The Honorable Dan Burton, Chairman
House Government Reform and Oversight Committee
C/o Milt Copulos/ Beth Clay
Room 2157 RHOB
Washington, DC 20515

Dear Congressman Burton,

I am sending you a form letter printed by The Life Extension Foundation as it expresses clearly what I would like to say. I am personally sick and tired of the FDA attempting to make end runs around the expressed will of the citizens of this country. The Nutritional Supplement Health and Education Act was passed by a huge majority in Congress and yet the FDA continues to try to use every extra legal means possible to nullify the will of the people and impose draconian limits on food supplements.

I am asking you to rein in this organization, if necessary, by applying criminal sanctions against any individual in the FDA who attempts extra legal or illegal means to subvert and ignore the law. Let the people have a measure of autonomy in providing for their health.

I am requesting that you inform me of your actions regarding the oversight hearing.

Thanking you for your attention and service,

I am,

Very Truly Yours

Mr. Paris E. Asta

Paris E. Asta
Dear Congressman Burton:

Prior to last September’s meeting of the Codex Committee on Nutrition and Food for Special Dietary Use, you and four other members of Congress strongly requested in writing that the FDA’s Dr. Yetley remove the second paragraph from the U.S. codex comments on agenda item #5 (vitamins and minerals), because it contradicted the first paragraph, and lent credence to the unscientific notion that “maximum upper potency limits” should be put on vitamins and minerals. Dr. Yetley not only ignored your written request, but John Hammell caught her doing so on videotape which has been put on the Life Extension Foundation’s website in the political section, along with footage of John being forced to stop taping by the German Codex Chairman (http://www.lef.org). A complete account of what happened is available at http://www.iahf.com under “breaking news.”

From a standpoint of safety, there is no justification for attempting to apply a “Risk Assessment” document which was designed for evaluating toxic pharmaceutical drugs, to dietary supplements, which have been well established through the National Association of Poison Control Centers, and numerous other sources to be extraordinarily safe, even when consumed in doses much higher than the RDA. Orthomolecular physicians such as Bonnie Camo, M.D. have seen doses as high as 3 grams per day of niacin used in complete safety, while the National Academy of Sciences and FDA are advocating a maximum upper potency limit of just 35 mg, just because a few highly sensitive individuals experience a tingling sensation known as the “niacin flush” when taking niacin in low doses. There is nothing unsafe about the niacin flush, which actually helps circulation and is considered pleasurable by some.

It is obvious to consumers around the world that the FDA is attempting to use the highly unscientific, and heavily prejudiced National Academy of Sciences document titled “A Risk Assessment Model for Establishing Upper Limits for Nutrients” as a means of moving beyond the consumer generated impasse at the Codex Committee on Nutrition and Foods for Special Dietary Use. The FDA has announced its intention to harmonize its regulations to emerging Codex standards in an Advance Notice of Proposed Rulemaking that was published in the Federal Register on July 7, 1997, vol. 62, #129 pp.36243-36248. You can view this at http://iahf.com/codx-fda.txt.

I urge you to call John Hammell, Bonnie Camo M.D., and other witnesses to a Hearing before your Committee, and I urge you to force the FDA to withdraw the second paragraph of its comments along with the NAS Risk Assessment document in keeping with current US Law. Congress has spoken clearly on this with the passage of DSHEA, and most recently again in October of 1997 when dietary supplements were specifically exempted from the harmonization language in the FDA Reform Bill.

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