

1 **SEC. ____ . COMMUNICATIONS REGARDING INTENDED USES**
2 **OF DRUGS AND DEVICES; SCIENTIFIC EX-**
3 **CHANGE.**

4 The Federal Food, Drug, and Cosmetic Act is amend-
5 ed by inserting after section 201 of such Act (21 U.S.C.
6 321) the following:

7 **“SEC. 201A. INTENDED USES OF DRUGS AND DEVICES.**

8 “(a) INTENDED USE.—For purposes of this Act, in-
9 cluding sections 301(d), 502(f)(1), 505, 510, 513, and
10 515, and for purposes of section 351 of the Public Health
11 Service Act, the intended use of a drug or device shall
12 not be determined by reference to—

13 “(1) actual or constructive knowledge of the
14 manufacturer or sponsor that such drug or device
15 will be used in a manner that varies from the use
16 approved, cleared, or licensed for marketing under
17 section 505, 510, 513, or 515 of this Act or section
18 351 of the Public Health Service Act;

19 “(2) non-public statements about the drug or
20 device that are not reflected in any claim, pro-
21 motional statement or material, or circumstances
22 surrounding the distribution of the drug or device
23 that involve interactions with third parties; or

1 “(3) communications meeting the criteria under
2 subsection (b) to be considered scientific exchange
3 safe harbor communications.

4 “(b) SCIENTIFIC EXCHANGE SAFE HARBOR.—

5 “(1) IN GENERAL.—A communication by a
6 manufacturer or sponsor, or a person acting on be-
7 half of a manufacturer or sponsor, about informa-
8 tion that is not included in the drug or device label-
9 ing required by this Act, constitutes a scientific ex-
10 change safe harbor communication if—

11 “(A) the communication is not advertising
12 or otherwise promotional in nature;

13 “(B) the communication is supported by
14 competent and reliable scientific evidence;

15 “(C) the communication clearly discloses
16 appropriate contextual information about the
17 data presented, including information about
18 limitations of the data, the scientific and ana-
19 lytical methodologies used, and any contradic-
20 tory data or information known to the manufac-
21 turer or sponsor;

22 “(D) the communication includes a con-
23 spicuous and prominent statement about such
24 information not being contained in the drug or
25 device labeling required by this Act; and

1 “(E) if the communication relates to a use
2 of a drug or device that has not been approved
3 or cleared for marketing under section 505,
4 510, 513, or 515 of this Act or section 351 of
5 the Public Health Service Act, the manufac-
6 turer or sponsor, or person acting on behalf of
7 the manufacturer or sponsor, makes no rep-
8 resentation that such use has been dem-
9 onstrated to be safe or effective.

10 “(2) RULE OF CONSTRUCTION.—Nothing in
11 this subsection shall be construed—

12 “(A) to authorize the Secretary to require
13 that a manufacturer or sponsor submit an ap-
14 plication, certification, or other such submis-
15 sion, or to seek the Secretary’s review or ap-
16 proval, before, during, or subsequent to engag-
17 ing in scientific exchange; or

18 “(B) to limit the ability of a manufacturer
19 or sponsor to engage in communications or ac-
20 tivities not specified in this subsection, but that
21 are otherwise permissible.”.