***ADDIS ABABA UNIVERSITY,***

***COLLEGE OF VETERINARY MEDICINE & AGRICULTURE,***

***DEPARTMENT OF MICROBIOLOGY, IMMUNOLOGY &VETERINARY PUBLIC HEALTH ( MIVP), BISHOFTU***

***Handouts for the Course, Zoonosis(VeLT-4162), VLT Year- III, Second Semester (2019/2020)***

***Instructor: Professor Ashwani Kumar (50% share)***

**\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_**

**Impact of Zoonoses on Health and Economy**

Human health is linked to animal health and production. This link between human and animal populations, and with the surrounding environment, is particularly close in developing regions where animals provide transportation, draught power, fuel and clothing as well as proteins (meat, eggs and milk). In both developing and industrialized countries, however, this can lead to a serious risk to public health with severe economic consequences.

**The negative effects of zoonoses:**

* Loss of man-hour and man-day
* Loss of labor productivity
* Monetary loss
* Adverse effect on morale of personnel
* Reduced travel and tourism to the affected areas
* Reduced livestock and food production
* Death and destruction of affected animals
* Restriction on and reduction in international trade of animals
* Loss of country’s economy

**Zoonoses and public health**

Zoonotic diseases and human health are matter of particular concern because people, especially in developing and underdeveloped countries, are in direct close contact with domestic animals, and often not very far from wild animals. Their activities such as working with animals, in their sheds, improper disposal of waste material from animal sheds, skinning of infected animals, slaughtering of diseased animals, disposal of infected diseased animals, and poor personal hygiene practices have been one of the important risk factors.

Infectious diseases account for 29 out of the 90 major causes of human mortality and morbidity and 25% of global deaths. About 75% of the new diseases that have affected humans have been caused by pathogens originating from an animal or from products of animal origin. Many of these diseases have the potential to spread through various means over long distances and to become global problems.

Every year millions of people get sick because of foodborne zoonoses such as Salmonellosis and Campylobacteriosis which cause fever, diarrhoea, abdominal pain, malaise and nausea. Other bacterial zoonoses are anthrax, brucellosis, infection by verotoxigenic*Escherichia coli*, leptospirosis, plague, Q fever, shigellosis and tularaemia. Every year, 2.2 million people in less developed countries ( LDCs), most of them young children, die from diarrhoeal diseases, chiefly of zoonotic origin, caused by Campylobacteror Salmonella.

Cysticercosis/Taeniasis is caused by a parasite which infects swine and can cause seizures, headache and many other symptoms in humans. Other parasitic zoonoses are trematodosis, echinococcosis/hydatidosis, toxoplasmosis and trichinellosis.

In addition a number of well-known and preventable animal diseases that can be transmitted to humans (i.e. zoonoses) such as rabies, brucellosis, leishmaniasis and echinococcosis continue to occur in many countries especially in the developing world where they mostly affect the poorest segment of the human population. They cause a serious amount of deaths and millions of people affected every year.

Almost all persons infected by rabid animals will die if not treated. Dogs are responsible for most human deaths.

Other viral zoonoses are avian influenza, Crimean-Congo hemorrhagic fever, Ebola and Rift Valley fever. Recent emerging zoonosis is infection due to Covid-19.

**Economic impact of zoonoses**

* Besides apart from mortality and morbidity, zoonoses are responsible for huge economic losses, particularly in dairy animals, meat, milk and other foods and products of animal origin.
* Many zoonoses cause disabling diseases in humans. A sick or disabled person cannot work as much or as well as a healthy person, hence will not earn enough money to buy proper food or healthy livestock. Unhealthy livestock produce less food. The result is less food to eat and less food to sell - less income again and not enough money.
* All major zoonotic diseases prevent the efficient production of food of animal origin, particularly of much-needed proteins, and create obstacles to international trade in animals and animal products. They are thus an impediment to overall socioeconomic development.
* The World Bank has estimated losses of USD 200 billion (direct and indirect) caused by zoonoses during 2000 and 2010. According to an OECD report (2011), a severe avian influenza pandemic could cause the death of 70 million people and decrease global GBP by 4.8%.
* The OIE estimates that morbidity and mortality due to animal diseases cause the loss of at least 20% of livestock production globally. This represents at least 60 million tones of meat and 150 million tons of milk with a value of approximately USD 300 billion per year.

**Social impact**

The social impacts can be expressed as indirect health consequences or behavioral changes, changes in societal values and changes in social standing and can be felt at the individual, family or community level.

Poverty is an important factor governing the risk of zoonotic diseases in rural and urban areas. Poverty can increase exposure to such infections and reduces the chances of getting rid of diseases. Poor communities are often trapped in a never-ending cycle known as "the devil's circle of poverty".

Poor people are more likely to suffer from zoonotic diseases for several reasons:

* Poor education: poor knowledge of disease existence, motivation to prevent them.
* Poor sanitary conditions: more chances of getting infection.
* They buy cheaper animals that are often less healthy animals. Cheaper meat is never the healthiest meat and often has not been inspected, therefore, can make the people ill.
* Poor veterinary and public health services: lack of veterinary services worsens the situation in rural communities for those who cannot pay for private services.
* Poor people are often less well-nourished and less healthy and are more susceptible to infectious disease in general and zoonoses in particular.
* Lack of money to set up a good health system for both animals and humans.

Close contact between domestic animals and humans is a characteristic of rural settings. Animals kept for food production in these areas. It is this close association that makes zoonotic diseases so vitally important in rural areas.

**Social customs:** Dogs and cats are kept as pets in both developing and developed countries. Often these animals are kept unrestricted and unsupervised, without adequate care and vaccinations. Rabies is the most important zoonosis transmitted especially though dog bites. Streets and parks in many cities in developed countries are polluted with dog faeces, posing major problems for city councils. Apart from dogs and cats, inexperienced owners often also keep exotic animals. These animals need to be kept under stringent hygienic conditions to prevent transmission of zoonotic diseases to their handlers (such as salmonella and parasitic diseases).

Socio-economic, cultural and religious factors have been observed to play an important role in the transmission of zoonotic diseases. The hydatid disease in man is more prevalent in Turkana people of North-western Kenya, where as per their religious customs the human dead bodies are exposed to hyenas and dog, thus perpetuating the transmission of the disease. Similarly, the Muslim belief of the unclean less of dogs have been responsible for the observed reduced incidence of the disease in Muslim Arabs as compared to Christian Arabs in Lebanon.

The disease is of much greater incidence in certain occupational workers like shoemakers and shoe-repairs in Lebanon, mainly related to their practices of dipping hides of animals in a decoction of dog faeces for preparing leather.

It has also been observed that certain modes of recreation in man (sea side sports, camping, tourism, mountaineering, hunting, fishing etc.) have resulted in bringing man in close contact with hydatid endemic foci and increasing the chances of his acquisition of the disease.

In the community setting in some African countries, new infections of Ebola virus were related to the ministration of funeral rites, which involve ritual cleansing of the cadaver and removal of hair, finger nails, toe nails and clothing before burial. People visiting or taking care of infected persons in their homes or in hospitals also risk being exposed to Ebola infections.

**Impact on productivity and animal health**

Among the factors influencing the productivity and profitability of livestock, animal diseases including zoonoses deserve special attention because they diminish the capacity of the animal to achieve its inherent potential level of production, for any given feeding and management regimen.

Animal diseases lead to significant losses and decrease productivity

Animal diseases kill around 18% of the livestock population in low-income

countries (World Organization for Animal Health [OIE]).

In LDCs (least developed countries), mortality rates in traditional village livestock systems average 20% to 22% for calves, 7% for cattle aged over 12 months, 22% to 24% for lambs and kids, 15% for adult sheep and goats, 40% to 50% for piglets and 50% for chickens aged 0 to 6 months. Cattle farmers lose as many animals as they sell.

Not only do animal diseases cause deaths, they also lead to production losses and indirect losses, which are generally far greater. Very few studies provide information on total losses in low-income countries caused by animal diseases. However,the information available for transmissible diseases confirms that losses are heavy.For instance,

An epizootic of Rift Valley fever in Somalia in the early 2000s had little impact on the country’s livestock but, when Saudi Arabia banned Somalian livestock imports to protect its human and animal population, it drove down livestock prices on local markets, causing Somalian livestock producers to lose US$47 to US$55 million.

***RICKETTSIA & ZOONOSES***

**General Features**

The rickettsia are bacteria which are obligate intracellular parasites. They are considered a separate group of bacteria because they have the common feature of being spread by arthropod vectors (lice, fleas, mites and ticks). The cells are extremely small (0.25 u in diameter) rod-shaped, coccoid and often pleomorphic microorganisms. *Rickettsia* cannot live in artificial nutrient environments and are grown either in tissue or embryo cultures (typically, chicken embryos are used).

The family Rickettsiaceae is taxonomically divided into three genera:

* *Rickettsia* (11 species)
* *Coxiella* (1 species)
* *Ehrlichia* (2 species)
* *Baartonella* (3 species)

***A. Rickettsia*** species are carried by many ticks, fleas, and lice, and cause diseases in humans. Based on serology, the *Rickettsiae are grouped* as below:

* Spotted fever Group
* Typhus Group
* Scrub typhus Group

All three of these contain human pathogens.

1. Spotted Fever Group

• Rocky Mountain spotted fever cuased by *Rickettsia rickettsii*

• Rickettsial pox: *Rickettsia akari*

• Canadian typhus: *Rickettsia canada*

• Mediterranean spotted fever:*Rickettsia conorii*

• Siberian tick typhus: *Rickettsia siberica*

• Queensland tick typhus: *Rickettsia australis*

2.Typhus Group

• Murine/Endemic typhus: Fleaborne typhus): Caused by *Rickettsia mooseri (typhi)*

• Epidemic typhus (Brill- Zinsser disease ) and Louse-borne typhus):Caused by

*Rickettsia prowazekii*

3.Scrub typhus (or Chigger fever): *Rickettsia tsutsugamushi*

**B. Genus Coxiella:** *Coxiella burnetii causes Q-fever.*

**Q Fever**

Q fever is an illness caused by the bacterium *Coxiella burnetii.* Q fever is spread to humans from infected animals. The bacteria survive for long periods in the environment as they are resistant to heat, drying and many disinfectants.Q fever is usually an acute (immediate) infection but it can sometimes lead to a chronic (long-term) illness. Acute Q fever can cause a severe flu-like illness that is sometimes associated with hepatitis (inflammation of the liver) and pneumonia. Chronic Q fever most commonly results in inflammation of the heart (endocarditis) and people who already have heart valve disease are at increased risk.

**Symptoms**

Many infected people have no or few symptoms. People who do become sick often have a severe flu-like illness. Symptoms begin about 2-3 weeks after exposure and typically include:

* high fevers and chills
* severe sweats
* severe headaches, often behind the eyes
* muscle and joint pains
* extreme fatigue (tiredness)

If untreated, symptoms can last from 2-6 weeks. Most people make a full recovery and become immune to repeat infections. Occasionally, people develop chronic infections which affect the heart (endocarditis) or the liver (hepatitis). Some people develop chronic fatigue (post-Q fever fatigue syndrome), which can last for many years after the initial infection. Symptoms of chronic Q fever may occur up to two years after the initial infection.

**Spread**

People usually get infected by breathing in infected aerosols or dust when working with infected animals, animal tissues, or animal products. The main carriers of the disease are farm animals such as cattle, sheep and goats but other animals such as kangaroos, bandicoots, domestic pets such as dogs and cats can also be infected. Pigs are not known to carry the disease.

Infected animals often have no symptoms and can shed the bacteria into their urine, faeces or milk. High concentrations of the bacteria are found in the placenta (birth by-products). Q fever can be contracted by inhaling dust from wool, hides, straw or grass that contains the Q fever bacteria. Common activities where people are exposed include birthing calves and shearing. There is also a potential risk of contracting Q fever by ingestion of unpasteurised milk from an infected animal.

Spread of Q fever from person to person has been reported but is extremely rare. Contaminated work clothing may be a source of infection.

**People at at risk**

* abattoir and meat workers (including contractors who visit these facilities)
* farmers and shearers
* stockyard workers and animal transporters
* veterinarians, veterinary assistants and veterinary students
* agriculture college staff and students (working with high-risk animals)
* laboratory workers (working with the bacteria or with high-risk veterinary specimens)

Horticulturists or gardeners in environments where dust, potentially contaminated by animal urine, faeces or birth products, is aerosolised (e.g. lawn mowing) may also be at risk.  
People commencing work in or visiting these industries are at high risk of contracting the disease. Some long-term workers in these industries become immune to the disease without becoming sick.

**Prevention**

A vaccine (Q-Vax®) is available to protect people against Q fever. Vaccination is recommended for all people who are working in, or intend to work in, a high-risk occupation Workplaces at risk should have a vaccination program.

**People must be screened and tested before they are vaccinated against Q fever.**

People who work with animals or materials that may carry the Q fever bacteria should use appropriate protective equipment and be aware of the steps required to stop the spread of the bacteria. The risk of Q fever can be further reduced by:

* washing the hands and arms thoroughly in soapy water after any contact with animals
* washing animal urine, faeces, blood and other body fluids from the work site and equipment, and disinfecting equipment and surfaces where practicable
* properly disposing of animal tissues including birthing products
* minimising dust in slaughter and animal housing areas
* keeping yard facilities for sheep and cattle well away from domestic living areas
* removing clothing that may carry the bacteria before returning to the home environment
* wearing a mask when mowing lawn or gardening in areas where there are livestock or native animals.  
  People who are unimmunised should not be allowed to visit high-risk work areas such as abattoirs.

**Diagnosis**

The initial suspicion of a Q fever diagnosis is based on clinical symptoms and signs. Blood tests are required to confirm the diagnosis with repeated testing after two weeks.

**Treatment**

Q fever is treated with antibiotics, usually in the tablet form. A cardiac assessment, which may include echocardiography, is required to assess whether there are underlying abnormalities of the heart valves which increase the risk of developing chronic Q fever endocarditis. Chronic Q fever infection requires prolonged treatment with antibiotics.

**Ringworm (Dermatophytosis)**

The dermatophytes are a group of closely related fungi that have the capacity to invade keratinized tissue (skin, hair, and nails) of humans and other animals to produce an infection, dermatophytosis, commonly referred to as ringworm. Infection is generally cutaneous and restricted to the nonliving confide layers because of the inability of the fungi to penetrate the deeper tissues or organs of immune-competent hosts & feed on [keratin](http://en.wikipedia.org/wiki/Keratin), the material found in the outer layer of skin, hair, and nails. Reactions to a dermatophyte infection may range from mild to severe as a consequence of the host’s reactions to the metabolic products of the fungus, the virulence of the infecting strain or species, the anatomic location of the infection, and local environmental factors.

**Etiology**

The etiologic agents of the dermatophytoses are classified as following in three genera,

***Epidermophyton* spp:-**The type species is *Epidermophyton floccosum*.

***Microsporum* spp:-**The type species is *Microsporum audouinii*.

***Trichophyton* spp:-**The type species is *Trichophyton tonsurans*.

**Transmission**

These fungi live for an extended period as spores in soil. Ringworm is highly contagious and can spread multiple ways. Humans and animals can contract ring-worm after direct contact with this soil. The infection can also spread through contact with infected animals or humans. The infection is commonly spread among children and by sharing items that may not be clean.

Ringworm is an example of a zoonotic disease. Cats are among the most commonly affected animal. Dogs, cows, goats, pigs and horses can spread ringworm to humans and other animals via direct contact or contact with objects the infected animal has touched such as bedding, grooming articles, saddles, furniture, carpeting etc.

**Risk factors**

As described previously, it is possible to acquire ringworm from a variety of places and circumstances. The greatest risk factor is coming in contact with an affected individual or a contaminated surface. Areas of frequent sweating as well as skin fold such as those in the groin or between the toes are commonly affected. Also from contaminated items as toilet articles, clothing, pool surfaces, showers and locker rooms, transmission is favorable.

**Symptoms**

The medical term for ringworm is *tinea*. Depending upon the site of infection, tinea has been named differently with different symptoms.

**Tinea corporis**: When fungus affects the skin of the body, it often produces the round spots of classic ringworm. Sometimes, these spots have an "active" outer border as they slowly grow and advance. Sometimes, scaling, crusting, raised areas, or even blister-like lesions can appear, particularly in the active border.

**Tinea cruris**: Tinea of the groin ("jock itch") tends to have a reddish-brown color and extends from the folds of the groin down onto one or both thighs.

**Tinea faciei (faciale)**: ringworm on the face except in the area of the beard. On the face, ringworm is rarely ring shaped. Characteristically, it causes red scaly patches with indistinct edges.

**Tinea manus**: ringworm involving the hands, particularly the palms and the spaces between the fingers. It typically causes thickening (hyperkeratosis) of these areas, often on only one hand.

**Tinea pedis**: Athlet’s foot may cause scaling and inflammation with itching and burning irritation in the toe webs, especially the one between the fourth and fifth toes. Another common form of tinea pedis produces a thickening or scaling of the skin on the heels and soles. This is sometimes referred to as the "moccasin distribution." Occasionally, tinea causes blisters between the toes or on the sole. Aside from athlete's foot, tinea pedis is known as tinea of the foot or, more loosely, fungal infection of the feet. Tinea pedis is an extremely common skin disorder. It is the most common and perhaps the most persistent of the fungal (tinea) infections. It is rare before adolescence. It may occur in association with other fungal skin infections such as tinea cruris (jock itch).

**Tinea barbae**: Ringworm of the bearded area of the face and neck, with swelling and marked crusting, is often accompanied by itching, sometimes causing the hair to break off. In the days when men went to the barber daily for a shave, tinea barbae was called barber's itch.

**Tinea capitis**: Ringworm of the scalp commonly affects children, mostly in late childhood or adolescence. This condition may spread in schools. Tinea capitis appears as scalp scaling that is associated with bald spots (in contrast to seborrhea or dandruf, for instance, which do not cause hair loss).

**Tinea unguium**: Finally, fungal infection can make the fingernails and, more often, the toenails yellow, thick, and crumbly. This is referred to as fungal nails or onychomycosis.

**Epidemiology**

These pathogenic fungi are found worldwide, and all domestic animals are susceptible. In developed countries, the greatest economic and human health consequences come from dermatophytosis of domestic cats and cattle. A few dermatophyte species are soil inhabitants (geophilic), and cause disease in animals that are exposed while digging or rooting. Other species are host-adapted to people (anthropophilic).

The zoophilic species are transmitted primarily by contact with infected individuals and contaminated fomites such as furniture, grooming tools, or tack.

The most important animal pathogens worldwide are *M. canis*(cat, dog), T. equinum *& M. equinum*(horse), *T. verrucosum*(cattle, sheep, dromedary, *M. gallinae*(fowl).These species are zoonotic, especially M canis infections of domestic cats and T. verrucosum of cattle and lambs.

**Diagnosis**

Often, the diagnosis of ringworm is obvious from its location and appearance. Circular bare patches on the skin suggest the diagnosis but no lesion is truly specific to the fungus (similar patches may result from allergies, sarcoptic mange, and other conditions)

Skin scrapings for microscopic examination and a culture of the affected skin can establish the diagnosis of ringworm. If the diagnosis is unclear, a potassium hydroxide (KOH) preparation of a skin scraping can be reviewed under the microscope to confirm the diagnosis of a fungus as given below.

**Direct microscopic examination** although false negative in 5 to 15% of cases in ordinary practice, is a highly efficient screening technique, of hairs or skin scrapings may enhance clinical suspicion by demonstrating characteristic hyphae or arthrospores in the specimen. The technique is more useful in diagnosing dermatophytosis in large animals than in small animals. Hairs (preferably white ones) and scrapings from the periphery of lesions are examined for fungal elements in a wet preparation of 20% KOH that has been gently warmed or incubated in a humidity chamber overnight. And examined under X400 magnification for fungal structures.

**Culture:-**Fungal culture is the most accurate means of diagnosis. Dermatophyte test medium (DTM) may be used in a clinical setting. Incubation at room temperature is sufficient except when culturing for T verrucosum from food and fiber animals. Dermatophyte growth is usually apparent within 3–7 days but may require up to 3 weaks on any type of DTM. Dermatophytes growing on DTM cause the medium to change to red at the time of first visible colony formation. Dermatophyte fungi have white to buff-colored, fluffy to granular mycelia. Saprophytic contaminant colonies are white or pigmented and almost never produce an initial color change on DTM.

Definitive diagnosis and species identification require removal of hyphae and macroconidia from the surface of the colony with acetate tape and microscopic examination with lactophenol cotton blue stain

Many typical isolates of common dermatophytes can be identified directly from primary isolation media, particularly, Sabouraud glucose agar and potato glucose or potato flake agar.Identification characters include colony pigmentation,texture, and growth rate and distinctive morphological structures,such as microconidia, macroconidia, spirals, pectinate branches, pedicels, and nodular organs .

**Treatment**

Ringworm can be treated topically (with external applications) or systemically (for example, with oral medications):

**Topical treatment**: When fungus affects the skin of the body or the groin, many antifungal creams can clear the condition in around two weeks. Examples of such preparations include those that contain [clotrimazole](http://www.medicinenet.com/clotrimazole/article.htm), [miconazole](http://www.medicinenet.com/miconazole/article.htm), [ketoconazole](http://www.medicinenet.com/ketoconazole/article.htm), econazole, naftifine, and [terbinafine](http://www.medicinenet.com/terbinafine/article.htm). It is usually necessary to use topical medications for at least two weeks.

**Systemic treatment**: Some fungal infections do not respond well to external applications. Examples include scalp fungus and fungus of the nails. To penetrate these areas and for particularly severe or extensive disease, oral medications can be used.

For a long time, the only effective antifungal tablet was [griseofulvin](http://www.medicinenet.com/griseofulvin-oral_tablet/article.htm). Now, other agents are available that are both safer and more effective. These include terbinafine, [itraconazole](http://www.medicinenet.com/itraconazole/article.htm) and [fluconazole](http://www.medicinenet.com/fluconazole/article.htm). Oral medications are usually given for a three-month course.

**Prevention and control**

[Fungi](http://en.wikipedia.org/wiki/Fungi) thrive in moist, warm areas, such as locker rooms, [tanning beds](http://en.wikipedia.org/wiki/Tanning_bed), swimming pools and in [skin folds](http://en.wikipedia.org/wiki/Skin_fold); accordingly, those that cause dermatophytosis may be spread by sharing sporting equipment, towels, or clothing.

* Avoid sharing clothing, sports equipment, towels, or sheets.
* Washing clothes in hot water with [fungicidal](http://en.wikipedia.org/wiki/Fungicidal) soap after suspected exposure to ringworm.
* Avoid walking barefoot; instead wear appropriate protective shoes in locker rooms and sandals at the beach.
* After being exposed to places where the potential of being infected is great, one should wash with an [antibacterial](http://en.wikipedia.org/wiki/Antibacterial) and [anti-fungal](http://en.wikipedia.org/wiki/Anti-fungal) soap or one that contains [tea tree oil](http://en.wikipedia.org/wiki/Tea_tree_oil), which contains [terpinen-4-ol](http://en.wikipedia.org/wiki/Terpinen-4-ol)
* Avoid touching pets with bald spots as they are often carriers of the fungus.

Control may be accomplished by educating infected individuals not to expose others by walking barefoot near swimming pools, locker rooms, and public showers and by not sharing footgear. Frequent cleaning and disinfecting floors of public baths, swimming pools, etc.

**ASPERGILLOSIS**

The fungus, Aspergillus is responsible for mycosis and mycotoxicosis. Mycosis is the development of fungus in or on the body of host. These fungi grow in different types of foods and produce mycotoxins.

* **Myco-toxicosis** is the ingestion of food stuffs contaminated with myco-toxins resulting into in-toxicosis.
* **Mycotoxins** are toxic metabolites that are produced by microscopic filamentous fungi.

**Aspergillus Mycosis**

**Infectious Agent** :*Aspergillus fumigatus* , Aspergillus parasiticus

**Host:** The fungus is ubiquitous. People with weakened immune systems or lung diseases are at a higher risk. Animal hosts in rehabilitation are generally wild birds such as raptors or waterfowl.

**Transmission:** *Aspergillus*, the mold that causes aspergillosis, is very common both indoors and outdoors, so most people breathe in fungal spores every day. While under stress in a captive facility, rehab animals are more likely to develop the fungal disease. They in turn shed the spores of the fungus and the workers may inhale them.

**Symptoms:** .

In birds, respiratory signs. Most healthy people have no trouble resisting infection. However, this is not true for anyone who has been debilitated by illness, other diseases, or has been on long term antibiotic, antimetabolite, or corticosteroid therapy.

In humans different types of aspergillosis are prevalent with different symptoms. The allergic bronchopulmonary aspergillosis (ABPA) are similar to asthma symptoms; symptoms of allergic *Aspergillus* sinusitis include:Stuffiness, Runny nose, headache, reduced ability to smell; aspergilloma (“fungus ball”) include cough, coughing up blood, shortness of breath, chronic pulmonary aspergillosis is manifested as weight loss, cough, fatigue, shortness of breath while invasive aspergillosis is manifested by fever, chest pain,cough,coughing up blood,chortness of breath,

**Diagnosis**

* Medical history, risk factors, symptoms, physical examinations, and lab tests to make diagnosing of aspergillosis.
* Imaging tests such as a chest x-ray or a CT scan of lungs or other parts of body depending on the location of the suspected infection.
* Tissue biopsy, in which a small sample of affected tissue is analyzed in a laboratory for evidence of *Aspergillus* under a microscope or in a fungal culture.
* A blood test can help diagnose invasive aspergillosis early in people who have severely weakened immune systems.

**Prevention**

* Good hygiene and good ventilation. At least 12 air exchanges per hour are recommended in any room where susceptible birds are housed.
* Waterfowl are not to be housed on wood shavings because the fungus will thrive in them when wet.
* Moldy grains and food stuffs are not fed or stored.
* Necropsies are performed with masks as are treatments on suspect patients.
* Spray necropsy birds down with a disinfectant to matt feathers and reduce aerosolized lint and debris.

**Aspergillus Mycotoxicosis**

**Three important mycotoxins are produced by the genus Asppergillus, Viz.,** Aflatoxin(*Aspergillus flavus; A. parasiticus),*  Ochratoxin(*A.ochraceus) and*  Sterigmatocystin (*Aspergillus nidulans, A. versicolor*) resulting in mycotoxicosis. Among these aflatoxicosis due to Aflatoxin occurs widely and is described as below.

**Aspergillosis**

**Types of foods involved:**

* Peanuts, barley, coconut, cotton seeds: invariabily mycotoxins are present.
* Corn, millet, rice, oat, sorghum, beans, dry fruits: mycotoins have been isolated.
* Soyabean: very rarely involved. Natural toxin production is rare. Presence of calcium, magnesium, boron in soyabeans inhibit toxin production by fungi.
* Peanuts are used for animals in the form of peanut cakes or pea nut butter. Oil is extracted from peanut and the remaining is concentrated as cakes to be fed to animals. Oil may have some toxin, but in cakes, the toxins are concentrated.
* Fortifying the food with peanuts cakes means fortifying the feed with toxin.

**Factors favoring toxin production by moulds.**

* Substrate
* Fungal strain variation
* Genetic susceptibility of host plant or commodity or composition of commodity:
* Structural integrity of commodity
* Temperature:Wide range of temperature for growth of fungi. 20-30oC good for growth but may not be optimum for toxin production.

**Major types of aflatoxins and their metabolites**

At least 14 different types of aflatoxin are produced in nature. Aflatoxin B1 is considered the most toxic, Aflatoxin G1 and G2 are produced exclusively by *A. parasiticus*.

Aflatoxins M1, M2 were originally discovered in the milk of cows that fed on moldy grain. These compounds are products of a conversion process in the animal's liver.

**Standard for allowable contamination of commodities destined for human and animal consumption.**

* Human foods are allowed 4-30 ppb aflatoxin, depending on the country involved.
* In contrast, grains for animal feed in the United States are allowed 300 ppb aflatoxin , because this concentration not only provides protection against acute aflatoxicosis but also is low enough to allow most of the grain produced to be traded.

**Illness due to aflatoxins**

**Humans**

High-level aflatoxin exposure produces an acute hepatic necrosis, resulting later in cirrhosis or carcinoma of the liver. Acute hepatic failure is made manifest by hemorrhage, edema, alteration in digestion, changes to the absorption and/or metabolism of nutrients, and mental changes and/or coma.

Chronic, subclinical exposure does not lead to symptoms as dramatic as acute aflatoxicosis. Children, however, are particularly affected by aflatoxin exposure, which leads to stunted growth and delayed development. Chronic exposure also leads to a high risk of developing liver cancer.

A strong synergy is observed between aflatoxin and hepatitis B virus (HBV) and hepatitis C virus (HCV) agents for liver cancer.

**Animals**

Turkeys are extremely susceptible to aflatoxicosis. Aflatoxin has potential to lead to liver disease in dogs; however, not all dogs exposed to aflatoxin will develop liver disease. Toxic level in dog food is 100–300 ppb and requires continuous exposure/consumption for a few weeks to months to develop aflatoxicosis.

.

**Prevention**

There is no specific antidote for aflatoxicosis.

Symptomatic and supportive care tailored to the severity of the liver disease may include intravenous fluids with dextrose, active vitamin K, B vitamins, and a restricted but high-quality protein diet with adequate carbohydrate content

**Possible intervention strategies**

**During Production**

* The management can be used to minimize contamination, and the practice of inoculating the fields with non-aflatoxigenic strains of fungi may shortly be a new tool in the battle to prevent economic loss.
* Insect damage in the field can be controlled by pesticides or by cultural practices;
* Harvesting is usually done without machinery, and drying should be carried out very efficiently.

**Storage**

* To preserve quality in storage, it is necessary to prevent biological activity through adequate drying (<10% moisture), elimination of insect activity that can increase moisture content through condensation of moisture resulting from respiration, low temperatures, and inert atmospheres.

**Processing**

Three main approaches exist: dilution, decontamination, and separation.

* Dilution: the easiest means of satisfying the requirement is to mix grain low in aflatoxin with grain exceeding the regulated limits.
* Decontamination: Treatment with ammonia, alkaline substances , and ozone can denature aflatoxins, but whether this change is permanent is not clear.
* Separation: separating contaminated grain from the bulk. This approach depends on the heavy contamination of only a small fraction of the seeds, so that removing those leaves a much lower overall contamination. A major portion (80%) of the toxin is often associated with the smaller and shriveled seed, and thus screening can lower the overall concentration in the bulk. Further removal of aflatoxin-contaminated seeds may be achieved by color sorting, which, in the case of peanuts, is most effective when the seeds are blanched.

**TAENIASIS:**

Parasitic infection of man due to a large Tapeworm that lives in small intestine of man. It is an obligatory parasite of man and thus an example of obligatory cyclozoonoses. It is a true zoonosis which means the life cycle of this parasite is not completed without man. Under natural transmission cycle, there is no other definitive host

**Causative Agent**

*Taenia saginata:* The adult worm which lives in small intestine of man is white and semitransparent; measuring 5-12 meters.

*Taenia solium:* The adult worm which lives in small intestine of man and is 2-6 meters.

Larval stage of *Teania saginata is Cysticercus bovis while of Teania solium*: *C. cellulosae*

**Types of Cysticercosis:** Infection due to larval stage ios called as Cysticercosis

* Bovine Cysticercosis
* Porcine Cysticercosis
* Human Cysticercosis
* *Cysticercus bovis* mainly occurs in cattle and found in the masseter muscles, shoulder muscles, heart, tongue, diaphragm, esophagus, adipose tissue, liver, lungs and lymph nodes.
* *Cysticercus cellulosae*, mainly occurs in pigs and commonly found found in the masseter muscles, shoulder muscles, heart, tongue, diaphragm, esophagus, adipose tissue, liver, lungs and lymph nodes. In the tongue, skeletal muscles and sometimes in organs.

**Shape of cysts**

* *Cysticercus* is round or oval in shape and when fully developed consists of scolex invaginated into small fluid filled vesicle
* Cysts may be viable or dead
* Dead degenerated or calcified cysticerci: clearly form identifiable white spots and have fibrotic lesions,
* Viable cysticerci : Pinkish-red in color.
* True viability of the cyst can be ascertained by keeping the cysts in bile of cattle overnight. Viable cysts evaginate while the dead ones remain intact.

**Porcine cysticercosis**

Cysticercus cellulosae: Mainly occurs in pigs

Semitransparent, opalescent white, and elongate oval in shape and may reach a length of 0.6 to 1.8 cm

Common Sites: Tongue, Heart, diaphragm, internal masseter, neck, shoulder, intercostals and abdominal muscles.

Occasionally: Liver, lungs, kidney, eye and brain

**Meat having cysticerci**

Pork having cysticerci (*C. cellulosae)is called as measly pork.*

Beef having cysticerci *(C.bovis) is called as* Measly Beef

**Human cysticercosis**

Due to autoinfection or poor personal and environmental hygiene, man may also be infected with the eggs of *Taenia solium*

*Eggs* develop into larvae which may migrate even to brain and produces a condition called as neurocyticercosis (NCC) which is worldwide prevalent

**Host Range**

*Definitive host: T. saginata* & *T. solium* live exclusively in small intestine of man.

Intermediate host:

*T. saginata* mainly cattle, (deer, reindeer, llama, buffalo, giraffe and antelope)

*T. solium*: Mainly pigs.

(Rats, cats, dogs, sheep, cattle, deer, monkeys and man)

**Clinical features of the disease in man**

* Due to adult parasite (Human Taeniasis)
* Due to larval stage of *T.solium(Human Cysticercosis)*

**Clinical manifestation of infection with the adult worm** are frequently asymptomatic but may lead to nervousness, insomnia, anorexia, loss of weight, abdominal pain and digestive disturbances. The mobile gravid segments may make their way to unusual sites such as appendix, uterus or biliary tract and may cause serious disorders.

**Clinical manifestation of infection with the larval stage:** ***Due to larvae of T. solium***

**Somatic disease**

* Cysts in the muscles cause myositis, fever, muscle swelling and later to atrophy and fibrosis
* Ophthalmic Cysticercosis : visual difficulties, retinal edema, hemorrhage, a decreased vision or even a visual loss
* Subcutaneous cysts :formion of firm, mobile nodules, which are sometimes painful.

4.Neurocysticercosis:

*Cysts in brain:*

* 60% of the patients are having these cysts in brain
* headaches, nausea, vomiting, lethargy and altered mental status.
* Psychic symptoms, including epileptic seizures
* The cysts can persist in brain from 2-10 years.

*Cysts in meninges*

**Clinical features in animals**

There are no specific features of the infection in animals and the disease is diagnosed during post mortem examination.

**In cattle and pigs**

No specific features of the infection and the disease is diagnosed during slaughter or at post-mortem.

However, in cattle, heavy infestation by the larvae may cause myocarditis or heart failure.

**Epidemiology**

Man is universally susceptible. The mode of transmission for T. saginata is by ingestion of raw or inadequately cooked beef containing the cysticerci. For T solium i) by ingestion of raw or inadequately cooked pork containing the infective larvae, or ii) by direct transfer of eggs in faeces of a person harbouring an adult worm to his own or another’s mouth, or indirectly through ingestion of food or water contaminated with eggs, resulting in somatic cysticercosis. Man can also be infected with T.solium eggs from contaminated vegetables.

T. saginata is not directly transmitted from man to man but T. solium may be; eggs of both species are transmitted in the environment as long as man harbours the worm in the intestine, sometimes 30-40 years; eggs may remain viable for months.

Besides contamination by man, birds especially sea birds and corpophagus bettles may disseminate the eggs.

A person can have mixed infection with T.saginata and T.solium.

**Diagnostic procedures in animals**

**1.Routine meat inspection:**

* Two deep cuts in the external and one deep cut in the internal muscles of mastication.
* The cut surfaces of the muscle and the tongue are inspected visually.
* The pericardial surface of the heart is inspected, then the heart muscle incised lengthwise to open the ventricles and to cut through the intraventricular septum.
* When one or more cysts are found there is a requirement for further cuts with specific reference to predilection sites e.g. diaphragm and inspection of offal.

**Judgment**

***Heavy*” *infestation***

* Lesions in two of the usual inspection sites i.e. masseter muscles, tongue, oesophagus, heart, diaphragm or exposed musculature .
* Two sites during incisions into the shoulder and into the rounds.

***Carcass and viscera of heavily infested animals are condemned***

**In moderate or light infestation small number of cysticerci including dead or degenerated cysts are present.**

***Localized infestation: there is a requirement to store the carcass at a temperature:***

* ***Not exceeding -7°C for not less than 21 days or***
* ***At a temperature not exceeding -10°C for not less than 14 days before release for human consumption.***

***the carcass is conditionally passed***

**2. Other methods of diagnosis in animals**

Immunodiagnostic methods like Indirect-haemagglutination test; indirect immune-fluorescence;

Skin reaction; Radioimmunoassay; Complement fixation test; ELISA; PCR

**Diagnostic procedures in man**

* Demonstration of *T.saginata* and *T.solium* eggs (oval, 30ux20u brown in color) and gravid segments in faeces.
* Radiologically, the cestodes may sometimes be demonstrated in the intestine.

**Treatment**

There is no specific chemotherapy for human cysticercosis; surgical excision is the only satisfactory treatment. For adult parasites, niclosamide, quinacrine hydrochloride are effective.

Prrrraziquintal and benzimidazole derivative (albendazole) have been found fairly promising particularly in cattle.

**Prevention and control**

**1.**Massive chemotherapy of infected individuals,

* Niclosamide,
* Quinacrine hydrochloride ,
* Praziquantel
* Traditional herbal remedy called ‘Kosso’.

2.Improving sanitation

3.Educating people

4.Avoid use of sewage effluent for pasture irrigation without adequate treatment.

5.Processing of meat

* Freezing
* <-5oC for >360 hr, i.e. 15 days/
* <-10oC for >216 hr, i.e. 9 days /
* <-15oC for >144 hr, i.e. 6 days.
* Heating
* >56oC core temperature >1 sec,
* Irradiation (100Krad death, 40Krad inhibition for development),
* Pickling meat in 25% salt solution for 5 days

Preventive measures in animals

* Restrict the access of the cattle to surface drinking water and by supplying them with fresh water
* Avoid the access of pigs to latrine or to human faeces.
* Competent meat inspection must be made compulsory
* Anthelmentic treatment of infected animals
* Premises disinfection

For human cysticercosis

* Surgical treatment or chemotherapeutic drugs or both should be applied.
* [Albendazole](http://en.wikipedia.org/wiki/Albendazole) is preferable over [praziquantel](http://en.wikipedia.org/wiki/Praziquantel) due to its lower cost.

Corticosteroids and anticonvulsants do not reduce CSF and brain drug levels.

* Surgical treatment includes direct excision of ventricular cysts, shunting procedures, and removal of cysts via endoscopy

**ECHINOCOCCOSIS**

Caused by a parasitic tapeworm: Echinococcus

Species causing human infection:

*E. granulosus*

*E. multilocularis*

*E.oligarthrus*

*E. vogelli*

The disease in man is due to its larval stage, hydatid cyst, which means cyst with water like fluid and thus the name, hydatidosis. It has following types:

**Classic hydatid disease**

* Caused by *E.granulosus.*
* This species is adapted to dogs and a variety of domestic and sylvatic animal intermediate hosts.
* A major public health and economic problem.

**Alveolar hydatid disease by *E.multilocularis*.**

* Final and intermediate hosts are foxes and rodents respectively.
* Potential exposure to man is not great.
* However, if it occurs, it is one of the most lethal parasitic infections to occur in humans.

**Polycystic form of hydatid disease by *E.vogeli.***

Life cycles of *E. oligarthus* and *E.vogeli* are limited to sylvatic animals. Limited to central and South America

**Hydatid cyst**

* Cyst is filled with fluid.
* Cyst consists of an inner germinative layer of cells supported externally by laminated membrane of variable thickness. Surrounding the cyst, there is another layer due to host reaction.
* Small secondary cyst, called brood capsules, bud internally from germinal layer which produces multiple protoscolices.
* A protoscolex is a scolex with the rostellum and suckers.

**Types of hydatid cysts**

* Sterile hydatid cysts: Brood capsules and protoscolices in the cysts are not formed.
* Fertile hydatid cysts:Cysts with brood capsules and protoscolices.
* Unilocular cysts: Cystic stage E. granulosus and most common in food animals
* Multilocular cysts : Cystic stage of E. multilocularis. Primary cyst gives to others by continuous exogenous budding, to produce a larval mass made of hundreds of contiguous vesicle that may occupy more than the one half of the invaded hepatic lobe.

**Size of the hydatid cyst**

* In animals: varies from a marble to small football, usually of the size of goose egg.Early forms appear as white nodules, as yet containing no fluid: may be seen in the liver 4 weeks after ripe eggs are ingested.The cysts are 2.5.mm in size and contain fluid : after 8 weeks of taking ripe eggs.
* Hydatid cysts are 15-20 mm in diameter: after 6 months and only then do they produce scolices and brood capsules and become infective.
* The hydatid cysts develop slowly, taking months to increase in size to 5-10cm although some, especially in man, may reach 50 cm in diameter

An average-sized hydatid cyst may contain as 2 million protoscolices.Bovine livers of 91-113 Kg and pig liver of 50 Kg have been recorded. Such liver are markedly cirrhotic and cause ascites.

**In humans,** the slowly growing hydatid cysts may attain a volume of many liters and contain many thousands of protoscolices. The fluid is highly allergic.

Hydatid cysts in liver is always associated with marked fibrous tissue reaction which may be 13 mm thick. Affected liver are enlarged depending upon number and size of the cysts.

**Sites of cysts**

Liver and lungs are the commonest sites. In sheep lungs are affected as often as liver, the commonest form being unilocular fertile cysts; in the ox lung is affected more ofen than liver, usually with small unilocular cysts; in pigs and horse liver is the most frequent site of infection.

**Source of infection to man**

* Ingestion of the ova of dog tapeworm
* Contamination of the hairs of the dog’s coat with ova from feces: most common source
* Dogs may pass ova from anus by licking
* Hand to mouth transfer of tapeworm eggs by handling sheep fleeces of sheep contaminated by sheepdog feces

**Clinical features in man:**

* The clinical signs are caused by mechanical pressure exerted by the cyst on the surrounding tissue and hence vary with the size and location of cyst.
* Cysts of moderate size are generally asymptomatic and may cause mild abdominal heaviness.
* Large cysts, however, cause mechanical obstruction.
* Physical examination reveals hepatomegaly
* Jaundice may ensue from pressure on major biliary ducts.
* Fever, malaise, headache and eosinophilia are generally associated with the clinical disease.
* In pulmonary form, there is often coughing with or without haemoptysis(spitting of blood) Coughing with membrane from a ruptured cyst.
* Liver lesions are sometimes painful
* Cranial cysts may produce nervous symptoms
* If the cyst is ruptured inside the body, there can be secondary infection and/or cause fatal anaphylaxis.

**Clinical features in animals**

Generally few, if any , clinical signs despite severe levels of infection.Large cyst in liver cause ascites. Liver becomes cirrhotic.Adult worms in the dogs rarely cause problems except enteritis in heavy manifestations.The disease is usually detected during slaughter.

**Epidemiology:**

It is a cyclo-zoonotic infection. In India, the infection is prevalent widely in all parts of the country but the prevalence rates vary from place to place and species to species. In some areas the disease is found to affect a high (89%) proportion of cattle population. The prevalence of echinococcosis shows a marked variation in different species of domestic animals: 17.82-31.9%(cattle), 11.3-48.1% buffaloes, 2.75-30.5% sheep, 2.6-21% goats and 3,52% swine. In dogs the parasite can be found in as many as 16.30% of the animals.

Reports concerning the occurenec of human hydatidosis in India have been appearing in the literature from time to time. The infection is high between 16-30 years of age. The infection ratio between human female and male is 30:40.

Socio-economic, cultural and religious factors have been observed to play an important role in the transmission of this disease in man. The disease is more prevalent in Turkana people of North-western Kenya, where as per their religious customs the human dead bodies are exposed to hyenas and dog, thus perpetuating the transmission of the disease. Similarly, the Muslim belief of the unclean less of dogs have been responsible for the observed reduced incidence of the disease in Muslim Arabs as compared to Christian Arabs in Lebanon.

The disease is of much greater incidence in certain occupational workers like shoemakers and shoe-repairs in Lebanon, mainly related to heir practices of dipping hides of animals in a decoction of dog faeces for preparing leather.

It has also been observed that certain modes of recreation in man (sea side sports, camping, tourism, mountaineering, hunting, fishing etc.) have resulted in bringing man in close contact with hydatid endemic foci and increasing the chances of his acquisition of the disease.

**Diagnosis:**

1.Clinically, abdominal radiography, choangiography and liver scanning together with individual history diagnose the disease.

2. Casoni’s skin test, and intradermic allergic test using filtered hydatid fluid as antigen is a screening test of some value for estimating the prevalence of this disease.

3. Serological tests, such as, CFT, the latex agglutination test, the fluorescent antibody test, immuno-electroporesis etc. are helpful for studying the Epidemiology of the disease**.**

**Diagnosis in animals**

In the food animals the diagnosis is normally made postmortem

Confirmation is made at PM or by demonstartion of adult *E.granulosus* in the feces of dogs purged with praziquantel or other taenicide.

**Prevention and control**

* Heath educational efforts for improving personal hygiene in the handling of dogs as pets and companion
* The prevention of dogs from gaining access to raw offal and the proper disposal of the offal is an essential control measure.
* Reducing the dog population (Control of stray dogs).
* Minimizing dogs’ role in the transmission by mass treatment (praziquantel: 5 mg/kg body weight).
* Control of environmental contamination by dog feces.
* Awareness of the public especially the farmers in relation to responsible dog ownership
* Disposal of carcass of dead animals, especially sheep, properly and immediately by deep burial or incineration

8.Efficient meat inspection procedure with effective control of rejected meat and offals is an essential control measure.

9.Surgical removal of the cysts is a conventional practice of treatment especially in case of human patients.

**TOXOPLASMOSIS**

* Disease caused by arc (toxon) shaped organisms
* *Toxoplasma gondii*, an intestinal coccidian of felids is the causative agent
* Cats are the key host
* Socioeconomically important because of human sufferings and long term care of children with mental retardation and blindness are enormous

Stages of parasite:

* Adult parasite
* unsporulated oocyst : passed in the feces
* sporulated oocyst: infective stage in the environment. Each is having two sporocysts and each of it has 4 banana shaped sporozoites
* Bradyzoites: Resting stage
* Tachyzoites: multiplying stage

**Life cycle**

Hosts:

Definitive: Cats, both wild and domestic, are the only definitive hosts

Intermediate : Animals like sheep, goat, pigs, poultry and humans

Life cycle in two stages

Asexual stage: in intermediate and definite host

Sexual stage: occurs in definite host

**Clinical features in man**:

* It may occur as a *congenital* or an *acquired infection.*
* Pregnant women -- not themselves unusually prone to *T. gondii* infection or clinical disease
* Congenital infection may occur following maternal infection acquired during the first trimester of pregnancy are more severe than those acquired in the second and third trimester.

*Congenital infection* occurs by the transplacental transfer of *Toxoplasma*. There may be severe fulminating infection resulting in abortion, miscarriage, stillbirth or the birth of a baby with manifestations of acute toxoplasmosis. In the benign forms the disease may become manifest in infancy, child hhod or even adult life. Infection in first trimester is associated with more severe lesions. Congenital infection may be manifested by anaemia, jaundice, hepatospleenomegaly, lymphadenitis, and retinochoroditis. In neurological from of congenital infection there may be retinochoroiditis, convulsions and intracranial calcification.

***Disease in congenitally infected children***

*(i)* Mild disease may consist of slightly diminished vision only

(ii)Severely diseased children: full tetrad of signs including retinochoroiditis, hydrocephalus, convulsions, and intracerebral calcification. Of these, hydrocephalus is the least common but most dramatic lesion of toxoplasmosis.

Testing of all pregnant women for T. gondii infection is compulsory in some countries

Postnatally Acquired Toxoplasmosis

By ingesting food and water contaminated with oocysts from infected cat feces, or by ingesting tissue cysts in under cooked or uncooked meat.

Lymphadenitis is the most frequently observed clinical form. Postnatally Acquired Toxoplasmosis

* Lymphadenopathy may be associated with fever, malaise, rash, fatigue, muscle pain, sore throatand headache. It is also a common cause of chorioretinitis
* Although the condition may be benign, its diagnosis is vital in pregnant women because of the risk to the fetus.
* Encephalitis is the most clinically important manifestation of toxoplasmosis in immunosuppressed patients.

Toxoplasmosis is a major cause of death among patients with AIDS.

Pregnant women -- while not themselves unusually prone to T. gondii infection or clinical disease, parasite invasion across the placenta can result in severe neurological abnormalities in the fetus; congenital toxoplasmosis is a leading cause of birth defects.

A minority may experience temporary parasitaemia and may have low-grade fever, malaise, and lymphadenopathy.

Acquired infection in man is usually less severe and can present with a varied clinical picture. Lymphadenopathy is the commonest manifestation and may be accompanied with fever, malaise, and rash. It is also a common cause of chorioretinitis.

**Disease in animals**:

Severe disease is caused by T*. gondii* in many species of animals.

This includes:

* Cats of any age can die of toxoplasmosis, kittens and those with depressed immunity are the most likely.
* embryonic death and resorption, fetal death and mummification, abortion, stillbirth and neonatal death in goats and sheep.
* Outbreaks of toxoplasmosis in pigs cause higher mortality in young pigs than in adult pigs.
* . Sporadic and widespread outbreaks of toxoplasmosis occur in rabbits, mink, birds and other domesticated and wild animals.
* Among companion animals, fatal toxoplasmosis may occur in dogs that are immunosuppressed following infection with concurrent distemper virus.
* No clinical disease has been observed in cattle

**Epidemiology:**

*T. gondii* has the distinction of being one of the most cosmopolitan parasites known with more than 180 countries reporting its occurrence.

Toxoplasmosis is transmitted in two ways: acquired or congenital. Animal houses, gardens, fields and pastures are frequently contaminated by Toxoplasma oocysts excreted by the infected cats. After primary infection, a cat can shed oocyst for about two weeks and millions of these can be present in a single sample of stool. Oocysts are resistant to most ordinary environmental conditions and can survive under moist conditions for months and even years. Coprophagous invertebrates like cockroaches and flies can spread oocsysts mechanically. The sporulated oocysts can also be carried by wind, water, insects, birds and fomites from one place to another. Oocysts are resistant to environmental stresses and possess a remarkable transmission due to their small size.

The edible parts of food animals especially those of sheep, goat and swine contain the tissue cysts of *T.gondii*. The viable cysts can persist for life in animals. Raw or undercooked meat from infected animals is therefore, a serious threat to human health. Milk of goats has been found to contain the parasite and can also act as a source of infection especially for the human infants. Cow milk, however, has not been found to pose a similar risk. Carnivores and man are usually infected by the ingestion of tissue cysts.

Congenital transmission involves passage of tachyzoites through placenta to the developing foetus. Infections due to laboratory and autopsy accidents and handling of infected meat in homes and slaughterhouses have been reported.

**Diagnosis:**

***Demonstration and isolation of parasite***

* Biopsy material (lymph nodes and muscles) or impression smears can be used for detection of the parasite.
* Toxoplasma can be isolated from patients by inoculation of laboratory animals. Mice are highly susceptible.
* The inocula used are secretions, heperanized blood, CSF, biopsy material and products of conception in cases of abortion.
* Samples suspected of contamination are treated with antibiotics (1000units/ml penicillin; 10ug/ml streptomycin) and allowed to stand at room temperature for an hour.

***Detection of T. gondii oocysts in the environment***

* Contamination of the environment by oocysts is widespread, as oocysts are shed by domestic cats and other members of the Felidae. Cats may excrete millions of oocysts after ingesting as few as 1 bradyzoite or 1 tissue cyst, and many tissue cysts may be present in one infected mouse.
* Oocysts are detected in the feces of infected cats by concentration methods (e.g.flotation in high density sucrose solution) because too few may be present to be detected by direct smear.
* For definitive identification, T. gondii oocysts should be sporulated and then bioassayed in mice to distinguish them from other related coccidians.
* T. gondii oocysts have been isolated from soil.
* Bioassay of soil samples by feeding to pigs and chickens may be more sensitive than direct determination of oocysts in soil since pigs can be infected by feeding as few as one oocyst .
* Detection of T. gondii oocysts in water which however, is more difficult .

***Serology*:**

Paired serological tests (methylene blue dye test, IHA, CF, agglutination, IF and ELISA) can determine the infection status of animal/man.

**Prevention and Control**:

1.Proper cooking of meat is essential. Heating meat to 66oC kills the organisms.

2. The hazards inherent in handling raw meat is in slaughter houses and in the homes are well recognized and measures such as hand washing or wearing of rubber gloves should be adopted.

3. Freshly passed oocsyts in cat’s faeces are non-sporulated and therefore non-infective. Proper disposal at this stage by incineration or flushing down the toilet obviates the risk of infection

4. Control of diet of pet cats (prevent ingestion of raw meat, wild birds and rodents) and elimination of strays.

5. Sand and soil represent a durable source of infection; therefore children’s play areas should be made inaccessible to cats.

6. Pet cats shedding oocysts can temporarily be isolated and treated with sulphadiazine and pyrimethaine. Immunoprophylaxis has been attempted in animals using phenol killed toxoplasma and irradiated toxoplasma with some success.

7. As the foetus and neonates represent the most vulnerable section, avoidance of infection by pregnant women is very important. Routine serological check up in antenatal clinics and treatment of infected mothers would be a valuable step in protecting the foetus.

**LEISHMANIASIS**

Leishmania : A protozoa

Two morphological forms:

Promastigote:

* Spindle shaped body having a flagellum near the anterior end.
* Occurs in the gut of sandfly and artificial culture.
* Infective stage which is introduced into the skin by the bite of fly.

Amastigote ( Leishman Donovon bodies, LD bodies):

* Ovoid shape with one blunt end.
* Found in the mononuclear cells of the skin or reticuloendothelial system in vertebrates

**Types of Leishmanisis**

*1*.Visceral leishmanisis( Kala- azar)

* Caused by *L. donovani*
* Lesions from skin metastasizes throughout the reticuloendothelial system

2.Cutaneous leishmaniasis

* Caused by *Leishmania tropica*
* Primary lesion in skin and infection limits to it.

3. Mucucutaneous leishmaniasis( Espundia)

* Caused by *L.brazileinsis*
* Metastasis in lymph glands, skin and mucocutaneous junction

Visceral leishmaniasis

Clinical features:

* I-period: few months to two years, optimum is 4 months
* Discomfort below the left costal margin from the enlarged spleen
* Weak and emaciated appearance but with good diet and clean tongue
* Temperature of 102F but quite unaware that he has fever
* Fever is like enteric fever but no toxaemia and appetite is good
* Onset of malaria like fever with chills and rigors but unaffected by antimalarial drugs
* Irregular fever may continue for a prolonged period.

Post Kala-azar Dermal Leishmanoid

* This condition appears after about one year of treatment of visceral leishmanisis with sodium antimony compounds.
* There are skin lesions: erythematous patches, depigmented patches or nodules over the face, forearms, inner aspects of thighs.
* LD bodies are present in the lesions
* The phenomenon is the result of immune response of the host.
* Eruptions disappear during relapse and reappear with recovery from infection.
* Delayed hypersensitivity is positive in such cases.

**Epidemiology**

* Man to man transmission by the bite of sand fly, *Phleobotomous aergentipee( vector)*.
* Young adults and adolescents mainly affected. Children can also suffer.
* Disease affect low-socio economic group of people
* It is house based
* Overcrowding, ill ventilation, collection of organic matter inside the house facilitate transmission
* Alluvial soil, dense vegetation, high altitutde and high humidity(above 70%) facilitate epidemics
* Reservoir: Canines( Dogs in urban region, jackals in sylvatic transmission)
* Rodents have been found in Kenya

**Laboratory Diagnosis**

* Direct Demonstration of parasite
* Microscopic examination of the bone marrow, spleen or liver biopsy for LD bodies.
* Culture of bone marrow aspirate, liver and spleen biopsy material to NNN medium, incubate at 22C for a week and examine for promastigote

**2. Indirect evidence**

* 1. Leucopenia with relative neutropenia.
* 2. Aldehyde test
* 3. Demonstration of antibodies

Aldehyde test

* Principal: Marked production of globulin by plasma cells in Kala-azar. The test detects these globulins.
* Method:
* 1 ml serum + one drop of 40% formaldehyde
* Mix and leave for 2 hrs.
* Positive test: Opacity of egg white color along with jelly formation within 20 min to 2 hrs
* The test is good for surveillance.
* Not good as diagnostic test because test is positive only after 2-3 months of infection and becomes negative after 6 months of cure.

Demonstration of antibodies

* Complement Fixation test
* Indirect Fluorescent Antibody test( IFT)
* ELISA
* Leishmania Skin Test( Delayed hypersensitivity reaction)

**Cutaneous Leishmaniais**

* Incubation period: Few weeks to six months. In some cases it may be one to two years.
* The lesion begins as a small nodule, which ulcerates in majority of the cases. The parasite is found along the red margin.
* The sore heals spontaneously.
* The sores are distributed on exposed parts of the body, particularly on the face and extremities.
* Two species of sandflies: *Phleobotomus paptasi* and *Phleobotomus sergenti* transmit this type leishmaniasis.

Dogs act booster animal during epidemic

Mucocutaneous Leishmaniais

* *Leishmania braziliensis* produces single or multiple, seldom self-healing ulcers in the primary cutaneous stage. The condition is accompanied by involvement of lymphatic. Untreated primary ulcers lead to mucocutaneous leishmaniais.
* The metastatic lesions spread along the lymphatic, producing nodules is essential form of the disease

Tapir Nose

* A condition in which nasal and pharyngeal cartilages are destroyed, lips and nose swell.
* The condition may be painful or painless with secondary infection.
* Mucocutaneous leishmaniais is a serious type of condition.
* The primary lesions are similar to cutaneous leishmaniais in the mucosa of upper respiratory tract.
* In two third of the patients, lesions restricted to nose.
* In rest of cases, pharynx, larynx, palate and lips also involved. The lesion progresses as: Small crust------ulceration------erosion-----destruction of tissues. Nasal septum may be destroyed.

**Prevention, Control and Eradication of Zoonotic Diseases**

**Control:** The reduction of disease incidence, prevalence, morbidity or mortality to a locally acceptable level as a result of deliberate efforts; continued intervention measures are required to maintain the reduction. Example: diarrhoeal diseases.

 **Prevention** primarily covers measures to prevent the occurrence of disease, such as by reductions of risk factors.

It also means to arrest disease progress and reduce its consequences once established. These are known as secondary and tertiary prevention that arrest or retard existing disease and its effects through early detection and appropriate treatment; or to reduce the occurrence of relapses and the establishment of chronic conditions through, for example, effective rehabilitation.

**Eradication:**Permanent reduction to zero of the worldwide incidence of infection caused by a specific agent as a result of deliberate efforts. Example: smallpox.

**Methods of prevention and control**

* Interruption of transmission
* Controlling reservoir of infection
* Protecting susceptible population

**1. Interruption of transmission**

**1.1.** Changing some components of environment to prevent the infectious agent from patient or carrier from entering the body of susceptible host, e.g., chlorination of water will interrupt cholera agent from water to host.

**1.2.** Food borne zoonotic diseases: Hand washing, adequate cooking, prompt refrigeration of food and withdrawal of contaminated food. Establishment of food hygiene like hygiene in animal production; hygiene of slaughter; hygiene in handling and processing food stuffs.

**1.3.**When vector borne: control measures primarily at the vector and its breeding places.

**1.4**. Droplets or droplet nuclei transmission: Early diagnosis and treatment, personal hygiene and proper handling of secretions and excretions.

**1.5.** Ensure safety of animal products (wool, hides, bones, fat, others) by hygiene during collection, storage, processing, transport.

**1.6.** Ensure safety or use of animal carcasses and washes by proper animal carcass disposal and excreta/garbage disposal.

**2. Controlling reservoir of infection:** The **reservoir** for infectious agents is the principal habitat where a specific infectious agent lives and multiplies. The reservoir is necessary for the infectious agent either to survive, or to multiply in sufficient amount to be transmitted to a susceptible host, e.g. soilis the reservoir for *Clostridium tetani* or **water**, the reservoir for *Legionella pneumophila*.

Elimination of the reservoir may be easy with animal reservoir ( e.g. bovine tuberculosis, brucellosis) but not possible in humans.

**General measures of reservoir control comprise:**

2.1.Early diagnosis: This is the first important step.

2.2.Notification: Once disease has been detected or even suspected, it should be notified to the local health authority that will control measures. Some diseases are notifiable which are considered to be of serious menaces (hazards) to public health.

2.3.Isolation: It means separation for the period of communicability, of infected persons or animals from others in such places as to prevent or limit the direct or indirect transmission. Isolation is also possible by “ring immunization” i.e.encircling the infecting population with a barrier of immune one through whom the infection is unable to spread.

2.4.Treatment: The objective is to kill the infectious agent when it is still in the reservoir, i.e.before it is disseminated. Treatment reduces the communicability of disease, cut short the duration of illness and prevent the development of secondary cases. Treatment can be individual treatment or mass treatment. Inadequate treatment may induce drug resistance and affects the control by chemotherapy.

2.5.Quarantine: It has been defined as the limitation of freedom of movement of such animals or persons exposed to disease for a period of time no longer than the longest usual incubation period of the disease. This will prevent the contact of infected from uninfected population.

**3.Protecting susceptible population**

3.1.Active immunization: Several vaccines are available for animals and human. Rabies is a good example.

3.2.Passive immunization: Useful only when exposure to exposure has just occurred or is possible within the next few days. Immunity is short and variable 1-6 weeks.

3.3.Chemoprophylaxis

3.4.Eliminating the pathogen from its animal reservoir(s). In some countries, livestock diseases such as bovine and porcine brucellosis and bovine tuberculosis have been eradicated, and the prevalence of *Salmonella* in poultry has been significantly reduced.

3.5.Vaccination.

3.6.Flea and tick control.

3.7.Periodic testing for enteric parasites or other pathogens, and other disease control measures in domestic animals can also protect people.

3.8.Non specific measures: Improvement in the quality of life like better housing, water supply, sanitation, nutrition.

**4.Surveillance** is defined as the ongoing systematic collection, analysis, interpretation, and dissemination of outcome-specific data essential to the planning, implementation, and evaluation of public health practice.

**Veterinary surveillance:** As frontline healthcare providers, veterinarians assist with the recognition, diagnosis, reporting, and **control of zoonotic disease in animals.** When an unusual zoonotic disease trend or outbreak is recognized, veterinarians can assist the investigation through enhanced surveillance for animal disease. Health alerts typically include information on veterinary occupational risks as well as symptoms, diagnosis, and reporting protocols for the disease in animals.There should be veterinary alert systems, as in developed countries, for rapid notification of zoonotic or animal disease outbreaks.