

AMENDMENT TO
RULES COMMITTEE PRINT 116-41
OFFERED BY MR. CÁRDENAS OF CALIFORNIA

Add at the end of title VIII the following new section (and amend the table of contents accordingly):

1 **SECTION 812. DEMONSTRATION PROJECT TO INCREASE**
2 **ACCESS TO BIOSIMILAR BIOLOGICAL PROD-**
3 **UCTS UNDER THE MEDICARE PROGRAM.**

4 (a) ESTABLISHMENT.—Beginning not later than 1
5 year after the date of the enactment of this Act, the Sec-
6 retary of Health and Human Services shall establish and
7 implement a 3-year nationwide demonstration project
8 under part B of title XVIII of the Social Security Act to
9 evaluate the benefits of providing a shared savings pay-
10 ment for biosimilar biological products furnished under
11 such part.

12 (b) PARTICIPATION.—

13 (1) IN GENERAL.—Participation under the
14 demonstration project shall be voluntary, and a par-
15 ticipating provider may terminate participation at
16 any time and the Secretary may terminate the par-
17 ticipation of such a provider at any time.

1 (2) APPLICATION AND SELECTION.—To partici-
2 pate under the demonstration project, an eligible
3 provider shall submit to the Secretary an application
4 in such form and manner and containing such infor-
5 mation as specified by the Secretary. Each eligible
6 provider who submits such an application shall be
7 selected by the Secretary for participation under the
8 demonstration project.

9 (c) COVERAGE.—Except as otherwise provided in this
10 section, payment may be made under the demonstration
11 project for a biosimilar biological product only if such prod-
12 uct is covered under part B of title XVIII of the Social
13 Security Act and such payment shall be made in the same
14 manner as payment is provided for such a product under
15 such part.

16 (d) ADDITIONAL PAYMENT.—Under the demonstra-
17 tion project, in addition to the payment that would other-
18 wise be made under part B of title XVIII of the Social
19 Security Act for a biosimilar biological product furnished
20 or dispensed by a participating provider to a Medicare
21 beneficiary, there shall be made an additional payment,
22 in an amount determined by the Secretary, that reflects
23 a portion of any difference between the that may result
24 from furnishing the biosimilar biological product, as com-
25 pared to the reference biological product.

1 (e) WAIVER AUTHORITY.—The Secretary may waive
2 such requirements of title XVIII of the Social Security Act
3 as may be necessary to carry out the demonstration
4 project.

5 (f) REPORTS.—

6 (1) INTERIM EVALUATION AND REPORT.—Not
7 later than 3 years after the date of enactment of
8 this Act, the Secretary shall submit to Congress a
9 report that contains an analysis of the appropriate-
10 ness of expanding or extending the demonstration
11 project and, to the extent such analysis determines
12 such an expansion or extension appropriate, rec-
13 ommendations for such expansion or extension, re-
14 spectively.

15 (2) FINAL EVALUATION AND REPORT.—Not
16 later than one year after the date of completion of
17 the demonstration project, the Secretary shall sub-
18 mit to Congress a report that contains a final anal-
19 ysis of the project and recommendations described in
20 paragraph (1).

21 (g) DEFINITIONS.—In this section:

22 (1) DEMONSTRATION PROJECT.—The term
23 “demonstration project” means the demonstration
24 project conducted under this Act.

1 (2) BIOSIMILAR BIOLOGICAL PRODUCT.—The
2 term “biosimilar biological product” means a biologi-
3 cal product approved under an abbreviated applica-
4 tion for a license of a biological product that relies
5 in part on data or information in an application for
6 another biological product licensed under section 351
7 of the Public Health Service Act (42 U.S.C. 262) .

8 (3) ELIGIBLE PROVIDER.—The term “eligible
9 provider” means a provider of services or supplier
10 that is eligible to receive payment under part B of
11 title XVIII of the Social Security Act for furnishing
12 or dispensing biosimilar biological products.

13 (4) MEDICARE BENEFICIARY.—The term
14 “Medicare beneficiary” means an individual who is
15 enrolled for benefits under part B of title XVIII of
16 the Social Security Act.

17 (5) PARTICIPATING PROVIDER.—The term
18 “participating provider” means an eligible provider
19 that has been selected for participation under the
20 project under subsection (b)(2) and with respect to
21 whom such participation has not been terminated.

22 (6) REFERENCE BIOLOGICAL PRODUCT.—The
23 term “reference biological product” means the bio-
24 logical product licensed under section 351 of the
25 Public Health Service Act (42 U.S.C. 262) that is

- 1 referred to in the application described in paragraph
- 2 (2) of the biosimilar biological product.

