AMENDMENT TO
RULES COMMITTEE PRINT 114-22
OFFERED BY MR. FITZPATRICK OF
PENNSYLVANIA

Page 235, after line 2, insert the following:

Subtitle R—Other Provisions

SEC. 2321. SENSE OF CONGRESS.

It is the sense of the Congress that any medical device that is approved by the Food and Drug Administration should be sufficiently evaluated for risk and patient safety, including any medical device cleared pursuant to section 510(k) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 360(k)).