CHRIS CANNON BRD DISTRICT, UTAH

COMMITTEE ON THE JUDICIARY Chairman, Subcommittee on Commercial and Administrative Law

COMMITTEE ON RESOURCES

COMMITTEE ON GOVERNMENT REFORM



Congress of the United States

House of Representatives Washington, DC 20515

August 11, 2003

WASHINGTON OFFICE. 118 Cannon House Office Building Washington, DC 20515 (202) 225-7751 • Fax: (202) 225-5629

UTAH OFFICES:

51 SOUTH UNIVERSITY Ανεινις, SUITE 319 Ρκάνο, UT 84806 (801) 379-2500 ΓΑΧ: (801) 379-2509

3600 South Constitution Bollevard West Valley City, UT 34115 (301) 955–3631 Fax: (801) 955–3632

www.house.gov/cannon Email: cannon.u03@mail.house.gov

The Honorable Mark B. McClellan, Commissioner Food and Drug Administration Dockets Management Branch (HFA-305) Food and Drug Administration 5630 Fishers Lane, Room 1061 Rockville, MD 20852

Docket No. 1996N-0417 Good Manufacturing Practices for Dietary Supplements

Dear Commissioner McClellan:

المامية والمستشرقين سنادات المتجاد بروتهم والرزا محدة مترسيا مامه

I wish to comment on proposed regulations establishing Good Manufacturing Practices (GMPs) for dietary supplements. These are very important to me because of the size of the dietary supplement industry in Utah. Any proposed regulations must take into account the realities of dietary supplement production, and in that connection, the expertise of Utah industry representatives is important.

The Utah Natural Products Alliance has submitted comments on the proposed regulation, and in general I wish to associate myself with those comments. They provide extensive information that the Agency can use in evaluating and modifying its proposals. In addition to those comments, I would like to address a couple of points as a Member of Congress.

The first is the legal basis for the regulations. While I was not a member of Congress in 1994, when the Dietary Supplement Health and Education Act was originally passed, I have since examined it carefully, particularly in light of my membership on the Government Reform and Oversight Committee, and as current Vice Chair of the Wellness and Human Rights Subcommittee on which I serve because of its importance to my congressional district. I am also aware of proposals for much stricter regulation of dietary supplements, and the view that the proposed regulations are a first step toward the eventual regulation of supplements as prescription drugs.

The Agency, of course, has jurisdiction over both food and drugs. It is important to keep that distinction in mind, and to avoid applying the stricter regulation entailed in drug marketing to products that are not drugs. The application of protocols appropriate to

96N-0417

PRINTED ON RECYCLED PAPER

.

C 195

drugs is not appropriate for such products as dietary supplements. Despite the attention paid to the effects of a few supplements, the safety record of natural supplementary products is exemplary, and the Agency should avoid overreaction.

At some point Congress may change its instructions to the FDA with respect to food additives, and I am sure you are aware that legislation has been introduced to do that. Until that legislation is acted on, the Agency should avoid using expansive definitions of such words as "model" to bootstrap itself into new realms of administrative and regulatory authority.

The second point I would like to make is with the lack of economic analysis provided to the public along with the proposed regulations.

To put it mildly, the economic effect of the regulations suggested in the proposals is orders of magnitude out of line with reality. UNPA has provided the Agency with several case studies refuting the assumptions that are the basis of the Agency's analysis. I strongly urge you to accept the expertise of the industry in re-evaluating the economic assumptions.

More importantly, and in my capacity as an overseer of the regulatory process, I am concerned that the proposals were rushed into print and a comment period established without the publication of some of the core economic analysis that undergirds them.

How did this happen? How common is such a divorce of proposals from the economic analysis that informs them at the FDA? I would appreciate answers to these questions and at your earliest convenience. These questions are outside the regulatory comment process.

I understand that an additional 29 days have been allocated for comments on the economic data from the Agency, at such time as those become available, and I reserve the right to comment additionally on them during that period. It is possible that the economic analysis will give rise to additional comments on other areas of the proposals, and I strongly urge the Agency to accept whatever comments it receives during that extension period.

Finally, I would like to comment on the use of the regulatory process as an anticompetitive tool.

It is no secret that industries and companies regulated by the Food and Drug Administration have very different attitudes about the legitimacy of competition from other companies and sectors of the industry. In developing regulations, it is important that the Agency not allow rent seekers from one side or another to use the power of the Agency to raise barriers to entry or expansion against new products, companies, and processes. The danger to the Agency of assuming the desirability of the status quo is high. Companies now in dominance will urge its maintenance. The Agency should use special care to avoid limiting the introduction, marketing, use and expansion of new

2004/004

products and services. Obviously, public safety is critical, but you and your fellow commissioners are dealing with a vibrant, dynamic industry, one that constantly challenges accepted ways of doing business and accepted ideas about obtaining and maintaining health. This has proven enormously beneficial to a generation of Americans who are healthier and fitter than any in the past.

The citizens of the United States owe an enormous debt to the medical and pharmaceutical professions. At the same time, those professions are subject to the same incentives as other actors in our economy, and the Agency has to insist that the marketplace be open to competing theories and practices. In the final analysis, the consumer is the supreme regulator, and the Agency must be careful not to restrict the consumer's access and choices for any reasons short of compelling issues of safety.

I look forward to your response to my questions, and to an opportunity to examine the economic data that will be forthcoming shortly.

Sincerely,

.

Chris Cannon Member of Congress

Facsimile CONGRESSMAN CHRIS CANNON



Utah's 3¹⁴ District

CONGRESS OF THE UNITED STATES OF AMERICA 51 S. University Avenue Suite #319 Provo, Utah 84606 (801) 379-2500 Fax (801) 379-2509 To: Commissioner McClellan, FDA From: Megan Faulkner Subject: Comments on Docket #1996N-0417 Hof Pages: 4

Messages:

Attached is a letter from Congressman Chris Cannon commenting on the Good Manufacturing Practices for dietary supplements.

Thank you,

This facsimile is from the Office of Congressman Chris Cannon and contains information that may be confidential or privileged. It is intended only for the individual or entity named above; for anyone else to disclose, copy, distribute, or use the contents of this message is prohibited. All personal messages, which are not to be attributed to Congressman Cannon, express the views of the sender, solely. Such messages may not be copied or distributed without this disclaimer. If you received this message in error, please notify us immediately via email at <u>cannon.ut03@mail.house.gov</u> or by phone (801) 379-2500. Thank you.