IDENTIFICATION OF INFECTIOUS DISEASE PROCESS

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Good morning. My name is Jessie Seidelman and I am one of the ID faculty from Duke. I am also one of the medical directors for Duke University hospital. I am very excited to speak with you today regarding identification fo the infectious disease process.

Disclosures

Royalties: UpToDate, Inc.

Acknowledgements:

Rebekah Moehring, MD



I will disclose that I receive royalties from UptoDate. I also want to acknowledge Dr. Rebekah Moehring for these slides

Objectives

- Infectious disease process (aka. Pathogenesis)
- Clinical signs/symptoms of infection (aka. Immune process)
- Diagnostics and laboratory reports including specimen handling
- Infection vs. colonization vs. contamination
- Antimicrobial Use



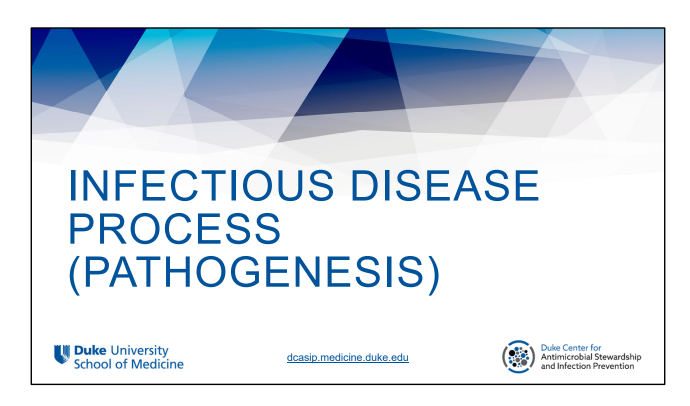
Here are the objectives of this talk. We will be covering pathogenesis, immune process, diagnostics and laboratory reports, true infection versus colonization or contamination and lastly antimicrobial use

Disclosures (2)

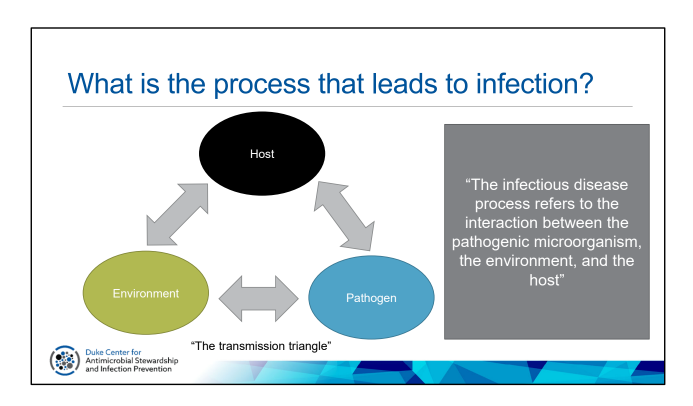
- I cannot teach you all of infectious diseases and microbiology in 90 minutes.
- I will address each core principle broadly and add context.
- To illustrate the core principles I'll add a specific scenario.
- Try to hit key pathogens from each of the (38!) chapters I'm supposed to cover
 - · Average 2.4 minutes per chapter



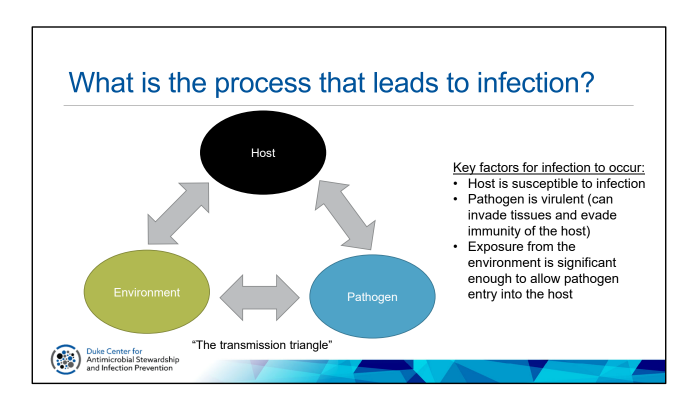
I have a few more disclosures for all of you. First, this is only a 90 minute sessions. People including myself spend years learning about infectious diseases and microbiology. As such, I am going to address each core principle broadly and add some context. I will do my best to hit key pathogens from all of the 38 other chapters in the study guide, but again, there is so much more to all of these topics than we can cover in a short session.



Ok, let's start with the infectious disease process or pathogenesis



So what lead to a person getting an infection? This is really an interaction between three key factors: the host, the pathogen and the environment, which is affectionately refered to as the transmission triangle



Specifically, the factors that contribute to infection acquisition are the host's immune system. Are they susceptible to infection. How virulent a pathogen is in terms of invasion and ability to evade the immune system. Lastly is the environmental exposure. Is the exposure significant enough to allow pathogen entry into the host and are there enough infection units acquired to cause an infection.

Environmental risk factors

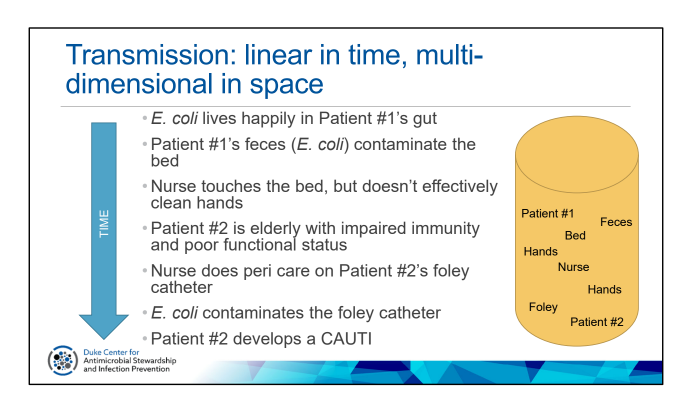
Environment

- People, places/physical structure/space, time
- Healthcare environments (e.g. Hospitals, OR, Clinic, ED, SNF/ALF) or community settings
- Each setting and practice/procedure has it's own unique environmental risks to consider
 - Examples:
 - · Likelihood of needle sticks?
 - · Likelihood of significant contamination events?
 - · Likelihood of an encounter with a returning traveler presenting with fever?
 - · What infections are circulating at this time of year?



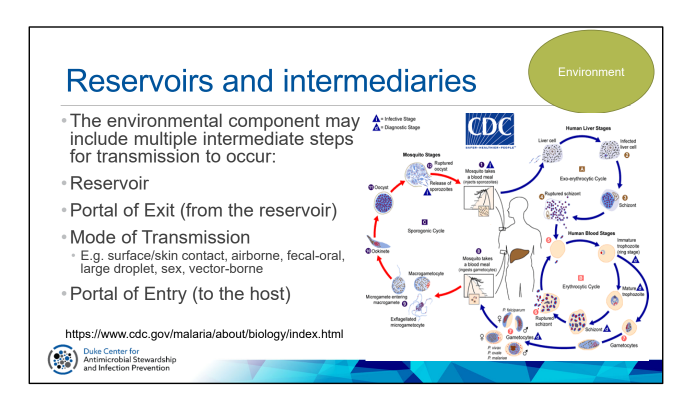
Let's specifically dive into the environmental risk factors. This specifically includes the people, places and time spend in a specific area. Another way to stratify the environment as it related to infectious diseases is healthcare versus community settings.

Each setting has it's own unique environmental risk to consider for example the likelihood of a contamination event? The likelihood of an encounter with a returning traveler with a potentially transmissible infection or even what time of the year should be considered when thinking about environmental risk factors.



So let's think about how transmission may occur via the environment.

E coli is part of patient 1's gut microbiome. Patient 1 had a BM that leads to E coli contamination of the bed. The nurse for patient 1 touches the bed, but doesn't effectively clean his or her hands. Patient 2 is elderly with impaired immunity and poor functional status. The nurse then does pericare on Patient 2's foley catheter. E coli contaminates patient 2's catheter and patient 2 then develops a catheter associated urinary tract infection.



However, the transmission process may not always be this linear. There may be multiple intermediate steps including reservoirs and intermediaries. The best examples of this phenomena are illustrated by the CDC life cycle diagrams. Here is one example of malaria where you can see the mosquito reservoir, how the parasite reproduces in the mosquito and exits the reservoir into the human. This is important in thinking about potential exposures and incubation periods.

The reservoir of an infectious agent is the habitat in which the agent normally lives, grows, and multiplies. Reservoirs include humans, animals, and the environment. The reservoir may or may not be the source from which an agent is transferred to a host. For example, the reservoir of *Clostridium botulinum* is soil, but the source of most botulism infections is improperly canned food containing *C. botulinum* spores.

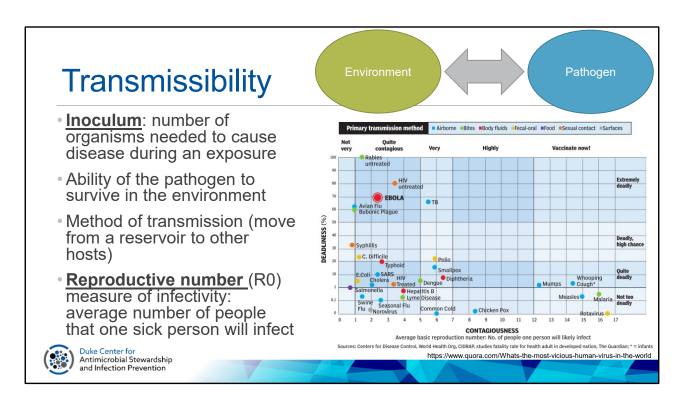
Many common infectious diseases have human reservoirs. Diseases that are transmitted from person to person without intermediaries include the sexually transmitted diseases, measles, mumps, streptococcal infection, and many respiratory pathogens. Because humans were the only reservoir for the smallpox virus, naturally occurring smallpox was eradicated after the last human case was identified and isolated

Humans are also subject to diseases that have animal reservoirs. Many of these diseases are transmitted from animal to animal, with humans as incidental hosts.

Plants, soil, and water in the environment are also reservoirs for some infectious agents. Many fungal agents, such as those that cause histoplasmosis, live and multiply in the soil. Outbreaks of Legionnaires disease are often traced to water supplies in cooling towers and evaporative condensers, reservoirs for the causative organism *Legionella pneumophila*.

Portal of exit is the path by which a pathogen leaves its host. The portal of exit usually corresponds to the site where the pathogen is localized. For example, influenza viruses and *Mycobacterium tuberculosis* exit the respiratory tract, schistosomes through urine, cholera vibrios in feces, *Sarcoptes scabiei* in scabies skin lesions, and enterovirus 70, a cause of hemorrhagic conjunctivitis, in conjunctival secretions. Some bloodborne agents can exit by crossing the placenta from mother to fetus (rubella, syphilis, toxoplasmosis), while others exit through cuts or needles in the skin (hepatitis B) or blood-sucking arthropods (malaria).

The portal of entry refers to the manner in which a pathogen enters a susceptible host. The portal of entry must provide access to tissues in which the pathogen can multiply or a toxin can act. Often, infectious agents use the same portal to enter a new host that they used to exit the source host. For example, influenza virus exits the respiratory tract of the source host and enters the respiratory tract of the new host. In contrast, many pathogens that cause gastroenteritis follow a so-called "fecal-oral" route because they exit the source host in feces, are carried on inadequately washed hands to a vehicle such as food, water, or utensil, and enter a new host through the mouth.



There are a few important definitions and descriptions to consider when it comes to talking about transmissibility of pathogens.

First, inoculum is the number of organisms needed to cause a disease during exposure. For example, norovirus is very infectious, only 10-100 viral particles may be sufficient to infect an individual.

The other thing to consider is how well the pathogen survives in the environment. Again, some pathogens can remain viable in the environment for weeks to months. For instance, C difficile spores are resistant to alcohol and can remain viable in the environment for months

Then there is the method of the transmission. infectious agent may be transmitted from its natural reservoir to a susceptible host in different ways. There are different classifications for modes of transmission. Direct versus indirect. In direct transmission, an infectious agent is transferred from a reservoir to a susceptible host by direct contact or droplet spread. **Direct contact** occurs through skin-to-skin contact, kissing, and sexual intercourse. Direct contact also refers to contact with

soil or vegetation harboring infectious organisms. Thus, infectious mononucleosis ("kissing disease") and gonorrhea are spread from person to person by direct contact. Hookworm is spread by direct contact with contaminated soil. **Droplet spread** refers to spray with relatively large, short-range aerosols produced by sneezing, coughing, or even talking. Droplet spread is classified as direct because transmission is by direct spray over a few feet, before the droplets fall to the ground. Pertussis and meningococcal infection are examples of diseases transmitted from an infectious patient to a susceptible host by droplet spread. **Indirect transmission** refers to the transfer of an infectious agent from a reservoir to a host by suspended air particles, inanimate objects (vehicles), or animate intermediaries (vectors).

Lastly is the reproductive number. This is the expected number of cases directly generated by one case in a population where all individuals are susceptible to infection. The definition assumes that no other individuals are infected or immunized

Practice Question

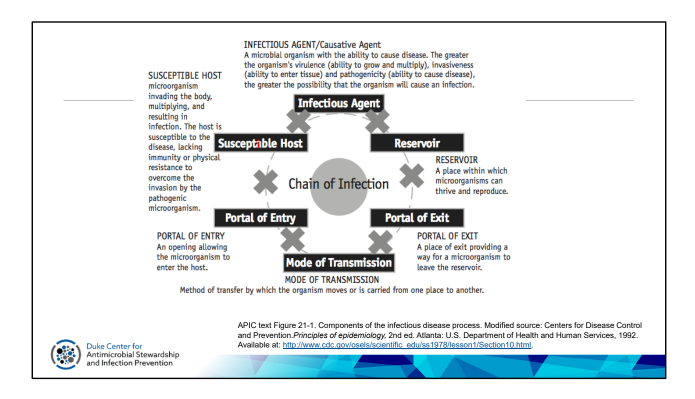
Which of the following statements about influenza is **FALSE**?

A. Influenza is primarily spread between individuals via respiratory secretions (droplets)

- B. Viral shedding starts 48-72 hours after infection and typically 48 hours before the onset of symptoms
- C. Viral shedding normally persists for less than 5 days but can be longer in children and in immunocompromised persons
- D. The typical influenza symptomology is not always predictive of influenza in elderly or immunocompromised persons.



B. Adults shed flu virus from the DAY BEFORE symptoms begin through 5-10 days after illness onset. Incubation for flu is 1 to 4 days (average 2 days).



Here is the diagram that really puts a lot of the concepts we just talked about together

- Think about the susceptibility of the host: does the host lack the immunity or physical resistant o overcome invasion from the pathogen
- Next, think about the actual pathogen. As we talked about previously this includes the virulence or infectiousness, the ability of the organism to survive in the environment, and how many bacteria, virions, parasites etc are able to casue an infection
- The next part is the reservoir. Where doe the pathogen reproduce and thrive
- What is the portal of exit from the reservoir
- Next is the transmission from the portal of exit to the host and finally the portal of entry into the host.

Mitigation of transmission risks

There are MANY things we can do to reduce transmission (examples):

- Environmental engineering, cleaning/disinfection
- Occupational health, avoidance of presenteeism
- Appropriate use of transmission based precautions
- Hand hygiene!
- Cohorting; staffing ratios

Some things are NOT modifiable with facility-level IP (examples):

- Host factors, complexity of patient population
- External factors, e.g. geographic and regional epidemiology
- Cannot wholly avoid risky things like surgery, chemotherapy, and central lines (aka competing risks)



So what can we do to mitigate transmission of pathogens to hosts? This includes environmental engineering, cleaning and disinfection, avoidance of presenteeism in other words staying home when you are sick. Appropriate use of transmission-based precautions, hand hygiene and patient cohorting and appropriate staffing ratios

Unfortunately, there are also factors that we cannot modify within infection prevention. That includes the hosts, geographic or regional epidemiology, and things like chemotherapy, invasive devices or surgery.

Practice Question

An IP is conducting an educational session to help the nursing staff understand infectious disease transmission. She explains that an initial element in transmission is the ability of an organism to survive in the external environment during transit between hosts. What is the second element?

- A. Secretion of enzymes that enhance spread through tissues
- B. A mechanism for transmission to a new host
- C. Invasion and dissemination in the host
- D. Avoidance of host resistance



B. As we talked about portal of entry

Host Defenses

Host

Non-specific defenses against invading pathogens

- Physiologic barriers: Secretions, Fever, normal flora
- Mechanical barriers: Mucosa or skin

Immune system

- Non-specific "Innate" immunity: Phagocytic cells (neutrophils, monocytes), hormones, fibronectin
- Complement system: protein pathways that poke holes and ramp up inflammatory response
- Pathogen-specific "Adaptive" immunity: Cellular immunity (T cells), Humoral immunity (B cells, antibodies)



Ok so now we are moving onto the host.

Nonspecific factors that defend against infection include the skin, mucous membranes, gastric acidity, cilia in the respiratory tract, the cough reflex, and nonspecific immune response. Factors that may increase susceptibility to infection by disrupting host defenses include malnutrition, alcoholism, and disease or therapy that impairs the nonspecific immune response.

When it comes to the immune system specifically tehre are three different components. An individual's genetic makeup may either increase or decrease susceptibility. For example, persons with sickle cell trait seem to be at least partially protected from a particular type of malaria. Specific immunity refers to protective antibodies that are directed against a specific agent. Such antibodies may develop in response to infection, vaccine, or toxoid (toxin that has been deactivated but retains its capacity to stimulate production of toxin antibodies) or may be acquired by transplacental transfer from mother to fetus or by injection of antitoxin or immune globulin.

Practice Question

The IP is teaching nurses how to assess infection risks in patients. Depletion of what cell type provides the best indication of susceptibility to most bacterial infections?

- A. monocyte
- B. eosinophil
- C. neutrophil
- D. lymphocyte



C. Neutrophil

Monocytes. They have a longer lifespan than many white blood cells and help to break down bacteria.

Lymphocytes. They create antibodies to fight against bacteria, viruses, and other potentially harmful invaders.

Neutrophils. They kill and digest bacteria and fungi. They are the most numerous type of white blood cell and your first line of defense when infection strikes.

Eosinophils. They attack and kill parasites and cancer cells, and help with allergic responses.

As a bonus WBC **Basophils.** These small cells seem to sound an alarm when infectious agents invade your blood. They secrete chemicals, such as histamine, a marker of allergic disease, that help control the body's immune response.

Immunity

Terms and numbers to know:

- Normal WBC: 4,000 10,000 cells/mm³
- Leukocytosis: WBC >10,000 cells/mm3
- Leukopenia: WBC <4,000 cells/mm3
- Neutropenia: PMN or band forms <500 cells/mm3 or absolute neutrophil count less than 1000 cells/mm3
 - Absolute count = total WBC count x % of PMN leukocytes
 - Polys: PMNs: mature or segmented neutrophils
 - Bands: Immature or nonsegmented neutrophils
- Infection risk is high when absolute neutrophil count is <500 cells/mm</p>



In terms of immunity here are some terms and numbers to know

A normal WBC range is from 4,000 to 10,000 cells per mm cubed. However, this is more often displayed as 4.0 to 10.0. when the WBC is higher than 10 it is referred to as leukocytosis. When the WBC is lower than 4 it is referred to as leukopenia. Neutropenia is referring to low neutrophils. The specific definition is when the the absolute neutrophils count is less than 1000 or when polys plus ands are less than 500. You can see below that some additional definitions of the absolute neutrophil count, the polys or PMNs and the bands.

Infection risk is high when the absolute neutrophil count is less than 500

Immunity = Defense

Host

Active

- Acquired through prior exposure (+/- resolved infection) or vaccination
- Memory of prior antigens and previously produced humoral reaction (antibodies)

Passive

- Antibodies acquired through other means than a patient's own immune system
 - Maternal (first 6mo of life)
 - IVIG (all types of immune globulin)
 - Can be specific: e.g. rabies Ig, VZIg



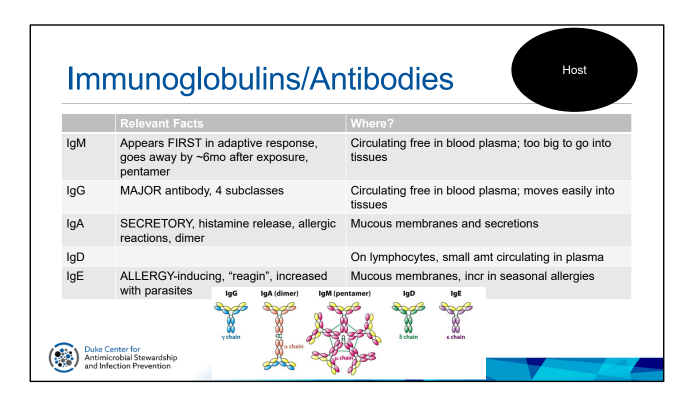
Two types of immunity exist — active and passive:

Active immunity occurs when our own immune system is responsible for protecting us from a pathogen. Active immunity is created by our own immune system when we are exposed to a potential disease-causing agent (i.e., pathogen). active immunity is important because it lasts a long time in the form of immunologic memory. Immunologic memory consists of B and T cells that can recognize a particular pathogen. Vaccines contribute to active immunity by providing us with a controlled way to create an immune response. When a vaccine is introduced, our immune system treats it like any other exposure. It works to stop the "assault" and, in the process, immunologic memory develops. Because vaccines are designed such that they do not cause illness, we gain the benefits of the exposure without the risks associated with fighting off a natural infection.

Passive immunity occurs when we are protected from a pathogen by immunity gained from someone else. Passive immunity, or immunity gained in a way other than from one's own immune system, can occur in a few ways and can be life-

saving. However, passive immunity is short-lived because the antibodies are not continually replenished as they would be in an individual whose immune system is responding directly. Passive immunity can occur in a couple of ways: maternal antibiotics, IVIG

A third category, community immunity, does not involve physical components of the immune system for protection, but is still worth discussion in this capacity. Community immunity occurs when people are protected by those around them. This type of protection is indirect in that it does not involve physical components of immunity, such as antibodies, but rather results when a pathogen is less likely to infect a susceptible person because of the high numbers of protected people around them. Because this immunity is not based on "products" of the immune system, it is the least reliable. However, for some in our communities, such as those too young to be immunized or those with weakened immunity due to illness or treatment, community immunity is the only way they can be protected. This includes the notion of herd immunity



There are five main classes of immunoglobulins—IgG, IgM, IgA, IgD, and IgE

IgM is predominantly found in the lymph fluid and blood and is a very effective neutralizing agent in the early stages of disease. Elevated levels can be a sign of recent infection or exposure to antigen.

IgG is the major immunoglobulin in blood, lymph fluid, cerebrospinal fluid and peritoneal fluid and a key player in the humoral immune response. IgG is produced in a delayed response to an infection and can be retained in the body for a long time. The longevity in serum makes IgG most useful for passive immunization by transfer of this antibody. Detection of IgG usually indicates a prior infection or vaccination. IgG has 4 subclasses. Determination of IgG subclasses can be a valuable tool in indicating a potential antibody deficiency. Selective IgG subclass deficiencies are associated with disease.

Immunoglobulin A (IgA) is the most abundant type of antibody in the body, comprising most of the immunoglobulin in secretions and a significant amount of circulating immunoglobulin. In secretions, it serves to protect the mucosal tissues

from microbial invasion and maintain immune homeostasis with the microbiota.

IgE and IgD are found in serum in much smaller quantities than other Ig classes. IgE primarily defends against parasitic invasion and is responsible for allergic reactions. Membrane IgD is a receptor for antigens found mostly on mature B-lymphocytes.

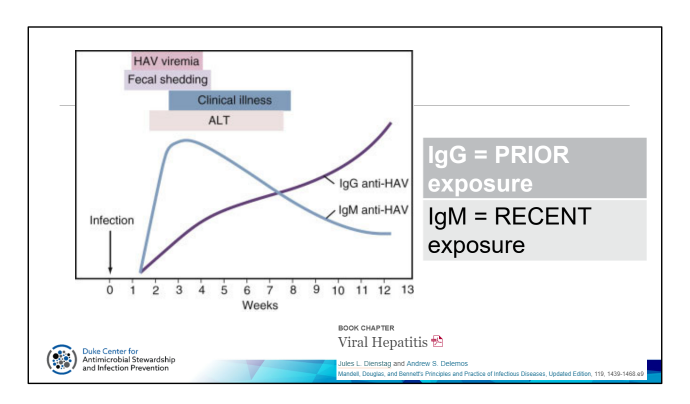
Practice Question

The first immoglobulin response after exposure to a communicable disease pathogen or vaccine is production of:

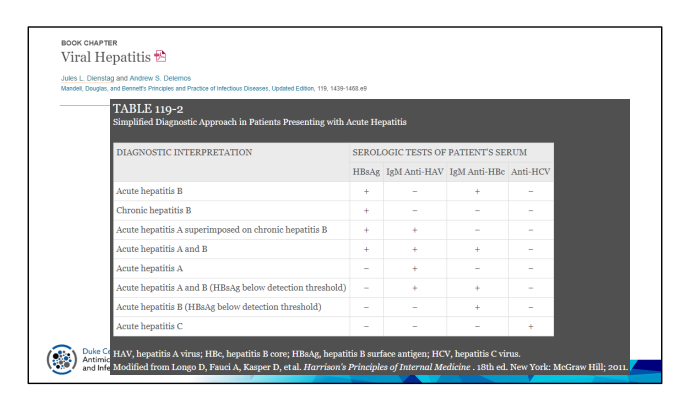
- A. Immunoglobulin G (IgG)
- B. Immunoglobulin M (IgM)
- C. Immunoglobulin A (IgA)
- D. Immunoglobulin C (IgC)



B. I always think M is for Immediate



Here's a slide to help show you when IgG and IgM develop in acute hepatitis A infection. You can see that the IgM peaks and then declines and the IgG slowly increases over time. Knowing the timing of when the body produces these immunoglobulins can be helpful to know if a patient is acutely or chronically infected.



Here is an example of a table that is one that you should save somewhere for reference. It is helpful in diagnosing if a patient ahs acute hepatitis, A, acute hepatitis B, acute hepatitis C, or coinfections.

I'll start by going through the serologies at the top. Hepatitis B surface antigen or HBSAg is A protein on the surface of hepatitis B virus that can be detected in high levels in serum during acute or chronic hepatitis B virus infection. The presence of HBsAg indicates that the person is infectious, except when it might be transiently positive within 30 days after a dose of hepatitis B vaccine (HepB). The body normally produces antibodies to HBsAg as part of the normal immune response to infection

IgM Anti-HAV is going to indicate an acute hepatitis A infection

IgM Anti-HBC Positivity indicates recent infection with hepatitis B virus (<6 mos). Its presence indicates acute infection. IgM anti-HBc should be ordered only when acute HBV infection is a concern.

Anti HCV is going to be antibodies against hepatitis C

Practice question

Higher morbidity rates in chronic hepatitis B virus carriers are associated with a co-infection of which of the following:

- A. Hepatitis A
- B. Hepatitis D
- C. Hepatitis C
- D. Hepatitis E



В.

Hepatitis D virus (HDV) infection is caused by a defective virus: the hepatitis D virus. HDV is often referred to as hepatitis delta virus or delta agent. However, the term HDV is preferred. Individuals with hepatitis D are always dually infected with HDV and hepatitis B virus (HBV). Although HDV can replicate autonomously, the simultaneous presence of HBV is required for complete virion assembly and secretion.

Practice Question

All of the following are descriptions of patients with immunocompromised status EXCEPT:

- A. HIV with CD4 count <200
- B. Leukemia or lymphoma
- C. Neutropenia (absolute neutrophil count <500/mm3)
- D. 1 year post bone-marrow transplant



D. We hope these guys have developed their donor's immunity at 1 year following bone marrow transplant. All of the other scenarios describe patients that are immunocompromised

"Susceptible" Host



- Can include a <u>large variety of factors</u>:
- No prior exposures and thus no adaptive immunity
- Invasive procedures (breaking through mechanical defenses)
- Immunocompromise (partial list)
 - Medications (e.g. high dose steroids, chemotherapy, transplant meds)
 - Malignancy (e.g. real or functional neutropenia)
 - Metabolic (e.g. diabetes, ESRD, ESLD)
 - HIV/AIDS
- Asplenia (e.g. s/p MVA + splenectomy, sickle cell disease)
- Inherited immune deficiency



Ok so what is a susceptible host. This can really include a large variety of clinical elements. It can be something as simple as the patient hasn't been exposed to a specific virus, so they don't have any adaptive immunity. They could have had an invasive procedure. By disrupting the gut or skin barreirs that can make you more susceptible to bacteremia translocating to the wrong place. There are also many medications and diseases that put patients at increased risk of infection. Specifically this could be medicaitons like high dose steroids or chemoterhapy. This could also be leukemias that reduce your numbe rof functional neutrophils. Even metabolic disorders such as diabetes can increase your risk of infection. HIV or AIDS puts you at risk of opportunistic infections. Similarly, if you have asplenia or functional asplenia from sick cell disease then your body cannot control the production and removal of blood elements. Lastly, inhereited immune deficiency disorders such as Severe Combined Immunodeficiency (SCID) can put an individual at risk for infections

Pathogen Virulence

Pathogen

Factors about the pathogen that can contribute to its ability to invade the host, evade host immunity, or survive:

Advantage	Example virulence factor	Pathogen/syndrome
Enzymes to increase local tissue damage/spread	Toxin production	S. pyogenes and necrotizing fasciitis
Invade, disseminate	Motility	E. coli swimming up a ureter
Evade host defenses	Biofilms Attach or adhere to surfaces Alter cell wall or membrane Capsule prevents phagocytosis	Coag-neg Staph on IV line S. aureus on a prosthetic knee HIV S. pneumoniae
Survive in harsh conditions	Spore-formation Lipid coat	C. difficile, Bacillus spp. M. tuberculosis
Anti-inabilit Chausadabia		



Ok so we have talked about the environment. We have talked about the host. Now it is time to bring it home and tlak about the pathogen.

So there are different factors that contribute to its ability to invade the host, evade host immunity or survive.

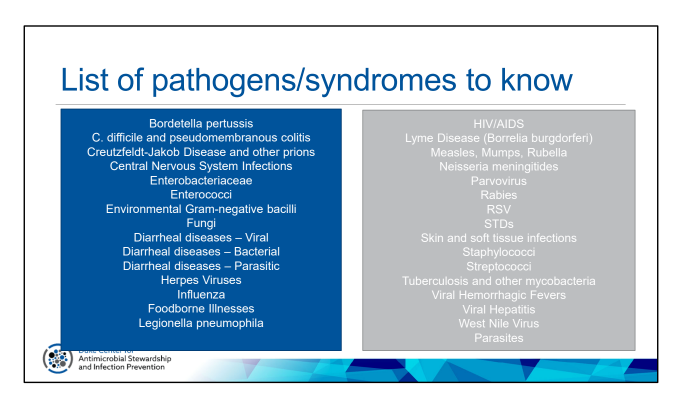
Certain bacteria may produce enzymes to increase local tissue damage to promote their spread such as strep pyogenes or group A strep toxin in the setting of necrotizing fasciitis. Some bacteria are very motile and in that way can invdde an easily disseminate such as E coli's ability to swim up a ureter. Biofilms are a great way that bacteria can evade host defenses. This happens a lot with indwelling prosthetic devices such as coagulase negative staph on an IV line or staph aureus in a prosthetic knee infection. Some bacteria can also live in very harsh conditions. For example in C difficiel or bacillus species they can live in a spore form which are hardy and can survive for weeks to months out in the environment.

Pathogen-specific features you must know

- Type of microorganism: bacteria, virus, fungus, parasite
- Clinical features of infection
- Laboratory diagnosis (e.g. culture, serology, PCR)
- Precautions recommended in healthcare setting (e.g. contact, airborne, droplet, standard)
- Key transmission data:
 - Mode of transmission
 - Timing: incubation period and "shedding"/contagious period (key for droplet, airborne, and some contact/viruses), typical duration of symptoms
 - Vectors



In terms of specific pathogen features that you need to know here is a brief list. You have to know whether the microorganism is a bacteria, virus, fungus, parasite or a prion. How do these pathogens clinically manifest themselves? What are the correct studies that we should send to the lab to evaluate fo rthem. Whata re the correct precautions that a patient should have in a healthcare setting if they are being treated for or suspected to have one of these pathogens. The timing and type of precautions really depends on how the microorganism is transmitted and when it is transmitted.



Here is a general list that you can reference when you studying for your exam. Again, you want to know what is causing the illness, how it presents, what is used to test for it, and what precatuions a patient might need and for how long.

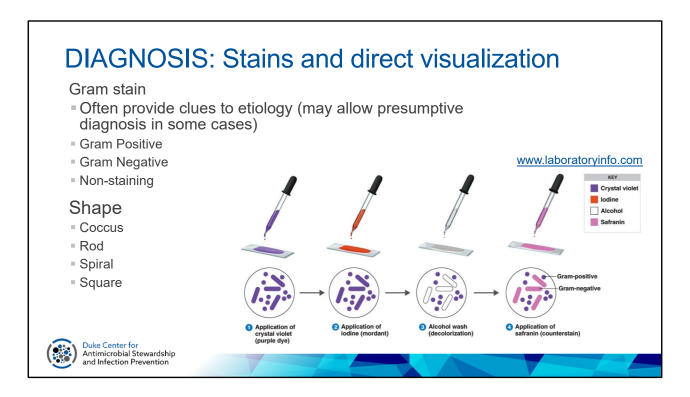
Quick and Dirty Micro Classification

- Gram positive skin, lung, guts, devices
- Gram negative guts, urine, some lung
- Atypicals lung, STIs, ticks
- Anaerobes –gas- and abscess-forming, bad odors, guts
- Less commonly encountered: Mycobacterium (lung), spirochetes (Syphilis and Borrelia)
- Fungal guts, devices, really bad in immunosuppressed hosts



Here are some quick microbiology correlations that may help you think about where the infection is coming from

When it comes to gram positive organisms, think about the skin, the lungs, the GI tract and implanted devices. When it comes to gram negatives, thing about the GI tract, urine, and some lung pathogens. Atypical pathogens that don't have true cell walls like legionella, mycoplasma, and chlamydia. Think about the lung, sexually transmitted infections and tick born illnesses. When it comes to anaerobic pathogens, think of gas and bascess forming clinical pictures, bad odor wounds and the GI tract. Less commonly encountered pathgoens may be mycobacteriaand spirochetes such as borrelia. I am not sure theat syphilis in generally is less commonly encountered but perhaps in the inpatient IP setting it may be. Lastly, fungal apthogens think of guts, devices, particularly if a aptient is receiving total parenteral nutrition and really immunocompormiesd individuals.



So how does the identification of organisms actually happen in the alb. Often times a gram stain is performed and this can give you information pretty quickly. The gram stain can tell you if the organism is gram positive (will be purple), gram negative (will be pink) or non-staining. You will also be able to see the shape of these pathogens that will give you good clues as to what you are dealing with. Si it a coccus or a circle, a rod, a spiral or a square

Significance of the Gram Stain

By knowing the shape and gram staining reaction of the organisms, along with the body site involved; clinicians can make a reasonable guess as to the causative agent.

The reasonable guess can guide early empiric antibiotic choices.



The gram stain is a very old technology but incredibly important to you as IPs and also to clinicians who are trying to empirically treat patients. Knowing the shape and gram stain can help guide early empiric antibiotic choices.

Practice Question

When reviewing the Gram stain of a person with a wound infection, the IP sees Gram-positive organisms in clusters. Which organism would this most likely represent?

- A. Streptococcus
- B. Enterococcus
- C. Corynebacterium
- D. Staphylococcus



Ok, time for a question break!

D. Remember staph is from the Greek word staphyle (bunch of grapes) and kokkos (berry),

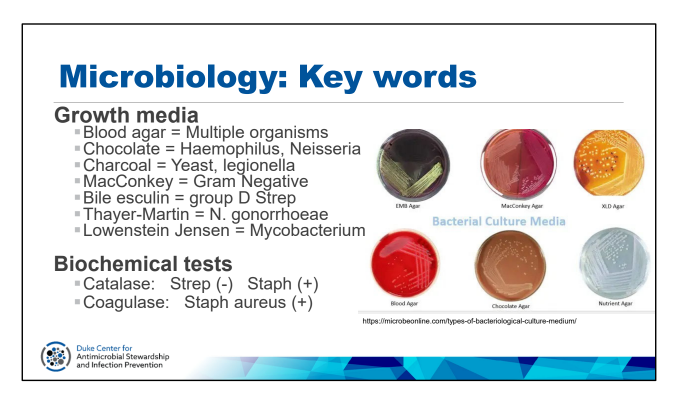
Microbiology

Physical requirements for growth of bacteria

- Nutrition (media)
- ■Temperature 35° for most bacteria
- Atmospheric conditions
 - Aerobic (needs oxygen to survive)
 - Anaerobic (needs <u>absence</u> of oxygen to survive)
 - Facultative anaerobes (with or without oxygen)
 - Microaerophilic



There are other things about the bacteria that you need to know if you are going to try and grow them up in the lab and isolate them. Specifically there are specific physical requirements for growth of bacteria. They need specific media to grow that contain various nutritional elements. There also needs to be controlled termpeature and atmospheric conditions. Aerobic bacteria need oxygen verus anaerobic bacteria that need there to be NO bacteria in order to survive. Facultative anaerobes can grow in either. Microaerophilic means they require lower levels of oxygen than are present in the atmosphere to survive.



So here is another good slide for reference. Here you have what growth media with support which pathogens. I will say that the standard agar is blood agar. MacConkey is for gram negative pathogens. The rest of them are selective agar meaning that they are only going to support specific pathogens for growth. The other thing I have on this slide is the biochemical tests. For the catalase test they basically add hydrogen peroxide to the plate and see if it produces bubbles. however it should never be performed on organisms that have been grown on blood agar (a medium that contains blood). This is because there is a catalase activity in blood that would produce a false positive result. The other test is the coagulase test. The coagulase test is one way to differentiate the highly pathogenic S. aureus from the other less pathogenic staphylococcal species. If positive you will basically see cells clumping together on the sldie

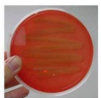
Microbiology: Key words

Hemolysis on Blood Agar

- Alpha = green
- Beta = clear
- Gamma = no hemolysis







Beta Hemoly

Alpha Hemolysi

https://microbiologyinfo.com/haemolysis-of-streptococci-

Gamma Hemolysis

Lancefield grouping = Strep A to O
Bile esculin = black pigment
Optochin inhibition: S. pneumoniae (+)



Some other key microbiology grouping is whether the bacteria causes hemolysis or not. The key difference between alpha beta and gamma hemolysis is that alpha hemolysis is the partial destruction of red blood cells in the blood and beta hemolysis is the complete destruction of red blood cells in the blood, while gamma hemolysis does not involve any breakdown of red blood cells. This equates to the visualization of the bacteria on the plates as clear (complete hemolysis), green (partial hemolysis) and red (no hemolysis)

Other keywords are the lancefield grouping. This is a system of classification that classifies <u>catalase</u>-negative <u>Gram-positive</u> <u>cocci</u> based on the carbohydrate composition of bacterial <u>antigens</u> found on their <u>cell walls</u>.

The bile-esculin test is widely used to differentiate enterococci and group D streptococci, which are bile tolerant and can hydrolyze esculin to esculetin, from non-group D viridans group streptococci, which grow poorly on bile.

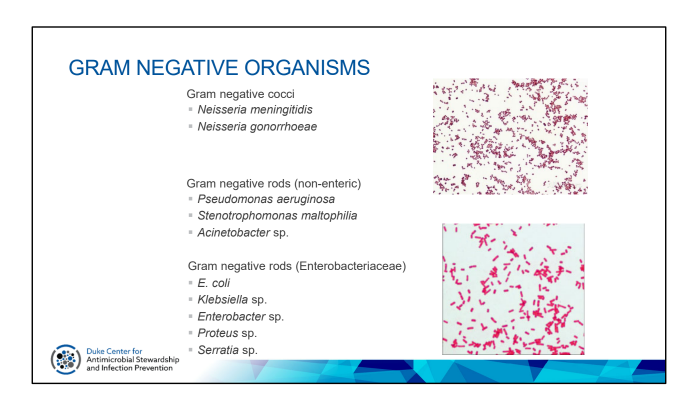
Optochin is a chemical that is toxic to some bacteria but harmless to others. It is useful in the identification of *Streptococcus pneumoniae*, the alpha-hemolytic

Streptococcus most commonly susceptible to this chemical.

GRAM POSITIVE ORGANISMS Gram positive cocci Staphylococcus aureus Coagulase negative staphylococcus Streptococcus pneumoniae Streptococcus sp. Enterococcus sp. Enterococcus sp. Gram positive rods Bacillus sp. (aerobes) B. antracis Clostridium sp. (anaerobes) Listeria

Lets talk a little bit about common gram positive organisms. Again, tehse are purple on the stain. When ti comes to gram positive cocci these include staph aureus (clsuters), coagulase negative staphylococcus, streptococcus pneumoniae, streptococcus, and enterococcus. Again, you can see why we may need some of those biochemical tests after we see purple cocci on the gram stain.

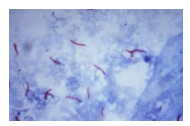
When it comes to gram positive rods, if they are growing in the aerobic tube think bacillus. If they are only in the anaerobic tube think clostridium. Listeria is a facultative anaerobe, so it can grow under both conditions



How about the gram negative pink-staining organisms. When you see gram negative cocci think Neisseria. The buzz word for this is gram negative diplococci. When it comes to gram negative rods we typically divide these into the enterics meaning they grow better in anaerobic conditions versus non-enterics that you more likely find in lungs and oxygen rich environments.

NON-STAINING/Special stain PATHOGENS

- Not stained by Gram's method
 - Legionella sp.
 - Chlamydia
 - Rickettsia
 - Mycobacteria
 - M. tuberculosis
 - Non-tuberculous mycobacteria



Ziehl-Neelsen Stain of TB

Aka. Acid Fast



How about the pathogens that are not going to stain on a standard gram stain? These are organisms such as legionella, chlamydia, rickettsia, and mycobacteria. Classically you need to do a ziehl neelsen stain to see mycobacterial organisms. Acid fast bacteria will be red, while nonacid fast bacteria will stain blue/green with the counterstain

Organism Diagnosis

Fungal

- Morphology
- Presence of hyphae
- Size of yeast
- Presence of capsule

Virology

- Direct- electron microscopy
- Antigen detection
- Virus isolation from culture
- Antibody detection/serology
- PCR testing

Parasitology

- Direct exam
- Microscopy (oocytes)
- Antigen detection/serology



How about diagnosing other patheogens? When it comes to fungi what is really helpful is how they look, do they have hyphae are they branching or not branching and if they branching do they branch at acute angles. Are there fungal capsules? What is the size of the yeast?

For viruses typically we will use PCR testing or antigen detection. Antigen detection his how many rapid covid tests work. Antibody detection and serology can also be helpful. Virus isolation from culture is very difficult as is direct electron microscopy.

When it comes to parasitology these are typically larger organisms that we look at under the microscope. You can also use antigen detection and serology.

Choice of Empiric Antimicrobials

What class of pathogen am I likely to be treating?

• (Bacterial? Viral? Fungal? Other?)

If bacterial, what organisms are most likely?

Gram positive? Gram negative? Anaerobe?)

What information can I get to guide treatment?

Microbiology data?

Do I need to order any other diagnostic tests?

How sick is my patient? How risky would it be if I miss?



When it comes to empiric antibiotics you have to ask yourself a few questions: what class of pathogens am I treating. If bacterial is this most likely gram positive? Gram negative? Anaerobic? What information can I get to guide my treatment? In other words what test should I order. And lastsly, how sick is my patient? How risky would it be if I missed the coverage of a potential pathogen?

General Indications for Antibiotics Empiric Prophylaxis: prevent infection Broad-spectrum EASY! Guidelines and ordersets E.g. surgical prophylaxis, pre-transplant protocol Empiric: when you suspect infection but don't exactly know with what Not easy. Local guidelines help (based on local micro data). Clinical syndrome guides choice Directed: pathogen known Only available for a small portion of folks treated for infection Moderately easy. Follow and interpret patient-specific micro data. Targeted Narrow-spectrum **Duke Center for** Antimicrobial Stewardship and Infection Prevention

IN terms of general indications for antibiotics you can think about it in 3 big buckets. First is prophylaxis. This means that we are trying to prevent infection. Think of surgical prophylaxis. We know that the surgical site gets contaminated by bacteria so we use these to prevent a subsequent SSI. The next big pucket is empiric antibiotics. You don't know what is causing the infection, but you are treating a specific clinical syndrome. For example, you suspect that a patient has cellulitis. You don't know what is causing it all the time, so you start with empiric vancomycin to cover gram positive organisms. The last one is the directed antibiotic therapy meaning that you know that causative pathogen such as an E coli UTI. Unfortunately, this probably represents the smallest proportion of folks we see in the hospital

De-escalation De-escalation is a core principle of Antimicrobial Stewardship. Target/narrow antibiotic therapies after more clinical data returns Stop therapy when infection has been ruled out Targeted Narrow-spectrum Targeted Narrow-spectrum

De-escalation Allows initial broad therapy to maximize initial effective therapy Then target/narrow antibiotic therapies after more data returns (microbiology, clinical progress, diagnostic tests)

Avoid unintended consequences of extra days of broad therapy.

DIAGNOSIS: Culture

- "Gold standard" to identify the pathogen
- Requires sampling of site of infection, best if collected *prior to* therapy
- Allows determination of antimicrobial susceptibility
- Can be "banked" for future tests if needed, e.g. outbreak investigations, strain typing, PFGE





So let's talk about cultures as a means of diagnosis. Cultures are really considered the gold standard. This requires sampling of the site of infection prior to therapy. Typically we say that patient should be off of antibiotics for 14 days prior to cultures in order to maximize culture yield. Cultures also allow determination of antimicrobial susceptibility and can help with outbreak investigations so we can compare genetics

DIAGNOSIS: Culture

- There are some key limitations to traditional culture methods:
 - Time and resource intensive
 - Typically have no info other than the stain for 2-3 days
 - Highly reliant on specimen collection techniques
 - Sometimes positive in absence of infection
 - Sometimes negative when infection is present (e.g. in the setting of antibiotics) or get contaminated/mixed flora result





However, there are some key limitations to cultures. Ti is time and resource intensive. You typically won't have a lot of information for the first couple of days aside from the gram stain and it requires folks in the micro lab to look at plates and perform biochemical tests. That being said it is very reliant on specimen collection techniques such as blood culture collection and can be positive in the absnce of infection such as is the case with contamination with blood culture collections. In addition, it can also be negative when infection is present such as in the case when patients are on antibiotics or when you can contamination or mixed flora.

Practice Question

Guidelines for transporting specimens include:

- 1) Transport within 2 hours of collecting a specimen
- 2) Transport in leakproof specimen containers and sealable leakproof bags
- 3) Transport specimen in the syringe used to collect it
- 4) Refrigerate all specimens prior to transport
 - A. 1,4
 - B. 2,3
 - C. 1,2
 - D. 3,4



C you want to transport the scimens within 2 hours of collection and in leakproof specimen contains and leakproof bags

Culture Contamination

- Inadequate specimen collection technique can lead to confusing results.
- Best example: Contaminated blood cultures with skin flora infection or not?
 - Solutions: Collect blood cultures in pairs. Avoid drawing from existing lines. Hire/educate phlebotomists.
- Other examples: Patients had clinical symptoms, are sick, but culture comes back "mixed flora" and pathogen remains unknown.
 Patient is treated with broad spectrum therapy.
 - Example: urine cultures from existing foley catheter (doh!)
- Example: lower respiratory cultures

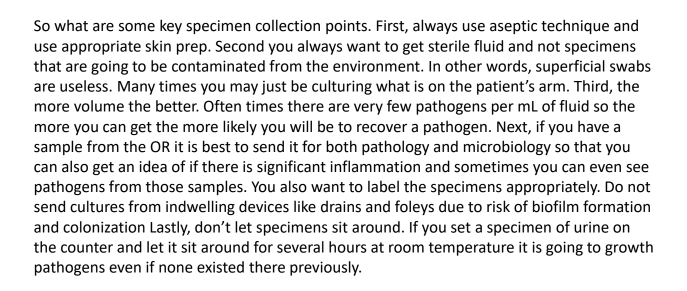


So lets talk a little bit more about contamination of cultures. First and foremost inadequate specimen collections can lead to confusing results. They make you think is this real? Is it not? The best example is contaminated blood cultures with skin for a. More specifically your patient gets 2 sets of Blood culture s and one of them grown coaguals enegative staph, which is a known skin colonizer. Is this real? This is why we always ask to collect blood cultures in pairs and avoid drawing blood cultures from indwelling lines. Other examples include urine cultures from an existing foley catheter. Remember each day a foley caltheter is in place you increase the risk of colonization by 5-7%, so you may just be getting the organism that is living in the catheter but not the true pathogen causing infection. The same thing goes with lower respiratory cultures. You have to get a good quality sample.

Specimen Collection: Key points

- Do not contaminate sterile specimens: Aseptic technique + sterile specimen carriers; Appropriate skin prep; Get more than 1
- Tissue > Fluid >>>>> Swab (avoid!)
- More volume is better (blood, fluids)
- Send tissue from the OR to BOTH path and micro
- Label appropriately and include key clinical clues for the lab, esp for pathogens that are more difficult to culture
- Don't send cultures from drains/foleys that are already in place
- Don't let specimens sit around (to lab within 2h preferred)





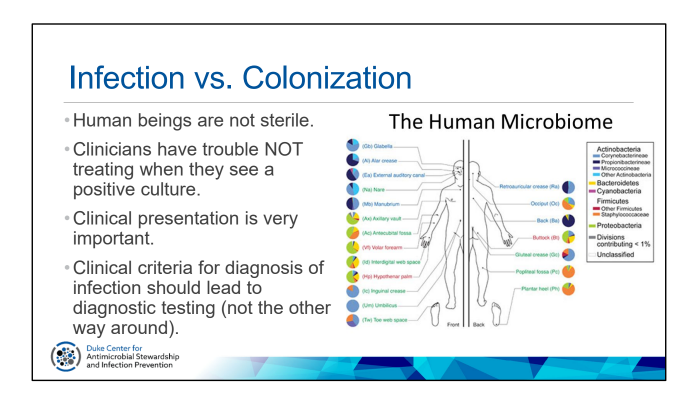
Practice Question

A patient has a nasal swab positive for methicillin-resistant Staphylococcus aureus (MRSA) in the absence of symptoms. This is an example of:

- A. Normal flora
- B. Colonization
- C. Asymptomatic infection
- D. Symptomatic infection



B. Colonization. MRSA is a very common colonizer of our nasal passages



So what exactly do we mean by infection versus colonization? Human beigns are not sterile. We are made up of communities of organisms that was affectionately refer to as the human microbiome. However, clinicians have trouble not treating when they see a positive culture. Clinical presentation is very important. In that same vein go fhtought clinical criteria for diagnosis should lead to testing not testing leading to clinical suspicion.

Infection vs. Colonization

- ?
- Diagnostic tests used indiscriminately lead to overdiagnosis, overtreatment, and associated negative consequences.
 - Exhibit A: Asymptomatic bacteriuria
 - Exhibit B: Patient colonized with *C. difficile* had only 1 loose BM after kayexalate = +PCR and classified as HO-CDI LabID event
- Questions to ask before sending diagnostic test:
 - "What is the pre-test probability that this patient has infection?"
 - "What would I do differently if the test comes back positive? Negative?"



By just sending off tests, this can lead to overdiagnosis, overtreatment and postentially negative consequences. The one we cite all the time is asymptomatic bacteruria. Patietns get antibiotics and then develop C difficile infection. So before you send off a diagnostic test you always want to ask yourself, what I the pre-test probability that this patient ahs an infection and how would this change my management if the test is positive or negative

	Establish Criteria for Testing Urine				
	Diagnoses	Urine Culture	Clinical Symptoms		
\$9.55 \$ 55	Acute, uncomplicated urinary tract infection	>100,000 bacteria, No more than 2 species of bacteria	Dysuria OR Fever AND 1 of the following: -Frequency -Urgency -Suprapubic pain -Incontinence* -Gross Hematuria**		
	Asymptomatic Bacteriuria	>100,000 bacteria, No more than 2 species of bacteria	No signs or symptoms referable to the urinary tract		
Duke C Antimic and Info Public Health Image Library	Stone et al. Infec Control Hosp *New or worsening of baseline i **I have never known hematuria mucosa, which can lead to urina	ncontinence to a sign of infection in an older adu	It. Rather, it seems to indicate trauma to the		

Change in inSo how do we differentiate between asymptomatic bacteruria and true cystitis. The big difference is in clinical symptoms. In a true infection a patient will have symptoms versus in asymtpaomtic bacteruria there are no symptoms. Note of what is not in clinical symptoms for cystitis and that is confusion. There is no study that links confusion to cystitis.

DIAGNOSIS: Antigen Tests

Identifies pathogen-specific proteins

- Very useful for diagnosing viral infections: HIV, HBV, COVID
- Occasionally useful for others: Cryptococcus antigen (CSF, blood), S. pneumoniae (urine), legionella (urine)



Ok so let's talk a bit about antigen tests. All of you should be very familiar with the test on the bottom left. These identify pathogen-specific proteins. They are particularly useful for viral infections like COVID, HIV, Hep B. It can also be useful for other infections such as cryptococcus, stpre pneumoniae and legionella.

DIAGNOSIS: Serologic testing



- Detects immune response to a pathogen, or prior exposure to a pathogen
- For bacterial infections, generally not useful in early diagnosis (may require acute and convalescent tests)
- For viral infections, IgM indicates early diagnosis or recent exposure (e.g., Hep A)
- Important for screening for prior exposure, documenting immunity, and ensuring vaccination
- e.g. Occupational health titers for varicella, HBV
- Once serology is positive, it is typically life-long



Ok so what about serologic tests. Tehse detect the immune response to a pathogen or a prior exposure to a pathogen. For bacterial infections these are not ypically useful. However, for viral infections we can look for that IgM positivity to indicate early diagnosis or recent exposure. These tests are also important for screening for prior exposure such as for varicells and hepatitis B. Once a serology is positive it is typically life-long.

DIAGNOSIS: Molecular tests

PCR and other "molecular" tests

- Increasingly used allows diagnosis of non-culturable pathogens (e.g., norovirus) and faster identification(e.g., pertussis, MRSA in blood);
- Subject to false positives due to sensitivity (e.g. *C. difficile*)



How about molecular tests? These are increasingly use to diagnose non-culturable pathogens such as norovirus and allow faster identification for pathogens such as MRSA or pertussis. However, they are very sensitive and thus can cause false positives as is the case with C difficile infection. You can have a positive PCR when it really only presents colonization as opposed to a true infection. Tha'ts why we often have to combine the PCR with a toxin-based test

DIAGNOSIS: Sterile fluid studies

- Evidence of infection due to inflammatory, chemical, and cellular changes in body fluids
- Examples: synovial fluid, CSF, pleural, and peritoneal fluid
- Typically combined with GS/culture (which takes a while)





When getting studies on sterile fluid we are looking for inflammatory, chemical and cellular changes. For example in synovial fluid or CSF we are looking for the number and types of cells in the fluid to help us know if there is infection. This is typically combined with gram stain and culture

Practice Question

An IP is reviewing the cerebrospinal fluid (CSF) result from a patient admitted the previous night. The CSF is cloudy and has an elevated White Blood Cell count (WBC), markedly elevated neutrophils, low glucose, and elevated protein. What type of meningitis should she suspect?

- A. Bacterial
- B. Viral
- C. Fungal
- D. Aseptic



Α.

Bonus – typical pathogens to suspect for community onset bacterial meningitis in adults Neisseria meningitides (contact and droplet)

- H. Influenza
- S. Pneumoniae

Listeria monocytogenes

	Opening Pressure	Glucose (Ratio of CSF to Serum)	WBC count	WBC type	Total protein	Stain
Bacteria, "Septic"	Elevated	Normal to decreased	≥1,000/m m ³	Neutrophils (early or partially treated may have lymphocyte predominance)	Elevated (mild to very)	Gram* stail may show GPC or GNC/GNR
Virus, "Aseptic"	Usually normal	Usually normal	<100 per mm ³	Lymphocytes	Normal to elevated	Gram stain negative
Fungi	Variable	Low	Variable	Lymphocytes	Elevated	India ink (Crypto), positive
ТВ	Variable	Low (can be extremely low)	Variable	Lymphocytes	Elevated	AFB stain, positive

Here's another table that you might want to keep as reference. This gives you can idea of what kind of meningitis you are looking at based ont eh opening pressure, the glucose, the number and types of white blood cells and the total protein.

Antibiotic susceptibility testing: Key Terms

- Antibiotic = A drug that kills or inhibits the growth of microorganisms
- Resistant = An antimicrobial will NOT inhibit bacterial growth at clinically achievable concentrations
- Susceptible = An antimicrobial WILL inhibit bacterial growth at clinically achievable concentrations
- Intermediate= An antimicrobial may not inhibit bacterial growth at typical doses



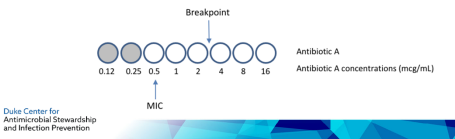
Ok let's talk a little bit about antibiotic susceptibility. An antibiotic is a drug that kills or inhibits the growth of microorganisms. If a pathogen is resistant that means that the antibiotic will NOT inhibit the bacterial growth at clinically achievable concentrations, so you shouldn't use that antibiotic to treat that infection. If a pathogen is susceptible to a specific antibiotic that means that it will inhibit bacterial growth at a clinically achievable concentration, so you can use that antibiotic treat the infection. Lastly, intermediate means that it may not inhibit bacterial growth at typical doses. We typically would not use this antibiotic alone or at standard doses but may use it in conjunction with another antibiotic that the pathogen is susceptible to or at higher doses.

Key Terms

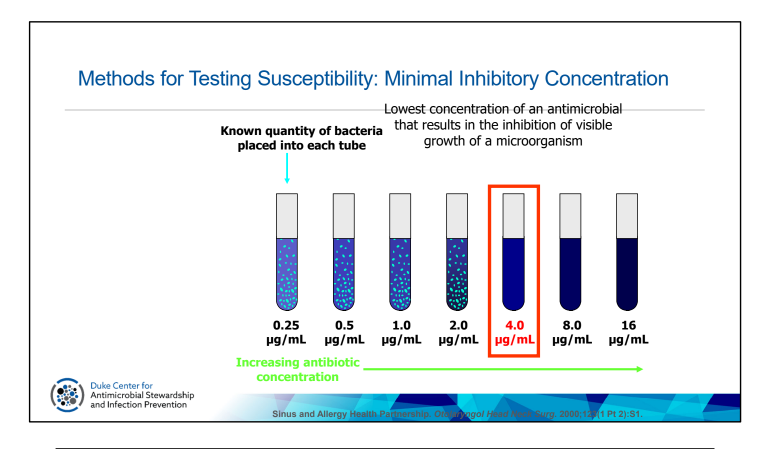
MIC = Minimal inhibitory concentration. Lowest concentration of antimicrobial that inhibits growth of bacteria. Commonly used in clinical lab

MBC = Minimal bactericidal concentration. Concentration of an antimicrobial that kills bacteria. Used clinically only in special circumstances

Breakpoint = The MIC that is used to designate between susceptible and resistant. Set by an expert committee (CLSI).



Ok so these are a little tricky and may take a bit of time to fully grasp. The MIC is the lowest concentration that an antibiotic inhibits growth of bacteria. The lower the better. The MBC is the concentration that kills the bacteria. We don't use this clinically. A breakpoint is the MIC number that is used to designate between susceptible and resistant and this is set by the CLSI committee. These breakpoints do change over time.



So how do they determine the MIC? identical doses of bacteria are cultured in wells of liquid media containing progressively lower concentrations of the drug. The minimum inhibitory concentration of the antibiotic is between the concentrations of the last well in which no bacteria grew and the next lower dose, which allowed bacterial growth

Methods for Testing Susceptibility

- Broth dilution = MIC testing (Automated system) = a number
- Disc Diffusion = Kirby Bauer (Manual) = a zone size
- E test = "Strip" (Manual) = a number











What are the methods for determining test susceptibility. The first one is broth dilatuion, which is the MIC testing we just spoke about. The next one is disc diffusion, which are the pictures in the middle. In diagnostic laboratories, the test is performed by inoculating the surface of an agar plate with bacteria isolated from a patient's infection. Antibioticcontaining paper disks are then applied to the agar and the plate is incubated. If an antibiotic stops the bacteria from growing or kills the bacteria, there will be an area around the disk where the bacteria have not grown enough to be visible. This is called a zone of inhibition. The susceptibility of the bacterial isolate to each antibiotic can then be semiquantified by comparing the size of these zones of inhibition to databases of information on known antibiotic-susceptible, moderately susceptible and resistant bacteria. In this way, it is possible to identify the most appropriate antibiotic for treating a patient's infection.[1][2] Although the disk diffusion test cannot be used to differentiate bacteriostatic and bactericidal activity, it is less cumbersome than other susceptibility test methods such as broth dilution. The last test on this slide is the E test. You set the reagent test strip with a predefined gradient of antibiotic, covering a continuous concentration range. It is applied to the surface of an agar plate inoculated with the test strain, where there is release of the antimicrobial gradient from the plastic carrier to the agar to form a stable and continuous gradient beneath and in nearby to the strip. After the test, the bacterial growth becomes visible after incubation

and a symmetrical inhibition ellipse centered along the strip is seen. The MIC value is read from the scale in terms of $\mu g/mL$ where the ellipse edge intersects the strip. After the required incubation period, the minimum inhibitory value is read where the edge of the inhibition ellipse intersects the side of the strip

Resistance Mechanisms

Intrinsic- inherited by the organism species

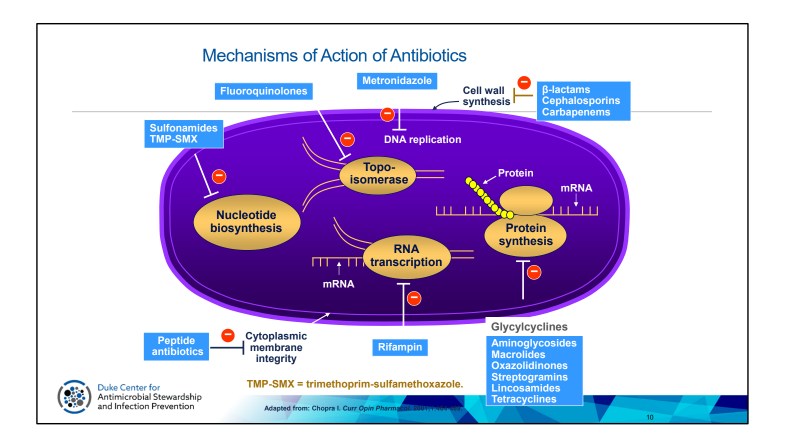
E.g. Serratia is intrinsically resistant to cefazolin

<u>Acquired</u>- results from altered cellular structure and physiology caused by changes in the genetic make-up

- Efflux pump
- Point mutation on a penicillin binding protein
- Beta lactamase production:
 - ESBL K. pneumoniae acquired resistance from a plasmid from E. coli
- Carbapenemase production



So what are resistance mechanisms. These can really be thought of in two braod categories. The first is intrinsic resistance meaning that is is inherited by the organism. All serratia are intrinsically resistant to cefazolin. You can also get acquired resistant from altered cellular structure and physiology. These can include things like efflux pumps to pump antibiotic out of the cell. Mutation son a penicillin binding protein making antibiotics unable to hook onto them. Beta lactamase production meaning that these enzymes can cut beta lactam rings and make them ineffective and carbapenemease production, which make carbapenems ineffective.



Slide 10.

Antibacterial drugs prevent bacterial growth by disrupting the function of a wide variety of molecular targets located within bacteria and at the cell surface. The penicillins, cephalosporins, and carbapenems all target cell wall synthesis by inhibiting transpeptidases required for peptidoglycan formation and the cross-linking of structures in the cell wall. Metronidazole interacts with DNA and inhibits nucleotide synthesis, leading to cell death. The fluoroquinolones block DNA synthesis by inhibiting DNA gyrase, which is responsible for folding and supercoiling the replicating DNA so that it does not become entangled with itself. The sulfonamides and trimethoprim-sulfamethoxazole (TMP-SMX) inhibit the metabolism of bacteria by inhibiting enzymes needed for the synthesis of folic acid. The peptide antibiotics (eg, vancomycin) form complexes with the peptidoglycans that prevent them from binding to the transpeptidases responsible for cross-linking cell membrane structures into a rigid matrix.

A large number of antibiotics, including the aminoglycosides, macrolides, oxazolidinones, streptogramins, lincosamides, and tetracyclines, interact with bacterial ribosomes to inhibit protein synthesis. In particular, the tetracyclines and glycylcyclines bind to the 30S subunit of the ribosome, inhibit binding of aminoacyl tRNA to the A site of the ribosome, and prevent the transfer of amino acids to newly forming protein chains. Glycylcyclines are a new class of antibiotics developed to overcome resistance to tetracyclines.

Reference

1. Chopra I. Glycylcyclines: third-generation tetracycline antibiotics. *Curr Opin Pharmacol.* 2001;1:464-469.



Practice Question:

The ED reports 3 cases of cramping, abdominal pain, and diarrhea within a 24-hour period. All persons are from the same community, and onset of symptoms was within 12 to 36 hours of a picnic they all attended. The IP suspects which of the following foodborne illnesses:

- A. Salmonella
- B. Hepatitis A
- C. Staphylococcus aureus
- D. Clostridium perfringens



Α.

Bonus: typhi vs. non-typhi (enteritidis)

S. Enteritidis is the most common cause of outbreak-related acute gastro in the US

Outbreaks assoc with many things: exotic pets, most recently pre-cut melons and honey Smacks!

Foodborne Diseases Foodborne diseases Etiology and Incubation period Staph aureus 30 min to 8 hrs, ave. 2-4 hrs. Ingestion of toxin Bacillus cereus 1-6 hrs vomiting, 6-24h diarrhea Salmonella 6-72 hrs, avg. 12-36 hrs 12-96 hrs, avg. 1-3 days

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Shigella

S. dysenteriae

Campylobacter

So here is another great reference slide. The timing from ingestion to symptom onset is very important. The most important takeaway form this slide is that if the toxin is causing the infection then the onset of symptoms is within hours versus bacteria it is typically days.

up to 1 week

Clostridium perfringens 6-24 hrs, avg. 10-12 hrs

1-10 days, avg. 2-5 days

A patient is admitted with pruritic lesions on the hands, webs of fingers, wrists, the extensor surfaces of the elbows and knees, and the outer surfaces of the feet, armpits, buttocks, and waist. The most likely diagnosis is:

- A. Scarlet fever
- B. Herpes zoster
- C. Scabies
- D. Measles



C. Scabies

The key here are the webs of her fingers. These are typically extremely pruritic.

A patient who was hospitalized for 2 days calls 3 days after discharge complaining that he has developed healthcare-associated scabies due to his recent inpatient stay. The IP knows that his scabies infestation is not healthcare-associated because:

- A. Scabies is only transmitted through contaminated linens, and the IP confirmed that all linens the patient came into contact with had been properly laundered
- B. the incubation period for scabies is longer than 5 days
- C. the incubation period for scabies is shorter than 3 days
- D. Scabies is only transmitted through direct contact and none of the healthcare personnel who cared for the patient are infested



В.

Might be as short as 10d, but is typically weeks (4-6)

He would have to be exposed prior to his hospitalization (by at least 5 days)

Quick and dirty on parasites

"any organism living within or on another living creature and deriving advantage from doing so while causing disadvantage to the host"

Multicellular

Lice Scabies Myiasis (maggots) Bed bugs

Protozoa

Water or food borne
Vector borne
Animal
Sexually transmitted

Worms

Round (Intestine) Round (Tissue) Tape Fluke



Ok so what is the quick and dirty on parasitse? A parasite is any organism living within or on another living creature and deriving advantage from doing so whiel causing a disadvantage to the host. Parasites can be thought of in three ig buckets. Multicellular such as lice. Protozoa such as malaria, and wroms such as tape worms.

The causative organism of Creutzfeldt-Jakob disease is a:

- A. helminth
- B. diphtheroid
- C. spirochete
- D. prion



D.

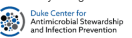
Prion Diseases

- "Transmissible neurodegenerative diseases" or "Transmissible spongiform encephalopathies (TSE)"
- Infectious protein replicates in the CNS and interrupts neuron functioning
- Rapidly progressive, ultimately fatal
- Spongiform appearance of neurologic tissues on path, no inflammatory response

Ways to acquire TSEs: Sporadic disease; latrogenic; Familial transmission (gene mutation); Ingestion of prions (e.g. bovine encephalopathies)

Transmission key features:

- Long incubation period
- Highly resistant to routine methods of disinfection/sterilization: requires prolonged cycle/increased temps and/or chemical disinfection for equipment exposed to infectious tissues
- Early recognition is KEY for the healthcare setting



<u>Humans</u>

Creutzfeldt-Jakob disease (CJD)

Kuru

Gerstmann-Straumlussler-Scheinker (GSS)

atal familial insomnia

<u>in animais:</u> Bovine spongiform encephalopathy (BSE, cattle)

Chronic wasting disease (deer, elk)

Prior diseases are rare but something you definitely should be aware of as an IP. Priors can cause transmissible spongiform encephalopathies or TSEs. The infectious protein replicates in the CNS and disrupts neuron functioning. These are rapidly progressive and ultimately fatal. You get a spongiform appearance eon fthe neurologic tissues but there is no inflammatory response. The way sin which to acquire TSEs are fro sporadic disease, iatrongenically, familial transmission or ingestion from infected cows. On the right hand side I have the infecitons in humans and in animals. The most famus is Creutzfeldt Jakob disease or CJD.

Which is TRUE about a tuberculin skin test (TST):

- A. Positive TST indicates active tuberculosis (TB) infection
- B. Negative TST rules out active TB infection
- C. Positive TST indications past exposure to TB
- D. Negative TST indicates past exposure to TB



C.

Some persons may react to the TST even though they are not infected with *M. tuberculosis*. The causes of these false-positive reactions may include, but are not limited to, the following:

Previous TB vaccination with the bacille Calmette-Guérin (BCG) vaccine Infection with nontuberculosis mycobacteria (mycobacteria other than *M. tuberculosis*)

Some persons may not react to the TST even though they are infected with *M. tuberculosis*. The reasons for these false-negative reactions may include, but are not limited to, the following:

Anergy

Recent TB infection (within the past 8 to 10 weeks)

Very young age (younger than 6 months)

The optimal time to collect a sputum specimen for acid-fast bacilli (AFB) testing to rule out TB would be:

- A. First thing in the morning
- B. After a respiratory treatment
- C. Prior to the patient going to bed
- D. Prior to a respiratory treatment



A.

Collection of early morning specimens is preferred because of the overnight accumulation of secretions

Mycobacterium tuberculosis

- Acid-fast bacilli
- Clinical features of pulmonary disease: subacute onset, cough/congestion (sputum may be bloody), weakness, fatigue, weight loss, chills, fever, night sweats
- Diagnostic testing: acid-fast smear/culture of sputum, PCR DNA probe, chest x-ray
- ■TST/PPD or IGRA is a screening test for latent disease or prior exposure



https://radiopaedia.org/cases/pulmonary-tuberculosis-29



So let's talk a bit about TB. This is an acid fast bacilli. Clinically it causes pulmonary disease with subacute onset. These folks are sick for weeks to months. The sputum is classically taught to be bloody and patietns often have systemic sympoms such as chills, fevres, and night sweats. TB is diagnosed by sputum culture, PCR DNA probe and CXR. The tuberculin skin test or IGRA is a screening tset for latent disease or prior exposure and not for active infection.

Mycobacterium tuberculosis

- Transmission: airborne by inhalation of droplet nuclei
- Prevention: negative pressure isolation room, N95 mask, direct observed therapy for all new cases
- Treatment: 4-drug therapy: isoniazid (INH), rifampin (RIF), pyrazinamide (PZA), ethambutol (EMB); others if drug-resistant (streptomycin, bedaquiline, FQ)



https://radiopaedia.org/cases/pulmonary-tuberculosis-29



A 14yo boy from rural Maryland was seen in the emergency department with fever, fatigue, chills, headache, and a large annular lesion on his left thigh. What is the most probable vector of this child's illness?

- A. tick
- B. mosquito
- C. flea
- D. louse



A.

Spirochetes: spiral shaped bacteria

Lyme (*Borrelia burgdorfori*): tick borne illness in the NE US (but expanding)

- Target lesions (erythema migrans), fever, headache, arthralgias; can also cause CNS disease and associated with Bell's palsy; cardiomyopathy
- Confusing diagnostics (serology and protein detection)
- Tick = Ixodes scapularis "black legged"
- Treatment = doxycycline, ceftriaxone for CNS







Ok so what about spirochestes? Lyme is cause by borrelia burgdorfori. It is a tick borne illness classically thought of in the NE US, but the region of infection is expanding. This classically causes target lesions called erythema migrans, fver, headache, arthralgias. Ti can also cause CNS diase and associated Bell's palsy and cardiomyopathy. Bells palsy is manifested by sudden weakness in the muscles on one half of the face. The diagnostics are a bit confusing and often ID clinicians get involved to help. The tick that carries the spirochete is the ixodes scapularis. The treatment is doxycycline, but can also be ceftriaxone for CNS involvement.

Spirochetes: spiral shaped bacteria

Syphilis (*Treponema pallidum*): "the great imitator" – sexually transmitted

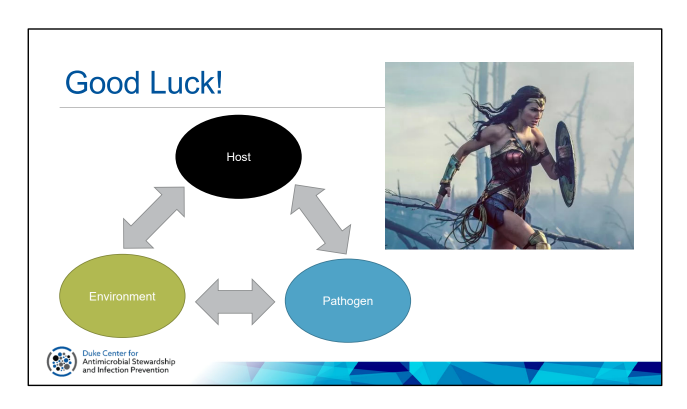
- Multiple stages of clinical disease: primary, secondary, early latent, late latent, tabes dorsalis, tertiary, ocular, congenital
- Darkfield microscopy is classical diagnostic, but never used now
- Serology IgG plus confirmatory testing for screening
- Trending of VDRL or RPR titers for response to therapy and detection of new exposures
- Treatment = penicillin



www.giantmicrobe.com



Anotehr spirochete is typhilis. Ti is called the great imitator as it can present very similar to a large variety of other diseases, which can sometimes complicate its diagnosis, especially in the later stages. It is sexually transmitted. It causes multiple stages of disease including primary, secondary, early latent, alte latent, tabes dorsalis, tertiary, ocular and congenital syphilis. Dkrkfiled microscopy is the classic diagnostic tool, but it is rarely if ever used. We typically use serology IgG plus a confirmatory test. RPR or VDRL titers are used for response to therapy and detection of new exposures. The atreatment is good olepenicillin.



That's it! We made it through! Please let me know if you have any additional questions!