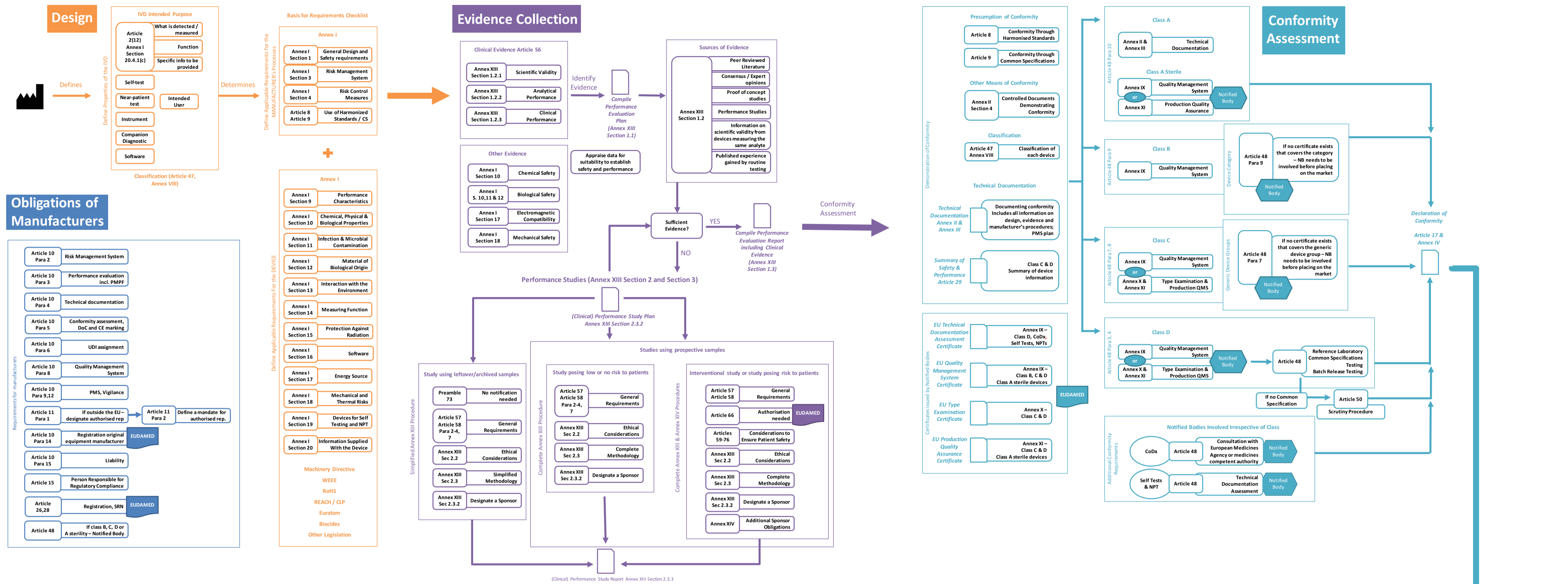


### Overview of requirements under the IVD Regulation

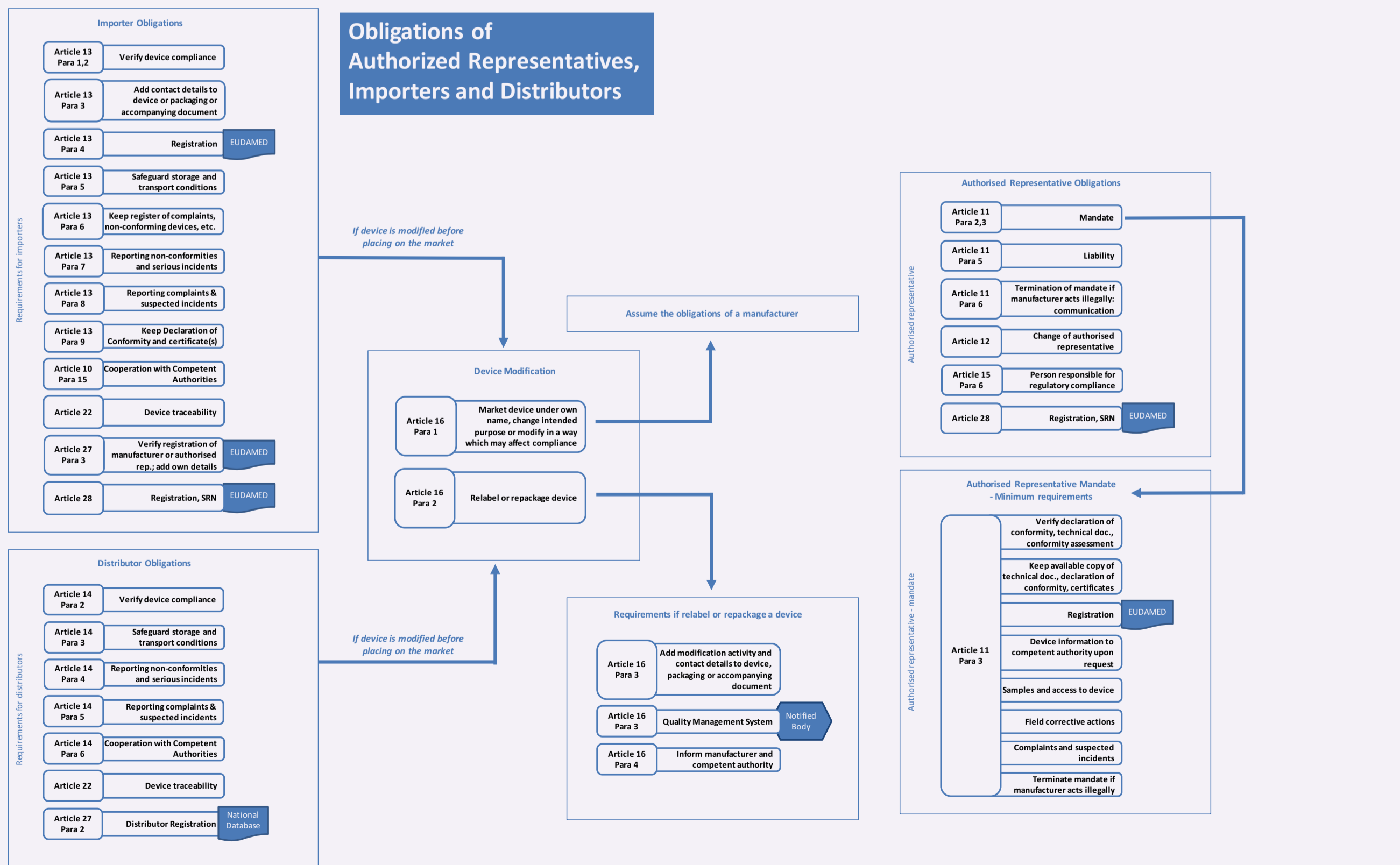
Regulation 2017/746/EU on In Vitro Diagnostic Medical Devices

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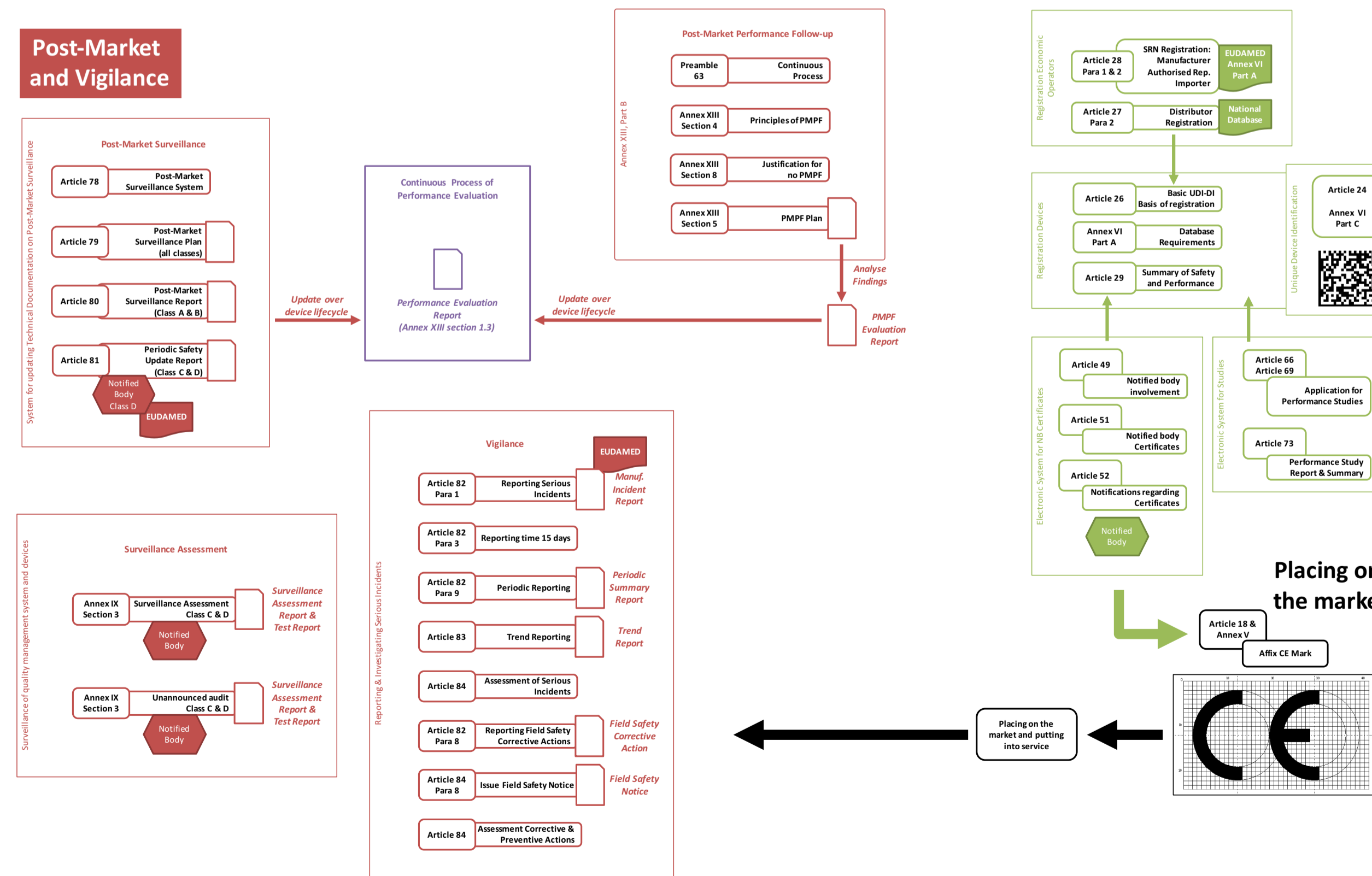
MedTech Europe reserves the right to change or amend the flowchart or any parts thereof at any time without notice. For more information please contact the regulations & industrial policy department: [regulatory@medtecheurope.org](mailto:regulatory@medtecheurope.org)



### Obligations of Authorized Representatives, Importers and Distributors



### Post-Market and Vigilance



# Overview of Regulation 2017/746/EU on *In Vitro* Diagnostic Medical Devices