Metabolic bone disease in lion cubs at the London Zoo in 1889: the original animal model of rickets

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Abstract
In 1889 Dr. John Bland-Sutton, a prominent London surgeon, was consulted about fatal rickets in over 20 successive litters of lion cubs born at the London Zoo. He evaluated the diet and found the cause of rickets to be nutritional in origin. He recommended that goat meat with crushed bones and cod-liver oil be added to the lean horsemeat diet of the cubs and their mothers. Rickets were reversed, the cubs survived, and subsequent litters thrived. Thirty years later, in classic controlled studies conducted in puppies and young rats, the definitive role of calcium, phosphate and vitamin D in prevention and therapy of rickets was elucidated. Further studies led to identifying the structural features of vitamin D.

Although the Bland-Sutton diet provided calcium and phosphate from bones and vitamins A and D from cod-liver oil, some other benefits of this diet were not recognized. Taurine-conjugated bile salts, necessary for intestinal absorption of fat-soluble vitamins, were provided in the oil cold-pressed from cod liver. Unlike canine and rodent species, felines are unable to synthesize taurine, yet conjugate bile acids exclusively with taurine; hence, it must be provided in the diet. The now famous Bland-Sutton “experiment of nature,” fatal rickets in lion cubs, was cured by addition of minerals and vitamin D. Taurine-conjugated bile salts undoubtedly permitted absorption of vitamins A and D, thus preventing the occurrence of metabolic bone disease and rickets.

Introduction
The story of the discovery of the cause and prevention of rickets is exciting and represents one of the most remarkable medical accomplishments of the 20th century [1]. The prevalence of rickets in children living at northern latitudes was staggering, as high as 60% to 80% in some areas [1-4]. A body of work over a 15-year period beginning in 1917 defined the biologic properties of a “substance” in cod liver oil that was antirachitic. This substance also appeared to be produced by ultraviolet irradiation of the skin [1,4]. The structure of this substance, vitamin D, was determined and resulted in a Nobel Prize for Dr. Adolph Windaus in 1928 [5]. Historically, the addition of irradiated ergosterol to milk at 400 IU per liter, done independently by Steenbock and Hess [6-8] led to the virtual elimination of rickets in humans. The recognition that vitamin D is a group of prohormones that are converted to the active form (1,25 (OH)2 D, or calcitriol) by a tightly regulated synthetic pathway is now recognized and accepted [9].

Relevance of animal models
Studying proper animal models provides an understanding of vitamin D action. The utility of animal models was obvious to basic and clinical science scholars of rickets because of the overwhelming prevalence of the disease [1,2,4,10,11]. Although overt rickets was more frequently seen in lower socio-economic groups and in crowded tenements, it was common at all levels of society. Surveys showed that fully 60% to 80% of children who lived in urban areas in northern Europe during the period of urban migration and industrialization...
had clinically obvious rickets. Skies were blackened by the smog from burning coal, and diets were often inadequate in calcium, phosphate and vitamin D content. By the mid-1920’s, there were many posited theories as to the cause of rickets, well summarized by Edwards Park in his 1923 review [4]. However, the need for carefully designed animal studies was obvious.

A somewhat obscure early model arose from the attempt to reverse rickets in lion cubs at the London Zoo in Regent’s Park. While this open label, non-controlled interventional trial reversed fatal rickets in cubs, its essential value was encouraging other investigators to design studies that explored the role of nutrients in the pathogenesis and treatment of rickets and osteomalacia [1,4].

It was from the studies of two groups (first, Edward Mellanby and co-workers in England, who evaluated the role of diet on the etiology of rickets in puppies, and second, the collaboration of Elmer V. McCollum, John Howland, Edwards A. Park and their team in Baltimore, MD, who studied rats fed restricted diets) that it was discerned that certain fats (cod-liver oil, butter and milk) could reverse the bone histologic and clinical features of rickets [2,4,9]. A substance in fat, far more concentrated in cod-liver oil than in butter, was the “antirachitic” factor. McCollum, who ultimately “discovered” vitamin D, was able to show that oxidized cod-liver oil (oxidation destroys vitamin A) still retained this antirachitic factor.

Impact of the Bland-Sutton study
To fully appreciate why the studies of Mellanby and McCollum led to the discovery of vitamin D, a more probing analysis of Bland-Sutton’s work with lion cubs is appropriate.

Three decades prior to Mellanby’s studies, in 1889, an “experiment of nature” occurred in the confines of the Zoological Garden in London. With the exception of a single litter, more than 20 consecutive litters of lion cubs perished. The surviving cubs were suckled by their dam for only two weeks, as she did not produce enough milk. The cubs were weaned very early onto lean horse-meat. They “invariably” developed extreme rickets and died [4]. Dr. John Bland-Sutton, a major figure in British surgery, was consulted and recommended a twice-weekly diet of goat flesh, goat bones and milk for both dams and cubs. He also suggested that the cubs lick cod-liver oil. He chose goat bones because they are softer than those of horses. Oddly, although Bland-Sutton wrote extensively, including his autobiography, he never recounted his experience in the zoo in a full publication. The curative diet was never formally described. There exist some conflicts in accounts of the exact diets fed the dams and cubs [4]. The message that Mellanby took away from Bland-Sutton’s work was that the curing of these lion cubs was related to the addition of fat to the diet. Mellanby used these ideas to design his famous dog studies in 1919 [12,13] and demonstrated the value of dietary fat, and in particular of cod-liver oil, in the prevention and treatment of rickets. Hence, the earlier studies of Bland-Sutton gained special importance. In contrast, Park, of the Baltimore team, felt that the true worth of Bland-Sutton’s work was somewhat overblown. He stated, “The so-called experiments of Bland-Sutton have had an influence which apparently they have not merited” [4]. In addition, Park pointed out that sometimes lions upon maturation “presented remarkable rickety changes in their skulls.” Despite Park’s reservations, numerous investigators of vitamin D deficiency in addition to Mellanby [2,4,9] recognized that the lion cubs in the London Zoo represented the initial animal experimental model of rickets.

Although cod-liver oil was curative of rickets in puppies and rats, the story in lion cubs is more complex and one in which numerous other factors are only now being recognized. An obvious interpretation of Bland-Sutton’s dietary manipulations is that the cubs were being provided with minerals (calcium and phosphorus) from the goat bones and vitamin D from the cod-liver oil. The initial diet was lean horse-meat, which has an inverse ratio of calcium and phosphorus compared to bone and is devoid of fat. The limited milk intake, which was the main source of calcium and phosphorus in proper ratios, no doubt contributed to the under-mineralization of bones in the cubs [2]. Thus, lactation failure on the part of the dam was also a huge component of the development of rickets [4].

Several other factors, however, are critical to the diet of big cats such as lions, tigers and leopards [14]. Cats do not synthesize vitamin D₃ adequately in the skin, and require a dietary source [15-18]. Fat-soluble vitamins work synergistically as well as antagonistically, particularly if imbalanced [19]. Felines also lack the ability to convert provitamin carotenoids, including β-carotene, into active vitamin A [20]. Vitamin A is important to the integrity of the epithelium of the respiratory and digestive tracts. Large cats that lack vitamin A can develop sinusitis, diarrhea, blindness, conjunctivitis and neurologic signs [14]. Clinical signs from vitamin A-deficient immature lions include incoordination, “star gazing”, blindness and intermittent convulsions [21]. Neurologic dysfunction accompanied by malformation of both the skull and the cervical vertebrae has been described in lions kept in captivity worldwide, and this dysfunction and malformation were most often related to vitamin A deficiency [22].

When big cats eat whole animals or are given bones they ingest calcium and phosphorus in a ratio of two
parts calcium to one part phosphorous. Chunk lean meat has a calcium:phosphorus ratio of 1:15 to 1:30. Great cats deficient in calcium intake can develop osteomalacia; thus, their diets should be fortified with calcium [14]. It is easier for cubs to chew softer bones, such as those of goats, rather than the hard bones of horses [4].

Cod-liver oil is replete with vitamin A and D, with a conventional value of 4000 to 5000 IU of A and 400 to 500 IU of D per teaspoonful (5 ml) [23]. However, in order to be utilized these fat-soluble vitamins must be absorbed from the intestine. Bile salts are essential in this biologic phenomenon. Cod-liver oil contains bile acids and, indeed, some bile salts. The conjugation of bile acids, especially those in herring [24] and cod [25], and in great cats, is dependent on the availability of taurine [23,24]. Because of the manner in which cod-liver oil is processed (extracted from fish liver at 82 °F under steam and then cold pressed to prevent emulsification), both taurine and bile acids are preserved [26]. This supplement (or medication) contains not only the vitamins but also the compounds that break down lipid micelles and permit intestinal fat absorption [26,27]. Moreover, cod, as with most marine fish or elasmobranchs, use taurine as an osmolyte, which is essential for cell volume regulation [27,28]. Typical supermarket cod contains 90 mg taurine per 100 g of raw fish [29]. The primary bile acids of cod, as with most fish, are cholic acid and to a lesser degree chenodeoxycholic acid [30].

The great cats, similar to their domestic cousins, are obligate carnivores. Felines have limited amounts of cysteinesulfonic acid decarboxylase, a rate-limiting enzyme in the synthesis of taurine from methionine and cysteine, hence taurine is an essential amino acid in cats. Cats fed a vegetable protein diet require taurine to prevent retinal degeneration and blindness [31,32]. A recent study has shown that the chunk horsemeat diet of exotic felines contains low taurine concentrations [33].

Cholic acid is also by far the dominant bile acid of lions [34]. Taurine conjugation of bile acids is mandatory, and glycine conjugation does not occur in either cod or lions [27,30,32,34]. Therefore, by providing cod-liver oil to the lion cubs, Bland-Sutton also increased their dietary intake of bile acids and taurine. This promoted absorption of fat-soluble vitamins [2,9].

Of interest, the goat can synthesize taurine, has a brisk enterohepatic circulation of bile acids, and its muscles contain ample taurine [35]. Thus, goat flesh was also a good source of taurine. Also, taurine is the most abundant free amino acid in goat milk [36].

The story of the elucidation of the pathogenesis of rickets obviously is also about the photocutaneous synthesis of vitamin D [1-4,9]. The remarkable observations of Palm [37] concerning the increased prevalence
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