

Your Quality & Regulatory Compliance Buddy

# REGULATORY COMPLIANCE FOR INDIAN & EUROPEAN MARKETS

RIFC aims to inform and assist **bio-entrepreneurs** in planning, seeking and securing regulatory approvals







Joint initiative of BIRAC & Venture Center under the BIRAC Regional Bio-Innovation Center (BRBC) Program



#### At RIFC

- Regulatory compliance for the Indian, European and US markets
- Competent team with diverse group of experts
- Specializes in Medical devices
- Regular events and clinics for our beneficiaries

## **Why Choose Us**

- 26 National/International Standards (E.g. ISO 13485, ISO 10993, ISO 14971, ISO 9001)
- 150+ ISO 13485 QMS & ISO 9001 Templates
- 250+ Dossier Templates (For India, EU and US regulations)
- Database of the Testing Labs and Certification Bodies
- 250+ startups impacted
- 15+ dedicated regulatory experts
- 11 ISO 13485 Certifications assisted out of which 5 certifications received



With the RIFC team guidance and support, you can confidently navigate complex regulatory requirements and empower your business with compliance.

RIFC team works to offer a comprehensive support & strategic guidance for your medical device startup, not only to ensure compliance but also to leverage regulatory frameworks as a competitive advantage for your business and enhance investor confidence.

## Services offered by RIFC team







Nº	ACTIVITIES	INDIAN	EUROPEAN	US FDA
1	General Advisory	Ø	Ø	Ø
2	Planning Regulatory Pathway	Ø	Ø	Ø
3	Standards Interpretation	Ø	Ø	Ø
4	ISO 13485 Document Preparation	$\bigcirc$	Ø	Ø
5	ISO 13485 Document Review	Ø	Ø	Ø
6	Risk Management	Ø	Ø	Ø
7	Clinical Trial Study Plan	Ø		
8	Clinical Trial Advice	Ø		
9	ISO 13485 Internal Audit	Ø		
10	ISO 9001 Document Review	$\bigcirc$		
11	HIPAA General Advisory			Ø





# CONTACT US TODAY AND SCHEDULE YOUR INTRODUCTORY MEETING

#### Connect with us





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