

117TH CONGRESS
2D SESSION

S. _____

To require summary approval information with respect to certain approved drugs and biological products.

IN THE SENATE OF THE UNITED STATES

Mr. HICKENLOOPER introduced the following bill; which was read twice and referred to the Committee on _____

A BILL

To require summary approval information with respect to certain approved drugs and biological products.

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 “Accelerated Approval
5 Transparency Act”.

6 **SEC. 2. SUMMARY APPROVAL INFORMATION.**

7 With respect to each new drug application for a new
8 molecular entity approved under section 505(c) of the
9 Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355(c))
10 or biological product licensed under section 351(a) of the

1 Public Health Service Act (42 U.S.C. 262(a)) pursuant
2 to accelerated approval under section 506(c) of the Fed-
3 eral Food, Drug, and Cosmetic Act (21 U.S.C. 356(c)),
4 the Secretary of Health and Human Services shall provide
5 for the drug or biologic action package a summary of the
6 basis for approval, including, as relates to such new molec-
7 ular entity, whether an advisory committee meeting was
8 held and a rationale for a determination by the Secretary
9 that a surrogate endpoint is reasonably likely to predict
10 clinical benefit.