

INTERNATIONAL SCHOOL OF MEDICINE

Department of Infectious Diseases

The topic of the lecture:

Trypanosomiasis

Professor Kutmanova A.Z.

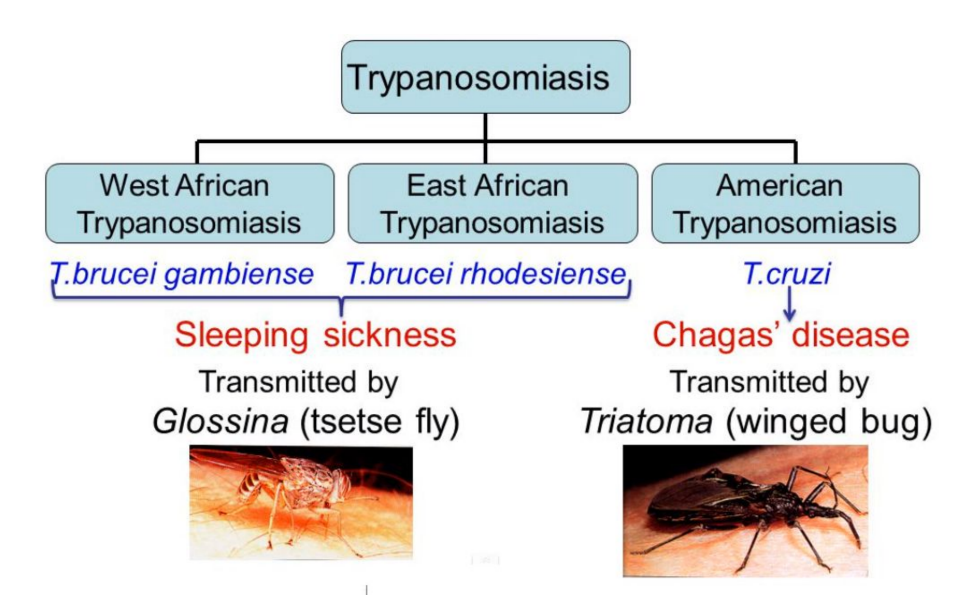
History

- its species was discovered by a scientist Valentine" in 1841.
- But these are found in mammals after 50 80 years later.
- Then further studied about their diseases.

Introduction

- The name is derived from Greek word,
 - trypano means (borer)
 - soma means (body)
- They are unicellular flagellate protozoa.
- Have spiral like motion.
- Need more than one host to complete its life cycle.
- Oftenly transmitted by a vector.
- Generally found in intestine, but some time found in blood stream or in heart.

Trypanosoma



Morphology of trypanosome

Morphology

Exist into 2 interchangeable forms:

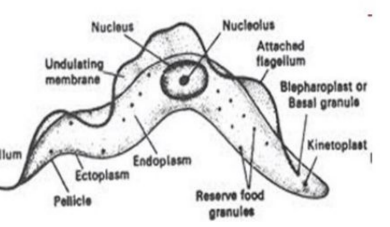
Trypomastigote in <u>Blood/ Lymph</u>/ tissue space of various organs & free flagellum <u>C.N.S is terminal & fatal</u>

Epimastigote in salivary gland of vector & Culture media.

Trypomastigote (Polymorphic Trypanosomes

Spindle shaped – Central nucleus – free flagellum – undulating membrane. <u>3 forms</u>

- 1- long Slender Form (30µ): active motile with free flagellum.
- 2- Short stumpy Form (15µ): sluggish without free flagellum.
- 3- Intermediate Form (20µ): with a short free flagellum.



Trypanosoma gambiense

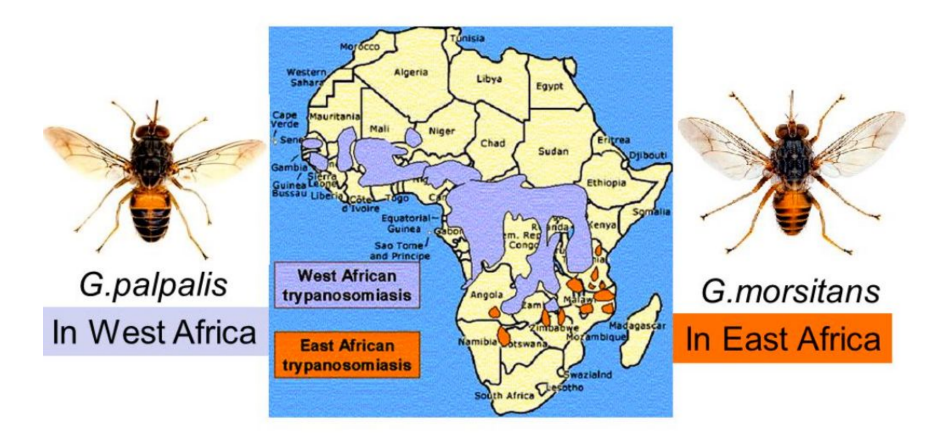
African Trypanosomiasis (Sleeping Sickness)

The parasites

- Belong to a group of closely related trypanosomes in the Trypanosoma brucei species complex. Three morphologically indistinguishable species are recognized:
- T. brucei infects game animals/livestock (causes nagana)
- T. rhodesiense causes E. African trypanosomiasis
- T. gambiense causes W. and Central African sleeping sickness

(Some authors consider these as subspecies: T. brucei brucei, T. b. rhodesiense, T. b. gambiense.)

Distribution

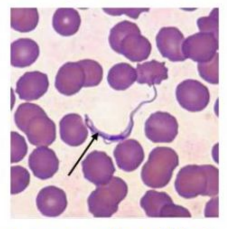


T. rhodesiense causes E. African trypanosomiasis

T. gambiense causes W. and Central African sleeping sickness

Trypanosoma brucei species

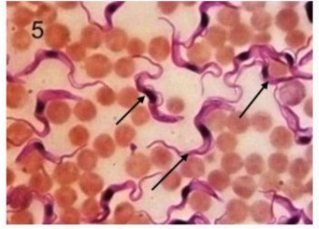
West Africa *T.brucei gambiense*



Less plentiful Cannot live in lab animals

Reservoir host: goats, cattle & pigs Transmitted by: *G.palpalis*

East Africa *T.brucei rhodesiense*



More plentiful <u>Can live</u> in lab animals Nucleus is shifted posteriorly <u>Reservoir host:</u> wild game animals Transmitted by: *G.morsitans*

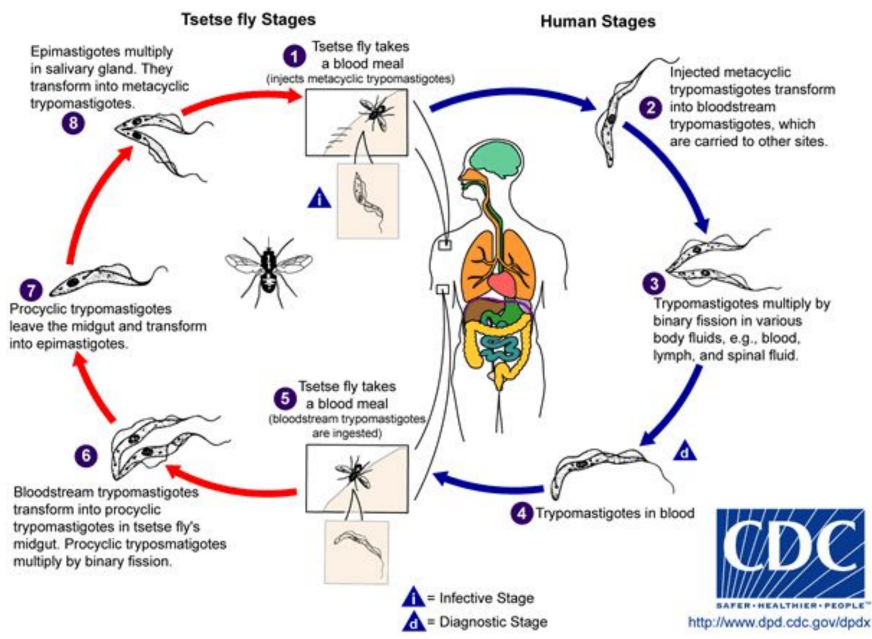
Major Differences Between African Trypanosome Species

Attribute	T. rhodesiense	T. gambiense
tsetse vector	G. morsitans group	G. palpalis group
ecology	dry bush, woodland	rainforest, riverine, lakes
transmission cycle	ungulate-fly-human	human-fly-human
non-human reservoir	wild animals	domestic animals
epidemiology	sporadic, safaris	endemic, some epidemics
disease progression	rapid, often fatal	slow (~1 yr) acute \Rightarrow chronic
parasitemia	high	low
asymptomatic carriers	rare	common

Transmission

- A bite from an infected tsetse fly causes African trypanosomiasis.
- Blood transfusions are a rare cause of parasitic transmission.
- In rare cases, accidental transmission in the laboratory has been implicated.

Life cycle



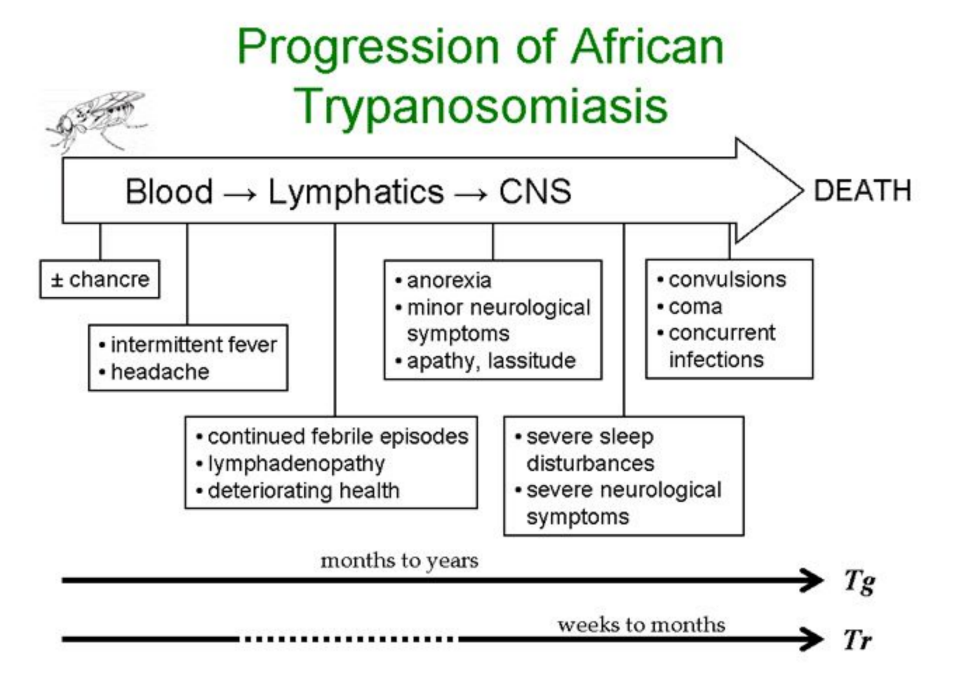
Pathogenesis

Incubation period (2 weeks) Trypanosoma chancre (at the site of bite) Via lymphatics: enlarged lymph nodes especially posterior cervical region. (Winterbottom's sign) Via blood stream: headache, fever(fluctuating), muscle & joint pain, irregular erythematous rash. Invasion of bone marrow (hypoplastic anaemia) Enlarged liver & spleen, generalized weakness. Invasion of CNS: severe headache, mental apathy, slow speech, deep sleep, coma & death In East African Trypanosomiasis: Disease runs more rapid & fatal course

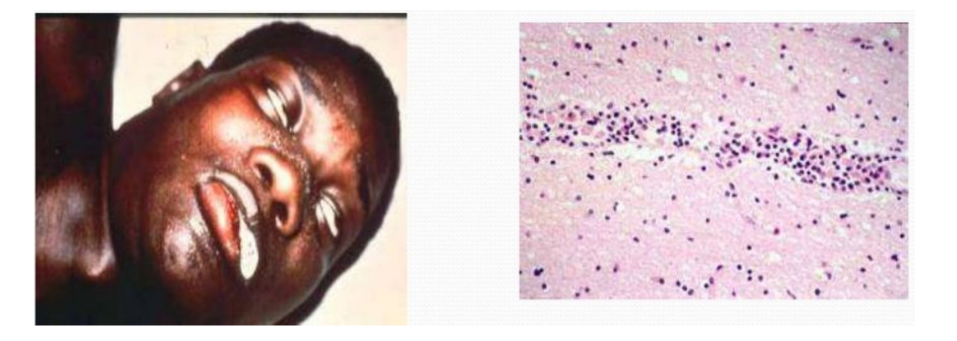
Clinical presentation

- Incubation period may be from few days to weeks.
- The first clinical manifestation of African trypanosomiasis chancre occurs at the site of inoculation .



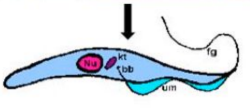


Clinical presentation



Diagnosis

- 1- Clinical picture
- 2- Demonstration of trypanosomes:
 - Microscopic examination of unstained or stained blood films
- Culture on suitable medium (N.N.N OR Weinmann's media to detect <u>Epimastigote</u>)



- Animal inoculation

N.B. in case of <u>T.brucei rhodesiense</u> injected in lab Animal produce a new form "Posterior Nucleus Shift

Polymorphic Trypanosomes





Diagnosis

General

- Anemia, hypergammaglobulinemia, low complement levels, elevated ESR, thrombocytopenia, hypoalbuminemia, but not eosinophilia or abnormal liver function.
- The total IgM level is higher in blood and CSF.
- A definitive diagnosis of infection requires actual detection of trypanosomes in blood, lymph nodes, CSF, skin chancre aspirates, or bone marrow.
- However, symptomatic improvement after empiric treatment is the usual confirmatory test in areas where diagnostic studies are not readily available.

Imaging Studies

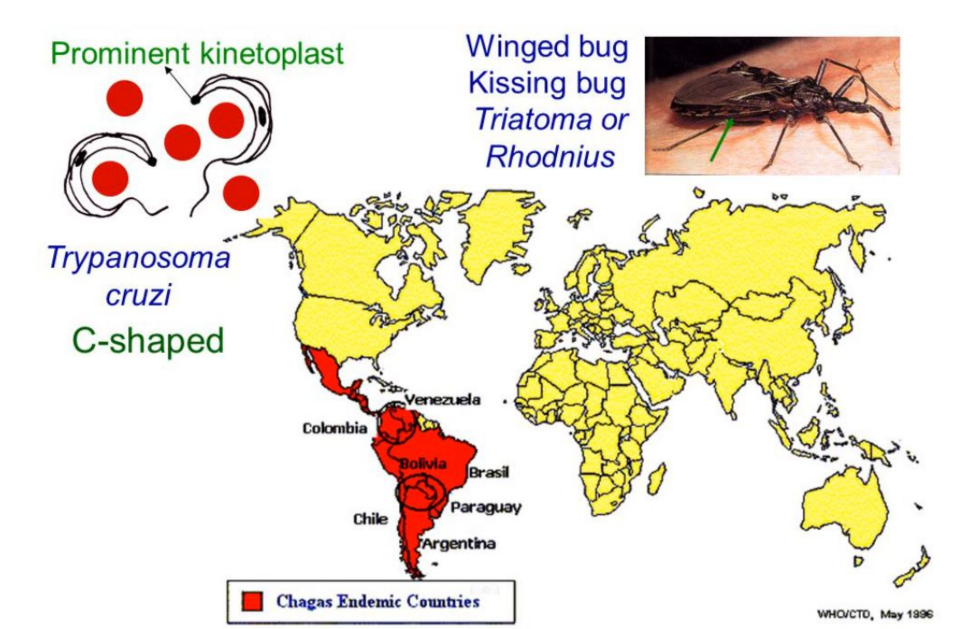
- CT scanning and MRI of the head: Both head CT scanning and MRI reveal cerebral edema and white matter enhancement, respectively, in patients with late-stage African trypanosomiasis.
- EEG in neurologic involvement usually shows slow wave oscillations (delta waves), a nonspecific finding

Treatment

Drug	Use	Drawbacks
Pentamidine	Effective against early-stage gambiense disease	 Adverse side effects Non-oral route
Suramin	Effective against early-stage gambiense and rhodesiense disease	 Adverse side effects Non-oral route
Melarsoprol	First line drug for late-stage gambiense and rhodesiense disease involving CNS	 Adverse side effects, especially encephalopathy Fatal in 1-5% of cases Parasite resistance Non-oral route
Eflornithine	Effective against late-stage gambiense disease involving CNS	 High cost Not effective against <i>T. rhodesiense</i> Non-oral route - has to be given intravenously (needs hospitalization for 14 days)

<u>American Trypanosomiasis</u> (<u>Chagas Disease</u>)

Distribution



Morphology

Morphology

Trypomastigote (Monomorphic)

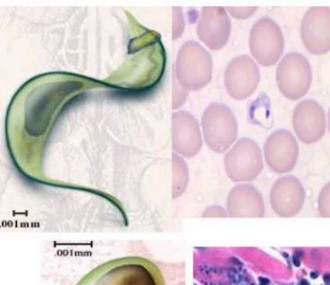
Slender shaped (20µ) – Central nucleus – C or U-shaped –Free flagellum 1/3 body-Large bulging peripheral kinetoplast

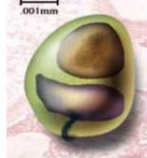
Amastigote

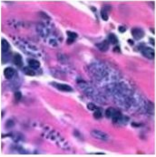
Obligatory intracellular – mainly in cardiac & Skeletal muscles – Brain meninges – Nerve ganglia – cells of GIT etc

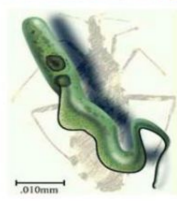
Epimastigote (Vector only)

Spindle shape– Kinetoplast anterior to central nucleus– Undulating membrane is short – terminal free flagellum



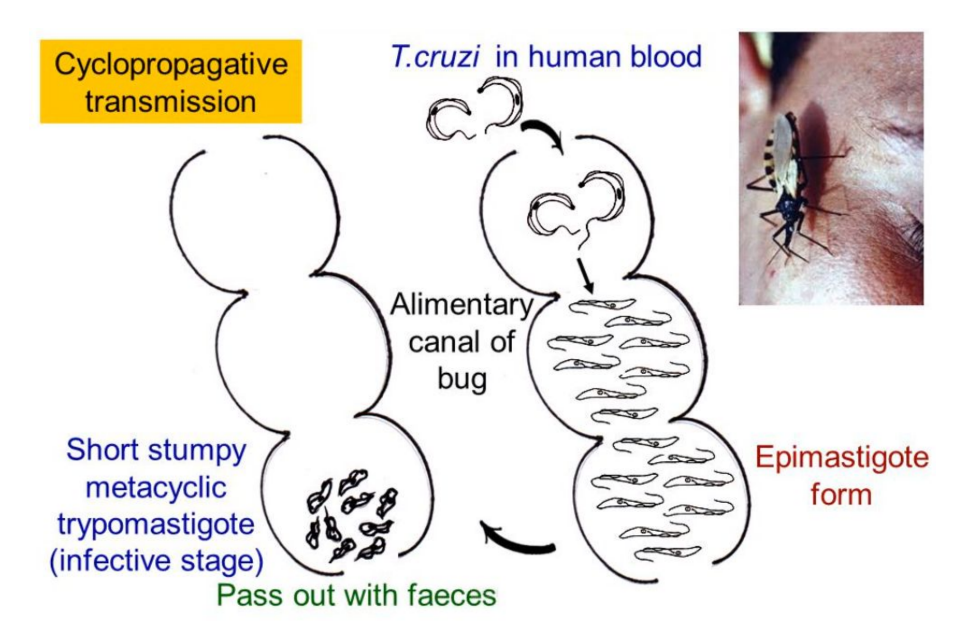




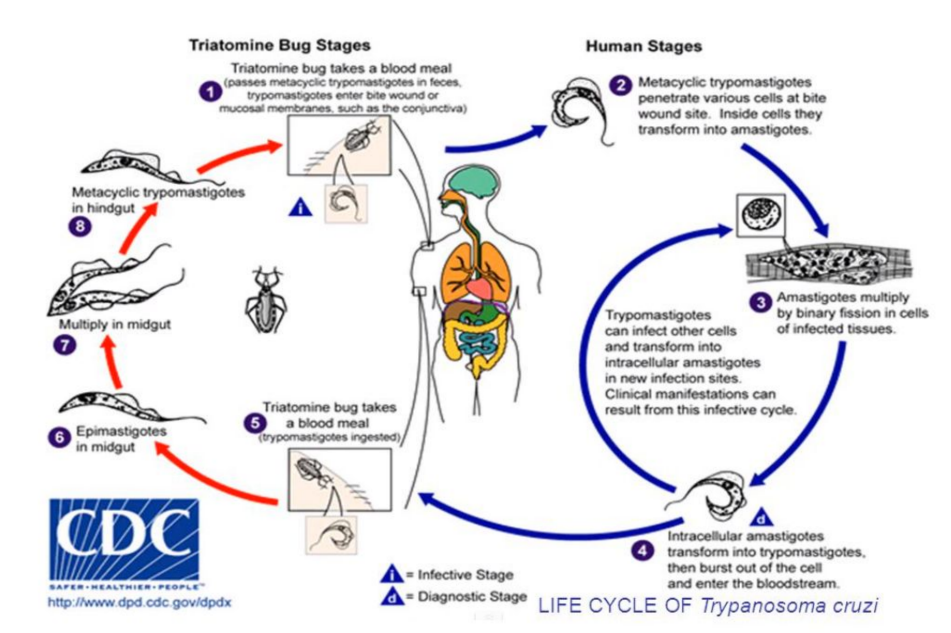




Transmission



Life cycle



Transmission

Mainly by

Contamination of skin abrasion by winged bug faeces

Cone nose Bug – kissing Bug – Assassin bug

Rarely by

Through infected blood transfusion Through infected mother's milk Through the placenta







Pathogenesis

I- Acute Form

Chagoma occurs at the site of bite. Parasite reaches regional lymph nodes Blood To. Organs and tissues Fever, enlarged lymph nodes, skin rash, enlarged liver & spleen. Romana's sign (Unilateral conjunctivitis appear suddenly

together with oedema of upper & lower eye lids & cheek)

Meningoencephalitis, heart failure

Death or pass to Chronic form







Pathogenesis

II- Chronic form

Parasite produces antigens similar to patient's self antigens: The body produces auto-antibodies that cause damage to:

Heart muscle fibres: congestive heart failure.

 Oesophageal muscle fibres: megaoesophagus and dysphagia. Destruction of Auerbach's plexus
 Colon muscle fibres: megacolon and constipation.

CNS or thyroid gland Exacerbation of infection in immunosuppressed patients.





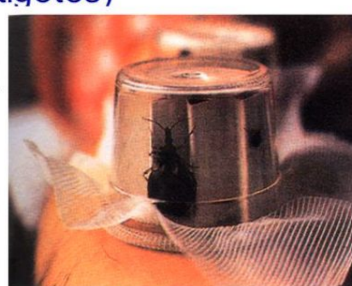
Diagnosis

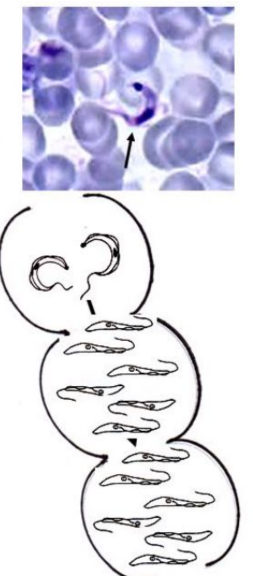
Finding the parasite in: <u>Blood film (C-shaped *T.cruzi*)</u> <u>Biopsy</u> from lymph node, liver or spleen (amastigotes) <u>Culture</u> (Epimastigotes)

<u>Xenodiagnosis</u>

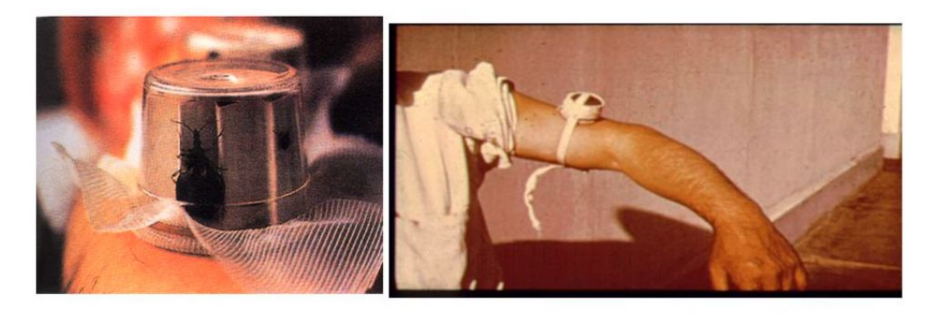
Serological tests Cruzin test (I.D.)

Molecular techniques





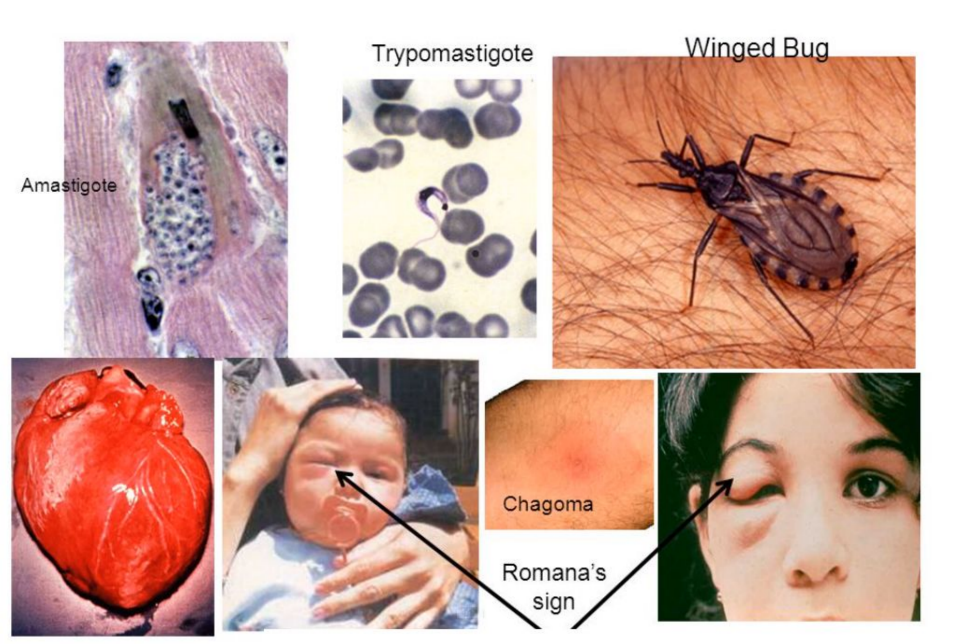
Diagnosis (Xenodiagnosis)



Highly efficient – demonstrate low level of parasite in blood Method:

A Laboratory bred winged bug is starved for 2 weeks then fed on suspected patient's blood – 30 days later, it faeces & gut examined for trypanosomes.

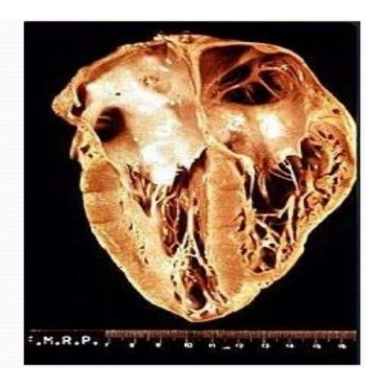
Diagnosis



Clinical presentation

- Chronic stage.
 - In chronic stage these parasites reside in heart and midgut, and some time in hind gut causing giantism of the midgut.





Clinical presentation

• Gigantism of midgut



Treatment

Sleeping Sickness

In early stage of the disease: Pentamidine OR Suramin

In late stages of the disease:

Tryparsamide

For both early and late stages of the disease:

> Eflornithine (DFMO) Ornidyl

Chagas Disease

Nifurtimox

- inhibits intracellular development.
- Drug of choice in acute and early chronic

OR

Primaquine destroys Trypanosoma in blood

Control

Sleeping Sickness

Treatment of patients

Control of vectors (Glossina)

Pentamidine as prophylactic drug Chagas' disease

Treatment of patients

Control of vectors (Triatoma)

Elimination of reservoir hosts