



DEPARTMENT OF HEALTH & HUMAN SERVICES

Reyo

*Archives
cc: Sopko
DMelson*

Food and Drug Administration
Rockville MD 20857

*PJung
KChapman*

• The Honorable John D. Dingell
Chairman
Committee on Energy and Commerce
House of Representatives
Washington, D.C. 20515-2215

JAN 25 2008

Dear Mr. Chairman:

Thank you for your letter of August 13, 2007, co-signed by Chairman Bart Stupak, Subcommittee on Oversight and Investigations, Committee on Energy and Commerce, regarding the circumstances surrounding a Food and Drug Administration (FDA) Warning Letter to Cordis Corporation concerning their CYPHER™ Sirolimus-eluting Coronary Stent in April 2004. We are enclosing a third partial response to your request that includes ten boxes of requested documents. The first two partial responses were sent on September 25, 2007 and November 27, 2007. Additional material is being compiled and will be sent as soon as possible.

Information in the enclosures includes information that is trade secret, commercial confidential or other information protected from disclosure to the public under the Freedom of Information Act (Title 5, *United States Code [U.S.C.] 552*), the Trade Secrets Act (18 U.S.C. 1905), and FDA regulations. The Committee should not publish or otherwise make public any such information. We would be glad to discuss with the Committee staff the protected status of any specific information.

Thank you again for your interest in this matter. If we can be of further assistance, please let us know. A similar letter has been sent to Chairman Stupak, without enclosures.

Sincerely,

Stephen R. Mason
Acting Assistant Commissioner
for Legislation

Enclosures