



COMMITTED TO YOUR SUCCESS IN CHINA

From Concept to commercialization and Beyond

WHO WE ARE



We are a global consulting firm that provides **Regulatory Affairs**, **Quality**, **Clinical** and **Safety services** to pharmaceutical and medical device companies worldwide.

With local offices in Shanghai and Beijing and local team of regulatory affairs, medical, quality and pharmacovigilance professionals we provide tailored services to our clients in China in compliance with the National Medical Products Administration (NMPA) regulations, the Local Health Authority (HA) of China.

WE ARE LICENSED by the NMPA to:



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Act as your Local Representative in China

and handle all your local regulatory, clinical, quality and pharmacovigilance activities. Register your products in China in compliance with NMPA regulations and Standards. Manage your your local clinical studies in line with NMPA requirements.

Import and release your products in China.

Our commercial team can help you define the perfect solution to access the China market by providing the following services:

- Market-access Strategy
- Pricing & reimbursement strategies
- Application for pricing
- Ouantitative & Oualitative market research
- Due-diligence and asset evaluation
- Evidence-based portfolio prioritization
- Review of Target Product Profile (TPP) and label claims
- Evidence and economic needs assessment and evaluation
- Commercial and marketing strategies



- Regulatory Strategy
- Regulatory Intelligence
- Registration of drugs and biologics
- Preparation of consultation meeting with HA
- DMF registration
- Medical writing
- Regulatory operations and publishing
- Orphan drugs designation
- Fast track application
- Preparation and submission of CTA
- Nutraceuticals and vitamins claims' review
- Nutraceuticals and vitamins formulation review
- Cosmetic packaging and labeling review & Registration



SERVICES



Global Regulatory Partners develops the appropriate regulatory strategies that:

CLINICAL

Local sponsor service

- PI recruitment
- Contracts with sites
- Protocol development
- eCRF and EDC
- ✓ IRB approval
- Import investigational products.
- Site initiation meetings
- Monitoring of clinical studies
- Data management Statistics
- Clinical study report (CSR) writing
- GCP audits

SAFETY

Receipt, review, and coding of adverse events

- Safety reports writing
- Safety reporting to different health Authorities
- Good Vigilance Practice (GVP)
- Product safety profile writing and updates
- Post-Marketing Surveillance (PMS) Studies
- Safety annual reports
- Medical literature screening
- Health authority PV database monitoring
- Risk Management Reports writing & submission to HA

QUALITY

- Quality System Management (QMS) implementation
- Preparation for HA inspection
- Standard Operating Procedures
- ✓ (SOP) writing
- Ouality manual writing
- Due diligence audits
- GXP quality audits
- CAPA management
- Mockup quality audits

EPRESENTATION

Products registrations and lifecycle maintenance

- Application for products pricing
- Local product release
- Pharmacovigilance
- Recalls and complaints handling D Distributors selections
- Local Labeling
- DMF registration and maintenance



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