IN THE HOUSE OF REPRESENTATIVES

Mr. TONKO introduced the following bill; which was referred to the Committee on

A BILL

To amend title XVIII of the Social Security Act to require the Secretary of Health and Human Services to add a new set of measures to the 5-star rating system under the Medicare Advantage program in order to encourage increased access to biosimilar biological products.

Be it enacted by the Senate and House of Representatives of the United States of America in Congress assembled,

SECTION 1. SHORT TITLE.

This Act may be cited as the “Star Rating for Biosimilars Act”.
SEC. 2. ADDITION OF NEW MEASURES BASED ON ACCESS TO BIOSIMILAR BIOLOGICAL PRODUCTS TO THE 5-STAR RATING SYSTEM UNDER MEDICARE ADVANTAGE.

(a) In General.—Section 1853(o)(4) of the Social Security Act (42 U.S.C. 1395w–23(o)(4)) is amended by adding at the end the following new subparagraph:

“(E) ADDITION OF NEW MEASURES BASED ON ACCESS TO BIOSIMILAR BIOLOGICAL PRODUCTS.—

“(i) IN GENERAL.—For 2022 and subsequent years, the Secretary shall add a new set of measures to the 5-star rating system based on access to biosimilar biological products covered under part B and, in the case of MA–PD plans, such products that are covered part D drugs. Such measures shall assess the impact a plan’s benefit structure may have on enrollees’ utilization of or ability to access biosimilar biological products, including in comparison to the reference biological product, and shall include measures, as applicable, with respect to the following:

“(I) COVERAGE.—Assessing whether a biosimilar biological prod-
uct is on the plan formulary in lieu of
or in addition to the reference biologi-
cal product.

“(II) PREFERENCING.—Assess-
ing tier placement or cost sharing for
a biosimilar biological product relative
to the reference biological product.

“(III) UTILIZATION MANAGEMENT TOOLS.—Assessing whether and
how utilization management tools are
used with respect to a biosimilar bio-
logical product relative to the ref-
erece biological product.

“(IV) UTILIZATION.—Assessing
the percentage of enrollees prescribed
the biosimilar biological product when
the reference biological product is also
available.

“(ii) DEFINITIONS.—In this subpara-
graph, the terms ‘biosimilar biological
product’ and ‘reference biological product’
have the meaning given those terms in sec-
tion 1847A(c)(6).

“(iii) PROTECTING PATIENT INTER-
ESTS.—In developing such measures, the
Secretary shall ensure that each measure developed to address coverage, preferencing, or utilization management is constructed such that patients retain equal access to appropriate therapeutic options without undue administrative burden.”.

(b) Clarification Regarding Application to Prescription Drug Plans.—To the extent the Secretary of Health and Human Services applies the 5-star rating system under section 1853(o)(4) of the Social Security Act (42 U.S.C. 1395w–23(o)(4)), or a similar system, to prescription drug plans under part D of title XVIII of such Act, the provisions of subparagraph (E) of such section, as added by subsection (a) of this section, shall apply under the system with respect to such plans in the same manner as such provisions apply to the 5-star rating system under such section 1853(o)(4).