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Getting Started: EU FMD Guide to Pharma Serialization



If your company is part of the pharmaceutical supply chain in Europe, preparing for the EU Falsified Medicines Directive should now be a priority. The regulation presents considerable challenges and responsibilities—serialization, government reporting, verification—so understanding the requirements and knowing how to prepare for the February 2019 deadline is a must.

Our latest infographic can get you started on the road to EU FMD compliance. It covers the regulations' essential facts, including which nations are affected—and which have a compliance grace period—what each supply chain company must do to comply, and other vital details about:

- Which products can be grandfathered.

- The 2D DataMatrix code format.
- The specific data that must be uploaded and maintained.

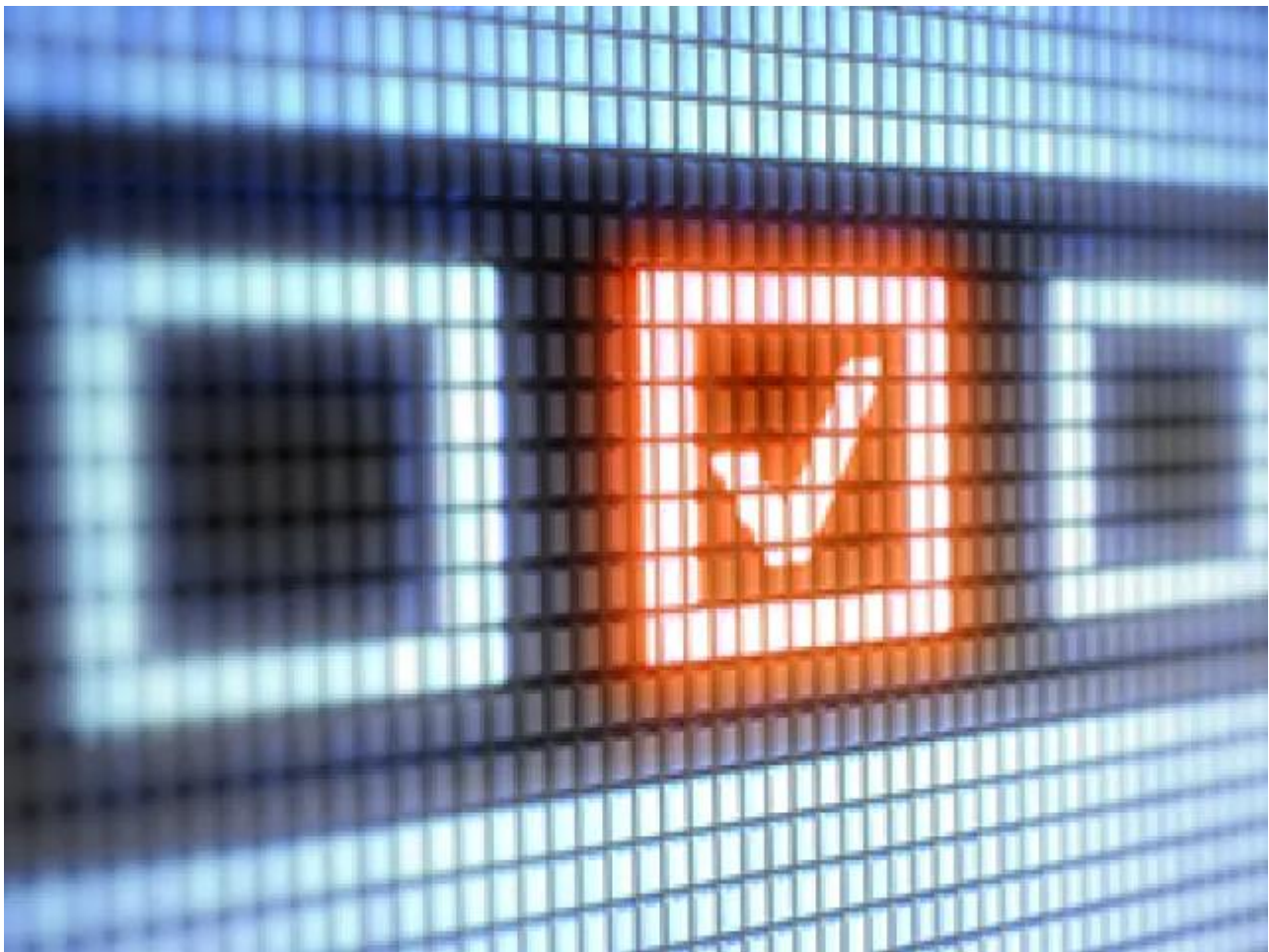
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More Serialization and Compliance Resources



Can You Afford a Manual Approach to EU FMD Compliance?

Manual data upload through the EMVO portal is simple in concept, but preparing and maintaining that data can be a challenge for smaller companies.

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HOSPITALS, PHARMACIES, AND DISPENSING DOCTORS:

The Impact of EU FMD on Workflow and Stock Management

WHAT DOES THE LAW MANDATE?

Under EU FMD, hospitals, pharmacies, and dispensing doctors must verify and decommission medicines against their local National System before they are dispensed to patients.

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graph LR; Manufacturer[Pharmaceutical Manufacturer] -- Product Flow --> Wholesaler[Wholesaler]; Wholesaler -- Product Flow --> Pharmacist[Pharmacist]; Pharmacist --> Patient[Patient];
```

Report Product Master Data and

Verification and Decommission

The Impact of EU FMD on Pharmacy Workflow and Stock Management

View an infographic on how EU FMD changes hospital and pharmacy workflows.

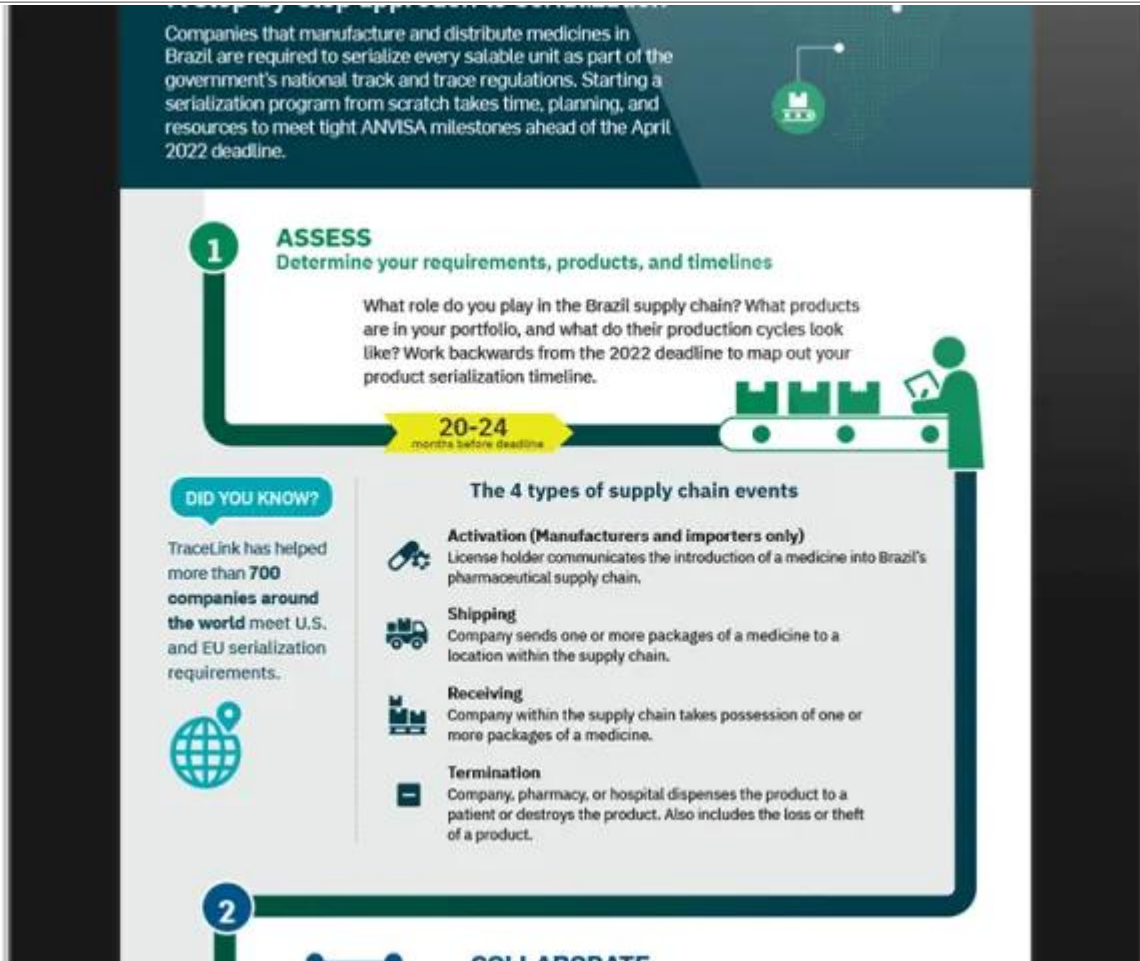
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What Are the 3 Major Requirements of EU FMD?

The EU Falsified Medicines Directive contains requirements for safety features and verification that details how companies must establish serialization and reporting.

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Brazil Compliance: A Step-by-Step Approach to Serialization

Begin your Brazil serialization journey and see why you need to start today to meet the April 2022 ANVISA deadline. Download the infographic.

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