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**A FIELD STUDY OF SOME NEMATODE PARASITES  
OF BOVINES IN A SEMI-ARID AREA, WITH SPE-  
CIAL REFERENCE TO THEIR BIOLOGY AND  
POSSIBLE METHODS OF PROPHYLAXIS.**

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**SUMMARY.**

Preliminary field investigations revealed the presence of Cooperia pectinata, C. dunstata, Haemonchus placei, Oesophagostomum radiatum and Bunostomum phlebotomum as the most common parasites of bovines in the North Western Cape.

Experiments were carried out under field conditions, at Armoedsvlakte Research Station near Vryburg in the North Western Cape, on the ecology and epizootology of these parasites.

Possible methods of prophylaxis based on experimental observations are described, and the possible use of strategic drenching using anthelmintics which gave promising results is included.

**INTRODUCTION:**

**(1) North Western Cape.**

The North Western Cape, formerly part of Bechuanaland is a vast semi-arid cattle ranching area. The annual rainfall varies from 10 inches in the West to 20 inches in the East, occurring mainly from mid-summer to autumn with very little if any rain in the winter and spring. At Armoedsvlakte Research Station (27°S, 25°W, altitude 4000 ft.) near Vryburg C.P., the average annual rainfall is 431.0 mm. (16.9 inches) and marked diurnal temperature variations occur from a mean maximum of 19.8°C (67.6°F) to a mean minimum of -0.5°C (31.1°F) in July and from a mean maximum of 32.1°C (89.8°F) to a mean minimum of 16.3°C (61.3°F) in January. Mean Relative Humidities vary from 36% in September to 59% in March and sunshine hours from 8.5 to

9.9 hours. Rates of evaporation are high.

The area lacks surface water of any kind. Fountains are few and far between; rivers fail to run other than in times of flood for short periods. Boreholes, powered by windmills and Diesel pumps must supply all water requirements. Subterranean water supplies are very deep, often brackish and frequently only 2-3 gallons a minute can be pumped up from 400-600ft. or more underground. Frequently on farms 5,000 morgen<sup>M</sup> in extent there is only one water supply and cattle must walk miles from their grazing to the drinking troughs.

The veld is a mixture of sweet grasses mainly annuals and edible bushes and trees e.g. Gravia spp., Tarshomanthus major and minor, Acacia giraffe etc. etc. After the rains in summer and autumn, the grass makes a natural hay in the veld in the winter and the edible bushes provide a protein supplement. Once the rains start grass seeds germinate and the growth is phenomenal (up to 5" in a week).

The grazing is of good quality but due to annual droughts of at least six to nine months or longer, light stocking of 1 beast per 5, in the Eastern to 1 per 15 morgen in the drier Western areas is necessary.

## (2) Farming Methods:

There are two main methods of farming in the district and modifications thereof.

### (a) Dairy Ranching.

On these farms cows are milked for cream and in some cases, milk for cheese factories. Calves are confined to calf pens and kraals<sup>KK</sup> at birth and

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<sup>M</sup> 1 morgen = 10,000 sq. yards = approx. 2½ acres.

<sup>KK</sup> Kraals are cow yards, or cattle pens.

separated from their dams. These calf pens are small fenced off enclosures within or adjacent to the kraal, having a wire gate giving access to the kraal. Both calf pens and kraals have floors consisting of manure accumulated over many years.

Cows graze in the veld at night and return to the kraal in the morning from 7 to 10 a.m. Milking is commenced by removing the calf from its pen to the kraal and once it has started suckling, securing the cows hind legs with a leather thong (riem), pushing the calf away and milking 2 or 3 teats. Thereafter the calf is allowed to suckle and left in the kraal with the cow until 2 p.m., when they are separated the calf remaining in the kraal, the cow returning to the veld. Calves from 2 weeks to 4 months of age are allowed into a small camp at night to graze with the older calves.

(b) Beef Ranching.

Cows graze with the herd in the veld, calve there and are not separated from the calves until the calves are weaned. The herd grazes from the late afternoon to the following morning, and comes to the drinking trough from about 8 a.m. in the morning. The animals lie down near the water supply during the heat of the day and from 2 to 3 p.m. or later, drink again and return to the veld. In most cases cows are not milked on beef ranches.

A more detailed description of these farming methods has recently been given by du Toit (1958) who modifies this classification.

(3) History of Verminosis in the N.W. Cape.

Farmers have stated that many sheep were kept in Bechuanaland 20 to 30 years ago but the presence of

worms forced them to abandon sheep farming in favour of cattle farming.

In 1930 a severe outbreak of gastro-intestinal nematodes in bovines occurred at Arnoedsvlakte (Mönnig 1931). In 1939 to 1941 Fourie (1942) described outbreaks in the district. In the annual reports of the State Veterinarian (Vryburg), verminosis was considered serious in 1946, 1947, 1950 and 1951. In 1954 another massive outbreak of helminthiasis in cattle occurred. This reached a crisis in the spring and summer under conditions of severe drought. The State Veterinarian instituted a dosing campaign and the Stock Inspectorate Staff dosed 150,888 bovines with tetrachloethylene emulsion with excellent results. Prior to dosing stock died off at an alarming rate, but as soon as dosing was instituted stock losses stopped immediately; stock began filling themselves in 4-5 days and milk yields improved.

In view of the critical position the Director of Veterinary Services gave instructions that a full scale investigation of bovine verminosis be carried out in semi-arid areas with head quarters at Arnoedsvlakte. The work commenced in January 1955 and some of the investigations are reported in this paper.

#### REVIEW OF THE LITERATURE.

Veglia (1928) considered the rainfall of 7.87 inches in 1922 at Arnoedsvlakte to be insufficient for the development of Oesophagostomum columbianum larvae in the faeces to the infective stage. According to Mönnig (1931), P.L. le Roux in 1930 diagnosed Haemonchus contortus (= ?H. placei), Cooperia spp., Hydrocotyle phlebotomum and Oesophagostomum radiatum in cattle. At that time the

climatic conditions were those of severe drought. Mönning in this same article showed that there was sufficient moisture in a dung heap for larvae to reach the infective stage in the absence of rain. Furthermore he recovered larvae from the soil next to the dung heap and suggested that grass growing around and through dung heaps may carry a severe infestation of worms into the animal.

Fourie (1942) described outbreaks of Verminosis along the Malope River in the Mafeking and Vryburg Areas. After floods in January and February of 1941, pools and vleis had remained which he considered assisted the spread of verminosis. Furthermore, he suggested that sprouting green grass near dung heaps, while usually not grazed would be grazed during times of drought and that sandy soils and the prevailing phosphate deficiency may have facilitated the spread of verminosis.

The embryonated eggs and infective larvae of Trichostrongylus spp. are very resistant to desiccation (Mönning 1930, Crofton 1948b). Sprent (1946b) showed that desiccation was the most important inhibitory factor in the development of Bunostomum phlebotomum and in the dry season these larvae could not develop in pastures apart from permanent swamps. Seddon (1950) reported that wet conditions were not essential for the development of heavy infestations in sheep. Although pasture was dry and scanty, green grass could grow due to subterranean moisture. Leaking drinking troughs, dampness under troughs lying on the soil, frontages on streams etc. provided sufficient moisture for the spread of verminosis.

Roberts (1951) working in Queensland, Australia stated that verminosis was absent in areas with less than 30 inches of annual rainfall, except during abnormally

wet years. Later investigators confirmed Mönig's (1931) observations that sufficient moisture was present in dung for larvae to reach the infective stage (Roberts, O'Sullivan and Riek 1952). Once dung pads had hardened Roberts (1951<sup>1/2</sup>) stated that 5 inches of rainfall a month was necessary for H. contortus ( = ?H. placei) to escape from dung pads, but Cooperia spp. and O. radiatus needed less rainfall. However, in later investigations they found that under drought conditions, when livestock concentrated on small areas in the vicinity of the waterholes calves became infested with H. contortus ( = ?H. placei), Traxsi, Ostertagia ostertagi, and Cooperia spp. (Riek, Roberts and O'Sullivan 1953.) They stated that under drought conditions selective grazing could not take place, and the animals fed to soil level and right up to the dung pad. Under such conditions larval migration became less important, but some migration was essential and this could be assisted by heavy dews which occurred in the winter. The absence of Bunostomum phlebotomus infestation of calves led them to the conclusion that the distribution of this parasite was governed by rainfall, and it would not survive in areas with less than 24-25 inches annually.

From the above review it will be seen that some investigators (Veglia 1928, Sprent 1946b, Roberts 1951, Roberts et al 1952) mentioned that verminosis could not be spread under drought conditions; other workers (Mönig 1931, Fourie 1942, Seddon 1950, Riek et al 1953) showed that verminosis did occur under severe drought conditions; the reasons for this occurrence were not investigated. In the North Western Cape such arid conditions occurred and it would appear that certain species of worms had found local conditions in which they were able to thrive and cause serious infestations of stock.

## PRELIMINARY INVESTIGATIONS.

Before embarking on experimental work at Armoedsvlakte it was essential that the species of worms in cattle in the area be diagnosed; the prevalence of these species under different methods of animal husbandry be investigated attempts made to ascertain at what season and age stock become infested, and which age groups, suffered most from the effects of verminosis.

### Species of Helminths Concerned.

The following species were diagnosed at post mortems on farms in the district, at Armoedsvlakte and at the Vryburg Abattoir:-

(1) Haemonchus placei (Place 1893) Ransom 1911.

The name H. placei has recently been resuscitated by Roberts, Turner and McKevett (1954) who have shown that this species in bovines is morphologically distinct both in adults and infective larvae from Haemonchus contortus (Rud. 1803) Cobb 1898 in ovines. The parasites found by the author at post mortems and the infective larvae were compared with Roberts et al descriptions and were found undoubtedly to be H. placei; this name will in future be used when referring to this parasite in bovines.

(2) Cooperia pectinata Rans. 1907.

(3) Oesopharostomum radiatum (Rud. 1803.) Raill. 1898.

(4) Bunostomum phlebotomus (Raill. 1900) Raill. 1902.

These four species were the most commonly recovered in order of their prevalence. Frequently, however, Cooperia pectinata was present in larger numbers than



Haemonchus placei, Cooperia punctata (v. Linst. 1907) was occasionally present and on rare occasions Trichouris globulosa (v. Lins. 1901) Rans. 1911. The only Cestode of note was Moniezia benedini (Moniez 189) which was fairly prevalent in calves. Trematodes were never recovered.

#### Field Observations:

Farms scattered over an area 200 miles from East to West and 280 miles from North to South were visited. Post mortems were carried out where farmers were prepared to sacrifice stock and faeces collected from different age groups. Notes were made on the size of the farm, stocking rate, available water supplies and systems of animal husbandry etc.

It soon became evident that calves and weaners were more heavily infested than adult stock, with the odd exception of the old cow. After weaning young stock lost condition rapidly and only improved at about 18-24 months of age after the change of teeth if suitable grazing was available.

On dairy ranches calves were more heavily infested than on beef ranches. Even calves confined to calf pens and kraals were severely infested with a mixed infestation of parasites. On beef ranches Cooperia spp. and H. placei were present with little other infestation. The infestations were usually milder than on the dairy ranch, and calves were in better condition. On farms where dairy ranching was practised, with as little as 7-10 inches annual rainfall, stock were more heavily infested than their counterparts on the beef ranch, under the same climatic conditions.

#### Helminthiasis at Arrodavilakte.



This research station consisted of two farms 11 miles apart known as Armoedsvlakte, 3,800 morgen and Biesjesvlakte, 3,200 morgen in extent stocked with approximately 1,000 head of cattle. Water supplies from boreholes were plentiful and both farms well divided into camps. The former was a dairy ranch stocked with Red Poll and Fries, the latter a beef ranch stocked with Afrikaner and Sussex cattle. Calving took place from the 1st of November to the 31st of January. At Armoedsvlakte calves were separated at birth from their dams, and at the twice daily milkings allowed two teats on which to suckle; the other two were milked. Cow byres, kraals and calf pens had concrete floors which were cleaned regularly. Three small camps were set aside for the calves and grazing alternated irregularly according to the state of the pasture. These calves had a mild infestation of Coonaria spp., H. placai and O. radiatum.

Biesjesvlakte on the other hand was run as a beef ranch. Cows calved in the veld and ran with the dams until weaning. The dams were herded daily into kraals and crush pens to be dosed with bone meal<sup>M</sup>. During the breeding season, from February to April cows were herded into small kraals to be served and left with the respective bulls for three hours every morning. The calves accompanied the cows both to the crush and kraals. The calves were more severely infested, than their counterparts at Armoedsvlakte, not only with the same species but with H. phlebotomus as well. Diametrically opposed results were observed when comparing the different worm burdens of stock on the farms with those on the experimental station.

<sup>M</sup>

Note: Due to the prevailing phosphate deficiency in the grazing all stock except suckling calves were dosed with bone meal every day.

The preliminary investigations had shown what species of nematodes were common in the district, and that calves and weaners were more heavily infested than adult animals.

### E C O L O G Y.

Before any rational means of prophylaxis could be attempted, the ecology and epizootology of these parasites had to be investigated.

#### A. HATCHING OF EGGS AND DEVELOPMENT OF LARVAE TO THE INFECTIVE STAGE UNDER FIELD CONDITIONS.

##### Materials and Methods -

- (1) Faeces were collected per rectum from 6 or more cattle suffering from a mixed nematode infestation. The faeces were well mixed and the number of eggs per gram established by taking the mean of two or more separate specimens, using Roberts and O'Sullivan (1950) technique with one modification. A 40% sucrose solution was used instead of NaCl solution. It was noted that sucrose solution gave better egg counts, and eggs were easier to see, than was the case with salt solution. More air bubbles were formed however with sucrose solution than with salt solution. This was overcome by adding a few drops of amyl alcohol as advocated by Roberts <sup>et al</sup> (1951).
- (2) Seventy five grams of the dung was used for a control culture which was incubated for eight days at 26°C.
- (3) The remainder of the dung was divided into equal quantities of 400 grams each; in two experiments, however, quantities of 1 Kg. each were used. These specimens were shaped into heaps similar to cattle dung pads and placed in a row next to each other in a paddock which had been ungrazed for years. They

were placed on the bare ground and the surrounding grass cut off to a height of 5 cms. In a few experiments the dung pads were placed in the shade of a large bush. Due to activities of dung beetles during summer it was found necessary to protect dung pads with wire gauze (mosquito netting) cages. This allowed free passage of air light and rainfall and only termites and ants could get at the dung which did not have much affect on the dung itself.

- (4) Dung pads were collected at intervals, weighed and the dung broken into small pieces and well mixed; sufficient dung was taken for egg counts and the remainder placed in the Baermann Apparatus for 24 hours. The maximum weight placed in any funnel was 50 grams, so that the thickness of dung did not exceed 1.5-2 cms.; poor results were obtained when this thickness was exceeded.

When the outer crust and the depth of the dung were examined separately, dung pads were weighed as before, the crust and depth separated, both weighed separately and kept apart for the rest of the examination.

- (5) After 24 hours 50 cc of fluid was tapped from the bottom of the funnels, the dung removed, mixed with dry sterile cattle faeces and incubated for eight days at 26°C. The larvae recovered from the funnels were allowed to settle in the collecting tubes for two to three hours, the supernatant fluid siphoned off and the sediment examined for larvae in the counting chambers of the Roberts and O'Sullivan (1950) counting slides. If the larvae were present in large numbers

in the sediment a dilution count technique was employed and the total number of larvae in 2 or 4 cc of a 50 or 100 cc dilution were counted in the counting slides as follows:

The larval suspension in the dilution tubes was shaken, and 0.5 cc pipetted into one of the chambers of the counting slide. The process of shaking and pipetting was repeated until all the chambers had been filled. A drop of sterile water was added to each chamber to give convex menisci on the outer edges of the fluid on the verandah of the counting chamber. If infective larvae were present they tended to swim towards the menisci and microscopy was facilitated thereby.

The chambers were examined microscopically with the low magnification (30 diameters) and larvae counted. As many larvae as were necessary were transferred from the sediment in the dilution tube on to glass slides heat killed and the species of infective larvae identified on a percentage basis. No attempts were made to identify the species of preinfective larvae and they were merely counted.

- (6) Larvae from control cultures and incubated dung were collected as follows:

A Baermann funnel was filled with water up to 2 inches from the brim. The lid of the culture jar was washed into this funnel to remove any larvae present. The culture jar was filled with water at 40°C, and a piece of plastic sheeting placed over the mouth of the jar, held firmly in position with the hands, the jar turned upside down and placed in the Baermann funnel containing water. The jar was then picked up slightly, the plastic sheeting gently removed and the jar allowed

to reseal itself in the funnel. While the plastic sheet was being removed care was taken that the mouth of the culture jar was below the level of water in the funnel. The plastic sheet was washed into a beaker to remove any larvae adherent to it and the beaker's contents poured into the funnel next to the culture jar.

The larvae were collected 20-24 hours later, by tapping the lower 50 cc into test tubes. Thereafter the larvae were examined in the same fashion as already described for larvae recovered from dung with the Baermann Apparatus.

(7) Prevailing - climatic conditions.

The daily variations in the climatic conditions were recorded for the experimental period.

EXPERIMENTAL OBSERVATIONS.

The observations are summarised in Table No. 1, in the appendix and unless otherwise stated wherever experimental numbers are referred to the results will be found in this table. A series of 22 groups of dung pads were placed in the veld from 15.8.56 to 3.6.57. Some of the data shown in the table were unreliable and have not been taken into account for the following reasons:-

- (1) Control culture of No. 3 and incubated culture from dung in No's. 1(e), 5(e), 21(e), 22(d) were not made.
- (2) Dung pads were damaged by dung beetles in Nos. 7, 9, 10 and 11.
- (3) Original egg per gram counts of Nos. 17 and 18 were obviously incorrect.

(A) Hatching of eggs in the field.

From September to March all eggs hatched within 4-8 days; on the average more than half the

eggs (61%) hatched within two days and nearly all

the eggs (87.8%) had hatched by the sixth day. In August all the eggs hatched within 8 to 9 days, but from April to June, with one exception, 58-96.3% of eggs hatched after dung had been in the veld for periods of 2-3 weeks. In this one exception all eggs hatched within 13 days in May. (No. 19b).

The rate of hatching is summarised in Table No. 1 below.

TABLE NO. 1.

Rate of Egg Hatching in Relation  
to Atmospheric Temperatures.

Percentage Hatched	Time in days	Maximum Temperatures		Minimum Temperatures	
		Mean °C	Range °C	Mean °C	Range °C
100%	4 - 8	31.1	28.4-32.6	12.8	10.1-15.2
100%	9 - 13	24.3	22.7-26.7	2.6	2.6-3.6
58-96.5%	14-21	22.6	18.0-27.2	3.2	1.5-4.4

At higher temperatures eggs hatched rapidly; at lower temperatures, however, there was a certain amount of inconsistency.

This was probably due to-

- (1) Marked diurnal fluctuations in temperature, particularly in the winter when differences of 21°C between the daily maximum and minimum temperatures were not unusual.
- (2) Thermograph readings give a truer picture of temperature fluctuations through the day than the mere recording of maximum and minimum temperatures. In the summer, temperatures at Arnoedsvlakte remained high for long periods and only fell below 20°C for a few hours i.e. from about 11 p.m. to sunrise. In the winter, however, temperatures rose above 15°C from about 10 a.m. to 4 p.m. and remained at a low level for the rest of the day; whereas in summer tempera-

tures exceeded  $20^{\circ}\text{C}$  for 18 hours every day, only for 6 hours daily did temperatures exceed  $15^{\circ}\text{C}$  in midwinter i.e. only from 10 a.m. to 4 p.m. During summer temperatures fell below  $20^{\circ}\text{C}$  for only 6 hours per day; in winter temperatures below  $15^{\circ}\text{C}$  prevailed for 18 hours daily. The inconsistencies of the rate of egg hatching in the winter is explained by the fact that temperatures were much lower for longer periods.

Eggs hatched more quickly in the sunlight than in the shade. In the winter dung pads were placed in shade or exposed to sunlight, but were otherwise identical. Those in the shade hatched and larvae developed more slowly than those in the sun (compare No. 19 with No. 20 and No. 21 with No. 22); Ground temperatures were frequently  $2-3^{\circ}\text{C}$  higher in the sunlight than in the shade.

Eggs hatched more rapidly in the crust of the dung pads than in the deeper lying parts. Whenever the crust and depth were examined separately fewer egg counts were recorded in the former than in the latter (Nos. 16(d-j) 19a, 20-22). This indicated more rapid hatching of eggs in the crust. Temperatures in the depth were often  $1-2^{\circ}\text{C}$  lower than in the crust.

Since evaporation of water was more marked from the outer surface of the dung, air was probably more plentiful in the crust and possibly assisted in egg hatching.

(B) The morphology of pre-infective larvae.

It was very important to distinguish the larval stages of these parasites from free-living worms. Pure cultures of the different species were examined microscopically, at regular intervals after incubation of eggs.



recovered from adult gravid female worms.

Little difficulty existed in distinguishing infective larvae from free-living Nematodes. These have been adequately described by Keith (1953). The pre-infective larvae however, were considerably more difficult and the following salient points were used to distinguish them from free-living Nematodes:-

(a) First stage larvae.

Morphologically the larvae belonged to the rhabditoid type. The form of the body was cylindrical, decreasing in size from the base of the oesophagus, gradually to the tail and to a lesser extent to the head. The length of all the species of larvae fell in the range of 292-500 microns; the maximum breadth at the base of the oesophagus was 19-21 microns.

The main point of differentiation was the oesophagus and the best description the author encountered in the literature, of this differentiation has been given by Loos (1911) in his epic work on Ankylostoma duodenale.

The description on page 349 of his publication is quoted:-

"The oesophagus presents division into three sections which is characteristic of the Rhabditidae. This fact has already been pointed out by the earlier writers; but in the Ankylostoma larvae the three sections are not nearly so sharply marked off, as in the Rhabditidae. The anterior section is more uniform in thickness (ca 0.009 mm) throughout its length, not distinctly conical as in the free living Nematodes, or even the non-parasitic generation of Strongyloides stercoralis. The transition to the thinner part takes place fairly gradually, as does the broadening to the terminal bulb, which is on the whole more slender and weaker than Rhabditidae. If these



differences are observed, confusion between young Ankylostoma larvae and those of Strongyloides stercoralis should hardly be possible."

Similarly Mönig (1926) made the observation that the first stage larvae of T. rugatus and T. instabilis had; "An oesophagus which is of the usual rhabditiform shape, but that the three regions are not as sharply marked off as e.g. in the larvae of Strongyloides papillosis."

These observations were confirmed. First stage larvae of the species of parasitic worms studied had an oesophagus of the rhabditoid shape but the three regions were not as sharply marked off as those of the larval stages of free-living worms. This information combined with the size and shape of the larvae described above facilitated in making a diagnosis.

(b) Second Stage Larvae.

These were more easily differentiated from the free-living nematodes than the first stage larvae. Second stage larvae of parasitic worms were cylindrical in shape, of equal diameter from the base of the oesophagus to the anus (20-21 microns); narrowed sharply to the tail. and the sheath of the tail, which was similar to the infective larvae particularly when the development of the second stage was more advanced. The larval length depended on the stage of development and varied with the individual species the maximum in B. phlebotomus being 500 microns, while the other species exceeded 700 microns.

The oesophagus became elongated, the corpus and isthmus more difficult to delineate and the bulbous more spatulate than the first stage. The cellular elements of the intestine became opaque usually being a dark brown colour. The larvae were more active.

The species of larvae of the parasitic worms were only differentiated from each other in their infective stages, for

counting purposes the pre-infective stages were differentiated from free-living worms.

(C) The development and survival of larvae.

Larvae developed rapidly to the infective stage from August to April being recovered after a minimum period of 5 days and more regularly within 8 days. In early winter they were recovered after 14 days but had not yet developed in mid-winter after 21 days.

TABLE NO. 2.

Mean percentages of larvae recovered  
from August to April.

No. of days in the field	Mean percentages recovered from field specimens.			Mean percentages recovered after incubation.
	1st Stage	2nd Stage	3rd Stage	3rd Stage
1	0.32%	-	-	30.7%
2	0.69%	0.07%	-	15.8%
3	2.96%	0.63%	-	19.8%
4	1.18%	1.08%	-	6.1%
5	1.9%	4.13%	0.52%	5.6%
6	0.8%	1.47%	1.03%	2.3%
7	0.21%	0.68%	4.74%	2.3%
8	0.5%	1.00%	1.40%	3.8%
9	0.41%	0.77%	2.79%	1.2%
30	-	-	1.01%	0.5%

The difference between the numbers of eggs per gram as revealed by egg counts on fresh faeces, and the numbers after various periods in the field was considered to represent the numbers of eggs which had hatched i.e. the number of larvae in the dung. The mean percentages of these larvae as recovered with the Baermann Apparatus and again after a period of 8

days incubation is shown in Table No. 2.

This table is of particular interest in that it reveals the very small percentage of larvae which reach the infective stage under field conditions which was probably due to a very high mortality of the pre-infective larvae. From the table it will be noted that first, second and third stage larvae were at their optimum on the third, fifth and seventh days respectively. Further that after collection of larvae with the Baermann Apparatus, the incubation of dung indicated that there was a dramatic drop in the percentage of infective larvae recovered after three days in the field.

The Baermann Apparatus is not as efficient in extracting pre-infective as infective larvae and this may account to some extent for the poor recoveries of the farmer. When this extracted dung was incubated both unhatched larvae and unrecovered pre-infective larvae were given ample chance to develop to the infective stage and these were then more easily recovered.

Egg hatching and the development of larvae to the infective stage were slower and more protracted in winter. In May and June first stage larvae were recovered from the fourth to the twenty-first days; second stage larvae from the fourth to the twenty-sixth day and infective larvae were not yet present even after twenty-one days. The number of specimens examined in winter was not sufficient to establish when the pre-infective larvae reached optimal development.

(D) Climatic factors affecting survival of larvae.

(1) Evaporation.

Due to lack of equipment evaporation could not be measured in the standard fashion. Dung pads were protected from the activities of beetles and since they were weighed before placing in the field and at the time of collection, the weight loss was probably due to evaporation.

This is expressed as a percentage in Table No. 3, in which three experiments were selected to indicate the effect of evaporation and other climatic factors, on the percentages of hatched larvae recovered from dung pads exposed to field conditions. The poor larval recoveries in experiment No. 14 when compared with No. 12 was due to desiccation of the larvae, brought about by water loss from the dung pad. In the period August to April if dung pads lost 65% of their weight by evaporation in the first 6 days in the field less than 0.27% of the hatched larvae reached the infective stage. The higher moisture content contributed markedly to the better larval recovery rates in Nos. 12 and 21 but it was noticeable that larval recoveries were falling off after 3 weeks in the latter when evaporation caused 62.5% decrease in weight of the dung.

In experiments summarised in Table No. I (appendix) it was marked feature that larval recoveries from the crust of the dung were poor in spite of more rapid egg hatching there (Nos. 16(d-j) 19a, 20-22). The crust was drier than the depth due to evaporation and probably accounted for the poor larval recovery rates due to either death of the larvae or possibly migration of the larvae to the moister depth of the dung pad.

## (2) Temperature.

In the period August to April infective larvae were recovered within 5-8 days. In June they were not yet present in dung after 21 days; if Table No. I (appendix) is consulted it appears that the infective larvae were not only conspicuous by their absence, having not yet developed after 3 weeks, but that specimens in the shade showed lower larval recovery rates than those in the sun, due to colder conditions there. In No. 20a (Table I) the mean maximum was 22.0°C and the mean minimum 2.8°C

TABLE NO. 3.

Climatic factors affecting the percentage of larvae recovered from hatched eggs.

Experimental No.	No. of days in field.	Total percentage of larvae recovered	Percentage weight loss of dung pads due to evaporation	Mean Maximum Temperature.	Mean Minimum Temperature.	Mean Relative Humidity.	Total Rainfall.	Days on which rain fell.
				°C	°C	%	M.M.	
12	2	2.9%	36%					
b	3	5.4%	34%				24.5	3rd
c	4	8.0%	30%	31.5	16.7	37.5	3.5	4th
d	5	9.3%	49%					
e	6	3.2%	47.5%					
f	7	14.9%	37.5%				21.5	7th
14	2	0.66%	50%					
b	4	0.11%	62.5%	33.1	12.7	41	0	0
c	6	0.02%	79%					
d	8	0.02%	84%					
21	4	1%	42.5%					
b	8	2.17%	42.5%				1.4	6th
c	10	0.02%	35%	18	1.5	65	21.6	9th
d	14	1.01%	51%					
e	21	0.33%	62.5%					

Nos. 12 and 14, January and February respectively.

No. 21 June.

over a period of 20 days and infective larvae had not yet developed. The maximum temperatures should have been adequate for larval development to the third stage. However as previously mentioned when egg hatching results were being described, temperatures in the winter remained below 15°C for most of the day and higher temperatures applied only for the briefest periods. Thus both egg hatching and larval development could proceed for short periods only in the winter.

Conversely higher temperatures applied for longer periods in the summer and probably accounted for the rapid development to the infective stage.

The few larvae recovered in No. 21 (Table No. 3) appeared to be due, therefore, to a combination of low temperatures, retarding development and the slowly increasing evaporation causing larval death by desiccation.

### (3) Relative Humidity.

There appeared to be little correlation between relative humidity of the atmosphere and larval survival if the results of Table No. 3 are consulted.

### (4) Rainfall.

In the summer the best recoveries of larvae occurred when rain fell during the first 5 days dung was in the field. The distribution seemed to be important. Optimal results were obtained when rain fell on the 3rd and 4th, or 2nd and 5th days followed by a further fall of rain (No.'s 8, 12, 13 and 15 Table No. I - appendix).

The absence of rain probably accounted for the poor results in No. 14 (Table No.3).

In autumn this did not appear to be so important, since rain only fell after 9 days and larval recoveries were reasonable (No. 16 Table No. I - appendix.)

In winter in spite of rainfall falling in the first 10 or 11 days larval recoveries remained at a low level

(Nos. 19, 20, 21, 22 Table No. I - appendix and No. 21, Tab. 3).

In spring rainfall on the first and third day, but none thereafter, assisted larval development to the sixth day but thereafter larvae decreased rapidly (No. 4, Table No. I - appendix.) Rain falling for the first time on the sixth day in spring was too late to have any effect on the survival of larvae (No. 6, Table No. I - appendix.)

(5) **Shade and Sunshine.**

Unfortunately this was only tested in the winter and eggs and larvae developed more rapidly in the sun than in the shade due to higher temperatures in the sunlight. Experiments described elsewhere (vide infra) however showed better results in the summer with specimens in the shade than those in the sun as far as the recovery of infective larvae were concerned.

Temperatures and rates of evaporation were thus intimately linked with shade and sunlight, the former lowering the latter exaggerating the effects of both temperature and evaporation.

(E) **Effects of environment on the different species.**

The effects of exposure to field conditions on the different species dealt with are summarised in Tables No. 4 and 5. The percentages of infective larvae recovered, both from dung in the Baermann Apparatus and after further incubation, were averaged out in 7 experiments in the period September to April and compared with the mean percentages of the control cultures and are summarised in Table No. 4. Similarly larvae recovered from 5 experiments in the winter (May and June) were averaged out and compared with the mean of the controls and summarised in Table No. 5.

On examination of Table No. 4 species exposed to field conditions in dung pads showed little change after 2 days, but



marked changes occurred after 4 days. Cooperia spp. showed a marked increase while H. placei decreased. This tendency continued until the 8th day, while O. radiatum and B. phlebotomum remained at almost the same level as the original controls.

In Table No. 5, Cooperia spp. and H. placei acted in the same fashion, but B. phlebotomum decreased rapidly and O. radiatum not so markedly over a 14 day period when the larval species were compared with the controls.

T A B L E N O. 4.

The effects of exposure to field conditions from spring to autumn on different species (see text.)

Days in Field	Mean larval species variations.			
	<u>Cooperia spp.</u>	<u>H. placei</u>	<u>O. radiatum</u>	<u>B. phlebotomum.</u>
Controls	46%	3%	12%	7%
2	44%	33%	13%	10%
4	57%	20%	14%	9%
6	60%	15%	11%	14%
8	72%	3%	11%	9%

T A B L E N O. 5.

The effects of exposure to field conditions in winter on different species. (see text.)

Days in Field.	Mean larval species variations.			
	<u>Cooperia spp.</u>	<u>H. placei</u>	<u>O. radiatum</u>	<u>B. phlebotomum.</u>
Controls	36%	44%	13%	7%
4	47%	37%	16%	4%
8	62%	20%	15%	3%
10	59%	23%	15%	3%
14	68%	27%	6%	1%



Before further elaboration of these results two points had to be cleared up:-

- (1) Infective larvae may have migrated from the dung pads before they were collected. This was not possible before the fifth day, because no infective larvae were recovered before five days exposure. As shown later on, migration could only occur after the fifth day provided adequate rain had fallen. This was possible in only two of the seven experiments, from which the data in Table No. 4 were calculated. The rainfall, however, fell only on the 7th and 8th days respectively in sufficient quantities to cause migration. By this time the species fluctuations were already marked and would not have materially affected the results.
- (2) The Baermann Apparatus was possibly more efficient in the collection of pre-infective larvae of H. placid than of the other species. Since the pre-infective larvae were merely counted and not identified this could be presumed. There was, however, no evidence to prove this. The low efficiency of the Baermann Apparatus in collecting pre-infective larvae is shown by the usually larger numbers of infective larvae recovered from incubated dung, after it had been in the Baermann Apparatus, than could be accounted for by the unhatched eggs present in dung exposed for a few days in the field. Furthermore in critical tests there was no evidence that infective H. placid larvae were more efficiently collected than the other species in the Baermann Apparatus. It was, therefore, unlikely, that less active pre-infective larvae would exhibit this tendency. There is, therefore, little likelihood that the results shown in Tables 4 and 5, are false reflections of the effects of environmental conditions on the pre-infective stages of the life cycle.

It appeared that Cooperia spp. were best adapted to the environment. In the summer pre-infective stages of O. radiatum and B. phlebotomum, <sup>were</sup> more resistant than H. placei (Table No. 5). However, in the winter B. phlebotomum particularly, rapidly decreased even after 4 days in the field whereas O. radiatum could withstand cold conditions for 10 days and showed a sharp drop by the 14th day in the field. Haemonchus placei appeared to be more resistant to cold, but Cooperia spp. again are predominant.

(6) Climatic Conditions:

Daily variations in the climatic conditions from 15.8.56 - 23.6.57 are shown in Fig. 2. Climatic data were important in these experiments while eggs were hatching and larvae developing to the infective stage. In summer hatching and development were completed in 7-10 days but in winter this process continued for longer than three weeks. The climatic data for the various experiments are summarised in Table No. 6: periods of 6-10 days from spring to autumn and 14-26 days in winter were used, to show prevailing climatic conditions, during periods of egg hatching and larval development.

TABLE NO. 6.

Prevailing Climatic Conditions during  
the periods of eggs hatching and lar-  
val development.

Experimental No.	Days in Field.	Mean Maximum Temperature.	Mean Minimum Temperature.	Mean Relative Humidity.	Total Rainfall	Days on which Rain fell.
		°C	°C	%	M.M.	
1	9	26	2.6	35	0	0
2	8	27	3.0	33.5	0	0
3	10	26.8	3.5	36	0	0
4	9	23.7	5.7	49	10.9	1st, 3rd.
5	9	23.4	3.6	33.4	0	0
6	6	31.6	12.6	28.5	1.5	6th
<sup>m</sup> 8	7	28.9	15.6	59.5	24.8	1st, 2nd, 3rd, 4th, 6th, 7th.
<sup>m</sup> 12	7	31.5	16.7	37.5	49.5	3rd, 4th, 7th.
13	8	31.3	16.7	39	51.3	3rd, 4th, 7th, 8th.
14	8	33.1	12.7	41	0	0
15	9	31.8	15.2	52	23.5	3rd, 4th, 8th.
16	9	24.8	4.4	43	0.3	9th
<sup>o</sup> 17	14	25.1	3.5	52	9.7	9th, 10th, 11th.
18	26	23.3	3.8	52	23.0	18th, 19th.
19	19	22.7	2.2	51	23.0	18th, 19th.
20	20	22.0	2.8	52	23.0	18th, 19th.
21	21	18.0	1.5	65	23.0	6th, 9th.
22	21	18.0	1.5	65	23.0	6th, 9th.

<sup>m</sup> Experimental Nos. 7, 9, 10 and 11 not included - due to affect of dung beetle activity.

<sup>o</sup> Nos. 17 - 22 were carried out during the winter, thence the longer periods.

## DISCUSSION.

The results shown on the previous pages and summarised in Table No. I (appendix) were generally poor. The climatic extremes under which these experiments were conducted were a marked feature of the investigations. While the different species of worms were undergoing the free-living stages of their life cycle in dung they were subjected to marked diurnal temperature variations, the mean relative humidities were generally low, rainfall was either absent, sparse or fairly well distributed and adequate (Table No. 6). In addition high rates of evaporation were common when dung lost up to 70% of its moisture in less than a week, particularly in spring and summer. It was, therefore, not surprising that results varied from one extreme to the other but were generally poor and indicative of unfavourable environmental conditions.

Temperature appeared to play an important role in egg hatching and larval development. It has been stated by Dinaburg (1944) that H. contortus eggs and larvae did not develop below a mean maximum temperature of 65°F (18.3°C) in the U.S.A. In Australia Gordon (1948) stated that a mean maximum temperature 63.9°F (17.7°C) was essential for larval development of this parasite. A similar temperature was given by Roberts et al (1952) for H. placei of cattle. Crofton (1948b) has shown that all eggs of T. raptorisformis parasitic in rabbits will hatch within 8 days at temperatures of 55°F (12°C) or more.

My findings did not confirm these observations. At mean maximum temperatures of 25.1°C (77.2°F) infective larvae took 2 weeks to develop (No. 17 Table No. I - appendix.) At a mean of 18°C (64.4°F) and even 22°C (71.6°F) larvae had not yet reached the infective stage, nor had all the eggs hatched after 21 and 20 days respectively (Nos. 21, 22 and 20 Table No. I); the mean maximum temperatures, however, were on

an average  $16.5^{\circ}\text{C}$  to  $21.6^{\circ}\text{C}$  higher than the mean minimum temperatures, (Table No. 6, Experiments 17 - 22.)

Recently work in Kenya has shown that H. contortis will develop to the infective stage in 8 to 12 days if the mean maximum air temperatures are  $74^{\circ}\text{F}$  ( $23.4^{\circ}\text{C}$ ) or more and the mean minimum temperatures are not less than  $52^{\circ}\text{F}$  ( $11.1^{\circ}\text{C}$ ) (Dinnik and Dinnik 1954-55). These workers pointed out that in the U.S.A. and Australia, which are countries with generally temperate climates, the temperatures limit of  $64^{\circ}\text{F}$  ( $17.8^{\circ}\text{C}$ ) can generally be applied, but this was not the case in the Kenya Highlands where there were marked diurnal temperature fluctuations. This was more in agreement with the observations at Armoedsvlakte than with those of the Australian, American and British workers.

In experiment No. 1, the mean maximum temperature was  $26^{\circ}\text{C}$  and mean minimum temperature  $2.6^{\circ}\text{C}$  whereas in experiment No. 20 the mean maximum and minimum temperatures were  $22.0^{\circ}\text{C}$  and  $2.8^{\circ}\text{C}$  respectively (Table No. 6). In the former larval development was completed in 8 days whereas in the latter no infective larvae were present after 20 days, (Table No. I - appendix). There are several possible reasons for this.

- (a) Thermograph readings in the winter showed prolonged periods of 16-18 hours a day when air temperatures were below  $15^{\circ}\text{C}$ . In spring thermographs showed a more even distribution of maximum and minimum temperatures.
- (b) Grass minimum temperatures were  $3-5^{\circ}\text{C}$  lower than minimum air temperatures in the winter, but seldom more than  $2^{\circ}\text{C}$  lower in spring.
- (c) Temperatures in the depth of dung were usually  $2^{\circ}\text{C}$  lower than temperatures at the surface of dung.

In spring, therefore, warmer conditions prevailed for a longer period and heavy frosts were infrequent; egg hatching and larval development was thus able to proceed for almost 12 hours a day. In the winter enough warmth for larval

development to proceed existed for only 6-8 hours a day on the dung's surface and for even shorter periods in the cooler depth of the dung.

Actual freezing of first and second stage larvae probably also occurred during the heavy winter frosts. This freezing probably killed pre-infective larvae as was shown for other species by Ransom (1906); Veglia (1915); Schwartz (1924) and Mönning (1930.) In the winter grass minimum temperatures of  $-7.3^{\circ}\text{C}$  ( $18.8^{\circ}\text{F}$ ) were recorded thus giving some idea of the severity of the frost.

At the other extreme all stages of the free-living life cycle were completed in 5 days when the mean maximum temperature was  $32^{\circ}\text{C}$  ( $89.6^{\circ}\text{F}$ ) and the mean minimum  $16.8^{\circ}\text{C}$  ( $62.3^{\circ}\text{F}$ ). Many other workers with a wide variety of parasitic nematodes have found rapid egg hatching and larval development at high temperatures (Theiler and Robertson 1915; Veglia 1915; de Blicck and Baudet 1926; Ortlepp 1925 and 1937; Mönning 1930; Crofton 1948(b) etc.) In the summer months minimum temperatures were only approached for a few hours in the early morning, and temperatures stayed above the minimal level for larval development throughout the rest of the day.

Drying and desiccation has been shown to be a major cause of larval death, particularly of the pre-infective larval stages (Veglia 1915 and 1923, Mönning 1930, Ortlepp 1937; Crofton 1948b; Dinnik and Dinnik 1954-55 etc.) Dung pads that lost more than 65% of their moisture in the first 5-7 days in the field, gave poor larval recoveries; in the period spring to autumn should rapid desiccation have occurred with no rainfall only 0.27% or less infective larvae were recovered. Eggs hatched during this period but few larvae were able to reach the infective stage. First and second stage larvae reached optimal development on the third and fifth days respectively. If moisture was inadequate during this period

larval death was very marked. Observations that enough moisture exists in a dung pad for larvae to reach the infective stage have been made by various workers (Mönnig 1931; Roberts 1951; Roberts et al 1952 and Rick et al 1953). These workers, however, did not show what percentage of hatched larvae reached the infective stage, although Mönnig only recovered a few larvae. This was probably also due to mortality of pre-infective larvae.

When the effect of evaporation was decreased by rainfall particularly when pre-infective larvae were present in large numbers (i.e. 3rd - 5th day), the percentage of larvae which reached the infective stage was considerably increased. When rates of evaporation were high the best results were obtained when on at least one, but preferably on two separate days, rain was recorded during the first five days in the field; optimal results being obtained when rain fell from the second or third day onwards.

In winter, due to extremely low temperatures, in spite of much lower rates of evaporation, pre-infective larvae seldom developed to the infective stage. The effects of desiccation were also observed in winter, where although egg hatching occurred more rapidly in the crust of the dung, fewer larvae were recovered there than in the moister depth of the dung pad. This may be due to either mortality of pre-infective larvae or their migration to the moister deeper layers.

Some interesting observations were made on reactions of the different species to the environment.

The pre-infective stages of Cooperia spp. were best adapted to withstand environmental extremes of heat and cold, and dry and comparatively dry conditions. Haemonchus placei on the other hand was much less resistant to heat, cold and arid conditions. In summer its pre-infective stages decreased even more rapidly than those of G. radiatus and B. phlebotomus but during the cooler winter months conditions were more



favourable and this species was able to maintain itself better. Once it had reached the infective stage however, it certainly was more resistant than either Q. radiatum or R. phlebotomum. This was also confirmed in experiments described elsewhere.

Gasonbagostomum radiatum while maintaining itself fairly well in the summer months was apparently sensitive to cold conditions and fewer larvae were able to reach the infective stage, in experiments conducted over periods of longer than 10 days in the winter, than the larvae of Cooperia spp. or H. placei.

Banostomum phlebotomum maintained itself fairly well in the warmer months of the year but was sensitive to cold conditions. According to Schwartz (1924) and Sprent (1946b) this species is also very sensitive to desiccation. This was not entirely confirmed in my experiments, as infective larvae were present when no rain was recorded and the dung very dry (2b) and (c), 5(d) Table No. I - appendix.) However, with well distributed rainfall over the first five to seven days, many more larvae reached the infective stage. Although dung pads yielded few of these larvae when the Baermann Apparatus was used, incubated dung showed the presence of large numbers of larvae. Possibly under flooding conditions the eggs did not hatch as readily as those of the other species due perhaps to increased moisture of the dung depriving the eggs of air. This could not be confirmed but according to Sprent (1946b) eggs of this species need air to hatch. Although this hypothesis may be feasible it was more likely that pre-infective larval stages, being very sensitive to desiccation survived under wet conditions and were recovered as infective larvae after incubation in the laboratory. Experiments on infective larvae described elsewhere (vide infra) tended to confirm the latter view.



Investigations were carried out on the hatching of eggs and development of larvae to the infective stage of *Coenoria* spp. *H. planci*, *O. radiatum* and *B. phlebotomum* under field conditions.

The following observations were made:

- (1) Eggs hatched within eight days and infective larvae were recovered after a minimum period of five days in summer but more frequently at seven days from spring to autumn. First and second stage larvae reached their optimum development on the third and fifth days respectively and infective larvae on the seventh day.
- (2) In winter 42% of eggs had not yet hatched after 21 days in the field and no larvae had reached the infective stage when temperatures ranged from a mean minimum of 1.5°C to a mean maximum of 18°C. Protracted development of the free-living stages was possibly due not only to diurnal temperature fluctuations but also to the long duration of cold conditions; lower temperatures in the shade than in the sun; in the depth of dung than on the surfaces; grass minimum temperatures as much as 3-5°C lower than air temperatures; heavy frosts possibly causing freezing and death of pre-infective stages.
- (3) Marked evaporation occurred from spring to autumn. When moisture loss due to evaporation caused a decrease of 65% or more in weight of dung within the first 5-7 days, very few larvae reached the infective stage.
- (4) Eggs hatched more rapidly but less infective larvae were recovered from the crust than the depth of the dung.
- (5) Rainfall on one but particularly on two separate days from the second to the fifth or sixth day, caused optimal conditions for larval development from spring to autumn. In the winter rainfall made little difference to larval development and temperatures appeared to be more important.

- (6) No correlation existed between relative humidity of the atmosphere and larval development. The affect of shade compared with sunlight in the winter had its main affect in the modification of temperatures and rates of evaporation.
- (7) Copestia spp. were well adapted to extremes of heat and cold, dryness and desiccation. H. placai were more sensitive to hot arid conditions but more resistant to cold. O. radiatum was resistant to hot dry conditions but more sensitive to cold winter conditions. B. phlebotomum although occasionally recovered under very dry conditions at high temperatures, was very sensitive to cold, and only present in large numbers when well distributed rains provided adequate moisture for larval development in the summer.

**B. ACTIVITY AND SURVIVAL OF INFECTIONAL LARVAE UNDER FIELD CONDITIONS.**

**MATERIALS AND METHODS:-**

(1) Infested dung was collected from the rectum of two or more infested animals, shaped into a dung pad and placed in various positions in the veld to simulate natural defaecation. Since dung beetles were very active from November to April most dung pads were placed in a protected position in the open for 4-5 days to harden before being placed in the veld. In the hard dry state dung did not attract the beetles so readily.

(2) To trace migrations both the infested dung pad and the surrounding habitat were examined for larvae. Specimens were therefore, collected and placed in separate labelled containers as follows:-

- A. Dung pad.
- B. Soil to a depth of 2 cm. under the dung pad.
- C. Grass adjacent to the dung pad.
- D. Grass roots adjacent to the dung pad.
- E. Soil 2 cm. deep adjacent to the dung pad. The order of collection was C, D, E, A, B.

(3) Notes were made on the day of collection as follows:-

- (a) Date the dung pad was placed in the veld and the date specimens were collected.
- (b) Number of days in the veld.
- (c) The dung was measured and expressed as the maximum and minimum diameters on the lower surface and the maximum height e.g. 20 x 18 x 8 cm.
- (d) The surrounding vegetation, height of grass, bushes etc. was noted.
- (e) Since dung beetles often removed manure to the underlying soil and dung and soil could not be separated, it was included in the specimen B.
- (f) Where horizontal or lateral migration was being investigated grass and soil furthest from the dung pad was

collected first, followed by specimens nearer the dung

pad.

When vertical migration was being investigated the tops of the grass were cut off first, thereafter at various heights above the soil surface to ground level.

(4) After transfer to the laboratory specimens were:-

(a) Weighed, placed in the Baermann Apparatus using a technique similar to that advocated by Mönning (1930). His technique was modified as follows:-

(1) Dung was broken into small pieces with a pestle and mortar; no specimen in any funnel exceeded 50 grams in weight, nor did the thickness of dung exceed 2 cm. Similar remarks applied to humus, but sand was not weighed and the thickness of material only, was taken into consideration.

(2) Large funnels with a 20 cm. diameter were used and mutton cloth in place of linen.

(3) Grass roots were cut into small pieces and only 30 grams were placed in funnels.

(b) After 24 hours in the funnels, 25 cc. of fluid was withdrawn from the bottom of the funnels and left for 2-3 hours to settle in the collecting tubes. The supernatant fluid was then siphoned off and 2-3 cc. of the sediment left for examination.

(c) The methods of examination of larvae have already been described in the previous experiment.

(5) Prevailing climatic conditions.

The daily variations in the climatic conditions from June, 1955, to August, 1956, were recorded.

#### EXPERIMENTAL OBSERVATIONS:-

A total of 58 experiments was carried out over the period 23/6/55 to 7/8/56 and dung pads were left in the veld for periods varying from 7-105 days.

The results of these experiments are summarised in Table No. II in the appendix and wherever reference is made to experimental numbers, the results of these experiments will be found in this

Table, unless otherwise mentioned.

(a) Development of larvae to the infective stage.

This has been more fully dealt with elsewhere (*vide supra*) but two important points as an addendum to those experiments were noted here.

In the summer months more larvae reached the infective stage in dung protected from sun by the surrounding vegetation, than those exposed to the direct sunlight.

Poor recoveries of larvae invariably occurred from dung pads that had been attacked by dung beetles; possibly due to the effects of increased evaporation giving rise to desiccation and death of the preinfective larval stages (Nos. 24-28, 30-33 and 35).

(b) The distribution of infective larvae in the dung pad.

In experiments described elsewhere (*vide supra*), it was shown that more larvae reached the infective stage in the depth of the dung pad than in the crust. In 25 experiments the hard outer crust and the moister, deeper layers were examined separately. In 22 of these experiments considerably more larvae were recovered from the depth than the crust, thus confirming the observations in the experiments previously described. In three cases however the opposite was true (Nos. 12, 51 and 55). A further experiment also yielded more larvae in the crust than the depth but in this case a large proportion of the depth was included with the crust (No. 58). Most of the specimens with more larvae in the crust were collected in the winter.

A further subdivision of dung pads was made in 5 experiments. The crust was examined as before and the centre portion or depth divided into two layers, upper and lower layer.

In four of the experiments the largest number of larvae were recovered in the upper layer, i.e. the portion lying just under the crust. (Nos. 17, 19, 38 and 48). In three experiments the lower layer had more larvae than the crust (Nos. 17, 38 and 48), but in one case the opposite was true (No. 19). The latter was artificially moistened by having water dripping next to it.

In one experiment, most/the larvae were recovered from the crust, less in the upper layer and least in the lower layer. In this experiment active vertical migration had taken place; the crust was very soft under the influence of heavy rains (No. 42).

(e) Migration of infective larvae from the dung pads.

Larval migration was governed by rainfall. In the absence of rain or where rainfall was very limited no migration occurred (Nos. 5, 12, 26, 52 and 56). Little or no migration occurred under the influence of 14.0 mm. (0.55 inches) of rain which had been preceded by a comparatively dry spell (Nos. 22 and 23).

Larval migration occurred when 32 mm. (1.26 inches) of rain fell in the first 48 hours followed by 6.5 mm. (0.26 inches) on the tenth day when specimens were collected, (No.29.). Similarly when 55.8 mm. (2.19 inches) fell on the second day followed by 11.1 mm. (0.43 inches) from the eighth to the eleventh day when specimens were collected, larval migration was found to have occurred (No.43).

In these two experiments most of the larvae migrated to the soil, but some vertical migration on grass blades also occurred. In an experiment conducted over 7 days where 19 mm. (0.75 inches) fell on the fifth day the rain caused migration, most migration being to the soil under the dung pad (No.39).

Massive migration occurred for limited periods in December, February and March. Well distributed heavy rains occurred during these periods. In all cases rain fell on one, two or three days in the first 8 days the dung was in the veld. In one of these experiments conducted over 8 days rain fell on the 5th, 6th and 8th days a total of 116.0 mm. (4.6 inches) being recorded, and 83% of all the larvae recovered were from the grass. In addition horizontal migration of over 50 cm. from the edge of the pad had occurred to the grass and soil (No.42). In another 4 experiments where rains were well distributed, the totals varying from as little as 40.4 mm. (1.59 inches) to as much as 127.7 mm. (5.04 inches) over the whole period the experiments were conducted, massive larval migration had occurred, and 41% - 90% of all larvae recovered

were on the grass (Nos. 30, 41, 44 and 45.)

An experiment which ran for 9 days, where the rainfall recorded from the 4th to the 6th day was 26.0 mm. (1.02 inches) followed by 25.5 mm. (1.01 inches) on the 9th day did not cause more than 17% of all the larvae to migrate. However, a few larvae were recovered from the grass and soil 80 cm. from the dung pad (No. 46).

Larvae were recovered from grass 20 cm. above the soil surface in an experiment which ran for 41 days. All the rainfall, a total of 172.0 mm. (6.78 inches) fell in the first 24 days, there being no rain recorded during the last 17 days of the experiment (No. 49).

(d) Migration of larval species.

The results of species migration are summarised in Table No. 7.

**TABLE NO. 7:** Percentage of experiments in which larval migration occurred.

Species	Site of Larval Migration.			
	B.	C.	D.	E.
<u>Cooperia</u> spp.	53.5%	40.3%	27.6%	43.1%
<u>H. placei</u> .	32.7%	25.9%	20.7%	24.1%
<u>O. radiatum</u> .	25.9%	17.2%	12.1%	27.6%
<u>B. phlebotomum</u> .	12.1%	-	-	5.2%

B = Soil under the dung pad.

C = Grass adjacent to the dung pad.

D = Grass roots adjacent to the dung pad.

E = Soil adjacent to the dung pad.

Cooperia spp:

These species were the most active migrators from the dung under extremes of heat, cold and drought. They were the most frequently recovered larvae of all species dealt with and were recovered 80 cm. in the soil, 70 cm. on the grass, from the edge of the dung pad (No. 46) and migrated over 20 cm. vertically on grass (No. 49 Table



Haemonchus placei:

As can be seen in Table No.7 this species was not as active a migrator as Cooperia spp. When good rains fell this species was particularly prominent but was also found to have migrated under less favourable conditions. Horizontally it was recovered from grass 80 cm. and from soil 70 cm. from the edge of the dung pad (No. 46). It migrated vertically over 20 cm. on grass (No.49).

Oesophagostomum radiatum:

This species did not migrate as frequently as the species mentioned earlier but was recovered more frequently than H. placei from the soil next to the dung pad (Table No. 7). However, it only migrated horizontally 30 cm. in soil (No.41) but was recovered from grass 50 cm. from the edge of the dung pad (Nos. 41 and 42). Vertically this species only migrated up to 5 cm. on grass (Nos. 35 and 43).

Bunostomum phlebotomum:

This species was not really an active migrator from dung and was only recovered in the soil, more frequently beneath, and less so adjacent to the dung pad. (Table No.7). The larvae although recovered from the soil adjacent to the dung pad were confined to the soil right next to the dung pad (Nos. 30, 33 and 42). This species was never recovered from grass or grass roots.

(e) The role of insects in larval migration.

Under field conditions coprophagous beetles, termites, ants, etc., frequently attacked dung pads and removed the faeces to the underlying and adjacent soil. These small pieces of manure could not be separated from soil specimens. It is feasible that infective larvae developed in these small pieces of dung. Where dung beetles had been active, soil beneath the dung yielded infective larvae when climatic conditions were unfavourable for migration (No's. 13, 18, 24, 48 and 57), as well as in the adjacent soil (Nos. 18 and 57). There was possibly enough moisture, even when minute quantities of rain fell to stimulate larval migration from the underlying soil containing this manure, to account for the recoveries of larvae on grass under extremely unfavourable



conditions (Nos. 10, 11 and 18). Under slightly more favourable conditions of rainfall for larval migration, it was difficult to assess, whether rainfall, or beetles, or both were responsible for migration (Nos. 20, 27, 28, 33 and 35).

(f) The survival of infective larvae.

The maximum number of days larvae survived after infested dung was placed in the veld is summarised in Table No.8.

Cooperia spp. were the most resistant larvae regardless of the site of larval recovery, followed by H. placei, C. radiatum and B. phlebotomum in that order. It was interesting to note the similarity between H. placei and C. radiatum in the maximum survival rates on grass, and the longer period C. radiatum survived on grass roots compared with H. placei.

**TABLE NO.8:** Maximum number of days infective larvae were recovered after experiments commenced.

Species	Site of Larval Recovery.				
	A	B	C	D	E
<u>Cooperia</u> spp.	105	105	93	104	105
<u>H. placei</u>	105	104	41	28	104
<u>C. radiatum</u>	105	42	41	84	67
<u>B. phlebotomum</u>	24	25	-	-	25

A. = Dung pad.

B. = Soil under dung pad.

C. = Grass adjacent to dung pad.

D. = Grass roots adjacent to dung pad.

E. = Soil adjacent to dung pad.

■ = Days.

Tables Nos. 7 and 8 were compared and it was noted that migration and survival followed essentially the same pattern, with a few minor differences when H. placei and C. radiatum were compared. In this connection the frequency of larval migration was more important than actual survival rates in isolated cases.

Poor recovery rates of larvae occurred when experiments ran over long periods (Nos. 9, 16, 21, 32 and 57) except in two experi-

ments; in one the dung mass was fairly large and rainfall was well distributed; the other was conducted in autumn and winter under colder conditions (Nos. 38 and 58).

Apart from the two exceptions mentioned, larvae were recovered in large numbers only when the experimental period did not exceed 24 days (Nos. 14, 17-20, 23, 29, 36, 40, 42-44, 46-48 and 53). A fair number of larvae were recovered in two experiments which ran over 40 and 41 days (Nos. 22 and 49 respectively).

In experiments described elsewhere (vide supra), the minimum period in which infective larvae developed in dung pads was five days. The results of these experiments indicate good recoveries of infective larvae up to the 24th day i.e. after infective larvae had been present for a maximum period of 19 days. Thereafter the mortality rate increased sharply and larvae were rarely recovered in large numbers after the 25th day, except in isolated dung pads, or in small numbers where migration had occurred.

(g) Migration of infective larvae to grass.

Distribution of rainfall has already been mentioned as an important factor in larval migration. As five days is the minimum period required for infective larvae to develop to the infective stage, rainfall in this period would assist larval development to the infective stage and thereafter assist larval migration.

In Table No.9, four experiments where larval migration to the grass occurred are compared. Heavy rainfall particularly after the fifth day explained the excellent migration in Experiment No.42. Although the actual total amount of rainfall for the period was less in No. 44, than in either No.29 or 46, a higher percentage of larvae migrated in the former when compared with the latter experiments.

Apparently the rainfall that fell after the fifth day accounted for better migration of larvae to the grass, although in this period only 10.2 mm. (0.40 inches) more was recorded in No. 44, than No. 46. It was interesting to note that as little as 6.5 mm. (0.26 inches) in No. 29 after the fifth day, was probably responsible for migration of 10 per cent of the infective larvae to grass.

**TABLE NO. 9:**

**Distribution and amount of rainfall required  
for optimal larval migration to grass.**

Experiment No.	Up to 5 days		After 5 days		Larvae on Grass		No. of days in field.
	No. of days rain fell.	Total rain-fall in mm.	No. of days rain fell.	Total rain-fall in mm.	Total No. Recovered.	Percentage of total No. from all sources.	
42	1	27.5	2	89.3	6,118	83%	8
44	2	3.6	4	36.9	586	46%	12
29	2	38.5	1	6.5	125	10%	10
46	2	24.8	2	26.7	266	10%	9

In experiments where horizontal migration was measured the largest percentage of larvae were recovered within 10-15 cm. from the edge of the dung pad (Nos. 41, 42, 44, 45-47). It was also noted that the largest proportion of larvae were recovered at heights of less than 10 cm. and decreased in numbers the higher the grass was cut off from the soil surface (Nos. 41, 44, 45, 47 and 49). whereas in one experiment 50% of the larvae were recovered below 10 cm. and a further 28% between 10 and 20 cm., only 22% were found in grass from 20 cm. upwards (No. 49). Migration on grass was most marked where the grass was thick and formed a "mat".

(h) Prevailing climatic conditions.

The climatic conditions prevailing during the experimental period (26/6/55 - 7/8/56) are shown in Fig. No. 1. The monthly rainfall is included in Table No. 10, and in Table No. II relevant climatic data for the period each experiment ran are also shown.

It will be noted that during the experimental period little or no rain fell except for 3 months in December 1955, February and March 1956, when rainfalls were good. Marked diurnal temperature variations occurred. In July the mean minimum was 0.3°C (32.5°F) and the mean maximum 20.7°C (69.2°F). In January the mean minimum was 15.9°C (60.6°F) and the mean maximum 31.6°C (88.8°F.) Mean relative humidity varied from 31.5% in September to 70.4% in March.

**TABLE NO.10: Monthly Rainfall at Armoedsvlakte from July, 1955 to June, 1957.**

M O N T H.	Number of days rain fell.	Total Rainfall.	
		mm.	inches
July, 1955.	0	0	0
August.	0	0	0
September.	1	0.2	0.01
October.	6	16.5	0.7
November.	6	40.4	1.6
December.	10	127.9	5.0
January, 1956	9	38.3	1.5
February.	7	150.3	5.5
March.	15	175.6	6.9
April.	0	0	0
May.	5	9.4	0.4
June.	0	0	0
<b>TOTAL.</b>	<b>59</b>	<b>548.6</b>	<b>21.61</b>
July, 1956	0	0	0
August.	0	0	0
September.	3	13.1	0.53
October.	6	18.3	0.72
November.	2	11.5	0.45
December.	11	67.4	2.66
January, 1957	7	82.7	3.26
February.	7	29.9	1.18
March.	10	47.7	1.88
April.	5	41.0	1.61
May.	4	10.0	0.39
June.	2	23.0	0.91
<b>TOTAL.</b>	<b>57</b>	<b>344.9</b>	<b>13.59</b>

**DISCUSSION:-**

Generally speaking the results of these experiments were disappointing. No larvae were recovered in 12%, less than a hundred in 35% and between a hundred and a thousand in 19% of the experiments respectively. Only in 34% of the experiments were more than one thousand larvae recovered, of which 24% had one to ten thousand, and 10% more than ten thousand larvae respectively. The poor results were probably due to adverse climatic conditions, the surrounding habitat, survival of larvae and secondary causes (e.g. dung beetle activities).

It has been shown in experiments described elsewhere that no larvae reach the infective stage in the cold winter months at temperature varying from a mean minimum of 2.6°C (36.7°F) to a mean

maximum of 22.7°C (72.9°F). When temperatures in spring were adequate for hatching and larval development, drying and desiccation due to evaporation caused death particularly of the preinfective larvae. Similar results were obtained in summer when no rain fell during the first 6-7 days dung pads was placed in the field. The negative and poor results in spring and in hot dry periods in summer were probably due to death by desiccation of preinfective larvae. This has been the experience of many other workers with different species of strongyloid worms (Hanson 1906; Veglia 1915; Theiler & Robertson 1915; Schwartz 1924; Mönig 1930; Ortlepp 1937; Crofton 1948 (b); Dinnik & Dinnik 1954-5, etc.).

It was noticed that dung which had no protection from the sun during the day gave very poor results. Partial and deep shade yielded more larvae under similar conditions. This was probably due to the temperature causing higher rates of evaporation in exposed dung pads. Dinnik and Dinnik (1954-5) working with Haemonchus contortus found similar results in the dry season. In winter in one experiment however large numbers of larvae were recovered from a dung pad exposed to the sun (No.53) and only one larvae recovered from dung lying in the shade (No.52). In these cold conditions exposure facilitated development whereas shade retarded development, due probably to higher temperatures in the former assisting development to the infective stage, whereas the temperatures in the shade were too low to allow the preinfective stages to reach the infective stage.

Poor recovery rates were recorded when experiments ran for long periods, with one major exception, the last experiment conducted over 105 days from April to August (No. 58). In this experiment larvae developed to the infective stage in autumn, i.e. they survived the whole winter, practically all being confined to the dung pad. The life expectation of larvae in other experiments throughout the year appeared to decrease sharply after the twenty-fourth day of exposure to field conditions. This observation was of importance in that it indicated how long heavy infestation could remain in a pasture, and would have to be taken into account in the prophylaxis of verminosis by pasture management.

Dung beetles are undoubtedly the ally of the cattle rancher in the control of verminosis. Wherever these beetles had been active larval recoveries from dung pads were very poor, due doubtless to the aeration of dung, assisting evaporation and accelerating the desiccation and death of preinfective and infective larvae. By the mechanical removal of dung particularly from the lower surface to the underlying and also adjacent soil, these beetles probably accounted for the large number of experiments in which larvae were present in the soil, particularly underneath, but also next to the dung pad. It was a feature of these experiments that wherever dung pads were hollowed out by beetles, larvae were recovered from the soil beneath the dung. Dung pads protected from beetles did not show this phenomenon to the same extent, unless rainfall had been present. It is suggested that beetle activity was the main reason why larvae were recovered from soil, and not the migratory habits and positive heliotropism of the larvae as suggested by Mönning (1931). In most cases Mönning found more larvae in the soil next to rather than underneath the dung pad. This observation was not confirmed in most cases in the experiments described above, in fact the reverse was observed.

From the observations that larvae could be recovered from soil underneath and adjacent to dung pads, it appeared reasonable to assume that a few would be found near the bases of grass tufts. This was actually the case. Very little rain would then be necessary to stimulate larvae to migrate on to grass, as shown in certain experiments (Nos. 10, 11 and 18).

In the dung pad itself most moisture is retained in the so-called depth of the dung pad, and more larvae were recovered in the depth than elsewhere. This was possibly due to either death of preinfective larvae in the drier crust or migration of these larvae to the moister and deeper layer.

Migration of larvae in the dung pads was however interesting. Only in one experiment was it noticed that active vertical migration was taking place in the dung pad, where rainfall had been very heavy indeed and the crust of the dung pads was softened thereby (No. 42). In most of the cases where active



migration occurred under the influence of rainfall larvae were recovered in large numbers in the soil, particularly beneath the dung pad. Possibly the hard crust of the dung did not allow passage of infective larvae from the depth but since the soil surface layers were softer they could escape to the ground. This migration did not include the effects of dung beetles, since it was present even when dung beetle activity had been excluded. In one experiment which ran over seven days and 19.0 mm. of rain was recorded on the fifth day, only downward migration occurred (No.39). In another two experiments a rainfall range of 11.1 - 19.0 mm., after the fifth day caused most of the infective larvae to migrate to the soil beneath the dung pad (Nos. 38 and 40). With slightly higher rainfall larvae were also recovered from the grass but the larger percentage of migrated larvae were still found in the soil under the dung pad. It appeared therefore that a rainfall range of 11-19 mm. (0.43 - 0.77 inches) stimulates downward migration of larvae into the soil; some larvae, however, were found on adjacent grass; marked larval migration to the grass was only noted when heavier and well distributed rains fell.

A significant observation was that even heavy dews in the absence of rain failed to cause larval migration to the grass, although 0.3 per cent of the larvae were found under the dung pad (No. 48). This is not in agreement with the views of Australian workers who stated that heavy dew would assist migration from dung pads (Riek, Roberts & O'Sullivan 1953). However, once larvae had migrated from dung pads dew would undoubtedly assist further migration.

Of great practical importance is the amount and distribution of rainfall necessary for significant larval migration to occur to the grass. The amount of rainfall necessary, for significant larval migration was from 40.4 mm. upwards. Of greater significance however was the distribution of rainfall. It has already been shown that rainfall during the first 5-6 days assisted larval development. Once infective larvae were present further rainfall would assist in larval migration. The important rainfall for larval migration would thus occur after the fifth day. This point i

shown in Table No.9. A rainfall range of 6.5 mm. to 26.7 mm. after the fifth day caused only 10% of the larvae to migrate to grass, whereas 36.9 mm. falling after the fifth day caused the migration of 45% of larvae to the grass, i.e. a mere 10.2 mm. of extra rain falling when infective larvae were freely available increased larval migration to the grass by 35%.

According to Rogers (1940) light intensities of 62 f.c. have been shown to be optimal for larval activity. These intensities exist under English conditions from half way down the grass to the base depending on the brightness of the day (Crofton 1948a). Light intensities at Armoedsvlakte during summer apart from cloudy days which were rare, were probably high due to the fact that grass tufts were scattered. Consequently specimens were collected from dawn to 7 a.m. which according to Rogers (1940) would be nearly optimal. However Crofton (1949) found more larvae on grass from 12 noon to 5 p.m. than in the early morning when dew was present. He stressed the fact that migration of larvae was impossible below 10°C (50°F) and it was fallacious to assume that moisture only would control larval activity. In the summer months at Armoedsvlakte minimum daily temperatures were usually in excess of 12°C (53.6°F) and as high as 20°C (68°F) so that temperatures at dawn were adequate for larval migration. Apparently in England this is not the case.

The vertical distribution of larvae on grass generally followed the experience of other workers who found most of the larvae near the base i.e. lower 2-3 inches of grass. (Taylor 1938, Crofton 1948a). In these experiments more larvae were recovered in the lower 5-10 cm. of grass than higher up although a few larvae were found in one case to have migrated as much as 20 cm. vertically. The number of larvae recovered from grass appeared to be greater where the grass was thicker. This agreed with Kausal's (1941) observation. On the other hand where optimal larval migration was taking place in five of these experiments (Nos. 30, 41, 42, 44 and 45) 41% - 90% of available larvae were recovered from the grass. This percentage is much higher than the 16% observed by Kausal.



The horizontal distribution of larvae on grass showed that 72-100% of the larvae were recovered within 15 cm. of the dung pad. The numbers of larvae decreased rapidly, as the distance from the dung pad increased. The maximum horizontal migration on the grass of H. placei was over 80 cm. whereas Cooperia spp. were only found up to 70 cm. from the dung pad i.e. within a radius of 89 and 79 cm. respectively from the centre of the dung pad (No. 46).

In this particular experiment the grass was fairly dense but did not form a continuous thick mass around the dung pad. Most of this migration was influenced by rainfall of 25.5 mm. (1.0 inches) which fell the last 2<sup>4</sup> hours before the specimens were collected. This may possibly have caused a little flooding and assisted in disseminating the larvae over a large area.

The behaviour of the various species of larvae differed in certain important aspects.

#### Cooperia spp.

These larvae were the most drought resistant, lived longer under various climatic conditions in the dung, soil, grass and grass roots and migrated and survived when no other species was found. This observation confirms work done in Australia (Roberts, Riek and O'Sullivan 1952). Since almost a pure Cooperia nectinata infestation was present it can be stated that this species can withstand cold better than H. placei.

#### Haemonchus placei.

Although not as resistant to drought as the previous species it was recovered from grass in small numbers under very unfavourable circumstances.

In comparison with Cooperia spp. it survived as long in dung and soil, although not as frequently, but died off more rapidly in the grass and grass roots than Cooperia spp. did. When conditions were optimal for larval migration it was however more prominent than the former species.

#### Oesophagostomum radiatum.

Both this species and B. phlebotomus were for unavoids-

ble reasons not available initially in infested dung to the same extent as the first two species mentioned. It would be wise therefore to treat the results of the behaviour of the infective larvae of these latter two species with reserve.

It was found that *O. radiatum* did not migrate as readily as either *Cooperia* spp. or *H. placei* but was recovered more frequently than the latter from the soil adjacent to the dung pad, but less frequently elsewhere. It survived longer on grass roots than *H. placei*, the same length of time on grass and dung but shorter periods in the soil.

#### *Bunostomum phlebotomum*.

This species was the most sensitive to adverse climatic conditions. It neither lived as long nor migrated to any appreciable extent when compared with the other species. Its maximum survival rate from the time dung was placed in the veld was 24 days in the dung pad and 25 days in the soil. It was recovered on a few occasions from the soil beneath the dung pad and only rarely in the soil next to the dung. It did not migrate to the grass, an observation also made by Sprent (1946b) who stated that they remained in the dung. He did not mention whether they migrated to soil. His first observation was confirmed by these observations but although more frequent in dung, larvae were also recovered from the soil in these experiments.

Australian workers have stated that *B. phlebotomum* will not survive in areas with an average annual rainfall of less than 24 to 25 inches (Riek et al 1953). This observation was not confirmed. Over a 14 month period in these experiments 21.6 inches of rain fell. This species was recovered in 25.8% of the experiments from October to May inclusive, in which period monthly rainfall varied from nil to 6.9 inches.

#### SUMMARY.

Studies were carried out on the activity and survival of infective larvae of *Cooperia* spp., *H. placei*, *O. radiatum* and *B. phlebotomum*, under field conditions. The following observations were made:-

- (1) Fifty-eight experiments were carried out over a period of fourteen months. Eleven of these months had little or no rain. Three months had monthly rainfalls varying from 127.9-175.6 mm. (5-6.9 inches).
- (2) Only thirty-four per cent of experiments gave satisfactory results. In most cases these experiments were carried out in summer in periods when rainfall was adequate, dung pads were placed in areas partially or wholly protected from the sun, the experimental periods were short and dung beetle activity excluded.
- (3) Sixty-six per cent of experiments gave unsatisfactory results due to one or more of the following factors.
  - (a) Adverse climatic conditions.
  - (b) Over-exposure to sunlight.
  - (c) Long term experimental periods.
  - (d) Dung beetle activity.
- (4) Larvae were usually recovered in dung pads in greater numbers in the depth, which lay between the outer crust and ground surface layers, than either the crust or ground surface layers.
- (5) Migration from dung was influenced by rainfall as follows.
  - (a) Under the influence of 14.0 mm. (0.55 inches). of rain after a fairly dry period larvae failed to migrate.
  - (b) When rainfall occurred in the first five days dung was in the field a rainfall range of 11-19 mm. (0.43-0.77 inches) thereafter stimulated migration of larvae mainly to the soil.
  - (c) In experiments which ran over periods of 9-12 days and rainfall had been recorded in the first five days, rainfall thereafter caused greater or lesser migration to the grass. When 6.5-26.7 mm. (0.26-1.05 inches) of rain fell after the fifth day only 10% of the larvae migrated to grass, but if 36.9 mm. (1.45 inches) or rain fell in this period 46% of the larvae migrated to the grass.

- (6) Cooperia spp. larvae migrated horizontally 70 cm. on grass, 80 cm. in soil and 20 cm. vertically on grass. They migrated more frequently and survived longer than any other species under field conditions.
- (7) Haemonchus placei larvae migrated horizontally 80 cm. on grass, 70 cm. in soil and 20 cm. vertically on grass. They migrated less frequently and survived as long in dung, almost as long in soil, but much shorter periods than Cooperia spp. on grass and grass roots.
- (8) Oesophagostomum radiatum larvae migrated horizontally 50 cm. on grass, 30 cm. in soil and 5 cm. vertically. They migrated less frequently, only survived as long in the dung as the previous species, and were also recovered from grass for the same maximum period as H. placei. Otherwise their survival rate was lower.
- (9) Bunostomum phlebotomus larvae did not migrate on to grass but only to soil underneath or right next to the dung pad. Their survival rate was extremely short by comparison with the other species.
- (10) Horizontal migration was more marked within 15 cm. of the dung pad. Vertical migration was more noticeable within 5-10 cm. of the soil surface on grass.
- (11) Dung beetles decreased the larval recoveries from dung pads by increasing aeration and evaporation causing death by desiccation of preinfective larvae. At the same time they assisted migration by their mechanical movement of the dung to soil, in which some larvae could reach the infective stage.

## A D D E N D U M.

### INSECT ACTIVITIES UNDER NATURAL CONDITIONS.

#### I N T R O D U C T I O N.

Subsequent to the completion of this paper, the experimental data were re-examined and the observations on the effects of dung beetles on the free-living stages of the life-cycle re-assessed. Since these insects play an important role on the potential infestation in the field, this addendum has been drawn up. Attention is drawn to experimental observations and their significance is discussed.

#### EXPERIMENTAL OBSERVATIONS.

##### A. Hatching of eggs and development of larvae to the infective stage.

These results have been summarised in Table No.I in the appendix and the experimental numbers referred to will be found in this table.

Dung beetles and other insects attacked dung pads during November 1956 and again from January to April 1957. Although afforded no protection against dung beetles, dung pads placed in the field from August to October, in December 1956 in May and June 1957 were not attacked by them.

During November 1956 in experiment 7 (a-f) six dung pads were placed in the field. The first dung pad 7(a) was collected after one days exposure to field conditions, and had not been attacked by dung beetles. The other five dung pads 7 (b-f) collected subsequently had been hollowed out by beetles. Two days after exposure to field conditions 7(b) and 7(c) were collected; the former weighed 150 gm. the latter 100 gm., the difference in weight being due to increase insect activity; egg counts were 120 and 20 eggs per gram respectively indicating more rapid hatching of eggs in the latter. Only 25 first stage and 12 second stage larvae were recovered in the former while 256 first stage larvae and no second stage larvae were recovered in the latter after extraction with the Böhmer Apparatus. However after a further 8 day incubation in the laboratory the specimens yielded 10,000 and 3,500 larvae respectively. More than

twice the number of infective larvae were recovered from the former, when compared with the latter although the weight of the dung in the latter at collection was two thirds of the former.

Another dung pad, 7(d), was examined on the third day; the dung only weighed 67 gm. and both egg counts and larval examination after extraction with the Bärman Apparatus were negative. After a further 8 day incubation in the laboratory only 550 infective larvae were recovered. This dung pad had been extensively hollowed out by insects and although it was 67% of the weight of 7(c) when collected in the field, the number of larvae collected from this specimen was only 15.7% of the number collected the previous day in 7(c). The activities of dung beetles as well as the longer exposure to field conditions were responsible for the marked drop in larval recoveries.

The remaining dung pads 7(e) and 7(f) exposed to field conditions for 30 and 64 days respectively, were also extensively damaged by dung beetles and yielded few larvae on examination.

In experiment No.9 six dung pads were placed in the field on 11/1/57. Three days later 9(a) was collected and examined. It was extensively hollowed out by beetles, and weighed 113 gm.: egg counts were negative; two first stage larvae were recovered with the Bärman Apparatus and 825 infective larvae after 8 days incubation.

The other five dung pads in this experiment were almost completely destroyed. Small heaps of loosened soil covered with one or two pieces of dung, the largest of which was 2 cm. in diameter indicated where the dung pad had been. On examination of these heaps minute pieces of manure were found to be inextricably mixed with the soil.

A large mass of dung 3 lb. in weight was also placed in the field on 11/1/57. This experiment, No.10, suffered the same fate as the five dung pads in No.9. In experiment No.11, four dung pads were placed in the field on 17/1/57. Within 48 hours only four small heaps of loosened soil indicated the site of these dung pads.

It was important to know whether infective larvae could develop in the small pieces of dung removed to the soil by these insects and be recovered from the soil. This was included in the study of infective larvae in the previous year and is reported hereunder.



**B. The activity and survival of infective larvae under field conditions.**

These results have been summarised in Table No.II in the appendix and the experimental numbers referred to will be found in this Table.

In sixteen experiments (No's. 7,10,11,13,18,20,24,25, 26,27,28,30,31,32,33, and 35) dung beetles had been active to a marked degree. In the remnants of the dung pads infective larvae were recovered in large numbers in only one experiment (No.18); in ten experiments (No's. 11,13,20,26,27, 28,30,32, 33 and 35) from 1 to 388 larvae were recovered, while no larvae were present in five experiments (No's. 7, 10, 24, 25, and 31).

Dung beetles removed manure to the soil underlying and adjacent to the dung pads. It was impossible to separate these small pieces of dung from the soil and they had to be examined together.

It was interesting to compare the number of infective larvae recovered from the soil beneath the dung pad with those in the dung pad itself. More larvae were recovered in the soil beneath the dung pad in five experiments (No's. 20,28,30, 33, and 35) and less in four experiments (No's. 13,18,27, and 32). In two experiments (No's. 24 and 31) a few larvae were recovered in the soil, while none were present in the dung pad. In two experiments (No's. 13 and 18) rainfall was inadequate to cause any migration; in a further three experiments (No's. 20, 27, and 28) the amount of rainfall only partly assisted downward migration from the dung to account for some of the infective larvae found there. Where rainfalls had been adequate, it was impossible to assess whether the presence of larvae in the soil was due to migration from the remnants of the dung pad, or whether they had developed in the small pieces of dung removed to the soil by dung beetles (No's. 30, 31, 32, 33 and 35).

Larvae were recovered in nine experiments (No's. 18, 20, 27, 28, 30, 31, 32, 33 and 35), from the soil adjacent to the dung pad. In one experiment (No.18) rainfall was inadequate to have caused this migration from the dung pad. In the other eight experiments the same remarks apply as mentioned in the previous paragraph, to account for the presence of larvae in the soil, in relation to rainfall and beetle activity.

On the grass next to the dung pad larvae were recovered in ten experiments (No's. 10, 11, 18, 27, 28, 30, 31, 32, 33 and 35). The most interesting observations were made in three of these experiments (No's. 10, 11, and 18). The rainfall recorded during all these experiments was less than 0.5 mm. for the periods of exposure to field conditions. Experiments described elsewhere showed that this amount of rain was inadequate for larval migration, from the dung pad itself. Dung beetles had removed small pieces of the dung to the soil at the base of the grass. It is more than likely that even very little rain was sufficient to cause migration, on to the grass, of infective larvae from these small particles.

In only four experiments (No's. 28, 30, 33 and 35) were larvae recovered from grass roots adjacent to the dung pad. They were usually present in smaller numbers than those recovered from other sources.

On examination of the results of twelve experiments which ran over short period of seven to nineteen days, it was interesting to note that in only two experiments (No's. 18 and 20), were large numbers of larvae recovered. In similar experiments (No's. 15, 17, 23 and 29), conducted at the same period, when dung beetles had not been active, considerably larger numbers of larvae had been recovered. These parallel experiments carried out under the same climatic conditions, afforded evidence that the end result of dung beetle activity, was a marked reduction in the number of infective larvae.

### C. The effect of low level phenothiazine dosing on dung beetle activity.

#### Introduction.

The observations reported previously showed that dung beetles were important in reducing the number of available infective larvae. Phenothiazine when dosed to stock in small quantities is excreted with the faeces and has been extensively used to control the pasture stages of the life cycle of round worms. It was important to know whether this drug when excreted in faeces during low level phenothiazine dosing had any deleterious effect on dung beetles. If this were the case low level phenothiazine dosing would counteract the insect activities in dung and could not be recommended. For this reason low level dosing of phenothiazine was carried out on two animals from which faeces were collected, and dung beetle activities in these faeces compared with their activities in faeces derived from an undosed control animal.



Materials and Methods.

Two cows were dosed daily with two and five grams of phenothiazine respectively. Three days after commencement of dosing faeces were collected from these two animals as well as from an undosed control cow. From each of these animals dung weighing 1 Kg. was placed in the field. These dung pads were collected after exposure to field conditions for periods of three to six days. In the laboratory they were weighed and the number of dung beetles in each specimen counted. In addition each dung pad was examined to see how extensively they had been hollowed out by insects. The insects collected were placed in bottles to see how long they would survive after removal from the dung. Dosing continued and specimens were collected over a period of two weeks.

Experimental Observations.

**TABLE:** A comparison between the activity and the number of dung beetles recovered from 1 Kg. samples of dung exposed to field conditions.

No. of days in field:	No. of insects recovered from dung of Cow No.1:	No. of insects recovered from dung of Cow No.2:	No. of insects recovered from dung of Cow No.3:	Remarks.
3	172	134	13	Dung in specimen (c) was more hollowed out by beetles than either (a) or (b)
3	260	70	174	Little damage was done to any of the dung pads by beetles.
3	174	48	12	More evidence of dung beetle activity in (b) and (c) than in (a).
4	74	55	8	Dung in (c) more extensively hollowed out than either (a) or (b).
5	18	32	5	Dung of (a) and (b) more hollowed out by beetles than (c).
6	35	18	3	All specimens extensively hollowed out by dung beetles.

\* (a) Cow. No.1. dosed daily with 2 gram of phenothiazine.  
 \* (b) Cow No.2. dosed daily with 5. grams of phenothiazine.  
 \* (c) Cow No.3. undosed control.

In the accompanying table the number of insects recovered from the various specimens showed no correlation whatever. Dung pads were frequently extensively hollowed out but very few insects were recovered in such specimens. In most cases more dung beetles were recovered from the dung pads derived from cows receiving low level phenothiazine dosing than from the dung of the control cow. The evidence that insects had been active varied to such an extent between the specimens, that although they were not necessarily still present when searched for in the dung in the laboratory, their absence did not indicate that other dung beetles had not been active prior to collection and had subsequently left the dung pad. There was no evidence that insect activity was more marked in dung from the undosed control than either of the dung pads from the two cows receiving daily doses of phenothiazine.

The insects that were recovered from these dung pads were placed in bottles in the laboratory. They lived for two to three days before they started dying regardless of whether they were derived from dung from the dosed cattle or the undosed control. They lived as long as six days before dying when if dung was placed in the bottles, regardless of the presence or absence of phenothiazine in the dung which was added to the bottles.

From these observations it is clear that low level phenothiazine dosing of cattle had no apparent affect on the activity of insects that attacked their dung, nor did it cause death of any dung beetles. Since it had no deleterious effects on dung beetle activity, this drug can safely be used in the control of verminosis in stock, as is at present advocated in many parts of the world.

## DISCUSSION.

The observations on the hatching of eggs and development of larvae to the infective stage in dung pads in the field, indicated that dung beetles caused increased aeration of dung pads, which speeded up the tempo of egg hatching. This confirms Mönnig's (1931) observation. In spite of the more rapid hatching of eggs, the recovery of larvae from such dung pads after incubation in the laboratory, was considerably less than from those dung pads in which the insects although present had not hollowed out the dung pad to the same extent. The death of the larvae was probably due to increased aeration, giving rise to more rapid desiccation and death of the preinfective larval stages.

During January 1957 eleven dung pads were placed in the field. Ten of these dung pads were completely destroyed within 48 to 72 hours leaving only small heaps of soil covered with a few small pieces of dung. The one dung pad left was extensively hollowed out and was examined as previously reported.

Throughout the rest of the summer and autumn of that year, repeated attempts to study the preinfective stages of the life-cycle, under completely natural conditions met with a similar fate. Once the dung beetles had destroyed the dung pads egg counts and the recovery of the various developing larval stages was impossible. Since the author was also studying the effects of other environmental factors on these stages of the life-cycle at the same time, he was compelled, albeit reluctantly, to protect the dung pads with wire gauze. Fortunately dung beetle activity ceased at the end of April, so that it was unnecessary to protect the dung pads in the winter and they could be studied under completely natural conditions.

In the previous year dung beetles had also been active, but not nearly to the same extent as in the autumn of 1957. The poor recoveries of larvae as a result of dung beetle activity reaching a peak in the autumn, offset to a marked degree the beneficial affects of increased rainfall on the development of infective larvae.

The poor recoveries of infective larvae, in fourteen of the sixteen experiments on infective larvae substantiated the statement made above, that insect activity facilitated the death of the preinfective larval stages. Dung beetles,

are attracted to fresh manure only, while it is still soft and very moist, i.e. in the first four days it is in the field. They burrow into the dung pads and remove manure to the soil particularly beneath but also next to the dung pad. These small pieces of manure were seen in the soil. The recovery of fairly large numbers of larvae in the soil mixed with manure granules below the remnants of the dung pad, under conditions where rainfall was inadequate for larval migration from the dung pad, must have been due to the presence of larvae in these small pieces of manure. Since they had been removed within the first few days the dung was in the field, it is only reasonable to assume that they must have developed to the infective stage after being removed by dung beetles to the soil. The movement of dung beetles in the soil loosens the soil, thereby introducing air and probably also increasing the rate of evaporation. This although stimulating egg hatching, would also increase the death rate by desiccation of the larvae. This possibly accounts for the poor larval recoveries from soil samples in most of the experiments where dung beetles had been active.

During the two year period of these investigations from July 1955 to June 1957, it was noted that dung beetles were most active from January to April in both seasons.

Rainfall data at Arnedsvlakte over a period of more than 30 years indicated that the rainy season was from December to April, the best month being March. Experiments described elsewhere showed that these months were the most optimal for the free living stages of the nematode parasites. Further more experiments on the transmission of verminosis proved that stock only became infested during this period.

It was very significant that during this same period dung beetles were most active. While other environmental factors facilitated the propagation of infestation in the late summer and autumn, the burrowing habits of insects in the dung had the opposite effect. Within two or three days of being in the field dung pads were frequently destroyed. Only the presence of rainfall would save any remaining larvae from desiccation and death.

The presence of frequent rains and nocturnal dews would be necessary to prevent the death of the preinfective stages. Although dew was a nightly occurrence it would only moisten the surface of the soil and any remnants of dung pads above the soil. Below the surface of the soil rain would be

required to supply sufficient moisture for larval survival. In this respect the erratic distribution of the rain meant that few larvae had this benefit and therefore died in the frequent relatively dry periods between rainfalls. Only at those periods when rainfalls were regular would more larvae reach the infective stage. Thus insect activity would be balanced out to a certain extent by the moisture supplied by rain. The interim periods between days on which rain fell would allow the full effects of dung beetle activity to be felt. Since these periods were more frequent than the days on which rain fell, few larvae would reach the infective stage during this period. In other words it was only during rainfall periods that the larvae really had a chance to survive and develop to the infective stage.

Dung beetle activity can safely be stated to be an important method of biological control of worm infestation in the field and an important ally of the farmer in worm control.

#### S U M M A R Y .

1. Four experiments were described on the hatching of eggs and development of larvae under field conditions where insects attacked dung pads.

2. In November 1956 five dung pads in one experiment were attacked by dung beetles. These insects caused increased aeration, stimulating egg hatching, but larval recoveries after incubation in the laboratory, were considerably less in dung pads where insect activity was most marked, indicating increased larval mortality.

3. In three experiments in January 1957, eleven dung pads were placed in the field; of these only one was suitable for examination after three days. The other ten dung pads were completely destroyed, within 48 to 72 hours, leaving small heaps of sand containing a few small pieces of manure inextricably mixed with the soil. No semblance of dung pads were left to examine for egg hatching or the recovery of larvae.

4. The effects of dung beetles were observed in sixteen experiments in which the activity and survival of infective larvae were being studied. In two experiments larvae were recovered in fairly large numbers. In fourteen experiments, larvae were either absent, or present in small or very moderate numbers.

5. The presence of infective larvae in soil underlying and next to dung pads, was concluded to be partly due to their development in the small pieces of manure mixed with the soil. These pieces of dung had been removed from the dung pad by insects in the first few days it had been in the field, and before infective larvae were present in the dung pads.

6. Where rainfall was inadequate to cause larval migration, the presence of larvae in the soil was probably due to the mechanical removal of dung by beetles.

7. Low level phenothiazine dosing has no deleterious effects on the activities of dung beetles.

### C O N C L U S I O N S .

Dung beetles hollow out dung pads which gives rise to increased rates of evaporation. This leads to death of the preinfective larvae. In the late summer and autumn, when climatic factors are optimal for larval development, these insects are most active and frequently destroy dung pads. Their activities are therefore an important method of biological control of infestation in the field. The administration of small doses of phenothiazine to cattle does not effect the subsequent activities of beetles in dung.



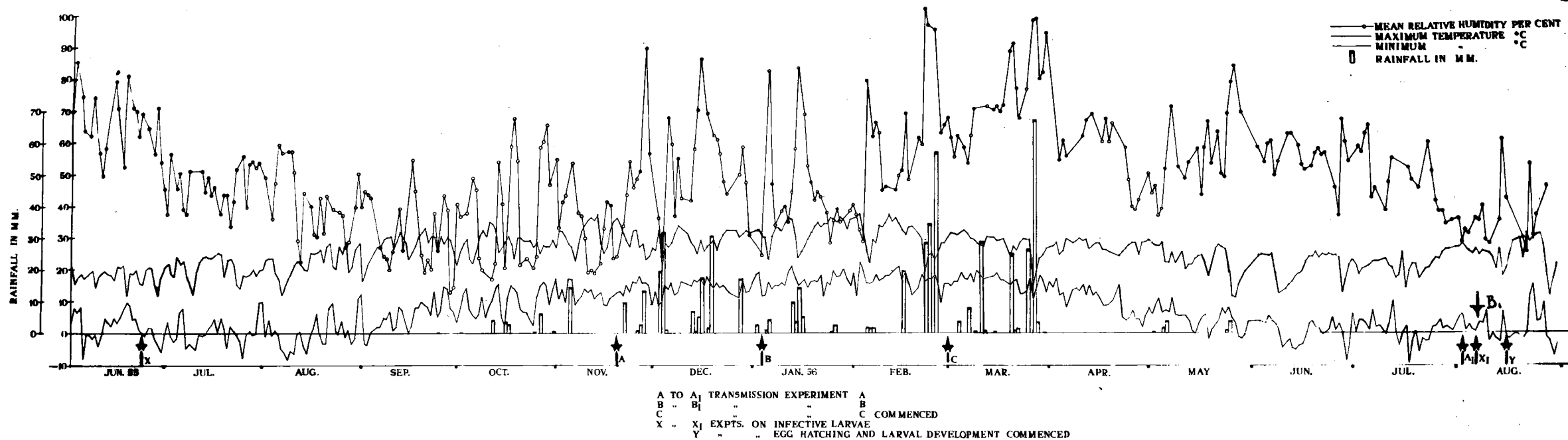


FIG. No.1 DAILY VARIATIONS IN THE CLIMATIC CONDITIONS AT ARMOEDSVLAKTE FROM 1/6/55 — 31/8/56





## EPIZOOTOLOGY.

### INTRODUCTION.

Based on field observations reported elsewhere (vide supra), four experiments were carried out on the transmission of verminosis to susceptible calves, reared by the various methods of animal husbandry practiced in the district.

The experiments were as follows:-

- A. Case report on calf No. 7180.
- B. Four groups of calves reared by different methods of animal husbandry from midsummer to winter.
- C. Five groups of calves reared by different methods of animal husbandry from autumn to midsummer.
- D. Seven groups of calves reared by different methods of animal husbandry from summer to early winter.

Note: In experiments C and D the work done in experiments A and B was repeated with the addition of extra groups.

### A. CASE REPORT ON CALF NO. 7180.

### MATERIALS AND METHODS.

1. Two oxen suffering from a massive mixed infestation of Cooperia spp., H. placei, O. radiatum and B. phlebotomus were introduced into a small camp, 5.4 morgen (about 1½ acres) in extent in May, 1955, followed by three artificially infested calves in October.

2. A kraal, with a surface area of approximately 1,500 sq. ft. was constructed and the infested stock herded into this kraal from 7 a.m. to 2 p.m. every day.

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\* A kraal is a cow yard or cattle pen.

The grass growing in the kraal and in a 3 ft. strip around the kraal was removed with a hoe. The kraal contained a raised water trough from which no leakage was allowed.

3. Calf No. 7180 was born on 15.11.55 and confined to the kraal from 19.11.55 until it was slaughtered on 2.8.56. Its dam grazed elsewhere but was allowed into the kraal from 7 a.m. to 2 p.m. daily to allow the calf to suckle. This was the calf's only food supply until February, when it was allowed access to an adjacent calf pen to feed on chaffed lucerne in a special stanchion constructed to prevent spillage on the ground. This calf pen will be described in experiment B.

4. Twice weekly initially but thereafter once a week faeces were collected from the calf per rectum and examined for worm eggs. At first a centrifugal flotation technique using  $ZnSO_4$ , S.G. 1.4 was employed, but as soon as the faeces were positive, egg counts, using the technique of Roberts & O'Sullivan (1950), were carried out. Positive faeces were mixed with sterile cattle dung, incubated for 8 days at  $26^{\circ}C$  and larvae collected, heat killed and identified on a percentage basis.

5. Manure was collected in the kraal at regular intervals in the following fashion. An area 10 sq. cm. was marked out on the surface, and manure removed to a depth of 2 cm., and placed in a container. A further specimen immediately below this specimen to a further depth of 2 cm. was removed and placed in a separate container. Specimens were taken to the laboratory, weighed and placed in Baermann funnels for 24 hours, larvae collected and examined as described in previous experiments (vide supra).

6. Prevailing climatic conditions were recorded daily.

7. At the conclusion of the experiment in August 1956 the calf was slaughtered, worms identified and counted.

## EXPERIMENTAL OBSERVATIONS.

This experiment is called a case report since only one calf was involved. The experimental period was from the 19th November 1955, to the 2nd of August 1956, when the calf was slaughtered. Due to circumstances beyond the authors control the post mortem was carried out 4 weeks after the faeces examination was concluded.

### (a) DEVELOPMENT OF INFESTATION.

(Results see Fig. No. 3).

Six weeks after the calf had been placed in the kraal faeces examinations were negative but the following week a sudden rise to 1,460 e.p.g.<sup>2</sup> was observed on 7.1.56. The egg counts reached their peak on 6.3.56 when 6,810 e.p.g. were noted and thereafter fell, and at the end of the experiment in July 240 e.p.g. were recorded.

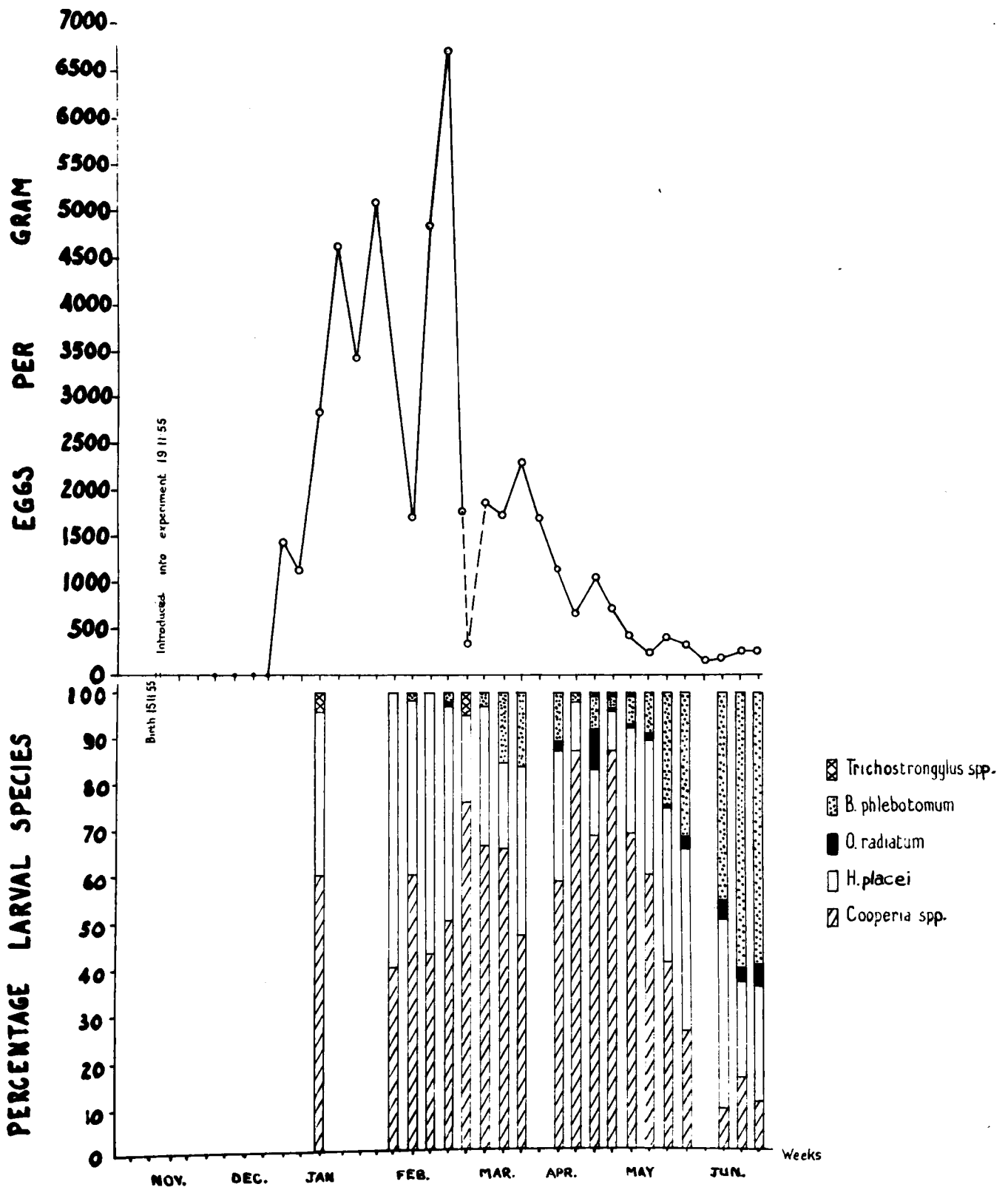
Larval cultures unfortunately were unsuccessful until 17.1.56 when Cooperia spp. H. placei and Tricho-strongylus spp. were diagnosed followed 5 weeks later by O. radiatum and B. phlebotomum. As the experiment progressed Trichostrongylus spp. larvae disappeared, Cooperia spp. and H. placei decreased, O. radiatum increased slightly and B. phlebotomum became the predominant species of larvae until the faeces examination was concluded in July.

### (b) POST MORTEM RESULTS.

At the post mortem of this calf on 2.8.56 five

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<sup>2</sup> e.p.g. eggs per gram.



**FIG. NO. 3. WEEKLY EGG PER GRAM COUNTS AND LARVAL SPECIES VARIATIONS OF**

**No. 7180**

**TABLE NO. 11. Number of worms recovered at Post mortem of calf No. 7180.**

Date of Post mortem.	<u>Cooperia pectinata.</u>	<u>Cooperia punctata.</u>	<u>Haemonchus placei.</u>	<u>Oesophagostomum radiatum.</u>	<u>Bunostomum phlebotomum.</u>
2.8.56.	504	6	35	24	140

6 There were only a few C. punctata worms present at post mortem and the total number of Cooperia spp. worms present at post mortem are included under C. pectinata.

c Including 5 Immature H. placei.

a Including 12 Immature O. radiatum.

species of worms were recovered. The species and their number are recorded in Table No. 11. Although larval culture results had revealed the presence of Trichostrongylus spp., until a few months before the post mortem, they were not recovered at post mortem.

**(c) EXAMINATION OF KRAAL MANURE -**

The results are summarised in Table No. III in the appendix, along with results pertaining to subsequent experiments. It will be noted that infective larvae of all four genera were recovered and were more frequent in March. During this month rainfalls were heavy.

**(d) PREVAILING CLIMATIC CONDITIONS -**

Daily variations in the climatic conditions are shown in Figure No. 1. Monthly rainfalls for the period are included in Table No. 10.

From Figure No. 1 and Table No. 10, it will be noted that heavy rainfalls were recorded in December, February and March. During or just after these rains larvae were

recovered from kraal manure. The role the dams teats and hair at the base of the teats plays in the transmission of verminosis to calves will be brought out in the discussion at the end of these experiments.

**B. FOUR GROUPS OF CALVES REARED BY DIFFERENT METHODS OF ANIMAL HUSBANDRY FROM MIDSUMMER TO WINTER.**

**MATERIALS AND METHODS.**

1. The same camp and kraal as described in the previous experiment were used in this experiment. However, next to the kraal in the paddock three calf pens were built as follows:

- a. A small calf pen (10 ft. x 15 ft.) was built next to the kraal. It had a ground floor and a gate leading into the kraal. In the calf pen in February, 1956, a special stanchion and trough was built. Once the calves heads were secured in the stanchion wetted chaffed lucerne was placed in the trough and there was no possibility of spillage on to the ground floor.
- b. and c. Next to the calf pen "a" two additional calf pens were built, with concrete floors. The former (b) was 15 x 20 ft. while the latter (c) was 10 x 15 ft. in size. Both had gates communicating with the paddock and lucerne and water troughs were provided.

2. Introduction of susceptible stock was unfortunately delayed awaiting the construction of the calf pens. Ten calves born between 8.11.55 and 22.12.55 were placed on concrete floors at the laboratory on 22.12.55.

Unfortunately seven of these calves had already become mildly infested and were treated with phenothiazine on 10.1.56 at the rate of 10 Grams/100 lbs. live weight with subsequent negative faeces. Before the calf pen construction had been completed 3 more calves were born and placed with the other 10 calves on concrete floors. The calf pen construction having been completed on 3.1.56 the calves were placed in

various groups. Calves born after the third of January were introduced into their groups at birth and by 3.2.56, nineteen calves were in the experiment. The calves were divided into groups as follows.

#### Group A.

Five calves (viz. Nos. 7176, 7216, 7290, 7329 & 7379) were introduced into calf pen "a". These calves were confined to the calf pen from 7 a.m. until 2 p.m.; from 2 p.m. until 7 a.m. the following morning they had access to the infested kraal by opening the communicating gate and mixed with calf No. 7180 in experiment A. Milk was fed to the calves twice daily in buckets and in February additional feed in the form of chaffed lucerne was fed in the special stanchion mentioned earlier.

#### Group B.

Five calves (viz. Nos. 7177, 7178, 7188, 7214 & 7331) were introduced into calf pen "b". These calves were confined to the calf pen and never allowed out.

The concrete floor was washed and scrubbed daily with brooms and water. The calves suckled their dams early in the morning in the calf pen for about 15-20 minutes. Thereafter the dams were herded into the infested kraal from 7 a.m. until 2 p.m. and returned to the calf pens to allow the calves to suckle for a further 15-20 minutes. After the calves had suckled, the cows were herded into a separate camp to graze overnight, returning to the calf pen just before 7 a.m. the following morning. The calves were also fed chaffed lucerne in troughs.

#### Group C.

Five calves (viz. Nos. 7193, 7227, 7292, 7234 & 7400) were introduced into calf pen "c". This was the control group. The calves were confined to the calf pen and never left it. The concrete floor was scrubbed daily with



brooms and water. Calves were fed <sup>milk</sup> in buckets twice daily as well as chaffed lucerne in troughs.

Group D.

Four calves (viz. Nos. 7207, 7262, 7353, & 7375) were introduced into the infested paddock in which the kraal and calf pens were situated. They grazed with the infested stock at night and suckled on their dams at 7 a.m. and 2 p.m. for about 15-20 minutes in each case. The calves remained in the infested paddock constantly, apart from a short period of about 20 Minutes once a week <sup>when</sup> they were herded into the crush adjacent to the kraal for faeces collection per rectum. The dams grazed elsewhere and were never allowed into the kraal.

3. Faeces examination of the calves and kraal manure examination was identical to the technique already described in the previous experiment.

4. Larval infestation of herbage was investigated from 29.2.56 onwards at regular intervals. Grass was collected from 2 plots, one adjacent to the kraal the other 300 yards away. The plots were labelled 1 and 2 and grass was collected and examined according to Taylor's (1939) technique except that in the laboratory Baermann funnels were used for larval collection and only 30 Grams of grass placed in each funnel. Larval counts were carried out using the technique already described in previous experiments. The formula as described by Taylor was used to estimate the number of larvae per lb. of grass.

5. Prevailing climatic conditions were recorded daily.

6. At the conclusion of the experiment in July, 1956, 10 calves were slaughtered for worm examination.

## EXPERIMENTAL OBSERVATIONS.

The experiments were conducted over the period 3.1.56 to 2.7.56. Due to circumstances beyond the authors control post mortems were delayed for three weeks after the faeces examinations were concluded; however all the calves used in post mortems were confined to their groups until slaughtered.

### a. DEVELOPMENT OF INVESTIGATION:

Group A. - Five hand reared calves confined to the infested kraal and adjacent calf pen.

(Results see Fig. No. 4).

Calves in this group developed infestation slowly and only after 20 weeks in this experiment did the majority of the group have egg counts exceeding 100 e.p.g. The highest individual egg count was 690 e.p.g. observed in calf No. 7176 after 22 weeks exposure to infestation.

Larval culture results revealed the presence of Cooperia spp., H. placei, O. radiatum and Trichostrongylus spp. before the average egg counts exceeded 100 e.p.g. It was significant that only when H. phlebotomus was well established did the egg counts rise to a level in excess of 100 e.p.g. for the majority of the group.

Group B. - Five suckling calves confined to a concrete floored pen; the dams were kept in the infested kraal from 7 a.m. to 2 p.m.

(Results See Fig. No. 4).

Infestation developed slowly but a sudden rise was noted 13 weeks after the experiment had commenced in the e.p.g. counts, and four weeks later all the calves had egg counts in excess of 100 e.p.g. One calf had 1,960 e.p.g. at this stage (vis. No. 7214). A week after the egg counts had reached their peak, i.e. 18 weeks after the

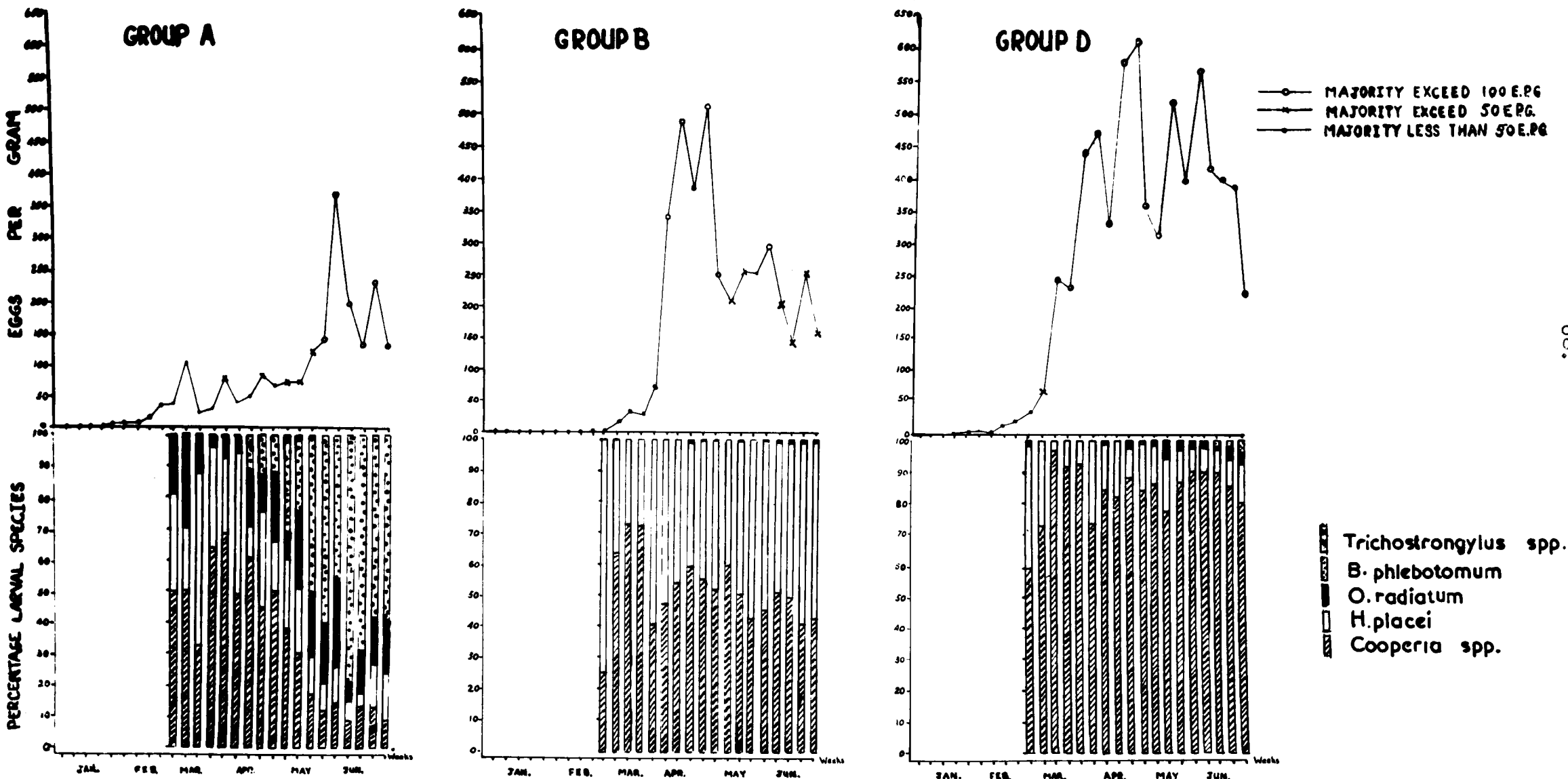


FIG. No. 4. WEEKLY MEAN EGG PER GRAM COUNTS AND LARVAL SPECIES VARIATIONS OF GROUPS A, B AND D.

68.

experiments commenced, egg counts fell dramatically and remained at a low level, most of the calves having egg counts between 50 and 100 e.p.g. until the experiment was concluded.

Larval culture results revealed a predominant Cooperia spp. and H. placei infestation. When egg counts were high more H. placei larvae were recovered from cultures but the reverse was the case when egg counts fell. Larval cultures revealed very little O. radiatum and even less B. phlebotomus infestation.

Group C. Five hand reared calves in a concrete floored pen.

Only one calf in this group developed infestation of any note (calf No. 7193), reaching a peak of 140 e.p.g. in May and falling to 20 e.p.g. at the end of the experiment. Another calf developed a very mild infestation and faeces examinations were negative for 5 weeks before the experiments ceased in July. The rest of the calves were negative throughout.

Larval cultures of Calf No. 7193 revealed the presence of H. placei, Cooperia spp. and O. radiatum. The cultures in most cases were very poor indeed and seldom yielded enough larvae to differentiate the species on a percentage basis.

The chaffed lucerne fed to these calves was found to have become infested with the species found in Calf No. 7193 and this was probably the source of infestation; chaffed lucerne fed to other groups was not infested.

Group D. Four suckling calves confined to the infested paddock.

(Results see Fig. No. 4).

Eleven weeks after the experiment commenced all the calves in this group had egg counts exceeding 100 e.p.g., which rose to a mean of 650 e.p.g. six weeks later and remained at a high level only falling to 255 e.p.g. at the end of the experiment.

Cooperia spp. larvae were dominant in cultures although the other species were also recovered.

**b. POST MORTEM RESULTS.**

The results of post mortems are summarised in Table No. 12, and confirmed the findings and observations of faecal examinations of the various groups.

**TABLE NO. 12 - Number of worms recovered at post mortem of calves in Experiment B.**

Group.	Calf No.	Date of Post Mortem.	<u>Cooperia pectinata.</u>	<u>Haemonchus placei.</u>	<u>Gesophagostomum radiatum.</u>	<u>Bunostomum phlebotomum.</u>	<u>Trichouris globulosa.</u>
A.	7216	24.7.56	5	-	4	2	2
	7290	26.7.56	-	6	-	22	2
	7329	26.7.56	-	-	-	98	-
B.	7177	31.7.56	29	1	-	1	-
	7178	31.7.56	9	10	-	-	-
	7188	31.7.56	8	9	-	-	-
C.	7193	8.8.56	30	5	5	-	-
D.	7262	3.8.56	742	-	5	25	-
	7353	7.8.56	900	1	10	19	-
	7375	8.8.56	425	-	6	58	-

**c. EXAMINATION OF KRAAL MANURE.**

The results of these examinations are summarised in Table No. III in the appendix; larvae of all four genera were recovered particularly in March.

**d. LARVAE PER POUND OF HERBAGE.**

These results are summarised in Table No. IV in the appendix. It will be noticed that larvae were recovered in large numbers in February, March and up to the 24th of April, particularly in Plot 1 next to the kraal. Larval recoveries in the winter months were very sporadic.

**e. PREVAILING CLIMATIC CONDITIONS.**

These are shown in Fig. No. 1 and Table No. 10. Heavy rainfalls were recorded in February and March less rain in January and little or no rain from April to May. Kraal manure and grass were more heavily infested with larvae in the months when rainfalls were good and calves showed evidence of increased egg counts in faeces examination a month or so after rains had started at the end of February.

**C. FIVE GROUPS OF CALVES REARED BY DIFFERENT METHODS OF ANIMAL HUSBANDRY FROM AUTUMN TO MIDSUMMER.**

**MATERIALS AND METHODS.**

1. The same paddock, kraal and calf pens were used as in the previous experiment and calf faeces, kraal manure and grass examination was carried out as mentioned previously. Climatic data were also recorded.

2. Introduction of calves into the groups in this experiment was carried out in a different fashion than in experiment B. Calves were not available from the large breeding herd on the station in the autumn because the breeding policy at Arcoedsvlakte only made provision for summer calving. The author, however, had been given 50 cows

for another experiment in April of the previous year. Two bulls were allowed to run alternately with the cows, each bull being with the herd for one month and resting for the following month until January, 1956. This led to irregular calving of the cows over a period of nearly seven months (i.e. February-September, 1956). At birth the calves were introduced into the various groups and ran with the calves in the previous experiments and by the time the previous experiments (A & B) were brought to a conclusion 75% of the calves used in this experiment were already born and placed in their respective groups.

3. Twenty-six calves were used in this experiment, divided into five groups; there were five calves in each group, apart from group 3 in which there were six calves. The division into groups 1 to 4 and treatments were the same as in previous experiments. Groups 1, 2 and 3 were the same as groups A, B and D in experiment B, while group 4 was a duplication of experiment A.

Group 5 was a new group. Five calves were confined to the infested kraal along with calves in group 4 from 7 a.m. to 2 p.m. and allowed to suckle on their dams there. From 2 p.m. to 7 a.m. they grazed with the infested stock and calves in group 3 in the infested paddock. Their dams grazed elsewhere. The full details of the division into groups is shown in Table No. V. in the appendix.

NOTE: Due to the shortage of calves, the approach of winter conditions and the fact that only two calves became mildly infested in experiment B in the control group (Group C) it was felt that no useful object would be served in continuing with a similar group in this experiment.

4. Four calves (No's. 7176, 7207, 7214 and 7379) that became infested in experiment B were not slaughtered for post mortem in July when the other calves in that



experiment were autopsied. They were kept to act as infested stock along with the other infested stock mentioned in experiment A under materials and methods (1 and 2). They ran with the older infested stock in the paddock and kraal and remained there until May of the following year (1957).

EXPERIMENTAL OBSERVATIONS.

The experimental period was from February, 1956, to January, 1957. The first calf (vis. No. 8227) was born on 29.2.56 and the last post mortem carried out on 30.1.57. Negative calves were, however, discharged on 15.1.57. In addition post mortems were also carried out in October and November.

(a) DEVELOPMENT OF INFESTATION.

Group 1. - Five handreared calves confined to the infested kraal and adjacent calf pen.

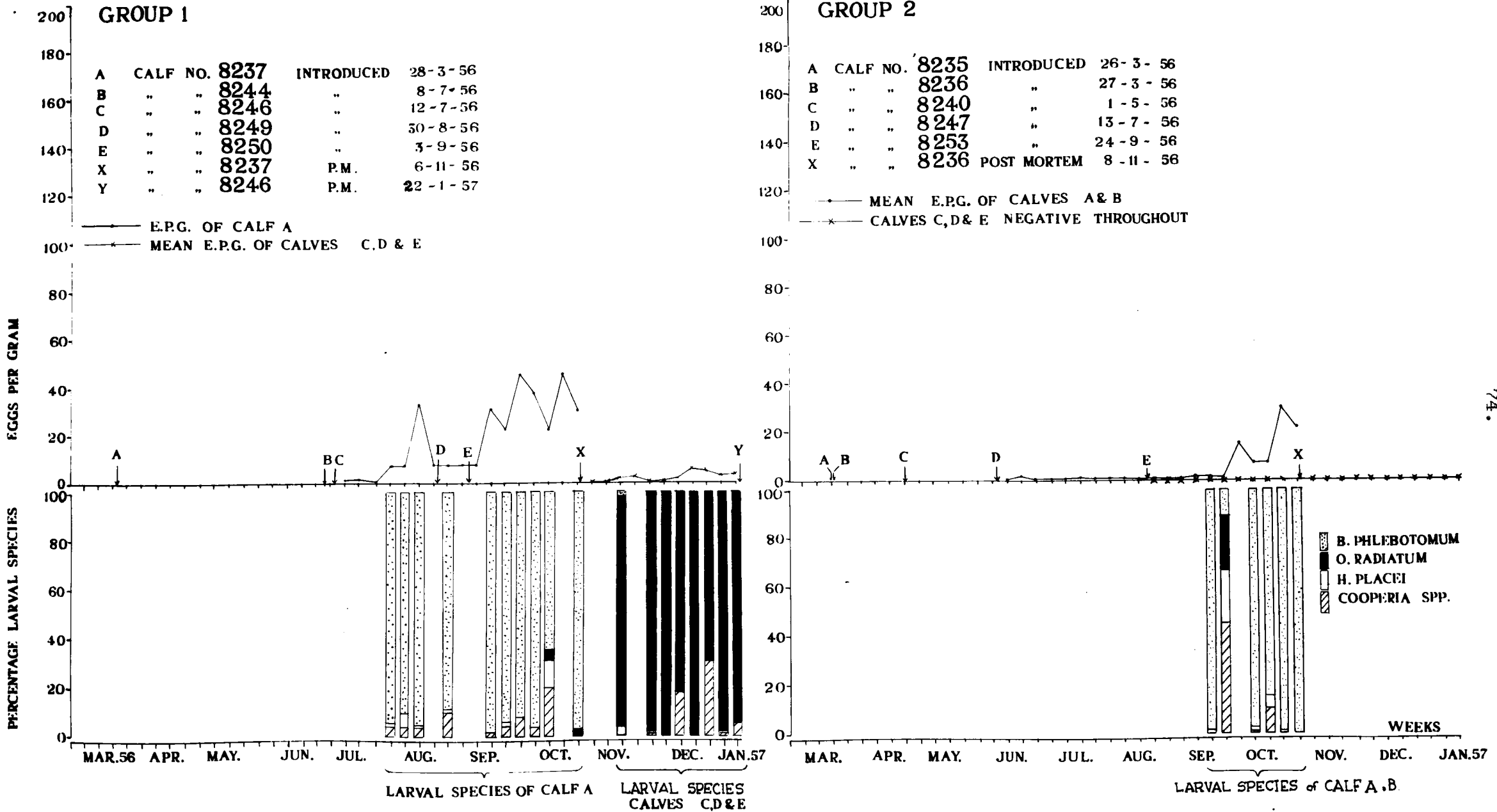
(Results see Fig. No. 5)

Of the five calves in this group one (No.8237) born in March became positive on faeces examination in the winter, egg counts reaching a peak in October and November. This calf was slaughtered in November for post mortem. The balance of the group were born from July to September and only three of the calves became mildly infested in the summer the maximum egg counts for any calf being 22 e.p.g. in Calf No.8246. One calf was negative throughout (No.8244).

Larval cultures of the calf showing infestation in winter were predominantly B. phlebotomus, while O. radiatum was the most common larval species recovered from cultures of calves that become infested in summer.

Group 2. - Five suckling calves confined to a concrete floored pen; the dams were kept in the infested kraal from 7 a.m. to 2 p.m.

(Results see Fig. No. 5).



Only two calves born in March in this group became infested the other three being negative throughout i.e. calves born from May until September. The two infested calves had very low egg counts reaching their peak in October when one calf became negative the other maintaining its infestation until slaughtered in November.

Larval cultures showed a predominant B. phlebotomus infestation.

Group 3. - Six suckling calves confined to the infested paddock.

(Results see Fig. No. 6).

Two of the six calves in this group born before the end of March became infested at 6-7 weeks of age, their egg counts rising until June and then falling to show another rise in October and November. These two calves were slaughtered at the peak of their egg production (viz. Nos. 8227 & 8229). The other two calves born in September showed a very low egg count in December and January.

Larval cultures from the calves infested in autumn showed the presence of Cooperia spp. at first, followed by H. placei. By August H. placei and B. phlebotomus were predominant. A. radiatus was diagnosed in September. Larval cultures of the two calves that became infested in the summer indicated the presence of H. placei and Cooperia spp.

Group 4. - Five suckling calves confined to the infested kraal and adjacent calf pen.

(Results see Fig. No. 6).

All five calves in this group were born in March and all of them became infested, the egg counts showing a sharp rise in July followed by another peak in October.

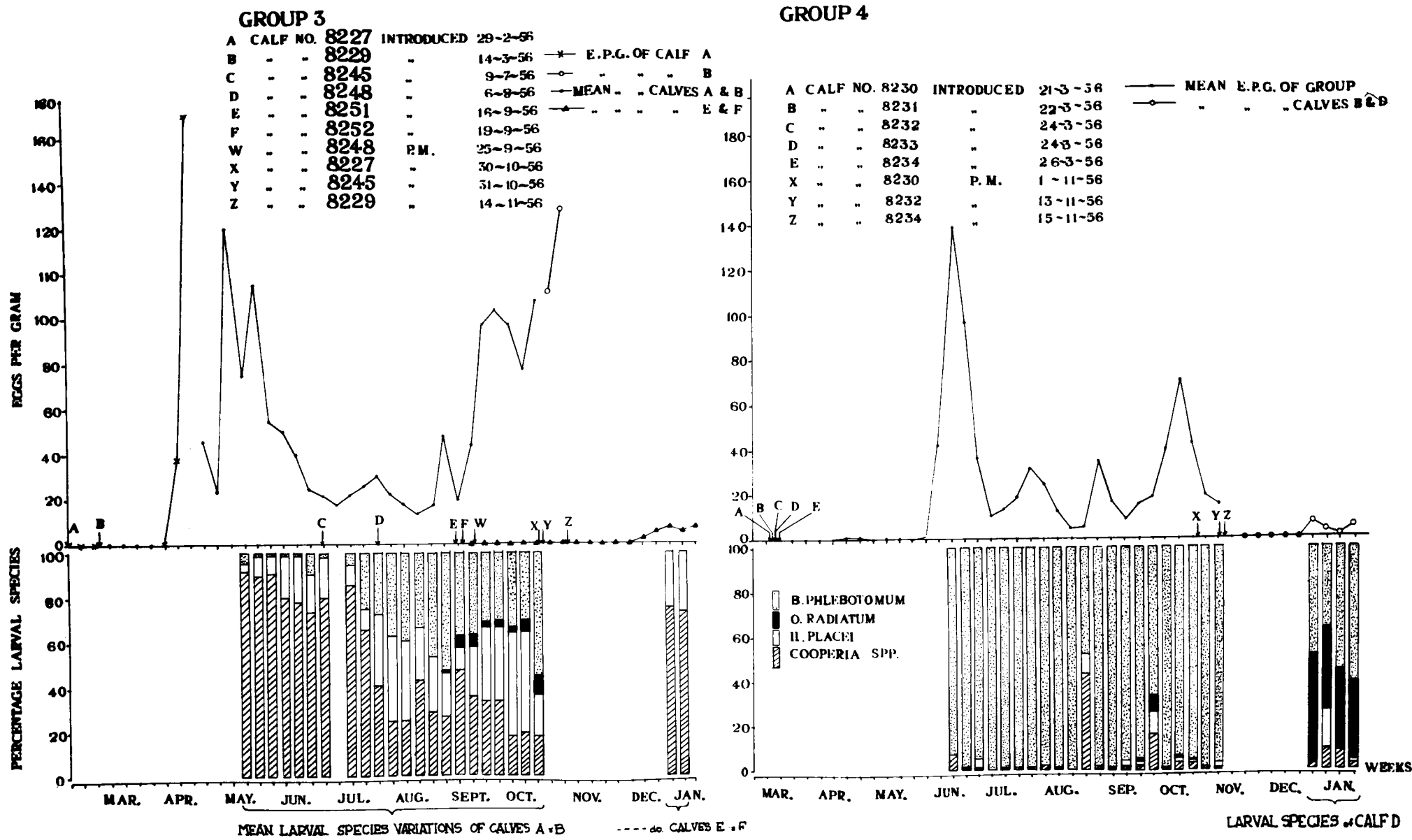


FIG. NO. 6 WEEKLY MEAN EGG PER GRAM COUNTS & LARVAL SPECIES VARIATIONS OF GROUPS 3 & 4

However, two of the calves became negative in October and November (Nos. 8231 & 8233); in one calf (No. 8231) this was maintained until the experiment was concluded, while the other calf (No. 8233) again became positive in December and January although the egg counts remained at a low level.

Larval cultures consisted almost entirely of B. phlebotomus in the winter but O. radiatus and other species of larvae were also recovered from faeces of No. 8233 in the summer.

Group 5. - Five calves that suckled<sup>in</sup> and were confined to the infested kraal from 7 a.m. - 2 p.m. and grazed in the infested paddock from 2 p.m. to 7 a.m. The dams grazed elsewhere.  
(Results see Fig. No. 7).

The calves in this group were born from 14/4/56 to 6/6/56. They were negative throughout the winter but all of them became infested within a short period of each other from the 18th of December onwards. Egg counts showed a steady rise until the experiments were concluded in January.

Larval cultures revealed the presence of Cooperia spp initially which were soon followed by H. placei; O. radiatus only made its appearance in January.

(b) POST MORTEM RESULTS.

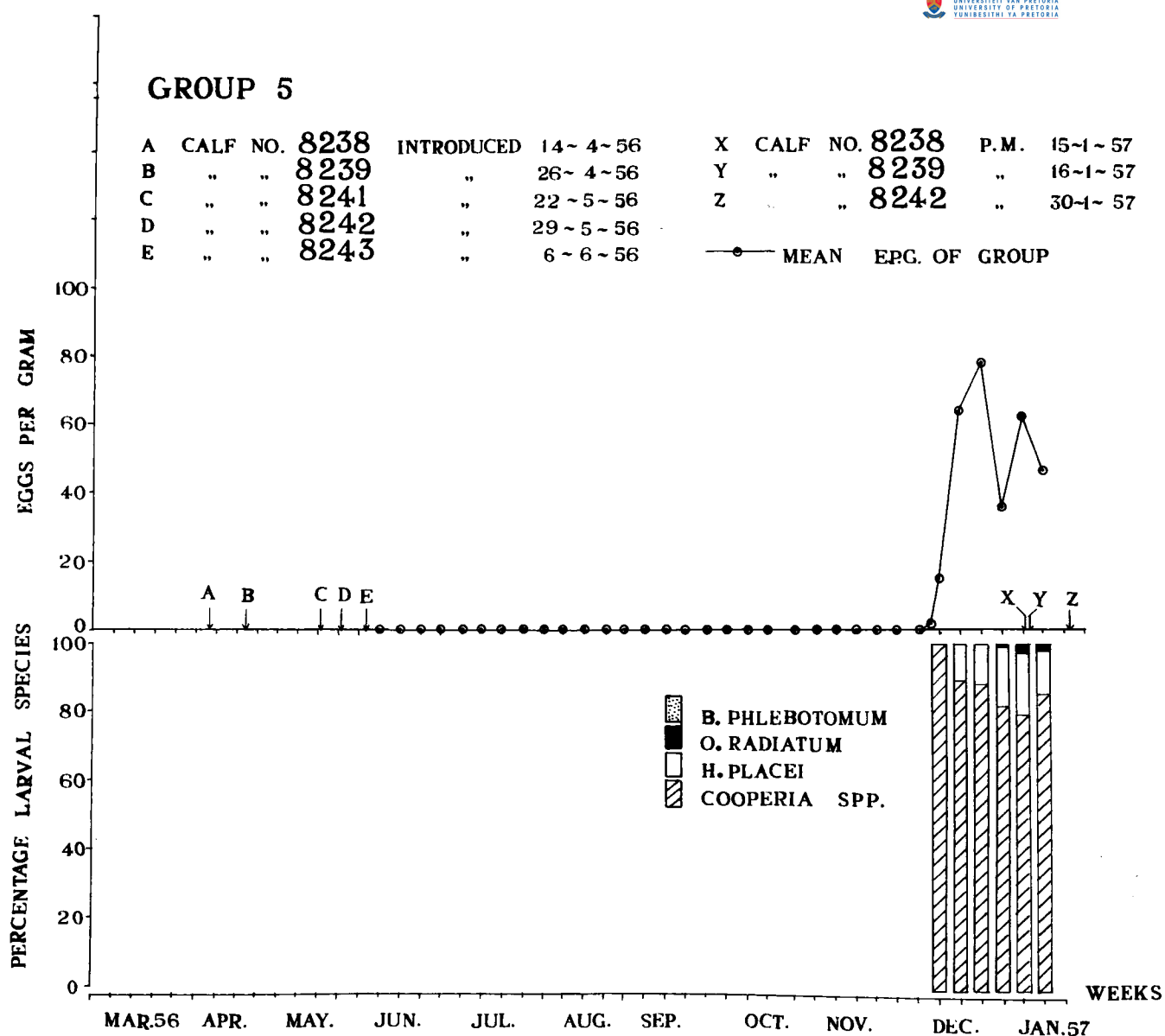
(cf Table No.13).

From the results summarised in Table No. 13 the most striking observations are those made on calves No's. 8227 and 8229 (Group 3). Although all four species were recovered from these calves at post mortem, the presence of immature H. placei worms was particularly interesting. The finding of these immature worms led to slaughtering of calf No. 8245

**GROUP 5**

A	CALF NO.	8238	INTRODUCED	14~4~56	X	CALF NO.	8238	P.M.	15~1~57
B	"	8239	"	26~4~56	Y	"	8239	"	16~1~57
C	"	8241	"	22~5~56	Z	"	8242	"	30~1~57
D	"	8242	"	29~5~56					
E	"	8243	"	6~6~56					

—●— MEAN EPG. OF GROUP



**FIG. NO. 7. WEEKLY MEAN EGG PER GRAM COUNTS & LARVAL SPECIES VARIATIONS OF GROUP 5**

78.

which had been negative on faecal examination. This calf in spite of diligent search, including microscopical examination of mucosal scrapings of the abomasum, was negative for parasites (Table No. 13). Another calf (No. 8248), in this group had died of torsion of the colon in September and was similarly negative at post mortem. Both these calves were born in July and August respectively (Table No. V in the appendix). The calves that showed both mature and immature H. placei worms at post mortem were born in February and March (Table No. V in the appendix). The remaining two calves (No's 8251 and 8252) in this group, born in September were negative for H. placei until the 31st of December.

Since only the calves that were born before the end of March showed immature worms at post mortem in October and November, whereas the other calves under identical conditions, but born later showed no worm infestation either by faecal examination or at post mortem, it would appear that the immature worms had been acquired prior to winter and had remained immature until the post mortems were carried out. Furthermore five calves in group 5 born from April to June were negative for H. placei until the end of December. These calves grazed in the same paddock as those in group 3. Their freedom from infestation substantiated, the supposition that the paddock was free of infestation from April to November.

(c) EXAMINATION OF KRAAL MANURE.

These results are summarised in Table III in the appendix. Larvae were consistently recovered from kraal manure in March and up to the 19th of April. Thereafter results were inconsistent; from the end of November to the early part of January, larvae were recovered every week.



**TABLE NO. 13 - Number of worms recovered at Post Mortem of Calves in Experiment C.**

Group.	Calf No.	Date of Post Mortem.	<u>Cooperia pectinata.</u>	<u>Haemonchus placei.</u>	<u>Oesophagostomum radiatum.</u>	<u>Bucestomum phlebotomum.</u>	<u>Trichouris globulosa.</u>
1.	8237	6.11.56	-	-	-	28	1
	8246	22.1.57	3	-	35	1	-
2.	8236	8.11.56				34	
	8248	25.9.56	-	-	-	-	-
3.	3227	30.10.56	2	*341	11	32	-
	8245	31.11.56	-	-	-	-	-
	8229	14.11.56	97	*667	9	60	-
4.	8230	1.11.56	-	-	-	27	-
	8232	13.11.56	-	-	-	74	-
	8234	15.11.56	-	1	-	32	-
5.	8238	15.1.57	27	171	-	-	
	8239	16.1.57	228	6	25	2	
	8242	30.1.57	280	24	3	9	

\* Including 50 Immature H. placei.

◆ Including 36 Immature H. placei.

.....

(d) LARVAE PER FOUND OF HERBAGE.

These results can be seen in Table No. IV in the appendix. The periods when infective larvae were more plentiful were similar to those mentioned under kraal manure investigations but the summer period was shorter; consistent results were only obtained for three weeks after the 20th of November, and rarely thereafter until the end of January.

(e) PREVAILING CLIMATIC CONDITIONS.

The daily variations in the climatic conditions will be noted in Figs. No. 1 & 2 and the monthly rainfalls for the period of this experiment are included in Table No. 10.

It will be noted that the only months the rainfall exceeded 50 mm. in the experimental period were March, December 1956 and January 1957.

D. SEVEN GROUPS OF CALVES REARED BY DIFFERENT METHODS OF ANIMAL HUSBANDRY FROM SUMMER TO EARLY WINTER.

MATERIALS AND METHODS.

1. The same paddock, kraal and calf pens were used as in previous experiments; grass was similarly examined and climatic data recorded daily.

2. Thirty five calves were used. Those born from 31/10/56 to 13/11/56 were confined to concrete floored pens at birth and placed in their various groups on 13/11/56. Calves born after 13/11/56 were immediately placed in their various groups. By 13/12/56 all the groups were complete

Information of the date calves were born and placed in groups etc., is given in Table No. VI in the appendix. Each group contained five calves.

- (a) Groups 1 to 5 were the same as Groups 1 to 5 in experiment C. Calves of similar groups in both experiments ran together until the end of January when experiment C was concluded.
- (b) Group 6 the control group was a duplication of Group C in experiment B.
- (c) Group 7: The calves in this group were confined to a calf pen with a concrete floor near the stables, away from the infested paddock. The dams udders were washed before the calves were allowed to suckle at 7 am and 2 pm. This was regarded as an additional control group.

3. Faeces collections were carried out as previously described, but the laboratory techniques were modified, in that 40% sucrose solution was used for both centrifugation and egg counting instead of  $ZnSO_4$  and NaCl solutions. Eggs were easier to see microscopically and higher egg counts were obtained when sucrose solution was compared with the NaCl solution on the same faeces. However more air bubbles were trapped in the emulsion after shaking when sucrose was used, but with the addition of a little amyl alcohol this was overcome. Before counting it was essential that slides be left for at least three minutes to allow the eggs to rise to the top. Larval examinations were carried out as described previously.

4. The examination of kraal manure was carried out as before until February. Thereafter the technique was modified as follows.

- (a) Manure was collected as described previously -
- (b) Specimens were brought to the laboratory, hard lumps broken into small pieces, and each specimen thoroughly mixed; two samples weighing 50 grams each from each specimen were placed in separate Baerman funnels.
- (c) Twenty grams of the residue of each specimen were thoroughly mixed; 50 grams of this mixture were well mixed with 1,000 active infective larvae and placed in a Baermann funnel to act as a control.
- (d) The larvae were collected, and examined with the microscopical technique described in previous experiments.
- (e) The number of larvae per kg. of manure were estimated using the same formula advocated by Taylor (1939) for the estimation of larvae per lb. of herbage.

5. Larvae adhering to the teats of cows and hairs at the base of teats were recovered in the following fashion:-

The teats and hair at the base of the teats of six cows in the infested Kraal were wiped off with a sterile cloth into a recently boiled container. Thereafter another sterile cloth was used to wash the teats into another sterile container. Both dry material and teat washings were placed in separate Baermann funnels, larvae collected and examined in the manner described previously.

#### EXPERIMENTAL OBSERVATIONS.

The experimental period was from 13/11/56 to 20/5/57

##### (a) DEVELOPMENT OF INFESTATION.

Group 1. - Five hand-reared calves confined to the

infested kraal and adjacent calf pen.

(Results see Fig. No. 8).

Infestation developed extremely slowly in this group and only after 11 weeks in the experiment were all the calves infested. One calf died of hoven on 27/2/57 and only a single hookworm was recovered from it. Egg counts reached their peak in March and by the end of April two of the calves were negative.

Larval cultures indicated the presence of four genera of Nematodes.

Group 2. - Five suckling calves confined to a concrete floored pen; the dams were kept in the infested kraal from 7 a.m. - 2 p.m.

The calves in this group remained negative throughout and were discharge in May.

Group 3. - Five suckling calves confined to the infested paddock.

(Results see Fig. No. 8).

Worm eggs were noticed in the faeces of two calves six weeks after the experiment commenced; four weeks later all the calves were infested; egg counts reached their peak at fourteen weeks, to <sup>f</sup>all thereafter until the experiment was concluded.

Cooperia spp. and H. placei larvae were the first species recovered from cultures and only towards the end of the experimental period were larvae of O. radiatum and B. phlebotomum recovered.

Group 4. - Five suckling calves confined to the infested kraal.

(Results see Fig. No.8).

Infestation developed very slowly in this group, only two calves being infested after the first six weeks;

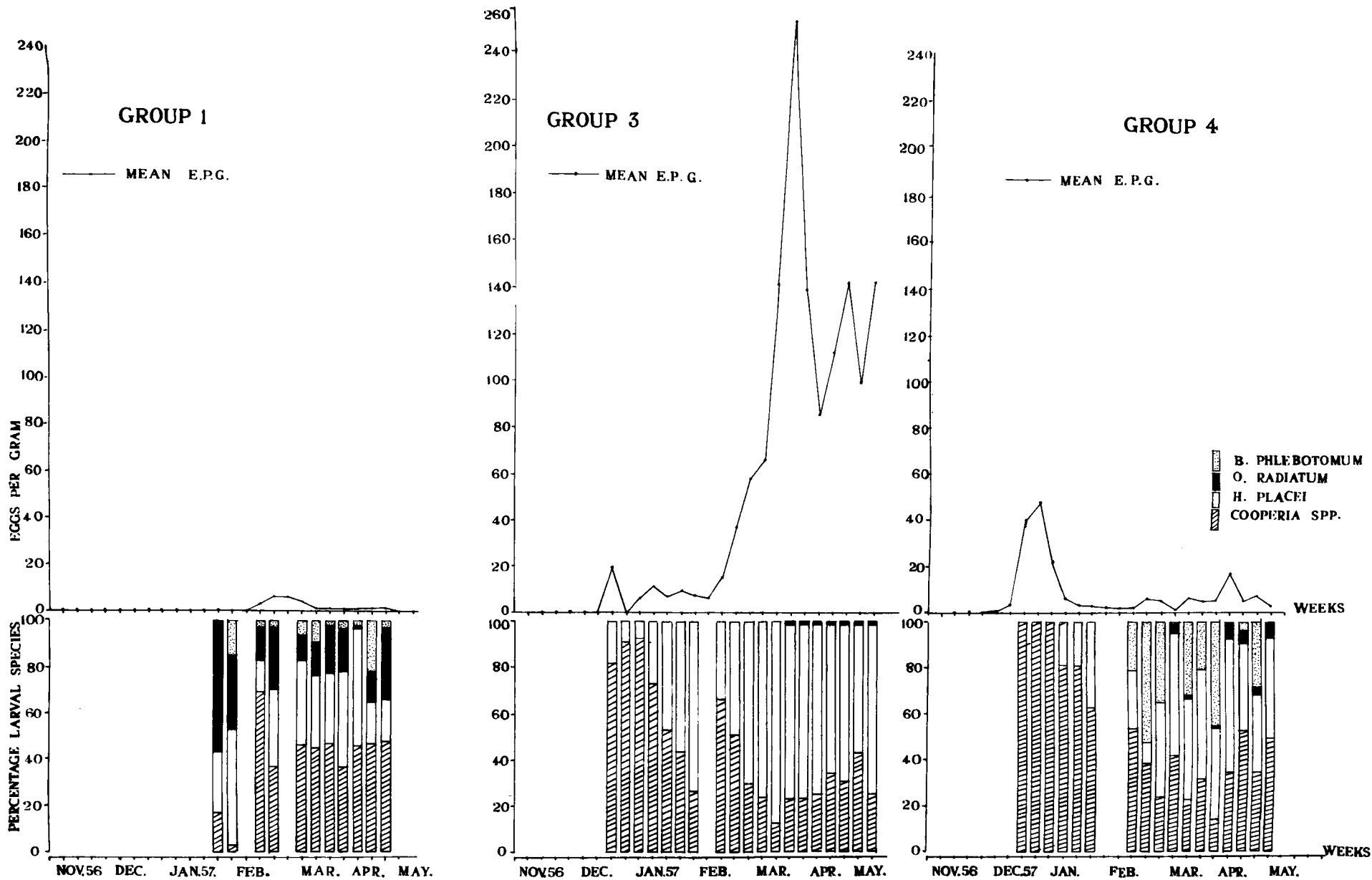


FIG. NO. 8. WEEKLY MEAN EGG PER GRAM COUNTS & LARVAL SPECIES VARIATIONS OF GROUPS 1 3 & 4

nine weeks later all the calves were infested. Egg counts never reached a high level although a slight rise was recorded a few weeks before the termination of the experiment.

Larval cultures at first only contained Cooperia spp; these were followed by H. placei; a few weeks later by B. phlebotomum and finally O. radiatum larvae were also present.

Group 5. - Five calves that suckled<sup>in</sup> and were confined to the infested kraal from 7 a.m. - 2 p.m., and grazed in the infested paddock from 2 p.m. - 7 a.m. The dams grazed elsewhere.

(Results see Fig. No. 9).

The infestation of calves in this group were first noted in two calves after five weeks; within another month all the calves were infested. Egg counts reached their peak after fourteen weeks and fell slowly until the experiments were concluded in May.

NB. The mean e.p.g. counts of this group were more than double those of any other infested group.

Cooperia spp. larvae were the first species recovered in cultures made on 18/12/56 from two calves; cultures made ten days later showed H. placei larvae. O. radiatum and B. phlebotomum larvae were recovered from cultures on 7/1/57 and 28/1/57 respectively.

Groups 6 and 7. - There were 5 calves in each group confined to concrete floors. Calves in the former group were hand reared; the latter suckled on previously washed udders.

Calves in these groups were negative throughout and were discharged in May at the conclusion of the experiment.



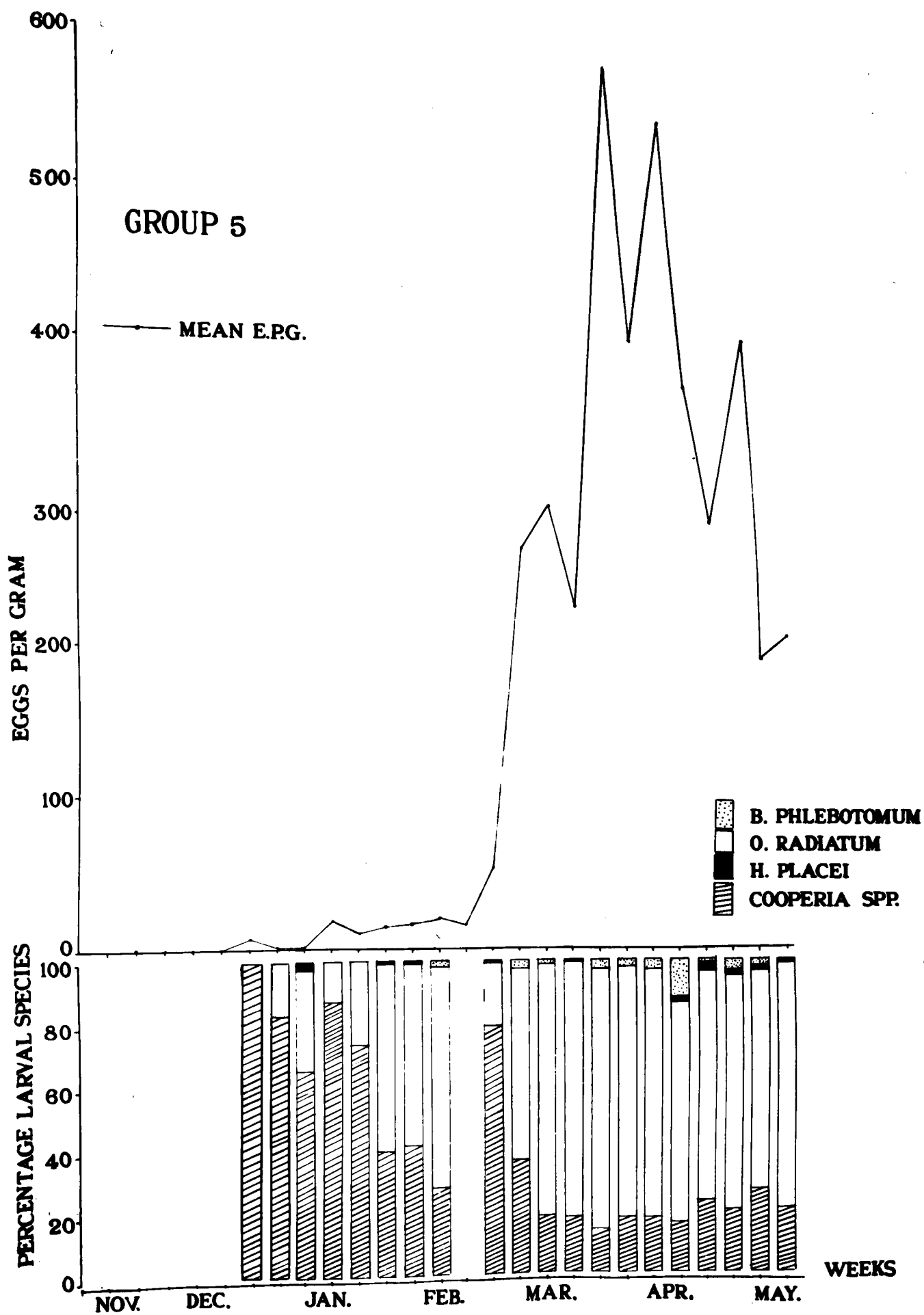


FIG. NO. 9. WEEKLY MEAN EGG PER GRAM COUNTS & LARVAL SPECIES VARIATIONS OF GROUP 5.

ERRATUM.

Page 87, Figure 9.

Where the key reads:

- O. RADIATUM
- H. PLACEI

It should read:

- O. RADIATUM
- H. PLACEI

TABLE NO. 14 - Number of worms recovered at post mortem of calves in Experiment B.

Group.	Calf No.	Date of post mortem.	<i>Cooperia pectinata.</i>	<i>Haemonchus placei.</i>	<i>Oesophagostomum radiatum.</i>	<i>Lunostomum phlebotomum.</i>	<i>Trichuris globulosa.</i>	<i>Moniezia benedini.</i>
1.	14	27/2/57	-	-	-	1	-	-
	15	1/5/57	-	-	1	4	1	-
3.	51	2/5/57	173	162	7	1	-	-
	52	8/5/57	13	29	-	-	-	-
	36	14/5/57	69	187	1	1	-	1
	59	15/5/57	308	164	3	2	-	-
	25	16/5/57	133	173	5	3	-	1
4.	50	1/5/57	3	1	-	19	1	-
	24	7/5/57	10	7	-	1	-	-
	26	8/5/57	7	8	5	4	-	-
5.	27	30/4/57	950	535	7	3	-	1
	6	6/5/57	95	478	2	12	-	-
	60	9/5/57	144	183	22	2	-	-
	87	13/5/57	103	169	3	18	-	-
	17	20/5/57	68	97	-	11	-	-

(b) POST MORTEM RESULTS.

These results are summarised in Table No. 14.

Apart from Calf No. 14 that unfortunately died of hoven in February, all the calves whose faeces were positive at the end of the experiment were slaughtered. It is not necessary to elaborate on these results, as the table clearly indicates the severity of infestation in the various groups, viz. Groups 5, 3, 4 and 1 respectively in order of severity.

(c) EXAMINATION OF KRAAL MANURE.

The results of these investigation using the old technique until the end of January are summarised in Table No. III in the appendix. Larvae were recovered regularly every week from specimens collected from the end of November to the middle of January.

From February the modified technique as described earlier was used and the results are summarised in Table No. IIIa in the appendix. Weekly examinations were carried out in the period 5/2/57 to 30/4/57; the mean number of larvae per Kg of kraal manure varied from 0 to 986.

(d) LARVAE PER POUND OF HERBAGE.

These results are summarised in Table No. IV in the appendix.

Infective larvae were only recovered from herbage irregularly in November and December, and from February to April.

(e) RECOVERY OF INFECTIVE LARVAE FROM THE TEATS OF COWS.

**TABLE NO. 15 - Infective larvae recovered from the teats of cows that had been confined to the infested kraal from 7 a.m. to 2 p.m.**

Date specimens collected.	No. of larvae recovered from dry material on the teats.	No. of larvae recovered from test washings.	<u>Cooperia</u> spp.	<u>H. placei</u> .	<u>O. radiatum</u> .	<u>B. phlebotomum</u> .
22/11/56	0	0	-	-	-	-
30/11/56	2	0	-	1	1	-
5/12/56	7	2	1	1	-	7
12/12/56	0	0	-	-	-	-

Some of the results of this examination are shown in Table No. 15. Seven further examinations at weekly intervals after 12/12/56 were negative and are not shown. However all species of infective larvae were recovered from the teats as shown in the Table. This point that larvae can adhere to teats, being proved, it was not necessary to continue this examination.

(f) PREVAILING CLIMATIC CONDITIONS.

Daily variations in climatic conditions are shown in Figure No. 2 and monthly rainfalls are included in Table No. 10. The total rainfall from 30/10/56 to 20/5/57, the complete period over which this experiment was conducted, was 290.2 mm. (11.43 inches).

(g) THE EFFECT OF RAINFALL ON THE TRANSMISSION OF VERMINOSIS.

An excellent opportunity existed, when this

experiment began to study the effects of rainfall on the transmission of verminosis, for the following reasons:-

(1) Prior to the commencement of this experiment on 13/11/56, conditions were very dry and only 13.4 mm and 18.3 mm of rain had been recorded for the months of September and October respectively (Table No. 10).

(2) The presence of calves born in the winter and early spring which were still worm free when this experiment commenced and remained worm free on faecal examination until the 18th of December (Experiment C).

(3) The introduction of 26 newly born calves between 13/11/56 and 21/11/56 in a dry period, before the summer rains began on the 22nd of November.

(4) The fact that some of the worm free calves of the previous experiment (C) and a few newborn calves of this experiment (D) become positive on faecal examination, on the same day, and with the same succession of species thereafter.

The abovementioned reasons showed that conditions must have been favourable to set up infestation.

The minimum period necessary for infective larvae of the different species to reach maturity in the host, i.e. the prepatent period was known (Table No. 16); consequently the probable date of infestation could be determined. The rainfall recorded after the calf birth and before the day the calf became infested, would then give the amount of rainfall necessary, to render infective larvae available to the host in numbers large enough to cause infestation (Table No.17).

## DISCUSSION.

### (a) ANIMAL HUSBANDRY METHODS.

The transmission experiments were conducted in a manner as closely allied as possible to the methods of animal husbandry practiced in the district. The various groups were attempts at separating the various facets, thought to be

possible sources of infestation to calves. The experiments showed that the sources of infestation were:

- (1) Herbage.
- (2) Kraal manure adhering to tests.
- (3) Kraal manure by direct contact.

(1) Herbage:

The paddock was the most constant source of infestation both during wet and comparatively dry years, (Group D Expt. B, and Group 3 Expts. C and D). At post mortem C. pectinata and H. placei were the most common species recovered, and to a lesser extent B. phlebotomum and O. radiatum.

TABLE NO. 16. The minimum period for infective larvae to reach maturity in calves.

Species	No. of days that infective larvae require to develop to adults in calves: - prepatent period	Authority.
<u>Cooperia punctata</u>	11	Bailey (1949)
<u>Cooperia pectinata</u>	16	Own observations
<u>Haemonchus placei</u>	19	Roberts (1957)
do. do.	26-28 (average)	Roberts (1957)
do. do.	14	Mayhew (1941)
do. do.	23-26 (average)	Mayhew (1941)
<u>Oesopharostomum radiatum</u>	35	Mayhew (1948)
<u>Bunostomum phlebotomum</u>	52	Mayhew (1948)
do. do.	56	Sprent (1946 a)

⊙ At post mortems of calves in experiments (C) and (D) no C. punctata worms were recovered.

✱ Minimum period between dosing pure cultures of C. pectinata larvae to calves at Armoedsvlakte and appearance of worm eg of this species in faeces. The calves were worm-free prior to dosing and kept under worm-free conditions until faeces were positive.

**TABLE NO. 17. The minimum rainfall necessary at Armoedsvlakte for infective larvae to be present, in numbers large enough to infest calves.**

<u>SPECIES.</u>	<u>TOTAL RAINFALL FROM DATE OF CALF'S BIRTH TO PROBABLE DATE ON WHICH THE CALF BECAME INFESTED.</u>		<u>DATE ON WHICH INFESTATION PROBABLY OCCURRED.</u>	<u>MINIMUM PREPARENT PERIOD IN DAYS.</u>	<u>DATE ON WHICH EGGS WERE FIRST DETECTED IN PAGES OF CALVES.</u>
	<u>mm</u>	<u>inches</u>			
<u>C. pectinata</u>	13.3	0.52	2/12/56	16	18/12/56
<u>H. placei</u>	13.3	0.52	2/12/56	26	28/12/56
do. do.	42.3	1.59	9/12/56	19	28/12/56
do. do.	72.8	2.87	14/12/56	14	28/12/56
<u>O. radiatus</u>	13.3	0.52	3/12/56	35	7/1/57
<u>B. phlebotomus</u>	42.3	1.59	7/12/56	52	28/1/57

H Three alternatives given for H. placei, see discussion.

x Note: Sprent's (1946 a) observation shown in Table No. 16 not included in this table - see discussion.

(2) Kraal manure adhering to teats.

Cooperia spp., H. placei, O. radiatus and B. phlebotomus larvae were recovered from both dry material on and washings of, the teats of cows that had been lying in the infested kraal (b Tab. No. 15). Larvae adhering to the teats transferred infestation from the infested kraal to suckling calves in a nearby concrete floored pen, during a particularly wet autumn (Group B. Expt. B. Group 2 Expt G), but not during a dry summer and autumn (Group 2 Expt. D).

b Tab. is Table.



Although all species of larvae were recovered from cultures (Fig 4) only C. pectinata, H. placei and B. phlebotomum were recovered at post mortem (Group B. Tab. No. 12), and in a later experiment (C) only B. phlebotomum (Group 2 Tab.No.13).  
(3) Kraal manure by direct contact.

Hand reared calves with access to the calf pen and infested kraal were predominantly infested with B. phlebotomum (Group A Tab. No. 12; Group 1 Tab. No's 13 & 14) and only mildly infested with C. pectinata, O. radiatum and H. placei (Group A. Tab. No. 12).

Suckling calves in the infested kraal not only walked in infested manure all day, but were forced to suckle on their dams teats after the cow had been lying in the infested kraal manure during the morning i.e. the combined effects of (2) and (3) above. At post mortem they had heavier worm burdens (Group 4 Tab. 13 and 14) when compared with hand reared calves in the infested calf pen and kraal (Group 1 Tab. No. 13 and 14). Under the influence of particularly good rains in summer and autumn, one suckling calf (viz. No. 7180) confined to the infested kraal, acquired a heavy mixed worm burden (Expt. A.\*Fig. No. 3 and Tab. . No. 11).

Roberts et al (1952) mentioned that infestation could occur at a very early age through the contamination of the teats of cows, or through the calves' habit of sucking various objects, but that infestation in this manner does not appear to be of any significance. This view was not confirmed, but it must be mentioned that the average kraal in the North western Cape and the kraal created in these experiments had a layer of at least six inches of manure as a floor. In the rainy season particularly, wet and semi dry manure adhered very easily to the udder and the ventral surface of the abdomen. In addition calves had the habit of

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\*Fig. 1 is Figure.

\*Fig. 1 is Figure.

licking themselves and each other, particularly about the navel. These unhygienic conditions probably did not apply to the farms which Roberts and his co-workers investigated and possibly this was not an important source of infestation there.

When all sources of infestation were combined and calves grazed in the infested paddock and suckled in the infested kraal, it was not surprising therefore that the experiment (D) indicated mean egg counts (Fig. No. 9), and worm burdens of all species at post mortem (Group 5 Tab. No. 14), more than double any other group. This practice was the most popular method of rearing calves in the district.

The provision of clean concrete floors in calf pens made a big difference in the incidence of verminosis in hand reared calves. Only two calves out of five become mildly infested in one experiment (Group C. Expt. B.) and none in other experiment (Group 6 Expt. D.). Although suckling calves on a clean concrete floor became infested in two experiments (Group B. Expt. B. Group 2 Expt. C) this was due to their dams lying in kraal manure during a particularly wet autumn and transporting the infestation to them via the teats. In a dry summer and autumn the same group did not become infested (Group 2 Expt. D.) nor did calves suckling on previously washed udders become infested when kept on clean concrete floors. Calf pen hygiene was responsible for their freedom from parasitism.

(b) SEASONAL INCIDENCE.

These experiments indicated that parasitism in calves was seasonal. During the summer and autumn depending on the rain and husbandry methods, stock became infested to a greater or lesser degree. During the dry months, i.e. from April to November, with one exception (Calf No. 8246, Group 1 Expt. C.), stock did not become infested. Calves

born between April and November (Expt. C and D), only became infested in December.

Since rainfall, played an important role in rendering infective larvae available, in numbers large enough to infest the host, it was important to know the minimum amount of rainfall necessary, to create these conditions.

In Table No.17 three minimum prepatent periods are given for H. placei larvae to reach maturity in the host, and using this information a rainfall range of 13.3 to 72.8 mm over periods of 10 to 22 days was shown to be the amount of rainfall necessary. However these larvae were recovered from kraal manure (Table No. III Appendix) and grass before this amount of rain had fallen at the end of November. Roberts (1957) although giving a minimum prepatent period of 19 days, reports an average period of 26-28 days for H. placei larvae to reach maturity in the host. Similarly Mayhew's (1941) results showed that in eight out of thirteen cases tested prepatent periods varied from 23 to 26 days. Both these workers showed an average of 26 days as the prepatent period. Since in my observations larvae of H. placei were diagnosed in faeces collected on 28/12/56 for the first time, it is reasonable to assume that infestation took place 26 days prior to this, i.e. on the same day Cooperia pectinata larvae infested calves. If that were the case only 13.3 mm of rain distributed over <sup>2</sup>10 days was necessary to render enough larvae available to infest the host.

In Table No. 16 two minimum prepatent periods of 52 and 56 days are shown for B. phlebotomus according to Mayhew's (1948) and Sprent's (1946a) observations respectively.

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<sup>2</sup>The first rain was recorded on 22/11/56 giving a 10 day period until 2/12/56.

In Table No. 17 however only the period given by Mayhew (1948) was used, and it was shown that 42.3 mm of rain was necessary to render enough infective larvae available to infest calves. This was done in view of the fact that observations reported elsewhere, as well as observations by other workers (Schwartz 1924 and Sprent 1946b) showed that the larvae of this species were very sensitive to drying and desiccation, and therefore unlikely to be available in large enough numbers to infest the host when only 13.3 mm of rain had fallen, which would be the case if the prepatent period of 56 days was correct.

Nocturnal dews in April and May 1956 were extremely heavy and no rain fell in April and only 9.4 mm in May. Five calves born in this period failed to become infested (Calf No. 8240 Group 1 and Calves No's 8238, 8239, 8241 and 8242 Group 5 Expt. C). Experiments reported elsewhere showed no larval migration from dung under the influence of dew so that the observation that calves failed to become infested was understandable. Riek et al (1953) and Roberts (1957) however, stated that heavy dew would assist in migration of larvae from dung pads. Previous observations (vide supra) and failure of calves to become infested in April did not confirm their observations.

(c) FLUCTUATIONS IN WORM EGG COUNTS.

Worm egg counts in faecal samples rose to a peak from March to May in calves born from November to January (Fig. No.4 & Fig. No's 8 & 9). Sudden rises in egg counts in summer and autumn were noted in certain groups of calves three to five weeks after well distributed, heavy rains fell (Group B & D Fig. 4). A spectacular rise from a negative examination the previous week to 1,460 e.p.g. was noted in one calf (No. 7180) on 7/1/56, five weeks after the rains had started in the previous month (Fig. No.3).

Egg counts in calves born in February and March showed two peaks, the most prominent one being in June and July, followed by a secondary peak in October and November (Fig. No.6) Calves born after the rains had ceased at the end of March, i.e. April to September calves, and kept under similar conditions<sup>S</sup> did not show infestation on faecal examination until <sup>M</sup>December and January (Group 3 Fig. No.6, Group 5 Fig. No.7). The calves born in February and March therefore acquired their infestation before the rains ceased at the end of March; the variations in the worm egg counts were due to that infestation and not to infestation acquired during the winter and early spring.

Post mortem results in the winter, spring and summer showed that worm egg counts were not a reliable index of the worm burdens in calves.

Roberts (1957) has stated: "egg counts may be regarded as a reasonably accurate index of the population during the period of susceptibility to infestation, but they are of doubtful value when the animal has acquired, or is acquiring, resistance. In other words, it would seem that whereas high egg counts are a safe index of heavy infestations, low egg counts do not necessarily imply that only a few worms are present."

The observations in these experiments are in complete agreement with Roberts' statement. Most calves were slaughtered when they were over seven months of age and had been infested with worms for periods of four months or longer. Some resistance to infestation must have been acquired by the host since post mortem results were very variable when compared with worm egg counts.

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<sup>M</sup>There was one exception viz. Calf No. 8246 (Group 1 Expt. C) which was born in July and became infested at the end of November.

(d) IMMATURE H. PLACEI WORMS AT POST MORTEM.

The post mortem results of Experiment C revealed the presence of mature and immature H. placei worms in two autopsies done in October and November (Table No. 13).

The nearest description of these immature H. placei the author found in the literature, was that by Roberts (1957) in his paper on reactions of calves to H. placei as "small adult worms with few or no eggs in the females." The females in my experiments had no eggs and both males and females were small. Roberts furthermore showed that immature worms, in the fourth and fifth stage particularly could live in the host for long periods. This has been observed with other species<sup>of</sup> nematodes by various workers. (Taylor and Michel 1952 and 1953; Gibson 1953; Soulsky 1957). On the other hand, other workers stated that the presence of immature worms at post mortem was due to continual infection (Mönnig 1931; Morgan, Arnell & Ryski 1951).

In the description of these post mortem results, under the experimental observations of Experiment C, it was shown that the presence of these immature worms must have been due to infestation acquired in March i.e. 7 months previously. This conclusion was drawn from the fact that a winter born calf from the same group was negative at autopsy. Furthermore seven other calves born between April and September, grazing in the same paddock, remained negative on faecal examination for this species at that time and for a further six to eight weeks.

SUMMARY.

Four experiments are described on the transmission of Cooperia spp., H. placei, O. radiatus and B. phlebotomus to susceptible calves reared by different

methods of animal husbandry. The following conclusions are drawn:-

(1) Calves become infested in the summer and autumn only, and fail to become infested in the winter and spring.

(2) A minimum of 13-40 m.m. of rain over a period of 10-15 days, is necessary for infective larvae to become available in numbers large enough to infest the host.

(3) It is shown that the sources of infestation are:-

(a) Infested herbage.

(b) Infested kraal manure adhering to cows teats.

(c) Infested kraal manure by direct contact.

(4) Different systems of calf rearing facilitate the spread of verminosis, not only when rainfall increases the availability of infective larvae but even when little rain falls in summer and autumn. In order of their importance, the transmission of verminosis is facilitated by the following systems of calf rearing:-

(a) Calves suckling in the infested kraal and grazing in the infested paddock.

(b) Suckling calves grazing in the infested paddock.

(c) Suckling calves confined to the infested kraal and adjacent infested calf pen.

(d) Hand reared calves confined to the infested kraal and adjacent infested calf pen.

(5) Under the influence of heavy autumn rains calves reared on clean concrete floors also became infested. The manner in which they became infested is:-

(a) Suckling on teats of cows that became contaminated in the kraal.

(b) Accidental infestation of lucerne-hay fed to hand reared calves.



(6) During a comparatively dry summer and autumn calves confined to clean concrete floored pens failed to become infested when reared as follows:-

(a) By hand.

(b) Suckling on previously washed or unwashed udders.

(7) Post mortem results confirmed the results already mentioned under the transmission of verminosis. Although four species of worms were recovered at post mortem in all the groups, the order of numerical predominance of the species was as follows:-

(a) Cooperia pectinata and Haemonchus placei

Where the source of infestation was the herbage these species were recovered in large numbers.

(b) Dunostomum phlebotomum.

This species was predominant where kraal manure was the source of infestation.

(c) Oesophagostomum radiatum.

This species was recovered in small numbers both where the herbage and kraal manure was the source of infestation.

RR/EDL/5/6/58.

## PROPHYLAXIS

In his paper on the epidemiology of parasitic nematodes of sheep, Gordon (1948) divided the life cycle of parasitic nematodes into two stages and two phases. "The stages are: Parasitic - the worms in the host animal and Free-living - eggs and larvae in faeces on the ground. The two phases are: Contamination - eggs passing from the host to the pasture and Infection - larvae passing from the pasture to the host".

The emphasis of the experiments described above has been on the free-living stage and the contamination and infection phases. Control measures described below are therefore mainly a description of improved methods of animal grazing management based on experimental observations, and to a lesser extent the rôle anthelmintics might play in controlling the parasitic stage of the life cycle.

Prophylaxis can therefore be viewed from two angles which will be discussed separately. They are:-

1. Animal and grazing management.
2. The use of anthelmintics.

### 1. Animal and Grazing Management.

Since two systems of farming i.e. dairy and beef ranching are practiced in this area they will be discussed separately.

#### (a) Dairy Ranching.

As mentioned earlier calves on dairy ranches are separated from their dams at birth and placed in calf pens. When the cows come into the kraal in the morning from 9 a.m. onwards, two teats are milked and the calves allowed to suckle on the other two. After the cow returns to the grazing between 3 and 4 p.m. calves are herded into a small camp or if very young left in the kraal.

These calves are invariably in extremely poor condition, due not only to verminosis and various gastro-intestinal disorders, but also due to the fact that they are starved because the dams milk supply is inadequate and the grazing in the calf camp usually very inferior. On many farms the calf camp is grazed by sheep during the day, with the result that very little good grazing is available for calves.

It has been shown that calves kept under unhygienic conditions in kraals and grazing on infested pastures become infested with heavier worm burdens than calves reared by other methods. The provision of a concrete floor in the calf pen was sufficient to reduce the worm burden to low levels and even to the state where calves were free from parasites. It is not only necessary to improve the hygiene in the calf pen and kraal, but also to provide good quality uninfested grazing for calves; attempts should be made to improve the nutritional state of these calves by allowing them adequate milk, higher quality grazing in the form of improved pastures, or at least enough good quality veld grazing, by providing more than one calf camp. According to Dr. C.E.M. Tidmarsh of the Division of Crops and Pastures (personal communication), veld in the North-Western Cape should be allowed a rest in late summer followed by a rest in early summer, once every four or five years. The important period to rest the pastures are during the growing season and Tidmarsh suggests the following rotation on a four camp system, shown in the schedule below:-

Year	Season	Camps			
		A	B	C	D
1	E.S.	G	G	G	R
	L.S.	G	G	R	G
	W.	G	G	G	G
2	E.S.	G	G	R	G
	L.S.	G	R	G	G
	W.	G	G	G	G
3	E.S.	G	R	G	G
	L.S.	R	G	G	G
	W.	G	G	G	G
4	E.S.	R	G	G	G
	L.S.	G	G	G	R
	W.	G	G	G	G

**NOTE:** Tidmarsh divides the year into three seasons:-  
 E.S. is early summer i.e. from mid-September to mid-December. L.S. is late summer i.e. from mid-December to the end of April. W. is winter i.e. from May to mid-September. Furthermore, G stands for graze and R for rest.

As can be seen from the schedule, three camps are grazed in the summer and four in the winter. Each camp is grazed for two and rested for four weeks in the summer, whereas each camp is grazed for two and rested for six weeks in the winter. Should more than 40 m.m. of rain fall in any fortnight calves must be moved every week to another camp, and return to a fortnight rotation when drier conditions return. This would

assist in verminosis control without detriment to the veld.

The size of these camps would be determined by the carrying capacity of the veld, the number of calves on the farm, the available water supplies etc., and in each case the farmer would have to be guided by the advice of the local Agricultural Extension Officer.

On farms where dairy ranching is practiced, bulls run with cows constantly, in attempts to make the cows calve through the year. Bonsma (1939) has shown that the main calf crop is in mid-summer - from November to January - and to a lesser extent in mid-winter - June and July. It is suggested therefore that cows be served only in February, March and April for the summer calf crop and from the middle of August to the middle of October for the winter calf crop.

If these calves are allowed to graze in special camps by themselves it is highly desirable to wean the summer calf crop by the end of August and the winter calf crop by the end of February. It was shown in experiments on the epizootology of verminosis that calves carried worm burdens from autumn through to the following summer. In spite of these calves being reared together with winter and spring calves, the younger calves remained free of infestation until the rains started in December. It is therefore important that the calves born in the previous summer be weaned and separated from calves born during or after June, before the rains start in December. If calves born in summer are weaned at the end of August, and given good grazing in a separate camp, they will not carry infestation to calves born in the winter, because they will have left the calf camps in the dry season, at least two months before the rains start. Furthermore, it is desirable that winter born calves, when weaned at the end of February, do not graze before the following winter with calves weaned at the end of the previous August.

This therefore entails at least four calf camps and two weaner camps. In brief, summer calves may become infested in summer and autumn but will not infest winter calves if the latter are born after June and the summer calves are weaned in August. Calves born in winter will be free of infestation and can run with calves born in the following summer until the former are weaned in February. If both August and February weaners are kept separate until the rains cease in autumn, they can join the main herd the following summer. By this time

their ages will vary from 18 to 24 months and the older heifers can join the breeding herd and both groups will be fairly worm free; if not resistant to verminosis, at least the effect of verminosis on them will not be so marked if they become infested thereafter.

Riek et al (1953) have suggested that the low incidence of verminosis in calves at Maleny, Queensland, Australia, where annual rainfalls are high (average 68 inches) was due to improved ley pastures, rotated every few months and other nutritional considerations, as well as the provision of concrete floored calf pens and cow yards.

The dairy herd at Armeedsvlakte was similarly lightly infested. Calves were born in summer only, kept in three camps that were rotated every six to eight weeks and after weaning in August kept in separate camps for another year. No sheep grazed in any of these camps. The calf pens, kraals and cow byres had concrete floors, and calves suckled on their dams after the udders had been washed. All these calves received more milk than was the practise in the district. Hygiene, improved nutrition and the separation of age groups all contributed to the low incidence of verminosis.

(b) Beef Ranching.

On farms where beef ranching is practiced cows calve in the veld and the calf is not separated from its dam until weaning time. The incidence of verminosis on these farms is of a much lower degree, than on the dairy ranch and calves are more healthy and better<sup>fed</sup> than their counterparts on the dairy ranch.

It would be advisable on these ranches to keep the breeding herd in separate camps from the rest of the herd, Furthermore, since milk is not the farmers source of income, there is no object in having more than one calving season. for more efficient management of the herd, unless fertility of the cows is at a low level. Since most calves are born in the summer, cows should be served from February to April so that calving can occur in November, December and January. This is unfortunate from the verminosis control point of view, and winter calving would be preferable, but calf crops in winter are seldom as plentiful as summer calf crops; management is also simplified with one calf crop.

All calves should be weaned in August and placed in their own camps. Since calves will normally lose condition at this time of the year after weaning, the best available grazing should be set aside for the weaned calves, and if at all possible weaners should be separated from the rest of the herd until they have reached the age of two years when the heifers join the breeding herd and the oxen join the older stock for fattening prior to slaughter. From the management point of view it would be desirable to separate heifers and oxen at weaning if separate camps are available for both groups.

It can be assumed that calves born in summer will become infested with parasites before the rains cease in autumn. Apart from the use of anthelmintics in worm control, which will be dealt with later, two additional methods are suggested for consideration. These methods are:-

(1) Grazing Rotation.

Grazing rotation as advocated for dairy calves would be highly desirable for the breeding herd on beef ranches. However, this may entail considerable expense both in fencing material and the provision of water supplies, particularly if the breeding herd is large. Where a four camp system is impracticable, Tidmarsh (personal communication) suggests the following rotation on a three camp system shown in the schedule below:-

		Camps		
Year	Season	A	B	C
1	E.S.	G	G	R
	L.S.	G	R	G
	W	G	G	G
2	E.S.	G	R	R
	L.S.	R	G	G
	W	G	G	G
3	E.S.	R	G	G
	L.S.	G	G	R
	W	G	G	G

**NOTE:** E.S. is early summer i.e. from mid-September to mid-December; L.S. is late summer i.e. from mid-December to end of April; W. is winter i.e. from May to mid-September. G. stands for graze and R for rest.

In the summer camps are grazed for three weeks and rested for three weeks; in winter camps are grazed for three and rested for six weeks. Verminosis control under these conditions is almost impossible and much less satisfactory than the four camp system mentioned earlier.



Due to the general lack of water in the North-Western Cape it is difficult to supply water to separate camps. This can be overcome by extending existing water supplies, and providing separate troughs for each camp. Division into camps would therefore entail fences meeting at or near a central point and water troughs being placed in each camp; the fence between the camps to divide the trough as well, so that stock have water in each camp. On many farms water troughs are divided in the manner suggested. By various methods existing reservoir tanks can supply more than one water trough, or the water can be pumped from tanks, to other reservoirs, to extend the distribution of existing water supplies. Provision is made under the Soil Conservation Act, for the subsidising of fencing material and piping for the extension of existing water supplies from boreholes to additional watering points.

(2) Removal or Disinfestation of Kraal Manure.

(a) Removal of manure:

Conditions similar to those existing in kraals are common around every drinking trough on farms owned by European farmers in the district. Many years of accumulated manure are found around watering points on almost every farm.

On the trust farms of the Native Affairs Department, no such manure accumulations exist, because the Native stock owners collect dung pads while still fresh and place them in heaps next to their huts for use as fuel when dry or even for building purposes. It was noted that Native owned stock were practically worm free, although calves were allowed to run with their dams in a similar fashion to European owned stock. In view of the investigations previously described the lack of manure at watering points must have been largely responsible for their freedom from parasitism.

Since cattle tend to congregate near the water troughs during the heat of the day and deposit a high concentration of manure there, it would be advisable to remove manure regularly, particularly in the rainy season, from the vicinity of water troughs. This is not practical where the soil is very sandy, since the manure becomes well mixed with the sand and it is impossible to separate manure from sand.



**(b) Disinfestation.**

The disinfestation of kraals was attempted in a kraal on a farm<sup>x</sup> in the district, using delta isomere of benzine-hexachloride. There were two adjacent kraals, used by two different herds, and kraal manure examination indicated that both kraals were infested. The one kraal was left as a control and the other kraal dused with 25 lbs of a 5% delta B.H.C. powder. The area of the treated kraal was approximately 20,000 sq.ft. and the treatment was carried out in March, 1956, when rainfalls were particularly heavy i.e. 120.0 mm for the month. Subsequent examinations showed that infective larvae were absent from the treated kraal for a period of two weeks only, whereas the control kraal remained infested throughout. This insecticide was therefore of temporary value only, in decreasing the number of infective larvae in kraals.

Laboratory trials of the effects of various chemicals as larvacides were carried out and the results of some of these trials are shown in Table No. 18.

Table No. 18.

The effect of various compounds at concentrations of one part per million parts in aqueous solution on infective larvae.

Compound	Percentage mortality of infective larvae after:-	
	(a) 24 hours	(b) 48 hours
x Malathion 25%	70%	80%
∂ Chlorthion 20%	40%	90%
Phenothiazine	35%	40%
gamma Benzine-hexachloride	30%	40%
delta Benzine-hexachloride	5%	20%
■ Polybor	8%	10%

x Malathion is O,O-dimethyl-dithiophosphate of diethyl-mercapto-succinate.

∂ Chlorthion is O,O-dimethyl-O-3 chloro-4-nitrophenyl thiophosphate.

■ Polybor's active ingredients are Sodium Pentaborate tetrahydrate 77% and Sodium Tetraborate pentahydrate 18%

Trials conducted by Levine et al (1956) showed that malathion and chlorthion killed or prevented the development of horse strongyle larvae at concentrations of 0.02 and 0.06 per cent respectively. The results summarised in Table No. 18

∞ Buckshee - owner Mr. Rex Butler

showed that these compounds were also very effective larvacides against infective bovine nematode larvae. The results with polybor were unsatisfactory (Table No. 18) although Hoerlein (1950 and 1951) showed that this compound destroyed Ancylostoma caninum larvae.

The effect of malathion on the development of larvae in cattle dung was tested by dosing three infested year old cattle with this drug much on the same principle as low level phenothiazine dosing. The toxicity of this drug was unknown and it was decided to incorporate toxicity trials with tests of the effects on the worms. The three animals were dosed with 5, 10 and 20 mgm. per Kg. body weight of malathion respectively every day for three months. Although this drug had no apparent toxic effect on the three animals tested, it was also of no value as a vermicide, and apparently not enough malathion passed through the alimentary canal with the faeces to act as a larvacide. When mixed with infested faeces in cultures, both malathion and chlorthion at one part per 10,000 parts of faecal culture medium, caused a 50 per cent reduction in the development of larvae to the infective stage when compared with controls.

These drugs may be of value in disinfecting kraal manure but were unfortunately not tested in the field. Malathion appears to be non-toxic for stock even at dosage rates of 20 mgm. per Kg. body weight for a period of three months. Before its use is attempted as a larvacide in kraals, its toxicity should be tested on new born calves, which are usually more susceptible to the effects of any poison than older stock.

At the moment, all that can be said is that the use of certain insecticides as larvacides in kraal manure needs further investigations before recommendations can be made.

Apart from these methods of decreasing the danger of manure as a potential source of infective larvae to susceptible calves, concrete floors can be laid around water troughs and regularly cleaned. It is doubtful whether any farmer will be able to carry this out with the scarcity of labour, or whether it is practical to surround every trough in the various camps with large areas of concrete, which must be cleaned regularly.

It is as well to mention that all water troughs should be well constructed and leak proof and at least a small area of concrete flooring be built next to the trough in case of overflow of water. It was frequently noted that due to leaking drinking troughs or the overflow of water due to faulty ball valves, the area next to the trough was a mixture of soft mud and manure. Such artificially created conditions obviously supplied more than enough moisture for larval development, and were dangerous, potential sources of infestation.

The remarks on leaking drinking troughs also apply to dairy ranches.

## 2. The use of anthelmintics.

The author having tested only two drugs viz. phenothiazine and tetrachlorethylene for their efficiency as anthelmintics, the recommendations made below are based on the observations of other workers as well. Recent investigations show that the following drugs are of value:

(a) Toluene: According to Riek and Keith (1957a) this drug is an effective anthelmintic against H. placei, B. phlebotomus and Cooperia spp. Cattle must be starved overnight and immediately prior to treatment, 60 c.c. of a 10% Sodium bicarbonate solution given to close the oesophageal groove (Riek 1954). The drug is then dosed at the rate of 10 c.c. per 100 lb body weight, in an emulsion prepared by the addition of an emulsifying agent. Apart from temporary anaesthetic effects this drug is relatively non-toxic according to these authors.

(b) 1:8 dihydroxyanthraquinone: Riek and Keith (1957b) have shown that this drug was highly efficient against H. placei, Cooperia spp. and O. radiatum administered per os as a suspension in water at 2.5 grams per 100 lb body weight. No premedication to close the oesophageal groove, nor starvation prior to dosing is necessary.

(c) Phenothiazine: This drug is effective against H. placei and O. radiatum according to various workers (Sprent 1946c; Riek 1951, Roberts 1955). These results were confirmed at Armoedsvlakte. The usual dosage rate is 0.2 grams per lb body weight according to Riek (1951) and Roberts (1955). The usual maximum dose is 40 grams for an adult.

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X This drug is sold under the proprietary names "Diaquone" "Altan" "Istin" and "Istisin".

(d) Tetrachlorethylene: This was mixed with an emulsifying agent and one part of  $C_2Cl_4$  was well shaken with two parts of water to make an emulsion. Immediately after dosing with 60 c.c. of 10% sodium bicarbonate solution, 25 c.c. of this emulsion per 100 lb body weight were dosed to a maximum of 100 c.c. i.e. 33.3 c.c. of  $C_2Cl_4$ . Results were not very satisfactory although the drug was effective against H. placei. When larger doses of 60 c.c. of emulsion were used i.e. 20 c.c.  $C_2Cl_4$  per 100 lb body weight, after premedication with 10% sodium bicarbonate solution, this drug was highly efficient against H. placei, but not very effective against other worms.

The most efficient drugs against the parasitic nematodes in cattle would appear to be toluene and 1:8 dihydroxy-anthroquinone although these were not personally tested. The price of the latter unfortunately is too high for it to be used where large herds have to be treated. The disadvantage of toluene is that animals must be starved overnight, and that this drug must pass directly into the abomasum to be efficient. It is effective against H. placei, Cooperia spp., and B. phlebotomum, however, according to Riek and Keith (1957a) and since it is cheap its use on a large scale is recommended. Phenothiazine appears to be the drug of choice for the treatment of O. radiatum and H. placei.

Strategic drenchings: It has been shown that stock only become infested in summer and autumn, and that different species of worms are present in stock in significant numbers at different times. Each species will be dealt with separately.

Cooperia spp.: Egg counts reach a maximum on faecal examination in autumn and post mortems of calves in the winter showed large recoveries of C. pectinata.

H. placei: There were two peaks in egg counts in this species, the most marked being in autumn and a secondary rise in spring. More worms of this species were recovered at post mortem in the winter and spring than in the summer.

O. radiatum: Egg counts only reach their maximum in winter and at post mortem this species was more plentiful in the winter than at any other time.

B. phlebotomum: Egg counts in this species were at their maximum in winter, whereafter they fell to show a secondary rise in the spring. More worms were recovered in the winter and spring than in mid-summer at post mortem.

It is therefore suggested that strategic drenching of stock be carried out on all stock apart from calves younger than two months of age or winter born calves. The best time to dose, and the drugs of preference would then be as follows:-

1. February: This drenching is aimed at Copperia spp. and H. placei, before the rise in egg output in autumn. Toluene would be the drug of preference.
2. May: This treatment is aimed at Cooperia spp, H. placei and B. phlebotomus which will be present in fair numbers by this time. Toluene would be the drug of choice.
3. July: This treatment is aimed at O. radiatum and phenothiazine would be preferable.
4. August: This treatment is aimed at the spring rise in eggs output of B. phlebotomus and H. placei and toluene would be the drug of choice.

To summarise, toluene would be dosed in February, May and August and phenothiazine in July.

Calves born in the winter need not be dosed until February and since they will be weaned in February, they must be dosed before going to the weaner camps. There is no object in dosing summer born calves before they are two months of age since their worm burdens are very low before this age.

Ranch cows due to calve in summer should be dosed in October. Since they seldom have more than a mild infestation of H. placei and very few other parasites phenothiazine should be used. In these animals it may be advisable to incorporate low level phenothiazine treatment from November to May by adding 10 per cent of this drug to the lick. The expense of low level phenothiazine dosing would hardly be warranted with other stock.

In the event of heavy rains occurring at any time, tactical drenching with toluene four to six weeks after the rains commenced would be advisable.

#### SUMMARY.

Possible methods of prophylaxis based on experimental observations are described and the possible use of strategic drenching using anthelmintics which gave promising results when tested by other workers is included.

The recommendations made are as follows:-

1. Separation of calves born before the winter, from younger calves born before the following summer.
2. The provision of concrete floored calf pens and kraals on dairy ranches as well as four separate grazing paddocks for calves; the paddocks to be regularly rotated.
3. The separation of breeding stock from the rest of the herd on beef ranches, and the provision of camps for these animals which are regularly rotated.
4. The provision of separate camps for weaners.
5. Strategic dosing of stock with anthelmintics using toluene in February, May and August, and phenothiazine in July.
6. Wherever possible, improving the nutritional state of calves and weaners.



A D D E N D U M.

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After completion of this paper the author obtained the following reference - RIEK R.F. and KEITH R.K. (1958). "Studies on anthelmintics for cattle : IV. The organic phosphorous compound O,O-dimethyl-2,2,2,-trichloro-1-hydroxyethyl phosphonate (Bayer L.13/59)". Aust. Vet. Jl. Vol. 34, No. 4 pp. 93-103.

This drug is known by the trade name of "Mevuvon". Riek and Keith observed no significant difference in the anthelmintic efficiency of the aqueous solution or as a 50% emulsion when tested at the same dosage rates.

They found the drug highly effective against H. placei and O. radiatum at dosage rates of 2 grams per 100 lb. body weight; at dosage rates of 5 grams per 100 lb. body weight it was effective against Cooperia spp., B. phlebotomum and even T. axei.

An extremely important observation was that this drug proved effective against the immature stages of most of the common nematodes of cattle.

Although relatively non toxic at the dosage rates recommended, calves on high protein diets showed severe symptoms, and these authors advise the use of this drug under veterinary supervision until more experience has been gained with its use.

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**A C K N O W L E D G E M E N T S.**

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R E F E R E N C E S.

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- BAILEY, W.S. (1949) "Studies on calves experimentally infected with Cooperia punctata (v. Linstow 1907) Hanson 1907" Amer. Jl. Vet. Res. Vol. 10, No. 35, pp.119-129.
- BONSMAN, J.C. (1939) "Breeding Seasons on Cattle Ranches" Farming in S. A. Vol. 14, No. 159, pp. 230 and 241.
- CROFTON, H.D. (1948a) "The ecology of immature phases of Trichostrongyle nematodes. (1) The vertical distribution of infective larvae of Trichostrongylus retortaeformis in relation to their habitat. Parasit. Vol. 39, No's 1/2, pp. 26-38.
- CROFTON, H.D. (1948b) "The ecology of immature phases of Trichostrongyle nematodes II. The effect of climatic factors on the availability of the infective larvae of Trichostrongylus retortaeformis to the host". Parasit. Vol. 39, No's 1/2, pp. 26-38.
- CROFTON, H.D. (1949) "The ecology of immature phases of Trichostrongyle nematodes III Larval populations on hill pastures". Parasit. Vol. 39, No's. 3/4, pp. 274-280.
- DE BLIECK L et Baudet E.A.R.F. (1926). "Contribution a l'étude du développement des Strongylidés (Schlerostomes) du gros entestine chez le cheval". Ann. de. Par. Hum. et Comp. Vol. 4, No. 1, pp. 87-96.
- DINABURG, A.G. (1944) "Development and survival under outdoor conditions of eggs and larvae of the common ruminant stomach worm, Haemonchus contortus". Jl. Agric. Res., Vol. 69, No. 11, pp. 421-433.
- DINNIK, J.A. & DINNIK, NN. (1954/55). "The development and survival of Haemonchus contortus larvae on pasture under the local condition of the high-lands of Kenya". East. Afr. Vet. Res. Ory. Ann. Rep 1954-1955 in corpor. Ann. Repts 1952 and 1953. pp. 76-84.
- DU TOIT, S.J. (1958) "Dairy ranching in the Malepo". Farming in S. Af., Vol. 33, No. 11, pp. 45-48.

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- FOURIE, J.M. (1942) "A note on acute verminosis in cattle in a semi-arid area". Jl. S. Af. Vet. Med. Ass., Vol. 13, No. 3, pp. 70-72.
- GIBSON, T.E. (1953) "The effect of repeated anthelmintic treatment with phenothiazine on the faecal egg counts of housed horses, with some observations on the life cycle of Trichostrongylus spp. in the horse". Jl. Helminth. Vol. 27, No's. 1/2, pp. 29-40.
- GORDON, H. Mel (1948) "The epidemiology of parasitic diseases, with special reference to studies of nematode parasites in sheep". Aust. Vet. Jl. Vol. 24, No. 2, pp. 17-44.
- HOERLEIN B.F. (1950) "The evaluation of various chemical agents in the treatment of soil infected with larvae of the dog hookworm Anoxylostoma caninum". N. Amer. Vet. Vol 31 pp. 253-262.
- HOERLEIN B.F. (1951) "Further studies on the chemical treatment of soil infected with larvae of the dog hookworm (Anoxylostoma caninum)". Vet. Med. Vol. 46 No. 9 pp. 350-355.
- KAUZAL, G.F. (1941) "Examination of grass and soil to determine the population of infective larval nematodes on pastures". Aust. Vet. Jl., Vol 17, No. 10, pp. 181-184.
- KEITH, R.K. (1953) "Differentiation of infective larvae of cattle nematodes". Aust. Jl. Zool., Vol. 1, No. 2, pp. 223-235.
- LEVINE N.D., IVENS V., KLECKNER W.C., BENDER J.K. (1956). "Nematocidal screening tests of organic phosphorus, nitrofurans, cadmium, and other compounds against horse strongyle larvae in vitro". Am. Jl. Vet. Res. Vol. 17 No. 62 pp. 119-120.
- LOOS, A. (1911) "The anatomy and life history of Archylostomum quadrinale Dub". A monograph Pt. II. Rep. Egypt. Gov. Sch. of Med., Vol. 4, pp. 159-313. Cairo.
- MAYHEW, R.L. (1941) "Studies on bovine gastrointestinal parasites V. Immunity to the stomach worm with a note on the prepatent period". Amer. Jl. Hyg., Vol. 33, No. 3, pp. 103-111.
- MAYHEW, R.L. (1948) "The parasites and parasitic diseases of cattle". Lesions State Agric. Exp. Sta. Bull., No. 423, 48 pages.
- MÖNNIG, H.O. (1926) "The life histories of Trichostrongylus inaequalis and T. rugatus of sheep in South Africa". Pt. I 11th and 12th Rep. Dir. Vet. Ed. & Res. Un. S. Afr., pp. 229-251.
- MÖNNIG, H.O. (1930) "Studies on the bionomics of the free living stages of Trichostrongylus spp., and other parasitic nematodes", 16th Rep. Dir. Vet. Serv. Anim. Ind. Un. S. Afr., pp. 175-193.

- MÖNNIG, H.O. (1931) "The development of nematode eggs and larvae in cattle dung - preliminary note". 17th Ann. Dir. Vet. Serv. An. Ind. Un. S. Afr. pp. 207-208.
- MORGAN, D.O.; PARWELL, I.V. & RAYSKI, C. (1951). "The seasonal variations in the worm burden of Scottish Hill Sheep". Jl. Helminth., Vol. 25, Nos. 3/4, pp. 177-219.
- ORTLEPP, R.J. (1925) "Observations on the life history of Tridontophorus tenuicollis, a nematode parasite of the horse". Jl. Helminth. Vol. 3, No. 1, pp. 1-14.
- ORTLEPP, R.J. (1937) "Observations on the morphology and life history, of Geigeria pachycoelis Raill & Henry 1910 : A hookworm parasite of sheep and goats". Ann. Jl. Vet. Sc. Ania. Ind. Vol. 9, No. 1, pp. 183-219.
- RANSON, H.M. (1906) "The life history of the twisted wire worm (Hagmonchus contortus) of sheep and other ruminants". Un. St. Paul. Agric. Exp. An. Ind. Circ. No. 93, 7 pages.
- RIEK, H.F.; ROBERTS, F.H.S., & O'SULLIVAN, P.J. (1953). "Further observations on the epidemiology of parasitic gastroenteritis of cattle". Aust. Vet. Jl., Vol. 29, No. 8, pp. 122-123.
- RIEK, H.F. (1954). "The influence of sodium salts on the closure of the oesophageal groove in calves". Aust. Vet. Jl., Vol. 30, No. 2, pp. 29-37.
- RIEK, H.F., & KEITH, P.K. (1957a) "Studies on anthelmintics for cattle". I The efficiency of toluene with special reference to the hookworm, Banostomum phlebotomus". Aust. Vet. Jl. Vol. 33, No. 7, pp. 162-165.
- RIEK, H.F., & KEITH, P.K. (1957b) "Studies on anthelmintics for cattle. II The efficiency of 1:8 Dihydroxyanthraquinone". Aust. Vet. Jl., Vol. 33, No. 7, pp. 169-173.
- ROBERTS, F.H.S., & O'SULLIVAN, P.J. (1950). "Methods for egg counts and larval cultures for strongyles infecting the gastrointestinal tract of cattle". Aust. Jl. Agric. Res., Vol. 1, No. 1, pp. 99-
- ROBERTS, F.H.S. (1951) "Parasitic gastroenteritis of cattle with particular reference to the occurrence of the disease in Queensland". Aust. Vet. Jl. Vol. 27, No. 10, pp. 274-282.
- ROBERTS, F.H.S.; O'SULLIVAN, P.J., & RIEK, H.F. (1952). "The epidemiology of parasitic gastroenteritis of cattle". Aust. Jl. Agric. Res., Vol. 3, No. 2, pp. 137-233.
- ROBERTS, F.H.S.; TURNER, H.T. & McKEVITT, W. (1954) "On the specific distinctness of the ovine and bovine "Strains" of Hagmonchus contortus (Rad.) Cobb (Nematoda; Trichostrongylidae)". Aust. Jl. Zool. Vol. 2, pp. 275-295.

- ROBERTS, F.H.C. (1955) "Field trials on the evaluation of tetrachlorethylene as an anthelmintic for cattle". Aust. Vet. J., Vol. 31, No. 7, pp. 165-169.
- ROBERTS, F.H.C. (1957) "Reactions of calves to the stomach worm *Haemonchus placei* (Place 1893) Ransom 1811". Aust. J. Agric. Res., Vol. 8, No. 6, pp. 740-767.
- ROGERS, W.P. (1940) "The effects of environmental conditions on the accessibility of third stage *Trichostrongyle* larvae to grazing animals". Parasit. Vol. 32, No. 2, pp. 208-225.
- SCHWARTZ, B. (1924) "Preparacitic stages in the life history of the cattle hookworm (*Bunostomum phlebotomum*)". J. Agric. Res. Vol. 19, No. 9, pp. 451-458.
- SKEDDON, H.H. (1950) "Diseases of domestic animals in Australia Part I. Helminth infestations". Serv. Rep. (Div. Vet. Hyg.) No. 5 Sydney. A.N. Pettifer Govt. Printer.
- SOULSBY, E.J.L. (1957) "Some immunological phenomena in parasitic infections". Vet. Rec., Vol. 59, No. 49, pp. 1129-1130.
- SPRENT, J.F.A. (1946a) "Studies on the life history of *Bunostomum phlebotomum* (Fairbairn 1900) a hookworm parasite of cattle". Parasit., Vol. 37, No's 3/4, pp. 192-201.
- SPRENT, J.F.A. (1946b) "Some observations on the bionomics of *Bunostomum phlebotomum* a hookworm of cattle". Parasit., Vol. 37, No's. 3/4, pp. 202-210.
- SPRENT, J.F.A. (1946c) "Critical anthelmintic tests in cattle". Vet. J., Vol. 102, No. 4, pp. 83-87.
- TAYLOR, E.L. (1939) "Observations on the bionomics of strongyloid larvae on pastures". Vet. Rec., Vol. 50, No. 40, pp. 1265-1272.
- TAYLOR, E.L. (1939) "Technique for the estimation of pasture infestation by strongyloid larvae". Parasit., Vol. 31, No. 4, pp. 473-476.
- TAYLOR, E.L. & NIGHEL, J.F. (1952) "Inhibited development of *Distomatina* larvae in the lungs of cattle and sheep". Nature, Vol. 169, p. 753.
- TAYLOR, E.L. & NIGHEL, J.F. (1953) "The parasitological and pathological significance of arrested development in nematodes". J. Helminth. Vol. 27, Nos. 3/4, pp. 199-206.
- THEILER, SIR A., & ROBERTSON, W. (1915) "Investigations into the life history of the wireworm of ostriches". 3rd and 4th Rep. Dir. Vet. Res. Un. S. Afr., pp. 291-345.
- CEM.  
TIMMARCHI, (1953) Personal communication.
- VEGLIA, F. (1915) "The anatomy and life history of *Haemonchus contortus* (Rud)". 3rd and 4th Rep. Dir. Vet. Res. Un. S. Afr., pp. 347-500.

- VEGLIA, F. (1923) \*Preliminary notes on the life history of  
Oesophagostomum columbianum\* 8th & 10th  
Rep., Dir. Vet. Ed. & Res., Un. S. Afr.  
....., pp. 809-823.
- VEGLIA, F. (1926) \*Oesophagostomiasis in sheep (Preliminary  
Note)\* 13th & 14th Rep., Dir. Vet. Ed. Res.  
Un. S. Afr., pp. 763-767.

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**RESULTS OF EXPERIMENTS ON EGG HATCHING AND DEVELOPMENT OF LARVAE TO THE INFECTIVE STAGE UNDER FIELD CONDITIONS.**

EXPERIMENTAL NUMBER.	POSITION OF DUNG PADS IN VELD.	DATE EXPERIMENT COMMENCED.	NUMBER OF DAYS IN VELD.	SPECIMENS EXAMINED.	ORIGINAL WEIGHT OF DUNG IN GRAMS.	WEIGHT OF DUNG AT COLLECTION IN GRAMS.	EGGS PER GRAM.	Larvae recovered with the Baermann Apparatus.			THIRD STAGE LARVAE RECOVERED AFTER INCUBATION.	Cooperia spp.	H. placei.	O. radia tum.	B. phlebotomum.	REMARKS.
								1st Stage	2nd Stage	3rd Stage						
1	Exposed to sun.	15.8.56		F.			160	-	-	-	-	-	-	-		
(a)	do.	do.	7	X.	75	2	-	-	-	3,200	72.5%	8%	9.5%	10%		
(b)	do.	do.	8	E.	400	100	20	8	16	40	33	-	-	7	1(b) No cultures results from incubated dung.	
(c)	do.	do.	9	E.	do.	100	0	1	31	10	9	19	7	1		
(d)	do.	do.	35	E.	do.	130	0	-	-	2	-	-	-	-		
										87	87	-	-	-		
2	Exposed to sun.	16.8.56		F.			110	-	-	-	-	-	-	-		
(a)	do.	do.	4	X.	75	2	-	-	-	2,550	71.5%	3%	12.5%	13%		
(b)	do.	do.	6	E.	400	175	10	150	50	812	83%	2%	15%	-		
(c)	do.	do.	8	E.	do.	140	20	11	33	125	89%	4%	7%	-		
										16	13	-	1	2		
3	Exposed to sun.	18.8.56		F.			130	-	-	-	-	-	-	-		
(a)	do.	do.	2	E.	400	195	40	12	-	5,250	93%	2.5%	2.5%	2%	No control culture made.	
(b)	do.	do.	10	E.	do.	85	0	11	100	13	13	-	-	-		
(c)	do.	do.	60	E.	do.	65	0	-	-	4	4	-	-	-		
										8	8	-	-	-		
4	Exposed to sun.	13.9.56		F.			410	-	-	-	-	-	-	-		
(a)	do.	do.	6	X.	75	2	-	-	-	2,375	48%	5%	8%	39%		
(b)	do.	do.	7	E.	400	195	10	500	2,500	5,000	71.5%	6.5%	9%	13%		
(c)	do.	do.	9	E.	do.	155	0	40	200	250	437	86%	5%	1%	50%	
(d)	do.	do.	30	E.	do.	125	0	-	350	1,100	-	84%	1%	2%	13%	
(e)	do.	do.	60	E.	do.	95	0	-	-	455	1,600	88%	-	3%	9%	
										800	1,050	94.5%	2.5%	4%	-	
										-	2,270	69%	1%	30%	-	
										-	-	95%	2%	3%	-	
5	Exposed to sun.	22.9.56		F.			410	-	-	-	-	-	-	-		
(a)	do.	do.	2	X.	75	2	-	-	-	2,175	38%	34%	15%	13%	Incubated dung culture in 5(c) accidentally discarded before examination.	
(b)	do.	do.	4	E.	400	210	20	1	1	15,720	60%	9%	22%	9%		
(c)	do.	do.	6	E.	do.	190	60	900	37	11,575	90.5%	2%	7%	0.5%		
(d)	do.	do.	9	E.	do.	105	20	250	388	-	-	-	-	-		
										-	4	44	-	-		
										175	38	2	-	4		
										-	91%	-	7%	2%		

X F. Fresh faeces examined for eggs before dung pads were placed in the field.  
 X. Control cultures incubated in the laboratory.  
 E. Entire Dung pad.





EXPERIMENTAL NUMBER.	POSITION OF DUNG PADS IN VELD.	DATE EXPERIMENT COMMENCED.	NUMBER OF DAYS IN VELD.	SPECIMENS EXAMINED.	ORIGINAL WEIGHT OF DUNG IN GRAMS.	WEIGHT OF DUNG AT COLLECTION IN GRAMS.	EGGS PER GRAM.	Larvae recovered with the Baermann Apparatus.			THIRD STAGE LARVAE RECOVERED AFTER INCUBATION.	<i>Cooneria</i> spp.	<i>H. pla. cei.</i>	<i>O. radiatum.</i>	<i>B. phlebotomum.</i>	REMARKS.
								1st Stage	2nd Stage	3rd Stage						
13	Exposed to sun	22.1.57		F. X. 75			140	-	-	-	-	-	-	-		
(a)	do.	do.	8	E. 1000	650	0	1,575	3,150	2,500	7,500	39.5%	24.5%	20%	16%	Compare with No. 12 (See remarks No. 12)	
(b)	do.	do.	30	E. do.	215	0	-	-	575	8,125	61%	12%	21%	4%		
											28%	20%	12%	40%		
											59%	5%	36%	-		
											86%	4%	10%	-		
14	Exposed to sun	12.2.57		F. X. 75			140	-	-	-	-	-	-	-		
(a)	do.	do.	2	E. 400	200	90	250	-	-	1,500	56%	36%	3%	5%		
(b)	do.	do.	4	E. do.	150	0	63	-	-	1,850	49%	41.5%	6.5%	3%		
(c)	do.	do.	6	E. do.	85	0	-	-	12	100	85%	12%	3%	-		
(d)	do.	do.	8	E. do.	65	0	-	-	9	13	12%	1%	-	-		
(e)	do.	do.	30	E. do.	70	0	-	-	2	27	9%	1%	-	-		
(f)	do.	do.	60	E. do.	85	0	-	-	105	15	15%	-	-	-		
											100%	-	-	-		
											7%	7%	-	-		
15	Exposed to sun	12.3.57		F. X. 75			400	-	-	-	-	-	-	-		
(a)	do.	do.	1	E. 400	340	360	-	-	-	400	40%	48%	9%	3%		
(b)	do.	do.	2	E. do.	240	60	1,350	-	-	20,000	51%	47.5%	2%	0.5%		
(c)	do.	do.	3	E. do.	200	285	9,200	1,150	-	33,125	52.5%	43.5%	3.5%	0.5%		
(d)	do.	do.	4	E. do.	205	195	3,025	2,825	-	7,125	41.5%	54%	4.5%	-	Small number of larvae recovered in control culture	
(e)	do.	do.	5	E. do.	180	150	1,200	5,400	700	2,500	57%	31%	9%	3%		
(f)	do.	do.	6	E. do.	135	120	800	2,850	2,300	1,375	65%	33%	1%	1%		
(g)	do.	do.	7	E. do.	125	30	250	1,150	7,500	2,500	50%	30%	12%	-		
(h)	do.	do.	8	E. do.	215	0	150	200	8,750	2,500	71%	19%	7%	3%		
(i)	do.	do.	9	E. do.	185	0	-	36	6,875	14,765	86%	12%	2%	-		
(j)	do.	do.	30	E. do.	105	0	-	-	5,000	3,162	69%	18%	12%	1%		
											71%	20%	9%	-		
											75%	6%	18%	1%		
											78%	16%	4%	2%		
											41.5%	14%	43%	1.5%		
											74%	20%	5%	1%		
											85%	4%	11%	-		
											96%	1%	3%	-		
16	Exposed to sun	24.4.57		F. X. 75			390	-	-	-	-	-	-	-	All dung pads collected from 16(d) onwards where divided into the crust or upper layers and depth or lower layers as indicated by C. (crust) and D. (depth) in the fifth column of the table. The crust and depth were weighed and examined separately.	
(a)	do.	do.	1	E. 1400	385	260	200	-	-	28,750	43%	31%	19.5%	6.5%		
(b)	do.	do.	2	E. do.	325	240	750	-	-	50,000	49%	31%	15%	5%		
(c)	do.	do.	3	E. do.	280	250	1,250	-	-	50,000	34%	37.5%	19.5%	9%		
(d)	do.	do.	4	C. 90 D. 180	90 180	30 375	100 425	125	-	40,000	43%	40%	12%	5%		
(e)	do.	do.	5	C. 75 D. 130	75 130	0 285	5 3,750	1,005	-	7,500	42%	32.5%	20.5%	5%		
(f)	do.	do.	6	C. 45 D. 130	45 130	0 375	1 3,750	4,750	-	20,000	49%	23.5%	20.5%	7%		
(g)	do.	do.	7	C. 35 D. 105	35 105	0 270	1 1,400	-	-	12,400	6%	4%	4%	1%		
											14%	10%	55%	21%		
											17%	12%	16%	55%		
											6.5%	1.5%	43.5%	48.5%		
											-	-	4%	19%		
											2.5%	3%	45%	49%		

C. Crust of the Dung Pad.  
 D. Depth of the Dung Pad.





TABLE NO. I. (continued).

EXPERIMENTAL NUMBER.	POSITION OF DUNG PADS IN VELD.	DATE EXPERIMENT COMMENCED.	NUMBER OF DAYS IN VELD.	SPECIMENS EXAMINED.	ORIGINAL WEIGHT OF DUNG IN GRAMS.	WEIGHT OF DUNG AT COLLECTION IN GRAMS.	EGGS PER GRAM.	Larvae recovered with the Baermann Apparatus.			THIRD STAGE LARVAE RECOVERED AFTER INCUBATION.	Cooperia spp.	H. placei.	O. radiatum.	B. phlebotomum.	REMARKS.
								1st Stage	2nd Stage	3rd Stage						
18.	In shade	9.5.57		F.			200									
(a)	do.	do.	2	C.	400	125	315	-	-	-	20,000	44.5%	32.5%	13%	10%	
(b)	do.	do.	4	D.	do.	200	315	-	-	-	3,250	49%	36%	13%	2%	
(c)	do.	do.	8	C.	do.	95	225	1	-	-	21,750	43.5%	39%	13.5%	4%	
(d)	do.	do.	11	D.	do.	200	570	1	-	-	7,500	14.5%	59.5%	21%	5%	
(e)	do.	do.	14	C.	do.	75	30	50	-	-	20,000	32%	42%	25%	1%	
(f)	do.	do.	18	D.	do.	125	255	850	50	-	450	38%	48%	6%	8%	
(g)	do.	do.	26	C.	do.	125	30	5	20	-	3,100	33%	44%	21%	2%	
(h)	do.	do.	14	D.	do.	135	75	1,250	1,250	-	675	50%	46%	10%	4%	
(i)	do.	do.	14	C.	do.	85	0	77	41	7	750	32%	12%	48%	8%	
(j)	do.	do.	14	D.	do.	130	30	2,250	1,250	3	487	40%	54%	4%	2%	
(k)	do.	do.	18	C.	do.	60	0	-	-	-	3	2	1	-	-	
(l)	do.	do.	18	D.	do.	110	30	1,520	750	-	1,325	41%	50%	5%	4%	
(m)	do.	do.	26	C.	do.	55	0	-	-	1	3	2	1	-	-	
(n)	do.	do.	26	D.	do.	55	0	=	50	225	0	1	-	-	-	
(o)	do.	do.	26	C.	do.	55	0	=	50	225	375	76%	24%	-	-	
(p)	do.	do.	26	D.	do.	55	0	=	50	225	375	90%	10%	-	-	
19.	Exposed to sun	22.5.57		F.			305									
(a)	do.	do.	5	X.	75	65	90	500	-	-	21,750	34%	45.5%	17.5%	3%	
(b)	do.	do.	13	C.	400	130	855	1,825	-	-	3,675	34%	29%	20%	17%	
(c)	do.	do.	19	D.	do.	45	0	100	-	-	46,250	55.5%	27.5%	14.5%	2.5%	
(d)	do.	do.	19	C.	do.	45	0	-	-	-	8	7	1	-	-	
(e)	do.	do.	19	D.	do.	40	0	2	-	-	262	80%	20%	-	-	
(f)	do.	do.	19	C.	do.	40	0	-	-	-	18	16	2	-	-	
(g)	do.	do.	19	D.	do.	40	0	2	-	-	13	13	-	-	-	
(h)	do.	do.	19	C.	do.	40	0	-	-	-	13	13	-	-	-	
20.	In shade	2.5.57		F.			305									
(a)	do.	do.	20	X.	75	100	0	-	-	-	21,750	34%	45.5%	17.5%	3%	
(b)	do.	do.	20	C.	400	100	150	1,050	100	-	75	75	-	-	-	
(c)	do.	do.	20	D.	do.	100	150	1,050	100	-	975	71%	23%	4%	2%	

Compare with No.17:  
Ignore original  
egg per gram counts  
of fresh faeces  
which were obvious-  
ly incorrect.

Compare the re-  
sults of No. 19  
with those of No.  
20, the dung being  
similar in every  
respect except that  
the former was ex-  
posed to the sun  
and the latter was  
in the shade.

Compare with No.19

TABLE NO. I. (continued.)

EXPERIMENTAL NUMBER.	POSITION OF DUNG PADS IN VEIL.	DATE EXPERIMENT COMMENCED.	NUMBER OF DAYS IN VEIL.	SPECIMENS EXAMINED.	ORIGINAL WEIGHT OF DUNG IN GRAMS.	WEIGHT OF DUNG AT COLLECTION IN GRAMS.	EGGS PER GRAM.	Larvae recovered with the Baermann Apparatus.			THIRD STAGE LARVAE RECOVERED AFTER INCUBATION.	<i>Cooperia</i> spp.	<i>H. placei</i> .	<i>O. radiatum</i> .	<i>B. phlebotomum</i> .	REMARKS.
								1st Stage	2nd Stage	3rd Stage						
21	Exposed to sun.	3.6.57		F.			140									Compare No.20 and No. 21. The former dung pads were exposed to the sun; the latter were in the shade.
(a)	do.	do.	4	X. 75 C. 400	95	150	150	25	-	9,350 5,000	28% 79%	55% 13%	8% 8%	9% -		
(b)	do.	do.	8	D. 135 C. 95	135	60	180	55	5	18,750 300	59% 77%	29% 2%	11% 18%	1% 3%		
(c)	do.	do.	10	D. 155 C. 105	155	30	180	375	5	2,575 100	76% 65%	11.5% 10%	12% 20%	0.5% 5%		
(d)	do.	do.	14	D. 155 C. 70	155	0	195	-	-	3,750 50	62% 50	25% -	12% -	1% -	Incubated dung cultures of 21(e) were unfortunately not examined.	
(e)	do.	do.	21	D. 125 C. 65	125	0	120	200	212	1,450	63%	18%	18%	1%		
				D. 85	85	90	100	-	-							
22	In shade	3.6.57		F.			140								Compare with No.20: (see remarks above)	
(a)	do.	do.	8	X. 75 C. 400	100	120	120	-	-	9,350 1,925	28% 89%	55% 2.5%	8% 7.5%	9% 1%		
(b)	do.	do.	10	M. 180 C. 115	180	150	225	-	-	3,500 2,200	83% 77%	8.5% 19%	8.5% 3%	- 1%	Unfortunately incubated dung cultures of 22(e) were not examined.	
(c)			14	D. 175 C. 95	175	30	195	-	-	6,300 1,150	67% 98%	21% -	11% 1%	1% 1%		
(d)	do.	do.	21	D. 155 C. 90	155	15	225	5	10	6,200	81%	18%	1%	-		
				D. 120	120	195	195	40	-							





Experimental No.	Date of collection of specimens in veld.	No. of days in veld.	DUNG PAD'S SIZE IN CM/ AND POSITION IN THE VELD.	CLIMATIC CONDITIONS				THIRD STAGE LARVAE RECOVERED.																REMARKS.										
				Mean Maximum Temp. in degrees C.	Mean Minimum Temp. in degrees C.	Mean Relative Humidity per cent.	Total Rainfall in Millimetres.	A DUNG PAD				B SOIL UNDER THE DUNG PAD				C GRASS SURROUNDING THE DUNG PAD.				D GRASS ROOTS AROUND THE DUNG PAD.					E SOIL NEXT TO THE DUNG PAD TO A DEPTH OF 2 CM.									
								Part examined.	<i>Cooberia</i> spp.	<i>H. placid.</i>	<i>O. radiatum.</i>	<i>E. phlebotomum.</i>	Total No. of larvae recovered.	<i>Cooberia</i> spp.	<i>H. placid.</i>	<i>O. radiatum.</i>	<i>E. phlebotomum.</i>	Total No. of larvae recovered.	Distance from A in cm.	Height above soil in cm.	<i>H. placid.</i>	<i>O. radiatum.</i>	Total No. of larvae recovered.		Distance from A in cm.	<i>Cooberia</i> spp.	<i>H. placid.</i>	<i>O. radiatum.</i>	<i>E. phlebotomum.</i>	Total No. of larvae recovered.	Distance from A in cm.	<i>Cooberia</i> spp.	<i>H. placid.</i>	<i>O. radiatum.</i>
1	19/7/55	26	Size: 17x9x1 Exposed to sun.	19.9	0.1	49.9	0	*E.	-	-	-	-	0	-	-	-	-	0-15	-	-	0	0-15	-	+	-	1	0-15	-	-	-	-	0	1	Atmospheric conditions were unsuitable for development.
2	19/7/55	12	Size: Not measured. Exposed to sun.	21.7	3.9	44.5	0	E.	-	-	-	-	0	-	-	-	-	do.	-	-	0	do.	-	-	-	0	do.	-	-	-	-	0	0	Ditto
3	19/7/55	12	Size: 9x8x6 in partial shade.	21.7	3.9	44.5	0	E.	-	-	-	-	0	-	-	-	-	do. 0-10	-	-	0	do.	-	-	-	0	do.	-	-	-	-	0	0	Ditto
4	12/8/55	36	Size: 13x9x2-5 Exposed to sun.	19.1	-0.5	47	0	E.	-	-	-	-	0	-	-	-	-	do.	-	-	0	do.	-	-	-	0	do.	-	-	-	-	0	0	Ditto
5	18/8/55	8	Size: 20x42x5 in partial shade.	20.0	-0.6	35	0	<sup>o</sup> C. <sup>o</sup> D.	+	+	+	-	67 87	-	-	-	-	do. 0-6	-	-	0	do.	-	-	-	0	do.	-	-	-	-	0	67	The first signs of the presence of larvae. The number of larvae recovered is not very significant.
6	20/8/55	26	Size: Not measured. In partial shade	19.9	-0.8	30.5	0	E.	-	-	-	-	0	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	0	No larvae recovered in the dung pad due to unfavourable atmospheric conditions. Specimens B, C, D, and E, accidentally discarded before examination.	
7	8/9/55	12	Size: 12x7x4 in partial shade.	25.6	2.0	37	0	E.	-	-	-	-	0	-	-	-	-	do. 0-20	-	-	0	do.	-	-	-	0	do.	-	-	-	-	0	0	The dung pad was hollowed out by dung beetles which may have accounted for the lack of development.
8	24/9/55	14	Size: 19x7x2-5 Exposed to sun.	28	6.2	30	0	C. D.	+	+	-	-	16 193 209 99.5%	+	-	-	-	do. 0-5	-	-	0	do.	-	-	-	-	do.	-	-	-	-	0	210	Atmospheric conditions were more favourable for larval development. N.B. More larvae were present in the depth of the dung pad than the crust.

\*E. - Entire dung pad.  
<sup>o</sup>C. - Crust of the dung pad.  
<sup>o</sup>D. - Depth of the dung pad.





Experimental No.	Date of collection of specimens in veld.	No. of days in veld.	DUNG PAD'S SIZE IN CM. AND POSITION IN THE VELD.	CLIMATIC CONDITIONS				THIRD STAGE LARVAE RECOVERED.																REMARKS.																
				Mean Maximum Temp. in degrees C.	Mean Minimum Temp. in degrees C.	Mean Relative Humidity per cent.	Total Rainfall in Millimetres.	A DUNG PAD				B SOIL UNDER THE DUNG PAD				C GRASS SURROUNDING PAD.				D DUNG					E GRASS ROOTS AROUND THE DUNG PAD TO A DEPTH OF 2 CMS.															
								Part examined.	<i>Copieria</i> spp.	<i>H. placei</i> .	<i>O. radiatum</i> .	<i>B. phlebotomum</i> .	Total No. of larvae recovered.	<i>Copieria</i> spp.	<i>H. placei</i> .	<i>O. radiatum</i> .	<i>B. phlebotomum</i> .	Total No. of larvae recovered.	Distance from A in cms.	Height above soil in cms.	<i>Copieria</i> spp.	<i>H. placei</i> .	<i>O. radiatum</i> .		Total No. of larvae recovered.	Distance from A in cms.	<i>Copieria</i> spp.	<i>H. placei</i> .	<i>O. radiatum</i> .	<i>B. phlebotomum</i> .	Total No. of larvae recovered.	Total No. of larvae recovered from all sources.								
9	29/9/55	84	Size: 15x6 in partial shade.	23.3	0.0	40	0.2	E.	-	-	-	-	0	-	-	-	-	0	0-10	0-20	-	-	-	-	0	0-10	-	-	-	+	1	0-10	-	-	-	-	0	1	The long period of exposure to field conditions as well as the unsuitable atmospheric conditions, when dung was placed in the veld in the winter probably accounted for the almost negative results.	
10	3/10/55	16	Size: 17x12x7 in deep shade.	27.6	8.6	29	0.4	E.	-	-	-	-	0	-	-	-	-	0	do. 0-2	+	+	-	-	3	do.	-	-	-	-	-	-	-	-	0	0	7	Lung beetles had been active. In spite of very little rain larvae were recovered on the grass.			
11	4/10/55	7	Size: 18x2-5 Exposed to sun.	26.2	8.0	19.5	0.2	C. D.	- +	- +	- -	- -	0 2	-	-	-	-	0	do. 0-2	-	+	-	-	N/C	do.	-	-	-	-	-	-	-	0	0	0	4	As in No. 10.			
12	6/10/55	49	Size: 28x26x0 in partial shade.	25.5	4.8	33.5	0.4	C. D.	+ +	+ -	+ -	- -	71 72	-	-	-	-	0	do. 0-5	-	-	-	-	0	do.	-	-	-	-	-	-	-	0	0-10	-	-	-	0	72	The number of larvae recovered in the dung pad was not very significant.
13	11/10/55	7	Size: 12x6 in partial shade.	27.5	6.9	33.5	0.4	C. D.	+ 72%	- 17%	- 11%	- -	11 377 388 91%	+	+	-	-	38 38 9%	do. 0-5	-	-	-	-	0	do.	-	-	-	-	-	-	-	0	do.	-	-	-	0	426	The soil under the dung pad included small pieces of sawdust left there by dung beetles. This may have accounted for the presence of larvae in the specimen B.
14	13/10/55	10	Size: 10x12x6 Exposed to sun.	27.7	8.4	30	4.0	C. D.	+ 94%	- 1%	+ 4%	- -	44 6,800 6,844 ± 100%	+	-	-	-	1 1 less than 0.001%	do. 0-5	-	-	-	-	0	do.	-	-	-	-	-	-	-	0	do.	-	-	-	0	6845	Excellent recovery of larvae from the dung. No migration occurred in spite of the 4.0 mm. of rain.
15	17/10/55	37	Size: 17x10x4 in partial shade.	27.6	7.9	29.5	10.4	C. D.	+ 83%	+ 0.5%	+ 16.5%	- -	17 3,000 3,017 99.2%	+	-	-	-	5 5 ± 0.3%	do. do.	+	-	-	-	16 16 0.5%	do.	-	-	-	-	-	-	-	0	do.	-	-	-	0	3038	Most of the larvae were confined to the dung. The number of <i>Copieria</i> spp. larvae recovered on the grass was not very significant.



Experimental No.	Date of collection of specimens in veld.	No. of days in veld.	DUNG PAD'S SIZE IN CM/ AND POSITION IN THE VELD.	CLIMATIC CONDITIONS				THIRD STAGE LARVAE RECOVERED.																	REMARKS.										
				Mean Maximum Temp. in degrees C.	Mean Minimum Temp. in degrees C.	Mean Relative Humidity per cent.	Total Rainfall in millimetres.	A DUNG PAD				B SOIL UNDER THE DUNG PAD				C GRASS SURROUNDING THE DUNG PAD				D GRASS ROOTS AROUND THE DUNG PAD				E SOIL NEXT TO THE DUNG PAD TO A DEPTH OF 2 CM.											
								Part examined.	Copieria spp.	H. placei.	O. radiatum.	E. phlebotomum.	Total No. of larvae recovered.	Copieria spp.	H. placei.	O. radiatum.	E. phlebotomum.	Total No. of larvae recovered.	Distance from A in cm.	Height above soil in cm.	Copieria spp.	H. placei.	O. radiatum.	Total No. of larvae recovered.		Distance from A in cm.	Copieria spp.	H. placei.	O. radiatum.	Total No. of larvae recovered.	Distance from A in cm.	Copieria spp.	H. placei.	O. radiatum.	E. phlebotomum.
16	17/10/55	90	Size: Not measured. In partial shade.	23.9	3.3	37	10.4	C. - D. -	-	-	-	-	0	-	-	-	-	0	0-10 0-5	+	-	-	1	0-10	+	-	-	1	0-10	-	-	-	0	2	The long period of exposure probably accounted for the practically negative results.
17	20/10/55	9	Size: 16x16x9 In deep shade.	27.2	10.3	44	10	C. 94% U.D. 95% L.D. 93%	6% 2.5% 5%	- 1.5% 1%	- 0.5% 1%	11900 135000 60000	+	-	-	-	15	do. 0-5 5-60	-	-	-	0	do. +	-	-	1	do. -	-	-	-	0	206916	The dung was well protected against the sun; it's mass was fairly large and all the rain recorded fell on the first five days the dung was in the veld. This probably accounted for the excellent recovery of larvae from the dung pad.		
18	25/10/55	7	Size: 18x10x6 In partial shade.	27.3	7.6	37	0.3	C. + D. 73%	- 4%	- 23%	- -	50 13,000 13,050	+	-	-	-	2	do. 0-5 5-60	+	-	-	11	do. -	-	-	0	do. +	-	-	-	10	13073	Dung beetles had removed some of the dung to the soil. This may have accounted for the presence of larvae in the soil. If this were the case the low rainfall probably assisted migration to the grass.		
19	27/10/55		Size: 20x19x7 Lay in the kraal on Southern side of drinking trough in partial shade. Water was allowed to drip next to dung for 48 hours before collection of specimens.	27.5	9.4	38	6.3	C. 91% U.D. 95% L.D. 93%	3% 2% 2%	6% 3% 5%	- - -	2,132 10,000 1,850	Species not identified		1,450																	8	14400	As mentioned in the fourth column of the table water was allowed to drip next to the dung pad for the last 48 hours before the specimens were collected. The soil under, and the lower surface of the dung pad, were very moist which may have accounted for the downward migration.	
20	5/11/55	11	Size: 28x18x2 In partial shade.	27.9	9.5	44	20.9	B. 78%	-	22%	-	300 300	73%	1%	26%	-	1200 1200	0-10 0-5	-	-	-	0	0-10	-	-	-	0	0-10	+	-	+	-	5	1505	As in No. 13, dung beetles had removed small pieces of manure to the underlying soil and this possibly accounted for the large numbers of larvae there.

U.D. - Upper layer of the depth of the dung.  
L.D. - Lower layer of the depth of the dung.





Experimental No.	Date of collection of specimens in veld.	No. of days in veld.	CLIMATIC CONDITIONS				THIRD STAGE LARVAE RECOVERED.																REMARKS.															
			DUNG PAD'S SIZE IN CM. AND POSITION IN THE VELD.				A DUNG PAD				B SOIL UNDER THE DUNG PAD				C GRASS SURROUNDING DUNG PAD.				D GRASS ROOTS AROUND THE DUNG PAD.					E SOIL NEXT TO THE DUNG PAD TO A DEPTH OF 2 CMS.														
			Mean Maximum Temp. in degrees C.	Mean Minimum Temp. in degrees C.	Mean Relative Humidity per cent.	Total Rainfall in Millimetres.	Part examined.	Cooperia spp.	H. placei.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	Cooperia spp.	H. placei.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	Distance from A in cm.	Height above soil in cm.	Cooperia spp.	H. placei.	O. radiatum.		Total No. of larvae recovered.	Distance from A in cm.	Cooperia spp.	H. placei.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	Distance from A in cm.	Cooperia spp.	H. placei.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	Total No. of larvae recovered from all sources.	
21	6/11/55	71	Size: 23x21x3 In deep shade.	23.5	7.5	36	31.3	C.	-	-	-	-	0	-	-	-	-	0	0-10	0-4	-	-	-	0	0-10	-	-	+	1	0-10	-	-	-	-	0	1	As in No. 16, the long period in the veld probably accounted for the practically negative results.	
22	6/11/55	40	Size: 15x14x5 In deep shade.	27.4	9.5	38.5	31.1	C.	+	+	+	-	33	-	-	-	-	0	do.	0-10	-	-	-	0	do.	-	-	-	-	0	do.	-	-	-	-	0	333	Although 14 mm. of the total of 31.1 mm. of rain fell in the last 24 hours before specimens were collected, this was insufficient to cause migration.
23	6/11/55	24	Size: 13x14x9 In partial shade.	27.3	10.8	44	16.9	C.	90%	4%	5%	1%	630	+	+	+	-	42	do.	0-15	+	-	-	1	do.	-	-	-	-	0	do.	-	-	-	-	0	7423	Rainfall was again insufficient to cause larval migration of any note.
24	21/11/55	16	Size: 18x16x6 In partial shade.	30.8	11.2	31	0	E.	-	-	-	-	0	+	-	-	-	1	do.	0-10	-	-	-	0	do.	-	-	-	-	0	do.	-	-	-	-	0	1	Dung beetles attacked the dung pad to such an extent that only a thin crust remained, probably accounting for the poor recovery of larvae.
25	23/11/55	9	Size: 12x8x6 In deep shade.	30.1	11.4	32	9.5	C.	-	-	-	-	0	-	-	-	-	0	do.	0-10	-	-	-	0	do.	-	-	-	-	0	do.	-	-	-	-	0	0	As in No. 24.
26	23/11/55	9	Size: 11x10x5 In deep shade.	30.1	11.4	32	9.5	C.	-	-	-	-	0	-	-	-	-	0	0-15	0-10	-	-	-	0	0-15	-	-	-	-	0	0-15	-	-	-	-	0	1	As in Nos. 24 and 25.
27	29/11/55	7	Size: 20x20x5 In partial shade.	27.6	15.2	54.5	16.3	E.	7%	47%	46%	-	112	-	+	+	-	15	0-15	0-5	+	-	-	1	0-15	-	-	-	-	0	0-15	-	-	+	-	5	143	As in Nos. 24 to 26, although more larvae were recovered.





Experimental No.	Date of collection of specimens in veld.	No. of days in veld.	DUNG PAD'S SIZE IN CM. AND POSITION IN THE VELD.	CLIMATIC CONDITIONS				THIRD STAGE LARVAE RECOVERED.																	REMARKS.												
				Mean Maximum Temp. in degrees C.	Mean Minimum Temp. in degrees C.	Mean Relative Humidity per cent.	Total Rainfall in Millimetres.	A DUNG PAD				B SOIL UNDER THE DUNG PAD				C GRASS SURROUNDING DUNG PAD.			D GRASS ROOTS AROUND THE DUNG PAD.			E SOIL NEXT TO THE DUNG PAD TO A DEPTH OF 2 CM.															
								Cooperia spp.	H. placcel.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	Cooperia spp.	H. placcel.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	Distance from A in cm.	Height above soil in cm.	Cooperia spp.	H. placcel.	O. radiatum.	Total No. of larvae recovered.	Distance from A in cm.		Cooperia spp.	H. placcel.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	Total No. of larvae recovered from all sources.						
28	29/11/55	7	Size: 21x21x4 In partial shade.	27.6	15.2	54.5	16.3	E.	24%	38%	37%	1%	$\frac{112}{112}$ 44%	16%	36%	42%	6%	$\frac{140}{140}$ 55%	0-15	0-5	-	+	+	$\frac{2}{2}$ 0.8%	0-15	+	-	-	$\frac{1}{1}$ 0.2%	0-15	+	-	+	-	$\frac{2}{2}$ 0.8%	257	As in No. 27.
29	13/12/55	10	Size: 10x10x4 In deep shade.	27.9	14.2	46	38.5	E.	22%	62%	14%	2%	$\frac{350}{350}$ 29%	28%	48%	24%	-	$\frac{112}{112}$ 9%	0-10	0-5	45%	42%	13%	$\frac{125}{125}$ 10%	0-10	+	+	+	$\frac{12}{12}$ 1%	0-10	12%	20%	68%	-	$\frac{622}{622}$ 51%	1221	This was the first experiment in which any migration of note had taken place to the grass. Of the total of 38.5 mm. of rain recorded, 32 mm. fell in the first 48 hours and only 6.5 mm. the last 24 hours before collection.
30	19/12/55	25	Size: 5x4x4 In partial shade.	27.4	14.1	54	127.7	E.	+	-	+	-	$\frac{8}{8}$ 5%	+	+	+	+	$\frac{25}{25}$ 14%	0-10	0-10	+	+	+	$\frac{75}{75}$ 41%	0-10	+	+	+	$\frac{24}{24}$ 13%	0-10	+	+	+	+	$\frac{50}{50}$ 27%	162	Under the influence of good rains, most of the larvae had migrated from the dung. The long period in the veld possibly accounted for the relatively poor recoveries of larvae. Dung beetles had also been active.
31	19/12/55	67	Size: 14x5x4 Exposed to sun.	28.2	12.3	43.5	164.1	E.	-	-	-	-	0	+	-	-	-	12	0-15	0-10	+	-	-	12	0-15	-	-	-	0	0-15	+	-	+	-	3	27	Dung beetles had been active and probably accounted for the poor recoveries of larvae. All the larvae had migrated from the dung due to good rains.
32	19/12/55	93	Size: 10x8x4 In deep shade.	28.2	10.9	39.5	168.5	E.	+	+	+	-	4	+	-	-	-	2	0-10	0-5	+	-	-	$\frac{4}{6}$ 10	0-10	-	-	-	$\frac{0}{0}$	0-10	+	-	-	-	$\frac{1}{1}$	17	Due to the long period of exposure most of the larvae had probably died. Although Cooperia spp. were the only larvae recovered on the grass, both H. placcel and O. radiatum larvae were present in the dung pad. Dung beetles had also been active.



Experimental No.	Date of collection of specimens in veld.	No. of days in veld.	DUNG PAD'S SIZE IN CM. AND POSITION IN THE VELD.	CLIMATIC CONDITIONS				THIRD STAGE LARVAE RECOVERED.																REMARKS.												
				Mean Maximum Temp. in degrees C.	Mean Minimum Temp. in degrees C.	Mean Relative Humidity per cent.	Total Rainfall in Millimetres.	A DUNG PAD				B SOIL UNDER THE DUNG PAD				C GRASS SURROUNDING PAD				D DUNG GRASS ROOTS AROUND THE DUNG P.D.					E SOIL NEXT TO THE DUNG PAD TO A DEPTH OF 2 CMS.											
								Part examined.	Cooperia spp.	H. placei.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	Cooperia spp.	H. placei.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	Distance from A in cm.	Height above soil in cm.	Cooperia spp.	H. placei.	Total No. of larvae recovered.		Distance from A in cm.	Cooperia spp.	H. placei.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	Distance from A in cm.	Cooperia spp.	H. placei.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered from all sources.
33	5/1/56	19	Size: 10x10x4 Exposed to sun.	24.3	14.2	49	51.5	E.	+	+	+	-	87 87 27%	12%	24%	64%	-	162 162 42%	0-20	0-1	+	-	-	7 11 18 5%	0-20	+	+	+	56 56 14%	0-20	+	+	+	62 62 16%	385	Most of the migration was to the soil underneath the dung pad, in spite of the fact that 19.1 mm. of rain fell on the dung over a period of 10 days before collection of specimens. Dung beetles had also been active.
34	13/1/56	8	Size: 12x12x5 in deep shade.	31.6	16.0	45	13.5	E.	-	-	-	-	0	-	-	-	-	0	0-15	0-4	-	-	-	0	0-15	-	-	-	0	0-15	-	-	-	0	0	Only 4 mm. of rain fell during the first 24 hours of exposure to field conditions, and the balance in the last 24 hours. The high temperatures and dry conditions probably caused mortality of the pre-infective larvae.
35	19/1/56	26	Size: 15x14x5.5 in partial shade.	31.4	16.5	45.5	52.1	E.	+	+	-	-	56 56 24%	+	-	+	-	80 80 34%	0-15	0-5	+	+	-	11 25 36 16%	0-15	+	+	-	38 38 17%	0-15	+	+	-	22 22 9%	232	In spite of attempts to protect dung pads from beetles, by placing in the court-yard for the first four days, before transferring to the veld, the dung pad was extensively damaged by beetles.
36	31/1/56	18	Size: 13x10x4 In deep shade.	31.8	16.0	45.5	30.8	E.	63%	-	35%	2%	1500 1500 99.97%	+	-	-	-	2 2 0.13%	0-15	0-5	-	-	-	0	0-15	-	-	-	0	0-15	-	-	-	0	1502	Most of the rain (28.1 mm) fell the first four days dung was exposed. This assisted development of the larvae to the infective stage, but the balance of 2.7 mm. was insufficient for migration.
37	6/2/56	18	Size: 21x18x6 In deep shade.	32.8	17.4	32.5	5.7	C. D.	- +	- -	- -	- -	0 6 6	-	-	-	-	0	0-15	0-5	-	-	-	0	0-15	-	-	-	0	0-15	-	-	-	0	6	The hot dry conditions when dung was first placed in the veld were unfavourable for larval development.

\*Since dung pads were attacked by dung beetles most of the dung pads from No. 34 onwards were placed in the open court-yard of the laboratory for about 4 days before being placed in the veld.





Experimental No.	Date of collection of specimens in veld.	No. of days in veld.	DUNG PAD'S SIZE IN CM. AND POSITION IN THE VELD.	CLIMATIC CONDITIONS				THIRD STAGE LARVAE RECOVERED.																				REMARKS.													
				Mean Maximum Temp. in degrees C.	Mean Minimum Temp. in degrees C.	Mean Relative Humidity per cent.	Total Rainfall in Millimetres.	A DUNG PAD				B SOIL UNDER THE DUNG PAD				C GRASS SURROUNDING THE DUNG PAD.				D GRASS ROOTS AROUND THE DUNG PAD TO A DEPTH OF 2 CM.				E GRASS NEXT TO THE DUNG PAD TO A DEPTH OF 2 CM.																	
								Part examined.	C. spp.	H. placei.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	C. spp.	H. placei.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	Distance from A in cms.	Height above soil in cms.	C. spp.	H. placei.	O. radiatum.	Total No. of larvae recovered.	Distance from A in cms.	C. spp.	H. placei.		O. radiatum.	Total No. of larvae recovered.	Distance from A in cms.	C. spp.	H. placei.	O. radiatum.	Total No. of larvae recovered.	Distance from A in cms.	C. spp.	H. placei.	O. radiatum.	Total No. of larvae recovered.	
																																									Total No. of larvae recovered.
38	16/2/56	18	Size: 14x12x7 Exposed to sun.	31.2	16.5	47	58.3	C. 54% U.D. 75% L.D. 51%	2% 6% 6%	44% 19% 43%	- - -	5,625 12,500 6,250	24,375	± 96.5%	60%	2%	38%	-	887	3.5%	0-25 0-5	- -	- -	0 0	0-15	+	-	+	2	0-15	-	-	+	-	3	25,267	Most of the larvae in the dung pad were recovered in the upper layer of the depth. The last 24 hours before collection of specimens, 19 mm. of rain fell which only stimulated migration to the soil beneath the dung, of any note.				
39	18/2/56	18	Size: 18x18x7 In partial shade.	31.8	17.2	51	19.0	C. 40% D. 46%	48% 49%	4% 1%	8% 4%	112 650	762	± 56%	32%	63%	1%	4%	525	39%	0-15 5-30	0-5 -	- -	0 0	0-15	-	+	-	1	0-15	+	+	+	-	69	1,357	As in No. 38, the rainfall only stimulated migration to the soil beneath the dung pad, of any note.				
40	23/2/56	23	Size: 20x22x15 In partial shade.	30.9	16.7	48.5	51.0	C. 44% D. 41%	56% 56.5%	- -	2.5% -	2,012 10,000	12,012	81%	32%	68%	-	-	2,500	17%	0-15 15-30	0-10 do.	++ ++	- -	65 99	164	1%	0-15 15-30	+	+	-	53 4	0-15 15-30	+	+	-	-	63 26	89	14,822	The last 24 hours before specimens were collected, 27.5 mm. of rain fell. This stimulated migration of larvae but most of the larvae that migrated were recovered from the soil under the dung pad.
41	25/2/56	28	Size: 18x18x5 In deep shade.	29.6	13.2	50.0	84.5	E. + - +	- -	- -	- -	39 39	78	29%	+	-	-	-	10	8%	0-15 30-50 15-50 50-55	0-10 0-50 0-50 0-50	++ ++ ++ ++	- - - -	39 25 4	10	78	58%	0-15	+	+	-	4	0-15 30-50 50-55	- - -	- - -	- - -	1 2 0	3	134	On the 22nd and 23rd of February, two days before the collection of specimens 61.0 mm. of rain fell. Marked horizontal migration took place and more larvae were recovered on grass than elsewhere.













Experimental No.	Date of collection of specimens in veld.	No. of days in veld.	DUNG PAD'S SIZE IN CM. AND POSITION IN THE VELD.	CLIMATIC CONDITIONS				THIRD STAGE										LARVAE RECOVERED.					REMARKS.		
				Mean Maximum Temp. in degrees C.	Mean Minimum Temp. in degrees C.	Mean Relative Humidity per cent.	Total Rainfall in millimetres.	A DUNG PAD				B SOIL UNDER THE DUNG PAD			C GRASS SURROUNDING THE DUNG PAD			D DUNG			E GRASS ROOTS AROUND THE DUNG PAD TO A DEPTH OF 2 CMS.				
								Part examined.	<i>C. coopersia</i> spp.	<i>H. placel.</i>	<i>O. radiatum.</i>	<i>B. phlebotomum.</i>	Total No. of larvae recovered.	<i>C. coopersia</i> spp.	<i>H. placel.</i>	<i>O. radiatum.</i>	<i>B. phlebotomum.</i>	Total No. of larvae recovered.	Distance from A in cm.	Height above soil in cm.	<i>C. coopersia</i> spp.	<i>H. placel.</i>		Total No. of larvae recovered.	Distance from A in cm.
48	9/4/56	9	Size: 30x29x9 In deep shade	26.7	14.3	52	0	C. 57% U.B. 60% L.D. 70%	37% 34% 28%	- 3% 2%	6% 3% -	625 13750 2500 16825 99.7%	+	+	-	+	48 48 0.3%	0-20 0-10 - - - do. 10-30 - - -	0 0 0	0-20 - - -	0 0 0	0-20 - - -	0 0 0	16,923	Notwithstanding the presence of heavy dews no horizontal migration took place and most of the larvae remained in the dung pad.
49	16/4/56	41	Size: 17x16x4 Exposed to the sun.	26.6	14.1	71	172.0	E. + + + -	-	-	61 61 55.4%	+	+	+	-	8 8 7%	0-20 0-10 + + + do. 10-20 + + + do. 20-65 + + -	18 10 8 36 33%	0-20 + + -	3 3 2.7%	0-20 + - - -	2 2 1.7%	110	All the rain recorded fell before the 30th of March. Although no rain fell the last 17 days before collection of specimens, a few larvae were recovered on the grass more than 20 cm. above the soil. However 50% of the larvae on the grass were on the lower 10 cm.	
50	23/4/56	83	Size: 14x10x4.5 In deep shade.	27.2	14.3	62	315.9	E. - - - -	-	-	0	-	-	-	0	0-20 0-10 + - do. 10-25 + -	3 3 1/4	0-20 + - - -	1 1	0-20 + - - -	1 1	6	More rain fell during the period of this experiment, than in any other experiment. However the long period of exposure to field conditions, probably accounted for the poor larval recoveries.		
51	9/5/56	12	Size: 14x10x4.5 In partial shade.	25.4	6.6	46	5	C. + + - - D. + + - -	-	-	7 3 10	-	-	-	0	0-20 0-14 - +	1 1	0-20 - - -	0 0	0-20 + - - -	1 1	12	Drier cooler conditions possibly accounted for the poor development of larvae.		
52	28/5/56	24	Size: 15x13x6 In deep shade.	21.3	3.5	56.5	9.3	C. - - - - D. + - - -	-	-	0 1 1	-	-	-	0	0-15 0-5 - - -	0 0	0-15 - - -	0 0	0-15 - - -	0 0	1	The fact that the dung was in the shade and the colder conditions, possibly retarded larval development to the infective stage.		



Experimental No.	Date of collection of specimens in veld.	No. of days in veld.	DUNG PAD'S SIZE IN CM. AND POSITION IN THE VELD.	CLIMATIC CONDITIONS				THIRD STAGE LARVAE RECOVERED.								REMARKS.																				
				Mean Maximum Temp. in degrees C.	Mean Minimum Temp. in degrees C.	Mean Relative Humidity per cent.	Total Rainfall in Millimetres.	A DUNG PAD				B SOIL UNDER THE DUNG PAD					C GRASS SURROUNDING PAD.				D DUNG GRASS ROOTS AROUND THE DUNG PAD.				E SOIL NEXT TO THE DUNG PAD TO A DEPTH OF 2 CMS.											
								Copieria spp.	H. placel.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	Copieria spp.	H. placel.	O. radiatum.		B. phlebotomum.	Total No. of larvae recovered.	Distance from A in cm.	Height above soil in cm.	Copieria spp.	H. placel.	Total No. of larvae recovered.	Distance from A in cm.	Copieria spp.	H. placel.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	Distance from A in cm.	Copieria spp.	H. placel.	O. radiatum.	B. phlebotomum.	Total No. of larvae recovered.	Distance from A in cm.
53	28/5/56	17	Size: 13x12x8 Exposed to the sun.	20.4	2.4	58	4.6	C. 83% D. 72%	14% 13%	3% 7%	- 8%	600 2250 3350 ± 100%	+	-	-	-	1 0.003%	0-15	0-5	-	-	0 0	0-15	+	-	-	-	1 0.003%	0-15	-	-	-	-	0 0	3,352	In contradistinction to No. 52, large numbers of larvae were recovered. This was possibly due to the exposure of the dung pad to the sun where temperatures were higher than in the shade.
54	13/6/56	25	Size: 18x13x7 Exposed to the sun.	18.1	2.2	59	4.0	C. + D. +	+ +	- -	- -	7 8 15	-	-	-	-	0 0	0-15	0-7	-	-	0 0	0-15	-	-	-	-	0 0	0-15	+	-	-	-	2 2	17	In spite of exposure to sunlight atmospheric temperatures were too low, to allow any appreciable development to the infective stage.
55	14/6/56	63	Size: 15x11x10 in partial shade.	22.5	4.2	54.5	9.4	C. + D. +	- -	+ -	- -	4 6	-	-	-	-	0 0	0-15	0-25	-	-	0 0	0-15	-	-	-	-	0 0	0-15	+	-	-	-	2 2	8	The long period of exposure to field conditions may have accounted for the poor larval recoveries.
56	26/6/56	32	Size: 20x13x8 in partial shade.	19.5	0.9	57	0	C. - D. +	- -	- -	- -	0 8 8	-	-	-	-	0 0	0-15	0-5	-	-	0 0	0-15	-	-	-	-	0 0	0-15	-	-	-	-	0 0	8	As in No. 52.
57	18/7/56	104	Size: 18x14x4 In deep shade.	22.5	3.8	47	9.4	C. - D. -	- -	- -	- -	0 0	+	+	-	-	7 7	0-10	0-25	-	-	0 0	0-10	+	-	-	-	5 5	0-10	+	+	-	-	10 10	22	Dung beetles had been active which may have accounted for the presence of larvae in the soil and on the roots of grass. Note the long period of exposure to field conditions.
58	7/8/56	105	Size: 16x14x12 Exposed to the sun.	21.4	1.9	50	9.4	C. & U.D. L.D.	42% + +	5% + +	53% - -	1,375 5 1,380 99.6%	+	-	-	-	4 4 0.28%	0-10	0-18	-	-	0 0	0-10	-	-	-	-	0 0	0-10	+	-	-	-	2 2 0.14%	1386	This was the longest period of exposure to field conditions and a fairly large number of larvae were recovered, most of them being confined to the dung pad. The dung pad was very thick in the centre and possibly this assisted in keeping the upper layer of the depth moist.



TABLE NO. III.

INFECTIVE LARVAE RECOVERED FROM KRAAL MANURE.

Date.	Number of larvae recovered from each specimen.				Total.	<u>Cooperia</u> spp.	<u>H. placei</u> .	<u>O. radiatum</u> .	<u>B. phlebotomus</u> .
	A	B	C	D					
29/2/56	0	0	0	3	3				
9/3/56	8	5	0	0	13				
22/3/56	3	9	29	3	44				
28/3/56	3	2	3	6	14				
3/4/56	24	1	2	1	28				
12/4/56	0	0	0	0	0				
19/4/56	12	8	1	0	21				
25/4/56	0	0	0	0	0				
4/5/56	0	0	0	0	0				
9/5/56	0	0	2	18	20				
16/5/56	0	0	1	0	1				
25/5/56	0	0	0	0	0				
30/5/56	0	0	0	1	1				
14/6/56	0	2	3	1	6				
4/7/56	0	0	0	6	6				
19/7/56	0	0	0	3	3				
9/8/56	0	0	0	0	0				
28/8/56	0	1	1	0	2				
12/9/56	1	1	0	5	7				
19/9/56	0	0	0	0	0				
25/9/56	0	0	0	0	0				
2/10/56	0	0	0	0	0				
11/10/56	0	0	12	0	12				
23/10/56	12	18	12	0	42				
31/10/56	0	0	0	0	0				
7/11/56	0	0	0	0	0				
20/11/56	25	0	0	0	25				
30/11/56	0	7	10	5	22				
5/12/56	2	300	250	0	552				
11/12/56	0	2	0	1	3				
19/12/56	1	0	0	4	5				
26/12/56	0	0	2	1	3				
1/1/57	0	0	3	7	10				
8/1/57	0	2	0	1	3				
15/1/57	0	0	0	0	0				
22/1/57	0	0	0	0	0				
27/1/57	0	0	0	0	0				



T A B L E N O. III. A.

MEAN NUMBER OF INFECTIVE LARVAE PER KG. OF  
MANURE IN THE INFESTED KRAAL.

<u>Date.</u>	<u>No. of larvae per Kg. of manure.</u>	<u>Cooperia spp.</u>	<u>H. placei.</u>	<u>O. radiatum.</u>	<u>B. phlebotomus.</u>
5/2/57	380	X	X	-	-
12/2/57	6	X	-	-	-
19/2/57	0	-	-	-	-
26/2/57	88	X	-	-	-
5/3/57	0	-	-	-	-
12/3/57	0	-	-	-	-
19/3/57	10	-	-	-	-
26/3/57	40	X	-	X	-
2/4/57	0	-	-	-	-
9/4/57	30	X	-	-	-
16/4/57	986	X	X	X	X
23/4/57	17	X	-	-	-
30/4/57	9	-	X	-	-

TABLE NO. IV MEAN NUMBER OF LARVAE PER POUND OF HERBAGE. PLOT 1 WAS NEXT TO THE KRAAL AND PLOT 2 WAS APPROXIMATELY 300 YARDS FROM THE KRAAL.

Date.	No. of larvae per lb. of herbage.		Date.	No. of larvae per lb. of herbage.	
	Plot 1	Plot 2		Plot 1	Plot 2
29/2/56	223	0	12/12/56	214	0
6/3/56	99	21	19/12/56	0	0
14/3/56	270	54	26/12/56	0	0
21/3/56	180	51	2/1/57	0	0
28/3/56	28	25	9/1/57	0	0
5/4/56	98	30	15/1/57	0	0
10/4/56	43	19	22/1/57	0	0
17/4/56	63	28	29/1/57	0	0
24/4/56	48	0	5/2/57	0	974
12/5/56	0	0	12/2/57	0	0
23/5/56	51	0	19/2/57	0	0
29/5/56	0	19	26/2/57	104	0
7/6/56	0	0	5/3/57	0	0
22/6/56	0	0	12/3/57	0	0
4/7/56	0	0	19/3/57	28	0
11/7/56	0	0	26/3/57	0	0
6/8/56	0	0	2/4/57	133	17
19/8/56	0	0	9/4/57	0	0
26/8/56	0	0	16/4/57	38	25
9/10/56	0	0	23/4/57	15	0
11/10/56	0	0	30/4/57	0	0
17/10/56	0	0			
23/10/56	0	0			
31/10/56	0	0			
7/11/56	0	0			
20/11/56	16	0			
30/11/56	29	0			
5/12/56	0	0			

**TABLE NO. V. - FIVE GROUPS OF CALVES USED IN EXPERIMENT G.**

Group.	No. of calf.	Date of birth and introduction into group.	Method calves were reared from birth.
1.	8237	28/3/56	Handreared and allowed access to the infested kraal from 2 p.m. to 7 a.m. From 7 a.m. to 2 p.m. confined to an adjacent calf pen with a ground floor covered with manure.
	8244	8/7/56	
	8246	12/7/56	
	8249	30/3/56	
	8250	3/9/56	
2.	8235	26/3/56	Confined to a calf pen with a concrete floor which was cleaned twice daily. Suckled on dams at 7 a.m. and 2 p.m. Dams confined to the infested kraal from 7 a.m. to 2 p.m.
	8238	27/3/56	
	8240	1/6/56	
	8247	15/7/56	
	8253	24/9/56	
3.	8227	29/2/56	Grazed in the infested paddock and suckled on dams there at 7 a.m. and 2 p.m. Dams grazed elsewhere. Once a week calves were herded in to the crush next to the kraal for faecal collections.
	8229	14/3/56	
	8245	9/7/56	
	8248	6/3/56	
	8251	16/9/56	
	8252	19/9/56	
4.	8230	21/3/56	Confined to the infested kraal with access to the calf pen used by calves of group 1 from 2 p.m. to 7 a.m. Suckled on dams in the infested kraal at any time from 7 a.m. to 2 p.m.
	8231	22/3/56	
	8232	24/3/56	
	8233	24/3/56	
	8234	26/3/56	
5.	8238	14/4/56	Grazed in the infested paddock from 2 p.m. to 7 a.m. and suckled on dams in the infested kraal at any time from 7 a.m. to 2 p.m.
	8239	28/4/56	
	8241	22/5/56	
	8242	29/5/56	
	8243	6/6/56	

**TABLE NO. VI - SEVEN GROUPS OF CALVES USED IN EXPERIMENT D.**

Group	No. of calf.	Date of Birth.	Date of introduction into group.	Method of calf rearing from birth.
1.	13	5/11/56	15/11/56	As in group 1, Experiment C.
	14	6/11/56	15/11/56	
	15	7/11/56	15/11/56	
	21	10/11/56	15/11/56	
	21	13/11/56	15/11/56	
2.	10	4/11/56	13/11/56	As in group 2, Experiment C.
	12	5/11/56	13/11/56	
	18	9/11/56	13/11/56	
	28	12/11/56	13/11/56	
	22	25/11/56	25/11/56	
3.	28	7/11/56	13/11/56	As in group 3, Experiment C.
	36	14/11/56	14/11/56	
	31	13/11/56	13/11/56	
	32	14/11/56	14/11/56	
	39	21/11/56	21/11/56	
4.	24	11/11/56	13/11/56	As in group 4, Experiment C.
	26	10/11/56	13/11/56	
	30	20/11/56	20/11/56	
	31	21/11/56	21/11/56	
	39	29/11/56	29/11/56	
5.	6	31/10/56	13/11/56	As in group 5, Experiment C.
	17	8/11/56	13/11/56	
	27	11/11/56	13/11/56	
	30	22/11/56	22/11/56	
	37	22/11/56	22/11/56	
6.	16	7/11/56	13/11/56	Handreared on concrete floors that were cleaned twice daily with water and brooms. Mucorine in starchiens was also fed to the calves. This was a control group and similar to Group C. Experiment B.
	19	9/11/56	13/11/56	
	20	9/11/56	13/11/56	
	23	11/11/56	13/11/56	
	32	15/11/56	13/11/56	
7.	88	22/11/56	22/11/56	Suckling calves kept on concrete floors which were cleaned twice daily with water and brooms. Before the calves suckled at 7 a.m. and 2 p.m. the dams udders were washed. This was also a control group.
	110	6/12/56	6/12/56	
	119	11/12/56	11/12/56	
	121	13/12/56	13/12/56	
	124	13/12/56	13/12/56	

**NOTE** - Calves born prior to 13/11/56 were placed on concrete floors and then placed in groups. Calves born after 13/11/56 were placed into groups at birth.