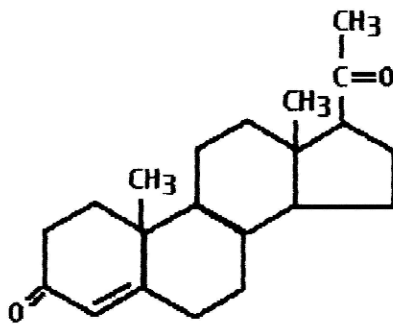


**THE JOURNAL OF  
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**Progesterone**

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Cover Molecule  
PROGESTERONE

Progesterone, the steroid hormone most notably associated with pregnancy, is the only naturally occurring human progestagen (also spelled progestogen). It is synthesized from pregnenolone, a derivative of cholesterol, and consists of four interconnected cyclic hydrocarbons. Progesterone is the precursor of several critical steroid hormones, including cortisol, aldosterone, testosterone, estrone and estradiol. The number and placement of various functional groups (aldehydes, ketones, hydroxyl and methyl groups) on the hydrocarbon rings distinguishes the various steroids from one another. Progesterone should not be confused with progestins, the synthetically produced progestagens.

Progesterone levels vary considerably during the menstrual cycle. More progesterone is secreted in the second half of the menstrual cycle, after ovulation, which serves to thicken the uterine lining and prepare the uterus to support a fertilized egg. If pregnancy does not occur, progesterone levels will drop sharply within a few days, initiating the breakdown of the uterine lining that we know as menstruation. If conception does occur, chorionic gonadotrophin produced by the embryo causes the corpus luteum to maintain progesterone production until the placenta itself can produce enough to maintain the pregnancy.

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## WON'T SOMEBODY THINK OF THE CHILDREN?

The National School Lunch Program was put in place to ensure that all students, even the poorest, would be served lunch while in school. Providing wholesome food to finicky children in an institutional setting is always a challenge, but turning over the job to fast food providers is not the answer. I recall that the school lunches from my childhood in the 1950s were a far cry from the pizza and “happy meals” of today. These plain lunches may have depended heavily on canned vegetables but the mashed potatoes were always made from scratch and a lot of the other food was “home cooking.”

A book such as *Brain-Building Nutrition: The Healing Power of Fats and Oils* by Dr. Michael Schmidt (see a review of this book in this issue) causes one to consider the impact of modern school lunches on the health and behavior of our children. In fact, a report on an “uncontrolled” experiment on diet and school behavior is available at [www.michaelfieldsagainst.org/Food%20Systems/ACACaseStudyFinalVersion.doc](http://www.michaelfieldsagainst.org/Food%20Systems/ACACaseStudyFinalVersion.doc).

This case study describes the experience of students and teachers who participated in the Appleton Central Alternative Charter High School’s Nutrition and Wellness Program in Wisconsin. With money from a local nutrition-oriented baker, the school over a period of a few years simply removed vending machines and

installed water coolers, and switched from providing sack lunches to providing healthy and natural foods.

The results were nearly miraculous. The staff reports that health complaints diminished substantially and students also seemed more able to concentrate. Impulsive behaviors, such as talking, fidgeting and the use of foul language, decreased. Most persuasive is the description of what happens on Junk Food Day, the once-a-year binge that allows sugar sweetened soda, chips, brownies, cookies, and candy bars. Not only are there more complaints of stomachaches, headaches, and feeling tired that day but attendance on the day following Junk Food Day is lower.

The hurdle to overcome before this program can be implemented in every school is money. Not only will federal funds have to be increased in order to provide the more nourishing food for students, school districts will have to get by without the money the vending machines within the schools have provided. Only when enough parents realize that good nutrition is well worth the cost can we expect a change in the school lunch program.

James Heffley, Ph.D., CCN, DANLA  
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## ANTINEOPLASTONS IN DAIRY PRODUCTS

Stanislaw R. Burzynski, M.D., Ph.D.<sup>1</sup>, Elwira Ilkowska-Musial, Ph.D.<sup>2</sup>,  
Maciej W. Klimczak Ph.D.<sup>2</sup>, Lester Musial, M.Sc.<sup>3</sup>

The compounds isolated by our team from plasma and urine, and named antineoplastons (ANP), decrease expression of oncogenes and activate silenced tumor suppressors. Four ANP ingredients, 3-phenylacetyl-amino-2, 6- piperidinedione (A10), phenylacetylglutamine (PG), phenylacetylisoglutamine (isoPG), phenylacetic acid (PN) and one pro drug, 4-phenylbutyric acid (PB), were determined in milk, farmer's and feta cheese and whey by a Shimadzu HPLC system. PG and isoPG occur in the highest concentrations in whey (29.0 and 6.0 mg/100 mL) and farmer's cheese (22.0 and 3.0 mg/100 g). A10 and PN were found in the highest concentration of 7.0 mg/100g and 4.0 mg/100g correspondingly in farmer's cheese. PB occurred in trace amounts in farmer's cheese. In conclusion, A10, PG, isoPG, PN and PB exist in small amounts in dairy products. Based on the previous studies, the supplements containing ingredients of antineoplastons may play an important part in prevention of cancer and anti-aging.

**Keywords:** Antineoplastons in food, chemoprevention, anti-aging, phenylacetate, phenylbutyrate, phenylacetylglutamine.

### INTRODUCTION

The first few years of the new century coincided with the 100<sup>th</sup> anniversary of radiation therapy and the introduction of new, exciting molecular-targeted therapies for cancer treatment (1, 2). Prominent oncologists believe that this is the end of conventional cancer treatment and the beginning of a new era of molecular therapies. The agents of targeted therapies, such as monoclonal antibodies or small molecules, correct the genetic changes which trigger the cancerous process, with only minimal adverse

reactions (3, 4). A similar approach is reflected in cancer prevention, with the introduction of the new generation of food supplements effecting gene expression by "turning off" over-expressed oncogenes and "turning on" silenced tumor suppressor genes (5-7). Study of the human genome revealed that only about 10% of our genes are active at any time in adult life (8). The genes are being turned on and off by the system of biochemical factors named epigenome (9,10). In case of a malfunction of this system, the genes which promote cancer—oncogenes—will be over-expressed and the genes, that protect against cancer—tumor suppressor genes—will be silenced. A disturbed balance in the expression of these genes leads to cancer (8). Silencing of important genes, such as WRN1 helicase, will lead to premature aging as documented in Werner syndrome (11). Cancer

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and aging are integral parts of life and can be traced to very primitive organisms. The most important genes and “molecular switches” involved in these processes can also be traced from humans to rodents, insects, worms, plants, and even yeast (12).

The proper use of these molecules in our diet can provide us with a defense against cancer and slow down genetic aging. These substances, isolated from plants, prevent cancer by turning off the signal in important oncogene pathways (6). The chemicals of human origin, which also were isolated from animal products, play an important part in reducing the activity of over-expressed oncogenes and in activation of silenced tumor suppressor genes (8, 13). Our team has isolated these compounds from human blood and urine since 1968 and named them antineoplastons (ANP) (13, 14). The emphasis was placed on ANP which were reproduced synthetically, including 3-phenylacetyl-amino-2, 6-piperidinedione (A10) (15), phenylacetylglutamine (PG) (16) and phenylacetic acid (PN) (17).

Early research data indicated that ANP are species-specific. It was found, initially, that the only animal species, in addition to humans, which conjugate substantial quantities of L-glutamine with phenylacetic acid (the first step in the synthesis of A10) are old world monkeys (18-23). The other animal species conjugate different amino acids with phenylacetic acid: mice-glycine, rats and dogs-L-glutamic acid, and ferrets-aurine (21-23). Sheep are an exception; they produce 4-phenylbutyric acid (PB), which, in the human liver, converts into PN and PG (24, 25). Studies performed by researchers from the USDA revealed that PG is a normal ingredient of cows' milk, raising the possibility that other antineoplastons and their prodrug (PB) may also exist in small quantities in dairy products (26).

The objective of this study is a quantitative determination of A10, PG, phenylacetylglutamine (isoPG), PN, and PB in dairy products.

## MATERIALS AND METHODS

### Materials

Farmer's cheese, feta cheese, fresh whole milk and condensed milk are commercially available products obtained from the store.

Curd and whey were prepared in our laboratory according to the procedure described below.

### Sample preparation

#### Farmer's cheese and feta cheese

The portion (approx. 40-60g) of raw farmer's or feta cheese was homogenized using Biohomogenizer model M113 (Biospec Products Inc.). The homogenized sample was weighted and then packed in extraction thimbles made of filter paper (Whatman No 4). Extraction was carried out in the Soxhlet apparatus, with methanol as the solvent, for approx. 25 hrs. 100 mL of methanol (HPLC grade, Fisher Scientific) was used for each extraction. When the extraction process was finished, methanol was partially evaporated from the extract on the rotary evaporator RE47 (Yamato Scientific Co. Ltd.). Then, a small amount of water was added to the mixture to enable freezing of the solution. The condensed extract was frozen, then lyophilized to dryness using a VIRTIS freeze dryer (The Virtis Company, Gardiner, N.J.). The process was carried out for approx. 48 hrs. Next, the dry residue was reconstituted with 1 mL of N,N-dimethylformamide (DMF) (HPLC grade, Fisher Scientific). Samples varied from one another, so, for some of them, 1 mL of the solvent was not enough. The dry residue extensively absorbed the solvent. In such cases more DMF had to be used. Due to the higher amount of dry residue, the feta cheese extract was reconstituted with 2 mL of DMF. In some experiments, DMF was replaced by methanol, water: methanol mixture (1:5), 0.01M sodium hydroxide (NaOH) or methanol: 0.01M NaOH mixture (5:1). Generally, changing the solvents did

not improve the solubility of the dry residue. The mixture was centrifuged for 20 min. at 13,000 rpm in order to separate insoluble ingredients from the extract. Some part always remained insoluble. Then the solution was filtered through a 0.45 $\mu$ m nylon syringe filter (Whatman).

#### **Milk and condensed milk**

100 mL of whole fresh or condensed milk was frozen. Then it was lyophilized to dryness using a VIRTIS freeze dryer. The dry sample was treated similarly to farmer's cheese.

#### **Curd and whey.**

A portion of whole fresh milk (1.8 L) was warmed up to 28 $^{\circ}$ C. It was stirred well and 1/2 enzymatic tablet (rennet) dissolved in 100 mL of distilled water was added to the warm milk. The mixture was stirred well for 15 min. Then it was left aside for 2 hrs at room temperature. Next, the formed curd was cut and drained through the Whatman paper filter No 2. The separated whey and curd were taken for further analyses. The whey was condensed on the rotary evaporator (approx. 10x) and 120 mL of concentrated whey was lyophilized. The dry residue was then loaded into the Soxhlet apparatus and the extraction process was carried out according to the described procedure.

The extraction was preformed in three replications for each sample. Additionally, the blank extraction was performed in each series.

#### **Preparation of standard solution**

A stock solution of antineoplaston standards was prepared by dissolving 0.100 g of each compound in 10 mL of DMF. Then, the standard solutions used for the calibration were prepared by diluting the stock solution with DMF to obtain the target concentrations of: 2.0 mg/mL, 1.0 mg/mL and 0.5 mg/mL.

#### **HPLC analysis**

HPLC was carried out using a Shimadzu

HPLC system containing two LC-AS pumps, an SCL-10A controller, SIL-10A autoinjector, SUS variable volume injector and SPD-10A UV detector. All chromatography buffers were made with HPLC water, filtered through a 0.2 $\mu$ m hydrophilic filter (Millipore) and degassed under vacuum. The reversed-phase column,  $\mu$  Bondpak C<sub>18</sub> (Waters) had dimensions of 300 x 3.9 mm I.D. and 10  $\mu$ m particle size. The mobile phase was water: methanol: acetic acid (78:22:1 v/v/v) during the first 70 minutes and further was in a 59:41:1 (v/v/v) ratio. 45 $\mu$ L of the sample was injected into the HPLC and separated at 0.9 mL/min at a temperature of 27 $^{\circ}$ C. The analyzed compounds were detected at 254 nm. ANP levels, expressed in mg/mL, were calculated from the peak areas, using the EZ-Chrom Data System software from Shimadzu. The actual concentrations of antineoplastons in the particular dairy products were recalculated, considering the weight of the specimen taken for extraction as well as the final volume of the reconstructed solution.

## **RESULTS**

The detection of PG in cow's milk (26) inspired us to study the other ANP components, i.e. A10, IsoPG, PN and the ANP prodrug PB. Several samples of farmer's cheese, originating from various batches, were studied in order to determine quantitatively the presence of the compounds. It was found that PG is the most abundant ANP component not only in farmer's cheese, but also in all studied dairy products. Nevertheless, the results obtained varied significantly. The highest concentration of PG was calculated as 22.0 mg/100g (average of three replicates), while the lowest concentration was 1.0 mg/100g. Moreover, the chromatograms differed significantly, suggesting that the specimens from different batches were not uniform. The typical chromatogram is presented in the Figure 1.

The other compounds, i.e. isoPG, A10 and

TABLE 1. Antineoplastons in Dairy Products

	MILK (mg/100mL)	CONDENSED MILK (mg/100mL)	FARMER'S CHEESE (mg/100g)	FETA CHEESE (mg/100g)	WHEY (mg/100mL)
Phenylacetylglutamine (PG)	6.0	18.0	22.0	3.0	29.0
Phenylacetylisoglutamine (isoPG)	0.1	0.1	3.0	1.0	6.0
3-phenylacetyl-amino-2,6- piperidinedione (A10)	0.1	0.1	7.0	0.1	3.0
Phenylacetic acid (PN)	Not found	Not found	4.0	Not found	0.3
4-phenylbutyric acid (PB)	Not found	Not found	traces	Not found	Not found

The results are averages from 3 analyses

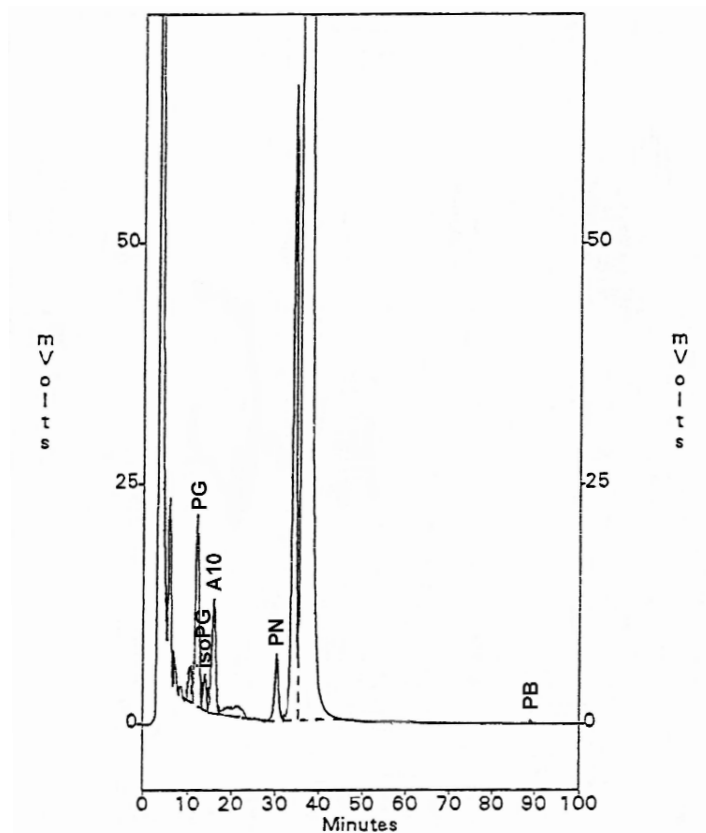


FIGURE 1. Typical HPLC chromatogram of antineoplaston ingredients in farmer's cheese.

A10—3-phenylacetyl-amino-2, 6-piperidinedione, PB—4-phenylbutyric acid, isoPG—phenylacetylisoglutamine, PG—phenylacetylglutamine, PN—phenylacetic acid.

PN were also found in farmer's cheese, but PB was not present. Their concentrations, however, were significantly lower (approx. 3.0-4.0 mg/100g) and did not vary so drastically between batches, as was the case of PG. Some analyses indicated trace amounts of PB, but the concentrations (less than a milligram per 100 g) were at the border of method sensitivity. The results are summarized in Table 1.

Our experiments confirmed the results of earlier studies on PG in cow's milk (26). In our studies we used fresh whole milk as well as condensed milk. We found that, similar to the farmer's cheese samples, milk contained mostly PG. The concentration of PG was approximately 3 times higher in the condensed milk than in the fresh milk. The differences in the PG concentrations can be explained simply by the more concentrated sample. The other compounds, i.e. isoPG and A10 were found in trace concentrations (less than a milligram per 100 g). Nevertheless, PN and PB were not found at all (see Table 1). The significantly lower concentrations of isoPG, and A10 can be caused by the process of the milk preparation, such as pasteurization.

Interesting results were obtained in the experiment with whey. The amount of PG determined in the whey was comparable to that present in farmer's cheese. These results were obtained for the whey sample, which was concentrated on the rotary evaporator and then deproteinized by addition of acetonitrile. The other compounds, isoPG, A10 and PN were found from trace amounts to 6.0mg/100g, comparable to those present in the other dairy products.

The analysis of feta cheese was more difficult to perform due to the high content of fat. PG was in concentrations approximately 7 times lower than in farmer's cheese. This may be explained by the different source of the product (sheep vs. cow) and the loss of some compounds during the preparation steps, especially during the fat extraction.

## DISCUSSION

A10 is present in human blood and urine (14, 27). Based on the findings described in this article it also exists, in small amounts, in farmer's cheese and whey and in trace amounts in cow's milk and feta cheese. A10 specifically intercalates with DNA and protects the sequences which are vulnerable to the effects of carcinogens, such as benzo [a] pyrene, urethane, and aflatoxin B<sub>1</sub> (28, 29). In animal tests, conducted at the Medical College of Georgia, University of Kurume Medical School in Japan and Burzynski Research Institute, mice and rats were protected from development of breast, lung and liver cancers when they were exposed to carcinogens and were fed a diet containing A10 (30-34). Human pharmacokinetics studies revealed that 70% of A10 is absorbed intact from the small intestine, but 30% is converted into PG and isoPG (35). In addition, PG is biosynthesized in the liver from glutamine and phenylacetate (22, 36). PG was detected in cow's milk by researchers from the USDA and by our team (as described in this paper) in milk, farmer's cheese, whey, and feta cheese (26). It is also derived from PB, which is present in lamb (25). PG exhibits antineoplastic activity across a wide array of cancer cell lines, including breast, liver and glioblastoma multiforme (37, 38). It inhibits the uptake of growth-critical amino acids such as L-glutamine and L-leucine in neoplastic cells (38). PG enters cells by the stereospecific amino acid transporters and works as a competitive inhibitor of these transporters. It also normalizes the pattern of genome-wide methylation, stabilizes the genes, decreases expression of oncogenes and promotes apoptosis (38).

PN is present in relatively low concentration in human blood, urine, and in dairy products (39, 40). It is produced by normal intestinal bacterial flora and is generated from metabolism of PB (25, 39). It works as a molecular switch, which turns off the electrical signal in one of the

most important oncogene pathways, the *RAS* oncogenes, and activates the tumor suppressor genes *TP53* and *p210* (41-44). PN inhibits farnesylation of the *p21<sup>ras</sup>* protein and causes down-regulation of *BCL-2* through inhibition of mevalonate 5-pyrophosphate decarboxylase (41, 43, 45). PN activates the *TP53* and *p21* tumor suppressor genes through inhibition of methyltransferases (42, 44). It also binds excessive amounts of L-glutamine, a promoter of cancer growth (46).

PB exists in lamb and in trace amounts in farmer cheese and it is partially converted in the liver to PN and PG (24, 25). It activates tumor suppressor genes through inhibition of histone deacetylase (47).

A10 and PG were formulated together with selected amino acids and vitamin B-2 into two supplements currently available in the United States and in the European Union (48). In addition to cancer preventive effects, which were documented in animals, there have also been a number of other positive effects observed by users. Among these were anti-aging effects reported by individuals who take these supplements. Positive effects included increased energy, improved healing, reduction of wrinkles and hyperpigmentation spots, reduction of cholesterol concentration in blood, improved cellular immunity (with a decreased frequency of common cold and viral infections), improvement of benign prostate hypertrophy and a decrease of benign nodules in breasts, as well as an anti-depressant effect (48).

### CONCLUSIONS

The ingredients of antineoplastons: A10, PG, isoPG, and PN and the pro-drug PB were found in dairy products. PG occurs in the highest concentration of 29 mg/100 mL in whey, 6.0 mg/100 mL in milk, 22.0 mg/100 g in farmer's cheese and 3.0 mg/100 g in feta cheese. The other compounds were present in much smaller amounts (from trace to 7.0 mg/100 g) and these

varied significantly between samples. A10 and PN were found in the highest concentration of 7.0 mg/100 g and 4.0 mg/100 g correspondingly in farmer's cheese. PB occurred only in trace amounts in farmer's cheese.

Prevention of cancer and age management by phytochemicals and non-nutritional components of the diet is now considered to be an inexpensive and acceptable practice. The system of small molecules, such as polyphenols in plants and amino acid derivatives, carboxylic acids, and peptides in animal products apparently protect various organisms from formation of cancer and has an impact on aging. Our research group postulated the existence of such a system in 1976 (49). As early as 1980, the NCI's Chemoprevention Programme began evaluating phytochemicals for safety and efficacy. In 1998, NCI's Division of Cancer Prevention started the Chemoprevention Implementation Group. NCI has more than 400 potential agents under investigation and is sponsoring over 60 chemoprevention trials (6). The European Union, through its EPIC program (European Prospective Investigation into Cancer and Nutrition), conducts larger and more ambitious investigations (7). Over 500,000 volunteers in ten European countries are included in the studies, which link dietary, biochemical, and genetic analysis. The chemopreventive effects of most dietary products are the sum of several distinct mechanisms. In many cases the concentration of active chemicals in food is not sufficient, which is typical for substances derived from animal products. Therefore, it will be necessary to take supplements in addition to a proper diet. The new term "neutrigenomics" has been introduced, with more and more attention devoted to food supplements, which affect the genes. It is expected that, in the near future, designer foods will be available containing a chemopreventive and anti-aging agents and they will have a substantial impact on reducing the incidence of cancer.

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## WILL THE REAL HYALURONAN PLEASE STAND UP?

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This review intends to inform readers about the identity of Hyaluronan (Hyaluronic Acid or HA) materials currently used in dietary supplement products. As contestants from the TV game show “To Tell The Truth” (1956-1978) were made to choose the real person from two other foils by asking questions of each, this review will answer questions so that the real HA can stand up. There are major differences between the three major types of HA materials used in dietary supplements. According to the voluminous scientific literature, these differences may lead to distinctly different biological effects. This review does not attempt to discuss the rationale for oral HA as opposed to injectable or topical uses – this is an obvious topic for further review. However, ingestible HA products are available to the general public, and their use is growing to the point that HA-containing products are available from almost every mass-market retail provider. Consumers and health care professionals alike are not given facts about the identity of HA materials used in dietary supplements. Informed choices for evaluation of HA-containing products cannot be made without an understanding of what materials are used for each product.

### HA PRIMER

HA is an extremely large molecule (normally 1-5 million daltons) of repeating subunits of N-acetylglucosamine and glucuronate (1-14). HA is classified as a glycosaminoglycan (GAG), and is present physiologically and commercially as the sodium salt (sodium hyaluronate). Single HA molecules exist as a hydrophilic, coiled mass approximately 200 nm in diameter, but form a mesh under typical *in vivo* conditions (8). HA is found everywhere in the body, but has especially important roles in joints. HA is the primary agent responsible for the viscosity and lubricating properties of synovial fluid (6, 8, 11-29). In cartilage, HA serves as the backbone of aggrecan proteoglycans, bound to core and link proteins (18, 30-49). Furthermore, HA is an impor-

tant and under-recognized regulator of cell and tissue functions, as shown by the presence of specific receptors on all cells for HA and its fragments (8, 11-14, 50-65).

The scientific and medical literature on HA and its effects on joints has had virtually no references to oral use of HA until very recently. Instead, HA has a prodigious literature on biological effects (6, 8, 11-14, 19-26, 28, 62, 66, 67), including a rather large number of *in vivo* animal and human clinical studies using intraarticular injections to treat osteoarthritis (OA) with purified HA (reviewed in 21, 22, 25, 26, 68-86). Thus, there is substantial knowledge of the biological effects of purified, high molecular weight HA on joint health.

### HYALURONAN AS AN ORAL DIETARY SUPPLEMENT

HA as an oral supplement entered mainstream consciousness after an ABC News pro-

gram aired a segment that attributed possible anti-aging effects to dietary HA in November 2002. Since then, HA has been appearing as an ingredient in an ever-increasing number of dietary supplement products. HA-containing products have focused on joint and skin health. At the beginning of 2005, several of the largest dietary supplement companies were selling products containing HA in major retail outlets throughout the United States. This is in addition to other products available through Internet sales. In short, HA has catapulted from obscurity to mainstream store shelves rapidly.

There are a few animal studies on joint health after intravenous or intraperitoneal administration of HA (87-89), which provide some insight into tissue uptake and possible effects of HA after its appearance in the bloodstream. Four recent reports investigated effects of oral HA on animal musculoskeletal tissues. One report from the Hyaluronan 2003 Proceedings by Matrix Biology Institute administered 100 mg HA orally to 12 racing thoroughbreds for 59 days (90). Horses given HA were examined for lameness less frequently than 13 control group horses, suggesting that oral HA prevented lameness in active horses. Another report by Stancikova *et al* administered HA (0.5-1 mg/kg) with molecular weights of 0.75 and 1.62 million daltons by oral routes for eight weeks to ovariectomized rats (91). Both sizes of HA showed increased serum concentrations of nitric oxide (NO), but only the larger HA showed decreased markers of bone resorption as well as an inhibition of bone mineral density loss. Two other animal studies were found on the Contipro website (92). Oral HA fed to rats at a dose of 0.1 mg/kg showed reduced swelling of hind leg and decreased NO production after arthritis induction by Freund's adjuvant. An open study from 7 veterinary clinics studied oral HA given to 53 racehorses with inflammatory and degenerative joint conditions. After 30 days of oral HA, symptoms were reduced and functionality improved. All reports used purified, high-mole-

cular weight HA from a microbial fermentation source (Nutrihyl®, Contipro Group Holding, Czech Republic).

There are unpublished reports of oral HA in humans for joint health from the Czech Republic. A product named Chondrorevit was reported to improve joint health in subjects with knee OA or after knee surgeries (92). Daily doses of 20 mg HA and 400 mg chondroitin sulfate were administered for 90 days. Improvements in joint function or recovery time after surgery were reported. Interestingly, this amount of chondroitin sulfate has not been shown to have significant efficacy in dose-finding studies from Europe (93), and thus, any benefits may be attributable to the HA or the combination of HA and chondroitin at a less than efficacious dose. However, these studies did not appear to be randomized, double blind or placebo controlled, and thus, their findings must be verified in controlled studies before results can be relied upon. Nevertheless, HA has been reported to have benefits for joint health after oral use in animal and human clinical studies. At this point in time, reports of oral HA for joint health have all used high molecular weight, purified HA, except for one study discussed later in this review.

#### **MOLECULAR WEIGHT AND BIOLOGICAL EFFECTS OF HYALURONAN: EVIDENCE SHOWING THAT SMALL HYALURONAN AND FRAGMENTS ARE DIFFERENT**

Because HA is a repeating polymer devoid of modifications (such as branching or sulfation), major differences in biological activities for HA are attributable to its molecular weight (size) (25, 65, 67, 94-98). There are clear cutoffs of molecular weight for lubricating properties (17, 19, 25, 99-106), cell signaling (107-117), and perhaps for efficacy of injectable intraarticular preparations for OA treatment (63, 67, 118-120). Molecular weight of less than 500,000 daltons seems to be

the cutoff for differing properties of HA. This concept is summarized by a comment from Camenisch and McDonald: “The work of Ohkawara and coworkers and numerous other observations suggest that there is something fundamentally different about the biological response to high (megadaltons) and lower molecular weight HA.” (95). In essence, bigger is better for health benefits from HA, and smaller fragments have very different or opposite properties than native, high molecular weight HA.

Very low molecular weight HA fragments cannot be expected to have the same biological effects as high molecular weight HA, as is clear from the growing literature on biological effects of HA. Very low molecular weight fragments either do not bind to specific cell membrane receptors that recognize native HA, or they transduce different signals to cells (65, 107, 108, 113, 121-126). For example, very low molecular

weight HA fragments (6.9 kDa), but not high molecular weight HA, have been associated with promotion of cancer cell neovascularization, migration and metastasis (112, 117, 122, 127-132). Low molecular weight HA fragments have either lost anti-inflammatory effects or instigate proinflammatory, catabolic properties compared to native HA in several biological systems (64, 110, 114, 116, 121, 125, 132-141). For example, HA fragments of 250,000-dalton molecular weight induced expression of inflammatory genes in macrophages (64, 65, 132, 135, 142). These effects are not due to contamination, as shown by the fact that the size range of HA tested *in vitro* is the same range as is found *in vivo* during inflammation (143). Some reviewers have postulated that HA fragments promote and propagate OA (65, 97, 136, 140, 144). These are obviously not desirable traits for dietary supplements purporting to support joint health.

TABLE 1: Identity and Characteristics of Hyaluronan Materials Used in Dietary Supplements

Hyaluronan Type	Source	Molecular Weight (daltons)	Purity	References
Hydrolyzed chicken sternal cartilage Type II Collagen	Chicken sternal cartilage	50–10,000 <sup>1</sup>	~10%	<ul style="list-style-type: none"> <li>• Ishaq S. Hyaluronic acid and chondroitin sulfate based hydrolyzed collagen type II and method of making same. United States Patent 6,780,841, August 24, 2004.</li> <li>• <a href="http://www.biocelltechnology.com">http://www.biocelltechnology.com</a></li> </ul>
Rooster Comb (Injuv™)	Chicken comb	50,000–200,000	9%	<ul style="list-style-type: none"> <li>• Udell RG, Naguib YMA. Hyaluronic acid in soft gel form. United States Patent 6,806,259, October 19, 2004.</li> <li>• <a href="http://www.injuv.us">http://www.injuv.us</a></li> </ul>
Sodium hyaluronate	Microbial fermentation (Streptococcus species) or chicken comb	700,000–1 million or higher	90+%	<ul style="list-style-type: none"> <li>• <a href="http://www.bioiberica.com/jointcare/hyaljoint.htm">http://www.bioiberica.com/jointcare/hyaljoint.htm</a></li> <li>• <a href="http://www.cpn-contipro.com">http://www.cpn-contipro.com</a></li> </ul>

<sup>1</sup>Patent 6,780,841 (column 6, line 12) stated: “The average molecular weight of the final product is between 50 and 10,000 daltons, preferably 5,500 daltons.” Neither the patents nor website specifically list a range of molecular weight for hyaluronan itself. Methods of analysis to determine molecular weight or identity of HA were not stated.

## HA MATERIALS

There are three major types of commercially available HA materials used in dietary supplements. Table 1 lists these types and their basic attributes. Information was obtained in the public domain from material supplier or supplement company websites. It is readily apparent that there are major differences in molecular weight of each type, yet all are labeled identically as Hyaluronic Acid or Sodium Hyaluronate on dietary supplement product labels (see Table 2). Only one type of HA matches what is normally present in the human body – high molecular weight, purified (protein-free) HA. The other two major types of HA materials have lower molecular weights.

### HYDROLYZED CARTILAGE HA (BIOCELL COLLAGEN II™)

HA from hydrolyzed cartilage has a patent describing the enzymatic degradation of Type II collagen from chicken sternal cartilage into very small fragments (50 to 10,000 daltons), with no high molecular weight material remaining (145). This process results in a material consisting of 10% Hyaluronic Acid, 20% depolymerized chondroitin sulfate, and around 63% Type II collagen fragments (145). This review is not concerned with the collagen or chondroitin contents of this material, which may have their own effects on joint health. However, the HA in this material is pertinent since it is used in dietary supplement products listed as a source of HA (see Table 2). It is not stated in the patent or in company website information what size range of HA fragments exist in this material. The patent further states (column 7, line 46):

“By extracting HA from chicken sternal cartilage, a low molecular weight HA may be obtained by hydrolysis after such extraction. This distinction is crucial because the beneficial therapeutic activity of HA is mostly

dependent upon the molecular weight of HA. Due to the low molecular weight of the HA found in the hydrolyzed collagen type II, the hydrolyzed collagen type II readily absorbs into the gastrointestinal tract and allows the rejuvenating constituents of HA and CSA to restore viscoelasticity to the skin, protect connective tissues, promote cartilage synthesis, retain skin moisture, heal wounds and improve the overall appearance of skin.”

The patent does not supply any data on the size of HA, the analytical method used to assess HA identity, or any effects of the material. In other words, there is no supportive data to substantiate the properties attributed to HA in the statement above. However, it is clear that the inventor assumes that the HA in this material is of low molecular weight.

As briefly discussed previously in this review, HA fragments do not have the same properties in biological systems as native, high-molecular weight HA does, and thus, this material does not fit the definitions and descriptions of HA agreed upon by textbooks, reviewers and experts in the field (8, 11-14, 19, 23-26, 28, 62, 113, 125, 126). In short, this material is not typical HA and thus should not be described as HA on dietary supplement labels. Rather, this material is more accurately described as very low molecular weight HA oligosaccharides or even pre-digested HA. This statement assumes that the HA in this material is indeed hydrolyzed, which at present is not substantiated.

Furthermore, HA in cartilage does not exist as free, unbound molecules, but rather, as the backbone of aggrecan proteoglycans (18, 30-49). As such, HA is bound to link and core proteins, and sometimes to other GAGs or chondrocytes themselves. Even assuming proteolytic enzyme treatment as described in the patents to prepare this material can cleave HA from protein (145-147), the presence of amino acids crosslinked to HA fragments are not synonymous with a definition of HA.

TABLE 2. Examples of How Dietary Supplement Products Containing Hyaluronan Are Labeled<sup>1</sup>

Product	Description of Hyaluronan in Supplement Facts	Amount per serving	Accompanying Label Claims
21 <sup>st</sup> Century Healthcare, Inc. Arthri-Flex	Hyaluronic Acid	Not stated—part of a Flexicol <sup>TM</sup> ) proprietary blend with four other ingredients totaling 645 mg	<ul style="list-style-type: none"> <li>• With Flexicol Premium Chicken Sternum Type II Collagen</li> <li>• Containing: Premium Chicken Sternum Cartilage Type II Collagen (naturally occurring Chondroitin Sulfate and Hyaluronic Acid)</li> <li>• Contains Highly Absorbable Hyaluronic Acid</li> <li>• <a href="http://www.21stcenturyvitamins.com">www.21stcenturyvitamins.com</a></li> </ul>
Arthritis Research Corp. Flex-a-min <sup>®</sup> Complete <sup>TM</sup>	Hyaluronic Acid (as Sodium Hyaluronate)	Not stated—part of a proprietary blend with three other ingredients totaling 400 mg	<ul style="list-style-type: none"> <li>• Hyaluronic Acid—The Glucosamine in Flex-a-min<sup>®</sup> Complete<sup>TM</sup> promotes the production of Hyaluronic Acid in the body, including the synovial fluid surrounding joints.</li> <li>• <a href="http://www.flexamin.com">www.flexamin.com</a></li> </ul>
Arthritis Research Corp. Flex-a-min <sup>®</sup> Triple Strength	Hyaluronic Acid (as Sodium Hyaluronate)	Not stated—part of proprietary blend with Silica totaling 40 mg	<ul style="list-style-type: none"> <li>• Plus, we added the popular ingredient Hyaluronic Acid to set this formula apart.</li> <li>• Hyaluronic Acid—The Glucosamine in Flex-a-min<sup>®</sup> Complete<sup>TM</sup> promotes the production of Hyaluronic Acid in the body, including the synovial fluid surrounding joints.</li> <li>• <a href="http://www.flexamin.com">www.flexamin.com</a></li> </ul>
Injuv <sup>TM</sup> Hyaluronic Acid	Hyaluronic Acid	12.6 mg (from 140 mg Injuv <sup>TM</sup> ) <sup>2</sup>	<ul style="list-style-type: none"> <li>• Stop the stiffness, stop the pain and increase your mobility with Injuv<sup>TM</sup> (<a href="http://www.jointlubricant.com/injuvworks.htm">www.jointlubricant.com/injuvworks.htm</a>)</li> <li>• The HA in Injuv<sup>TM</sup> stops pain and inflammation of joints, because it helps to hydrate and lubricate the synovial cell membrane. (<a href="http://www.jointlubricant.com/injuvworks.htm">www.jointlubricant.com/injuvworks.htm</a>)</li> <li>• Injuv<sup>TM</sup> has an exclusive patent on the process that breaks down HA so that it can be taken orally.</li> <li>• The hyaluronic acid molecule in Injuv has been enzymatically cleaved into smaller fragments. (<a href="http://www.injuv.us">www.injuv.us</a>)</li> </ul>
Nature's Bounty Hyaluronic Acid	Hyaluronic Acid (as Sodium Hyaluronate)	20 mg per capsule	<ul style="list-style-type: none"> <li>• Hyaluronic Acid is found in almost all adult connective tissue. Hyaluronic acid is a natural component of the body.</li> <li>• <a href="http://www.NaturesBounty.com">www.NaturesBounty.com</a></li> </ul>
Schiff <sup>®</sup> Move Free Triple Strength	Hyaluronic Acid (in Other ingredients list)	Not stated (Consumer Service line stated 3.3 mg per serving)	<ul style="list-style-type: none"> <li>• Now with Joint Fluid (Hyaluronic Acid)</li> <li>• <a href="http://www.schiffvitamins.com">www.schiffvitamins.com</a></li> </ul>
Product			

TABLE 2. Continued

Product	Description of Hyaluronan in Supplement Facts	Amount per serving	Accompanying Label Claims
Spring Valley Glucosamine & Collagen with Chondroitin & HA (Hyaluronic Acid)	Hyaluronic Acid	40 mg	<ul style="list-style-type: none"> <li>• In a base of amino acids (Hydrolyzed Collagen Type II)</li> <li>• Collagen Type II 500 mg</li> <li>• Chondroitin Sulfate (Avian, Porcine, Bovine) 100 mg</li> <li>• Collagen is the most plentiful protein and building block in the body and is a natural source of Chondroitin, a nutrient shown to regenerate joint tissue, and Hyaluronic Acid (HA), a revolutionary compound that retains moisture and functions as a lubricant between connective tissues.*</li> <li>• Hyaluronic acid is fast absorbing in the blood and thus available for joint tissues.</li> <li>• Lot number 4MA0236</li> </ul>
Swanson Ultra® Hyal-Joint™	Hyal-Joint™ Hyaluronic Acid	20 or 50 mg per capsule.	<ul style="list-style-type: none"> <li>• The unique hyaluronic acid formulation in Hyal-Joint has the proven ability to be absorbed in the intestinal tract and has demonstrated significant joint-protective properties in numerous studies.</li> <li>• <a href="http://www.bioiberica.com">www.bioiberica.com</a></li> </ul>
Windmill™ Health Products Glucoflex™ Hyaluronic Acid	Hyaluronic acid sodium salt	10 mg per tablet	<ul style="list-style-type: none"> <li>• Advanced Joint Lubricating Formula with Glucosamine</li> <li>• Glucoflex™ Hyaluronic Acid formula with Glucosamine supplies the most bioavailable form of HA (Hyaluronic Acid) for the most direct delivery of this amazing nutrient.</li> </ul>

<sup>1</sup>Products were obtained from store shelves from the United States or from internet website information. Products are listed in alphabetical order according to manufacturer name.

<sup>2</sup>Some products containing Injuv™ HA list the total amount of material instead of the actual amount of HA, misleading consumers into believing there is 70 or 140 mg of HA per softgel.

Another issue apparently not considered by some marketers of hydrolyzed cartilage HA is the integrity of borrowing literature describing properties of HA to support such product claims (see Table 2). Since hydrolyzed HA has quite different physical and biological properties compared to high molecular weight HA, it is a different entity. For products with very low molecular weight HA (hydrolyzed cartilage) it is inappropriate to make claims, such as lubricating properties, which rely on literature describ-

ing high molecular weight HA. This is especially true when there is also evidence of very low molecular weight HA not having the properties claimed (such as lubricating properties). Thus, it would be more appropriate to base any claims for joint health with this material on studies using this material.

Recently, a late-breaking abstract presented at the 2004 FASEB meeting in Washington DC described the results of a human clinical study using hydrolyzed cartilage material (148). A press

release from BioCell Technology, LLC posted on their website on April 2003 also reiterated the results of the abstract (149). In the abstract, the material was described as “. . . a hydrolyzed type II collagen product (avian source: Biocell™; HC2)” without mention of HA. A dose of 2000 mg daily was given to eight subjects with knee or hand osteoarthritis for two months, and compared to a placebo group of eight subjects in a randomized, double blind, placebo-controlled study. WOMAC scores were improved significantly greater in the HC2 group compared to placebo, but no data were presented in the abstract or press release. This amount of material would contain approximately 200 mg of hydrolyzed HA daily, according to the supplier's website and patent. However, approximately 1600 mg of hydrolyzed Type II collagen and 400 mg of chondroitin sulfate fragments were also present in this material. Thus, the contribution of HA to observed results is unclear. Furthermore, human studies with subjective questionnaire endpoints (WOMAC) and with only eight subjects per group run a high risk of low statistical power, making results preliminary until confirmed in more powerful studies. Most other studies that mixed subjects with knee and hand osteoarthritis have been criticized since the arthritic process in hand osteoarthritis is distinctly different than in knees.

In addition, a press release by BioCell Technology from February 2, 2004 described a 36-hour peak absorption study conducted in an unknown number of human subjects with hydrolyzed cartilage material (150). HA levels were determined after a single dose and after 28 days of administration. Levels of HA in blood were claimed to be 7008.62% above control levels after 12 hours. Also, two HA metabolites 1/600th the size of the ingested HA were found. After 28 days, HA levels in blood were reported to be 3542.58% above control, and metabolites at 11,890.15% above control. No details on blood collection or analytical methods were given, although the reporting of HA metabo-

lites with two decimal places implies a high degree of precision for the method of analysis. Normal human serum levels of HA are 0.01-0.1 mg/L (mcg/ml) (8). Assuming the press release was describing serum HA levels, a 7000% increase in blood HA levels would produce levels of 0.7-7 mcg/ml, which is within the range seen for certain cancers, liver conditions and inflammatory diseases (8). One possible interpretation of this study is that this material elevates serum levels of HA fragments, along with native HA levels, although more experimental details are needed to confirm this interpretation because the effects of activity and food intake are known to increase plasma HA levels (151). Absorption of HA from the material is not proven by these results, since HA can rapidly enter circulation, especially in inflammatory diseases (8). Thus, another interpretation of these results is that this material is pro-inflammatory, which in and of itself would elevate serum levels of HA, congruent with findings of other researchers on pro-inflammatory effects of very low molecular weight HA fragments.

Assay of HA fragments, especially in a mixture with chondroitin sulfate or its hydrolysis products, is problematic. In fact, assay of products containing very low molecular weight HA fragments by validated assays utilizing HA binding protein technology do not find any measurable HA (Bucci, L., unpublished results). This indicates that if present, HA fragments are less than eight subunits in length. These assays also indicate that there is no high molecular weight HA present, so claims for this material on product labels or websites based on properties of high molecular weight HA cannot be substantiated by borrowing from the literature describing high molecular weight HA.

Therefore, HA derived from hydrolyzed chicken sternal cartilage is characterized by extremely low molecular weights, a lack of identity as native HA itself, a probable difference in biological effects (assuming uptake and distribution to tissues), a lack of transparency on how

the HA content was verified, and inability to show presence of HA in validated assays for HA using specific HA binding protein methodology. In addition, some products do not describe this material as hydrolyzed (see Table 2), which misleads consumers into thinking that they are getting native HA. Consumers expect to receive high molecular weight HA since this is what media, medical usage and website information all describe as characteristic of HA. This material is easy to identify in products because of the presence of Type II Collagen, chicken (avian) sternal cartilage source, and simultaneous presence of HA, chondroitin sulfate and Type II collagen naturally. These facts argue strongly that this material should not be labeled as HA in dietary supplement products as the identity and putative biological effects are much different from that of high molecular weight HA.

#### **ROOSTER COMB HA (INJUV™)**

The second major type of HA used in dietary supplement products is prepared from dried rooster comb, a rich natural source of HA (7500 mg/L) (8). This material is claimed to be enzymatically denatured, with a molecular weight of 50,000 to 200,000 daltons in order to facilitate uptake, without any supporting evidence (152). Injuv™ HA is identified by validated HA binding protein assays (Bucci L, unpublished data). However, experts in the field regard HA with a molecular weight of 50,000-200,000 daltons to be low molecular weight HA. Thus, HA in this size range has lost its ability to form networks in solution, and thus, has lost viscosity and lubricating properties.

Inspection of HA in intact rooster combs shows tight association of HA with proteins (153-155). It is unclear if conditions used to prepare rooster comb HA with enzymatic hydrolysis would actually dissociate the tightly bound proteins from the HA, since details of preparation are lacking in the published patent. High molecular weight HA prepared from rooster combs for

injectable use, such as Hyalgan® Sodium Hyaluronate from Sanofi-Synthelabo Inc., has been extensively treated to remove proteins in order to prevent reactions after injection, and has not shown appreciable adverse effects after intraarticular injections (156). Such material would fit into the third type of HA material.

HA with molecular weights of less than 500,000 daltons exhibit decreased viscosity and lubricating properties in solution, as stated earlier in this review. HA in this size range has lost its ability to form networks in solution. Therefore, claims about lubricating properties are inappropriate with this material unless specific studies are performed to investigate its lubricating effects in joints. Also, signals transduced by HA in the molecular weight range of 100,000-200,000 are different from those of HA at 1 million daltons, and generally are proinflammatory. The presence of tightly bound proteins in rooster comb HA has uncertain implications for uptake of this type of HA. Digestion and uptake of rooster comb HA may be different from native, purified HA. So this material, while clearly low molecular weight HA, is different from native, high molecular weight HA and has been shown to have different biological effects.

#### **PURIFIED HA**

The third major type of HA available for dietary supplement products is essentially purified HA (Table 1). Typical commercial sources are from microbial fermentation (*Streptococcus* species) or rooster combs. Molecular weight is around 1 million daltons, which corresponds to native HA in cartilage and in synovial fluid. Protein is absent or very low in these preparations. HA from microbial fermentation sources can be declared as vegetarian, whereas HA derived from any cartilage or chicken comb sources cannot. Since purified HA for food use is produced by manufacturers that also produce purified HA for pharmaceutical use, there is more certainty that these products have a tradition of quality

TABLE 3. Biological Properties of High Molecular Weight Hyaluronan and Hyaluronan Fragments

Molecular Weight (daltons)	Biological Property	Reference
Very low molecular weight oligosaccharides (800-5000)	<ul style="list-style-type: none"> <li>• Angiogenic—Induced angiogenesis in rat skin grafts, healing wounds, chick chorioallantoic membranes</li> <li>• Angiogenic—Induced rapid transient up-regulation of the immediate early genes c-fos, c-jun, jun-B, Krox-20 and Krox-24 that control angiogenesis</li> <li>• Angiogenic—Increased tyrosine kinase, protein kinase C cascades in endothelial cells</li> <li>• Angiogenic—Induced proliferation of bovine brain and aortic endothelial cells</li> <li>• Antiinflammatory—Tetrasaccharides, but not larger sizes of HA, induced heat shock protein 72 in K562 cells and prevented cell death of PC12 cells after serum deprivation.</li> <li>• Antioxidant—Exhibited antioxidant activity against superoxide, but less than higher molecular weight HA; did not scavenge hydroxyl radicals (unlike high molecular weight HA)</li> <li>• Catabolic—Did not affect sulfate incorporation in chondrocyte cultures</li> <li>• Catabolic—Blocked binding of aggrecan HA to chondrocytes, resulting in loss of staining for proteoglycans (disrupted cartilage structure)</li> <li>• Catabolic—Caused chondrolysis (loss of chondrocytes) when hexasaccharides exposed to cartilage tissue slices and chondrocyte cultures</li> <li>• Inflammatory—High molecular weight HA cleaved at sites of inflammation into small fragments</li> <li>• Inflammatory—Activated macrophages and dendritic cells in healing wounds</li> <li>• Inflammatory—Induced maturation of dendritic cells to tissue macrophages</li> <li>• Inflammatory—Induced production of IL-1beta, TNF-alpha, IL-12 and iNOS in dendritic cells and macrophages</li> <li>• Inflammatory—Did not inhibit arachidonic acid release after bradykinin from synovial fibroblasts from osteoarthritic subjects</li> <li>• Inflammatory—Did not inhibit MMP-1 and RANTES expression or production in normal and osteoarthritic human chondrocytes in culture</li> <li>• Inflammatory—Induced expression of matrix metalloproteinases, but not TIMPs, in normal rabbit and bovine cartilage chondrocytes</li> <li>• Inflammatory—Induced MMP-9, MMP-13 expression and production in murine fibroblasts and 3LL cells</li> <li>• Inflammatory—Stimulated MAP kinase and urokinase-type plasminogen activator in human HCS-2/8 chondrosarcoma cells</li> <li>• Inflammatory—Did not inhibit neutrophil adhesion or aggregation under conditions of shear and turbulent flow</li> <li>• Inflammatory—Stimulated tyrosine phosphorylation and c-Met expression in human chondrosarcoma cells</li> </ul>	7, 8, 11, 15, 20, 28-32, 35, 38, 39, 42, 46, 52, 53, 57, 59, 60-63, 67-69, 71-73

TABLE 3. Biological Properties of High Molecular Weight Hyaluronan and Hyaluronan Fragments (*continued*)

Molecular Weight (daltons)	Biological Property	Reference
24,000	<ul style="list-style-type: none"> <li>• Catabolic—Inhibited tumor growth, induced apoptosis</li> <li>• Inflammatory—Induced expression and production of inflammatory mediators from murine macrophages (IL-1, TNF-alpha, IL-12, iNOS, metalloelastase)</li> <li>• Lubrication—Reduced elasticity &amp; viscosity of 1.3MDa HA solutions (competed for meshwork formation)</li> </ul>	13, 23, 36, 37, 43, 44, 54
33,000	<ul style="list-style-type: none"> <li>• Antiangiogenic—Did not induce angiogenesis of rat skin grafts</li> </ul>	35
<50,000	<ul style="list-style-type: none"> <li>• Catabolic—Did not stimulate HA synthesis or inhibited synthesis (high concentrations) of HA from human osteoarthritic synovial fibroblasts</li> <li>• Inflammatory—Did not inhibit proliferation of human mononuclear cells</li> </ul>	47, 58
50,000	<ul style="list-style-type: none"> <li>• Inflammatory—Did not inhibit MMP-1 and RANTES expression or production in normal and osteoarthritic human chondrocytes in culture</li> <li>• Inflammatory —Induced expression and production of inflammatory mediators from murine macrophages (IL-1, TNF-alpha, IL-12, iNOS, metalloelastase)</li> </ul>	23, 36, 37, 43, 44, 61
90,000	<ul style="list-style-type: none"> <li>• Catabolic—did not enhance bone mineralization in rat calvarial cells</li> <li>• Inflammatory—Did not inhibit phagocytosis by mouse peritoneal macrophages at same viscosity as higher molecular weight HA</li> <li>• Lubrication—Osmotic pressure opposing fluid drainage in rabbit joints reduced, facilitates escape of fluid and HA from joints</li> </ul>	6, 10, 24
80,000–200,000	<ul style="list-style-type: none"> <li>• Antiinflammatory—Did not induce matrix metalloproteinase production in murine fibroblasts and 3LL cells</li> <li>• Antiinflammatory—Did not induce IL-1beta, TNF-alpha or IL-12 in human dendritic cells or macrophages</li> <li>• Antioxidant—Scavenged hydroxyl radicals in vitro, but was less effective than high molecular weight HA for protecting tendon fibroblasts from hydroxyl radicals</li> <li>• Inflammatory—Did not inhibit cell proliferation of rabbit synovial cells</li> <li>• Inflammatory—Induced expression and production of inflammatory mediators from murine macrophages (IL-1, TNF-alpha, IL-12, iNOS, metalloelastase)</li> </ul>	8, 16, 23, 36, 37, 43, 44, 49, 62, 63
200,000	<ul style="list-style-type: none"> <li>• Inflammatory—Markedly stimulated iNOS mRNA in liver endothelial and Kupffer cell types</li> <li>• Inflammatory—Intravenous HA (250,000 daltons) did not inhibit proinflammatory cytokine levels in serum after liver injury in rats</li> <li>• Inflammatory—In human eosinophils, increased production of intercellular adhesion molecule-1, TGF-beta, protein secretion, and transformed cells from round to spindle shapes</li> </ul>	19, 40, 45, 51

TABLE 3. Biological Properties of High Molecular Weight Hyaluronan and Hyaluronan Fragments (*continued*)

Molecular Weight (daltons)	Biological Property	Reference
	<ul style="list-style-type: none"> <li>• Inflammatory—Human peritoneal mesothelial cells produced more inflammatory cytokines (MCP-1, IL-8)</li> </ul>	
280,000	<ul style="list-style-type: none"> <li>• Anticatabolic—Inhibited cartilage degradation by neutrophils, but significantly less than 950,000 or 2.0 million dalton HA</li> <li>• Inflammatory—Produced inflammatory cytokines RANTES, IL-12, MIP from murine macrophages</li> </ul>	21, 65
350,000	<ul style="list-style-type: none"> <li>• Lubrication—Formed poor meshworks with reduced viscosity</li> <li>• Lubrication—Osmotic pressure opposing fluid drainage in rabbit joints reduced, facilitates escape of fluid and HA from joints</li> </ul>	6, 54
460,000–2.8 million	<ul style="list-style-type: none"> <li>• Antiinflammatory—Dose-dependent inhibition in phagocytosis from mouse peritoneal macrophages (steric hindrance)</li> <li>• Antiinflammatory—Did not induce matrix metalloproteinases in normal rabbit and bovine cartilage chondrocytes</li> <li>• Antinociceptive—Molecular weight-dependent improvement in abnormal gait in rats acute arthritis, indicating nociceptive effects with higher molecular weight HA</li> </ul>	1, 10, 17, 46
Native HA (approx. 800,000 to 6 million)	<ul style="list-style-type: none"> <li>• Anabolic—Stimulated synthesis of high molecular weight HA from human osteoarthritic synovial fibroblasts</li> <li>• Anabolic—Increased proteoglycan synthesis from IL-1Beta treated chondrocytes</li> <li>• Anabolic—Stimulated fibroblast proliferation in a collagen lattice, but not in fibroblast monolayer</li> <li>• Anabolic—Enhanced proteoglycan synthesis in human cartilage explants after fibronectin fragment treatment</li> <li>• Anabolic—Enhanced chondrocyte proliferation and chondroitin sulfate synthesis in rabbit chondrocytes embedded in collagen gels</li> <li>• Anabolic—Forms pericellular matrix (with aggrecan) in presence of chondrocytes</li> <li>• Anabolic—Improved bone mineralization in rat calvarial cells</li> <li>• Antiangiogenic—Inhibited stimulation of angiogenesis and early response genes by low molecular weight HA fragments</li> <li>• Antiangiogenic—Inhibited or did not induce angiogenesis of chick chorioallantoic membranes or endothelial cells</li> <li>• Antiangiogenic—Did not cause proliferation of bovine brain and aortic endothelial cells</li> <li>• Anticatabolic—Decreased reduction in proteoglycan synthesis from fibronectin fragments</li> <li>• Anticatabolic—Decreased cartilage degradation from fibronectin fragments</li> <li>• Anticatabolic—Decreased expression of matrix metalloproteinase 3 in human cartilage explants after fibronectin fragment treatment</li> <li>• Anticatabolic—Inhibited leukocyte elastase and lysozyme release</li> <li>• Anticatabolic—Inhibited neutrophil-mediated cartilage degradation (reduced release of sulfated GAGs from bovine nasal cartilage explants)</li> <li>• Antiinflammatory—Did not induce iNOS mRNA from liver cell types</li> </ul>	1-8, 10, 12, 15-18, 20-22, 24-29, 32-34, 38, 39-44, 45, 47-51, 53, 58, 59-63, 66, 68, 70-72, 74, 75

TABLE 3. Biological Properties of High Molecular Weight Hyaluronan and Hyaluronan Fragments (*continued*)

Molecular Weight (daltons)	Biological Property	Reference
	<ul style="list-style-type: none"> <li>• Antiinflammatory—Reduced IL-1beta, TNF levels in rat air pouches</li> <li>• Antiinflammatory—Did not induce IL-1beta, TNF-alpha or IL-12 in human dendritic cells or macrophages</li> <li>• Antiinflammatory—Inhibited PGE2 production by human osteoarthritic synovial cells and normal rabbit articular chondrocytes after IL-1alpha induction</li> <li>• Antiinflammatory—Did not induce differentiation of human dendritic cells into macrophages</li> <li>• Antiinflammatory—Decreased mitogen-dependent activation of human mononuclear lymphocytes</li> <li>• Antiinflammatory—Inhibited proliferation of rabbit synovial cells in culture</li> <li>• Antiinflammatory—Inhibited carrageenin-induced edema in dose-dependent manner</li> <li>• Antiinflammatory—Inhibited adjuvant arthritis in dose-dependent manner</li> <li>• Antiinflammatory—Inhibited arachidonic acid release induced by bradykinin in synovial fibroblasts of osteoarthritic humans</li> <li>• Antiinflammatory—Inhibited advanced glycation endproducts from activating NF-kappaB, IL-1alpha, IL-6, TNF-alpha</li> <li>• Antiinflammatory—Inhibited MMP-1, RANTES expression and production in normal and osteoarthritic human chondrocyte cultures</li> <li>• Antiinflammatory—Did not enhance MAP kinase, urokinase-type plasminogen activator, tyrosine phosphorylation or c-Met in human HSC-2/8 chondrosarcoma cells</li> <li>• Antiinflammatory—Enhanced production of TIMP-1 in normal bovine chondrocyte culture, lowered stromelysin/TIMP-1 ratios</li> <li>• Antiinflammatory—Decreased synovial PGE2 and bradykinin in arthritic rats after intraarticular injection</li> <li>• Antiinflammatory—Inhibited neutrophil aggregation and adhesion under conditions of shear and turbulent flow</li> <li>• Antiinflammatory—Intravenous HA (&gt;780,000 daltons) decreased proinflammatory cytokines and liver enzymes after liver damage in rats</li> <li>• Antiinflammatory—Did not cause eosinophils to produce inflammatory cytokines</li> <li>• Antiinflammatory—Did not induce inflammatory cytokines RANTES, IL-12, MIP from murine macrophages</li> <li>• Antioxidant—Decreased free radical formation from IL-1Beta treated chondrocytes</li> <li>• Antioxidant—Scavenged superoxide and hydroxyl radicals dose-dependently</li> <li>• Antioxidant—Protected tendon fibroblasts in culture from damage by hydroxyl radicals</li> <li>• Lubrication—High molecular weight HA (&gt;500,000 daltons) caused outflow buffering in rabbit joints, reducing drainage and loss of HA and fluid</li> <li>• Regulatory—Inhibited sulfate incorporation into chondrocyte cultures</li> </ul>	

TABLE 3. *Continued*

Note: Filion and Phillips determined that some (2/7) of both high (0.5-2 million daltons) and low (10,000-300,000) molecular weight preparations of HA were inflammatory (produced IL-12 and TNF from human monocytes) due to contaminating DNA (based on reduction of activity after DNase treatment). This suggested that a low percentage of individual HA preparations have potential to cause inflammatory events not due to the molecular weight of HA. However, many of the studies cited in Table 3 examined different molecular weights from the same source of HA, and noticed an effect dependent on molecular weight, indicating contamination was not involved. The large number of studies showing an effect dependent on molecular weight from a wide variety of investigators, preparations and biological sources also argues strongly that DNA contamination is not operative in at least a majority of studies, or else no differences dependent on molecular weight would have been found. Scott and Heatley provide tertiary structure evidence that smaller HA fragments are different from high molecular weight HA, helping to explain the differences in biological activities of denatured HA.

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control, toxicological testing and reproducibility than the other two types of HA material.

As mentioned previously in this review, HA with high molecular weight of 1 million daltons has viscosity and lubricating properties, and sends antiinflammatory, anabolic signals to cells. Indeed, one mechanism of action proposed for intraarticular HA treatments of OA is the normalization of cell signaling to the synovium from high molecular weight HA (66, 71). Table 3 lists some of the biological properties associated with high molecular weight HA as well as HA fragments. Products using high molecular weight HA match the identity of native HA in the body and from the scientific literature. Claims relating properties and actions of HA from products containing high molecular weight HA appropriately borrow evidence from the scientific literature. Thus, high molecular weight, purified, native HA is the most logical choice for use as HA material in dietary supplements.

## ORAL ABSORPTION OF HA

Of course, one prerequisite for any beneficial effects from HA in dietary supplement products is uptake into bloodstream and joint tissues after oral intake. Of the three types of HA materials, only one type, purified HA, has published evidence of oral absorption. An abstract from the 2004 FASEB meeting in Washington DC described uptake of labeled HA into the bloodstream and localization of the label into the joints and salivary glands of rats and dogs several hours after oral administration (157). The HA in this study was derived from microbial fermentation, and was 1 million daltons. Since 99m-technetium was used as the label, and control experiments showed that the label was not removed or exchanged from HA molecules (Bucci, L., unpublished data), the presence of the label in the joints is most likely explained by the presence of the HA absorbed after oral

administration. This study is the first published report of oral absorption and tissue uptake of oral HA and provides the rationale for HA in dietary supplements.

Another source of purified HA from rooster comb has stated on the company website that absorption of their material has been found (158). However, no details or data have been presented or published. Hydrolyzed chicken sternal cartilage material is likewise claimed to have not only absorption, but superior absorption to native HA because of its smaller size. This claim is also on product labels or accompanying text (see table 2 and previous sections of this review). Again, website information from the manufacturer describes that absorption was found in a study, but no details or data have been presented or published in a scientific forum. No head-to-head comparisons to other HA materials concerning uptake have been reported, rendering claims of superior absorption unsubstantiated and unsupported. Only one type of HA—purified, high molecular weight HA from microbial fermentation—has published evidence for oral uptake and distribution to connective tissues.

#### **BENEFICIAL PROPERTIES OF HA FOR JOINT HEALTH**

Oral HA products are mostly targeted to joint and skin health, according to structure and function claims on product labels and accompanying literature (for a sampling, see Table 2). These two uses coincide with the prevalence of HA in the human body (8). Almost all beneficial properties for joint health attributable to HA have been documented for high molecular weight HA unbound to protein. Extensive *in vitro* studies using purified HA have uncovered beneficial properties for joints (see table 3). In addition, publication of a large number of human clinical studies using intraarticular injection of HA has demonstrated direct benefits for joint health (reviewed in 21, 22, 25, 26, 68-86).

Of particular interest are the human studies showing repair or normalization of synovium and cartilage structure and architecture for long time periods after a course of injections to persons with OA (66, 71, 159-164). These changes were documented by before and after tissue biopsies, along with biochemical measurements long after the injected HA disappeared from joints. These studies show direct evidence in humans that presence of purified, high molecular weight HA in joints has long-term benefits months after exogenous HA was given. The results of the oral uptake study presented at FASEB (157) show that oral HA has the potential to reach joints, and provide a strong rationale for use of oral HA in dietary supplements. However, at this time, this rationale applies only to purified, high molecular weight HA from microbial fermentation sources.

#### **SUMMARY**

It is interesting to speculate that continued oral administration of HA might produce similar or superior effects to intermittent injectable HA, as has been seen for both glucosamine and chondroitin sulfate. However, dietary supplements containing HA are not equivalent due to the inherent properties of the three major types of HA commercially available as dietary supplement materials. Consumers and health care professionals need to be aware of the different types of HA and their very large differences in properties (even before ingestion). One source, hydrolyzed chicken sternal cartilage, is clearly unlike native HA, does not match the biological properties of native HA, and consequently should not be represented as HA to consumers on product labels. Rooster comb HA is significantly smaller than native HA (about 10-20% the size of native HA), and is tightly bound to connective tissue proteins. This type of HA does not match the literature on properties and benefits of native HA because of its smaller size and tight binding to proteins. Finally, high molecu-

lar weight, purified HA is available from vegetarian-compatible and animal sources, virtually identical to native HA and pharmaceutical preparations of HA. This material is analogous to the HA used to generate the extensive body of knowledge on HA, both its roles and its therapeutic effects in OA. HA from the microbial fermentation method of preparation has published evidence for uptake into joints after oral administration. From all considerations, purified, high molecular weight HA from microbial fermentation in dietary supplement products has the highest likelihood for matching the known attributes of HA. The real HA—purified, high molecular weight—can stand up now.

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## DEDICATION TO DR. MILDRED S. SEELIG— OBITUARY (1920–2005)

The field of nutrition, and particularly the role of magnesium in health and disease, has lost one of its most respected colleagues and scientist-physicians during this year, namely Dr. Mildred S. Seelig. Dr. Seelig was a prime mover in the magnesium field ever since she published the now classic paper on “the requirement of magnesium by the normal adult” in 1964 (1). She was involved in studying, advancing and teaching about this essential mineral, all over the world, for more than four decades. We should, therefore, recall Mildred’s undiminished and unselfish efforts to advance knowledge and application of the role of magnesium in clinical medicine.

Mildred incessantly explored and researched the latest literature and brought new, young promising scientists and physicians into the field. She befriended them and engaged them in ongoing discussions and exchange of ideas. One could say that Dr. Seelig was the “mover and shaker” of the magnesium field, or she might be referred to as the “backbone” of this field. Up to her last days, Mildred was a dynamo of ideas,

- (1.) Seelig MS: The requirement of magnesium by the normal adult. *Am J Clin Nutr* 14: 342-390, 1964.
- (2.) Seelig MS: *Magnesium in the Pathogenesis of Disease: Early Roots of Cardiovascular, Skeletal and Renal Abnormalities*. Plenum Press, NY, 1980.
- (3.) Seelig MS, Rosanoff A: *The Magnesium Factor: Prevention Treatment and Reversal of Cardiovascular Disease*. Avery, Penguin Putnam, NY, 2003.

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thoughts and a constant source of intellectual stimulation.

Among Dr. Seelig’s many endeavors, the Gordon Research Conference on “Magnesium in Health and Biochemical Processes” was initiated by her in 1978 in Plymouth, New Hampshire. She organized and chaired the second such conference in 1979. Since that time, Dr. Seelig was always consulted and she helped to organize the succeeding Gordon Conferences on Magnesium up to the past one, held in February 2005 in Ventura, CA, just days after her untimely passing. She was the planner and chairperson of the first annual meeting of The American College of Nutrition in 1975 as well as the College’s subsequent meetings in 1977 and 1978. Dr. Seelig also was the coordinator for the Second and Third International Symposia on magnesium held in Montreal and Baden Baden, respectively, in 1976 and 1981. She was the President and Organizer of the fourth international symposium on magnesium held in Blacksburg, VA. Dr. Seelig served as a major, continuing consultant for The National Institutes of Health, The National Science Foundation, The Department of Health and Welfare of Canada, as well as The Ministry of Health in Switzerland.

In addition to these various activities, Mildred Seelig was the founder and editor-in-chief of the *Journal of the American College of Nutrition*, 1982-1994. She was an Associate Editor of the *Magnesium Bulletin* and *The Journal of Nutritional Medicine*. Dr. Seelig was often sought for guest lectureships in numerous universities and medical centers, both here and abroad. She was the first Master of the American College of Nutrition, an honor few have attained. Mildred was the author and co-author

of approximately 100 original papers and numerous book chapters. Radio commentators sought her out to be an expert-guest on their programs.

Among many goals she set for herself, two always stood out: 1) to write the definitive text-compendium on magnesium in health and disease for the physician-scientist, and 2) to write a book to educate the layman on the role and importance of magnesium in health and disease. Both of these goals were met. After spending 10 years, she published "Magnesium in the Pathogenesis of Disease: Early Roots of Cardiovascular, Skeletal and Renal Abnormalities" in 1980 (2). Very recently, working with Dr. Andrea Rosanoff as co-author, she published the layman's text, "The Magnesium Factor" (3). Even today, the text she published in 1980 still remains the source for researchers in the field, widely utilized and consulted. As to the "Magnesium Factor", hope remains that it will fulfill Dr. Seelig's dream to become the incentive for better health worldwide.

Mildred received her bachelors' degree from the then prestigious Hunter College in 1942 and at her husband's urging, Dr. Alexander Seelig, D.D.S. ( Professor of Dentistry), she went to Medical school and received her M.D. degree in

1945 from New York Medical College. She went on to receive a Masters degree of Public Health from the Columbia University School of Public Health in 1950. Mildred served as an Adjunct Assistant through Associate Professor of Pharmacology at New York Medical College from 1964-1977 and Attending Physician at Goldwater Memorial Hospital, New York University Medical Center, 1973-1983. Mildred also served as Adjunct Professor of Nutrition at the University of North Carolina at Chapel Hill from 1985 until her death this year. Several awards have been established in her honor: The Mildred Seelig Award Lectureship of The Great Lakes Medical Association (2000); the Seelig Award for Lifetime of Meritorious Research on Magnesium (Gordon Research Conf. on Magnesium in Health and Biochemical Processes) (2005). In 2000, she helped to establish The Seelig Award Lectureship of the American College of Nutrition, named originally in honor of her husband, but now to be called the Alexander and Mildred Seelig Award Lectureship.

The most recent endeavor she took on was an invited lectureship tour to India, Taiwan and Thailand, in November 2004. She leaves behind a daughter, Dr. Beth Seelig, and a son, Dr. Charles Seelig, and six loving grandchildren.

## LITERATURE BRIEFS

### **Lutein and AMD**

Lutein consumption is inversely related to eye diseases such as age-related macular degeneration (AMD) and cataracts. Supplementation results in improved vision in patients with AMD and other ocular diseases and increases lutein in macular pigment. It may also serve to protect skin from UV-induced damage and may help reduce the risk of cardiovascular disease. Crystalline lutein is readily absorbed from foods and from dietary supplements, whereas, to enter the bloodstream, lutein esters require de-esterification by intestinal enzymes.

Alves-Rodrigues A, Shao A. The science behind lutein. *Toxicol Lett.* 2004; 15:150:57–83.

### **Grape seed extract and weight**

Energy intake was reduced by 4% after grape-seed extract supplementation 30–60 min prior to each meal, as compared to placebo treatment, in a sub-population of subjects with higher than average energy requirements. These findings suggest that grape seed could be effective in reducing 24 hour energy intake in normal to overweight dietary unrestrained subjects, and could play a significant role in body-weight management.

Vogels N, Nijs IM, Westerterp-Plantenga MS. The effect of grape-seed extract on 24 h energy intake in humans. *Eur J Clin Nutr* 2004; 58:667–73.

### **Probiotics and irritable bowel disease**

Probiotics are live microbial food supplements that confer a health benefit beyond inherent basic nutrition. Multiple potential beneficial effects have been attributed to the probiotic use of lactic acid bacteria, bifidobacteria and other non-pathogenic commensals. An improved understanding of the normal commensal flora

and host-flora interactions has the potential to open up new therapeutic strategies for inflammatory disorders of the gut.

Shanahan F. Probiotics in inflammatory bowel disease—therapeutic rationale and role. *Adv Drug Deliv Rev* 2004; 56:809–18.

### **B-vitamins and orofacial clefts**

Periconceptional folic acid supplementation has been suggested to prevent orofacial clefts (OFCs). Two hundred six mothers of a child with OFC and 203 control mothers filled out a general questionnaire and a food frequency questionnaire around 14 months postpartum as an indication of periconceptional nutrient intake. The periconceptional intake of thiamine, niacin and pyridoxine was significantly lower in mothers of an OFC child. Risk reduction was only demonstrated in women using folic acid supplements but supplement users tended to consume a diet richer in B vitamins. Periconceptional intake of thiamine, niacin and pyridoxine, in addition to folic acid, may contribute to the prevention of OFC.

Krapels IP, van Rooij IA, Ocke MC, van Cleef BA, Kuijpers-Jagtman AM, Steegers-Theunissen RP. Maternal dietary B vitamin intake, other than folate, and the association with orofacial cleft in the offspring. *Eur J Nutr* 2004; 43:7–14.

### **Food supplements for children in India**

According to the National Nutrition Monitoring Bureau of India, over 50% of apparently healthy looking children in India have subclinical or biochemical deficiencies of vitamin A, vitamins B2, B6, folate and vitamin C and over 67% have clinical evidences of iron deficiency. The full genetic potential for physical growth and mental development may be compromised

due to these subclinical deficiencies. Children with subclinical deficiency of micronutrients are also more vulnerable to more frequent and more severe infections due to an under-functioning immune system. All efforts should be made to provide preschool children a balanced and nutritious home-based diet. However, it has been shown that it is not possible to meet 100% recommended dietary allowances of micronutrients from dietary sources alone. Most preschool children need nutritional supplements to optimize their genetic potential.

Singh M. Role of micronutrients for physical growth and mental development. *Indian J Pediatr* 2004; 71:59–62.

#### **Omega-3 fatty acids and chronic inflammatory disease**

Eating fish or taking omega-3 fatty acid supplements can decrease the risk and severity of cardiovascular disease. Such supplements also provide symptomatic relief for rheumatoid arthritis patients. Recent research suggests that asthma, another highly prevalent, chronic inflammatory disease, may also respond to fish oil supplements.

Stephensen CB. Fish oil and inflammatory disease: is asthma the next target for n-3 fatty acid supplements? *Nutr Rev* 2004; 62:486–9.

#### **Calcium supplements for female collegiate athletes and BMD**

Calcium intake in adolescent and young adult female athletes is often inadequate to optimize peak bone mass, an important determinant of future osteoporosis risk. A calcium citrate supplement with vitamin D (providing 1000 mg calcium and 400 IU vitamin D) increased total calcium intake from about 750 mg/day to about 1400 mg/day based on 70% adherence to protocol. An overall non-significant 0.8% average increase in BMD was found. Only basketball players had a significant (1.5%) BMD increase.

Mehlenbeck RS, Ward KD, Klesges RC, Vukadinovich CM. A pilot intervention to increase calcium intake in female collegiate athletes. *Int J Sport Nutr Exerc Metab.* 2004; 14:18–29.

#### **Food supplements and psychiatric disorders**

Several studies have demonstrated that psychiatric symptoms such as depression, mood swings, and aggression may be ameliorated by supplementation with broad-based nutrient formulas containing vitamins, minerals, and essential fatty acids. These findings have been reported in young criminal offenders as well as in adults with mood disturbance and other psychiatric disorders. An open-label trial of nutrient supplements for 9 children revealed decreases in various measures of psychiatric symptoms, suggesting that formal clinical trials of broad nutritional supplementation are warranted.

Kaplan BJ, Fisher JE, Crawford SG, Field CJ, Kolb B. Improved mood and behavior during treatment with a mineral-vitamin supplement: an open-label case series of children. *J Child Adolesc Psychopharmacol* 2004; 14:115–22.

#### **Minerals, vitamins K1 and D and vascular elasticity**

Measurement of several vessel blood wall characteristics in 108 postmenopausal women following supplementation containing minerals and either vitamin D or vitamin K1 showed that the elastic properties of the common carotid artery in the group given minerals plus vitamin K1 improved in comparison to placebo and the group given minerals plus vitamin D.

Braam LA, Hoeks AP, Brouns F, Hamulyak K, Gerichhausen MJ, Vermeer C. Beneficial effects of vitamins D and K on the elastic properties of the vessel wall in postmenopausal women: a follow-up study. *Thromb Haemost* 2004; 91:373–80.

## BOOK REVIEWS

**A Natural Guide to Pregnancy and Postpartum Health** Dean Raffelock, D.C. and Robert Roundtree, M.D. Avery, a member of Putnam Penguin, Inc, New York, 2002. Paperback, 294 page, \$15.95. ISBN 1-58333-138-7

With the premise that a newborn baby's whole body is derived from its mother, the authors outline a "pregnancy recovery program" to restore the resources that flowed from mother to baby in the nine months of gestation. Ideally the program begins even before pregnancy, building reserves and forming habits that will contribute to a healthy pregnancy and avoid postpartum nutrient depletions. The program aims to avoid the postpartum depression that is common in today's society, and in addition to produce healthier and happier babies that are easier for a new mother to care for.

Information is presented at many levels of sophistication. Simple "food choice" advice is given, along with hints on breastfeeding, how to exercise effectively and sleep better. There is also instruction on the appropriate use of food supplements, achieving hormone balance, restoring digestive function, and even on choosing and interpreting laboratory tests.

Sprinkled in with the cerebral guidance, you will find poignant stories of the way real people have benefited from the insight of these two practitioners. The result is a book that promises to ease the passage of mothers-to-be through one of the most intense experiences a woman can have.

James Heffley, Ph.D., CCN, DANLA  
Austin, TX

**Brain-Building Nutrition: The Healing Power of Fats and Oils, 2<sup>nd</sup> Ed** Michael A. Schmidt. Frog, Ltd Berkeley, CA. 2001. Paperback, 286 pages, \$16.95. ISBN 1-58394-048-0

Perhaps the most significant insight in the still-evolving field of nutrition over the last few years is the realization that the type of fat in our diets is at least as important as the amount of fat. For decades all dietary fat, irrespective of quality, was condemned as being unhealthy. Dr. Schmidt has brought together the enormous amount of information that has recently arisen relating, in particular, the omega-3 family of fatty acids to all aspects of human health and well being.

The profound influence of omega-3 fatty acids is nowhere more significant than in their effect on the developing brains of infants. Beginning even before birth, and continuing for years after birth, the brain needs omega-3 fatty acids to build structure that will last a lifetime. Dr. Schmidt devotes an entire chapter to the "window of time" in which the reduced availability of omega-3 fatty acids has its most ominous consequences on health. This it not to say that omega-3 fatty acids are not important at every stage of life, just that deficiency at certain stages results in apparently irreversible damage.

Dr. Schmidt is an experienced author and teacher with an easy-to-comprehend writing style. His previous books have been well received by both traditional and avant-garde nutritionists, no doubt due in no small part to his impeccable documentation of his ideas. Combining a sterling intellect with a genuine talent for expressing in writing ideas that are sometimes difficult to understand makes for an outstanding book.

James Heffley, Ph.D., CCN, DANLA  
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## BOOKS RECEIVED FOR REVIEW

**Handbook of Nutrition and Immunity** M. Eric Gershwin, M.D., Penelope Nestel, Ph.D. and Carl L. Keen, Ph.D., Eds. Humana Press, Totawa, NJ, 2004. Hardback, 365 pages, \$89.50. ISBN 1-58829-308-4

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