

# The Western-Style Diet, Calcium Deficiency and Chronic Disease

#### Muhammad Nadeem Aslam and James Varani<sup>\*</sup>

The Department of Pathology, University of Michigan, Ann Arbor, MI 48109, USA

\*Corresponding author: James Varani, Department of Pathology, University of Michigan, 1301 Catherine Road/Box 5602 Ann Arbor, MI 48109, USA, Tel: 7346150298; E-mail: varani@umich.edu

#### Received date: March 21, 2016; Accepted date: April 14, 2016; Published date: April 21, 2016

**Copyright:** © 2016 Muhammad Nadeem Aslam, et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

## Abstract

The term "Western-style diet" refers to an eating pattern that includes a high content of saturated fat, a large amount of processed carbohydrate and excess total calories. The Western-style diet contributes to the growing epidemic of obesity and several age-related, chronic illnesses seen in the United States and throughout the world. In addition to its high content of fat and sugar, the Western diet is also characterized by a deficiency in calcium (and, undoubtedly) other trace minerals that are nutritionally associated with calcium. While epidemiological evidence suggests that the lack of adequate dietary calcium contributes to several chronic ailments associated with the Western-style diet, studies in experimental animals provides direct evidence. Rodents on a high-fat, low-calcium diet suffer many of the same chronic illnesses that are seen in humans. When the calcium concentration is increased to the level found in rodent chow diets, the ill-effects are mitigated. While calcium alone is protective, a combination of calcium and additional trace elements has been shown, in some studies, to induce even better protection. The implication is that providing an adequate supply of essential minerals (including calcium, of course, but also other trace elements that support calcium's beneficial activities), either through dietary modification or as a supplement if dietary modification fails should be considered as part of an overall strategy for counteracting the negative effects of the Western-style diet.

**Keywords:** Calcium; Cancer; Cationic trace elements; Chronic disease; Western-style diet

## Introduction

The term "Western-style diet" refers to an eating pattern that includes a high content of saturated fat, a large amount of processed carbohydrate and too many calories. Red meat, processed meat products, refined grains and starch (potatoes) are mainstays of the Western-style diet. High-fat dairy products are another component of the typical diet consumed in many Western countries. Fruits, vegetables, legumes, fish, other seafood and whole grains are, generally, under-consumed. While the Western-style diet is commonly assumed to be "unhealthful," there are numerous variations in what is actually consumed by any given individual (as is true of any diet). Not all eating patterns are equally bad. In, perhaps, its worst form, added fat from seed oil extractions and added highly-refined carbohydrate, i.e., sugar-(empty calories) make up a significant percentage of the overall calorie intake. At one time largely associated with individuals in North America, Europe and countries of European descent, the Western-style diet is now a world-wide phenomenon. The Western-style diet underlies the growing epidemic of obesity in the United States and throughout the world [1,2]. It is thought to contribute to the increasing incidence of several chronic illnesses including cancer (especially colon and liver), cardiovascular disease and metabolic disease (e.g. type II diabetes) [1,3,4]. The Western-style diet has been linked to chronic kidney disease [5], non-alcoholic fatty liver disease (NAFLD)/nonalcoholic steatohepatitis (NASH) [6,7] and osteoporosis [8]. A recent study has shown an association between the Western diet and earlyonset dementia [9]. A role in inflammatory skin diseases (acne, psoriasis, atopic dermatitis) has also been suggested [10].

There is little doubt of the relationship between the Western-style diet and the chronic diseases indicated here. What is not clear, however, is how the food components that make up the Western-style diet (either individually or together) bring about their detrimental effect on health. Is it entirely related to the high content of saturated fat and processed carbohydrates? The hypothesis we put forward here is that the Western-style diet, with its high content of processed fat and carbohydrate and, most concomitantly, its relative lack of unprocessed fruits, vegetables and whole grains, leads to a deficiency in essential minerals along with the vitamin co-factors necessary for proper mineral metabolism. The Western-style diet, we hypothesize, is detrimental to health as much by its lack of essential minerals as by what it contains.

#### The western-style diet, calcium-deficiency and chronic illness

While the Western-style diet includes a large amount of saturated fat and processed carbohydrates, there are additional nutritional features associated with this diet. Fiber, folic acid and choline are under-represented. With a deficiency in methyl group donors, the diet has characteristics of the "choline-deficient, amino acid-defined (CDAA) diet [11]. Additionally, and perhaps most importantly, the Western-style diet is deficient in calcium and vitamin D. With regard to calcium, per se, the recommended intake for adolescents and adults is in the range of 1000-1300 mg per day [12-14]. However, the average intact for many individuals in Western society is lower. Kudlacek et al. [15] reported in 2003 that the average calcium intake among a sample of over one thousand individuals in Austria was 560 mg per day. Many individuals, of course, had much lower intake. A similar pattern in the United States was reported a few years later [16]. The authors of both studies concluded that low calcium intake was more wide-spread than previously thought. In the ensuing decade, the situation has not changed for the better. In its most recent summary of critical nutrients,

the USDA concluded that a high percentage of individuals in multiple age groups failed to reach minimal recommended calcium intake [17].

A calcium deficiency is independently associated with numerous chronic diseases-i.e., with many of the same diseases noted here [18-22]. While bone loss and osteoporosis is the obvious example, the relationship between calcium intake and colon cancer may be particularly instructive. Interventional trials have demonstrated that calcium supplementation can lower the incidence of recurrent colon polyp formation, although the reduction is modest [23-25], and not all studies have confirmed reduced incidence [26,27]. Numerous epidemiological studies have shown a correlation between higher calcium intake and reduction in colon cancer risk [28-32]. While not every study has established a statistically-significant protective relationship, meta-analysis of past findings supports a positive correlation (i.e., reduced polyp incidence with increased calcium intake) [33] and a recent analysis suggest that protection extends to colon cancer itself [34].

Work with epithelial cells in culture provides mechanistic insight [35-38]. Epithelial cells from various sources (including the colon) proliferate optimally over a broad range of low-calcium concentrations (0.05-0.5 mM). Under these conditions, cells do not express features of the differentiated state. As the calcium concentration is increased above 0.5 mM, differentiation is induced. Key features include induction of E-cadherin synthesis; its translocation from the cytoplasm to the cell surface; and formation of the cell surface adhesion complex. This process is readily reversible. When calcium is removed, cells revert to an undifferentiated state. This is depicted in Figure 1.

Two consequences of calcium-induced differentiation include: i) reduced proliferation and ii) formation of the epithelial barrier. In regard to proliferation,  $\beta$ -catenin is sequestered in the adhesion complex along with E-cadherin. This leads to decreased  $\beta$ -catenin movement into the nucleus where it otherwise functions as a Wntpathway (growth-promoting) enhancer [36-38]. The end-result is decreased proliferation. Equally important, E-cadherin - mediated cellcell cohesion allows the differentiated epithelial cells to form a cohesive cell sheet (Figure 1). This is essential for barrier protein synthesis and formation of barrier structures (tight junctures and desmosomes) [39]. Defective barrier function in the gastrointestinal tract and chronic inflammation go "hand in hand". Commonly, it is thought that chronic inflammation is responsible for barrier breakdown, but it is more likely that poor barrier function contributes to the tendency toward inflammation [40]. In the absence of an effective barrier, bacteria, bacterial products, toxins and food allergens can all gain access to the interstitium. Inflammation in the gastro-intestinal tract and carcinogenesis in the colon are linked [41] and decreased inflammation can contribute to reduced tumor incidence with calcium. While calcium-induced tumor suppression could reflect a direct action on intracellular (growth-regulating) signaling pathways or result from inhibition of chronic inflammation in the gastrointestinal tract, these are not the only ways in which calcium might act. Calcium may be anti-carcinogenic by altering luminal pH with an effect on the microbial community [42] or by precipitating carcinogenic bile acids in the gastrointestinal tract [43]. These mechanisms are not mutually exclusive.



**Figure 1:** The cartoon depicts a cohesive sheet of epithelial cells in the presence of 1.5 mM calcium. When the extracellular calcium level is reduced to 0.15 mM, there is a rapid change in cell shape and a loss of cell-cell cohesion. This change can be seen within minutes. Subsequently, some of the cells undergo proliferation. Other cells undergo injury. In addition, there is a loss of barrier function. Microbes, toxins, allergens etc. penetrate the interstitium. The lower panels of the figure demonstrate changes in E- cadherin production/expression by epithelial cells in response to alterations in extracellular calcium. A rapid loss of total E-cadherin from the cell surface (right panel) occurs, resulting in dis-cohesion.

There is little doubt of the importance of calcium to effective growth regulation (by whatever mechanism) in the colon. Ultimately, however, the question is not whether calcium has beneficial properties, but whether the level of calcium in the Western-style diet is sufficiently low as to obviate any of these potential mechanisms by which calcium could affect tumor formation in the colon. Studies in experimental animals have begun to address this issue. Newmark et al. [44-46] maintained C57BL/6 mice on a rodent version of the Western-style diet for 18-24 months. A higher incidence of precancerous colon polyps was observed in these animals than in littermates maintained on a rodent chow diet for the same period (29% incidence versus 12%). When calcium (reduced from 5.25 mg/kg in the rodent chow diet to 0.41 mg/kg in the Western diet) was replenished to the level of the rodent chow diet, the adenoma incidence was reduced to nearbackground levels. When other modifications of the Western diet (low fiber, folate and choline; and replacement of methionine with cysteine) were adjusted to conditions in the control diet, there was no mitigation of polyp formation. In a subsequent study using mice containing a mutated adenomatous polyposis coli (APC) gene, a similar trend was observed; that is, there was a higher incidence of colon polyps (as well as tumors in other sites) in Western diet-fed mice than in controls. As seen with C57BL/6 mice, the combination of calcium and vitamin D provided significant protection [47]. The same combination has also

been shown to suppress formation of pre-neoplastic colon lesions induced by the strong carcinogen, azoxymethane [48].

Using a similar approach to Newmark's, our own studies examined the effects of a calcium-rich, multi-mineral natural product (Aquamin<sup>°</sup>) derived from the skeletal remains of red marine algae on colon tumor formation in C57BL/6 mice. Consistent with the findings of Newmark et al. [44,46] we also demonstrated a significant reduction in tumor incidence with mineral supplementation [49,50]. In our studies, animals maintained on a rodent chow diet, had a colon polyp incidence of 18% (16 of 90 mice).

In animals fed a Western-style diet without the mineral supplement, the incidence of polyp formation was 29% (26 of 90) while in littermates fed the Western-style diet with supplementation, the incidence was 2% (2 of 90). Of note, when the tumors were examined histologically, several of the lesions in the Western diet-fed mice proved to be invasive carcinomas. No invasive tumors were seen in mice fed either the rodent chow diet or the calcium-supplemented Western-style diet. These data, thus, suggest that supplementation may affect tumor progression as well as tumor formation. Also of note, all of the diets in our studies contained vitamin D (120 IU/kg), suggesting that in the absence of an adequate supply of calcium, this amount of vitamin D, by itself, was not effective.

Although suppression of growth-regulating signaling pathways or effects on carcinogenic bile acids might explain anti-carcinogenicity in the colon, a reduction in chronic inflammation could have broader effects. Our own studies not only demonstrated reduced colon polyp formation but also showed that mineral supplementation protected mice against bone loss [51,52] and reduced the incidence and severity of ulcerative dermatitis [53]. Perhaps most interesting, during the course of our studies, we observed a high incidence of liver tumor formation in mice on the Western-style diet [54]. Unlike what was observed with colon polyps (where both males and females were susceptible), virtually all of the liver tumors were in males. When these lesions were examined histologically, they encompassed a wide range of presentations - from large non- regenerative and regenerative hyperplastic nodules to premalignant hepatic adenomas and fullymalignant hepatocellular carcinomas. Other manifestations of liver injury, i.e., inflammation, and ballooning degeneration of hepatocytes, along with areas of necrosis and fibrosis - were also observed. In male mice on the mineral-supplemented Western diet, tumor formation was substantially reduced (48% incidence without supplement versus 12% incidence in supplement-fed mice, against a background incidence of 16% for male mice on the rodent chow diet). Inflammation and hepatocyte necrosis were also reduced.

As part of the study, serum calcium levels were assessed in mice from each diet group. In male mice on the supplemented Western-style diet, the average calcium level was  $10.1 \pm 0.7$  mg/dl, while in mice on the un-supplemented diet; the average was  $9.5 \pm 1.3$  mg/dl. Since serum calcium levels are tightly controlled between approximately 9-10 mg/dl, these values put the calcium-supplemented animals at the high end of the normal range while the un-supplemented mice were at a level midway between the upper and lower normal range values. In the same study, male mice maintained on rodent chow diet (containing a comparable amount of calcium to that in the supplemented Western diet [5.25 mg/kg of diet]) also had serum calcium levels at the upper end of the normal range ( $9.9 \pm 1.3$  mg/dl). Of interest, female mice had higher levels of serum calcium than males under all conditions. In females, serum calcium levels were  $10.9 \pm 1.3$  mg/dl,  $10.9 \pm 1.0$  mg/dl and  $11.4 \pm 0.8$  mg/dl on the un-supplemented and supplemented Western-style diets and rodent chow diet, respectively). Thus, liver disease and low serum calcium values appear to be correlated.

Of interest, while liver injury was confined almost entirely to males, both male and female mice on the Western-style diet gained excess weight, and demonstrated serum chemistry abnormalities. These were not affected by calcium-supplementation.

Thus, we hypothesize that it is not up-stream consequences of the Western diet that are mitigated by calcium, but the later events that produce overt injury. While our studies may have been the first to document the beneficial effects of calcium in the liver, previous studies have shown a reduction in liver fibrosis by vitamin D [55]. The beneficial effects of vitamin D were presumed to reflect interference with transforming growth factor- $\beta$  signaling, with little regard to its role in calcium uptake and utilization.

If these findings can be extrapolated to humans, they open up a new avenue for prevention of liver injury occurring as a consequence of poor nutrition. A public health strategy that focuses on preventing the consequences of fatty liver disease rather than targeting the formation of fatty changes per se may prove to be more effective.

Certainly, targeting down-stream consequences of steatosis along with steatosis, itself, should be considered. Finally, while these findings are in the context of the Western- style diet, liver injury due to viral infection as well as alcoholic liver disease may also be amenable to a similar interventional approach. The up-stream initiators of tissue damage are different, but all share common, down-stream pathophysiological mechanisms [56].

How circulating calcium protects the liver is not fully understood. We postulate a similar mechanism to what has been suggested in the colon - i.e., that in the presence of calcium, hepatocyte differentiation occurs, limiting excessive proliferation (and injury), while promoting barrier formation (Figure 1). This, of course, will need to be established experimentally.

## Calcium supplementation to mitigate health consequences of the western-style diet: Possibilities and limitations

To the extent that the Western-style diet is a problem of calciumdeficiency, the solution would seem obvious - provide a sufficient amount of calcium, preferably as part of a healthful diet, but as a supplement where dietary improvement fails. The use of calcium supplements (alone and in conjunction with other nutrients) is already widespread. Their primary use is in prevention of bone loss and osteoporosis, but people utilize calcium supplements to reduce risk of colon polyp formation or to mitigate other health concerns. Without minimizing the value of calcium supplementation, there are a number of issues that should be considered. Beyond the usual - bioavailability and tolerability - is the potential for adverse consequences at high doses. For example, a meta-analysis of calcium supplement use data concluded that a risk of cardiovascular events did exist for the highest doses of supplement use [57]. An association between calcium supplement intake (self-reported) and macular degeneration in the elderly has also been reported [58]. Perhaps more troubling is the positive correlation in some studies between calcium intake and prostate cancer [59-61]. Whether the benefits of calcium supplement use outweigh potential risks has to be determined; sometimes on a case by case basis. Equally important is the reality that no critical nutrient, including calcium, functions in a vacuum. How well calcium from any source performs depends on the presence (at appropriate levels) or absence of other nutrients. Importance of vitamin D to calcium uptake from the gastrointestinal tract and at the cellular level is well-known [18-21].

interest of evidence-based medicine, additional studies will be needed to address this issue.

Less well-known but, perhaps, equally important is the level of other important minerals. Magnesium, for example, has little chemopreventive activity by itself, but the ratio of magnesium to calcium has been shown to be important for calcium chemoprevention in the colon [62]. Magnesium is probably not unique. However, since magnesium is present in substantial amounts, it is possible to establish this interaction by creating an experimental deficiency and measuring the consequences. This is not the case with other potentially important divalent or trivalent cationic trace elements; some of which are present in truly "trace" amounts. The lanthanide elements constitute one such group. The lanthanides, because of their similarity to calcium in terms of orbital size and electronic configuration [63,64], interact with calcium-binding sites on proteins, often with higher affinity than calcium itself. Calcium channel proteins [65-67] and proteins that are part of calcium-exchangers [68] have been shown to bind lanthanide elements - leading to either enhanced or inhibited function (altered calcium influx-efflux). The extracellular calcium-sensing receptor (CaSR) is another calcium-binding protein capable of high-affinity lanthanide binding [69-71]. This protein, which is sensitive to tiny changes in the extracellular calcium concentration, plays a critical role in colon epithelial cell growth control [72-76]. Our own past studies have shown that in the presence of gadolinium (lanthanide family member), there is a "left-shift" in the response to calcium. That is, CaSR is up-regulated [73,74] and growth-suppression occurs at lower calcium concentrations than would otherwise occur [77].

While many experimental approaches have utilized gadolinium as a representative of the lanthanide family, we conducted a survey study in which all 14 naturally-occurring lanthanide elements were compared for ability to suppress epithelial cell proliferation [78]. Only three members in the entire family (terbium, dysprosium and ytterbium) failed to have significant activity at a concentration of 100 µM. At the other extreme, the most potent lanthanides (thulium, gadolinium and samarium) had activity at 5-10 µM. The capacity to modify response to calcium was not seen with several other divalent or trivalent cationic trace elements including aluminum, iron (ferrous and ferric), cobalt, copper, nickel, magnesium, manganese and zinc. Thus, the lanthanides appear to function through a mechanism that is not shared by many other cationic trace elements. This is not to suggest, however, that the lanthanides are unique. Two relatively abundant cationic elements (barium and strontium) are CaSR activators [70,71,79]. Of interest, it appears that strontium activation of CaSR and activation by calcium do not lead to identical signaling events-providing a rationale for potential co-operativity [80].

The question is not whether certain minor trace elements can modulate responses to calcium, but whether they are present (as a group) at circulating levels or tissue levels sufficient to accomplish this task *in vivo*. With the lanthanides, at least, this question will be difficult to address since the *in vivo* levels of individual lanthanide elements are low and not routinely measured. One can assume, perhaps, that since many of these trace elements are nutritionally associated with calcium, a diet that is deficient in calcium might also be deficient in these other trace elements, as well. The implication is that the mineral composition of a healthy diet cannot easily be duplicated in a supplement; no matter how well-thought-out it is. Alternatively, there are multi-mineralcontaining natural products available, and it would be premature to suggest that these cannot provide benefit if used appropriately. In the

## Conclusion

The Western-style diet has a number of features that make it unhealthy. While the focus of this work is on minerals (in particular, calcium), there is no doubt that the high content of saturated fat and processed carbohydrate underlies much of what is wrong with the diet. Our intent is not to minimize this. Rather, the intent is only to point out that in addition to saturated fat and processed carbohydrates; the Western-style diet is also lacking an adequate amount of calcium (and, presumably, other trace elements that are found in the same foods as calcium). A calcium-deficiency is an independent risk factor for many of the same chronic illness associated with the Western-style diet, and it is not unreasonable to hypothesize that the lack of dietary calcium (and, perhaps, other essential trace elements) may contribute to several of the chronic diseases associated with the Western-style diet. That having been said, what is the potential for mitigating these health issues by providing a supply of essential minerals (calcium, of course, but also other trace elements that support calcium's beneficial activities), either through dietary modification or as a supplement if dietary improvement fails? That, in our opinion, is still an open question, and one worth addressing.

## References

- 1. Ogden CL, Yanovski SZ, Carroll MD, Flegal KM (2007) The epidemiology of obesity. Gastroenterology 132: 2087-2012.
- NCD Risk factor Collaboration (2016) Trends in adult body-mass index in 200 countries from 1975 to 2014: a pooled analysis of 1698 populationbased measurement studies with 19.2 million participants. Lancet 387: 1377-1396.
- Heidemann C, Schulze MB, Frando OH, van Dam RM, Mantzoros CS, et al. (2008) Dietary patterns and risk of mortality from cardiovascular disease, cancer and all- causes in a prospective cohort of women. Circulation 118: 230-223.
- 4. Steyn NP, Mann J, Bennett PH, Temple N, Zimmet P, et al. (2004) Diet, nutrition and the prevention of type 2 diabetes. Public Health Nutr 7: 147-165.
- Odermatt A (2011) The Western-style diet: a major risk factor for impaired kidney function and chronic kidney disease. Am J Physiol Renal Physiol 301: F919-931.
- 6. Cohen JC, Horton JD, Hobbs HH (2011) Human fatty liver disease: old questions and new insights. Science 332: 1519-1523.
- Vernon G, Baranova A, Younossi Z (2011) Systematic review: the epidemiology and natural history of non-alcoholic fatty liver disease and non-alcoholic steatohepatitis in adults. Alimentary Pharmacol Thera 34: 274-285.
- 8. Prentice A (2004) Diet, nutrition and the prevention of osteoporosis. Public Health Nutr 7: 227-243.
- Shakersain B, Santoni G, Larsson SC, Faxén-Irving G, Fastbom J, et al. (2016) Prudent diet may attenuate the adverse effects of Western diet on cognitive decline. Alzheimers Dement 12: 100-109.
- 10. Melnik BC (2013) Western-diet mediated mTORC1-signaling in acne, psoriasis, atopic dermatitis, and related diseases of civilization; Therapeutic role of plant- derived natural mTORC1 inhibitors. Chapter 37 In: RR Watson and S Zibadi (eds.) bioactive Dietary factors and Plant Extracts in Dermatology, Nutrition and Health; Springer Science +Business Media, New York.
- Denda A, Kitayama W, Kishida H, Murata N, Tsutsumi M, et al. (2002) Development of hepatocellular adenomas and carcinomas associated in C57BL/6 mice given a choline-deficient, L-amino acid-defined diet. Jpn J Cancer Res 93:125-132.

- Food and Nutrition Board (2011). Institute of Medicine (USA), Dietary reference intakes for calcium and vitamin D. In; Ross AC, Taylor CL, Yaktine AL, del Valle HB (eds), Washington DC, National Academy Press.
- 13. Medical Scientific Advisory committee (2015) Osteoporosis Australia Calcium factsheet.
- 14. Scientific opinion on dietary references for calcium (2015) Euro Food Safety Authority 13: 4101.
- 15. Kudlacek S, Schneider B, Peterlik M, Leb G, Klaushofer K, et al. (2003) Assessment of vitamin D and calcium status in healthy adult Austrians. Eur J Clin Invest 33: 323-331.
- 16. Ma J, Johns RA, Stafford RS (2007) Americans are not meeting current calcium recommendations. Am J Clin Nutr 85: 1361-1366.
- 17. US Department of Agriculture (2015) 2015-2020 Dietary Guidelines for Americans. (8th ed) Washington DC.
- Peterlik M, Boonen S, Cross HS, Lamberg-Allardt C (2009) Vitamin D and calcium insufficiency-related chronic diseases: an emerging worldwide public health problem. Int J Environ Res Public Health 6: 2585-2607.
- 19. Heaney RP (2003) Long-latency deficiency disease: insights from calcium and vitamin D. Am J Clin Nutr 78: 912-919.
- 20. Peterlik M, Cross HS (2005) Vitamin D and calcium deficits predispose for multiple chronic diseases. Eur J Clin Invest 35: 290-304.
- Peterlik M, Cross HS (2009) Vitamin D and calcium insufficiency-related chronic diseases: molecular and cellular pathophysiology. Eur J Clin Nutr 63: 1377-1386.
- 22. Beto JA (2015) The role of calcium in human aging. Clin Nutr Res 4: 1-8.
- 23. Baron JA, Beach M, Mandel JS, van Stolk RU, Haile RW, et al. (1999) Calcium supplements for the prevention of colorectal adenomas. Calcium Polyp Prevention Study Group. N Engl J Med 340: 101-107.
- Wallace K, Baron JA, Cole BF, Sandler RS, Karagas MR, et al (2004). Effect of calcium supplementation on the risk of large bowel polyps. J Natl Cancer Inst 96: 921-925.
- 25. Grau MV, Baron JA, Sandler RS, Wallace K, Haile RW, et al. (2007) Prolonged effect of calcium supplementation on risk of colorectal adenomas in a randomized trial. J Natl Cancer Inst 99: 129-136.
- Baron JA, Barry EL, Mott LA (2015) A Trial of Calcium and Vitamin D for the Prevention of Colorectal Adenomas. N Engl J Med 373: 1519-1530.
- 27. Pommergaard HC, Burcharth J, Rosenberg J, Raskov H (2016) Aspirin, Calcitriol, and Calcium Do Not Prevent Adenoma Recurrence in a Randomized Controlled Trial. Gastroenterology 150: 114-122.
- Cho E, Smith-Warner SA, Spiegelman D, Beeson WL, van den Brandt PA, et al. (2004) Dairy foods, calcium, and colorectal cancer: a pooled analysis of 10 cohort studies. J Natl Cancer Inst 96: 1015-1022.
- 29. Flood A, Peters U, Chatterjee N, Lacey JV Jr, Schairer C, et al. (2005) Calcium from diet and supplements is associated with reduced risk of colorectal cancer in a prospective cohort of women. Cancer Epidemiol Biomarkers Prev 14: 126-132.
- 30. Kesse E, Boutron-Ruault MC, Norat T, Riboli E, Clavel-Chapelon F (2005). Dietary calcium, phosphorus, vitamin D, dairy products and the risk of colorectal adenoma and cancer among French women of the E3N-EPIC prospective study. Int J Cancer 117: 137-144.
- Larsson SC, Bergkvist L, Rutegard J, Giovannucci E, Wolk A (2006) Calcium and dairy food intakes are inversely associated with colorectal cancer risk in the Cohort of Swedish Men. Am J Clin Nutr 83: 667-673.
- Park SY, Murphy SP, Wilkens LR, Nomura AM, Henderson BE, et al. (2007) Calcium and vitamin D intake and risk of colorectal cancer: the Multiethnic Cohort Study. Am J Epidemiol 165: 784-793.
- Shaukat A, Scouras N, Schünemann HJ (2005) Role of supplemental calcium in the recurrence of colorectal adenomas: a metaanalysis of randomized controlled trials. Am J Gastroenterol 100: 390-394.
- Keum NN, Aune D, Greenwood DC, Ju W, Giovannucci EL (2014) Calcium intake and cancer risk: Dose-response meta-analysis of prospective observational studies. Int J Cancer 135: 1940-1948.

- 35. Behrens J, Vakaet L, Friis R, Winterchger E, Van roy F, et al. (1993) Loss of epithelial differentiation and gain of invasiveness correlates with tryrosine phosphorylation of the E-cadherin/beta-catenin complex in cells transformed with a temperature-sensitive v-SRC gene. J Cell Biol 120: 757-766.
- 36. Mariadason JM, Bordonaro M, Aslam F, Shi L, Kuraguchi M, et al. (2001) Down-regulation of beta-catenin TCF signaling is linked to colonic epithelial cell differentiation. Cancer Res 61: 3465-3471.
- Brembeck FH, Schwarz-Romond T, Bakkers J, Wilhelm S, Hammerschmidt M, et al. (2004) Essential role of BCL9-2 in the switch between beta-catenin's adhesive and transcriptional functions. Genes Dev 18: 2225-2230.
- Conacci-Sorrell M, Simcha I, Ben-Yedidia T, Blechman J, Savagner P, et al. (2003) Autoregulation of E-cadherin expression by cadherin-cadherin interactions: the roles of beta-catenin signaling, Slug, and MAPK. The J Cell Biol 163: 847-857.
- **39.** Tunggal J, Helfrich I, Schmitz A, Schwarz H, Gunzel D, et al. (2005) Ecadherin is essential for in vivo epidermal barrier function by regulating tight junctions. EMBO J 24: 1146-1156.
- Cario E, Gerken G, Podolsky DK (2007) Toll-like receptor 2 controls mucosal inflammation by regulating epithelial barrier function. Gastroenterology 132: 1359-1374.
- Itzkowitz SH, Yio X (2004) Inflammation and Cancer: IV. Colorectal cancer in inflammatory bowel disease: the role of inflammation. Am J Physiol: Gastrointestinal Liver Physiol 287: G7-G17.
- Newmark HL, Lupton JR (1990) Determinants and consequences of colonic luminal pH: implications for colon cancer. Nutr Cancer 14: 161-173.
- Bernstein H, Bernstein C, Payne CM, Dvorakova K, Garewal H (2005) Bile acids as carcinogens in human gastrointestinal cancers. A Review. Mutation Res. 589: 47-65.
- 44. Newmark HL, Yang K, Lipkin M, Kopelovich L, Liu Y, et al. (2001) A Western-style diet induces benign and malignant neoplasms in the colon of normal C57BI/6 mice. Carcinogenesis 22: 1871-1875.
- 45. Yang K, Kurihara N, Fan K, Newmark H, Rigas B, et al. (2008) Dietary induction of colonic tumors in a mouse model of sporadic colon cancer. Cancer Res 68: 7803-7810.
- 46. Newmark HL, Yang K, Kurihara N, Fan K, Augenlicht LH, et al. (2009) Western- style diet-induced colonic tumors and their modulation by calcium and vitamin D in C57BL/6 mice: a preclinical model for human sporadic colon cancer. Carcinogenesis 30: 88-92.
- 47. Yang K, Lamprecht SA, Shinozaki H, Fan K, Yang W, et al. (2008) Dietary calcium and cholecalciferol modulate cyclin D1 expression, apoptosis, and tumorigenesis in intestine of adenomatous polyposis coli1638N/+ mice. J Nutr 138: 1658-1663.
- 48. Myung YJ, Seol JK, Nam SY, Yun YW, Kim JS, et al. (2015) Effects of calcium on the formation of preneoplastic lesions in a mouse model of colon carcinogenesis. J Prev Vet Med 39: 15-22.
- 49. Aslam MN, Paruchuri T, Bhagavathula N, Varani J (2010) A mineralized extract from the red algae, Lithothamnion calcerum, inhibits polyp formation and inflammation in the gastrointestinal tract of normal mice on a high-fat diet. Integrative Cancer Therapies, 9: 93-99.
- 50. Aslam MN, Bergin IU, Naik M, Paruchuri T, Dame M, et al. (2012) A multi-mineral natural product from the red marine algae, lithothamnion calcareum reduces colon polyp formation in mice: comparison of the multi-mineral supplement with calcium alone. Nutrition & Cancer 64: 1020-1028.
- 51. Aslam MN, Kreider JM, Paruchuri T, Bhagavathula N, DaSilva M, et al. (2010) A mineral-rich extract from the red marine algae Lithothamnion calcareum preserves bone structure and function in female mice on a Western-style diet. Calcif Tissue Int 86: 313-324.
- 52. Aslam MN, Bergin I, Jepsen K, Kreider JM, Graf KH, et al. (2013) Preservation of bone structure and function by Preservation of bone structure and function by Lithothamnion sp. derived minerals. Biol Trace Elem Res 156: 210- 220.

- 53. Hampton AL, Aslam MN, Naik MK, Bergin IL, Allen RM, et al. (2015) Ulcerative dermatitis in C57BL/6NCrl mice on low-fat and high- fat diets with and without a mineralized red algae supplement. JAALAS 54: 1-10.
- Aslam MN, Bergin I, Allen R, Kunkel SL, Rush H, et al. (2012) A multimineral natural product inhibits liver tumor formation in C57BL/6 mice. Biol Trace Elem Res 147: 267-274.
- 55. Kitson MT, Roberts SK (2012) D-livering the message: The importance of vitamin D status in chronic liver disease. J Hepatol 57: 897-909.
- Anstee QM, Goldin RD (2006) Mouse models in non-alcoholic fatty liver disease and steatohepatitis research. Curent Status Review. Int J Exp Pathol 87: 1-16.
- 57. Bolland MJ, Avenell A, Baron JA, Grey A, MacLennan G, et al. (2010) Effects of calcium supplements on risk of myocardial infarction and cardiovascular events. Brit Med J 341: c3691.
- Kakigi CL, Singh K, Wang SY, Enanoria WT, Lin SC (2015) Self-reported Calcium Supplementation and Age-Related Macular Degeneration. JAMA Ophthalmol 133: 746-754.
- Allen NE, Key TJ, Appleby PN, Travis RC, Roddam AW, et al. (2008) Animal foods, protein, calcium and prostate cancer risk: the European Prospective Investigation into Cancer and Nutrition. Br J Cancer 98: 1574-1581.
- 60. Ahn J, Albanes D, Peters U, Schatzkin A, Lim U, et al. (2007) Dairy products, calcium intake, and risk of prostate cancer in the prostate, lung, colorectal, and ovarian cancer screening trial. Cancer Epidemiol Biomarkers Prev 16: 2623-2630.
- 61. Park Y, Mitrou PN, Kipnis V, Hollenbeck A, Schatzkin A, et al. (2007) Calcium, dairy foods, and risk of incident and fatal prostate cancer: the NIH-AARP Diet and Health Study. Am J Epidemiol 166: 1270-1279.
- 62. Dai Q, Shrubsole MJ, Ness RM, Schlundt D, Cai Q, et al. (2007) The relation of magnesium and calcium intakes and a genetic polymorphism in the magnesium transporter to colorectal neoplasia risk. Am J Clin Nutr 86: 743-745.
- 63. Evans CH (1990) Biochemistry of the lanthanides. In: Biochemistry of the Elements, New York: Plenum Press.
- 64. Gschneidner Jr KA, Eyring L (2000) Handbook on the Physics and Chemistry of Rare Earths. Elsevier Science & Technology Books. Amsterdam, The Netherlands.
- 65. Lansman JB (1990) Blockade of current through single calcium channels by trivalent lanthanide cations. Effect of ionic radius on the rates of ion entry and exit. J Gen Physiol 95: 679-696.
- Caldwell RA, Clemo HF, Baumgarten CM (1998) Using gadolinium to identify stretch-activated channels: technical considerations. Am J Physiol 275: C619-621.
- 67. Docherty RJ (1988) Gadolinium selectively blocks a component of calcium current in rodent neuroblastoma X glioma hybrid (NG108-15) cells. J Physiol 398: 33-47.

- Squier TC, Bigelow DJ, Fernandez-Belda FJ, deMeis L, Inesi G (1990) Calcium and lanthanide binding in the sarcoplasmic reticulum ATPase. J Biol Chem 265: 13,730-13,737.
- 69. Pidcock E, Moore GR (2001) Structural characteristics of protein binding sites for calcium and lanthanide ions. J Biol Inorg Chem 6: 479-489.
- 70. Huang Y, Zhou Y, Castiblanco A (2009) Multiple Ca(2+)-binding sites in the extracellular domain of the Ca(2+)-sensing receptor corresponding to cooperative Ca(2+) response. Biochemistry 48: 388-398.
- 71. McLarnon SJ, Riccardi D (2002) Physiological and pharmacological agonists of the extracellular Ca2+-sensing receptor. Eur J Pharmacol 447: 271-278
- 72. Kalley E, Baina E, Wrba F, Kriwanek S, Peterlik M, et al. (2000) Dietary calcium and growth modulation of human colon cancer cells: role of the extracellular calcium-sensing receptor. Cancer Detection Prevent. 24: 127-136.
- 73. Chakrabarty S, Radjendirane V, Appelman H, Varani J (2003) Extracellular calcium and calcium sensing receptor function in human colon carcinomas: Promotion of E- cadherin expression and suppression of ?-catenin/TCF activation. Cancer Res 63: 67-71.
- 74. Chakrabarty S, Wang H, Canaff L, Hendy GN, Appelman H, et al. (2005) Calcium sensing receptor in human colon carcinoma: Interaction with Ca2+ and 1,25- Dihydroxyvitamin D3. Cancer Res. 65: 493-498.
- 75. Bhagavathula N, Hanosh AW, Nerusu KC, Appelman H, Chakrabarty S, et al. (2007) Regulation of E-cadherin and beta-catenin by Ca2+ in colon carcinoma is dependent on calcium-sensing receptor expression and function. Int J Cancer 121: 1455-1462.
- 76. Singh N, Aslam MN, Varani J, Chakrabarty S (2015) Induction of calcium sensing receptor in human colon cancer cells by calcium, vitamin D and Aquamin: Promotion of a more differentiated, less malignant and indolent phenotype. Mol Carcinogenesis. 54: 543-553.
- Attili D, Jenkins B, Aslam MN, Dame MK, Varani, J (2012) Growth Control in Colon Epithelial Cells: Gadolinium Enhances Calcium-Mediated Growth Regulation. Biological Trace Element Research 150: 467-476.
- Jenkins W, Perone P, Walker K, Bhagavathula N, Aslam MN, et al. (2011) Fibroblast Response to Lanthanoid Metal Ion Stimulation: Potential Contribution to Fibrotic Tissue Injury. Biological Trace Element Research 144: 621-635.
- Coulombe J, Faure H, Robin B, Ruat M (2004) In vitro effects of strontium ranelate on the extracellular calcium-sensing receptor. Biochim Biophys Res Commun. 323: 1184-1190.
- 80. Hurtel-Lamaire AS, Mentaverri R, Caudrillier A, Cournarie F, Wattle A, et al. (2009) The calcium-sensing receptor is involved in Strontium Ranelate-induced osteoclast apoptosis: New insights into the associated signaling pathways. J Biol Chem 284: 575-584.