

ABSTRACT

MANGANESE NUTRITION OF THE PIG

By

Henry B. Kayongo-Male

Four experiments involving 104 pigs were conducted to study different aspects of manganese (Mn) nutrition in swine.

In the first experiment, a basal diet (16.2 ppm Mn) and basal diet supplemented with 10 ppm of Mn from $\text{MnSO}_4 \cdot \text{H}_2\text{O}$, MnCO_3 or MnO were compared for Mn availability to the growing pig. Growth rates were equal on all diets. Mn availability as measured by Mn balance data and tissue Mn concentrations indicated that Mn from the supplemented diets was more available than that from the basal diet. Mn retention, as a percent of intake, was higher on Mn from the supplemented diets than the basal diet. Regardless of dietary Mn source, over 90% of excreted Mn was recovered in the feces. Within the supplemented diets, Mn was essentially equally available to the growing pig. Hemoglobin, hematocrit, serum Mn and serum alkaline phosphatase did not differ significantly due to dietary treatment.

In the second experiment, flux patterns across the wall of the gastrointestinal tract of Mn from different sources was studied. Net absorption of Mn from the basal diet was evident in two sections of the gut, the stomach and the cecum, whereas Mn from the supplemented diets was apparently absorbed in the stomach, cranial small intestine and cecum.

The net cecal absorption of Mn from the basal diet was higher than that of Mn from the supplemented diets. Net Mn secretion in the caudal small intestine and the rectum was much higher on the supplemented than on the basal diet, but this trend was reversed in the colon. The pH values of the gut contents from different sections of the tract were not significantly different between dietary treatments.

In the third experiment, two ratios of Ca to P, two levels of Ca and P and two levels of Mn were studied using a factorial feeding trial. Mn supplementation significantly increased heart Mn levels and significantly depressed rib Ca and Mg values. Mn supplementation did not affect serum Ca, inorganic P, Mg and alkaline phosphatase levels. Dietary Mn levels had no significant effect on rib and metacarpal physical measurements, breaking strength and related parameters. A 2 to 1 Ca to P ratio significantly ($P < 0.05$) depressed rib Mn content. The increased levels of Ca and P supplementation significantly ($P < 0.01$) increased rib, pancreas and serum Mn levels but significantly ($P < 0.01$) depressed metacarpal Mn concentration. There was a significant ($P < 0.01$ to $P < 0.05$) interaction between levels of Ca and P and ratios of Ca to P on the levels of serum, liver and pancreas Mn, and on metacarpal Mn values. High levels of Ca and P in both ratios had a depressing effect on metacarpal Mn concentration. Feeding Mn along with Ca and P, in a 2 to 1 ratio, increased liver Mn. Metacarpal Mg was depressed when Ca and P were given in a 1 to 2 ratio.

The interaction between Ca and P levels and Mn levels was significant ($P < 0.05$) with respect to rib ash content, Ca and P, metacarpal Mn and serum inorganic P. With lower Ca and P levels, Mn supplementation increased metacarpal Mn and serum inorganic P but depressed rib ash, Ca and P concentration. The significant effects of Mn supplementation on

rib Mg, metacarpal internal vertical diameter and heart and serum Mn disappeared when Ca and P supplements were also fed. The 3-way interaction between level of Ca and P, ratio of Ca to P and level of Mn was significant ($P < 0.05$) relative to rib and serum Mn levels, pancreas dry matter and metacarpal Mg content and elasticity. With high or low Ca and P levels in a 2 to 1 ratio, Mn supplementation increased rib and serum Mn and pancreas dry matter but depressed metacarpal elasticity. With low Ca and P levels in a 1 to 2 ratio, Mn supplementation increased serum and metacarpal Mn, but high Ca and P levels in the same ratio depressed rib and serum Mn, metacarpal Mg and pancreas dry matter and increased metacarpal elasticity.

Mn supplementation produced more nearly normal histologic structure of the epiphysis than the basal diet, but animals on high Mn levels had significantly ($P < 0.05$) less compact bone in the diaphysis. However, the thickness of the epiphyseal cartilagenous plate was not affected. There was a significant ($P < 0.05$) interaction between Ca to P ratio and Mn level on the histology of the epiphysis. The interaction of diet Mn with Ca and P levels was significant ($P < 0.05$) in relation to the thickness of the epiphyseal cartilagenous plate. These changes were not typical of rickets but were changes in which there was failure of production of compact bone in the region of the diaphysis. However, the deleterious effects on weight gain, feed efficiency and histology of bone of a low dietary P level (0.35%) from soybean meal were much more pronounced than the effects of excessive dietary levels of Ca and P or of an inverse Ca to P ratio, regardless of dietary Mn supplementation.

In the fourth experiment, the Mn requirements of the baby pig born of sows fed a low Mn diet were determined using three dietary Mn concentrations. Growth, Mn balance data and serum Mn concentration were used

as measures of sufficiency. Average daily gain, serum Mn, Mn retention, fecal Mn excretion and urinary Mn excretion as percent of intake were significantly ($P < 0.01$ or $P < 0.05$) different between dietary treatments. The average daily gain and feed efficiency were highest on 2.67 ppm Mn in the diet. Mn intake was highly correlated with serum Mn and serum alkaline phosphatase activity but not with average daily gain and feed efficiency. Mn retention as percent of intake had very high negative correlations with feed efficiency and urinary Mn excretion as percent of intake. Fecal and urinary Mn excretion as percent of intake was significantly higher on the basal diet (0.46 ppm Mn). There was a negative Mn retention on this diet. Serum Mg levels substantially declined in pigs on the basal diet 28 days after the start of the experiment. Average daily gain was positively related to Mn retention as percent of intake and negatively related to fecal Mn excretion as percent of intake. Mn retention and fecal Mn excretion, both as percent of intake, were much more highly correlated with growth rate than absolute Mn intake, excretion and retention and the serum parameters examined.

Based on all criteria examined, the dietary Mn requirements of the baby pig on a semipurified diet are probably between 3 and 6 ppm.

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By

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A DISSERTATION

**Submitted to
Michigan State University
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Dedicated to

My Beloved Parents

... provided the necessary facilities... comfortable and fruitful.

I must express my sincere thanks to Dr. [Name]... the statistical analysis will always be remembered... graduate students and laboratory technicians... deal of assistance during the course of this study... and Miss Phyllis Whetter were particularly outstanding... thank also Dr. E. W. Wilkinson and Mr. S. [Name] of the Dept. of Chemical Engineering for their tremendous assistance... Fuller for her skillful preparation of this manuscript.

My parents have been a great inspiration throughout my entire program. Their encouragement and support have been my mainstay. I love them. Most important, I am indebted to my wife Diane, whose love, help and encouragement were outstanding.

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My parents have been a great inspiration throughout my study program. Their encouragement and support have been my motivating force. I love them.

Most important, I am indebted to my wife Diane, whose sacrifice, help and encouragement were outstanding.

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INTRODUCTION

Nutritional studies with manganese (Mn) were stimulated by the discovery that two poultry diseases, perosis and chondrodystrophy, were caused by inadequate intakes of Mn and could be prevented by Mn supplementation. Since then Mn has been assigned a growing number of biological functions vitally important to the organism. This element is now associated with normal activation of enzymes and co-enzymes associated with feed utilization. Manganese plays an important role in liver function, bile production, energy metabolism, reproduction and proper skeletal development. The occurrence of Mn in all tissues and the small range of Mn concentrations indicate a role of this element in general cell metabolism as opposed to metabolic processes characteristic of particular tissues.

Although Mn is an indispensable micro-element of considerable practical importance in animal feeding, its metabolism in swine has had limited research, with little attention to its availability, site of absorption and requirement at different stages of development. Manganese interactions with other elements and vitamins, shown in other species, have not been studied in swine.

Manganese compounds have different chemical and physical properties which are important in determining their availability to animals. Since the cost of Mn compounds varies widely, it is important not only to define accurately the nutritive requirement of Mn for swine at

LITERATURE REVIEW

A. Manganese: An Essential Trace Element

1. Growth and Development

The first experiments with rats and mice attempting to demonstrate the essentiality of Mn using purified diets were suggestive but not conclusive. In 1931 Kemmerer *et al.* found better growth rates of female mice when 0.1 mg of Mn was added daily to a milk diet containing additional Fe and Cu. Since that time many workers have demonstrated the essentiality of Mn for optimal growth in rats. Boyer *et al.* (1942) reported that young rats from Mn-deficient dams showed a markedly poorer growth rate and poorer feed utilization than similar rats on the same diet supplemented with Mn. Randoin (1944) found similar subnormal growth, followed by a rapid decline and death. Holtkamp *et al.* (1950) showed that a subcutaneous injection of colloidal Mn improved growth rate. Supplements of Mn improved growth rate and feed efficiency in guinea pigs and rabbits (Everson, 1970; Ellis *et al.*, 1947). Rabbits fed a Mn-free diet showed signs of anorexia (Rudra, 1944a) and, when kept on this diet for a long time, the animals ceased to grow normally and died. Postmortem examination revealed hemorrhages of the lungs, liver and intestinal capillaries.

Wachtel *et al.* (1943) reported that Mn deficiency in rats caused poor bone formation and Frost *et al.* (1959) observed delayed skeletal maturation in rats on Mn-deficient diets. O'Dell (1961) described short

different stages of their development but also to define the less expensive but equally effective ingredients to use in the mineral premix.

The Mn flux pattern across the gastrointestinal mucosa in swine has not been extensively studied. Such influx-efflux studies can provide information on the extent of absorption of the mineral and its secretion, in various sections of the digestive tract.

Manganese interactions with Ca and P have been studied in some detail in poultry and laboratory animals, but most of those findings have not been examined in swine. Interactions of this element with other elements, such as copper and iron, and vitamins have not been studied in swine. Verification of any Mn interactions in swine is important not only to the understanding of the utilization of Mn but also of other elements in the presence of low or high dietary levels of Mn.

The National Research Council has suggested 20 ppm Mn as the requirement for a young growing pig. Manganese requirements that have been reported in the literature vary widely. There is a definite need to define accurately the Mn requirement of the baby pig. Modern management practices, increased knowledge of other nutrient needs and ever-changing breeding programs in the swine industry all call for reevaluation of Mn requirement data. The development of more sensitive and specific analytical methods also necessitates this reexamination.

The experiments to be reported in this dissertation were designed to study the following aspects:

1. The availability of Mn from different Mn sources for the growing pig.

2. The flux pattern of Mn across the gastrointestinal tract in the pig.
3. The effect of high level supplementation of Ca and P and a Ca-P ratio less than 1.0 on Mn utilization.
4. The Mn requirement of the baby pig born of a Mn-deficient sow.

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bowed forelegs, extra sternebrae and fusion of the sternal and vertebral segments of Mn-deficient rats. Hurley *et al.* (1961a) confirmed the shortening of the long bones, both absolutely and relative to body length, and reported that the radii and tibia were greatly thickened and distorted; longitudinal growth of the skull was reduced, and the width and height of the skull were slightly less in the deficient animals on an absolute basis, but in proportion to length of the skull they were greater. They also reported that ribs were either missing or deformed and the chest was anteriorly-posteriorly flattened. Abnormal bone development due to Mn deficiency has been described in rabbits by Smith *et al.* (1944) and Ellis *et al.* (1947). Only the front legs were bowed; the ulna and the humerus were significantly reduced in size, density, Mn content and breaking strength. X-ray and microscopic studies showed changes distinctly different than those seen in rickets. The narrowing of the zone of provisional calcification and the epiphyseal plate, and a deficiency of spongy bone indicated a suppression of osteogenesis (Smith *et al.*, 1944).

Concrete evidence that Mn is involved in bone formation was obtained by Parker *et al.* (1955), who showed that a greater concentration of injected radioactive Mn was found in the regions of greater bone formation, by Leach (1967), who showed that the changes in epiphyseal cartilage of growing bone of Mn-deficient chick embryos were due to a reduction in mucopolysaccharide production, and by Tsai and Everson (1967), who reported a similar reduction in mucopolysaccharides due to a reduced hexose utilization for their synthesis in Mn-deficient guinea pigs. The conclusion by Caskey *et al.* (1939) that abnormal bone development was probably the result of disturbance in Ca and P metabolism in the absence of Mn was disproved by Parker *et al.* (1955), who showed that

the quantities of radioactive Ca and P deposited in bone were not affected by Mn intake.

Shils and McCollum (1943) reported ataxia, incoordination and poor equilibrium in young rats from Mn-deficient dams. Hurley and Everson (1959) and Hurley *et al.* (1961b) found that the offspring of Mn-deprived mothers were slower to acquire reflexes and showed inadequate development of the bony labyrinth. Hurley *et al.* (1961b) found that the brain weights of Mn-deficient rats were absolutely smaller, but relative to the body weight, larger than the controls; the cerebrospinal fluid pressure did not differ significantly. Assays of tissues for a number of enzymes, including acetylcholine esterase in the brain of Mn-deficient rats and guinea pigs did not reveal any biochemical faults (Van Reen and Pearson, 1955). Shrader and Everson (1967) and Hurley (1969) concluded that abnormal development of the otoliths was responsible for the congenital ataxia present in rats and guinea pigs when the maternal diets were Mn-deficient. Shrader and Everson (1967) reported abnormal curvatures of the semi-circular canals and misshapen ampullae in Mn-deficient guinea pigs. However, hematoxylin and eosin sections of the vestibular apparatus of ataxic rats failed to show physical lesions (Hill and Holtkamp, 1954). Erway *et al.* (1970) showed that the incidence of ataxia increased in proportion to the severity and duration of the Mn deprivation of the mother.

There is a depressed growth rate in chickens on low dietary Mn (Gallup and Norris, 1939a; Litricin, 1967). Improved growth rates due to Mn supplementation of low Mn basal diets have been reported by Settle *et al.* (1969), Belincenko (1968), and Van Reen and Pearson (1955). Since Wilgus (1936, 1937) showed that addition of 25 ppm of Mn to a basal diet containing 10 ppm of Mn completely prevented perosis,

other workers have confirmed this finding (Underwood, 1971). The occurrence of perosis on low Mn diets has been reported in ducklings (Van Reen and Pearson, 1955) and in turkey poults (Vohra and Heil, 1969). Wolbach and Hegsted (1953) found a suppression of the epiphyseal cartilage developmental sequence, with immature cartilage cells, followed by retarded tunnelling and abnormal matrix in the zone of growth. Leach (1968) reported a reduction in width of epiphyseal plate and metaphysis. Creek *et al.* (1960) reported that bone deformities of the hock joint in perosis were aggravated by weight applied to the leg.

Lyons and Insko (1937) reported that nutritional chondrodystrophy was caused by inadequate intakes of Mn, and Litricin and Andrejevic (1966) found that in Mn deficiency the long bones of the legs and wings were significantly smaller, had reduced diameter and length but did not thicken. Caskey and Norris (1940) did not find any significant effect on the sternum and metacarpus due to dietary Mn levels. Ataxia in the offspring of Mn-deficient chickens was first observed by Caskey and Norris (1940). Caskey *et al.* (1944) showed that ataxic chicks from Mn-deficient parents grew normally on Mn-supplemented diets. They described the ataxia as a tetanic spasm of the opisthotonic type. Histologically, the brain was normal, but chemical changes in lipid, total phosphorus and phosphatase content were noted.

Manganese is involved in egg shell formation. Pullets fed rations containing low Mn with high levels of Ca and P produced eggs with inferior shell characteristics (Lyons, 1939; Longstaff and Hill, 1971). Gutowska and Parkhurst (1942) showed that feeding chickens Mn-deficient diets significantly reduced the eggshell breaking strength, but Chubb (1954) found no detectable difference in eggshell quality due to Mn supplementation. Recently Hill and Mathers (1968a), Mathers *et al.*

(1971) and Longstaff and Hill (1970) have shown that eggshell thickness was depressed by a low Mn diet when given before laying but not when given from the point of lay. They also showed that low Mn diets depressed significantly the acid mucopolysaccharide content of the shell matrix.

The nutritional significance of Mn in ruminant feeding has not been clearly verified. Bentley and Phillips (1951a) showed that heifers fed diets containing 7 to 10 ppm Mn were slower to mature, and more of their calves were born with weak legs and pasterns at first calving. Grashuis *et al.* (1953) has reported leg deformities with overknuckling and poor growth in calves of cows grazing Mn depleted pastures. Rejas *et al.* (1965) reported reduced breaking strength and length of the humerus of deformed calves from Mn-deficient cows. Hartmans (1970) did not find any Mn deficiency symptoms in grazing animals in Holland. Anke and Groppe (1970) reported that there were no growth differences in female goats fed a ration containing 20 ppm Mn in the first year and 6 ppm in the second year as compared to the controls consuming a similar ration supplemented up to 100 ppm of Mn. Lassiter and Morton (1968) have reported that Mn-deficient sheep developed poor bones with low ash concentration in the femur and reduced Ca, P and Mn content in the ash, implying an effect of Mn on bone ossification.

Manganese deficiency has been shown to cause poor growth and lameness in growing swine. Miller *et al.* (1940) reported the occurrence of lameness in pigs at 27 kg when they were fed high-corn diets containing 14 ppm Mn. The condition was characterized by a slight halting gait, progressing into enlarged hocks and crooked legs, becoming painful for the pigs to rise to their feet. Bone analysis of the Mn-deficient pigs showed normal mineral deposition. Higher Mn supplementation did

not cure the stiffness but did prevent it. This work was confirmed by Keith *et al.* (1942), but in addition they showed that Mn-deficient pigs were non-ataxic and had good appetites. Sandstedt and Carlquist (1951) showed similar bone deformities, characterized by swollen streaks on the posterior external contours of the hocks in pigs fed Mn-deficient diets. Johnson (1943) obtained satisfactory growth rate and no early lameness on purified diets containing less than 0.5 ppm of Mn. However, after long periods of time on this diet some signs of lameness appeared. Leibholz *et al.* (1962) found that 0.4 ppm Mn in the diet of baby pigs was sufficient to support maximum growth. Johnson (1944) reported satisfactory growth rates from weaning to market weight on low Mn diets, and Mn supplementation was only slightly beneficial when the ash content of the ration was raised above 10 percent. Giessler and Kirchgessner (1959) found that Mn supplementation did not improve weight gains, feed intake or efficiency of conversion of Swabian-Hall pigs fattened from 20 to 96 kg on a diet of barley with 250 g of a standard German protein concentrate daily. But Williams and Noland (1949) found that Mn supplementation, along with other trace elements, improved both the growth rate and feed efficiency in swine.

Grummer *et al.* (1950) fed pigs a corn-soy basal diet containing 12 ppm Mn supplemented with 40, 80, and 160 ppm Mn. Pigs consuming diets supplemented with 40 ppm gained significantly faster than those on the basal diet but performance of the pigs was not improved with higher levels of Mn. There was a slight depression of bone ash of the pigs on the basal diets. Plumlee *et al.* (1956) showed no significant difference in growth rate and feed efficiency between groups of Duroc boars fed on diets ranging from 0.5 to 40 ppm Mn. Pigs on the low Mn diets showed a tendency towards sickle-hocks; and X-ray photographs of the legs of

Mn-deficient gilts showed that growth of the radii was prevented by closure of the distal epiphyseal plate. Plumlee *et al.* (1956), feeding gilts from Mn-deficient sows on Mn-deficient diets, observed signs of Mn deficiency manifested by pain and weakness of the legs at 27 kg liveweight, and shortened and thickened front legs becoming bowed at 45 kg. The deficient pigs were short in the body and excessively fat. Neher *et al.* (1956) reported similar defects in young pigs on purified diets of low Mn content. They showed generalized rarefaction of leg bones, but the lameness and rarefaction disappeared after 335 days on experimental diets, whereas the other signs of Mn deficiency persisted.

2. Reproduction

Kemmerer *et al.* (1931) reported that mice reared on cows' milk supplemented with Fe and Cu failed to ovulate normally, but mice on the same diet with 0.01 mg of Mn added daily exhibited normal estrous cycles. Orent and McCollum (1931) fed young rats on Mn-free diets; the females exhibited normal estrous cycles, but all their litters died, apparently due to deficient lactation. The males displayed testicular abnormalities. Boyer *et al.* (1942) reported irregular or no estrous cycles in female rats raised on low Mn diets, and there was a marked delay in the opening of the vaginal orifice; the males had testicular degeneration and complete sterility owing to lack of spermatozoa production when fed a similar diet. Shils and McCollum (1943) reported that when Mn-deficient females were mated to normal males, only 1.5 percent of the young survived to weaning as compared to 56 percent of the supplemented group. Male rats on low Mn diets from weaning did not show differences in sperm mortality and testicular weight. There were increased stillbirths and depressed survival rates when male and female Mn-deficient

albino rats were mated (Barnes *et al.*, 1941). Hurley *et al.* (1958) confirmed these effects and reported that they were aggravated in subsequent generations. Smith *et al.* (1944) and Ellis *et al.* (1947) also reported testicular degeneration and lack of libido in male rabbits on low Mn rations, and reduction in size of ovaries and uteri in females. In guinea pigs, the omission of Mn from the maternal diet resulted in a decrease in litter size and an increase in percentage of young born prematurely or ataxic or dead (Everson *et al.*, 1959).

Gallup and Norris (1939b) showed that Mn deficiency in the diet of chickens resulted in decreased egg production, fertility and hatchability. Others have reported improved reproductive performance when basal diets were supplemented with Mn (Underwood, 1971; Atkinson *et al.*, 1967). Christiansen *et al.* (1940) reported that Mn and flavin were the critical factors involved in the subnormal hatchability associated with soybeans. Hoogendorn (1940) found that with birds kept indoors, the addition of Mn to the diet improved egg production and hatchability. Gutowska and Parkhurst (1942) and Hill and Mathers (1968b) did not find any improvement in egg production, fertility and hatchability due to Mn supplementation. Chubb (1954) found that pullets fed on low Mn diets produced twice as many infertile eggs and dead-in-shell embryos as those on 50 ppm Mn diets. An increased incidence of head retractions in newly hatched chicks and chondrodystrophic embryos was associated with lines of birds producing eggs of low Mn content (Bolton, 1957).

Bentley and Phillips (1951a) showed that Mn-supplemented Holstein calves came into heat earlier than those on low Mn diets, suggesting a stimulating effect of Mn on sexual maturity; the number of services per conception and number of calves born and calves born dead were unchanged

by dietary treatments, but the calves born of Mn-supplemented cows in the second generation were noticeably heavier at birth. Experimental and field data show that Mn is necessary for fecundity in cattle (Rojas *et al.*, 1965; Bentley *et al.*, 1951). Bentley *et al.* (1951) found a significantly lower Mn content in the ovaries of repeat-breeders, but there was no direct relationship between the content of Mn in feed, tissue and organs, and infertility. Werner and Anke (1960) showed a relationship between the Mn supply and number of services per conception. Dyer (1960, 1961) and Dyer *et al.* (1964) reported a positive relationship between a low Mn intake by gestating cows and the incidence of neonatal deformities in their calves. Pastures with 15 ppm Mn or below resulted in greater incidence of infertility and abortion. Bonomi (1966) and Wilson (1966) reported that high dietary levels of CaO and P_2O_5 seemed to cause Mn deficiency, which precipitated functional infertility in cattle. Manganese supplementation has been shown to improve the fertility of cattle in Europe (Grashuis *et al.*, 1953; Wilson, 1965, 1966; Krolak, 1968). Anke and Groppe (1970) found that goats on low Mn diets came into estrus late, the symptoms of estrus were weak, and more inseminations per conception were needed. The low Mn group produced more male kids but also had more abortions. Similar observations were reported in guinea pigs by Everson *et al.* (1959).

Manganese involvement in reproductive processes of cattle was further proven by Anke and Groppe (1970), who showed a higher concentration of the element in the ovaries after an injection of radioactive Mn, and in Archibald and Lindquist (1943), who showed that there was a substantial transfer of Mn into the milk by the ovine mammary gland.

Johnson (1940) reported that bred sows fed a semi-purified diet containing 0.3 ppm Mn produced apparently normal pigs; however, the pigs

contained one tenth as much Mn as those fed 6.0 ppm Mn. Sows fed diets containing 6.0 ppm Mn raised their litters satisfactorily. Johnson (1944) showed satisfactory reproduction through two generations on a ration of 8.6 ppm Mn and 1.78 percent ash. Grummer *et al.* (1950) reported that sows on diets containing 12 ppm Mn tended to be less fertile, hard to settle and gave birth to smaller and abnormal litters. Supplementation of the basal diets with 40, 80, or 160 ppm Mn for sows during reproduction and lactation resulted in a significant difference in performance of their litters. Gligor *et al.* (1966) reported no significant effect on weight of sows at farrowing or weaning or 60 days later, or on number and weight of piglets. Speer *et al.* (1952) showed that pigs from sows supplemented with 70 to 90 ppm Mn during gestation and lactation made highest gains during the growing and fattening periods. Plumlee *et al.* (1956) fed female swine either low or high Mn diets throughout growth, gestation and lactation, and reported that the growth rate and feed efficiency of their litters were satisfactory at both levels. In another experiment, Plumlee *et al.* (1956) reported that when female pigs were fed on either low or high Mn diets, the differences in number of pigs farrowed or viability or birth weights of litters could not be attributed to Mn levels given. But when gilts were depleted of Mn by feeding Mn-free purified diets, the reproductive organs were histologically normal, and they ovulated normally but showed irregular estrus and sometimes would not accept the boar. When mated, they farrowed weak ataxic litters, unable to suck, and most of them died even when transferred to Mn-supplemented diets. The litters from supplemented gilts were normal, vigorous and healthy. Boars from gilts getting either low or high Mn levels in their diets, all reared on 1.3 ppm Mn ration, performed equally well and showed normal

spermatogenesis (Plumlee *et al.*, 1956). Neher *et al.* (1956) reported that sows on a low Mn diet farrowed abnormal pigs with reduced birthweights. The above findings were not sustained by the results of Leibholz *et al.* (1962) and Newland and Davis (1961), who found, independently, that sows on low Mn diets farrowed or produced normal fetuses without apparent reduction in birthweights. The involvement of Mn in swine reproduction is further shown by the findings that there is a rapid, unlimited transfer of the element to the fetus, colostrum and milk (Plumlee *et al.*, 1956; Newland and Davis, 1961; Leibholz *et al.*, 1962).

B. Biological Roles of Mn

1. Enzyme Activation

Lehninger (1970) lists many enzymes involved in intermediary metabolism that require Mn ions for activation. Mn is an integral part of many enzyme systems of the body. Some of the specific enzymes for which Mn is known to be essential as an activator are acetyl CoA carboxylase (Wells and Remy, 1965), cytochrome oxidase (Vorob'eva, 1970), enolase (Wacker *et al.*, 1964; Babin *et al.*, 1964), fructose 1,6 diphosphatase (Pontromoli *et al.*, 1969), heparinase (Dietrich *et al.*, 1972) and succinic dehydrogenase (Babin *et al.*, 1964). Other divalent cations, especially Mg, with similar properties, can replace Mn in the activation of many enzyme systems.

Rubenstein *et al.* (1962) showed that high levels of Mn induced severe hypoglycemia due to increased enzyme activation, and Johnson *et al.* (1959) reported that preincubation with Mn ions increased considerably the proteolytic activity of rat and chick pancreatic homogenates. Chatterjee *et al.* (1960) found that Mn ions restored the conversion of

gluconate to ascorbic acid by goat liver microsomal enzymes. Skinner and McHargue (1944) found that Mn supplementation of rats increased co-carboxylase activity. Reineke and Turner (1945) showed that enzymes involved in the iodination of tyrosine to diiodotyrosin and subsequent oxidation to thyroxine were best catalyzed by Mn, especially colloidal MnO_2 . Van Reen and Pearson (1955), studying a number of enzymes in Mn-deficient and Mn-supplemented ducks, showed that Mn had no effect on the activity of liver diphospho-pyridine nucleotidase, cytochrome oxidase, catalase and isocitrate dehydrogenase. Leach (1967) found that the enzymes, polymerase and galactotransferase, involved in the chondroitin sulfate synthetic system are activated by Mn.

In vitro studies indicated that Mn ions activated phosphatase, but addition of Mn ions to an enzyme preparation from the bones and blood of a perotic chick did not raise the activity to nearly the same level as that found for a chick without perosis (Weise *et al.*, 1939). Wachtel *et al.* (1943) found that Mn deficiency caused a significant increase in blood serum phosphatase but not bone phosphatase. But Ellis *et al.* (1947), Leibholz *et al.* (1962) and Rojas *et al.* (1965) showed a depression of bone alkaline phosphatase activity on low Mn diets. Combs *et al.* (1942) found an intimate relationship between the phosphatase activity of the bones and Mn deficiency. Lowering of the phosphatase activity retarded bone development. Phosphatase activity was greatly reduced by withdrawing Mn. Lassiter *et al.* (1970) reported that subnormal bone alkaline phosphatase activity does not invariably occur with Mn deficiency. Hurley and Everson (1959) showed that both Mn-deficient rats and controls showed the same pattern of enzyme activity and there were no significant differences between them. Nielsen and Madsen (1942) showed that blood phosphatase significantly increased in perotic

turkeys, but Van Reen and Pearson (1959) found that the enzyme activity of livers of Mn-deficient ducks was only one-half that of the livers of supplemented birds. Lassiter *et al.* (1970) found that serum alkaline phosphatase activity of lambs fed 1 ppm Mn was significantly below that of controls receiving 29 ppm Mn, but kidney alkaline phosphatase activity was increased. Leibholz *et al.* (1962) showed that the alkaline phosphatase activities of the kidney, liver and serum of baby pigs given 0.4 ppm Mn in the diet were not affected.

Boyer *et al.* (1942), Rehner and Stelte (1970) and Rehner and Cremer (1970) reported the arginase activity of liver preparations from Mn-deficient animals was greatly increased by Mn additions. Mn supplementation increased urea in blood, urine and saliva of cattle (Zerebeov *et al.*, 1970; Rozybakiev, 1966). Others have shown a depression in arginase activity of rats and guinea pigs on Mn-deficient diets (Ellis *et al.*, 1947; Everson, 1970). But Leibholz *et al.* (1962) found that Mn treatments did not significantly affect the liver and kidney arginase activity in baby pigs.

2. Energy Metabolism

Lehninger (1970) lists a number of key enzymes involved in the glycolytic pathway, gluconeogenesis and beta-oxidation requiring Mn ions for activation. Early investigations by Ray and Deysach (1942) showed that subcutaneous injection of Mn into guinea pigs in small doses raised oxygen consumption, but higher doses up to 100 mg per kg of bodyweight progressively depressed oxygen consumption. Wachtel *et al.* (1943) found no Mn effect on basal metabolic rate (BMR) in rats.

Bentley *et al.* (1951b) found that phosphorylation of chicken liver homogenates was increased 41 percent by Mn above that on a choline-

deficient, Mn-deficient ration. Hurley *et al.* (1970) showed a depression in oxidative phosphorylation in Mn-deficient mice. Buccellato (1953) found *in vitro* that a compound formed between pyridoxine and colloidal Mn had an active role in carbohydrate metabolism. Scrutton *et al.* (1966) found that pyruvate carboxylase is a Mn metalloprotein, and Mildvan *et al.* (1966) showed that Mn functions in the transcarboxylation part of the pyruvate carboxylase reaction.

Manganese involvement in glucose utilization was shown by Everson and Shrader (1968) and Shrader and Everson (1968). Newborn guinea pigs, severely deficient in Mn, exhibited a marked hypoplasia of all cellular components of the pancreas, with fewer and less intensely granulated beta cells than the controls. When glucose was administered orally or intravenously to young adults which were congenitally Mn-deficient, they showed glucose responses resembling the diabetic subject, whereas control animals always returned promptly to normal glucose levels. Mn supplementation of deficient animals completely reversed the reduced glucose utilization in guinea pigs and cattle (Everson and Shrader, 1968; Zerebeov *et al.*, 1970). The administration of this element to diabetic subjects has a hypoglycemic effect (Mehrolera *et al.*, 1964; Rubenstein, 1962); and both pancreatectomy and diabetes have been correlated with decreased Mn levels in blood and tissues (Konseko, 1965).

Manganese supplementation was shown to reduce liver and bone fat in Mn-deficient rats (Amdur *et al.*, 1946). Plumlee *et al.* (1956) showed the Mn-deficient gilts were excessively fat by 25 to 40 kg liveweight. Curran (1954) showed that Mn stimulates the hepatic synthesis of cholesterol and fatty acids in rats. Mn ions are necessary for the conversion of mevalonic acid to squalene by mevalonic kinase (Amdur

et al., 1957); and the phosphorylated derivative of mevalonic acid, necessary for this reaction, requires Mn for its synthesis. Barron (1966) showed that Mn was a necessary co-factor for mitochondrial fatty acid synthesis together with NAD and citrate; and others have reported that Mn inhibits lipoamide dehydrogenase (Lehninger, 1970).

3. Hemoglobin Formation

Manganese has been shown to either increase or decrease or have no effect on hemoglobin values in a variety of animal species. Wachtel *et al.* (1943) and Smith *et al.* (1944) found that hemoglobin levels were not significantly affected by lack of Mn. But Skinner and McHargue (1946a), using dry or milk diets, showed that Fe, Cu and Mn gave higher hemoglobin values than Fe and Cu added alone.

High levels of Mn (100 to 3000 ppm) had a small but significant depression on hemoglobin levels of calves (Cunningham *et al.*, 1966). Hemoglobin regeneration was greatly retarded and serum iron depressed in anemic and normal lambs, rabbits and pigs by feeding high levels of Mn. The levels causing reduction of hemoglobin ranged from 50 to 5000 ppm (Robinson *et al.*, 1960; Matrone *et al.*, 1959; Hartman *et al.*, 1955). With baby pigs, Matrone *et al.* (1959) showed that the regeneration of hemoglobin, when Fe intake was low, was depressed by excess Mn intakes but the depression was overcome by extra Fe. When the anemic baby pigs, given 25 ppm Fe and 5 ppm Cu in the diet, were supplemented with 125, 250 or 2000 ppm Mn, all the levels of Mn depressed hemoglobin formation and the drop was sharp and significant. Moinuddin and Lee (1960) found a decline in hemoglobin concentration, a reduction in red blood cell count and an increase in white blood cell count due to feeding high Mn levels. In 1960, Sullivan noted similar changes in rats

given a manganese edetate supplement. The changes were greater in young rats.

4. Ascorbic Acid Synthesis

Rudra (1939) found that rat and guinea pig livers were able to synthesize ascorbic acid when incubated with mannose or galactose in the presence of Mn which acts as a co-enzyme in the conversion. Later he showed that intraperitoneal injection of 20 mg of mannose in the presence of 0.04 percent Mn to guinea pigs gave a small increase of ascorbic acid in body tissues and protected against scurvy (Rudra, 1944b). He concluded that Mn was essential for the synthesis of vitamin C in animals and that failure to synthesize it is due to insufficiency of the metal at the site of ascorbic acid synthesis.

Injections of mannose plus Mn given to scorbutic guinea pigs did not stimulate ascorbic acid synthesis from mannose *in vivo* (Skinner and McHargue, 1946b). The Mn involvement in ascorbic acid synthesis is complicated by vitamin E (Caputto *et al.*, 1958). Using enzyme preparations from vitamin E-deficient rats, 70 to 90 percent less ascorbic acid was produced than from a preparation taken from animals given sufficient vitamin E, and addition of Mn to *in vitro* systems increased the ascorbic acid produced by vitamin E-deficient preparation 315 percent.

5. Genetics, Disease and Immunity

A difference in Mn requirement to prevent perosis among various breeds of birds has been documented (Golding *et al.*, 1940; Pilla, 1958). Caskey *et al.* (1944) reported that offspring of ataxic female and male chickens grew normally on a diet supplemented with Mn, suggesting the ataxia was not complicated by the inheritance of a simple

recessive character influencing Mn retention. Hurley (1969) showed that ataxia caused by maternal Mn deficiency is indistinguishable from that caused by certain genes in mutant mice. A supplement of 1000 ppm Mn to pregnant females on a diet that normally produced 68 percent ataxic young completely prevented the condition and, when the normal offspring were mated and fed a normal diet, the normal offspring subsequently produced 68 percent ataxic young. Thus, the high level of Mn prevented expression of the genetic abnormality without influencing genetic constitution.

Hoogendoorn (1940) reported that adding 12 mg of manganese sulfate to 100 kg of poultry meal afforded additional resistance to disease. Marot and Durand (1944) found that the Mn content of malignant and benign tumors was much lower than that of healthy tissues. Sandstedt *et al.* (1951) showed that daily supplementation with 0.5 g of manganese sulfate gave rapid recovery from acetonemia in cattle, and preventive treatment over the years greatly reduced the incidence of the disease. Samofal (1961) found that the level of Mn in the diet was a decisive factor in the occurrence of goiter. Kamchatrov (1959) and Hakimova *et al.* (1969) showed that excess dietary Mn affects thyroid metabolism. Kaellis (1970) showed a depression of the protein-bound iodine fraction. Earlier Kamchatnov (1953) had noted a relationship between Mn content of feed items and the distribution of enzootic and non-enzootic goiter regions. Angelico *et al.* (1965) and Antanova (1968) have reported that Mn supplementation significantly delayed the death of rats and increased the survival rate after nitrogen mustard poisoning and coliform bacterial infection.

Weinberg (1964) discovered that certain bacteria had specific requirements for Mn, in excess of that needed for growth, in order to

produce antibiotic, bacteriophage, and protective antigens. Antanova (1968) and Antanova *et al.* (1968) reported that agglutinin response and phagocytic activity were greater with higher Mn intakes in rabbits when immunized with coliform and typhoid bacteria. Muraleedharan and Pande (1968) found that with Mn-deficient diets, infection with *Prosthogonimus ovatus* seemed to hasten death.

C. Mn Metabolism

1. Absorption

Little is known about the mechanism of Mn absorption from the gastrointestinal tract. Scott *et al.* (1958) showed that a Mn compound capable of dissolving and being converted to manganese chloride in the acid medium of the gastrointestinal tract can be absorbed.

The uptake of Mn by the intestinal mucosa is very rapid (Miller *et al.*, 1972), and the element is bound to the serum beta-globulin fraction in all species studied (Panic and Ekman, 1967). While Saltman *et al.* (1956) thought that simple diffusion constituted the driving force for the transport of Mn, Rothstein *et al.* (1958) and Weed and Rothstein (1958) presented evidence for active transport for Mn. Gutowska *et al.* (1941) showed that the amount of Mn absorbed from a solution of manganese sulfate was proportional to its concentration; an average of a third of the amount injected in the intestines was absorbed in two hours. There was no significant sex difference in absorption of Mn in chickens. Many reports have indicated that only 3 to 4 percent of an oral dose of Mn is absorbed (Britton and Cotzias, 1966; Watson *et al.*, 1973). But Gamble *et al.* (1971) and Brown and McCracken (1965) have reported substantial Mn absorption. Pregnant sows absorbed up to 28 percent of ingested Mn (Gamble *et al.*, 1971).

Manganese absorption is affected by various factors present in the diet. Pollack *et al.* (1967) and Hill and Holtkamp (1954) reported that Mn was more readily absorbed at a lower than at a higher dietary Mn concentration. Cotzias and Greenough (1958) and Zajcev (1959) found that Mn absorption was not affected by its valency state in the compounds used. Many workers have reported that high dietary levels of Ca and P aggravated Mn deficiency due to reduction in Mn absorption (Underwood, 1971). Intestinal absorption of Mn was increased in rats made iron deficient (Pollack *et al.*, 1965). Saltman *et al.* (1956) found that Mn competitively inhibited Fe uptake and release by liver slices, which indicated that both Fe and Mn have a common pathway. Suso and Edwards (1968) reported that Mn absorption was enhanced by diethylenetriamine pentaacetic acid (DTPA) and decreased by ethylenediamine tetraacetate (EDTA). Leibholz *et al.* (1962) found that pigs fed casein diets containing 0.4 and 4(4 ppm Mn grew at a more rapid rate on less food per pound of gain than did pigs fed soybean protein rations containing 11.8 and 51.8 ppm Mn. Davis *et al.* (1962) found a factor in soybeans which tends to bind Mn and makes Mn unavailable.

Settle *et al.* (1969) found no appreciable binding of Mn in feather meal diets. Gilbert (1957) reported that thiamine, given in excess, precipitated a Mn deficiency, but Holtkamp *et al.* (1950) found no evidence of antagonism between dietary Mn and thiamine.

2. Excretion

Everson and Daniels (1934) observed that total Mn urinary excretion is virtually constant, irrespective of age, and that fecal Mn excretion varies directly with age and therefore with total dietary Mn intake. Many workers have reported that most of Mn administered

orally or intraperitoneally quickly appeared in bile and was excreted in the feces, and very little was excreted in the urine (Kent and McCance, 1941; Mahoney and Small, 1968; Starodubova, 1968). The predominance of the fecal route for Mn excretion has been verified in simple stomached animals (Britton and Cotzias, 1966; Miller, 1973), in sheep (Watson *et al.*, 1973), in cattle (Miller *et al.*, 1973), and in man (North *et al.*, 1960; Cotzias and Greenbough, 1958).

Under normal conditions, the bile flow is the principal regulatory mechanism of Mn excretion, and the concentration of Mn in bile can be increased tenfold or more by the animal (Underwood, 1971). The other routes of Mn excretion include pancreatic juice (Papavasiliou *et al.*, 1966; Burnett *et al.*, 1952) and secretions of the duodenum, jejunum and, to a smaller extent, the terminal ileum (Bertinchamps *et al.*, 1966). Excretion via the kidney is negligible normally or during jaundice or after a marked oral dose. The administration of chelating agents such as EDTA produces a marked rise in urinary excretion of Mn (Maynard and Fink, 1956).

Lassiter *et al.* (1970) reported that the body pool is small and the body does not ordinarily accumulate Mn. Total body excretion is continuous and very nearly equal to intake, and much of the Mn in the body pool is replenished daily. Animals placed on a low Mn diet continue to excrete Mn, suggesting an obligatory loss of Mn (Zajcev, 1959; Starodubova, 1968). The excretion of Mn administered parenterally was much lower on a 1.0 percent Ca diet than on a 0.6 percent Ca diet, but raising P levels had no comparable effect on the excretion or retention of Mn (Lassiter *et al.*, 1970). Underwood (1971) noted that Ca influences Mn metabolism by affecting its absorption. The normal excretion of Mn from the body is prevented by rectal and biliary ligation (Papavasiliou

et al., 1966), and there is no appreciable interdependence of these routes of excretion (Bertinchamps *et al.*, 1966).

3. Retention

Everson and Daniels (1934) reported that Mn retention varies inversely with age of children and therefore intake. North *et al.* (1960) showed that college women retained about a third of the absorbed Mn, and pullets retained about the same amounts (Brown and McCracken, 1965). Gamble *et al.* (1971) showed that 7 days after the administration of a radioactive dose of Mn, pregnant sows retained 26 percent of the oral and 78 percent of an intravenous dose. A highly significant correlation between intake and retention has been found (Mathers and Hill, 1967; Murty, 1957), but Hughes *et al.* (1966) reported that only a small change in tissue Mn level can be effected by a large change in dietary Mn intake. There is a linear relationship between Mn turnover and level of dietary Mn intake, and the half-life of body Mn is decreased with increasing Mn intake (Britton and Cotzias, 1966). The absorbed Mn is found principally in the liver and bone. At lower Mn intakes the liver retains more Mn than intestinal tissue, but at high intakes the latter retains more (Underwood, 1971). Passive transendothelial transport occurs immediately after intravenous administration of Mn, and about 70 percent of the blood Mn leaves the circulation each minute and is mainly taken up by the liver (Cotzias, 1958; Thomas, 1970). Borg and Cotzias (1958) showed that most of the endogenous Mn exists in highly labile intracellular combinations.

Tissue Mn concentrations generally are remarkably constant even though consumption levels vary greatly (Underwood, 1971). These might be regulated through variable excretion rates (Britton and Cotzias,

1966), and absorption differences (Howes and Dyer, 1971; Miller, 1973). Lassiter *et al.* (1970) showed that 0.9 percent dietary P caused significantly higher Mn retention of orally administered Mn than did 0.4 percent P, but there were no comparable effects on intraperitoneally administered Mn. Hughes and Cotzias (1961) found that administration of exogenous glucocorticoid hormone markedly affected the tissue distribution of radioactive Mn, but adrenalectomy did not affect retention (Hughes *et al.*, 1966). Hill (1967) found that vitamin D reduced Mn turnover, and Suso and Edwards (1968, reported that EDTA reduced the retention of intravenously injected Mn, and greatly increased Mn transport (Sahagian *et al.*, 1967). Gamble *et al.* (1971) reported that pregnancy in swine had no significant effect upon maternal tissue retention, organ distribution or turnover rate of Mn. Fournier *et al.* (1972) reported increased Mn retention in all organs as a result of lactose ingestion.

D. Manganese Requirement

1. Factors Affecting Mn Requirement

There is a very wide margin of safety between the minimum and toxic levels of Mn for all species. The minimum dietary requirements of Mn depend upon the species, the criteria of adequacy employed, the chemical form in which the element is ingested, and the nature of the rest of the diet (Underwood, 1971).

Manganese requirements differ between and within species. Chickens require more Mn than any other species (Thomas, 1970). A level of 13 ppm Mn in the diet resulted in signs of perosis in White Leghorns and New Hampshires but not in Rhode Island Reds (Pilla, 1958). Chondrodystrophy was confined to Barred Rock chicks only and did not affect

White Leghorns, and the response to Mn supplementation was greatest in New Hampshires and least in Leghorns (Golding *et al.*, 1940). Gutowska *et al.* (1941) and Mathers and Hill (1967) reported no significant sex difference in Mn utilization in chickens. But Barnes *et al.* (1941) reported that female chicks were more sensitive to Mn deficiency in the mother's diet than male chicks. And Pilla (1958) found that the response to Mn was greater in males than females. There is evidence showing that Mn levels necessary for growth are less for reproduction (Rojas *et al.*, 1965; Grummer *et al.*, 1950; Bentley *et al.*, 1951a).

In attempting to define dietary requirements of Mn, the composition of the diet is crucial. There are interactions that occur in the food or in the gastrointestinal lumen. Manganese interacts with Ca and P within the digestive tract. In 1939, Caskey and Norris observed that Mn was made unavailable by high levels of Ca and P in the diets of chickens. High dietary levels of Ca and P have been shown to decrease the growth rate and increase the incidence of perosis in chickens on diets having Mn levels ordinarily adequate to prevent perosis, and additional Mn prevented the development of perosis (Underwood, 1971). Wilgus and Patton (1939) explained the perosis-producing action of calcium phosphate as being due, at least in large part, to the removal of Mn from solution in the intestinal tract by adsorption or chemical combination. High levels of ferrous citrate also increased the incidence of perosis, presumably through similar intraluminal action (Wilgus and Patton, 1939). Addition of ferric oxide to a Mn and Cu mixture seemed to offset its beneficial effect on hatchability (Hoodengoorn, 1940). In intracellular fractions, the concentration of Mn and Fe have been shown to have a reciprocal relationship (Thiers and Vallee, 1957). There is some evidence of dietary Mg-Mn and Zn-Mn interactions

(Blakemore *et al.*, 1937; Cotzias, 1960; Sahagian *et al.*, 1966). Diez-Ewald *et al.* (1968) and Kolomijceva and Veznesenskaja (1968) have reported interrelationships between Fe, Cu and Mn metabolism.

In some other circumstances, high levels of dietary thiamine intake increase body Mn storage (Hill and Mathers, 1954), but Sandberg *et al.* (1939) found that the state of thiamine deficiency caused great increases in Mn retention. Anderson and Parker (1955) found no thiamine level effect on liver and heart Mn content. Holtkamp *et al.* (1950) found no evidence of antagonism between thiamine and Mn. Perla and Sandberg (1939) reported that high thiamine levels causing low reproduction rates can be counteracted by increasing Mn intakes. Riboflavin, at high levels, was shown to complicate perosis in ducklings (Turton, 1953).

The protein source of the diet may affect Mn metabolism. The availability of Mn to chicks was better when either fishmeal or dried skim milk was the protein source as compared to soybean meal (Morimoto *et al.*, 1959; Kealy and Sullivan, 1966). Davis *et al.* (1962) showed that soybean protein contains a component which combines with Mn leading to conditioned deficiency. Settle *et al.* (1969) found no such binding of Mn when feathermeal was fed as the source of protein.

The Mn compound fed may determine Mn availability and requirements. Bandemer *et al.* (1940) showed that precipitated $MnCO_3$ protected against perosis at a lower level than the naturally occurring carbonates which proved useless even when used in large amounts due to low solubility. Schaible and Bandemer (1942) reported that chemical forms of Mn compounds as diverse as carbonates, oxides, sulfates and chlorides were equally valuable as sources of Mn in poultry rations. Anke *et al.* (1967) and Watson *et al.* (1970) have shown that the chloride and sulfate are better utilized than the oxides in ^{chickens} cattle. Lyons (1939) and

Quereshi *et al.* (1963) found that Mn supplied in rice bran appeared to be as well utilized as the inorganic sulfate. The method of administration of Mn is important in determining its effectiveness. Caskey and Norris (1939) showed that small amounts of Mn injected intraperitoneally were more effective than ingested materials against perosis.

A number of feed additives have been shown to either depress or enhance Mn availability to the animal. The inclusion of chlortetracycline in low Mn diets for chickens reduced the incidence of perosis in the pullets but not in their progeny (Pepper *et al.*, 1952, 1953). Bolton (1955) showed that giving estradiol increased plasma and liver Mn. Hart (1953, 1954) reported that giving Vevoron, an antithyroid preparation containing methylthiouracil for fattening, significantly depressed the Mn content of the liver. Hydrazine administration can cause symptoms in animals similar to those of Mn deficiency (Thomas, 1970).

There are other factors which may influence Mn requirements. Christiansen *et al.* (1939) concluded that sunlight exerts a sparing action on the hens' Mn requirements. Urban (1959) showed that hepatectomy aggravated Mn deficiency.

2. Mn Requirement of Swine

There are conflicting data in the literature on the exact Mn requirements of pigs for proper growth, skeletal development and reproduction. Keith *et al.* (1942), using a high corn ration fortified with minerals and containing 11 to 14 ppm Mn, found that the growth of pigs was not impaired but skeletal development was poor, and a supplemental level of 50 ppm Mn prevented skeletal deformities but did not cure them. Miller *et al.* (1940) had a similar response with 60 ppm supplementation.

Johnson (1940) reported that 6 ppm was sufficient for successful reproduction of sows; and satisfactory growth was obtained on 0.3 ppm Mn, but reproduction was unsatisfactory and, at this level, tissue Mn content was significantly depressed. Johnson (1944) showed that pigs grew well from weanling to market weight on rations containing 7 to 10 ppm Mn. Grummer *et al.* (1950) reported that a diet containing 12 ppm Mn was adequate for skeletal development and growth but not adequate for optimum reproductive performance. When the same diet was supplemented with 40, 80 and 160 ppm Mn, the highest average daily gains were obtained on the 40 ppm level. Higher supplementation had no added benefit. Plumlee *et al.* (1956) reported that dietary Mn concentrations ranging from 0.5 to 34 ppm Mn did not show significant differences in pig performance. Speer *et al.* (1952) showed that pigs from sows supplemented with 70 to 90 ppm Mn performed best during growing and fattening as compared to those from unsupplemented groups. Leibholz *et al.* (1962) defined the baby pig Mn requirement at 0.4 ppm Mn for maximal growth rate. Leibholz *et al.* (1962) and Newland *et al.* (1961) found independently that sows farrowed or produced normal fetuses when fed 89 to 117 ppm Mn or 6 to 100 ppm Mn in their diets, respectively. The Mn requirements for growth of pigs are extremely low, well below the levels ordinarily found in practical swine diets, although the National Research Council (NRC, 1973) recommended 20 ppm Mn in the diet.

3. Mn Requirement of Other Species

The Mn requirements of many species have been estimated by many investigators and reviewed by Thomas (1970) and Underwood (1971). Birds have higher Mn requirements than mammals. The best evidence indicates that to prevent deformities in the fetus, cows should receive

a diet containing 20 ppm Mn. In calves 8.6 mg per day were not sufficient but 36 mg per day were optimum for growth. Poultry requirements have been put at 40 to 70 ppm Mn in the diet. Children require at least 1.2 mg per day; and older humans 3 to 5 mg. Requirements for the rat have been set at 0.5 to 1.0 mg per day, and the low amount of 1 ppm in the diet produces fetal abnormalities. Rabbits require about 1 mg per day, and sheep 50 to 60 ppm on a feed dry matter basis.

E. Mn Toxicity

Mn is one of the least toxic of the trace elements to mammals and birds. Richards (1930) fed pigs 3.5 g of manganese citrate daily for nine months without any adverse effects. Mussehl and Ackerson (1939) showed that turkeys can tolerate 385 ppm Mn in their diets and Insko *et al.* (1938) reported that hens tolerated 600 to 1000 ppm Mn. However, others have reported adverse effects due to high level feeding of Mn. Kamimura (1938) found that rabbits fed 0.5 to 6 g per kg bodyweight daily were stunted and their bone development impeded. Similar effects have been shown in rats and cattle (Chornock *et al.*, 1942; Cunningham *et al.*, 1966). Wessinger *et al.* (1943) found that injections of solutions containing 180 to 975 ppm Mn as $MnCl_2$ into white rats caused enamel hypoplasia, interrupting amelogenesis in the apical quarter of the zone of matrix formation. Heller and Penquite (1937), feeding a ration containing 4800 ppm Mn, showed the element to be highly toxic to young chickens.

High level feeding of Mn interferes with Fe, Cu and P metabolism (Underwood, 1971). Urinary Cu excretion is depressed and Cu retention in the tissues is increased, causing a microcytic, hypochromic anemia in rats fed very high Mn levels (Gubler *et al.*, 1954). Earlier, Edgar

(1942) supplemented sheep rations with 25 mg of Mn as the sulfate and 25 mg of Cu as the sulfate daily and showed large but non-significant increases in liver Cu levels over those of the controls receiving copper sulfate alone. Experimentally, dietary Mn levels causing a suppression of hemoglobin formation are 1000 to 2000 ppm for anemic lambs (Hartman *et al.*, 1955), 5000 ppm for normal lambs (Robinson *et al.*, 1960), 1250 ppm for mature rabbits (Matrone *et al.*, 1959), and 2000 ppm for baby pigs (Matrone *et al.*, 1959). Hartman *et al.* (1955) and Cunningham *et al.* (1966) reported that high levels of dietary Mn resulted in decreased concentrations of Fe in liver, kidney and spleen of ruminants, and depressed hemoglobin formation.

Blakemore *et al.* (1937) found that the Mn content of pastures in districts where lactation tetany was prevalent was 700 ppm on a dry matter basis as compared to 50 ppm of pastures on which tetany had never been recorded. The feeding of Mn to rabbits, sheep and cows in amounts supplied in the pastures on which tetany occurred brought about transitory falls in levels of Mg in blood. Robinson *et al.* (1960) and Cunningham *et al.* (1966) found that cattle fed high Mn levels produced less rumen propionic acid. Keith *et al.* (1942) showed that growing pigs fed on diets containing 2000 ppm Mn grew poorly, lost weight and appetite. They vomited and had nausea, diarrhea and dermatitis. Grummer *et al.* (1950) showed that pigs do not tolerate high levels of dietary Mn since 500 ppm in the diet reduced growth rate, feed efficiency and depressed appetite of growing and finishing swine. But Leibholz *et al.* (1962) showed no toxicity signs in baby pigs fed 4000 ppm Mn; however, there was evidence of reduced growth rate at 4000 ppm. This work showed a high tolerance by the baby pig and a considerable margin of safety between levels of Mn likely to be

ingested in the diet and detrimental levels. In man the contamination of air and water by large amounts of Mn causes *locula manganica*, disturbance of the extrapyramidal tract and atrophy and disappearance of nerve cells of the globus pallidus (Cotzias, 1958; Belani *et al.*, 1967). Cotzias (1968) and Mena *et al.* (1968) showed a decreased turnover of Mn in patients suffering from chronic Mn poisoning.

EXPERIMENTAL PROCEDURES

A. Introduction

Four experiments involving 104 pigs were conducted to study Mn metabolism in swine. These were:

Experiment 1. The relative availability of Mn from a 16% corn-soy basal diet and the basal diet supplemented with 10 ppm of Mn from $\text{MnSO}_4 \cdot \text{H}_2\text{O}$, MnCO_3 or MnO for the growing pig.

Experiment 2. Study of the gastrointestinal flux pattern of Mn from different Mn sources using chromic oxide (Cr_2O_3) as an indicator.

Experiment 3. The effect of high level Ca and P supplementation and an inverse Ca-P ratio on Mn utilization by the growing pig.

Experiment 4. The Mn requirement of the baby pig from sows fed a low Mn diet.

Yorkshire, Hampshire and Yorkshire-Hampshire crossbred pigs from the Michigan State University herd were used. All trials were conducted at the university swine farm facilities.

B. Experiments

1. Experiment 1

Before the diets for this experiment were made, three Mn compounds (manganous sulfate, manganous carbonate and manganous oxide) were subjected to a 0.4% HCl availability study.

A liter of 0.4% HCl was heated in a 1500 ml beaker to a constant temperature (37°C) in a gyrotory water bath shaker, Model G76.¹ One gram of each compound, ground to pass through a 100 mesh screen, was added; the temperature and agitation were maintained for one hour. The resulting solution was filtered at once through a dry Whatman #42 filter paper, discarding approximately the first 50 ml. The filtrate was thoroughly shaken, sub-sampled into three 20 ml aliquots, and Mn was determined by atomic absorption spectroscopy.

Twelve crossbred weanling pigs, 9 males and 3 females, from two litters, were allotted to the four diets shown in Table 1, equalizing for sex, litter and weight. The supplemented diets were made using the same compounds used in the above study. The basal diet was supplemented with 10 ppm of Mn from $\text{MnSO}_4 \cdot \text{H}_2\text{O}$, MnCO_3 or MnO to make diets 2, 3, and 4, respectively.

The pigs weighed about 8 to 10 kg initially. They were housed in individual stainless steel metabolism cages for a 7-day adjustment period followed by two balance trials. The first was a replicated Latin square design in which all animals were fed all the diets during four collection periods. The second was a split-plot design in which three repeated collections were made on the animals maintained on the same diets. The pigs were removed from the cages three times daily and individually fed an amount of food and water which could be consumed within a 5- to 10-minute period. The feed was mixed with water to make a slurry for quick consumption. Following feeding, the pigs' mouths were wiped clean to avoid contamination of excreta. The pigs were then returned to the cages.

¹New Brunswick Scientific, New Brunswick, N.J.

TABLE 1. DIETS USED IN EXPERIMENT 1

Ingredient	Diet			
	Basal	+ $\text{MnSO}_4 \cdot \text{H}_2\text{O}$	+ MnCO_3	+ MnO
Corn, shelled, ground	79.2	78.2	78.2	78.2
Soybean meal, dehulled solvent (49% CP)	17.9	17.9	17.9	17.9
Salt	0.5	0.5	0.5	0.5
Limestone, grd (38% Ca)	0.9	0.9	0.9	0.9
Dicalcium phosphate	1.0	1.0	1.0	1.0
VTM premix ¹	0.5	0.5	0.5	0.5
$\text{MnSO}_4 \cdot \text{H}_2\text{O}$ premix		1.0		
MnCO_3 premix			1.0	
MnO premix				1.0
	100	100	100	100
Level of CP (calculated), %	16	16	16	16
Level of Mn (analyzed), ppm	16.2	26.0	24.8	27.0

¹See Appendix A, Table A-1.

Fecal and urine collections were made over a 3-day period, with 3-day intervals in between each collection. Feces were separated from urine by means of a fine screen placed over the urine collection funnel. The pigs consumed near *ad libitum* quantities of feed daily, and constant daily feed intakes were maintained throughout the balance periods. Unconsumed feed was collected after each feeding during the 3-day collection period to get an accurate measure of feed intake during the balance trial.

Feces were oven-dried for 24 hours, equilibrated to room temperature for 12 hours, weighed, ground and stored in sealed plastic containers. Refused feed was air-dried, weighed and discarded. Urine was collected in polyethylene containers and acidified with 6N HCl. Following the collection period, the urine volume was recorded and 100 ml aliquots were stored in acid-washed polyethylene bottles at 4°C.

The animals were bled weekly from the anterior vena cava before the start of each collection period for the determination of serum alkaline phosphatase and serum Mn concentration. All the animals were killed after the second balance trial. The following tissues were collected, weighed and stored in a freezer in polyethylene bags: liver, kidneys, spleen, testes, heart, left femur, 8th left rib, pancreas and ham muscle from the left leg. Hair samples were collected from the animals at the beginning and at the end of the experiment from the loin region of each animal.

2. Experiment 2

The gastrointestinal flux pattern of Mn from different Mn sources. Chromic oxide was added (at 0.3%) to each of the diets used in Experiment 1. Pigs were fed these diets for a period of 4 days before they were

killed. Feces voided were weighed and grab samples of the feces were taken for further analyses.

Pigs were killed one and one-half to three hours postfeeding. The alimentary tract was quickly exposed and sectioned into the stomach, cranial small intestine, caudal small intestine, cecum, colon and rectum. The sections were ligated to prevent movement of the contents. Digesta were removed quantitatively from each section, weighed, and the pH determined on a small portion. The remainder was thoroughly mixed and divided into 20 g portions. These were placed in polyethylene bags and frozen for later analysis.

The indicator method for determining digestibility was used to estimate absorption and secretion of Mn along the gastrointestinal tract. The following equation was used:

$$\% \text{ digestibility} = 100 - \left(\frac{\% \text{ indicator in feed}}{\% \text{ indicator in feces}} \times \frac{\% \text{ nutrient in feces}}{\% \text{ nutrient in feed}} \right) 100$$

(Crampton and Harris, 1969). In order to measure the net absorption and secretion in the different sections of the alimentary tract as the digesta moved posteriorly, two consecutive sections of the tract were used in relation to the above equation. The digesta in a given section was considered to be the feed for the next posterior section. The digesta in the second section would be equivalent to the feces for the calculation using the ratio technique equation. Thus net absorption or secretion was calculated in the following alimentary sections as indicated:

1. Stomach - using feed as feed and stomach digesta as feces.
2. Cranial small intestine - using stomach digesta as feed and cranial small intestine digesta as feces.

3. Caudal small intestine - using cranial small intestine digesta as feed and caudal small intestine digesta as feces.
4. Cecum - using caudal small intestine digesta as feed and cecal digesta as feces.
5. Colon - using cecal digesta as feed and digesta in the colon as feces.
6. Rectum - using colon digesta as feed and the feces voided as feces.

With this method, a positive value indicated net absorption from the section and a negative value, net secretion into that section of the gastrointestinal tract.

3. Experiment 3

The effect of high levels of Ca and P supplementation and an inverse Ca-P ratio on Mn utilization by the growing pig.

A feeding trial using a 2^3 factorial design was conducted. Eighty weanling pigs weighing approximately 8 kg were randomly allotted to 8 treatments shown in Table 2, equalizing for sex, litter and weight. During the trial, the animals were housed in confinement on slotted concrete floors. Feed and water were provided *ad libitum*. The pigs were weighed every 2 weeks and 2 animals per lot, selected at random, were bled every 4 weeks to monitor changes in blood constituents. Blood samples were taken from the anterior vena cava.

After 10 weeks, the experiment was terminated when the animals weighed an average of 60 kg. Four animals from each lot, including the 2 animals that were regularly bled, were slaughtered at the university Meats Laboratory and tissue samples collected for chemical, physical and histopathological analyses. The tissues collected included bones,

TABLE 2. DIETS USED IN FEEDING TRIAL (EXPERIMENT 3)

Item	1	2	3	4	5	6	7	8
Ca, % (calculated)	.7	.7	.35	.35	.7	.7	1.4	1.4
P, % (calculated)	1.4	1.4	.7	.7	.35	.35	.7	.7
Mn, ppm (analyzed)	15.3	56.4	15.9	55.7	15.2	56.1	16.0	55.8
Corn	753	743	789	779	791	781	761	751
Soybean meal de- hulled solvent, (49% CP)	188	188	183	183	181	181	186	186
Salt	5	5	5	5	5	5	5	5
Limestone					17	17	23	23
Dicalcium phosphate	26	26	12	12			19	19
VTM premix ¹	5	5	5	5	5	5	5	5
Mn premix ²		10		10		10		10
Antibiotic ³	1	1	1	1	1	1	1	1
Monosodium phosphate	22	22	5	5				
	1000	1000	1000	1000	1000	1000	1000	1000

¹MSU vitamin - trace mineral premix without manganese. See Appendix A, Table A-1.

²Containing 4000 ppm Mn from $\text{MnSO}_4 \cdot \text{H}_2\text{O}$ reagent grade.

³Containing 22 g of chlortetracycline per kg.

heart, pancreas, liver and the left kidney. The bones taken included the first left rib and the lateral and medial metacarpals from the left leg. The medial metacarpals were split longitudinally, fixed in buffered 10% formalin, sectioned and stained with hematoxylin and eosin for histopathologic examination. Two sections were taken from each bone, one through the epiphyseal cartilagenous plate from the proximal end of the bone and one section through the diaphysis. Sections of the epiphysis and diaphysis were coded as follows: 1, normal, if they had very little cartilage and osteoid; 2, very slight change, cartilage and osteoid persisting in the bony spicules distal to the epiphyseal cartilagenous plate or distal from the periosteum of the diaphysis; and 3, slight change, 4, moderate change, or 5, severe change, depending on the degree of persistence of cartilage in bony spicules.

4. Experiment 4

The Mn requirement of the baby pig from sows fed a low Mn diet. Eight one-month pregnant first litter gilts were fed a low Mn diet shown in Table 3 throughout the remainder of the gestation period. After farrowing, 20 baby pigs were taken from the sows at 5 days of age and placed in stainless steel rearing cages equipped with stainless steel feeders and water troughs. The room temperature was maintained at 30°C for the period of the trial. The pigs were weaned to the basal dry purified diet (no. 1) shown in Table 4. Pigs were taught to consume the feed by placing small amounts in the animal's mouth. Some pigs readily adapted to the dry feed, but many were affected with diarrhea, became weak, refused to eat and eventually died.

After adapting to the diets, 12 healthy pigs were randomly allotted to the three dietary treatments shown in Table 4. The pigs were fed

TABLE 3. LOW MINERAL RATIONATION DIET (EXPERIMENT 4)

Ingredient	Amount, kg ²
Corn	1915
Limestone	20
Dicalcium phosphate	30
Salt	10
VTM premix ¹	10
Vit. E premix	5
Lysine (50%)	<u>10</u>
	2000

¹See Appendix A, Table A-1 supplied all the trace minerals except Mn.

²Level of manganese 11.2 ppm as analyzed.

TABLE 4. PURIFIED DIETS USED IN EXPERIMENT 4

Item	Diets		
	1	2	3
Casein	30	30	30
Cerelose	55	52	46
Cellulose	3	3	3
Lard	5	5	5
Vitamin premix ¹	1	1	1
Mn premix ²		3	9
Mineral premix ³	<u>6</u>	<u>6</u>	<u>6</u>
	100	100	100
Mn concentration, ppm (analyzed)	0.46	2.67	6.34

¹See Appendix A, Table A-3.

²Manganese premix containing 100 ppm Mn from $\text{MnSO}_4 \cdot \text{H}_2\text{O}$, J. T. Baker reagent grade.

³See Appendix A, Table A-1.

ad libitum and had free access to water, which was changed 3 times a day. All feed was weighed daily and individual feed consumption was recorded. Pigs were individually weighed every 4 days throughout the 30-day trial. One pig (234-4F) on the basal diet died suddenly and the cause of death was determined to be heart failure. The pigs were bled on the 9th, 21st, and 28th day of the feeding trial.

After the feeding trial, the pigs were used for a Mn balance study. Another pig (241-11) on diet 3 was eliminated from the study due to an infection in the left rear leg. The pigs were housed, treated, and fed as described in Experiment 1. The collection and treatment of the feces, urine and refused feed was similar to the procedures described for Experiment 1.

C. Analyses

1. Hematological Determinations

a. Hemoglobin. In Experiments 1 and 3, hemoglobin was determined by the cyanmethemoglobin method of Crosby *et al.* (1954). A Coleman Junior II spectrophotometer was used to determine optical density at 540 nm.

b. Hematocrit. Hematocrit was determined by the micro method (McGovern *et al.*, 1955) in Experiments 1 and 3. Blood samples were centrifuged for 5 minutes at 10,000 rpm in an International "Hemacrit" centrifuge.

2. Physical Determinations

a. Bone. In Experiment 1, the left femur and the 8th left rib were removed and cleaned of all connective tissue and periosteum and

stored in air-tight polyethylene bags. The ribs were used for specific gravity determinations and the femurs for breaking strength and related parameters.

In Experiment 3, the 5th metacarpal was removed from the left foot, cleaned of connective tissue and stored in a cold room in air-tight polyethylene bags. The first left rib was obtained and treated as the metacarpal. Bone strength and density tests were made on both the ribs and the metacarpals.

Specific gravity was determined according to the following formula:

$$\text{Specific gravity} = \frac{\text{weight in air}}{\text{weight in air} - \text{weight in water}}$$

In Experiments 1 and 3, the strength of the metacarpals, femurs, and ribs was determined using an Instron Testing Instrument, Model-TT CML¹ equipped with an FM-compression load cell having 100 kg full scale. The cross-head speed and chart speed were 0.2 cm/minute and 2 cm/minute, respectively. Metal fulcra were used to support the metacarpals and ribs and the distance between fulcra was maintained constant at 3 and 4 cm, respectively. The femurs varied in size and therefore the distance between supporting fulcra varied from bone to bone. The formulas for calculating the various strength characteristics were those described by Miller *et al.* (1962) and are as follows:

Maximal bending moment	$M = Wl/4$
Moment of inertia	$I = (Bd^3 - bd^3)/64$
Maximal stress	$S = MD/2I$
Elasticity	$E = Wl^3/48Iy$

¹Instron Engineering Corporation, Canton, Mass.

W = maximal load

l = distance between fulcra

B,b = outer and inner horizontal diameter

D,d = outer and inner vertical diameter

y = deflection at center of bone when load W is applied

3. Chemical Determinations

a. Blood. Upon withdrawal, the blood was placed in acid-washed centrifuge tubes, allowed to clot and centrifuged at 2000 g for 20 minutes. The cell-free serum was then harvested and placed in acid-washed vials. Small amounts were stored in the cold room (4°C) for determination of alkaline phosphatase activity, and the remaining serum samples were frozen for subsequent mineral analysis.

(1) Serum alkaline phosphatase. Determination of serum alkaline phosphatase activity was made according to the procedure described in Sigma Technical Bulletin No. 194 (1963). The Sigma 104 phosphatase substrate was used in the enzyme activity determination. A Beckman Model DU spectrophotometer was used for optical density determinations in Experiments 1, 3 and 4.

(2) Serum calcium and phosphorus. To one milliliter of serum from Experiment 3 was added 4 ml of 12.5% TCA. Serum proteins precipitated by the 12.5% TCA were centrifuged out at 2000 g for 15 minutes, and the resulting protein-free supernatant was diluted 1:1 with strontium mixture A¹ to suppress phosphate interference. Calcium was

¹61.0 g strontium chloride (SrCl₂ · 6H₂O) + 10.0 g sodium chloride (NaCl).

then determined at 422.6 nm by atomic absorption spectrophotometry using the Jarrell-Ash¹ Model 82-516 spectrophotometer equipped with a Hetco total consumption burner and an air-hydrogen flame, as described by Ullrey *et al.* (1967).

For phosphorus determinations the Gomorri (1942) method was used. The optical density was determined on a Coleman Junior II spectrophotometer at 700 nm after a 45-minute incubation period.

(3) Serum magnesium. Cell-free serum samples from Experiments 3 and 4 were diluted 1:100 with strontium mixture B² to avoid phosphate interference. Magnesium was determined with a Jarrell-Ash atomic absorption spectrophotometer at 285.2 nm in Experiment 3, and in Experiment 4 an Instrumentation Laboratories, Inc.,³ Model 453 atomic absorption spectrophotometer was used.

(4) Serum manganese-neutron activation

(a) Principle. Mn in serum from Experiments 1 and 3 was determined by modification of the neutron activation procedure of Hahn *et al.* (1968). The principle involved exposure of a small amount of serum to a thermal neutron flux in a nuclear reactor and the following nuclear reaction occurred:



¹Jarrell-Ash Co., Waltham, Mass.

²30.5 g strontium chloride (SrCl₂·6H₂O) + 5.0 g sodium chloride (NaCl).

³Instrumentation Laboratories, Inc., Lexington, Mass.

The induced ^{56}Mn has a 2.56 hour half-life and emits gamma rays of 0.85 Mev. The activity of ^{56}Mn can be measured by using gamma-ray spectrometry, comparing the areas of the 0.85 Mev gamma-ray photopeaks with those of the induced standards (Anong-Nilubol *et al.*, 1968).

(b) Sample preparation. One milliliter of cell-free serum was pipetted into an acid-washed 250 ml Phillips beaker and 5 ml of concentrated HNO_3 acid were added. The contents were heated gently to boiling and evaporated to dryness. Three milliliters of 1N HNO_3 were added and heated again gently to boiling. After cooling, the volume was made up to 5 ml. A like method was used to prepare the standards and nitric acid blanks.

(c) Irradiation. Four milliliters were pipetted into polyethylene vials,¹ 1.5 cm in diameter. The vials were thoroughly cleaned (Jacobson *et al.*, 1961)² before loading. The vials were heat-sealed, loaded into and activated by the MSU Triga Mark 1 reactor.³ The samples were subjected to a thermal neutron flux of $2 \times 10^{12} \text{ n/cm}^2/\text{sec}$ for a period of 15 minutes. The samples were removed from the reactor and the radioactivity of the polyethylene vials and samples was in the mrem/hour range and presented no special handling problem.

(d) Radiochemical separation. Radiochemical separation was performed according to the method of Hahn *et al.* (1968). The irradiated samples were transferred to separatory funnels. The vials

¹1.5 cm diameter - Olympia Plastics, Los Angeles, Calif.

²See Appendix A, Table A-4.

³Gulf General Atomic, Inc., San Diego, Calif.

were rinsed with 8 ml of deionized, distilled water. Two drops of 0.1% brilliant yellow were added and the pH was adjusted to neutral using 5N NH_4OH . Then 2 ml of 5.7% sodium diethyldithiocarbamate and 5 ml of carbon tetrachloride were added. The contents were shaken for exactly 3 minutes and 4 ml of the organic layer were pipetted into clean vials for counting.

(e) Counting procedure. Counting was done about one hour after the samples had been removed from the reactor on a 5.2 multi-channel analyzer - computer series one-thirty¹ utilizing a 3" X 3" NaI well crystal.² The voltage was set at 1100 volts. The counts of the integrated areas under the peaks were corrected for both background, decay time and blank. Sample counts were compared to standard counts for quantitation.

(5) Serum manganese - flameless atomic absorption. Mn in the serum samples from Experiment 4 was determined by flameless atomic absorption spectrophotometry on the Instrumentation Laboratories, Inc., Model 355 accessory to the Model 453. Serum samples were diluted 20% v/v. The diluted serum was divided into three 25 μl aliquots. To all but one of these were added known amounts of manganese, 10 μl of either 5 or 10 ngm Mn/ml. The solutions were then placed on the tantalum ribbon and analyzed at 279.4 nm.³ The samples were dried, pyrolyzed for 90 to 120 seconds and analyzed at a higher temperature than the one used during pyrolysis. The Mn in the serum was calculated using the method of additions (Slavin, 1968).

¹Tektronic, Inc., Portland, Ore.

²Packard Model, 1212 WSP serial 101-769.

³See Appendix A, Table A-5.

b. Bone. The cleaned ribs and metacarpals from Experiments 1 and 3 used in the strength and density studies were used in the chemical analyses as well.

(1) Bone ash. The bones were cut in small pieces with a power hand saw, wrapped in cheesecloth and extracted 24 hours with absolute ethanol and 24 hours with anhydrous diethyl ether in a Soxhlet extractor to remove water and fat. The dry, fat-free bone was ashed in a muffle furnace at 600°C for 18 hours. The percent ash was calculated from the following formula:

$$\frac{\text{weight of ashed bone} \times 100}{\text{weight of dry, fat-free bone}} = \% \text{ ash on a dry, fat-free basis}$$

(2) Bone magnesium, manganese, calcium and phosphorus. The ashed bone was finely ground and approximately 300 mg of the powdered bone ash were dissolved in 5 ml of 6N HCl. Two milliliter aliquots of the resulting ash solution were diluted 1:1 with strontium mixture A and Mn was determined on a Jarrell-Ash atomic absorption spectrophotometer at 279.4 nm. Bone Mn was expressed as ppm on a dry, fat-free basis.

The remainder of the ash solution was diluted 1:20 with deionized, distilled water for further mineral analysis. The resulting solutions were further diluted 1:100 with strontium mixture B and Ca and Mg were determined by atomic absorption spectrophotometry as previously described for blood serum. Phosphorus was determined by the colorimetric method previously described for serum inorganic P. Bone Ca, P and Mg were expressed as percent of the dry, fat-free bone.

c. Feed, feces and digesta

(1) Manganese. A wet ashing procedure was used. A 0.5 to 1.0 g sample was weighed into an acid-washed 250 ml Phillips beaker. One milliliter of strontium mixture A was added and the contents digested in 60 ml of concentrated HNO_3 acid on a hot plate to near dryness and cooled. A second digestion with 7 ml of 72% perchloric acid was performed. The contents were protected from excessively rapid evaporation by a small water glass. They were heated to near dryness, cooled and samples diluted to volume with deionized, distilled water. Standards were prepared in a like manner. Mn content was determined by atomic absorption spectroscopy using a Jarrell-Ash Model 82-516 spectrophotometer for feed and fecal samples from Experiments 1 and 3. For digesta, feed and fecal samples from Experiments 2 and 4, Mn was determined with the Instrumentation Laboratories, Inc., Model 453 atomic absorption spectrophotometer.

(2) Chromium. Chromium was determined by the method of Bolin *et al.* (1952). A 100 to 500 mg sample of feed, feces, or digesta was weighed into a 50 ml Erlenmeyer flask and 5 ml of oxidizing reagent¹ were added. The flask was then heated on a hot plate to digest the mixture until it was clear. The mixture was cooled and 2 ml of 72% perchloric acid added and reheated. The flask was cooled to room temperature and diluted to 50 ml using distilled, deionized water. The samples were then read at 470 nm using distilled, deionized water as a blank. The standard curve was prepared by oxidizing known amounts of the reference chromic oxide and diluting as described above.

¹10.0 g sodium molybdate + 150 ml distilled deionized water + 150 ml concentrated sulfuric acid and 200 ml 72% perchloric acid.

d. Urine. A twofold concentration and digestion procedure was devised due to the low Mn content in urine. Twenty milliliters of urine were pipetted into a Phillips beaker, 1 ml of strontium mixture A was added and the contents were digested with 50 ml of concentrated HNO_3 acid to near dryness. The flasks were cooled and diluted to 10 ml with deionized, distilled water. Manganese was determined by atomic absorption spectroscopy on the Jarrell-Ash unit for urine from Experiment 1 and on the Instrumentation Laboratories, Inc., Model 453 atomic absorption spectrophotometer for urine from Experiment 4. The method and specifications of determination were as described previously.

e. Tissues. The tissues included liver, kidney, pancreas, heart, spleen, testes and muscle from both Experiments 1 and 3. Magnesium and P were determined on only the liver, kidney, pancreas and heart samples from Experiment 3, and Mn was determined on all the samples from both experiments.

(1) Tissue dry matter. Approximately 2 g samples were weighed into disposable aluminum dishes and dried in a vacuum oven for 24 hours at 90°C. Dry matter was calculated as follows:

$$\text{Percent dry matter} = \frac{\text{tissue dry weight} \times 100}{\text{tissue fresh weight}}$$

(2) Minerals. Tissue homogenates containing about 1 to 2 g of the tissues were pipetted into 250 ml Phillips beakers and a wet ashing procedure was used as described previously for feed. The digesta were appropriately diluted and Mn, Ca and Mg were determined by atomic absorption spectroscopy and P by the colorimetric method described previously.

f. Hair. Hair samples were soaked for 15 to 20 minutes in deionized, distilled water, drained on filter paper and then placed in 95% ethanol for 15 to 20 minutes to remove adhering contaminant materials. After removing from the ethanol, the samples were air-dried and weighed into a 250 ml Phillips beaker and wet-ashed. Mn was determined by atomic absorption spectrometry as reported previously.

D. Statistical Analysis

All data from Experiments 1, 2, 3 and 4 were subjected to analysis of variance on a CDC¹ 3600 computer at the Michigan State University Computer Laboratory Center. The same computer was used to calculate simple correlations. Differences between means were determined by Tukey's test for non-additivity.

¹Control Data Corporation, Minneapolis, Minn.

RESULTS AND DISCUSSION

A. Experiment 1: The relative availability of Mn from 16% corn-soy basal diet supplemented with 10 ppm of Mn from either $\text{MnSO}_4 \cdot \text{H}_2\text{O}$, MnCO_3 or MnO for the growing pig

1. In vitro Solubility of Manganese Compounds

The solubility of manganous sulfate monohydrate ($\text{MnSO}_4 \cdot \text{H}_2\text{O}$), manganous carbonate (MnCO_3) and manganous oxide (MnO) is shown in Table 5. The sulfate was soluble in water but the carbonate and oxide were practically insoluble. The sulfate and the carbonate were equally soluble in 0.4% HCl but the oxide was slightly less soluble. These findings are in agreement with those of Watson *et al.* (1971).

TABLE 5. CHEMICAL COMPOSITION AND SOLUBILITY OF DIFFERENT MANGANESE COMPOUNDS¹

Compound	Mn %	Water			0.4% HCl		
		% Solubility	pH		% Solubility	pH	
			Initial	Final		Initial	Final
$\text{MnSO}_4 \cdot \text{H}_2\text{O}$	32.6	96.3	7.0	4.15	100.0	1.03	1.06
MnCO_3	44.3	0.91	7.0	7.70	100.0	1.03	1.04
MnO	65.0	0.12	7.0	7.65	89.5	1.03	1.10

¹Water-bath was maintained at 37°C.

The pH dropped sharply after the sulfate dissolved in water (Table 5), presumably due to dissociation of manganous sulfate and the formation

of a predominantly acid medium. With the carbonate and oxide, the pH rose slightly. The pH changes after these compounds were dissolved in 0.4% HCl were very small.

2. Mn Availability Studies Using a Split-Plot Design (where animals were fed their respective experimental diets for three consecutive Mn balance trials and three blood collections)

Weight gains, physical bone measures, and tissue Mn concentrations were not affected significantly by source of Mn (Tables 6 and 7). Animals receiving supplemental MnCO_3 had significantly ($P < 0.05$) heavier livers as a percent of bodyweight than those of animals receiving supplemental $\text{MnSO}_4 \cdot \text{H}_2\text{O}$. The kidneys, hearts and testes as a percent of body weight of animals on the basal diet were slightly heavier than those of animals on the other diets (Table 8).

The physical bone measures presented in Table 9 did not show any consistent pattern of variation with respect to Mn sources. Mn from all sources was equally well utilized for bone formation. Weight gains and tissue Mn concentration (Table 10) showed that Mn from all the compounds studied was equally utilized by the pig. This study also suggests that Mn requirements of young pigs for growth are not higher than 16.2 ppm, based on weight gains on all the diets. The low Mn levels of tissues from pigs on the basal diet may be a reflection of its low Mn content. In this study only liver, pancreas, spleen and testes Mn concentrations showed a substantial response to Mn supplementation.

Hemoglobin, hematocrit, serum Mn concentration and serum alkaline phosphatase activity did not differ significantly due to source of Mn; but serum Mn, serum alkaline phosphatase and hematocrit did differ significantly ($P < 0.01$) within the different Mn sources due to time of

TABLE 6. ANALYSIS OF VARIANCE FOR MN SOURCE EFFECTS ON BONE AND BLOOD PARAMETERS (SPLIT-PLOT DESIGN)

Category	Mean	Min.	Max.	P-level of F-statistic
<u>Femur, left leg</u>				
Weight, g (fresh basis)	72.6	63.9	85.5	0.40
Length, cm ^{1/}	9.2	8.5	9.5	0.12
External diameter, ^{2/} cm				
Horizontal (B)	1.42	1.31	1.5	0.65
Vertical (D)	1.35	1.25	1.5	0.93
Internal diameter, cm				
Horizontal (b)	1.02	0.81	1.31	0.89
Vertical (d)	0.98	0.80	1.20	0.79
Breaking strength, kg	165	140	187	0.67
Inertia, cm ⁴	1.15	1.03	1.26	0.07
Stress at the center, kg/cm ²	972	856	1089	0.06
Elasticity, 1000 kg/cm ²	27.6	21.6	34.2	0.33
<u>Rib, 8th left</u>				
Weight, g (fresh basis)	4.40	3.20	5.49	0.36
Ash, % of dry, fat-free rib	58.6	57.9	59.8	0.19
Specific gravity (fresh basis)	1.25	1.20	1.31	0.70
<u>Blood parameters</u>				
Serum Mn, mcg/100 ml	1.65	0.91	2.71	0.28
Hemoglobin, g/100 ml	10.8	9.0	12.2	0.68
Hematocrit, %	33.4	30.2	37.7	0.70
Serum alkaline phosphatase, Sigma units	7.12	5.0	11.4	0.72

¹Measured from the mid-medial condyle to the fovea.

²The horizontal and vertical diameters were measured at mid-shaft when the bone was positioned in such a way that the medial and lateral condyles were facing downwards.

TABLE 7. ANALYSIS OF VARIANCE FOR MN SOURCE EFFECTS ON MN BALANCE, TISSUE MN CONCENTRATIONS, ORGAN WEIGHTS AND GROWTH DATA (SPLIT-PLOT DESIGN)

Category	Mean	Min.	Max.	P-level of F-statistic
<u>Balance data</u>				
Mn intake, mg/day	8.70	5.94	10.1	<0.0005
Mn excretion, mg/day				
Fecal	3.82	1.77	5.63	0.001
Urinary	.05	.04	.07	0.355
Mn retention, mg/day	4.84	2.52	6.15	<0.0005
Mn retention, % of intake	55.3	41.8	69.6	0.56
Mn excretion, % of intake				
Fecal	44.1	29.6	57.4	0.61
Urinary	.60	.39	.99	<0.0005
<u>Tissue, Mn, ppm (dry matter basis)</u>				
Liver	6.93	5.14	9.03	0.11
Kidney	6.60	5.55	7.87	0.12
Pancreas	4.92	3.93	6.02	0.24
Spleen	2.38	1.94	2.76	0.83
Bone	1.12	0.77	1.54	0.52
Muscle	1.84	1.50	2.22	0.22
Heart	2.51	2.33	2.77	0.80
Testes	3.88	3.22	4.39	0.82
Hair				
Initial value	1.00	0.80	1.33	0.59
Final value	1.17	0.96	1.44	0.22
<u>Organs, % of body weight</u>				
Liver	2.12	1.93	2.33	0.05
Kidney	.33	.29	.37	0.35
Pancreas	.14	.11	.20	0.49
Spleen	.20	.14	.26	0.08
Heart	.49	.44	.58	0.96
Testes	.16	.10	.25	0.36
<u>Growth data</u>				
Average daily gain, g	187	176	225	0.73

TABLE 8. THE EFFECT OF MN SOURCE ON THE PHYSICAL CHARACTERISTICS OF DIFFERENT ORGANS¹

Item	Diet								SE ²
	Basal		+MnSO ₄ ·H ₂ O		+MnCO ₃		+MnO		
Mn conc., ppm	16.2		26.0		24.8		27.0		
No. of pigs	3		3		3		3		
	Abs. ³	% BW ⁴	Abs.	% BW	Abs.	% BW	Abs.	% BW	
Liver	324	2.10	287	1.90	311	2.20 ^{6/}	315	2.10	.03
Kidney	51	0.34	48	0.32	47	0.33	49	0.33	.08
Pancreas	23	0.15	20	0.13	18	0.12	22	0.15	.01
Heart	80	0.53	74	0.49	70	0.48	69	0.46	.08
Spleen	32	0.21	23	0.15	30	0.21	32	0.22	.06
Testes ⁵	34	0.22	18	0.12	20	0.13	20	0.14	.07

¹Based on fresh basis.

²Statistical analyses made on percent bodyweight only.

³Absolute weight in grams.

⁴As percent of body weight.

⁵Data based on two males on diets 1, 3 and 4 and 3 males on diet 2.

⁶Significantly (P<0.05) greater than least value.

TABLE 9. THE EFFECT OF MN SOURCE ON THE PHYSICAL MEASUREMENTS OF THE FEMURS AND RIBS, BREAKING STRENGTH AND RELATED PARAMETERS OF THE FEMURS

Item	Diet			
	Basal	+MnSO ₄ ·H ₂ O	+MnCO ₃	+MnO
Mn conc., ppm	16.2	26.0	24.8	27.0
No. of pigs	3	3	3	3
<u>Femur, left</u>				
Weight, g (fresh basis)	73.2	77.0	68.3	71.7
Length, cm ^{1/}	9.3	9.0	9.0	9.5
External diameter, ^{2/} cm				
Horizontal (B)	1.46	1.43	1.37	1.45
Vertical (D)	1.37	1.35	1.32	1.36
Internal diameter, cm				
Horizontal (b)	1.05	1.03	1.03	0.96
Vertical (d)	1.05	0.94	0.94	1.00
Breaking moment, kg	171	166	166	157
Inertia, cm ⁴	1.12	1.17	1.21	1.09
Stress at center, kg/cm ²	1049	951	909	978
Elasticity, 1000 kg/cm ²	27.6	24.4	30.7	27.8
<u>Rib, 8th left</u>				
Weight, g (fresh basis)	4.9	4.6	3.8	4.5
Ash, % of dry fat-free rib	58.9	58.5	58.6	58.3
Specific gravity (fresh basis)	1.27	1.23	1.26	1.23

¹Measured from the mid-medial condyle to the fovea.

²The horizontal and vertical diameters were measured at mid-shaft when the bone was positioned in such a way that the medial and lateral condyles were facing downwards.

TABLE 10. THE EFFECT OF MN SOURCE ON THE AVERAGE DAILY GAIN (ADG) AND TISSUE MN DISTRIBUTION

Item	Diet			
	Basal	+MnSO ₄ ·H ₂ O	+MnCO ₃	+MnO
Mn conc., ppm	16.2	26.0	24.8	27.0
No. of pigs	3	3	3	3
Avg. daily gain, g	200	186	180	182
Tissue Mn, ppm ¹				
Bone, 8th left rib	1.16	1.07	0.95	1.32
Liver	5.7	7.4	7.3	7.3
Kidney	6.7	6.7	6.1	6.9
Heart	2.5	2.6	2.5	2.5
Pancreas	4.5	5.1	4.8	5.3
Spleen	2.0	2.6	2.4	2.5
Testes ²	3.4	4.1	4.1	4.0
Muscle	1.8	2.0	1.7	1.9
Hair	1.1	1.2	1.3	1.1

¹Manganese expressed on dry matter basis except for hair and bone. Bone Mn was expressed on dry, fat-free basis and hair on ethanol-cleaned, air-dried basis.

²Based on 2 males on diets 1, 3 and 4 and 3 males on diet 2.

sampling (Table 11). Mn retention, fecal excretion and urinary excretion, all as percent of intake, showed significant ($P < 0.01$) Mn source effects. Time of sampling, but not Mn source, had a significant ($P < 0.05$) effect on urinary Mn excretion. Absolute Mn retention and urinary Mn excretion, as percent of intake, did not differ significantly source X time interaction. Only absolute urinary Mn excretion and urinary Mn excretion as a percent of intake showed a significant ($P < 0.05$) source X time interaction.

The hematocrit value at the second sampling was significantly ($P < 0.05$) greater than that at the third sampling for animals receiving supplemental $MnCO_3$ (Table 12). Serum alkaline phosphatase activity at the initial sampling of animals on the basal, basal supplemented with $MnSO_4 \cdot H_2O$ and basal supplemented with $MnCO_3$ were significantly ($P < 0.05$) greater than at the third sampling. The hematocrit and serum alkaline phosphatase values dropped on the second and third sampling on all diets. Serum alkaline phosphatase levels were lowest on the basal diet. Serum Mn increased with time of sampling on the basal diet and basal diet supplemented with $MnSO_4 \cdot H_2O$. On diets supplemented with $MnCO_3$ and MnO , the levels rose on the second sampling and dropped substantially on the third.

The Mn balance data are summarized in Table 13. The low Mn intake and excretion on the basal diet were a reflection of the lower Mn content of the basal diet as compared to the supplemented diets. The absolute urinary excretion of Mn was the same on all diets. The absolute Mn retention was significantly ($P < 0.01$) lower on the basal diet, essentially equal on diets supplemented with $MnSO_4 \cdot H_2O$ and $MnCO_3$, and slightly higher on the diet supplemented with MnO . When Mn retention was expressed as a percent of intake, the supplemented diets showed a

TABLE 11. THE EFFECT OF MN SOURCE AND TIME OF SAMPLING ON THE BLOOD AND BALANCE DATA¹ (SPLIT-PLOT DESIGN)

Item	Mn Source	Time of sampling	Source X time interaction
Serum alkaline phosphatase	0.72	<0.0005	0.41
Serum Mn	0.48	0.007	0.40
Hemoglobin	0.68	0.16	0.09
Hematocrit	0.70	0.001	0.59
Mn intake	<0.0005	0.009	0.28
Mn excretion, absolute			
Fecal	0.001	0.43	0.82
Urinary	0.36	0.028	0.03
Mn retention, absolute	<0.0005	0.38	0.87
Mn retention, % of intake	0.56	0.23	0.59
Mn excretion, % of intake			
Fecal	0.61	0.23	0.59
Urinary	<0.0005	0.13	0.03

¹P-level of F-statistic.

TABLE 12. EFFECT OF MN SOURCE AND TIME OF SAMPLING ON SOME BLOOD PARAMETERS

Item	Diet											
	Basal			+MnSO ₄ ·H ₂ O			+MnCO ₃			+MnO		
Mn conc., ppm	16.2			26.0			24.8			27.0		
No. of pigs	3			3			3			3		
Samplings ^{1/}	1	2	3	1	2	3	1	2	3	1	2	3
Hemoglobin, g/100 ml	10.9	10.4	10.2	11.1	10.1	11.2	11.0	10.7	10.6	10.9	11.3	10.9
Hematocrit, %	33.7	33.4	31.8	34.9	33.7	32.8	33.3	34.1	31.4	34.1	33.8	32.9
Serum alkaline phosphatase, Sigma units	7.9	6.0	5.3	9.3	6.4	6.3	9.0	6.8	6.3	8.1	7.3	7.2
	<u>2/</u>			<u>2/</u>			<u>2/</u>					
Serum Mn, mcg/100 ml	1.1	1.2	1.3	1.5	1.6	1.7	2.0	2.1	1.5	2.0	2.9	1.9
											<u>2/</u>	

¹Blood sampling weekly.

²Values under the same dietary source significantly (P<0.05) different than the least value.

TABLE 13. RETENTION AND ROUTES OF EXCRETION OF MN FROM BASAL DIET AND BASAL DIET SUPPLEMENTED WITH MANGANOUS SULFATE ($\text{MnSO}_4 \cdot \text{H}_2\text{O}$), MANGANOUS CARBONATE (MnCO_3) OR MANGANOUS OXIDE (MnO)

Item	Diet			
	Basal	+ $\text{MnSO}_4 \cdot \text{H}_2\text{O}$	+ MnCO_3	+ MnO
Mn conc., ppm	16.2	26.0	24.8	27.0
No. of pigs	3	3	3	3
Mn intake, mg/day	6.0	9.6 ^{3/}	9.2 ^{2/}	10.0 ^{4/}
Mn excretion, mg/day				
Fecal	2.7	4.2 ^{1/}	3.8	4.2 ^{1/}
Urinary	0.1	0.1	0.1	0.1
Mn retention, mg/day	3.2	5.3 ^{2/}	5.3 ^{2/}	5.7 ^{2/}
Mn retention, % of intake	53.4	55.1	57.4	57.0
Mn excretion, % of intake				
Fecal	44.9 ^{4/}	43.8	41.3	41.9
Urinary	1.7 ^{4/}	1.0	1.1	1.0

¹Significantly ($P < 0.05$) greater than least value.

²Significantly ($P < 0.01$) greater than least value.

³Significantly ($P < 0.01$) greater than least two values.

⁴Significantly ($P < 0.01$) greater than least three values.

slight advantage over the basal diet. Mn from $\text{MnSO}_4 \cdot \text{H}_2\text{O}$ was retained to a lesser extent than that from the other two compounds. Fecal Mn excretion as a percent of intake was slightly higher on diets supplemented with $\text{MnSO}_4 \cdot \text{H}_2\text{O}$ than those supplemented with MnCO_3 and MnO.

Urinary Mn excretion as a percent of intake was significantly ($P < 0.01$) different between the basal and the supplemented diets, but not within the supplemented diets. Fecal and urinary Mn excretion, both as a percent of intake, on the basal diet were higher than that on the supplemented diets. Regardless of dietary Mn source, over 90% of the excreted Mn was found in the feces.

3. Mn Availability Studies Using a Replicated Latin Square Design

(where all animals were subjected to all experimental diets in four collection periods)

The Mn source had no significant effect on the blood parameters (Table 14). There was a slightly lower serum Mn on the basal diet, presumably as a reflection of the lower Mn content of the diet. All sources were equally effective in maintaining the levels of hemoglobin, hematocrit and serum alkaline phosphatase (Table 15). Mn retention, fecal and urinary Mn excretion, all as percent of intake, were significantly ($P < 0.01$) different between treatments (Table 16). Mn intake and absolute fecal Mn excretion and retention were significantly ($P < 0.01$) different, but not absolute urinary Mn excretion.

The absolute fecal Mn excretion on the basal diet was significantly ($P < 0.01$) lower than on the supplemented diets. Absolute fecal Mn excretion was slightly higher on diets supplemented with $\text{MnSO}_4 \cdot \text{H}_2\text{O}$ and MnO than those supplemented with MnCO_3 . The absolute urinary Mn excretion was equal on all diets but, when expressed as a percent of intake, the

TABLE 14. EFFECTS OF MN SOURCE ON BLOOD AND MN BALANCE MEASURES
(REPLICATED LATIN SQUARE DESIGN)

Item	Mean	Min.	Max.	P-level of F-statistic
<u>Blood parameters</u>				
Hemoglobin, g/100 ml	10.74	8.17	14.92	0.181
Hematocrit, %	33.95	31.00	44.80	0.528
Serum alkaline phosphatase, Sigma units	7.42	4.70	11.40	0.559
Serum Mn, mcg/100 ml	1.94	1.26	3.34	0.160
<u>Balance data</u>				
Mn intake, mg/day	7.08	4.65	9.41	<0.0005
Mn excretion, mg/day				
Fecal	3.63	2.54	4.98	<0.0005
Urinary	.030	.008	.067	0.264
Mn retention, mg/day	3.39	1.49	5.72	<0.0005
Mn retention, % of intake	47.37	30.00	67.02	<0.0005
Mn excretion, % of intake				
Fecal	52.14	32.82	69.11	<0.0005
Urinary	.14	.39	.88	<0.0005

TABLE 15. EFFECT OF MN FROM DIFFERENT SOURCES ON SOME BLOOD MEASUREMENTS

Item	Diet			
	Basal	+MnSO ₄ ·H ₂ O	+MnCO ₃	+MnO
Mn conc., ppm	16.2	26.0	24.8	27.0
No. of pigs	12	12	12	12
Hemoglobin, g/100 ml	11.1	10.4	10.9	10.6
Hematocrit, %	34.5	34.3	33.7	33.4
Serum alkaline phosphatase, Sigma units	7.56	7.39	7.67	7.06
Serum Mn, mcg/100 ml	1.70	2.15	1.76	2.25

TABLE 16. RETENTION AND ROUTES OF EXCRETION OF MN FROM BASAL DIET AND BASAL DIET SUPPLEMENTED WITH MANGANOUS SULFATE ($MnSO_4 \cdot H_2O$), MANGANOUS CARBONATE ($MnCO_3$) AND MANGANOUS OXIDE (MnO)

Item	Diet			
	Basal	$MnSO_4 \cdot H_2O$	$+MnCO_3$	$+MnO$
Mn conc., ppm	16.2	26.0	24.8	27.0
No. of pigs	12	12	12	12
Mn intake, mg/day	5.17	8.31 ^{4/}	7.93 ^{2/}	8.63 ^{6/}
Mn excretion, mg/day				
Fecal	3.10	4.26 ^{2/}	3.91 ^{2/}	4.24 ^{2/}
Urinary	0.030	0.034	0.030	0.028
Mn retention, mg/day	2.04	4.02 ^{2/}	3.99 ^{2/}	4.36 ^{2/}
Mn retention, % of intake	38.8	48.0 ^{1/}	50.1 ^{2/}	50.3 ^{2/}
Mn excretion, % of intake				
Fecal	60.6 ^{4/5/}	51.6	49.5	49.4
Urinary	0.56 ^{6/}	0.39	0.36	0.31

¹Significantly ($P < 0.05$) greater than least value.

²Significantly ($P < 0.01$) greater than least value.

³Significantly ($P < 0.05$) greater than least two values.

⁴Significantly ($P < 0.01$) greater than least two values.

⁵Significantly ($P < 0.05$) greater than least three values.

⁶Significantly ($P < 0.01$) greater than least three values.

pigs on the basal diet excreted significantly ($P < 0.01$) more urinary Mn than the pigs on supplemented diets. Within the supplemented diets, urinary Mn excretion was not significantly different. Absolute Mn retention was significantly ($P < 0.01$) higher on the supplemented diets than on the basal diet. Mn from the diet supplemented with MnO was retained in somewhat greater amounts than Mn from those diets supplemented with MnCO_3 and $\text{MnSO}_4 \cdot \text{H}_2\text{O}$, but these differences were not statistically significant. When expressed as a percent of intake, Mn retention on the basal diet was significantly ($P < 0.05$ or $P < 0.01$) lower than that on supplemented diets. Fecal Mn excretion, as a percent of intake, on the basal diet was significantly ($P < 0.05$ or $P < 0.01$) higher than that on the supplemented diets, but the differences between the supplemented diets were not significant. Irrespective of Mn source, the main route of Mn excretion was fecal, and urinary Mn excretion was very small and constant on all diets.

4. Discussion of the Results of Experiment 1

The low Mn retention on the basal diet and the high Mn excretion on the same diet agrees with the findings of Morimoto *et al.* (1959), who showed poor Mn availability to chicks when soy protein was used in the diet, and of Davis *et al.* (1962), who reported that soy protein contained a component which combined with Mn, making it unavailable. In this particular study, most of the Mn in the basal diet was supplied by soy protein. The dominance of the fecal route as a means of Mn excretion and the small but constant urinary Mn excretion found in this study have been reported by others in other species (Underwood, 1971; Thomas, 1970; Miller, 1973). The 40 to 50% Mn retention by the pig shown in this study parallels Mn retention in the human reported by North *et al.*

(1960). Brown and McCracken (1965) found that chickens retained 32% of the absorbed Mn.

The conclusion that Mn in $\text{MnSO}_4 \cdot \text{H}_2\text{O}$, MnCO_3 , and MnO is equally available to a pig for growth agrees with the findings in other species. Schaible *et al.* (1938) and Gallup and Norris (1939) found that the sulfate, carbonate and oxide were equally effective in preventing perosis in chickens. Cotzias and Greenough (1958) and Zajvec (1959) have reported that the absorption of Mn was not affected by its valency state in the compounds used. But more recently others have reported that Mn oxides are not as available as the sulfate, carbonate and chloride (Anke *et al.*, 1967; Watson *et al.*, 1970, 1971).

Serum alkaline phosphatase and serum Mn levels were equal on all diets, which probably indicates that Mn from all sources was equally available to the growing pig or that the levels of Mn provided in all diets were high enough to sustain normal levels of serum alkaline phosphatase and serum Mn. Mn deficiency has been shown to reduce serum alkaline phosphatase and serum Mn levels in swine as well as other species (Plumlee *et al.*, 1956; Rojas *et al.*, 1965; Swaney and Kehar, 1958; Hawkins *et al.*, 1955; Ugnenko, 1972). The effect of time of sampling (age of the pig) on hematocrit and serum alkaline phosphatase values found in this study is supported by the findings of Miller *et al.* (1961), who showed a similar drop at a similar age of pigs. Long *et al.* (1965) also reported high serum alkaline phosphatase values in young pigs which gradually dropped with age.

In this study, liver, testes, and spleen Mn concentration increased with dietary Mn levels, which is in agreement with the reports of Leibholz *et al.* (1962), Johnson (1943, 1944), Grummer *et al.* (1950), and Underwood (1971). The response of spleen to Mn supplementation has

been shown in rats by Ugnenko (1972). Hair and bone, which were reported to respond to Mn supplementation by Leibholz *et al.* (1962), did not do so in this study. This may be due to the fact that Leibholz *et al.* (1962) used a 100-fold margin between the basal diet and the supplemented diet as compared to a twofold margin used in this study.

B. Experiment 2: Study of the gastrointestinal flux pattern of Mn from different Mn sources using chromic oxide (Cr_2O_3) as an indicator

The results of this study are summarized in Tables 17, 18 and 19 and Figures 2.1 through 2.4. The cranial small intestine was a major route of absorption for Mn from the supplemented diets but not for Mn from the basal diet. There was no net Mn absorption by the caudal small intestine. Hendricks (1967) showed that pigs fed a 16% soy protein diet supplemented with Mn from $\text{MnSO}_4 \cdot \text{H}_2\text{O}$ displayed net absorption in the stomach and cecum only. Since there is a large amount of Mn in the bile (Kent and McCance, 1941; Mahoney and Small, 1968; Starodubova, 1968), and bile is secreted into the cranial small intestine, it is apparent that in the case of Mn from the supplemented diets, the cranial small intestine must be absorbing Mn faster than it is being secreted into this section of the gut. This finding makes the cranial small intestine a very important and efficient homeostatic mechanism for regulating Mn levels in the body. Britton and Cotzias (1966) concluded that it is the variable excretion rates rather than regulated absorption that seem to maintain constant tissue Mn levels. Howes and Dyer (1971) and Miller (1973) reported that absorption differences are the main homeostatic control mechanisms for body Mn levels.

Although digesta pH did not differ significantly between treatments (Table 18), there was a substantial treatment pH difference in the

TABLE 17. NET ABSORPTION AND SECRETION OF MN IN THE DIFFERENT SECTIONS OF THE GUT BY THE GROWING PIG FED DIFFERENT MN SOURCES¹

Section of gut	Diet							
	Basal		+MnSO ₄ ·H ₂ O		+MnCO ₃		+MnO	
	Abs. ²	Sec. ³	Abs.	Sec.	Abs.	Sec.	Abs.	Sec.
Stomach	60		61		49		52	
Small intestine ⁴								
Cranial		129	22		62		60	
Caudal		185		560 ^{5/}		879 ^{6/}		967 ^{6/}
Cecum	52		41		19		34	
Colon		39		33		24		12
Rectum		64		138		13		119

¹Net flux as percent of Mn in feed or digesta in previous gut section.

²Net absorption.

³Net secretion.

⁴Divided roughly in two halves.

⁵Significantly (P<0.01) greater than least value.

⁶Significantly (P<0.01) greater than least two values.

TABLE 18. GUT CONTENT WEIGHTS AND pH¹

Section of gut	Diet			
	Basal	+MnSO ₄ ·H ₂ O	+MnCO ₃	+MnO
<u>pH values</u>				
Stomach	2.75	3.36	3.19	3.07
Small intestine ²				
Cranial	6.52	6.25	5.59	6.40
Caudal	6.61	6.13	6.77	6.70
Cecum	5.97	6.38	5.80	6.14
Colon	6.09	6.19	6.08	6.10
Rectum	6.45	6.42	6.39	6.39
<u>Gut contents, g (wet basis)</u>				
Stomach	459 ^{5/}	348	351	331
Small intestine				
Cranial	314	347	323	292
Caudal	361	323 ^{4/}	330	358
Cecum	442	498 ^{4/}	372	458 ^{3/}
Colon	74	91	163	125
Rectum	72	81	62	61

¹Based on three animals per treatment.

²Divided roughly into two halves.

³Significantly (P<0.05) greater than least value.

⁴Significantly (P<0.001) greater than least value.

⁵Significantly (P<0.05) greater than least three values.

TABLE 19. MEAN, MINIMUM AND MAXIMUM VALUES OF pH, WET GUT CONTENTS AND NET ABSORPTION AND SECRETION IN DIFFERENT SECTIONS OF THE GUT

Item	Mean	Min.	Max.	P-level of F-statistic
<u>pH</u>				
Stomach	3.09	1.88	3.88	0.82
Small intestine ²				
Cranial	6.21	5.02	7.20	0.29
Caudal	6.55	5.36	7.34	0.39
Cecum	6.07	5.38	6.83	0.50
Colon	6.12	5.72	6.38	0.92
Rectum	6.41	6.05	6.90	0.99
<u>Gut contents, g</u>				
Stomach	372	286	496	0.01
Small intestine				
Cranial	319	265	386	0.25
Caudal	343	278	405	0.75
Cecum	113	42	182	0.02
Colon	443	285	553	0.13
Rectum	69	50	93	0.21
<u>Net absorption (+) or secretion (-)¹</u>				
Stomach	56	41	78	0.63
Small intestine				
Cranial	6	-224	78	0.01
Caudal	-648	-1176	2	0.02
Cecum	36	5	73	0.06
Colon	-21	-85	22	0.21
Rectum	-1002	-264	-1	0.24

¹Net flux as percent of Mn in feed or digesta in previous gut section.

²Divided roughly into two halves.

Figure 2.1. Summary of net Mn flux from sections of the gastrointestinal tract of the growing pig fed the basal diet.

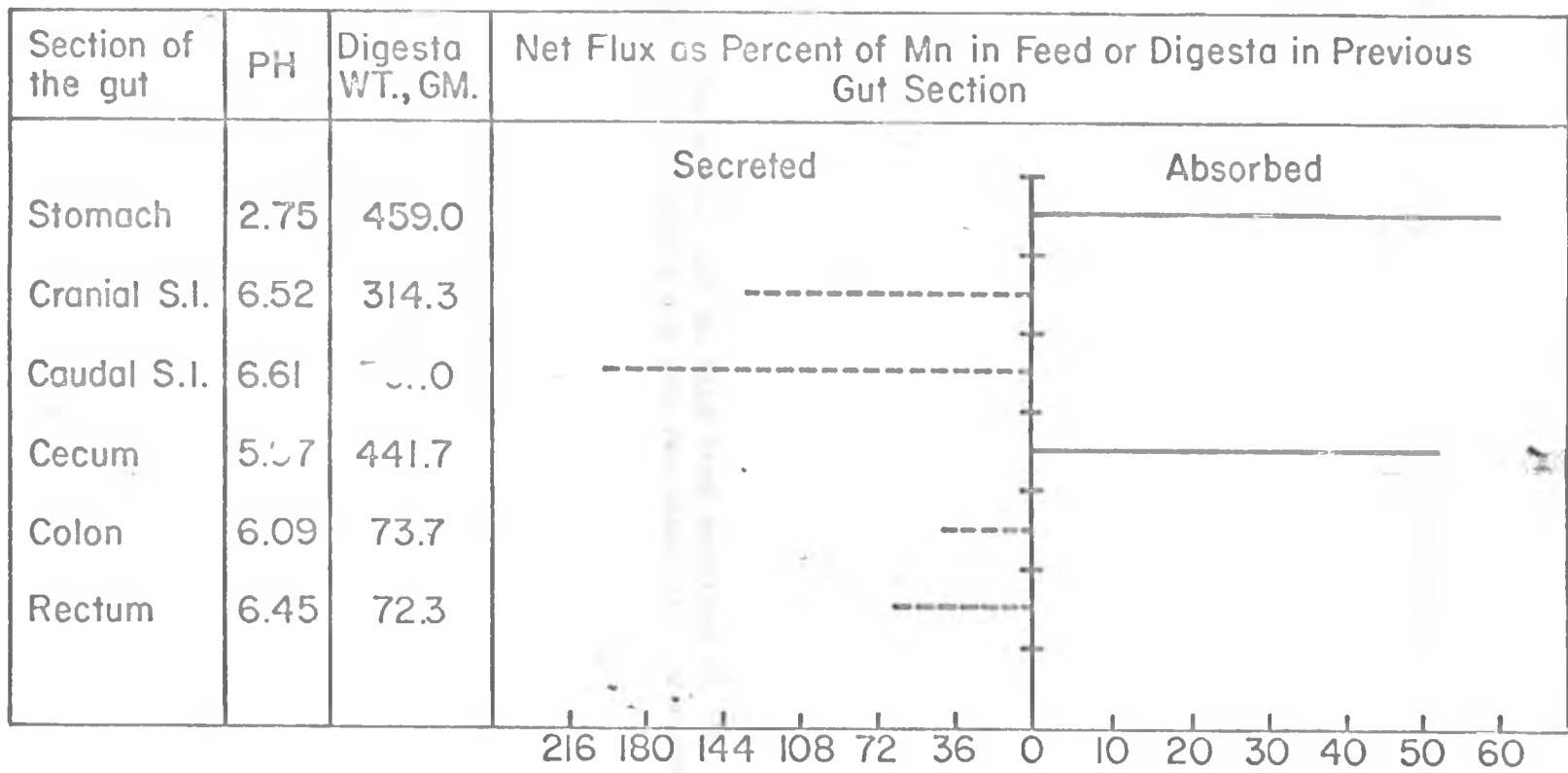


Figure 2.1.

Figure 2.2. Summary of net Mn flux from sections of the gastrointestinal tract of the growing pig fed the basal diet supplemented with $\text{MnSO}_4 \cdot \text{H}_2\text{O}$.

Section	Net Mn Flux (mg/day)
Small Intestine	5.00
Large Intestine	5.00
Stomach	5.00
Colon	5.00
Rectum	5.00
Total	25.00

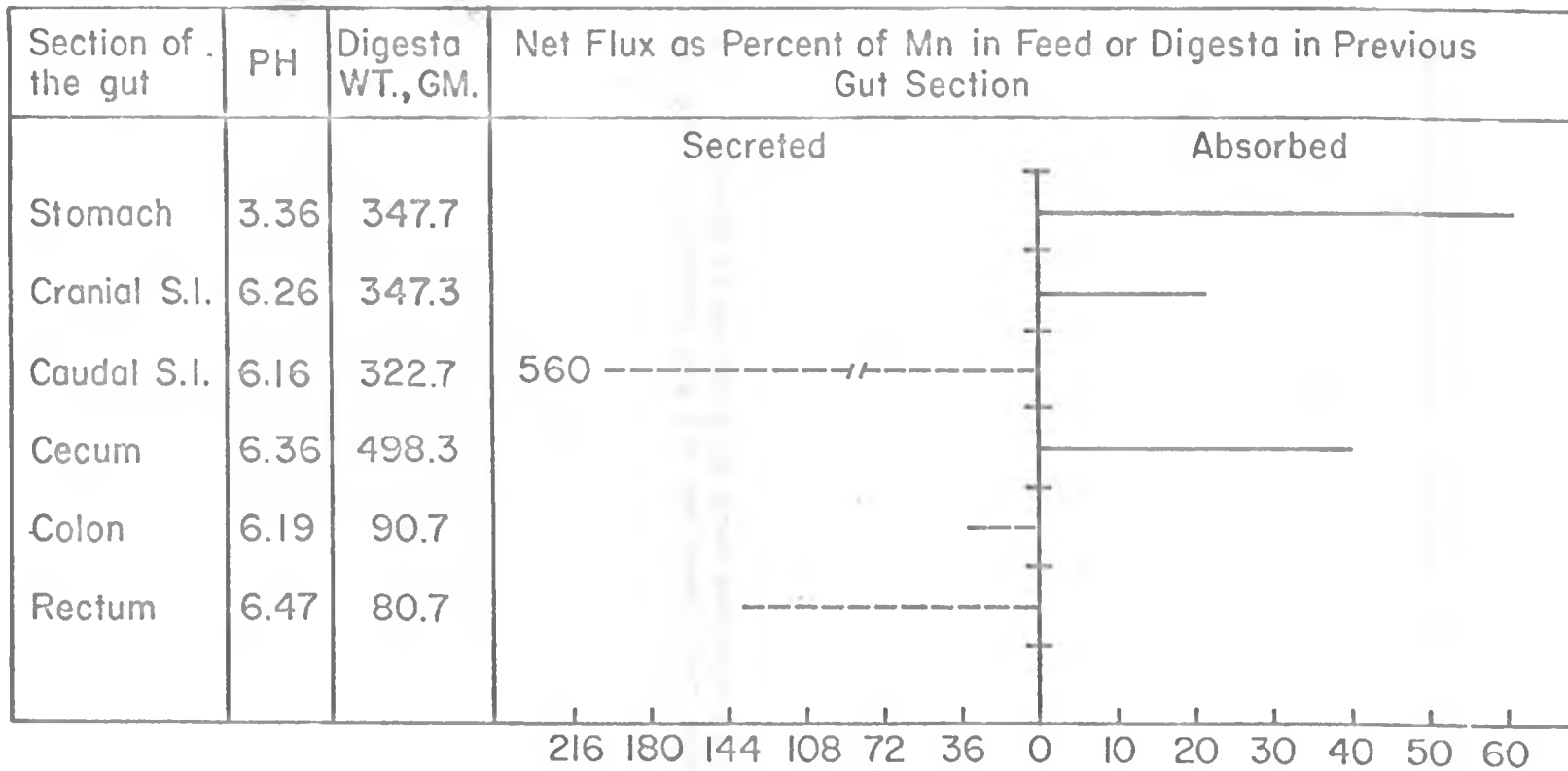
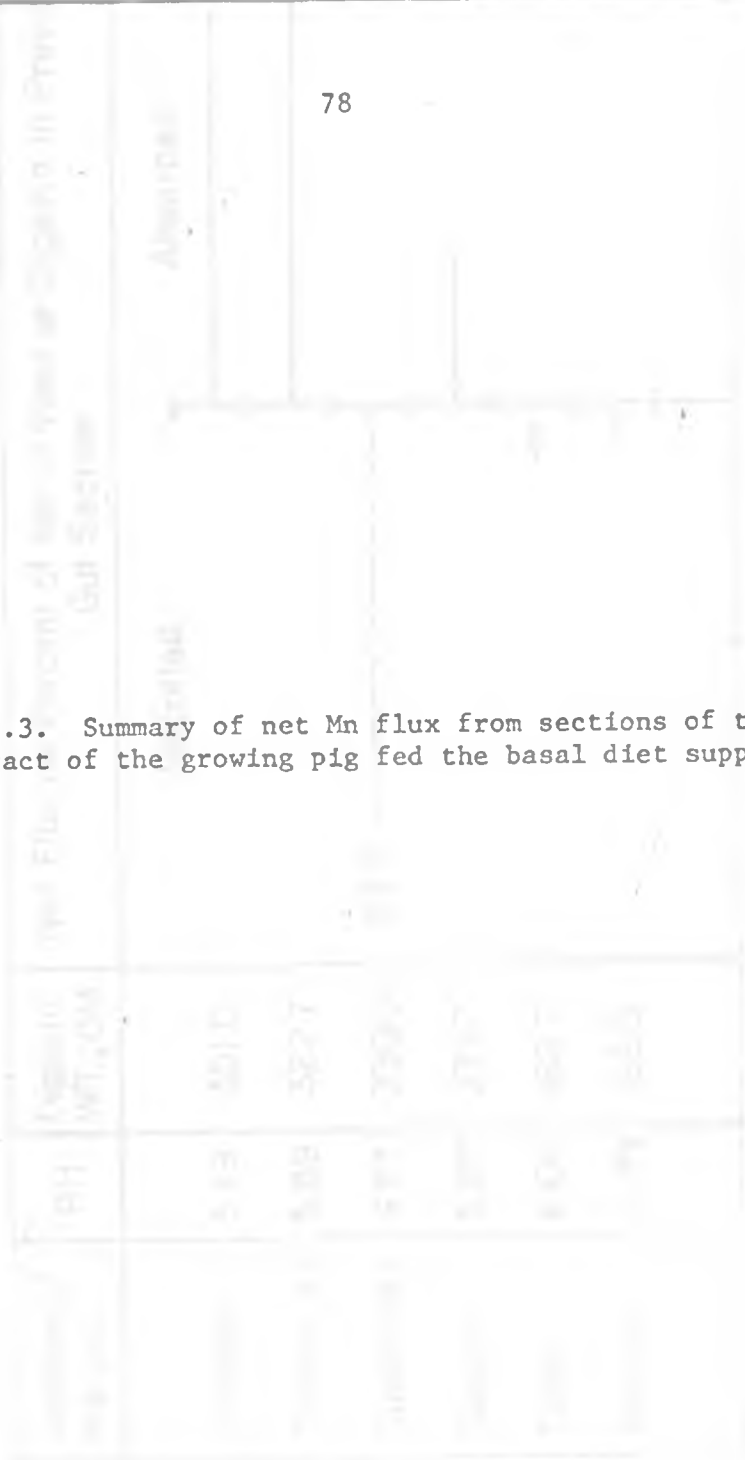


Figure 2.2.

Figure 2.3. Summary of net Mn flux from sections of the gastrointestinal tract of the growing pig fed the basal diet supplemented with $MnCO_3$.



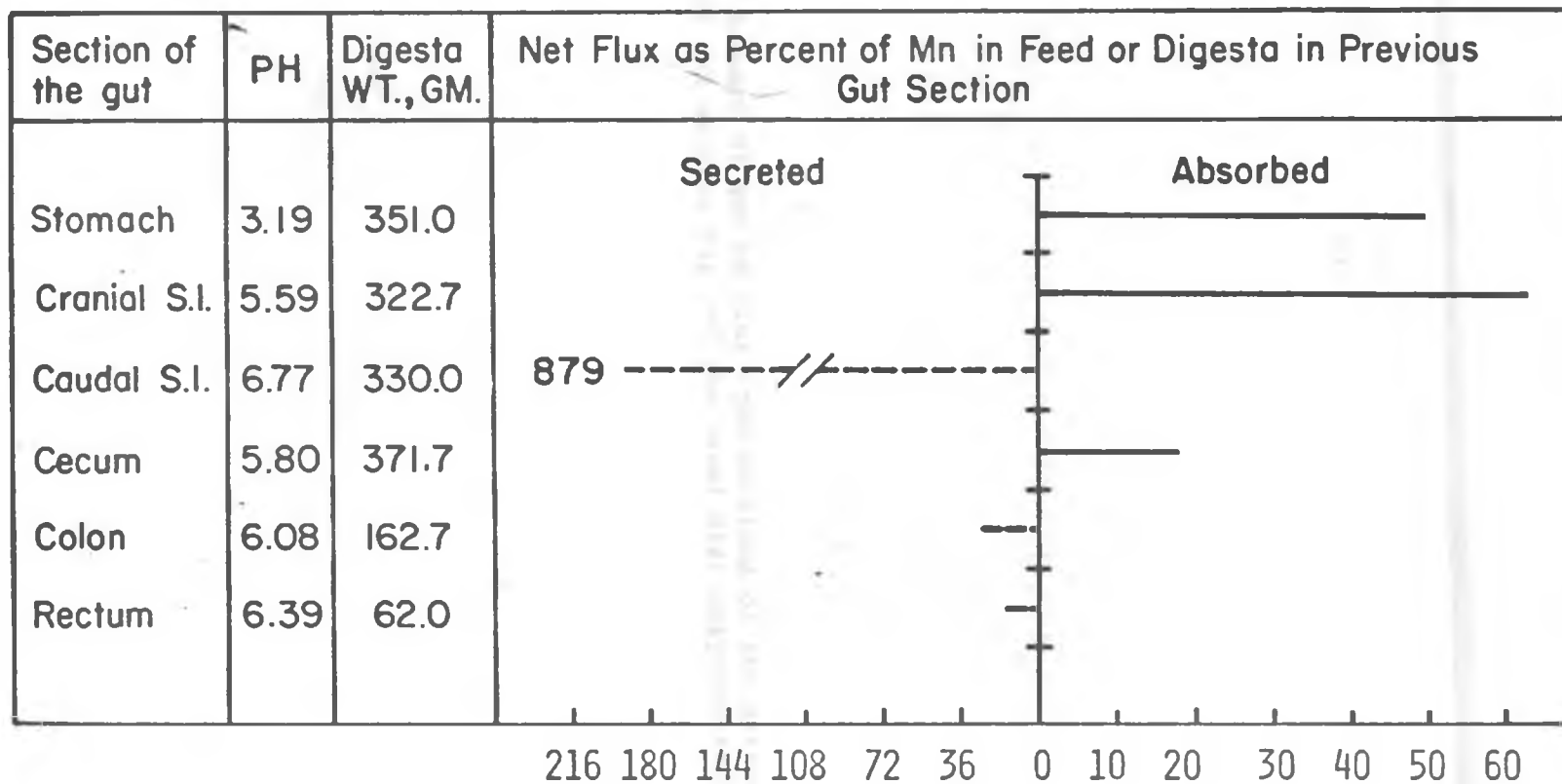
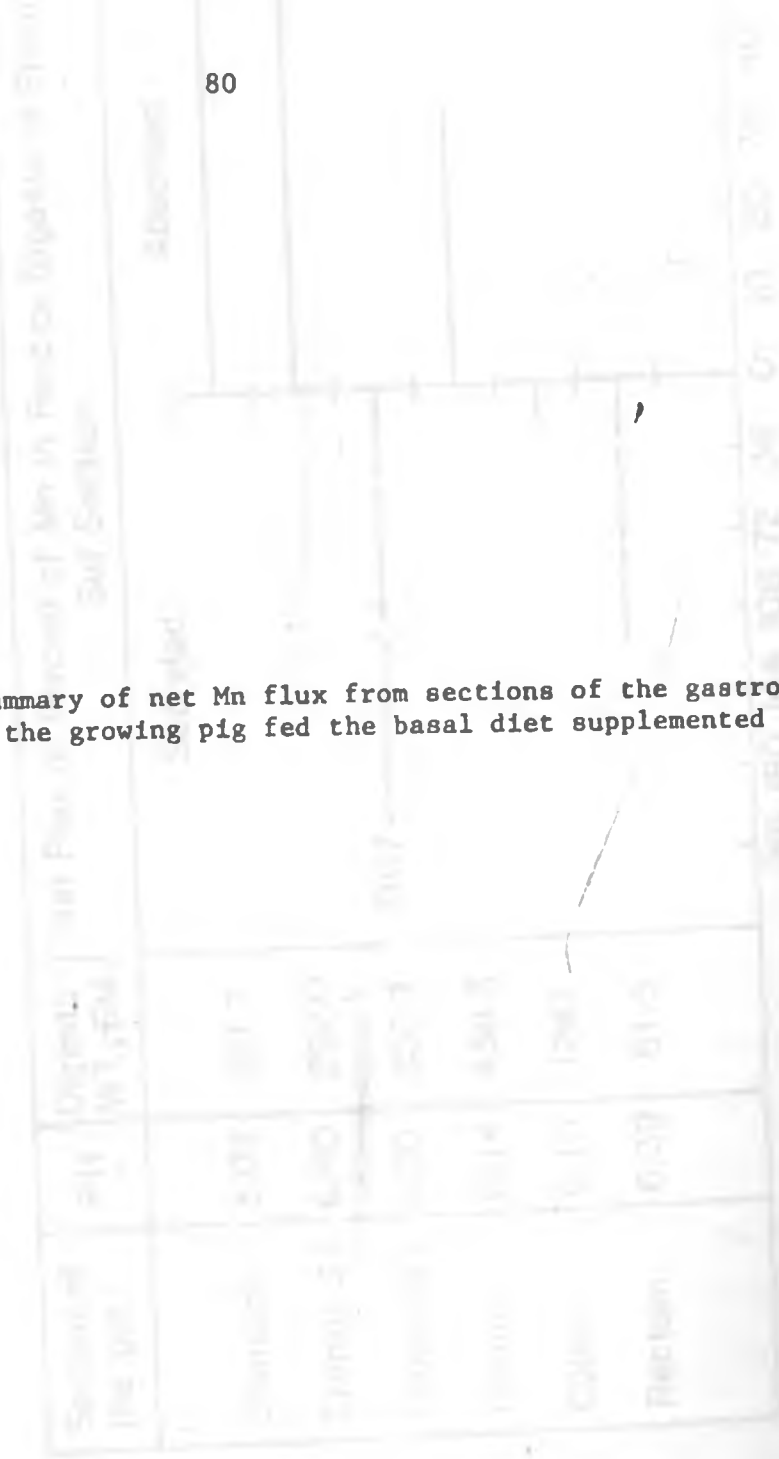


Figure 2.3.

Figure 2.4. Summary of net Mn flux from sections of the gastrointestinal tract of the growing pig fed the basal diet supplemented with MnO.



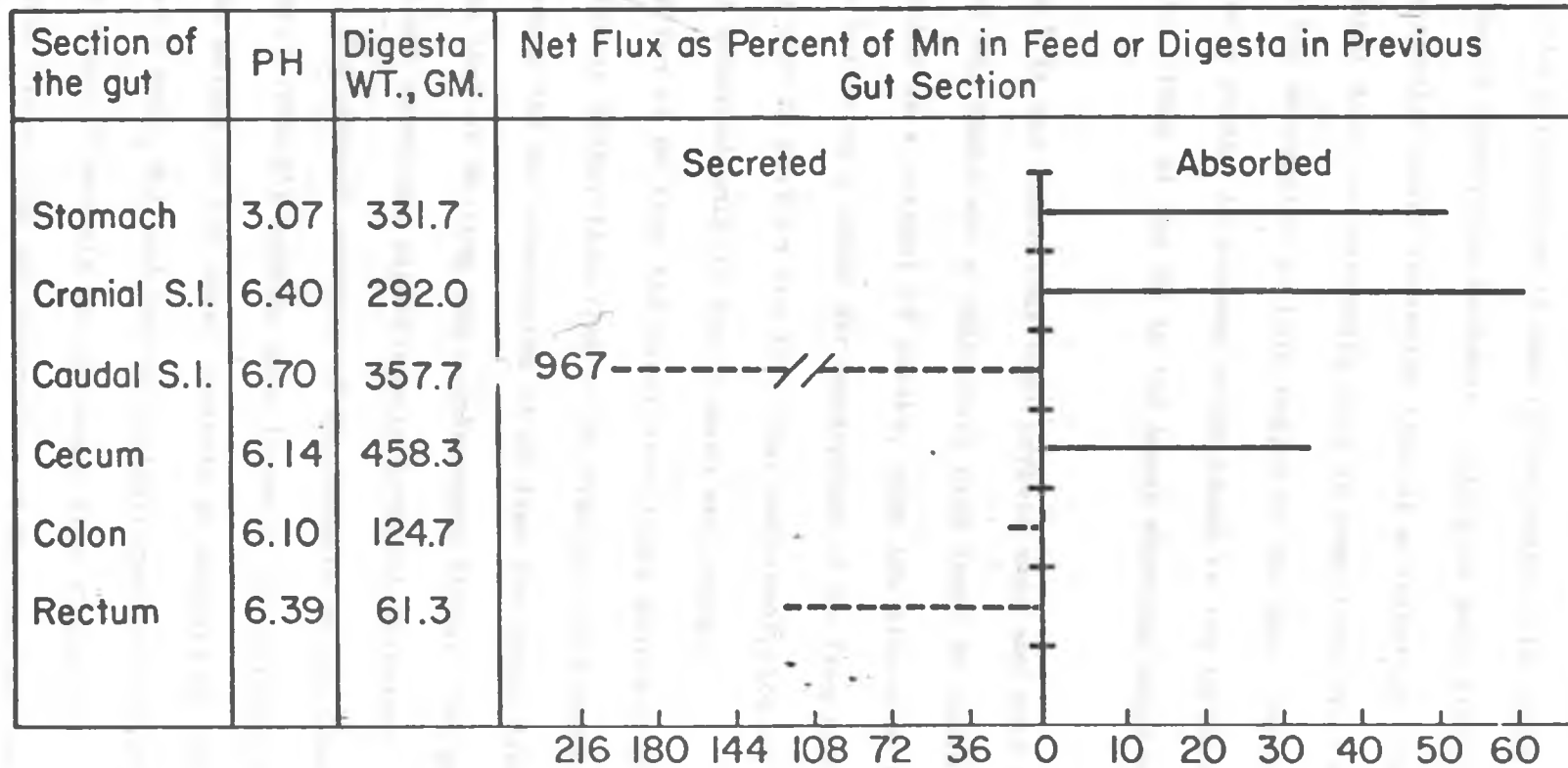


Figure 2.4.

cranial small intestine, being more acid in the case of pigs fed the supplemented diets than those fed the basal diet (6.08 vs 6.52). This might account for the differences in absorption rates. It could be postulated that the Mn absorption mechanism might be more efficient at a lower pH in the cranial small intestine than at a higher pH. Manganese from the basal diet is presumably held in some form which makes it less available for absorption in this region of the gut. Davis *et al.* (1962) reported a factor in soybean which tends to tie up Mn and make it unavailable. Most of the Mn in the basal diet was supplied by soy protein.

In Experiment 1 it was shown that when soybean meal and corn were the sole sources of Mn, there was a relatively high fecal Mn excretion and a low Mn retention as a percent of intake, plus low tissue Mn concentrations. This indicated a lower net absorption of Mn from the basal diet, perhaps due in part to the fact that net absorption of Mn from the basal diet occurred only in the stomach and cecum.

The net absorption of Mn from the basal diet (feed sources) in the stomach was slightly higher than that of Mn from the supplemented diets, but in the cecum the net absorption of Mn from the basal diet was much higher than that of Mn from the supplemented diets. The gut fill in the stomach and cecum was significantly ($P < 0.05$) different between treatments. The stomach contents of the animals on the basal diet were significantly ($P < 0.05$) heavier than those of the animals on other diets. The wet weight of the cecal contents of animals on the diets supplemented with $\text{MnSO}_4 \cdot \text{H}_2\text{O}$ and MnO were significantly ($P < 0.05$ or $P < 0.01$) heavier than those of animals on the basal diet (Table 18). Within the supplemented diets, the net absorption of Mn from the diet supplemented with $\text{MnSO}_4 \cdot \text{H}_2\text{O}$ was highest and that from the diets

supplemented with MnCO_3 lowest in both the stomach and cecum. In the cranial small intestine, the net absorption of Mn from the diets supplemented with MnCO_3 was highest while that from the diets supplemented with $\text{MnSO}_4 \cdot \text{H}_2\text{O}$ was lowest. The net absorption of Mn from diets supplemented with MnO was intermediate in the stomach, cranial small intestine and cecum.

The net secretion of Mn was significantly ($P < 0.01$) higher on the supplemented diets when compared to the basal diet in the caudal small intestine, slightly higher in the rectum but slightly lower in the colon. Within the supplemented diets, the net Mn secretion was significantly ($P < 0.01$) higher on diets supplemented with MnO and MnCO_3 than on diets supplemented with $\text{MnSO}_4 \cdot \text{H}_2\text{O}$ in the caudal small intestine (Table 17). The net Mn secretion on diets supplemented with MnO or MnCO_3 was not significantly different. Net Mn secretion in the rectum was much greater on diets supplemented with $\text{MnSO}_4 \cdot \text{H}_2\text{O}$ and MnO than on the diets supplemented with MnCO_3 . In the colon, the net Mn secretion was highest on diets supplemented with $\text{MnSO}_4 \cdot \text{H}_2\text{O}$ and lowest on diets supplemented with MnO. Hendricks (1967) reported net Mn secretion into the cranial small intestine, caudal small intestine, and colon when pigs were fed a 16% soy protein diet supplemented with Mn from $\text{MnSO}_4 \cdot \text{H}_2\text{O}$. He showed a high net Mn secretion in the colon, a low net Mn secretion in the cranial and caudal small intestine, and an absence of Mn secretion in the rectum.

C. Experiment 3: The effect of high level Ca and P supplementation and an inverse Ca-P ratio on Mn utilization by the growing pig

The effects of different Ca to P ratios, Ca and P levels and Mn supplementation on physical and chemical composition of pig tissues are summarized in Tables 20, 21, and 22. A 2 to 1 ratio of Ca to P significantly ($P < 0.05$) depressed rib Mn content and slightly increased

heart, pancreas and serum Mn as compared to a 1 to 2 ratio. The increased levels of Ca and P supplementation significantly ($P < 0.01$) increased rib and pancreas Mn concentration but also significantly ($P < 0.01$) depressed Mn concentration of the metacarpal bone. The high levels of Ca and P slightly increased heart Mn and significantly ($P < 0.05$) increased serum Mn concentration. This finding is at variance with those of Hawkins *et al.* (1955), who reported a suppression of serum Mn with high Ca and P intakes in cattle. Lassiter *et al.* (1970) reported that rats given a 0.9 percent P in the diet caused significantly higher retention of orally administered ^{54}Mn than did 0.4 percent P.

Mn supplementation increased heart Mn significantly ($P < 0.01$) and slightly increased rib, pancreas and serum Mn, kidney Ca and serum inorganic P. High Mn levels in the diets significantly ($P < 0.05$) depressed rib Ca and Mg and slightly depressed metacarpal Mn, P and Mg, serum Ca, and rib and kidney P. The slight increase of Mn in serum following increased Mn intakes is in harmony with other observations in swine (Plumlee *et al.*, 1956), and with some reports in cattle (Rojas *et al.*, 1965; Hawkins *et al.*, 1955), in rats (Ugnenko, 1972), and in poultry (Bolton, 1955). Other reports, however, are in contrast to these findings (Krieg, 1966; Bentley and Phillips, 1951a). In poultry, Nielsen and Madsen (1942) observed no appreciable difference in blood Ca concentration due to dietary Mn levels, and reported that acid soluble P of the blood and inorganic P of plasma did not differ significantly due to Mn supplementation.

Although Mn has been implicated in bone formation (Underwood, 1971), the results of this study showed no significant effect of Mn supplementation on rib and metacarpal physical measurements, breaking strength and related parameters (Tables 20 and 21). Dietary Mn levels did not

TABLE 20. EFFECT OF THE CA-P RATIO, CA AND P LEVELS AND MN SUPPLEMENTATION ON PHYSICAL AND CHEMICAL PARAMETERS OF THE FIRST, LEFT RIB

Item	Ca-P ratio		Ca-P level		Suppl. Mn, ppm		SE
	0.5	2.0	1X	2X	0	40	
Physical measurements							
Weight, g (fresh basis)	12.5 ^{2/}	10.9	9.4 ^{1/}	14.0	11.4	12.0	0.36
External diameter (B), cm	1.71	1.66	1.60 ^{1/}	1.77	1.70	1.67	0.04
Internal diameter (d), cm	.58 ^{1/}	.51	.57	.52	.56	.53	0.02
Specific gravity (fresh basis)	1.22	1.21	1.15 ^{1/}	1.27	1.22	1.21	0.01
Inertia, cm ⁴	.24	.22	.18 ^{1/}	.28	.24	.21	0.02
Breaking moment, kg	26.8	28.8	15.1 ^{1/}	40.5	28.4	27.2	1.04
Chemical measurements³							
Ash content, %	56.9	52.9	50.5	59.3	55.5	54.3	0.42
Calcium, %	23.0 ^{1/}	21.7	20.5 ^{1/}	24.2	22.6 ^{1/}	22.0	0.18
Phosphorus, %	10.9 ^{1/}	9.9	9.5 ^{1/}	11.2	10.5	10.3	0.09
Magnesium, %	.36 ^{1/}	.31	.30	.36	.35 ^{1/}	.32	0.01
Manganese, ppm	1.07 ^{1/}	.99	.95 ^{1/}	1.11	1.02	1.04	0.03

¹Numbers on the same line under the same subheading, i.e., Ca-P ratio, Ca-P level, or Mn supplementation level, are significantly (P<0.05) different.

²P<0.01.

³Expressed on dry, fat-free bone.

TABLE 21. EFFECT OF THE CA-P RATIO, CA AND P LEVELS AND MN SUPPLEMENTATION ON PHYSICAL, CHEMICAL AND HISTOPATHOLOGICAL PARAMETERS OF THE 5TH METACARPAL

Item	Ca-P ratio		Ca-P level		Suppl. Mn, ppm		SE
	0.5	2.0	1X	2X	0	40	
<u>Physical measurements</u>							
Weight, g (fresh basis)	14.6	13.9	12.8 ^{2/}	15.8	14.4	14.1	0.39
External diameter (D), cm	1.38	1.38	1.33 ^{2/}	1.43	1.39	1.37	0.02
External diameter (B), cm	1.36	1.32	1.28 ^{2/}	1.40	1.35	1.34	0.02
Internal diameter (b), cm	.97	.96	.94	.99	.99 ^{1/}	.94	0.02
Specific gravity (fresh basis)	1.23	1.21	1.18 ^{3/}	1.25	1.22	1.21	0.01
Inertia, cm ⁴	1.04	1.12	.87 ^{3/}	1.30	1.08	1.09	0.09
Breaking moment, kg	42.6	43.0	29.2 ^{2/}	56.3	44.3	41.3	1.80
Stress, kg/cm ²	299	263	229 ^{2/}	333	292	270	20.01
Elasticity, ₂ 1000 kg/cm	13.7 ^{2/}	7.3	10.4	10.7	10.6	10.5	0.57
<u>Chemical measurements</u> ^{3/}							
Ash content, %	58.1 ^{2/}	54.5	52.7 ^{2/}	59.9	56.8	55.8	0.57
Phosphorus, %	10.2 ^{2/}	9.4	9.1 ^{3/}	10.5	9.9	9.8	0.12
Manganese, ppm	.99 ^{2/}	1.04	1.09 ^{2/}	.94	.96 ^{1/}	1.07	0.03
Magnesium, %	.34 ^{2/}	.29	.29 ^{2/}	.34	.32	.30	0.01
<u>Bone histology</u>							
Epiphysis _{4/}	1.50	1.63	1.3 ^{2/}	1.81	1.75 ^{1/}	1.38	0.12
Thickness of compact bone of the diaphysis _{5/}	96.2 ^{2/}	51.8	45.8 ^{2/}	102.3	91.6 ^{1/}	56.4	10.59
Thickness of epiphyseal cartilagenous plate _{5/}	65.6	64.1	70.3 ^{2/}	59.4	62.5	67.2	2.33

¹Numbers on the same line under the same subheading, i.e., Ca-P ratio, Ca-P level, or Mn supplementation level, are significantly (P<0.05) different.

²P<0.01.

³Expressed on dry, fat-free bone.

⁴1 = normal; 2 = very slight change; 3 = slight change; 4 = moderate change; 5 = severe change.

⁵Percent of a 20X objective by 10X ocular field.

TABLE 22. EFFECT OF THE CA-P RATIO, CA AND P LEVELS AND MN SUPPLEMENTATION ON ORGAN AND BLOOD COMPOSITION

Item	Ca-P ratio		Ca-P level		Suppl. Mn, ppm		SE
	0.5	2.0	1X	2X	0	40	
<u>Organ composition, dry matter basis</u>							
Kidney calcium, ppm	258 ^{2/}	315	283	290	274	299	10.57
Kidney phosphorus, %	1.35 ^{1/}	1.40	1.38	1.37	1.39 ^{2/}	1.36	0.02
Heart manganese, ppm	2.80	2.86	2.76 ^{2/}	2.90	2.43 ^{2/}	3.23	0.12
Pancreas manganese, ppm	6.03	6.11	5.70 ^{2/}	6.45	5.90	6.25	0.17
<u>Blood parameters</u>							
Hemoglobin, mg/100 ml							
2nd bleeding	11.2	11.1	11.7 ^{1/}	10.6	11.1	11.2	0.23
3rd bleeding	11.7	11.8	12.7 ^{1/}	10.9	11.7	11.8	0.31
Hematocrit, %							
2nd bleeding	34.5	33.6	35.5 ^{1/}	32.6	34.0	34.1	0.56
3rd bleeding	35.1	36.2	38.8 ^{1/}	32.5	35.5	35.9	0.91
Serum alkaline phosphatase, Sigma units							
2nd bleeding	5.0 ^{1/}	6.2	6.1	5.2	5.4	5.8	0.27
Serum Mn, mcg/100 ml							
1st bleeding	1.94	2.32	1.85 ^{1/}	2.40	1.96	2.25	0.09
Serum Ca, mg/100 ml							
3rd bleeding	8.9 ^{1/}	10.6	9.9	9.7	10.0	9.6	0.04
Serum inorganic P, mg/100 ml							
1st bleeding	9.9 ^{2/}	12.1	11.6 ^{2/}	10.4	11.1	10.9	0.01
3rd bleeding	9.7 ^{2/}	7.2	8.4	8.5	8.2	8.6	0.03

¹Numbers on the same line under the same subheading, i.e., Ca-P ratio, Ca-P level, or Mn supplementation level, are significantly ($P < 0.05$) different.

² $P < 0.01$.

affect Ca and P metabolism in the bones, which is in accord with the findings of Parker *et al.* (1955) that Mn intake did not affect the quantities of radioactive Ca and P deposited in the bones of chicks. Ellis *et al.* (1947) reported no significant difference in breaking strength or Ca content of the humerus of rabbits due to dietary Mn levels, but this report is contrary to the reports of Smith *et al.* (1944), who showed a significant decrease in weight, density, length, breaking strength and ash content of the ulna and humerus of rabbits due to dietary Mn levels. Hemoglobin and hematocrit values were not affected by Mn supplementation. Sawney and Kehar (1958) reported similar findings in cattle. Similarly Mn supplementation did not significantly affect serum Ca, inorganic P, Mg, Mn and alkaline phosphatase. This work is in accord with the findings of Hawkins *et al.* (1955), Fain *et al.* (1952) and Sawney and Kehar (1958) in cattle, but Hawkins *et al.* (1955) and Fain *et al.* (1952) showed a depression of serum Mg due to Mn supplementation.

The slight increase in serum alkaline phosphatase activity due to Mn supplementation has also been reported by some (Lassiter *et al.*, 1970) but not others (Leibholz *et al.*, 1962). There was a time of sampling effect on serum Ca, inorganic P, Mn, Mg and alkaline phosphatase. In general, their levels increased within the first month except in the case of P, which declined drastically throughout the experimental period on diets 5 and 6 with low P content (0.35%) derived exclusively from corn and soybean meal. The serum Mg values tended to be higher on these same diets than on other diets.

A two-way interaction between ratio and level of Ca and P on Mn metabolism is shown in Tables 23 and 24. This interaction was significant ($P < 0.01$ or $P < 0.05$) relative to levels of serum, liver and pancreas

TABLE 23. EFFECT OF THE INTERACTION BETWEEN CA-P RATIO AND LEVEL OF CA AND P ON MN METABOLISM AND SOME OTHER BONE PARAMETERS⁵

Item	Ca-P Ratio				SE
	0.5		2.0		
	Ca-P level		Ca/P level		
	1X	2X	1X	2X	
<u>Rib, 1st left</u>					
Weight, g (fresh basis) ^{2/}	11.6	13.4	7.2	14.7	0.51
External diameter (D), cm ^{1/}	.75	.72	.65	.76	0.03
Specific gravity ^{2/} (fresh basis)	1.21	1.24	1.10	1.31	0.01
Inertia, cm ^{4 2/}	.22	.26	.13	.31	0.02
Breaking moment, k ^{2/}	23.0	30.7	7.3	50.2	1.60
Ash content, % ^{2/}	55.1	58.6	45.9	59.9	0.59
Calcium, % ^{2/}	22.1	23.8	18.9	24.5	0.26
Phosphorus, % ^{2/}	10.6	11.1	8.4	11.3	0.13
Magnesium, % ^{2/}	.39	.34	.22	.39	0.01
<u>Metacarpal, 5th</u>					
Weight, g ^{2/} (fresh basis)	14.8	14.4	10.7	17.2	0.56
External diameter (D), cm ^{2/}	1.39	1.36	1.27	1.49	0.03
External diameter (B), cm ^{2/}	1.36	1.37	1.20	1.44	0.03
Specific gravity ^{2/} (fresh basis)	1.22	1.23	1.14	1.28	0.01
Inertia, cm ^{4 2/}	1.16	.93	.58	1.67	0.13
Breaking moment, kg ^{2/}	41.7	43.4	16.7	69.3	2.55
Ash content, % ^{2/}	57.1	59.1	48.2	60.7	0.81
Calcium, % ^{2/}	21.0	20.8	18.2	21.2	0.71
Phosphorus, % ^{2/}	10.1	10.4	8.1	10.7	0.16
Magnesium, % ^{2/}	.36	.31	.22	.36	0.01
Manganese, ppm ^{2/}	1.00	.98	1.18	.91	0.04
Elasticity, 1000 kg/cm ^{2 2/}	15.33	12.12	5.44	9.17	0.81
<u>Bone histology</u>					
Diaphysis <u>3/1/</u>	1.38	1.25	1.13	2.13	0.24
Epiphysis <u>3/2/</u>	1.63	1.00	1.38	2.25	0.17
Thickness of epiphyseal cartilagenous plate <u>4/2/</u>	65.63	75.00	65.63	64.06	3.29

¹Significant (P<0.05).

²P<0.01.

³₁=normal; ₂=very slight change; ₃=slight change; ₄=moderate change; ₅=severe change.

⁴Percent of a 20X objective by 10X ocular field.

⁵Chemical constituents expressed on dry, fat-free bone.

TABLE 24. EFFECT OF THE INTERACTION BETWEEN CA-P RATIO AND LEVEL OF CA AND P ON ORGAN AND SERUM MN METABOLISM, AND SOME OTHER MEASURES

Item	Ca-P Ratio				SE
	0.5		2.0		
	Ca-P level 1X	2X	Ca/P level 1X	2X	
<u>Organ chemical composition</u> (dry matter basis)					
Liver Mn, ppm ^{1/}	13.76	13.42	12.93	15.50	0.58
Kidney P, % ^{1/}	1.39	1.31	1.37	1.43	0.03
Pancreas Mn, ppm ^{1/}	5.41	6.66	5.99	6.23	0.23
<u>Serum parameters</u>					
Serum alkaline phosphatase (Sigma units)					
1st bleeding ^{1/}	5.88	5.00	4.45	6.20	0.34
2nd bleeding ^{1/}	5.08	6.83	7.98	6.15	0.39
3rd bleeding ^{1/}	4.75	5.25	7.40	5.08	0.38
Serum Mn, mcg/100 ml ^{1/}					
3rd bleeding	2.29	2.89	3.21	2.47	0.10
Serum inorganic P, mg/100 ml					
1st bleeding ^{2/}	9.7	10.1	13.5	10.7	0.01
3rd bleeding ^{2/}	10.4	9.0	6.4	8.0	0.04

¹Significant (P<0.05).

²P<0.01.

Mn, and metacarpal Mn levels. High levels of Ca and P in a 1 to 2 ratio depressed pancreas Mn content appreciably and slightly depressed liver and metacarpal Mn content, but increased serum Mn. High levels of Ca and P supplementation in a 2 to 1 ratio increased liver and pancreas Mn but depressed the serum and metacarpal Mn concentration. Hawkins *et al.* (1955) reported that high intakes of Ca and P tended to suppress blood Mn.

The interaction between the ratio of Ca to P and Mn level on Mn and Mg metabolism is shown in Table 25. Feeding Mn along with Ca and P in a 1 to 2 ratio slightly depressed liver Mn but increased it when Ca and P were given in a 2 to 1 ratio. Although Mn supplementation alone did not significantly affect metacarpal Mg, feeding Ca and P in a 1 to 2 ratio did increase metacarpal Mg significantly ($P < 0.05$). The interaction between Ca to P ratio and diet Mn level upon serum Mn concentration was not significant, contrary to Hawkins' *et al.* (1955) findings in cattle.

The interaction between diet Ca and P levels and Mn levels on some bone and serum parameters is shown in Table 26. This interaction was significant ($P < 0.05$) relative to rib ash content, Ca and P, metacarpal Mn levels and serum inorganic P. With low Ca and P levels, Mn supplementation increased metacarpal Mn and serum inorganic P but depressed rib ash, Ca and P content. Mn supplementation, coupled with high Ca and P levels, did not affect metacarpal Mn, slightly increased rib ash, Ca and P, but depressed serum inorganic P. The interaction between diet Ca and P levels and Mn levels was not significant relative to Mg levels in the tissues. The significant effect of Mn supplementation on the metacarpal internal vertical diameter, rib Mg and heart Mn levels was suppressed when high Ca and P levels were also given in the diets.

TABLE 25. EFFECT OF THE INTERACTION BETWEEN CA-P RATIO AND MN LEVELS ON MN AND MG METABOLISM

Item	Ca-P ratio				SE
	0.5		2.0		
	Mn-level 0	Mn-level 40	Mn-level 0	Mn-level 40	
<u>Metacarpal, 5th left</u>					
Mg, % <u>1/</u>	.36	.31	.29	.29	0.01
<u>Liver</u>					
Mn, ppm <u>1/</u>	13.8	13.4	12.9	15.5	0.58
<u>Bone histology</u>					
Epiphysis <u>2/1/</u>	1.88	1.63	1.13	1.63	0.17

¹Significant (P<0.05).

²1 = normal; 2 = very slight change; 3 = slight change; 4 = moderate change; 5 = severe change.

TABLE 26. EFFECT OF THE INTERACTION BETWEEN CA-P LEVELS AND MN LEVELS ON SOME PHYSICAL PARAMETERS

Item	Ca-P level				SE
	1X		2X		
	Mn-level 0	40	Mn-level 0	40	
<u>Rib, 1st left</u>					
Ash content, % ^{1/}	51.81	49.13	59.14	59.42	0.59
Calcium, % ^{1/}	21.08	19.86	24.17	24.18	0.26
Phosphorus, % ^{1/}	9.77	9.31	11.15	11.22	0.13
<u>Metacarpal, 5th left</u>					
Manganese, ppm ^{1/}	.98	1.19	.94	.94	0.04
Serum inorganic P, mg/100 ml ^{1/}					
3rd bleeding	7.48	9.25	9.00	7.98	0.60
<u>Bone histology</u>					
Thickness of epiphyseal cartilagenous plate ^{2/1/}	71.88	53.13	68.75	65.63	3.29

¹Significant (P<0.05).

²Percent of a 20X objective by 10X ocular field.

Table 27 summarizes a three-way interaction between level of Ca and P, ratios of Ca to P and Mn levels on some measured parameters. The interaction was significant ($P < 0.05$) with respect to rib and serum Mn content, pancreas dry matter, and metacarpal Mg content and elasticity. With low Ca and P levels in a 1 to 2 ratio, Mn supplementation increased serum and metacarpal Mn values, slightly increased pancreas dry matter and did not change metacarpal Mn and elasticity. High Ca and P levels in a 1 to 2 ratio with Mn supplementation depressed rib Mn, metacarpal Mg, pancreas dry matter and serum Mn, but increased metacarpal elasticity. With low Ca and P levels in a 2 to 1 ratio, Mn supplementation depressed rib Mn content, pancreas dry matter and serum Mn, and increased metacarpal elasticity but did not change metacarpal Mg. With high Ca and P levels in a 2 to 1 ratio, Mn supplementation increased rib and serum Mn levels and pancreas dry matter, and slightly increased metacarpal Mg but depressed metacarpal elasticity. The deleterious effects on weight gain and feed efficiency of a low dietary P level (0.35%), derived exclusively from corn and soybean meal, were much more pronounced than the effects of excessive dietary levels of Ca and P or of an inverse Ca to P ratio, regardless of dietary Mn concentration.

1. Histopathology¹

a. Histopathologic examination. The histology of the epiphyseal cartilagenous plate was considered normal in all bones from all pigs regardless of treatment group. There was some persistence of cartilage

¹Performed by Dr. K. K. Keahey, Department of Pathology, Michigan State University.

TABLE 27. EFFECT OF THE INTERACTION BETWEEN CA-P RATIO, CA AND P LEVEL AND MN LEVEL ON SOME MEASURED PARAMETERS

Category	0.5				2.0				SE
	1X		2X		1X		2X		
	0	40	0	40	0	40	0	40	
<u>Rib, 1st left</u> Mn conc., ppm ^{1/}	0.98	1.06	1.15	1.10	0.91	0.85	1.05	1.15	.05
<u>Metacarpal, 5th left</u> Mg content, % ^{2/}	0.36	0.37	0.37	0.26	0.22	0.21	0.35	0.38	.02
Elasticity, 1000 kg/cm ² ^{1/}	15.4	15.3	11.1	13.2	5.0	5.9	11.0	7.4	1.15
Pancreas, dry matter % ^{1/}	28.4	29.6	28.2	25.6	28.1	26.7	27.4	28.4	.97
Serum Mn, mcg/100 ml ^{1/} 1st bleeding	1.25	2.01	2.36	2.10	2.21	1.95	2.05	3.00	.24

¹Significant (P<0.05).

²P<0.01.

in the spicules. The most striking histologic change was in the diaphysis of the bone. Pigs that had a thickness of compact bone equivalent to 70% of the diameter of the microscope field¹ or greater had normal histologic structures. Those with less than 70% had incomplete ossification of the Haversian canal system. The center of the Haversian canal was occupied by a blood vessel which was surrounded by a significant zone of loose connective tissue and osteoid. The compact bone, therefore, appeared histologically like spongy bone instead of normal compact bone. Examination of a record kept at the time of trimming of tissues indicated that most of the soft bones had either a very thin layer of compact bone in the diaphysis or an absence of compact bone. It was concluded that the condition observed in the bones from pigs from some dietary regimes were not changes of rickets but were changes in which there was a failure of production of compact bone in the region of the diaphysis. This conclusion agrees with that of Smith *et al.* (1944), who, using X-ray pictures and microscopic studies, reported bone changes in Mn deficiency were distinctly different from those seen in rickets. Neher *et al.* (1956) also reported generalized bone rarefaction in Mn deficient pigs.

b. Statistical inferences. Manganese supplementation was associated with a significantly ($P < 0.05$) higher histological normalcy score of the epiphysis as compared to no Mn supplementation. The histology of the diaphysis was not affected by diet Mn level. Animals on high Mn level diets had significantly ($P < 0.05$) less compact bone in the diaphysis, but these levels did not significantly affect the thickness of the epiphyseal cartilagenous plate (Table 21). The ratio of

¹10X eyepiece and 20X objective.

Ca to P, level of Ca and P and the interaction between the two had a significant effect (Table 23) on the bone histologic structure as reported in many studies of Ca and P on bone metabolism (Underwood, 1971). The interaction between the Ca-P ratio and Mn levels was significant ($P < 0.05$) relative to the histologic structure of the epiphysis (Table 25). With a 2 to 1 Ca to P ratio, Mn supplementation increased the incidence of change in bone histologic structure. Mn interaction with the Ca and P levels significantly ($P < 0.05$) affected the thickness of epiphyseal cartilagenous plate (Table 26). Production of the normal histologic structure of the epiphyseal bone by Mn supplementation is in agreement with the established concept that Mn is involved in bone formation (Leach, 1967; Tsai and Everson, 1967). However, the deleterious effects of a low dietary P level (0.35%) derived exclusively from corn and soybean meal on the histology of bone were much more pronounced than the effects of high dietary levels of Ca and P or of an inverse Ca to P ratio, regardless of dietary Mn supplementation.

D. Experiment 4: The manganese requirement of the baby pig from sows fed a low manganese diet

Table 28 summarizes the results of the analysis of variance. Average daily gain was significantly ($P < 0.01$) different between treatments but feed efficiency was not. Absolute Mn intake, and absolute fecal and urinary Mn excretion, were significantly ($P < 0.01$ or $P < 0.05$) different between treatments, but absolute Mn retention was not. Mn retention, fecal and urinary Mn excretion, all as percent of intake, were significantly ($P < 0.01$ or $P < 0.05$) different between dietary treatments. Of the serum parameters studied, serum Mg levels on the 28th day

TABLE 28. ANALYSIS OF VARIANCE OF EFFECT OF DIETARY MN LEVEL ON GROWTH, BALANCE AND BLOOD MEASURES

Item		Mean	Min.	Max.	P-level of F-statistic
<u>Serum parameters</u>					
Magnesium, mg/100 ml	9 ¹	2.38	2.00	2.85	0.83
	21	2.57	1.87	3.05	0.49
	28	2.04	1.26	2.90	0.03
Alkaline phosphatase (SAP), Sigma units	9	5.8	3.4	8.4	0.04
	21	7.0	5.3	10.3	0.10
	28	6.2	3.0	10.7	0.11
Manganese, mcg/100 ml	9	1.20	.90	1.90	0.348
	21	0.94	.60	1.50	0.129
	28	0.90	.40	1.70	0.005
<u>Growth data</u>					
Average daily gain, g		253	206	299	0.02
Feed/gain		1.33	1.18	1.45	0.238
<u>Balance data</u>					
Mn intake, mg/day		1.82	0.18	4.06	<0.0005
Mn excretion, mg/day					
Fecal		1.40	0.24	3.45	0.006
Urinary		0.015	0.01	0.02	0.049
Mn retention, mg/day		.44	-.30	1.59	0.240
Mn retention, % of intake		3.6	-72	73	0.016
Mn excretion, % of intake					
Fecal		94	26	167	0.02
Urinary		2.1	0.3	5.6	<0.0005

¹Days on experimental diets.

of the experiment, serum alkaline phosphatase (SAP) on the 9th day and serum Mn on the 28th day showed significant ($P < 0.01$ or $P < 0.05$) treatment effects.

There were no treatment effects on serum Mg levels 9 days after the start of the experiment, a slight rise on the 2.67 ppm Mn diet (diet 2) after 21 days, and a significant ($P < 0.05$) drop on the basal diet by the end of the experiment (Table 29, Figure 4.1). Manganese supplementation appeared to sustain normal serum Mg levels. The SAP levels were high on the 6.34 ppm Mn diet (diet 3) and low on the basal diet (0.46 ppm Mn) and diet 2 throughout the duration of the experiment. On the 9th and 21st days, the animals on the basal diet had higher SAP levels than those on diet 2 and those values dropped slightly by the end of the experiment (Table 29, Figure 4.2). The serum Mn values were significantly ($P < 0.01$) lower on the basal diet than on the diet containing 6.34 ppm Mn on the 28th day of the experiment. There was an appreciable and consistent drop of serum Mn values on the basal diet and diet 2 as the experiment progressed. In animals on the 6.34 ppm Mn diet, serum Mn values dropped slightly by the 21st day but returned to previous levels by the 28th day (Table 29, Figure 4.3).

Serum Mn response to dietary Mn levels has been reported in swine fed practical diets (Plumlee *et al.*, 1956; Newland and Davis, 1961), in ruminants (Hawkins *et al.*, 1955; Rojas *et al.*, 1965), and in chickens (Bolton, 1955). A rise in SAP levels in response to Mn supplementation has also been reported by some (Wachtel *et al.*, 1943; Lassiter *et al.*, 1970) but not by others (Leibholz *et al.*, 1962). The big drop of serum Mg and serum Mn levels by the 28th day on the basal diet may indicate that when animals are fed low Mn diets (0.46 ppm) for a long period their Mn stores are depleted and Mg substitutes in the functions that

TABLE 29. THE EFFECT OF DIETARY MN LEVELS ON SERUM AND GROWTH PARAMETERS

Item		Diets		
		1	2	3
Mn conc., ppm		0.46	2.67	6.34
No. of pigs		4	4	4
Serum Mg, mg/100 ml	9 ³	2.44	2.38	2.32
	21	2.38	2.72 ^{1/}	2.62
	28	1.47	2.40 ^{1/}	2.25
SAP, ⁴ Sigma units	9	5.6	4.6	7.3 ^{1/}
	21	7.0	5.8	8.3
	28	4.5	5.5	8.6
Serum Mn, mcg/100 ml	9	0.98	1.33	1.31
	21	0.73	0.95	1.15
	28	0.41	0.88	1.28 ^{2/}
Average daily gain, g		224	283 ^{1/}	251
Feed/gain		1.37	1.27	1.35

¹Significantly ($P < 0.05$) greater than the least value on the same line.

² $P < 0.01$.

³Days on experimental diets.

⁴Serum alkaline phosphatase.

usually involve Mn. Magnesium and Mn can replace each other in a number of enzymatic reactions since they have similar properties (Lehninger, 1950, 1970). Johnson (1943) found a lower body Mn content of pigs born of sows depleted of Mn. In this study, the correlations between serum Mg and serum Mn were positive and significant ($P < 0.05$) (Table 32).

Average daily gain was significantly ($P < 0.05$) greater on diet 2 than on diet 1 (Table 29, Figure 4.4). The average daily gain was also higher on diet 3 than on diet 1, but the difference was not statistically significant. Food efficiency was greater on diet 2 than on the other diets but those differences were not statistically significant. Many reports have shown increased growth rate with optimal Mn supplementation in the species studied (Underwood, 1971). The results of this study support those of Leibholz *et al.* (1962), who concluded that Mn requirements for baby pig growth were very low. For maximal growth, these data would indicate that 3.0 ppm Mn in the diet are adequate. This level is approximately equal to that reported by Plumlee *et al.* (1956).

Mn intake, fecal Mn excretion and Mn retention all reflected the dietary level of Mn (Table 30, Figure 4.). The absolute Mn intakes of the three diets were significantly ($P < 0.01$) different; the absolute fecal Mn excretion on diet 3 was significantly ($P < 0.01$) greater than that on diets 1 and 2 but the difference between diets 1 and 2 was not statistically significant. Diets 2 and 3 resulted in significantly ($P < 0.05$) greater absolute urinary Mn excretion than diet 1 but the absolute urinary Mn excretion on diets 2 and 3 was equal and did not reflect the differing dietary Mn levels. Other workers have also reported this small but virtually constant urinary Mn excretion in other species irrespective of Mn ingested and the dominance of the feces as the principal excretion route of this element (Britton and Cotzias,

TABLE 30. RETENTION AND EXCRETION ROUTES OF MN FROM DIETS CONTAINING DIFFERENT LEVELS OF MN

Item	Diets		
	1	2	3
Mn conc., ppm	0.46	2.67	6.34
No. of pigs	4	4	4
Mn intake, mg/day	0.19	1.67 ^{2/}	3.63 ^{2/4/}
Mn excretion, mg/day			
Fecal	.30	1.05 ^{2/}	2.85 ^{2/4/}
Urinary	.01	.02 ^{2/}	.02 ^{2/}
Mn retention, mg/day	-.11	.60	.77
Mn retention, % of intake	-58	36 ^{1/}	22 ^{1/}
Mn excretion, % of intake			
Fecal	154 ^{3/}	63	78
Urinary	5.3 ^{4/}	1.2	0.6

¹Significantly (P<0.05) greater than least value.

²P<0.01.

³Significantly (P<0.05) greater than least two values.

⁴P<0.01.

1966; Miller, 1973). Absolute Mn retention did not differ significantly between treatments. There was a negative absolute Mn retention on the low dietary Mn level (-0.11 mg/day). This finding agrees with the results of Zajcev (1959) and Starodubova (1968), who showed an obligatory loss of Mn when animals were placed on very low Mn diets. Mn retention, as a percent of intake, was significantly ($P < 0.05$) lower on the basal diet than on diets 2 and 3. Mn retention as a percent of intake on diets 2 and 3 did not differ significantly. Hill and Holtkamp (1954) reported a greater net absorption at a low Mn concentration in the diet than at high concentrations, and Gutowska *et al.* (1941) showed that net Mn absorption was proportional to dietary Mn concentration. Fecal and urinary Mn excretion, as percent of intake, were significantly ($P < 0.01$ or $P < 0.05$) higher on the basal diet than on diets 2 and 3. The fecal and urinary Mn excretion, as percent of intake, on diets 2 and 3 were not significantly different.

Dietary Mn was significantly ($P < 0.01$) related to levels of serum Mn on the 9th day, SAP on the 28th day and serum Mg on the 28th day of the experiment (Table 31). Mn intake was significantly ($P < 0.01$) correlated to serum Mn on the 9th day, serum Mn on the 28th day, and SAP on the 28th day of the experiment. Krieg (1966) found no parallel between Mn in the diet and that in blood. There was a poor positive correlation between Mn intake and average daily gain, and feed efficiency. Average daily gain and feed efficiency were poorly correlated to all the serum parameters except SAP on the 9th day. Serum Mg on the 28th day was related to feed, urine and tissue Mn concentration. Absolute fecal Mn excretion had a significant ($P < 0.01$) positive correlation with serum Mn on the 9th day and SAP on the 28th day of the experiment. Mn retention as percent of intake was significantly ($P < 0.01$ or $P < 0.05$)

TABLE 31. CORRELATIONS BETWEEN MN BALANCE MEASURES, SERUM PARAMETERS AND GROWTH⁴

Item	Day ³	Manganese level				RET	Growth data		% of intake		
		Feed	Intake	FE	UE		ADG	F/G	RET	FE	UE
SMn	9	.76 ^{1/}	.78 ^{1/}	.70 ^{1/}	.48	.55	.39	-.38	.71 ^{1/}	-.72 ^{1/}	-.88 ^{1/}
SMn	21	.49	.63 ^{2/}	.48	.56	.61 ^{2/}	.61 ^{2/}	-.39	.84 ^{1/}	-.84 ^{1/}	-.85 ^{1/}
SMn	28	.48	.81 ^{1/}	.61 ^{2/}	.41	.59 ^{2/}	.22	.09	.59 ^{2/}	-.59 ^{2/}	-.74 ^{1/}
SAP	9	.46	.39	.51	.15	.11	-.47	.76 ^{1/}	-.14	.15	-.08
SAP	21	.47	.36	.31	.28	.20	-.29	.58 ^{2/}	.05	-.05	-.09
SAP	28	.84 ^{1/}	.80 ^{1/}	.72 ^{1/}	.58 ^{2/}	.37	-.03	.20	.32	-.31	-.54
SMg	9	-.18	-.09	.00	.17	-.19	.01	.04	-.19	.19	.14
SMg	21	.59 ^{2/}	.11	.15	.62 ^{2/}	.50	.52	-.48	.40	-.40	-.22
SMg	28	.87 ^{1/}	.47	.12	.63 ^{2/}	.79 ^{1/}	.57	.30	.85 ^{1/}	-.85 ^{1/}	-.71 ^{1/}
ADG		.24	.31	.05	.57	.56	---	---	.65 ^{2/}	-.65 ^{2/}	-.61 ^{2/}
F/G		-.06	.15	.14	-.16	-.57	-.70 ^{1/}	---	-.61 ^{2/}	.61 ^{2/}	.39

¹Significant (P 0.01).

²Significant (P 0.05).

³Days after start of the experiment.

⁴Abbreviations are as follows: SMn = serum manganese; SAP = serum alkaline phosphatase; SMg = serum magnesium; ADG = average daily gain; F/G = feed/gain; FE = absolute fecal Mn excretion; UE = absolute urinary Mn excretion; RET = absolute Mn retention.

positively related to serum Mn throughout the experiment, serum Mg on the 28th day and average daily gain. This may indicate that a high retention of Mn enhances growth rate. Fecal and urinary Mn excretion, both as a percent of intake, were significantly ($P < 0.01$ or $P < 0.05$) negatively correlated to serum Mn throughout the experiment, and serum Mg on the 28th day. Feed efficiency was significantly ($P < 0.05$) negatively related to Mn retention as percent of intake. Mn retention and fecal Mn excretion, both as percent of intake, had high correlations with average daily gain and were better indicators of growth rate than absolute Mn intake, excretion or retention, or any of the serum parameters studied. Up to the 21st day of the experiment, none of the serum parameters were significantly correlated with each other. On the 28th day, serum Mn had significant ($P < 0.05$) positive correlations with serum Mg and SAP (Table 32).

Correlations within the balance measures are summarized in Table 33. Dietary Mn level was positively correlated to Mn intake, excretion and Mn retention as percent of intake. However, these correlations were significant ($P < 0.01$) only with Mn intake and absolute fecal Mn excretion. Urinary Mn excretion as a percent of intake was significantly ($P < 0.01$) negatively correlated to dietary Mn levels. Woerpel and Balloun (1964) and Hill and Holtkamp (1954) reported that retention was highly related to dietary Mn levels. Mn intake and absolute fecal and urinary Mn excretion were all positively correlated to Mn retention as percent of intake and absolute Mn retention. Mn intake was significantly ($P < 0.01$) positively correlated to urinary Mn excretion as percent of intake. Absolute Mn retention had high correlations with Mn retention as percent of intake ($r = + 0.85$) and fecal Mn excretion as percent of intake ($r = - .85$). There was a poor correlation between

TABLE 32. CORRELATIONS WITHIN THE SERUM PARAMETERS

Period ¹	Comparison ²	r ³
9th day	SMn vs SAP	0.31
	SMn vs SMg	-.13
	SMg vs SAP	0.09
21st day	SMn vs SAP	0.31
	SMn vs SMg	0.08
	SMg vs SAP	0.03
28th day	SMn vs SAP	0.57 ^{4/}
	SMn vs SMg	0.60 ^{4/}
	SMg vs SAP	0.39

¹Days after the start of the experiment.

²SMn, SMg, SAP (Serum Mn, Mg and alkaline phosphatase, respectively).

³Correlation coefficient.

⁴Significant (P<0.05).

TABLE 33. CORRELATIONS WITHIN THE MN BALANCE MEASURES¹

	Feed Mn	Mn Intake	FE	UE	RET	REPI	FEPI	UEPI
Feed Mn	1.00	.99 ^{2/}	.87 ^{2/}	.52	.50	.54	-.52	-.84 ^{2/}
Mn intake		1.00	.88 ^{2/}	.54	.50	.56	-.54	-.86 ^{2/}
FE			1.00	.45	.03	.19	-.17	-.67 ^{3/}
UE				1.00	.31	.51	-.50	-.63 ^{3/}
RE					1.00	+.85 ^{2/}	-.85 ^{2/}	-.60 ^{3/}
REPI						1.00	-1.00 ^{2/}	-.80 ^{2/}
FEPI							1.00	.79 ^{2/}
UEPI								1.00

¹Abbreviations are as follows: FE = fecal Mn excretion; UE = absolute urinary Mn excretion; RE = absolute Mn retention; REPI = Mn retention as percent of intake; FEPI = fecal Mn excretion as percent of intake; UEPI = urinary Mn excretion as percent of intake.

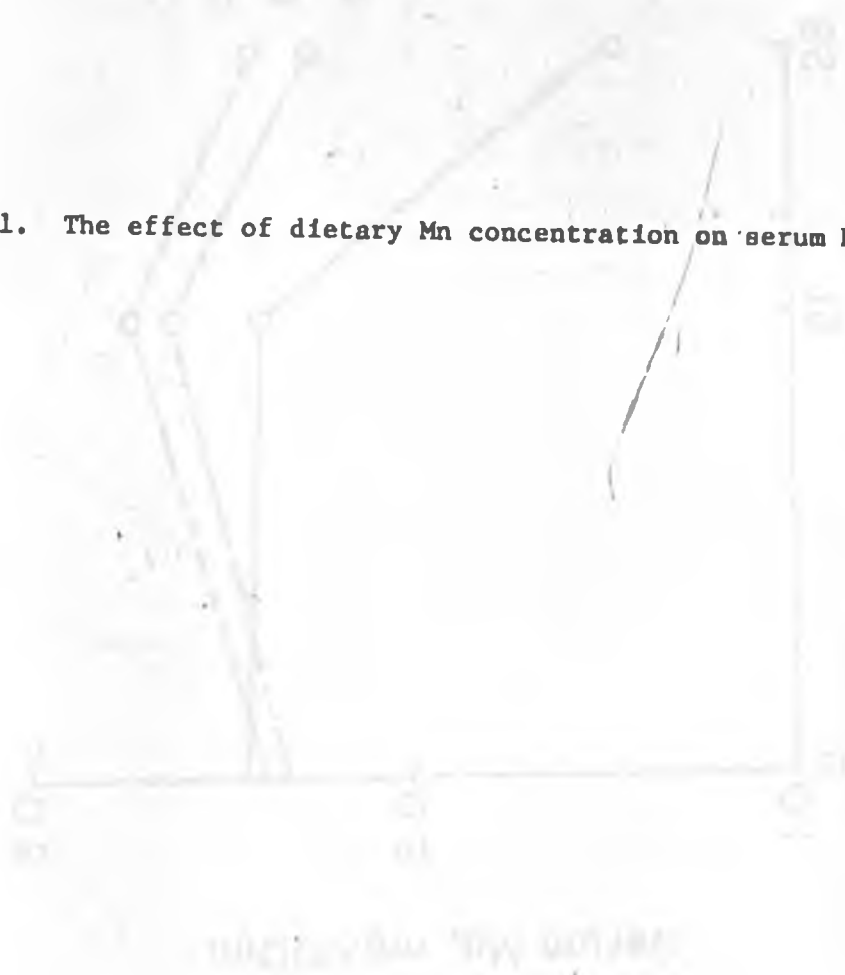
²Significant at $P < 0.01$.

³Significant at $P < 0.05$.

absolute fecal Mn excretion and absolute Mn retention. Mahoney and Small (1968) reported a good relationship between fecal Mn excretion and body retention. Mn retention as percent of intake had a perfect ($r = -1.0$) negative correlation with fecal Mn excretion as percent of intake and a highly significant ($P < 0.01$) correlation with urinary Mn excretion as percent of intake. Urinary and fecal Mn excretion, both as percent of intake, were significantly ($P < 0.01$) positively related. In this study, there was no significant positive correlation between Mn intake and absolute Mn retention or Mn retention as percent of intake, which is contrary to Murty's (1957) findings in sheep and to Mathers' and Hill's (1967) findings in chickens.

DIET 1 10.46
DIET 2 12.67
DIET 3 15.88

Figure 4.1. The effect of dietary Mn concentration on serum Mg concentration.



Boys on Filtered Diet

Figure 4.1

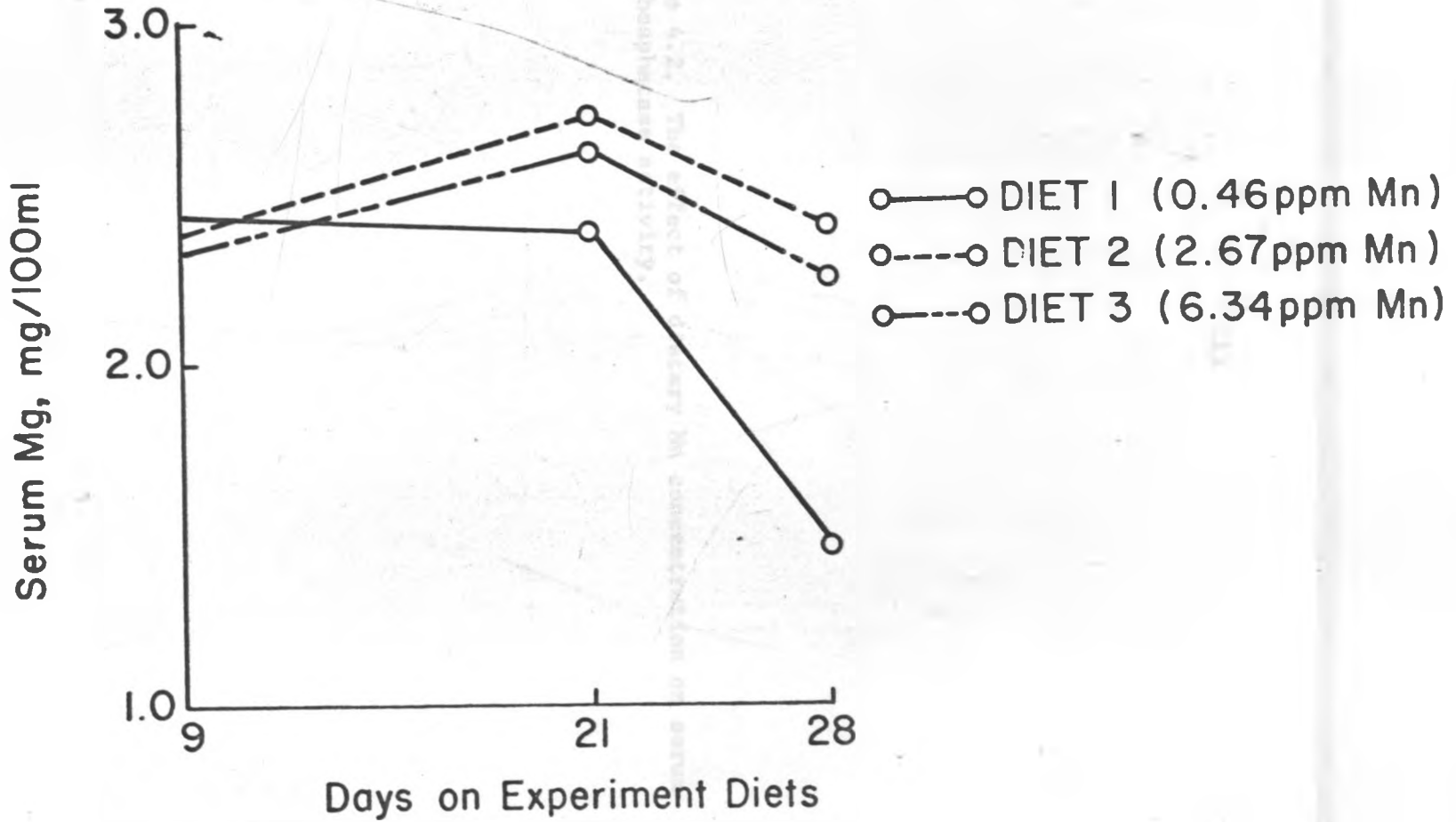


Figure 4.1.

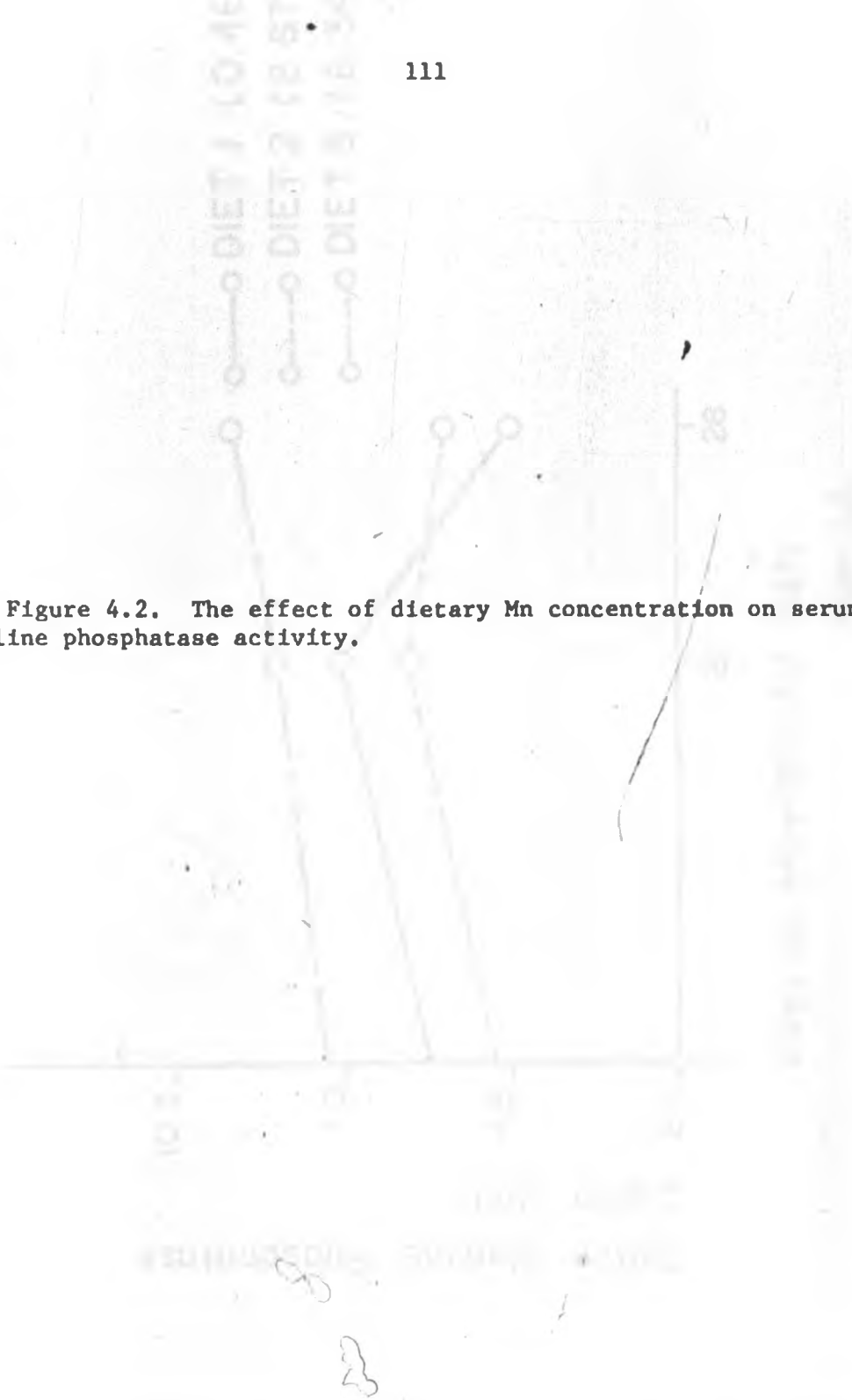


Figure 4.2. The effect of dietary Mn concentration on serum alkaline phosphatase activity.

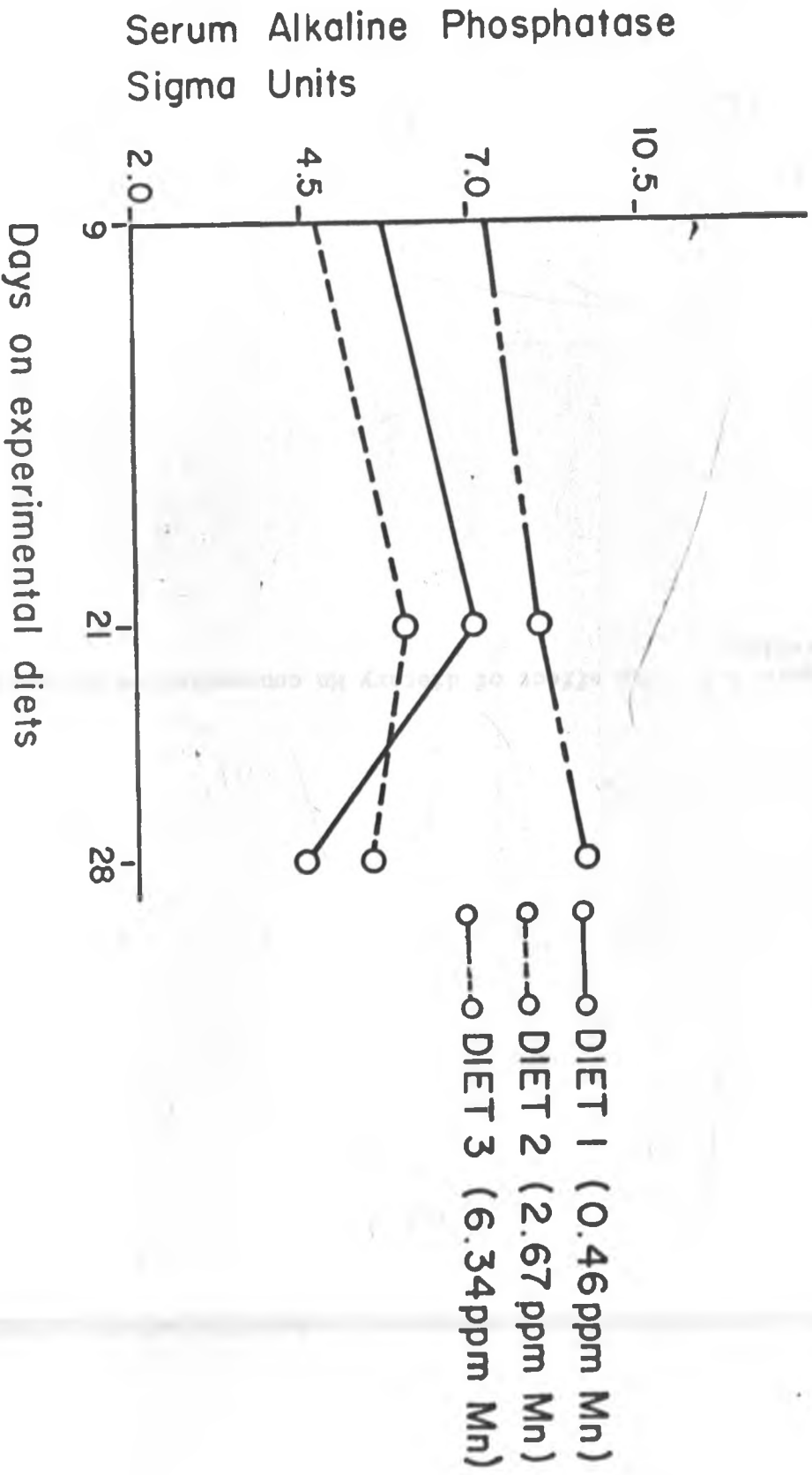


Figure 4.2.

○—○ DIET 1 (0.48)
 ○—○ DIET 2 (2.67)
 ○—○ DIET 3 (6.33)

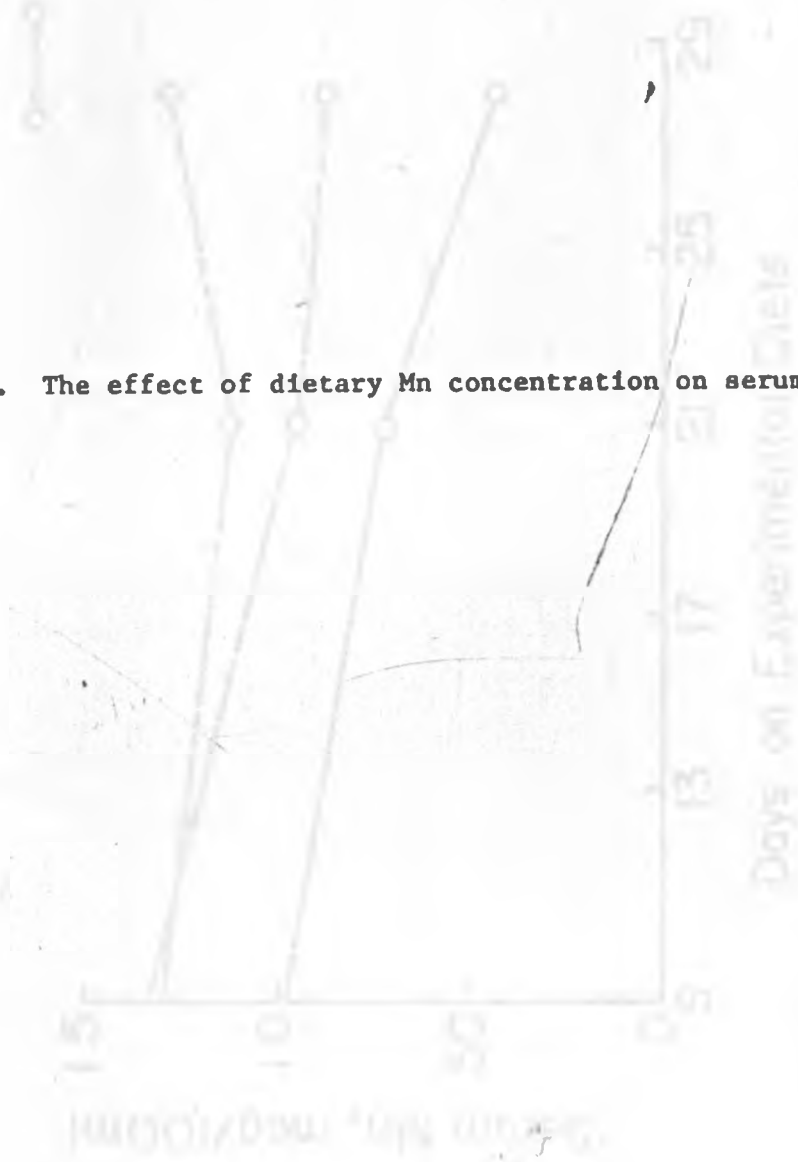


Figure 4.3. The effect of dietary Mn concentration on serum Mn concentration.

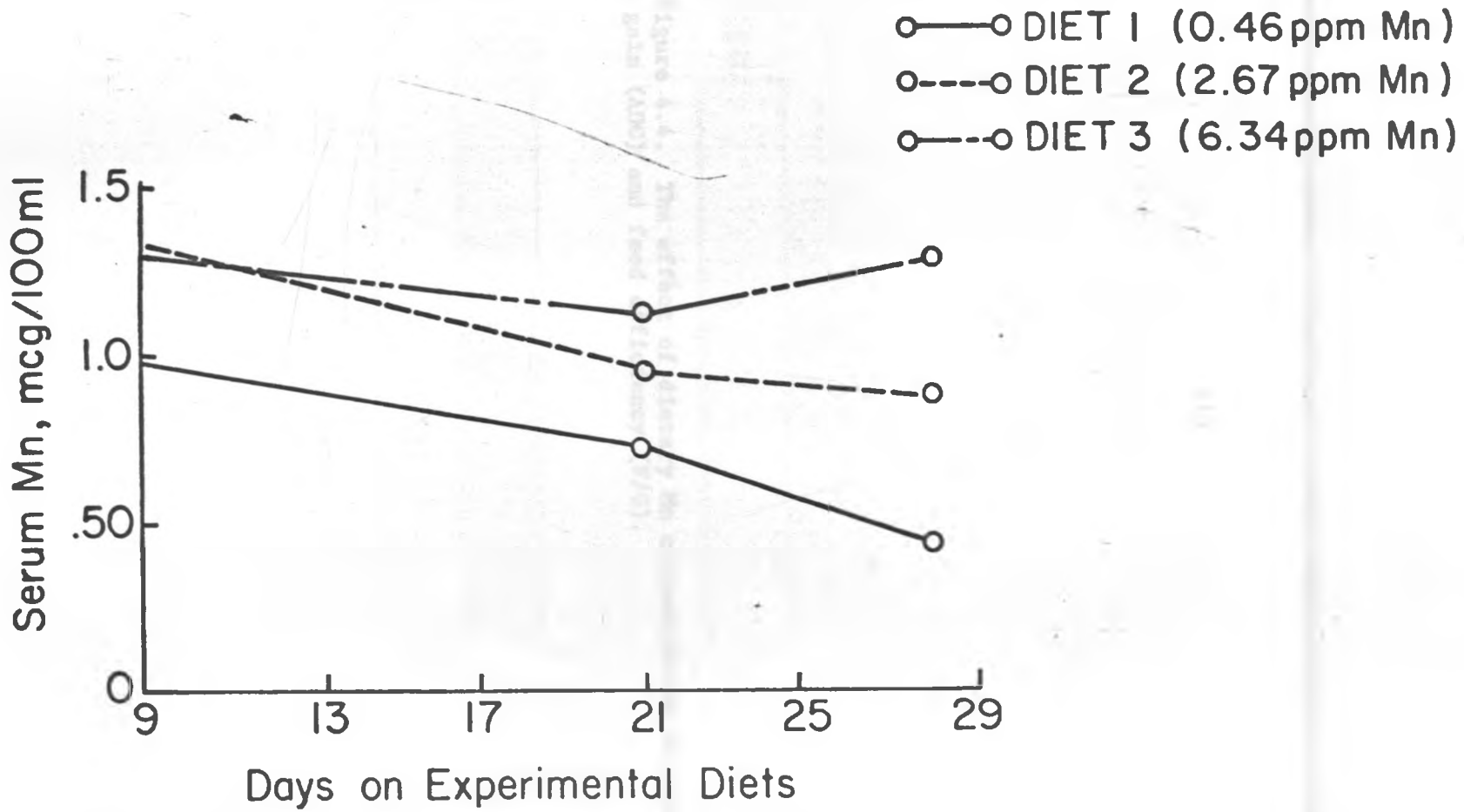


Figure 4.4.

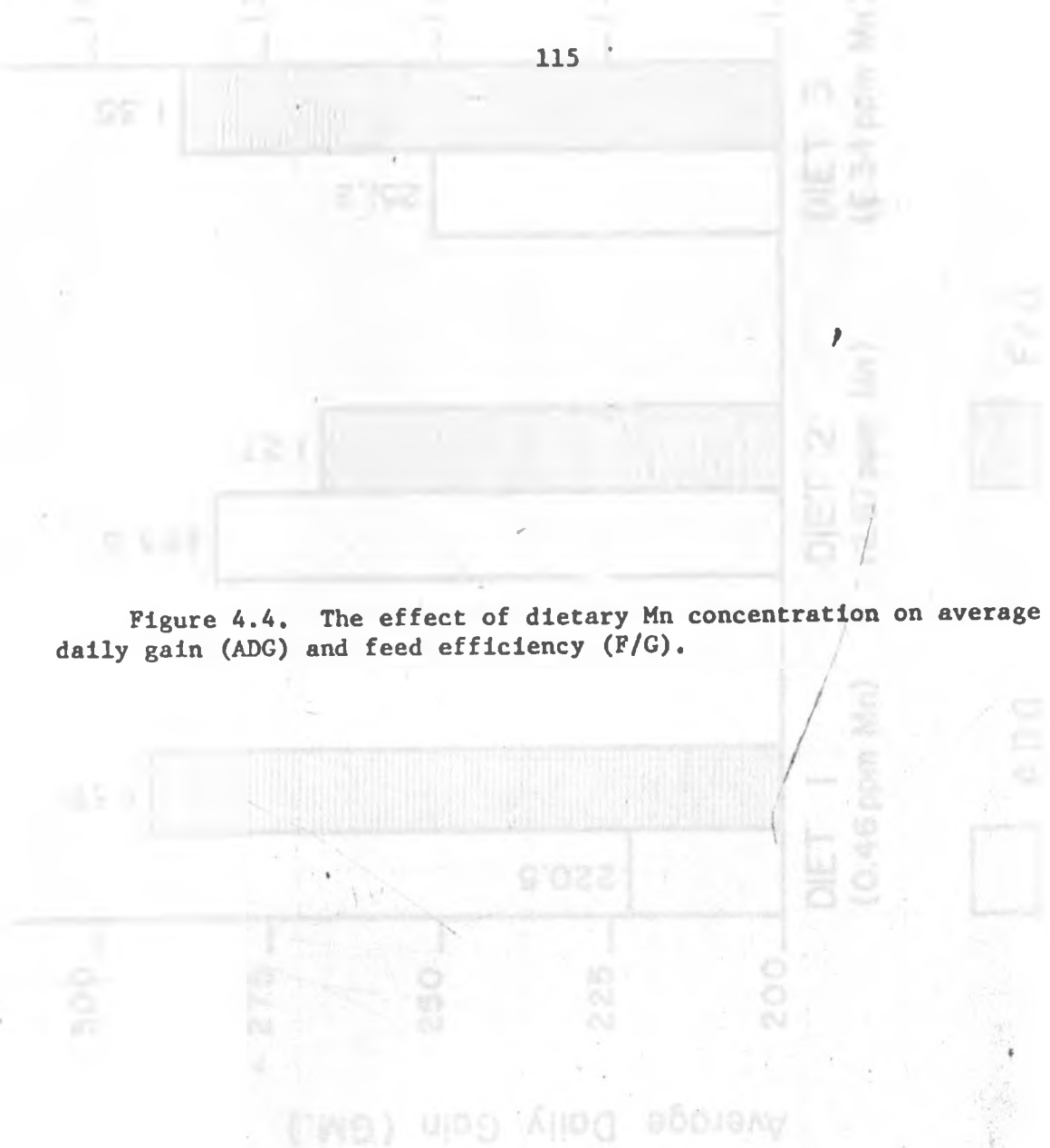
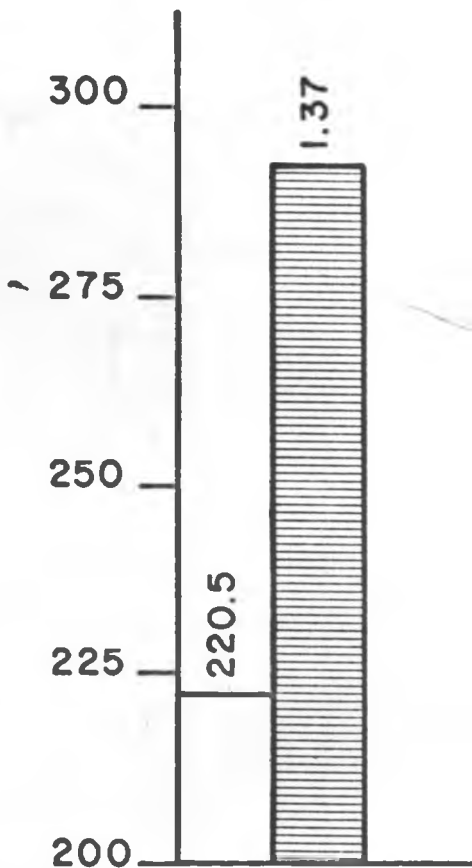


Figure 4.4. The effect of dietary Mn concentration on average daily gain (ADM) and feed efficiency (F/G).

Average Daily Gain (GM.)



DIET I
(0.46 ppm Mn)



ADG

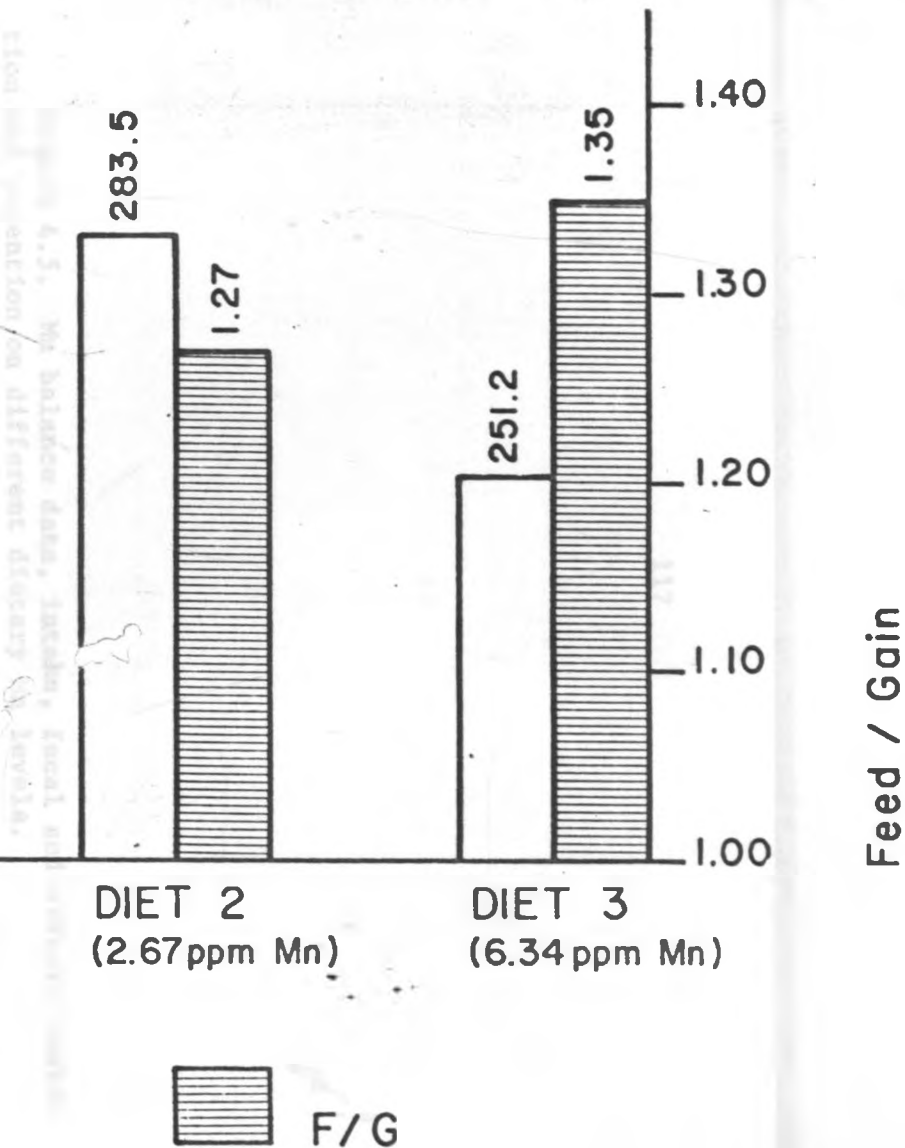


Figure 4.4.

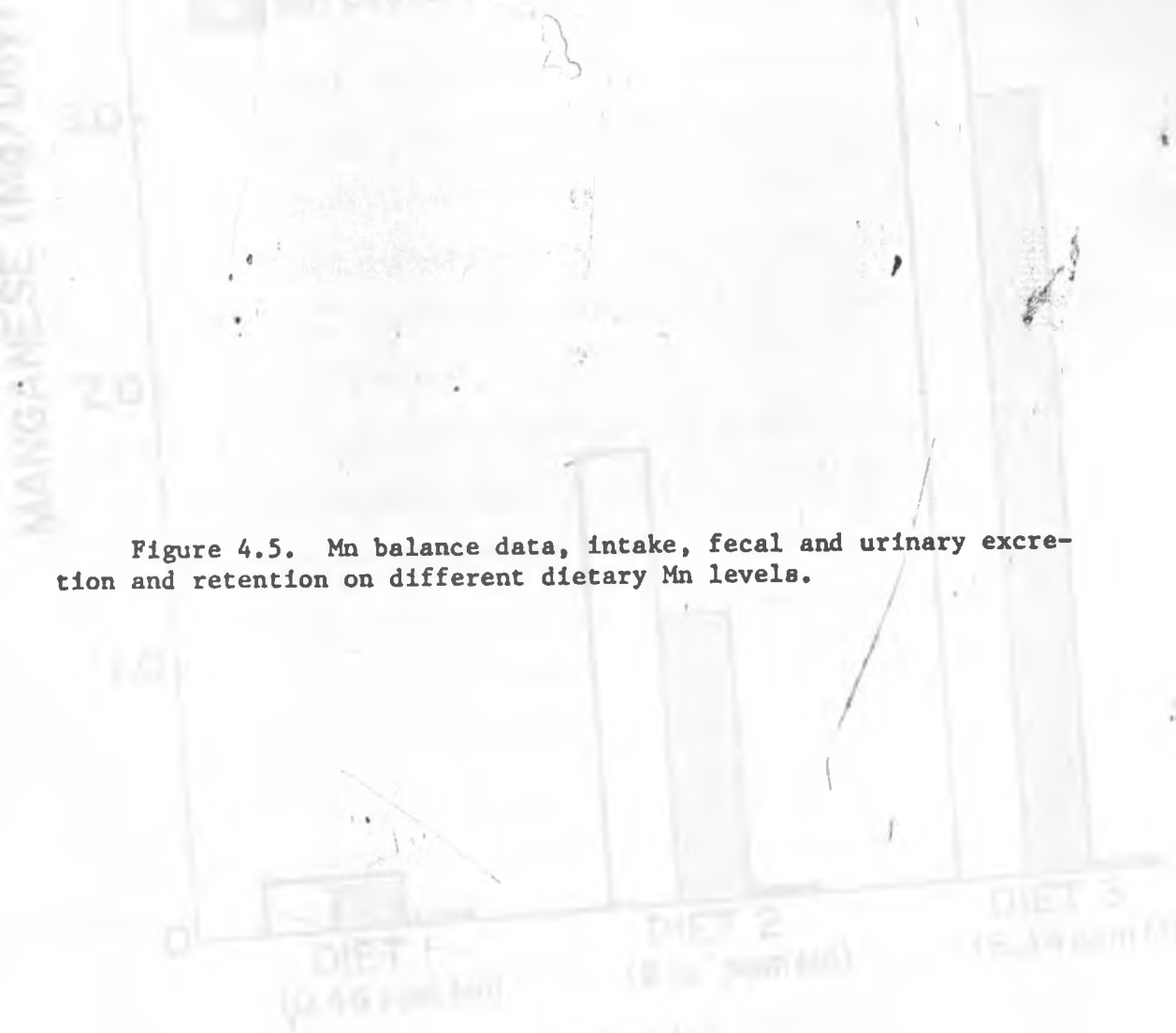


Figure 4.5. Mn balance data, intake, fecal and urinary excretion and retention on different dietary Mn levels.

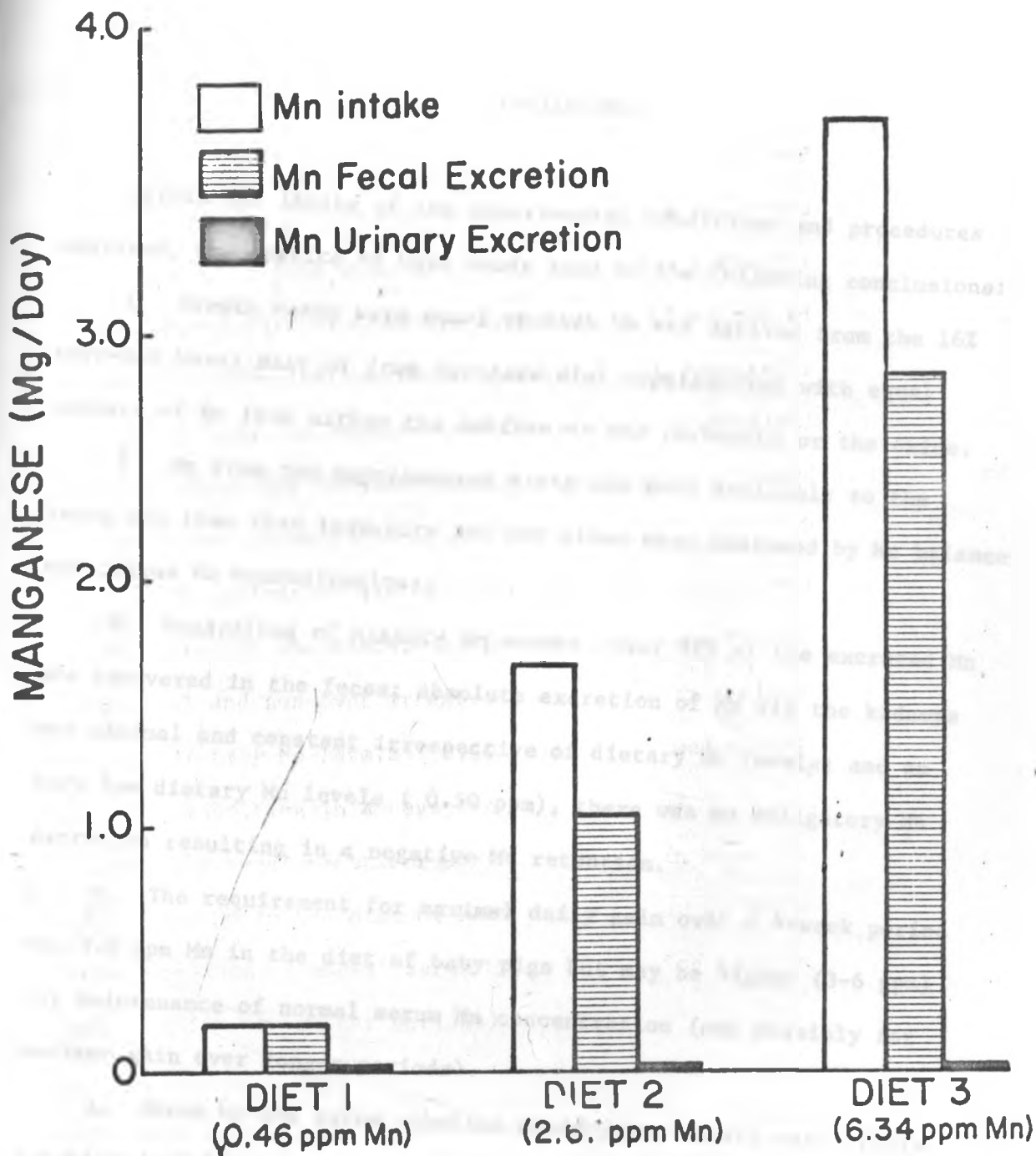


Figure 4.5.

CONCLUSIONS

Within the limits of the experimental conditions and procedures employed, the results of this study lead to the following conclusions:

1. Growth rates were equal whether Mn was derived from the 16% corn-soy basal diet or from the same diet supplemented with equal amounts of Mn from either the sulfate or the carbonate or the oxide.

2. Mn from the supplemented diets was more available to the young pig than that from corn and soy alone when assessed by Mn balance and tissue Mn concentrations.

3. Regardless of dietary Mn source, over 90% of the excreted Mn was recovered in the feces; absolute excretion of Mn via the kidneys was minimal and constant irrespective of dietary Mn levels; and on very low dietary Mn levels (0.50 ppm), there was an obligatory Mn excretion resulting in a negative Mn retention.

4. The requirement for maximal daily gain over a 4-week period was 3.0 ppm Mn in the diet of baby pigs but may be higher (3-6 ppm) for maintenance of normal serum Mn concentration (and possibly for maximum gain over longer periods).

5. Serum Mn and serum alkaline phosphatase levels were highly correlated with Mn intake, but average daily gain and feed efficiency were not. Mn retention and fecal Mn excretion, as percent of intake, had high correlations with average daily gain and were better indicators of growth rate than absolute Mn intake, excretion and retention, and the serum parameters.

6. Mn from the supplemented diets was apparently absorbed in the stomach, cranial small intestine and cecum, whereas Mn from corn and soy was apparently absorbed in the stomach and cecum only. Net cecal absorption of Mn from the basal diet was higher than that of Mn from supplemented diets, and net secretion of Mn into the caudal small intestine on the supplemented diets was higher than on the basal diet.

7. Mn supplementation of a 16 ppm Mn diet did not significantly affect hemoglobin and hematocrit values, serum Ca, Mg, and inorganic P levels, bone physical measurements, breaking strength or related parameters.

8. Although Mn supplementation produced a more normal histologic structure of the epiphysis, it reduced the thickness of compact bone in the region of the diaphysis.

9. There was a significant metabolic interaction between Ca, P and Mn levels in the diets of pigs, which produced changes in bone structure that were distinctly different from those seen in rickets.

It should be emphasized that the deleterious effects on weight gain, feed efficiency and the histology of bone of a low dietary P level (0.35%) from soybean meal were much more pronounced than the effects of excessive dietary levels of Ca and P or an inverse Ca to P ratio, regardless of dietary Mn supplementation.

Popov, 1968. Effect of
the diet on immunological

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APPENDIX A

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TABLE A-1. MICHIGAN STATE UNIVERSITY VITAMIN-TRACE MINERAL PREMIX

Nutrient	Amount in 10 lbs of premix ¹
Vitamin A, million	3.0 I.U.
Vitamin D ₂ , million	0.6 I.U.
Vitamin E, thousand	10.0 I.U.
Riboflavin	3.0 gm
Nicotinic acid	16.0 gm
Choline chloride	100.0 gm
D-pantothenic acid	12.0 gm
Vitamin B ₁₂	18.0 gm
Zinc	68.0 gm
Manganese	34.0 gm
Iodine	2.5 gm
Copper	9.0 gm
Iron	54.0 gm
Antioxidant ²	45.0 gm

Carrier (ground yellow corn) to bring total to 10 lbs

¹Ten pounds of premix mixed in 1 ton of feed.

²Butylated hydroxyanisole (BHA) and/or butylated hydroxytoluene (BHT).

TABLE A-2. MINERAL MIXTURE USED IN SEMI-PURIFIED EXPERIMENTAL DIETS

Nutrient	Percentage in diet
KCl	10.0
KI	0.0002
FeSO ₄ ·2H ₂ O	0.7
CuSO ₄	0.1
COCO ₃	0.1
MnSO ₄ ·H ₂ O	0.0, .009 ^{1/} , .027 ^{2/}
ZnSO ₄	0.4
MgHCO ₃	2.0
CaHPO ₄ ·2H ₂ O	36.0
CaCO ₃	12.5
Cerelose	13.098, 13.089 ^{1/} 13.071 ^{2/}
NaHCO ₃	25.0

^{1,2}For 3 and 9 ppm Mn supplementation of the diets 2 and 3.

TABLE A-3. VITAMIN MIXTURE USED IN SEMI-PURIFIED EXPERIMENTAL DIETS

Nutrient	
	<u>ppm in diet</u>
Thiamine mononitrate	3
Riboflavin	6
Nicotinamide	40
Calcium pantothenate	30
Pyridoxine HCl	2
Para-amino benzoic acid	13
Ascorbic acid	80
D- α -tocopheryl acetate	10
Inositol	130
Choline chloride	1300
	<u>ppb in diet</u>
Pteroyl glutamic acid	260
Biotin	50
Cyanocobalamin	100
2-methyl-1, 4-naphthoquinone	40
Vitamin A palmitate	1500
Vitamin D ₂	12.5

TABLE A-4. CLEANING POLYETHYLENE VIALS

Polyethylene vials were cleaned successively with the following solutions:

1. Xylene
2. 95% ethyl alcohol
3. Distilled deionized water
4. (0.1N) dilute nitric acid
5. Distilled deionized water

The vials were:

1. Dried in a draft oven at 100°C immediately prior to loading
2. Numbered by engraving letters using a stainless metal rod

TABLE A-5. IL MODEL 335 FLAMELESS SAMPLER

1. Instrumental parameters		
Mode of operation	A-B	Channel B
Hollow cathode	45472	H ₂ continuum
Lamp current	5mA	#IL 63490
Photomultiplier	R456	20mA
P.M. voltage	620V	
Slit width	320 μ m	
Wavelength	279.5	
2. Recorder (full scale)		
Response time	0.5 sec	
Range	10. mV	
3. Parameters for 355		
Mode	Manual	
Purge gas	Argon	
Gas flow rate	10 SCFH	
Dry setting	4 turns	
Pyrolyze setting	5 turns on	
	Analyse Scale	
Height of measurement	2 mm	
Analyse setting	10 turns	
