

Kind Attention

Regulatory

Quality | Regulatory

Clinical | Drug Safety

Clinical | Teams

& Engineering

INDIVIRTUS HEALTHCARE SERVICES PVT. LTD.
INDIVIRTUS GLOBAL CRO PVT. LTD.
INDIVIRTUS ECOLOGICAL SERVICES PVT. LTD.



PHARMACOVIGILANCE

BA/BE STUDIES & CLINICAL TRIALS

BROWNFIELD PROJECTS

VALIDATION (CSV/ EQUIPMENT/ HVAC)

CALIBRATION SERVICES (NABL ACCREDITED LAB)

TECHNOLOGY TRANSFER GMP,
SCHEDULE M
COMPLIANCE
& TRAINING

ADE/PDE
OEL/OEB &
RISK ASSESSMENT
OF IMPURITIES

REGULATORY AFFAIRS

About Us:

Indivirtus (Indian company with virtuousness, character, courage, transparency & truthfulness) is a group of companies offering cost effective solutions wrapped with quality.

Indivirtus Global CRO is the 5th group company, under the umbrella of Indivirtus. The company will have focused approach in the field of Clinical Research i.e. BA/BE Studies and Clinical Trials.

BA/BE STUDIES & CLINICAL TRIALS

- Bioanalytical Service (Small Molecule PK Studies)
- BA/BE Studies in Healthy Volunteers & Patients
- Biostatistics and Statistical programming
- Early Phase (Phase-I) & Late Phase (Phase II/III/IV/PMS)
 Clinical Trials
- Data Management and Electronic Data Capture (EDC)
- Medical Affairs & Writing



For any query, please contact



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PHARMACOVIGILANCE

- Pharmacovigilance Database Setup
- Medical Information Call Centre (MICC)
- PSMF and SOPs Preparation
- ICSR Case Processing and ADR Reporting
- Literature Monitoring
- PSUR/ PBRER/ PADER
- Risk Management Plan
- Signal Management

TOXICOLOGICAL RISK ASSESSMENT

- Toxicological Risk Assessment of Medical Devices
- ADE/PDE/OEL & OEB Calculation for Cleaning Validation
- Genotoxic (Mutagenic) Evaluation of impurities as per ICH Q3
 A (R2), ICH Q3 B (R2), ICH Q3 C (R8) & ICH M7 guidelines.
- Training on Cleaning Validation
- Safety Assessment Report of Finished Formulations
- Preparation of MSDS/SDS











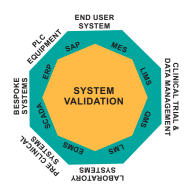


REGULATORY COMPLIANCE AND MEDICAL DEVICE REGISTRATION

- Preparation, Submission and Review of the CTD/ ACTD Dossiers
- Medical device (Class-A, B, C & D) Filing & Submission to CDSCO as per MDR-17 rule
- CEP/DMF Filing & Submission to EDQM/FDA/Health Canada.
- Clinical/Non-Clinical Overview and Bio Waiver Preparation
- Managing Regulatory Response & Remediation Activities
- M.A.H. representation, Qualified Person (Q.P.) & Batch Releasing & Testing Site for Europe Submission

VALIDATION (CSV/Equipment/HVAC)

- Laboratory Systems (HPLC/GC/KF/UV/FTIR/POLARIMETER)
- Process Automation Systems (PLC/HMI/SCADA/IPC)
- Environment Monitoring Systems (EMS)
- Enterprise Applications (SAP/ERP/LIMS/DMS/QMS/LMS etc.)
- Temperature Mapping (SIP/ Autoclave/ Tunnel/ Cold Room/ Warehouse)
- HVAC (DOP/Air Velocity/NVPC/Filter Integrity)



GMP APPROVALS FROM REGULATORY AUTHORITIES

EU-GMP USFDA WHO-GMP & PICS TGA Health Canada GCC ANVISA COFEPRIS (Mexico)

- Vender Assessment Audits for Plant & Product Approvals
- Assessment of
 - Plant Layout Design, Equipment, Infrastructure, Process Flow, Flow of Man & Material, Raw Material and FG Store, Documentation
 - Validation, Calibration & Qualification
 - CAPA and Risk Management with Appropriate Mitigation and Contingency Plan
 - GAP Assessment as per Schedule M Guideline







TECHNOLOGY TRANSFER

- R&D to GMP site transfer
- Formulation Development & Technology Transfer
- Know How Based Technology Transfer (1800+ products)
- Analytical Method Development & Validations

GREENFIELD/ BROWNFIELD PROJECTS

Turnkey Pharmaceutical Project of

- Finished Dosage Forms
- Active Pharmaceutical Ingredients
- Intermediates





TRAININGS ON

- Cleaning Validation
- GxP (GLP, GMP, GCP, GDP etc.)
- Revised Schedule M
- WHO TRS guidelines and Annexures
- Pharmaceutical Quality Systems like Data Integrity, Quality Culture etc.
- Quality Management Systems

CALIBRATION LAB (NABL ACCREDITED)

Thermal: Temperature & Relative Humidity (RH), IR Thermometers, Furnace, Oven etc. Electro Technical: AC/DC Current, Voltage, Resistance, Inductance, Time, Frequency etc.

Mechanical: Dimension, Speed & Acoustics, Length etc.

Pressure/Vacuum: Pressure/Vacuum/Magnehelic Gauges, Transmitter, Safety Valves etc.

Mass & Volume: Weights, Weighing Balance, Pipettes, Burettes etc. Others: pH Meters, ORP, Conductivity Meters, TDS, Flow Meter etc.



OUR

- **U** nique
- **S** elling

- Team with Experience of Performing 600+ BA/BE Studies
- Team with experience of conducting 100+ Phase III & 120+ Phase IV Trials
- End to End Pharmacovigilance Services
- Only company with a team of 6 Certified Toxicologists
- Experience of preparing 13000+ PDE/ADE & OEL Reports
- Experience of preparing 2500+ Genotox (Mutagenic & Risk Assessment Reports of Degradation/Nitrosamine/Elemental/Extractable Leachable Impurities
- P roposition Served 680+ clients in 55 countries
 - Experienced & Certified Auditors to Conduct GMP and Vendor Audits
 - Experienced Team to perform Technology Transfer
 - Vast portfolio of products, readily available for Technology Transfer i.e. 850+ Products for Regulated Markets & 1800+ products for ROW markets



GLOBAL OFFICES

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For Business Queries

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