

# Implementation of the EU MDR (2017/745) Requirements



**Founded in 2000**



**Expertise backed  
by over 500 SME's**



**Headquartered  
in Southeastern  
Wisconsin**



**Florida, Colorado  
and Eastern  
Europe**



**ISO 9001:2015**

At RCA, we provide worldwide services to the pharmaceutical, medical device, biologics, combination products, and compounding pharmacies industries for resolution of compliance and regulatory challenges.

Our backgrounds include every facet of R&D, operations, regulatory affairs, quality and manufacturing. We are used to working on the front lines and thriving in the scrutiny of FDA and globally regulated companies.



As your partners, we can assist with the following services related to the EU MDR requirements:

**Review of  
Product Portfolio**

**Staff  
Augmentation**

**Gap Assessment  
of QMS**

**Remediation and  
Implantation**

### Perform a Comprehensive Review of Client's Entire Product Portfolio

- Confirm whether Products have been affected by the MDR Definitions and/or Reclassification of certain types of products
- Prepare Product Rationalization Strategy Reports
- Check the status of MDD Certificate expiry dates and prepare an EU MDR Certificate Compliance Strategy for each product



### Remediation and Implementation of EU MDR Compliance Plan

- Supply Chain Requirements
- Development/Implementation of an EU MDR Compliance Plan for systems and devices
- Remediation of EU MDR Compliance Gaps
- Development/Implementation of EUDAMED Database compliance plan



### Staff Augmentation

- Readiness Audits / Pre-Certification Audits and/or Assessments

### Perform a Gap Assessment of the current Quality Management System (QMS) and its Current Processes and Procedures Compliance

#### Technical Documentation Requirements

