ANTIPROTOZOAN DRUGS
AT A GLANCE

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INSTRUCTIONAL OBJECTIVES

- List drugs used to treat trypanosomosis
- Drugs used to treat piroplasmosis prevalent in Nigeria.
- Have idea of mechanism of actions of drugs used, side effects, and Pharmacology.
- Vividly list other drugs used to treat giardiasis, cryptosporidiosis, toxoplasmosis.
- Therapeutic rational used in anticoccidials
ANTIPROTOZOAN AGENTS

Protozoal infections are common in tropical and sub tropical countries where sanitary conditions, hygiene practices, and control of the vectors of transmission are inadequate. Two types of infections caused by the major types of protozoa of veterinary importance are the haemoparasitic e.g. trypanosome, babesia, theileria, and the common enteric coccidian, toxoplasma and giardia.

ANTI TRYPANOSOMAL DRUGS:

**Diamidines** – Chemistry – the trypanocidal action of diamidines is related to the Amadine or Guanyl structure. Examples are Dinizanzen aceturate, Phenamidine, Stilbamidine and Pentamidine.

Diminazene aceturate -

- It is odourless
- Yellow powder
- Soluble in water
- Slightly soluble in organic solvents.

**Mechanism of action** –

- It binds irreversibly but not directly. It binds to the groove between the complementary strands of DNA – regular intervals and thus distorts the helical structure.
- It affects the phospholipids synthesis
- It is said to displace magnesium ion and it inhibits polyamines in the parasite.
- The drug is said to have dyskinetoplastic effect in the parasites.
- The drug interferes with glycolytic pathway of the parasite.

**Indication and Uses**

- Trypanosomosis in early stage
- Babesicidal effect or in babesia infection.
- It has bactericidal effect against boucella and streptococcus.

**Limitations**

- The drug is not effective in late stages of trypanosomosis
- There are report of resistance to the drug and relapse of infection
- It is ineffective against *Trypanosome evansi* in camels at 3.5mg/kg
- Special dose regimen is required for *T.brucei* infection.

**Side Effects**

- Local reactions in horses and rats might occur at site of injection.
- Neurotoxicity in dogs especially exotic breeds, ataxia, convulsion.
- Nephrotoxicity may be induced by the drug.
- Hepatic impairment
**PHARMACOKINETICS**

- It is poorly absorbed orally
- But the drug is rapidly absorbed intramuscularly and subcutaneously.
- Distribution of the drug in the tissues is rapid and wide
- Dimidines accumulate in the liver for months, like wise in the kidney and the adrenal glands respectively

**Dose**

For babesia and trypanosome infection a single intramuscular injection of 3.5mg/kg of 7 percent freshly prepare aqueous solution.

*T. brucei* infection requires 7mg/kg against *Babesia cabau* and *B. equi.*, two doses (5 and 12mg/kg ) are given apart

Today, Dimirazene aceturate is formulated with phenazone (8.75%), an antipyretic and analgesic to reduce the pain at the site of the injection.

**PENTAMIDINE.**

Pentamidine isethionate is preferred drug for prevention and treatment of haemolympathic stage of human trypanosomosis because the drug does not pass the blood – brain barrier; it therefore is not used to treat CNS involvement.
STILBAMIDINE ISETHIONATE

This drug has anti protozoal and antifungal activity. It is effective in blastomycosis and it is used in the treatment of human visceral leishmaniasis and early stage of sleeping sickness. It is more toxic then pentamidine.

PHENANTHRIDINES

History:

Phenanthridinium derivatives were introduced in 1938

The first active agents are phenidium and dimidium. This drugs exhibited photosensitization and other toxicities. Other agents replaced this drugs that are associated with toxicities, the less toxic homidium amd isometamidium are now in use.

Mode of Action

The phenanthridines bind strongly with D.N.A. especially kinetoplast D.N.A.

- They interfere with glycosomal functions.
- They interfere with function of unusual adenosine monophosphate – binding protein.
- They may cleave K.D.N.A. – topoismerase complexes this result to dyskinetoplastic trypanosome
Homidium is mutagenic and trypanosomes exposed to it for one hour may retain motility for 24 hours, but are no longer infective.

**PHARMACOKINETICS**

- They are poorly absorbed orally
- They are rapidly absorbed when injected intramuscularly
- It is eliminated in 24 hours
- Isometamidium administered it could form a tissue depot at the site of injection. The drug therefore is very slowly absorbed giving effective protection for up to 6 months
- Distribution is wide and prolonged the drug accumulates in the liver.

**Uses:**

Homidium is active against

- *T. vivax*, *T. congoïdense* and less active against *T. brucei*.
- *T. evansi* and *T. cruzi* has been reported to respond to carlidium another member of the group of Homidium salt.

**Dose**

Homidium is available as the bromide and the more water – soluble chloride salts.

A single dose of 1.0 mg/kg of a 2% solution is given 1/m. in to the neck or
ISOMETAMIDIIUM.

Originally known as methamidium or trypamidium

Mechanism of Action.

- It inhibits D.N.A. synthesis in a similar manner as diminazene aceturate.
- It modifies the mitochondrial membrane
- It modifies the glycoprotein structure in surface of the endoplasmic reticulum

Chemotherapeutic uses.

- It is effective against *T. vivax* and congoense.
- It is effective when trypanosomes are resistant to other conventional drugs.
- It has narrow safety margin and rather a narrow spectrum of activity.
- It is used prophylactically to prevent *T. congoense* and *T. brucei* in dogs.
- It is used in animals during long trek through the tsetse infested sore.
- It is used to confer protection against trypanosomal infection in endemic areas at 3-6 months.

Limitations of the Drug.

- It has narrow safety margin
- It is reported that there is relapse after the use of the drug
- There is severe local reaction at site of injection.
Dose

0.5-1mg/kg body wergilds administered deep intramuscularly or 0.25-0.5mg/kg

Quinapyramine Compounds.

Examples are quinapyramine chloride, quinapyramine sulphate, and suramin these compounds are dimethyl chloride is absorbed slowly.

In preparation of the drug 3 parts of dimethlysulphate and 2 parts of dimethyl chloride this is called “antrycide prosalt” and this is used for therapy and prophylexis some-times it is given in combination with suramin another quinapyramine compound.

Spectrum of Activity.

- It is effective against *T.congolense* and *T.vivax* in cattle and other animals.
- It is also effective against *T.bruceti* and *T.evansi*

Limitations of the Drugs.

- It is poorly tolerated by horses.
- It cause serious local reactions at the site and should be given in two or more divided doses at 6hours interval using 5% solution and 10% subcutaneously
Mechanism of Action

- It is a trypanostatic in action and therefore the host defence mechanism is very important in overcoming the infection.
- It causes a kinetoplastic D.N.A. condensation.
- It causes the loss of ribosomes with possible aggregate formation with large number of lysosomes.

Dosage

- It is given at 4.4mg/kg in heavier animals
- 150-200kg : 1g.
- 200-350kg : 1.5g
- over 350kg
- over 350kg weight : 2g

Contraindication

- The drug should not be used in young stock because it causes sweating, salivation, polypnoea, tachycardia and death might occur.

The Use of the Drug in Horses

- It is used prophylactically in horses especially in the breeding season at an interval 90days between injections is satisfactory.
• Transmission during service can be prevented using quinapyramine 18 days before service and it is used to prevent Dourines disease.

SURAMIN

Chemistry:- Surmin is a complex water-soluble derive of urea.

• It is complex aromatic organic compounds
• It is hydrosopic powder
• It has lower solubility in water.

Mechanism of Action

• The drug bind avidly to proteins and inhibits many enzymes among them, are those involved in energy metabolism (e.g. glycerophosphate dehydrogenase). This mechanism is correlated with the trypanocidal activity.
• It also distorts the intracellular membrane in lysosomes

Uses

• It is used prophylactically and curatively
• It is use against T.evansi the cause surra in horses, trypanosomosis in cattle and dogs.
• It is potentiated by phenanthridium and quinidine derivatives
Dosages

- Horses 7-10mg/kg bwt
- Camels 8-12mg/kg bwt
- Cattle 12mg/kg bwt

The dose in horse may be repeated three times weekly interval

Limitation of Surmin

- Narrow margin of safety
- It does not cross the blood-brain barrier so it could not be used in chronic or late stage of trypanosomosis.
- Camel trypanosomosis are quite resistant

Toxicity and Adverse Effects

The toxicity is frequent and severe this thus poses as nephrotoxicity, hepatotoxicity damages to the spleen and adrenal gland.

Synergistic Property of Suramin

Suramin would be as a supergistic potentiatior of other drugs (e.g suramin /quinapyramine) (homidium suraminate)
ORGANIC ARSENICALS

An example of organic arsenicals is melarsomin, or melarsoprol.

Organic arsenicals are used in treatment of late–stage of human African trypanosomosis

- It is used in haemolympathic stage of the disease. the drug is effective in late stage of the disease and it can pass to the blood-brain barrier to cause the therapeutic effect, when the trypanosomes are in the cerebro-spinal fluid and in the C.N.S

Limitations of Organic Arsenicals

- It is restricted to I.V. administration by the W.H.O. to avoid reactions
- There is relapse in melarsonyl than melarsorprol.
- Encephalopathy might occur.

Mechanism of Action

- It combines with the enzyme system in trypanosome trypanothione oxidase reductase system.
- Arsenicals acts by interacting with S.H group which is essential for intracellular metabolic process.
- It also act on the glycolytic enzymes
OTHER TRYPANOSOMAL DRUGS

- Antimony and potassium tartrate used I.V at 3.5mg/kg in horse and cattle and 1-3mg/kg for dogs.
- Stibophen.
- Trypan red
- Trypan blue.

ANTIPIROPLASMAL COMPOUNDS

The clinically important proplasms are anaplasmosis, babesiosis, cowdriosis, theileriosis, ehrlichriosis, hepatozoonosis and in avians spirochaetosis

AMICARBALIDE ISETHIONATE

Chemistry:- chemically made of complex urea compounds.

Uses:-
Used in babesiosis and theileriosis namely *B.divergens, B.cabali* and *Theileria pavae*

Route of Administration
1/m and I.V.

Dosage
5-10mg/kg between.
Toxicity: It might cause local irritation and localized swelling at the site of administration.

Treatment Regimen for Anaplasmosis

In treatment of anaplasmosis tetracycline and Imidocarb are of value in treatment, prophylaxis, and elimination of carrier–state a single l/m inj of oxytetracycline at 10mg/kg between will produce cure at 5% conc 2-3 daily doses may be necessary

- If long acting 20mg/kg is needed
- To eliminate the carrier-state, oxytetracycline is administered at a daily 1/m or 1/v dose rate of 11mg/kg for 10-14 days.
- Oral chlortetracycline is administered at 45-60 days or long acting of oxytetracycline is administered twice at 20mg/kg between l.m. 7 days apart
- Imidocrab at 3.5mg/kg birth weight, I.M; the dose is repeated 10-14 days.
- To eliminate the carrier-state, tow intramuscular or subcutaneous doses, each at 4mg/kg, 24 hours.

Treatment of Theileriosis

Like anaplasmosis, there is no specific treatment for theileriosis. But treatment with buparvquone, halfuginone, menoctone, parvaquone and tetracycline.

Infection detected at early stage could be treated using short acting at a dose of 15mg/kg I.M. for 5 consecutive days. The long acting oxytetracycline is administered once at 20mg/kg I.M.
Imidocarb (Imidocarbdipropionate)

Physical properties

- It has a white coloured appearance
- It has a melting point greater than 200°C

Chemistry

Imidocarb is an carbanilide dimidines

Uses:

- It is efficacious against babesiosis in dogs.
- Anaplasmosis (in cattle)
- Ehrlichiosis in dogs.
- It could be used prophylactically and therapeutically.

Pharmacokinetics:

After I.V. injection in sheep the drug would reach its peak in the plasma level of 10.8mg/ml\(^{-1}\), this would drop to 1.9mg/ml\(^{-1}\) in an hour.

It can be detected in the blood for 4 weeks.

The drug is detected in urine and faeces in an unchanged form.

Dosage:

Imidocarb is administered either I.V., I/M or S/c.

Babesiosis Therapy

Cattle - 1.2mg/kg birth weight
Horse - 2.4mg/kg birth weight.

Dogs - 6mg/kg birth weight

**Anaplasmosis Therapy**

Cattle – 3mg/kg birth weight

**Safety and Toxicity**

It has a very high safety margin in rats and dogs.

It has low safety margin in cattle.

**Withdrawal period.**

Withdrawal period of the drug is 28 days.

Sometimes 90 days withdrawal period might be required after last treatment.

**QUINURONIUM SULPHATE**

Quinuronium sulphate is used against *Babesia cabali, B. bouis, B. ouis, B. molasi*.

The drug is used in febrile stages in 24 - 48hrs a second treatment might be necessary. The course of treatment might be repeated for 2 weeks but preferably for 3 months.
**Premunity and Quinoronium**

The drug should not be used for a long duration. This might cause animal susceptibility to piroplasms. Therefore it is preferred to inoculate the animal with virulent strain of the parasite, but at a dose lower than the dose that will cause the disease.

**Dosages:**

0.3-0.5mg/kg birth weight for cattle, sheep, pigs and in rats. 0.5mg/kg birth weight for dogs, 0.25mg/kg birth weight. The drug should be diluted by 120 times i.e. 0.5%, but officially the drug is concentrated at 5%.

**Toxicity**

- Tremor
- Salivation
- Urination
- Defecation
- Shock might occur due to fall of the blood pressure.

Other Anti-proplasmosis drugs are

- *Trypan Blue*
- *Diaminazene*
- *Trypan red*
ANTIHISTOMONIASIS

- Aminonitrothiazole
- Nithiazide

*Pls read on your own this group of drugs.*

TREATMENT OF GIARDIASIS

- The main drugs used for treatment of giardiasis are follows: Metronidazole, dimetridazole, pronidazole, tinidazole, nimorazole these are know as 5 – nitroimidazoles

Spectrum of Activity

- It possesses a broad spectrum of activity.
- It is effective against trichomonads, amoebae and giardia and bacteria (anaerobic cocci and bacilli).

Mechanism of Action

It disrupts D.N.A. synthesis in protozoans and bacteria.

Pharmacokinetics

The oral bioavailability of metronidazole varies from 50-100%. If given with food absorption is enhanced in dogs.

- The absorption is due to increased of bile secretion that helps to dissolve the drug.
- Peak blood levels reaches in 1 hour of dosing
• Distribution is wide and rapid due to lipid solubility
• The drug is metabolized by glucuronide and several oxidation products that may darken the urine.
• Elimination half-life is 3-5 hours in dogs and horses.
• Excretion takes place in 24 hours, the drug's metabolite and unchanged from the drug are excreted in faces and urine.

Dose

• Canine giardiasis 25mg/kg po.IV or SC bid
• Equine trichomoniasis 20mg/kg by slow infusion
• Bovine trichomoniasis 75mg/kg IV bid
• Topical application 5% ointment plus urethral douche; to irrigate wound

The course of treatment is 5-7 days. When treating birds or rodents metronidazole is added in drinking water in amoebiasis.

Adverse Effect of Metronidazole

• Nausea, vomiting, abdominal cramp
• High doses in dogs may produce neurological disturbances characterized by tremor, weakness, muscle spasm, ataxia and convulsion
• Experiments in rats show that it is mutagenic if used for a prolonged time.
Other drugs used in gastrointestinal protozoan infections

Examples:

1) *Toxoplasma gondii* is treated using
   - **Sulfadiazine** (15-20mg/kg).
   - **Atavaguine** and **spiramycin** are used in difficult cases of toxoplasmosis
   - **Clindamycin** at 10-40mg/kg used in dogs 20-50mg/kg but in cats

2) **Amoebiasis** :- caused by *entamoeba hystolitica* is not common in animals but *E. invadei* in reptiles is treated using metronidizole at 10-25mg/kg bid orally for /52 or one week **furanolidine** 2-4mg/kg orally t/d

3) **Cryptosporidiosis**:- In neonates eg calves, kids, lambs piglets, it is usually caused by *Cryptosporidium paroum paromomycinsulphate*

**ANTICOCCIDIAL DRUGS.**

The major drugs used are classified as

- Sulphonamides
- Quinazolines
- Quinolones
- Symmetrical triazinones
- Thiamine antagonists
SULPHONSMIDES

The sulphonamides used are:

- Sulphadimethoxine
- Sulphaquinoxaline
- Sulphaclozine

Usually sodium salt of sulphadimethoxine (0.1%) or sulphaquinoxaline (0.02 per cent) is given in drinking water. Medication may continue for 3.5 days intermittently.

Precaution to avoid toxicity “the intermittent method” is preferred this consists two medical periods, each of **3 days-2 days apart another 3 days**, when normal food and drinking water is provided.

**Sulphonamide** preparations incorporating diaminopyrimidine potentiators are available for use in small animals (e.g. Sulphadimidine with *trimethoprim* or *ormethoprim*, *sulphadiazine* with trimethoprim).

LIMITATIONS OF SULPHONAMIDES IN TREATMENT OF COCCIDIOSIS

- None of the sulphonamides are broad spectrum for coccidiosis.
- They are previously not active against early asexual coccidian parasites.
Dosage

- In turkeys, achieved when 125ppm daily in food or water for 8weeks. Also treatment using 500ppm for 7days preferably in water.

In Rabbits

Prophylactic treatment is 250ppm daily in feeds as premix for 7days or treatment 1000ppm in water for 7days preferably in water

In Cattle

Prevention is achieved using 13mg/kg

Withdrawal Time in Days

- 28days before point of lay
- 75days before slaughter of animals.

Quinazolines

Example halofuquinone derived from febrifugine an extract of plant

- It is potent drug.
- It can be used in avian species
- Usually a steep dose – response curve is achieved when using the drug.

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Use:
Usually used in turkey and chicken

**Dosage:** chicken 3ppm in feed

Turkey the drug is same as chicken and should be used for 12 weeks.

**QUINOLONES.**

Examples:
- Decoquinate
- Methylbezoquate
- Nequinate

**Quinolones:** Their activity is essentially coccidiostatic used against invading sporozoites.

- Should be used in early stage or prophylactically the drug would not be effective if delayed.

**Mechanism of action.**

The quinolones selectively inhibit the electron transport chain in the *emeria* parasite.

**Use:**

Decoquimte could be used in food as premix or in water as powder for prevention of coccidiosis in broiler chickens.
Dosage:

Broilers: 20-40 ppm in food

Ewes: 100ppm for 28days

Cattle: 500ppm in feed

SYMMETRICAL TRIAZINONES

Example: Toltrazuril

This is a drug produced for its coccidiostatic property used against sporozoites

- It is also potent schizogony and gametogony
- In the use of the drug the drug is usually interpreted for 2-3 days of medication.

Spectrum of Activity

It is effective against *E. tenella*

It is used in turkeys, rabbits and chicken

Dosages

- Broilers 25ppm in drinking water
- Rabbit 10-15ppm

Contraindication

Poultry formation are different from rabbits except otherwise

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THIAMINE ANTAGONIST

e.g Amprolium

History: First used in the 60’s and was the leading drug until mid-70s

Spectrum of Activity

- It is used in confirmed *E. acervulina* and *E. tenella*
- When mixed with *ethopabate* it broadens its spectrum against *E. brunetti* and *E. Maxima*
- Sometimes used with sulphaquinoxaline to potential its activity
- It is used in chickens, rabbits and ruminants.

Dosage

- 125-150ppm in feed continuously
- 5mg/kg in water in calves for 21 days for treatment and used for 5 days for prophylaxis

Dosages with other drugs

125ppm Amprolium + 8ppm ethopabate or 100ppm amprolium + 5ppm ethopabate + 60ppm sulphaquinoxaline.
**Contraindication**

The drug “Amprolium” should not be used for a long time without using vitamin supplement, because it might predispose the flock to thiamine deficiency.

**OTHER ANTICOCCIDIALS**

**Pyridines**

Examples: clopidol

Clopidol does not allow natural immunity to develop

It is used in turkeys rabbits and chicken.

**Dosage**

In chicken 125ppm in feed

In rabbits 20ppm in feed

**Contraindications**

It should not be used with other drugs.

**Ionophores (polyether antibiotics)**

Examples: used in coccidials are

- Monensin
- Lasalocid
**Monensin** is produced from *strep cinnamomensis* Lasalocid produced from *strep lasaliensis*

**Dosage of Monensin**

- Chicken layer at 100 – 120ppm
- Turkey 100ppm
- Cattle 16.3 – 33ppm in feed
- Sheep 11-33ppm in feed

**Mechanism of Action**

Polyethers of monesin and lasalocid the polyethers interfere with transport of ions through membranes causing influx of positively charged ion cations this distorts the osmotic balance of the parasite so it dies.

Other anticoccidials are nitrobenzamides: eg dinitolmide, alkomid nitromide
REFERENCES


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