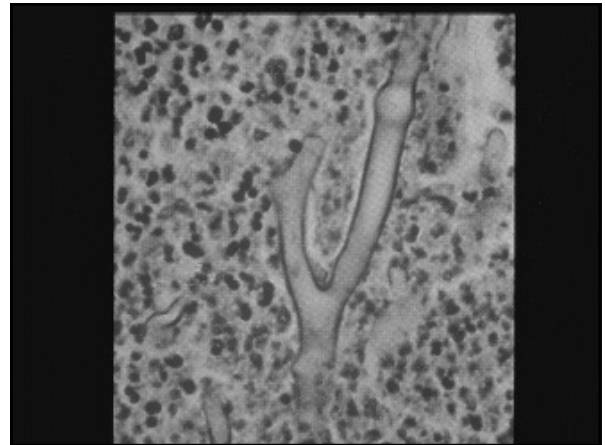
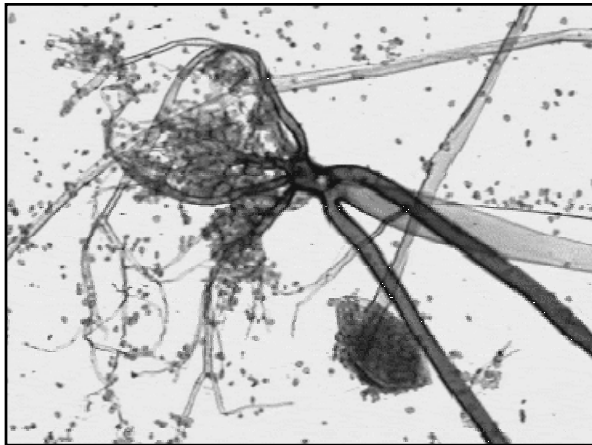
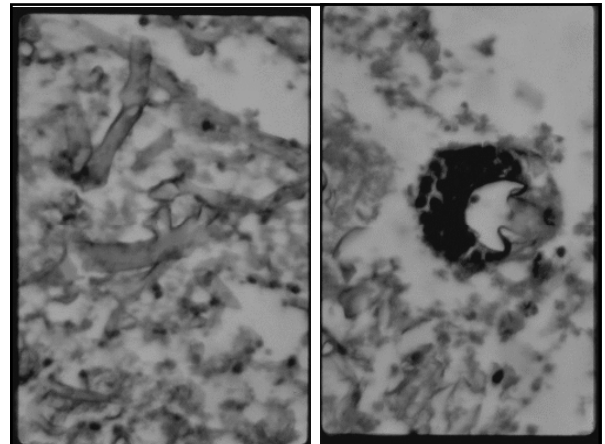
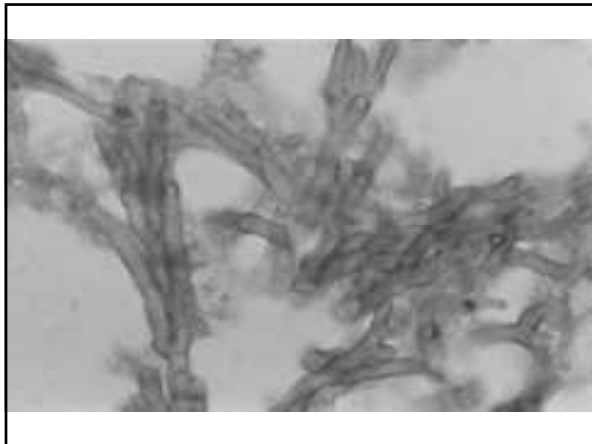




Mucormycosis

- Organism: species of Mucorales, genera *Rhizopus* and *Mucor* (Zygomycetes)
 - Molds without a yeast phase
- Habitat: everywhere, worldwide
- Pathogenesis: inhalation of spores





Mucormycosis

Pathophysiology:

Alveolar MPH/PML clear organisms, BUT:

- Metabolic acidosis
- Diabetes
- Neutrophil dysfunction
- Iron overload

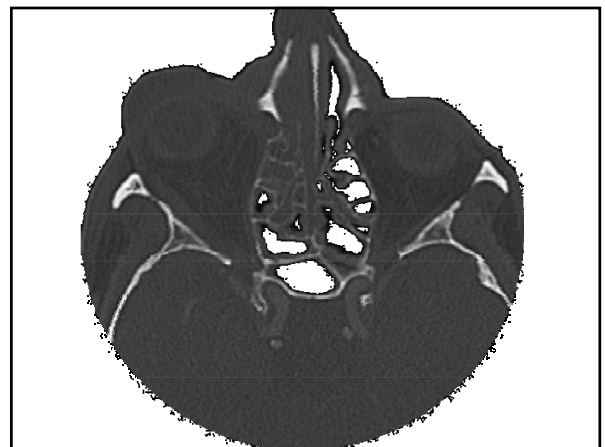
May enable relentless growth

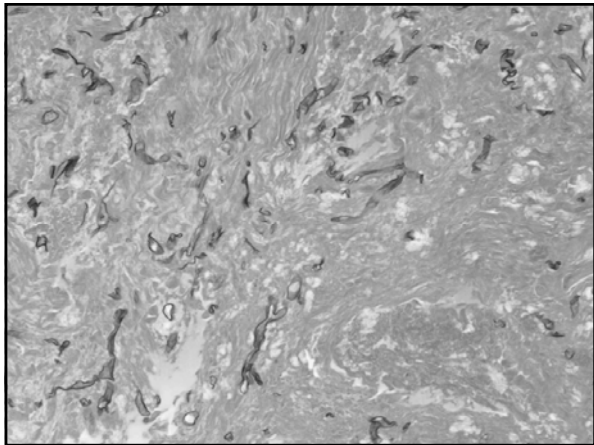
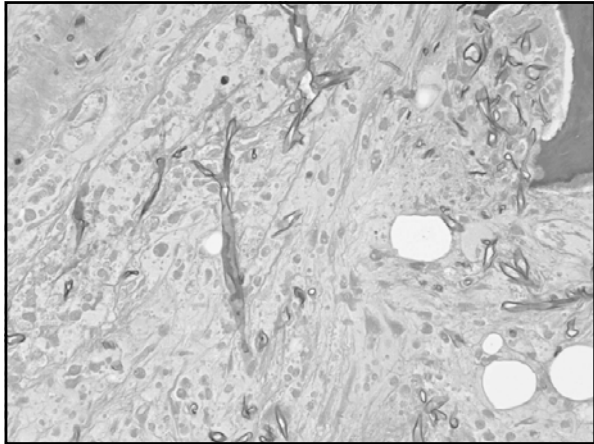
Clinical:

The most acute and fulminant fungal infection known

Lower airways: pneumonia progressing to infarction

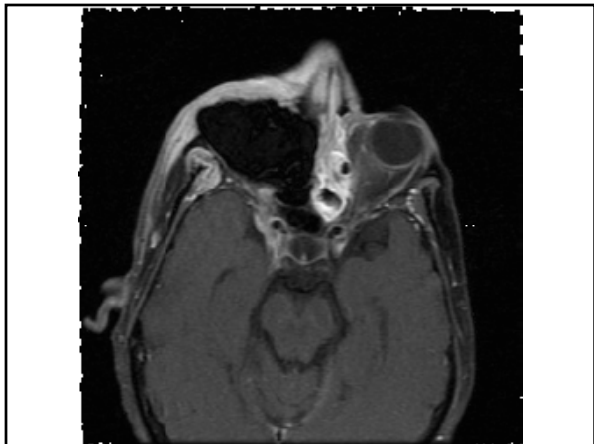
Upper airways: sinusitis progressing to brain abscess





Summary: Fungal “opportunism”

- Metabolic (dermatophytes, *M. furfur*, mucor)
- Dimorphism (sporo, histo, blasto, cocci)
- Capsule (cryptococcus)
- Adherence (candida)



Summary: antifungal defenses

- Intact skin (dermatophytes, *Candida*)
- Lymphocyte function (dimorphs, cryptococcus, candida)
- Neutrophil function (candida, aspergillus, mucor)
- Body milieu (candida, mucor)