

HR 2483

Quality Systems Certification Act of 2017

Congress: 115 (2017–2019, Ended)

Chamber: House

Policy Area: Health

Introduced: May 17, 2017

Current Status: Referred to the Subcommittee on Health.

Latest Action: Referred to the Subcommittee on Health. (May 19, 2017)

Official Text: <https://www.congress.gov/bill/115th-congress/house-bill/2483>

Sponsor

Name: Rep. Hudson, Richard [R-NC-8]

Party: Republican • State: NC • Chamber: House

Cosponsors (1 total)

Cosponsor	Party / State	Role	Date Joined
Rep. Bucshon, Larry [R-IN-8]	R · IN		May 17, 2017

Committee Activity

Committee	Chamber	Activity	Date
Energy and Commerce Committee	House	Referred to	May 19, 2017

Subjects & Policy Tags

Policy Area:

Health

Related Bills

Bill	Relationship	Last Action
115 S 1183	Related bill	May 18, 2017: Read twice and referred to the Committee on Health, Education, Labor, and Pensions.

Quality Systems Certification Act of 2017

This bill amends the Federal Food, Drug, and Cosmetic Act to require the Food and Drug Administration (FDA) to establish a third-party quality system assessment program to accredit persons to assess whether a medical device manufacturer's quality system can ensure the safety and effectiveness or substantial equivalence of an approved medical device after certain changes, including changes in manufacturing or changes to enhance device safety.

Device manufacturers with quality systems that have been certified by an accredited person are allowed to make changes to a device without submitting to the FDA the premarket notification, 30-day notice, or premarket approval supplement that would otherwise be required.

An accredited person who assesses a device manufacturer's quality system must submit a summary of their assessment and, as appropriate, a certification of the quality system to the FDA within 30 days of the assessment. An assessment summary and certification is deemed accepted by the FDA 30 days after submission unless the FDA determines that additional information is needed to support certification, the assessment or certification is unwarranted, or an action other than acceptance of the certification is otherwise justified.

Device manufacturers who make changes to devices without submitting a premarket notification must describe the changes in an annual summary submitted to the FDA. Changes made without submitting a 30-day notice or a premarket approval supplement must be described in a periodic report.

Certifications accepted by the FDA remain in effect for two years.

The FDA must report on this quality system assessment program no later than January 31, 2022. The program is terminated at the end of FY2022.

Actions Timeline

- **May 19, 2017:** Referred to the Subcommittee on Health.
- **May 17, 2017:** Introduced in House
- **May 17, 2017:** Referred to the House Committee on Energy and Commerce.