
**Induced Cobalt Deficiency in Lambs**

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The object of the experiments described below was to induce cobalt deficiency in lambs by giving a diet composed of foodstuffs sufficiently remote from pasture to avoid the criticism that some unknown pasture factor other than cobalt might be involved in the causation of the marasmus, now called ‘cobalt deficiency’, and to prove conclusively that a lack in the diet of cobalt *per se* is responsible for ‘pining’ in lambs.

**Experimental**

*First experiment, 1946*

**Basic ration.** The basic daily ration used was 1100 g. flaked maize, 250 g. hay and 60 g. of a mineral mixture composed of ground limestone, steamed bone-flour and crude rock salt. The introduction of hay was necessary to ensure that the rumen processes would be normal, as it was not considered satisfactory to use wood pulp or a similar source of cellulose, since it might cause upsets in rumination, regarding which little is known. The hay was kept to a minimum (250 g./day) and was obtained from a field of which the soil was deficient in cobalt.

The diet was adequate in starch and protein equivalent for a pregnant or lactating ewe. According to Woodman (1948), a 120 lb. ewe requires 10 lb. starch equivalent (s.E.)/week for maintenance and 4 lb. starch equivalent (s.E.)/week/gal. milk; and 0.46 lb. protein equivalent (p.E.)/week for maintenance and 1 lb. protein equivalent (p.E.)/week/gal. milk. The starch equivalents of the flaked maize and hay were 84 and 30 respectively, and the protein equivalents were about 10 and 3. Thus the diet outlined above supplied the energy requirements for maintenance and for at least 1 gal. weekly of milk, the average weight of the ewes being 115 lb.

The cobalt content of the ration was assayed spectrographically with the following result:

| Flaked maize | 0.025 p.p.m. |
| Hay         | 0.10 p.p.m.  |
| Mineral mixture | 0.27 p.p.m. |
Thus the daily intake of cobalt/ewe was about 0.0687 mg. Little is known of the daily requirements of sheep for cobalt. Australian research workers (Filmer & Underwood, 1937) have reported cure and prevention of cobalt deficiency by dosing with as small an amount of cobalt as 0.1 mg./day. The cobalt content of the Scottish pastures on which lambs pine ranges from 0.10 to 0.03 p.p.m. (Stewart, Mitchell & Stewart, 1941), and if lambs eat as much as 2.5 lb. dry matter daily they would be ingesting 0.1135-0.034 mg. cobalt. It was considered possible that the experimental diet containing 0.0687 mg./day might be low enough to produce pine in lambs.

![Fig. 1. Live-weight curves of lambs in the 1946 experiment.](https://www.cambridge.org/core/cover/54.70.40.11, on 10 Oct 2019 at 00:59:28, subject to the Cambridge Core terms of use, available at https://www.cambridge.org/core/terms).

**Sheep.** On 1 January 1946 nine in-lamb ewes were divided into three equal groups and placed in boxes with cement floors covered with peat-moss litter. They were fed the above ration, but each member of one group received weekly 10 mg. of cobalt, as cobalt chloride.

The ewes ate the ration readily, but several deaths occurred during the last week of pregnancy due to 'lambing sickness', probably brought about by the nature of the diet, highly digestible flaked maize, and by the lack of exercise due to restraint of the stalls.

**Lambs.** There were eventually on experiment five lambs, three, nos. 980, 981 and 988, from control ewes receiving the diet and 10 mg. cobalt/week and two, nos. 978 and 993, from the ewes receiving the diet only. Nos. 980, 981 and 988 received 10 mg. cobalt/week from birth and, when weaned at 10 weeks of age, were given 800 g.
flaked maize, 80 g. hay and 25 g. mineral mixture daily, which supplied 0.0347 mg. cobalt. The ration was steadily increased till at 16 weeks of age the lambs were being offered the full amount given to their mothers. This appeared to be the limit of appetite and occasionally food was left uneaten, so that the cobalt ingested from the diet was never above 0.0687 mg./day.

Further experiments, 1947–8

Basic ration. The basic ration was the same as in the first experiment.

Despite the foodstuffs being from new sources the cobalt content was still about 0.0687 mg./day, compensation for any increase in the cobalt content of the feeding-stuffs being made by altering slightly the amount of mineral mixture fed. Several of the ewes died of ‘lambing sickness’ in both years, but in similar experiments in 1949 and 1950, when great care was taken to exercise the ewes once a week by allowing them to run in concrete yards, no deaths occurred, the ewes all remaining healthy and lambing successfully. It is concluded that in the previous years the deaths in ewes on this diet were caused by the enforced lack of exercise rather than by the composition of the ration. Several lambs died in both 1947 and 1948 in the neonatal stage from a pyaemia caused by a staphylococcal infection.

Lambs. Lambs nos. 334, 338, 339, 340, 768, 780 and 806 were from ewes receiving cobalt supplement and lambs nos. 335, 336, 345, 785, 787, 788, 800 and 816 were from ewes receiving the diet only.

Lamb no. 816 within a few days after birth showed signs of a staphylococcal infection, and shortly afterwards its mother died. From this stage it was raised on cow’s milk and transferred to the control group receiving 10 mg. cobalt weekly after the 3rd week of life.

The lambs were managed exactly as in 1946 but fed individually.

RESULTS

First experiment, 1946. Lambs nos. 980, 981 and 993, receiving 10 mg. cobalt/week, made steady gains in weight, as seen in Fig. 1. Their live-weight curves correspond to those of lambs on good pasture. Lambs nos. 978 and 993 were offered the same diet as the others, and it will be seen from Fig. 1 that they showed similar weight gains until about the 14th week of age, no. 978 being the heaviest of all the five lambs at that age. From the 15th week no. 978 began to lose weight steadily and died at the 25th week, showing all the symptoms of cobalt deficiency and weighing only 38 lb. The weight of no. 993 from the age of 14 weeks fluctuated round 40 lb. and at 31 weeks was only 46 lb., as compared with 66, 72 and 75 lb. for nos. 980, 981 and 988 respectively. From this age no. 993 was dosed with 10 mg. cobalt weekly. Its weight, after falling for the 1st week, rose steadily and at 48 weeks of age was 68 lb., an increase of 27 lb. in 17 weeks, the live-weight curve being parallel to that of the control lambs.

Lambs nos. 978 and 993 had smaller appetites than the controls. They were continually leaving food, and various methods were tried to persuade them to eat more. Neither ever ate the full ration of flaked maize, hay and mineral mixture, so that their
cobalt intake was always less than 0.0687 mg./day. When lamb no. 993 was given cobalt at 31 weeks old it began to eat its ration more quickly, but even at 48 weeks it still left a quantity in the feeding trough.

Further experiments, 1947–8. The live-weight curves of the lambs, with the exception of nos. 787 and 788, are shown in Fig. 2. Lamb no. 816 made little progress for the first 10 weeks but from that age made normal live-weight gains, reaching 75 lb. at the 35th week. Lamb no. 340, which had never made normal live-weight gains, died at 18 weeks of age. A post-mortem examination failed to reveal the cause of death.

![Fig. 2. Live-weight curves of lambs in the 1947 and 1948 experiments. ---, lambs receiving cobalt-deficient diet and a cobalt supplement; ..., lambs receiving cobalt-deficient diet only.](https://www.cambridge.org/core)

All the control lambs receiving 10 mg. cobalt/week from birth made normal live-weight gains, with the exception of no. 338 which weighed only 74 lb. at the 40th week. Lamb no. 806 fluctuated in weight after the 15th week, but by the 25th week resumed gaining weight in a normal manner. The lambs on deficient diet only showed the effects of this diet at the 15th week. Lamb no. 785 continued on the deficient diet, but died at the 38th week with all the clinical symptoms of cobalt deficiency and weighing only 38 lb. The other lambs in this group received cobalt at different stages once they had obviously shown that their weight curves were on the decline, and immediately made spectacular increases in weight. Anomalous behaviour was shown by lamb no. 335. This animal at 25 weeks of age weighed only 60 lb. and appeared likely to become a typical case of cobalt deficiency if left on the deficient diet. No cobalt was given but, despite this, it began to gain weight and at 50 weeks of age...
weighed 85 lb. There is no accountable reason for this behaviour of no. 335 except that the diet of 0.0687 mg. cobalt must be near the margin of adequacy and that for exceptional animals this amount is sufficient to allow them to keep their appetite and so maintain, or even gain, weight.

Lambs nos. 787 and 788 were twin lambs of a ewe on the deficient diet only. Their weight curves are shown in Fig. 3.

They made identical weight gains till 15 weeks of age, when each weighed 34 lb. At this age no. 788 received 10 mg. cobalt weekly and increased its weight by 21 lb. in 8 weeks, whereas no. 787, receiving no cobalt, lost 6 lb. in the same period. At the 23rd week no. 787 was also given cobalt at the rate of 10 mg. weekly and in the next 17 weeks added 32 lb. to its weight, whilst no. 788 added 35 lb.

DISCUSSION

Three postulates must be fulfilled to justify the statement that a deficiency of a particular constituent of the diet is the cause of a definite disease syndrome in a certain species of animal.

(1) It should be possible to correlate every case of the disease with a real deficiency of the particular constituent in the diet under consideration.

(2) Addition of the particular constituent to the deficient diet should cure or prevent the disease.

(3) The disease condition should be capable of induction in healthy animals by the feeding of an artificial diet, adequate in every essential constituent except the particular constituent, which should be of a content comparable to that in the diet of naturally occurring cases of the disease.
Of these postulates, no. 3 has been often ignored of recent years, with the result that we are still in doubt regarding the causal role of many dietary constituents in well-established disease syndromes, and this is especially so with regard to the trace elements. It is not sufficient to use the criterion of (1) low intake, (2) depletion of the animal, as shown by tissue analysis, and (3) dramatic response after addition of a constituent to the diet or by injection of the animal, since the interrelationships of trace elements one with another have confused the true significance of the part played by several trace elements in disease syndromes, e.g. copper and molybdenum in ‘teart’ disease, copper and cobalt in ‘coast’ disease and the anomalous position of copper in the prevention of swayback in lambs in Great Britain, which usually occurs on pastures of normal copper content.

Several Australian workers (Marston, 1949) have produced the syndrome of cobalt deficiency by cutting herbage from a suspected cobalt-deficient area and feeding it to sheep kept indoors, and Bowstead and his colleagues in Canada (Bowstead & Sackville, 1939; Bowstead, Sackville & Sinclair, 1941–2), by maintaining sheep on a basal diet of non-leguminous hay and ground oats, reproduced a condition of unthriftiness which, after many curative attempts with a large number of substances, all without real effect, responded to cobalt therapy, but again the foodstuffs fed were all harvested off the suspected soil. Therefore neither of the above types of investigation fulfils postulate no. 3, since the ration used does not rule out the presence of some still unknown factor causing the physiological upset curable by cobalt supplements, similar to the action of copper in preventing molybdenum poisoning in ‘teart’ disease.

The experiments described in this paper therefore were attempts to fulfil the third postulate as far as cobalt deficiency was concerned as a causal factor of ‘pine’ in lambs and, although hay from a cobalt-deficient area was used, it was kept to a minimum, being only one-sixth of the total dry matter fed.

The majority of the experimental animals receiving the basal diet and cobalt made the normal live-weight gains to be expected from the starch equivalent and protein equivalent of the ration, reaching approximately 100 lb. live weight in about 40 weeks. Those receiving no cobalt supplement made normal progress till about the 10th–14th week when they had reached a weight of 40 lb., then their weight remained steady or decreased and, if continued on the deficient basal diet, the lambs showed eventually the signs of marasmus identical with that attributed to cobalt deficiency. The live-weight curves of both the control and the experimental animals were very similar to those obtained in experiments carried out in the field (Stewart, 1946). There was a dramatic response with every deficient lamb given cobalt, the two outstanding examples being the twins, nos. 787 and 788.

It would appear that the diet designed for these experiments which allowed the lambs a maximum cobalt intake of 0.0687 mg./day will produce the symptoms of cobalt deficiency, and that an immediate recovery response is obtained by allowing the marasmic lambs to ingest cobalt per se.

This type of diet might be of great value in the production of cobalt deficiency in experimental lambs and should facilitate work being carried out on the physiological action of cobalt and other research on cobalt deficiency.
SUMMARY

1. In-lamb ewes were fed a diet of 1100 g. flaked maize, 250 g. hay and 60 g. mineral mixture daily for the last 3 months of pregnancy. This diet allowed of a daily ingestion of 0.0687 mg. cobalt. Their lambs when weaned at 10 weeks of age were given 800 g. flaked maize, 80 g. hay, 25 g. mineral mixture daily (0.0347 mg. cobalt), increasing steadily to the quantities in the ewe’s diet at 16 weeks of age. These lambs all eventually showed the clinical signs of cobalt deficiency but made immediate dramatic response if fed 10 mg. cobalt/week.

2. Lambs given the same basal diet with a supplement of 10 mg. cobalt from birth all gained weight normally.

3. The necessity for this type of experimentation in trace-element research is stressed.

4. The usefulness of a diet such as that described for the production of cobalt deficiency to facilitate work on the physiological action of cobalt and other research on cobalt deficiency is demonstrated.

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REFERENCES