Subtitle R—Other Provisions

SEC. 2321. GAO REPORT ON FDA’S IMPLEMENTATION OF MEDICAL DEVICE REPORTING REQUIREMENTS.

Not later than 6 months after the date of enactment of this Act, the Comptroller General of the United States shall complete a study and submit a report to the Congress on the Food and Drug Administration’s implementation of the medical device reporting requirements under part 803 of title 21, Code of Federal Regulations, including with respect to problems involving reporting on power morcellators.