^{109TH CONGRESS} 2D SESSION H.R.4718

To require prescription drug manufacturers, packers, and distributors to disclose certain gifts provided in connection with detailing, promotional, or other marketing activities, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

FEBRUARY 8, 2006

Mr. DEFAZIO introduced the following bill; which was referred to the Committee on Energy and Commerce

A BILL

- To require prescription drug manufacturers, packers, and distributors to disclose certain gifts provided in connection with detailing, promotional, or other marketing activities, and for other purposes.
 - 1 Be it enacted by the Senate and House of Representa-
 - 2 tives of the United States of America in Congress assembled,

3 SECTION 1. SHORT TITLE.

- 4 This Act may be cited as the "Drug Company Gift
- 5 Disclosure Act".

SEC. 2. DISCLOSURE BY PRESCRIPTION DRUG MANUFAC TURERS, PACKERS, AND DISTRIBUTORS OF CERTAIN GIFTS.

4 Section 503 of the Federal Food, Drug, and Cos5 metics Act (21 U.S.C. 353) is amended by adding at the
6 end the following:

7 "(h)(1) Each manufacturer, packer, or distributor of
8 a drug subject to subsection (b)(1) shall disclose to the
9 Commissioner—

"(A) not later than June 30, 2007, and each
June 30 thereafter, the value, nature, and purpose
of any—

"(i) gift provided during the preceding calendar year to any covered health entity by the
manufacturer, packer, or distributor, or a representative thereof, in connection with detailing,
promotional, or other marketing activities; and

"(ii) cash rebate, discount, or any other financial consideration provided during the preceding calendar year to any pharmaceutical
benefit manager by the manufacturer, packer,
or distributor, or a representative thereof, in
connection with detailing, promotional, or other
marketing activities; and

25 "(B) not later than the date that is 6 months
26 after the date of enactment of this subsection and
•HR 4718 IH

each June 30 thereafter, the name and address of
 the individual responsible for the compliance of the
 manufacturer, packer, or distributor with the provi sions of this subsection.

5 "(2) Subject to paragraph (3), the Commissioner
6 shall make all information disclosed to the Commissioner
7 under paragraph (1) publicly available, including by post8 ing such information on the Internet.

9 "(3) The Commissioner shall keep confidential any 10 information disclosed to or otherwise obtained by the Commissioner under this subsection that relates to a trade se-11 12 cret referred to in section 1905 of title 18, United States 13 Code. The Commissioner shall provide an opportunity in the disclosure form required under paragraph (4) for a 14 15 manufacturer, packer, or distributor to identify any such 16 information.

17 "(4) Each disclosure under this subsection shall be18 made in such form and manner as the Commissioner may19 require.

"(5) Each manufacturer, packer, and distributor described in paragraph (1) shall be subject to a civil monetary penalty of not more than \$10,000 for each violation
of this subsection. Each unlawful failure to disclose shall
constitute a separate violation. The provisions of paragraphs (3), (4), and (5) of section 303(f) shall apply to

such a violation in the same manner as such provisions
 apply to a violation of a requirement of this Act that re lates to devices.

4 "(6) For purposes of this subsection:

5 "(A) The term 'covered health entity' includes
6 any physician, hospital, nursing home, pharmacist,
7 health benefit plan administrator, or any other per8 son authorized to prescribe or dispense drugs that
9 are subject to subsection (b)(1), in the District of
10 Columbia or any State, commonwealth, possession,
11 or territory of the United States.

"(B) The term 'gift' includes any gift, fee, payment, subsidy, or other economic benefit with a
value of \$50 or more, except that such term excludes
the following:

16 "(i) Free samples of drugs subject to sub17 section (b)(1) intended to be distributed to pa18 tients.

"(ii) The payment of reasonable compensation and reimbursement of expenses in connection with any bona fide clinical trial conducted
in connection with a research study designed to
answer specific questions about drugs, devices,
new therapies, or new ways of using known
treatments.

"(iii) Any scholarship or other support for medical students, residents, or fellows selected by a national, regional, or specialty medical or other professional association to attend a significant educational, scientific, or policy-making conference of the association.".

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