



**Kind Attention**  
Quality, Regulatory,  
Clinical, Drug Safety  
& Engineering Teams

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

  
**680+ Happy Clients**  
in 55 Countries

PHARMACOVIGILANCE

BA/BE  
STUDIES &  
CLINICAL  
TRIALS

GREENFIELD/  
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 5<sup>th</sup> 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

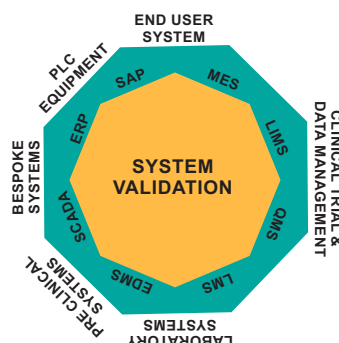


## REGULATORY COMPLIANCE AND MEDICAL DEVICE REGISTRATION

- 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

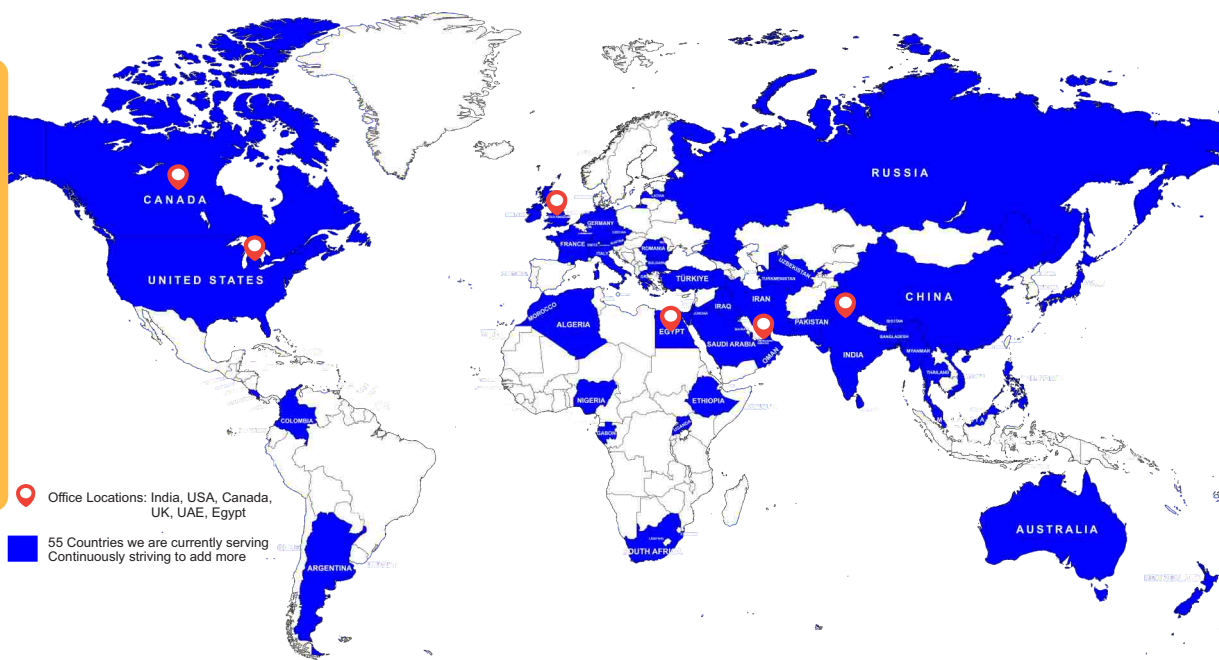
**U**nique

**S**elling

**P**roposition

- 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
- 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 FOOTPRINTS



#### Head Office

521-522, Top Floor,  
Taj Plaza, TDI City,  
Sector-118, Airport Road,  
Mohali, Punjab, India-160059

#### Raipur

404- Fourth Floor, Avinash Time  
Square, Sector - 19, Nawa Raipur,  
Chhattisgarh, India-492018

### GLOBAL OFFICES

#### Bhubaneswar

Plot No-249, Patrapada Khorda,  
Near Police Academy AIIMS Nagar,  
Bhubaneswar, Khordha,  
Odisha, India-751019

#### Bhopal

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To find out more,  
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