

How to declare if you are a manufacturer or distributor of in vitro diagnostic medical devices

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Post-market surveillance and vigilance

Manufacturer is required to submit vigilance reports to the ANSM for any incidents occurred in France involving in vitro diagnostic medical devices, in accordance with Regulation (EU) 2017/746.

The manufacturer must also take appropriate safety action when required.

Further information about [reporting adverse incidents and corrective actions to ANSM](#) is available for manufacturers of in vitro diagnostic medical devices.

We encourage manufacturers to use European Medical Device Nomenclature (EMDN).

Manufacturer Incident Report (MIR)

- Manufacturers should notify the ANSM of MIR using the [MIR Manufacturer Incident Report \(MIR\) form](#).
- Manufacturers should sent MIR (pdf and xml format) by mailbox: reactovigilance@ansm.sante.fr.
- To monitor the registration of MIR, manufacturers have an access via [Vigimater](#) to put ANSM's reference.

New MIR form - mandatory as from 4 months after the date of the publication of the next update of MIR 7.3.1. PDF and related files

A new version of MIR 7.3.1 will be applicable.

Documentation related to this update is available on the [European Commission's website](#).

[Access to Vigimater](#)

Field Safety Corrective Actions (FSCAs)

- Manufacturers should notify the ANSM of FSCA using the [FSCA Report Form](#).
- Manufacturers should sent FSCA by mailbox: reactovigilance@ansm.sante.fr.

Trend report

- Manufacturers should notify the ANSM of trend report using the [Manufacturer's Trend Report Form](#).
- Manufacturers should sent Trend Report by mailbox: reactovigilance@ansm.sante.fr.

Periodic Summary Report (PSR)

- Manufacturers should notify the ANSM of Periodic Summary Report using the [Manufacturer's PSR Report Form](#).

- Manufacturers should sent PSR by mailbox:reactovigilance@ansm.sante.fr.

Documents

Reporting

- [Manufacturer Incident Report \(MIR\) form](#)
- [Manufacturer incident report Helptext 2020](#)
- [FSCA Report Form](#)
- [Field safety notice template](#)
- [FSN customer reply](#)
- [FSN distributor/importer reply](#)
- [Manufacturer's Trend Report Form](#)
- [Vigilance related of medical device \(MD\) or in vitro diagnostic medical devices \(IVD\) : Q&A on reading anomalies of the UDI carrier](#)