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Engineering partnership towards patient centric ecosystem

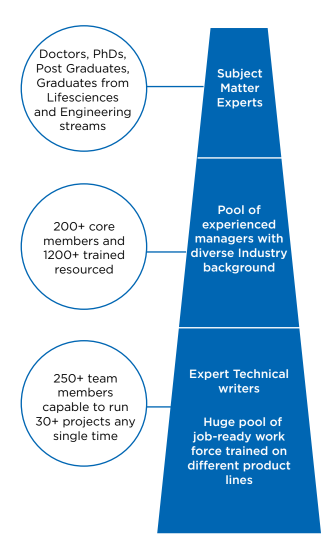
Acceleration | Compliance | Value

Regulatory affairs capabilities

With highly experienced 200+ **regulatory affairs** specialists and 1200+ trained resources, established relationship with notified bodies, experience in supporting remediation of QMS dor medical devices customers, and expert physicians within the clinical team, our Regulatory and Clinical CoEs bring in right domain expertise to provide simple and efficient approach to ensure **regulatory compliance** and quality.

HCL REGULATORY CENTRE OF EXCELLENCE

Values delivered



8 of 10 Top Medical Device firms and 4 of 5 Top IVD Device Firms

Interacting with 40+ Sites, working in 4 Geos, Multilingual like English, German, French, Japanese

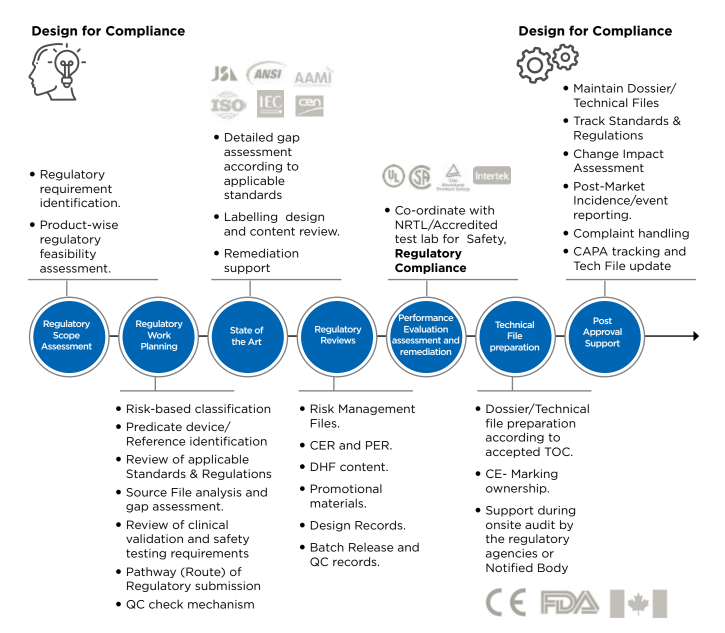
7,000+ Tech Files supported for assessment and remediation

20+ QMS Assessment - Effectiveness Analysis and Remediation of procedures and templates

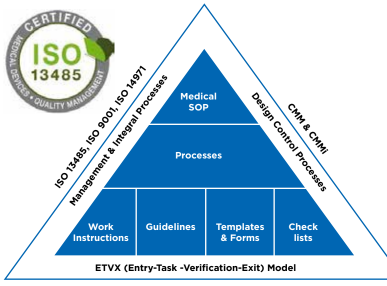
Process Maturity: CMMI Level 5 | ISO:13485 | BS 7799 | ISO:27001

Successful support in implementation of process and product quality standards (e.g. ISO 13485, ISO 4971, ISO 60601x, ISO 10993, IEC 61010x, 61326x IEC 62304, IEC 62366x and UDI)

At every step of your regulatory affairs journey



MQMS, our principle and approach



IEC 62304, 21 CFR Part 820

Deployment Tools

- Intranet based system
- Process database
 - Risk / Reusable components
 - Best Practices
 - Learning's

Deployment Mechanism

- Induction Training
- Calendar Based Trainings
 - MQMS Training
 - ISO 13485, ISO 14971, IEC 62304 & QSR awareness Trainings
- Role based Training PQA, CC
- Process update Training

- ISO 13485 Medical Devices-QMS-Regulatory Requirement
- 21 CFR Part 820 US FDA Quality System Regulations (QSR)
- 21 CFR Part 11 Electronic Records; Electronic Signature
- **ISO 14971** Risk Management for Medical Devices
- ISO/IEC 62304 Medical Device SDLC Process
- IEC 62366 MD Application of Usability Engineering
- AAMI TIR45 Guidance on the use of AGILE practices in the development of medical device software
- USFDA Guidance Documents
- **GAMP 5** (Good Automated Manufacturing Practice, by ISPE)
- **GDP** Good Documentation Practice
- ISO 9001 QMS General Requirement
- MHLW Ministerial Ordinance No.169: Ordinance on Standards for Manufacturing Control and Quality Control of Medical Devices and In Vitro Diagnostic Reagents
- **CMMI** Capability Maturity Model integration (Ver 1.3) Level 5
- ISO/IEC 27001 Information Security Management System (ISMS)

CASE STUDY: QMS assessment and update for MDSAP



- Total Team Size : 7 No's
- Gen Regulatory Lead (Offshore): 2 No's
- Gen Regulatory Engineer (Offshore) : 5 No's
- Duration of the Project : 12 Weeks

- Preparation of process wise work sheet mapping with MDSAP requirement
- Collect the required inputs from client
- Conduct the gap assessment as per companion document (MDSAP) requirements
- Address & Remediate the Gaps Identified
- Review & Querv Resolution

- Early meeting of project timelines
- 100% QMS Coverage
- Customer Satisfaction

CASE STUDY: Complaint Management – Data Analysis and Trend Analysis

Project Background:

Medical Device Customer was looking for someone who can review the complaint records of about 70K numbers and make determination if the content has sufficient info to determine if this is a complaint or if additional info is needed to make a decision. Also the person would need to review adverse event survey completed by complaint call center to determine if the survey appears to be completed correctly.

Project Scope:

To understand the Customer Complaint handling procedure and work closely with the PMS Team to evaluate the end Customer Product Complaints, verify whether the Adverse Event Survey is sufficiently aligned with the complaint text, review and bring it to Closure.

Accomplishments:

- Complete ownership of Intake data which includes (Initiation, Field Dispatch, Escalation to Management)
- Complete ownership on creation of Quality Notifications from Investigation.
- Supporting the Customer PMS team on the Closure which includes (Closed solved in Field and Waiting for Approval)

Stages:

- Complaint Evaluation and Processing
- Daily Reporting
- Quality Notification Creation
- Complaint Allocation
- Tracking and Trending
- Regional Reporting
- Seamless Communication
- Initiation
- Field Dispatch
- Escalation to management

60,000 + Complaints reviewed

40 + Complaints/day/head

100 + Quality Notification Creation

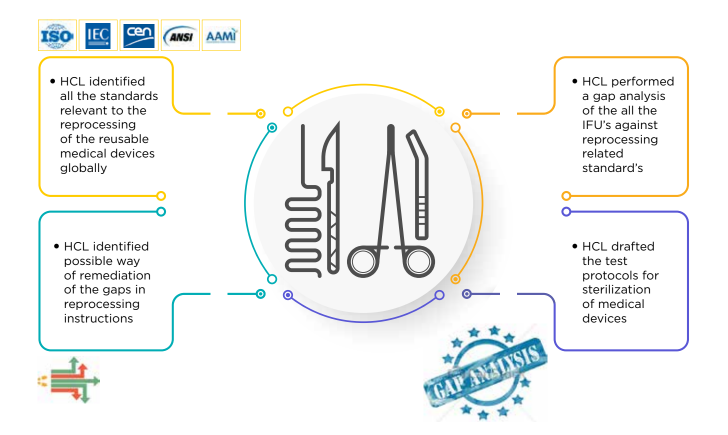
Process Understanding **Evaluating** & Processing Additional Information Quality **Notifications Tracking and** Trending of Metrics

CASE STUDY: CAPA for Cleaning and Sterilization of reusable Medical Device

Business Statement:

The USFDA audited OEM and issued an inspectional observation (#483) for one of their reusable product. OEM engaged HCL for analysis, CAPA proposal and implementation.

FDA requirements for the reprocessing of the reusable medical devices has been changed in March 2015



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