

## HR 1108

Recall Unsafe Drugs Act of 2017

**Congress:** 115 (2017–2019, Ended)

**Chamber:** House

**Policy Area:** Health

**Introduced:** Feb 16, 2017

**Current Status:** Referred to the Subcommittee on Digital Commerce and Consumer Protection.

**Latest Action:** Referred to the Subcommittee on Digital Commerce and Consumer Protection. (Feb 17, 2017)

**Official Text:** <https://www.congress.gov/bill/115th-congress/house-bill/1108>

### Sponsor

**Name:** Rep. DeLauro, Rosa L. [D-CT-3]

**Party:** Democratic • **State:** CT • **Chamber:** House

### Cosponsors (4 total)

| Cosponsor                                 | Party / State | Role | Date Joined  |
|---|---------------|------|--------------|
| Rep. Doggett, Lloyd [D-TX-35]             | D · TX        |      | Mar 8, 2017  |
| Rep. Langevin, James R. [D-RI-2]          | D · RI        |      | Mar 8, 2017  |
| Rep. Raskin, Jamie [D-MD-8]               | D · MD        |      | Mar 8, 2017  |
| Rep. Slaughter, Louise McIntosh [D-NY-25] | D · NY        |      | Oct 25, 2017 |

### Committee Activity

| Committee                     | Chamber | Activity    | Date         |
|-------------------------------|---------|-------------|--------------|
| Energy and Commerce Committee | House   | Referred to | Feb 17, 2017 |

### Subjects & Policy Tags

#### Policy Area:

Health

### Related Bills

*No related bills are listed.*

## **Recall Unsafe Drugs Act of 2017**

This bill amends the Federal Food, Drug, and Cosmetic Act (FFDCA) to require producers of medications to notify the Food and Drug Administration (FDA) of the identity and location of a medication if the producer has reason to believe: (1) that the medication is adulterated or misbranded; and (2) there is a reasonable probability that the use or consumption of, or exposure to, the medication will cause a threat of serious adverse health consequences or death to humans or animals.

The FDA may: (1) request that the distributor of a medication that is in violation of the FFDCA voluntarily recall the medication; (2) require the distributor of a medication that may cause serious adverse health consequences to immediately cease distribution of the medication; (3) recall a medication for which distribution has been ceased after giving the distributor an opportunity for an informal hearing; and (4) immediately recall a medication that presents an imminent threat of serious adverse health consequences. Distributors may appeal these FDA orders.

In the case of a recall, the FDA must notify consumers and state and local health officials to whom the medication was, or may have been, distributed.

Medication distributors must have a recall plan in effect.

## **Actions Timeline**

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- **Feb 17, 2017:** Referred to the Subcommittee on Digital Commerce and Consumer Protection.
- **Feb 16, 2017:** Introduced in House
- **Feb 16, 2017:** Referred to the House Committee on Energy and Commerce.

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