

107TH CONGRESS  
2D SESSION

# H. R. 4014

To amend the Federal Food, Drug, and Cosmetic Act with respect to the development of products for rare diseases.

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## IN THE HOUSE OF REPRESENTATIVES

MARCH 20, 2002

Mr. FOLEY (for himself, Mr. WAXMAN, Mr. SHIMKUS, Mr. BROWN of Ohio, Mrs. ROUKEMA, Mr. RUSH, Mr. KING, Mr. GREENWOOD, and Mr. DINGELL) introduced the following bill; which was referred to the Committee on Energy and Commerce

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## A BILL

To amend the Federal Food, Drug, and Cosmetic Act with respect to the development of products for rare diseases.

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 “Rare Diseases Orphan  
5 Product Development Act of 2002”.

6 **SEC. 2. FINDINGS AND PURPOSES.**

7 (a) FINDINGS.—Congress makes the following find-  
8 ings:

1           (1) Rare diseases and disorders are those which  
2           affect small patient populations, typically popu-  
3           lations smaller than 200,000 individuals in the  
4           United States. Such diseases and conditions include  
5           Huntington’s disease, amyotrophic lateral sclerosis  
6           (Lou Gehrig’s disease), Tourette syndrome, Crohn’s  
7           disease, cystic fibrosis, cystinosis, and Duchenne  
8           muscular dystrophy.

9           (2) For many years, the 25,000,000 Americans  
10          suffering from the over 6,000 rare diseases and dis-  
11          orders were denied access to effective medicines be-  
12          cause prescription drug manufacturers could rarely  
13          make a profit from marketing drugs for such small  
14          groups of patients. The prescription drug industry  
15          did not adequately fund research into such treat-  
16          ments. Despite the urgent health need for these  
17          medicines, they came to be known as “orphan  
18          drugs” because no companies would commercialize  
19          them.

20          (3) During the 1970s, an organization called  
21          the National Organization for Rare Disorders  
22          (NORD) was founded to provide services and to  
23          lobby on behalf of patients with rare diseases and  
24          disorders. NORD was instrumental in pressing Con-

1       gress for legislation to encourage the development of  
2       orphan drugs.

3               (4) The Orphan Drug Act created financial in-  
4       centives for the research and production of such or-  
5       phan drugs. New Federal programs at the National  
6       Institutes of Health and the Food and Drug Admin-  
7       istration encouraged clinical research and commer-  
8       cial product development for products that target  
9       rare diseases. An Orphan Products Board was estab-  
10      lished to promote the development of drugs and de-  
11      vices for rare diseases or disorders.

12              (5) Before 1983, some 38 orphan drugs had  
13      been developed. Since the enactment of the Orphan  
14      Drug Act, more than 220 new orphan drugs have  
15      been approved and marketed in the United States  
16      and more than 800 additional drugs are in the re-  
17      search pipeline.

18              (6) Despite the tremendous success of the Or-  
19      phan Drug Act, rare diseases and disorders deserve  
20      greater emphasis in the national biomedical research  
21      enterprise.

22              (7) The Food and Drug Administration sup-  
23      ports small clinical trials through Orphan Products  
24      Research Grants. Such grants embody successful  
25      partnerships of government and industry, and have

1 led to the development of at least 23 drugs and four  
2 medical devices for rare diseases and disorders. Yet  
3 the appropriations in fiscal year 2001 for such  
4 grants were less than in fiscal year 1995.

5 (b) PURPOSES.—The purpose of this Act is to in-  
6 crease the national investment in the development of  
7 diagnostics and treatments for patients with rare diseases  
8 and disorders.

9 **SEC. 3. FOOD AND DRUG ADMINISTRATION; GRANTS AND**  
10 **CONTRACTS FOR THE DEVELOPMENT OF OR-**  
11 **PHAN DRUGS.**

12 Subsection (c) of section 5 of the Orphan Drug Act  
13 (21 U.S.C. 360ee(c)) is amended to read as follows:

14 “(c) For grants and contracts under subsection (a),  
15 there are authorized to be appropriated such sums as al-  
16 ready have been appropriated for fiscal year 2002, and  
17 \$25,000,000 for each of the fiscal years 2003 through  
18 2006.”.

19 **SEC. 4. TECHNICAL AMENDMENT.**

20 Section 527(a) of the Federal Food, Drug, and Cos-  
21 metic Act (21 U.S.C. 360cc(a)) is amended in the matter  
22 following paragraph (2)—

23 (1) by striking “, of such certification,”; and

1           (2) by striking “, the issuance of the certifi-  
2           cation,”.

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