



1 ducing or distributing counterfeit, adulterated, mis-  
2 branded or unapproved medical products with the intent  
3 to defraud or mislead the consumer.

4 (c) REPORT.—Not later than 1 year after the date  
5 of the enactment of this Act, the Secretary of the Treas-  
6 ury, acting through Director of the Office of Foreign As-  
7 sets Control, and having consulted with any other agencies  
8 as the Director of the Office of Foreign Assets Control  
9 determines appropriate, shall submit to House Commit-  
10 tees on Financial Services, Foreign Affairs, and Energy  
11 and Commerce and the Senate Committees on Foreign Re-  
12 lations, Health, Education Labor and Pensions, and  
13 Banking, Housing, and Urban Affairs, a report that ad-  
14 dresses whether existing sanctions authorities of the Office  
15 of Foreign Assets Control are sufficient to address the  
16 production and distribution of counterfeit, adulterated,  
17 misbranded or unapproved medical products.

