

# Di Renzo® Regulatory Affairs

In 1985, *Di Renzo®* Regulatory Affairs began its regulatory consulting for medicines for human and veterinary use, food supplements, Presidi Medico Chirurgici (PMC) and then biocides, medical devices, In Vitro Diagnostics (IVDs), cosmetics and a range of other related services. As a result of the development of national and international standards, ever increasing business needs, and thanks to the technical-scientific and administrative structure which *Di Renzo®* Regulatory Affairs has adopted, more and more companies are entrusting

Di Renzo® Regulatory Affairs with numerous activities that were previously performed in-house.

Di Renzo® Regulatory Affairs collaborates with an international network of regulatory agencies and consulting firms in all countries of the European Union and the main non-European countries.



### Areas of interest

#### Regulatory Affairs

- o Medicinal products for human use
- o Medicinal products for veterinary use
- o Food supplements, Food for Specific Groups (FSG), Foods and Novel foods
- o Cosmetics
- o Medical Devices (MDs) and In Vitro Diagnostics (IVDs)
- o Presidi Medico Chirurigici (PMC) and Biocides

#### Vigilance

- o Pharmacovilance for medicinal products for human use
- o Surveillance and vigilance for MDs and IVDs
- o Cosmetovigilance
- o Foodvigilance and phytosurveillance

- Scientific information and advertising
- Clinical trials
- Quality Services
- Audits
- Legal and notary services
- Translations
- Publications in the Italian Official Journal



#### REGULATORY AFFAIRS

# Medicinal products for human use

- Feasibility studies for business projects
- **Update** on national and international regulations
- Due diligence, gap analysis and preparation of chemical, clinical and pharmaco-toxicological expertise
- Preparation of applications for marketing authorizations for national, Mutual Recognition,
   Decentralized and Centralized procedures initiating from either Italy or another EU country
- Preparation of dossiers in CTD and eCTD
- Preparation of variations, extensions, transfers of ownership and marketing authorization renewals
- Drafting, check and translation of Summaries of Product Characteristics (SmPCs), labels and Package Information Leaflets (PILs) in accordance with current regulations
- Readability Test for the Package Information Leaflet (PIL)
- Preparation and check of artworks for immediate, secondary packaging and PILs
- Traceability of medicinal products, application of optical antifraud stickers and serialization
- Consultancy and preparation of price dossiers, Health Technology Assessment (HTA) and pricing negotiations
- Request for Certificates of Pharmaceutical Product (CPPs) and other documents required for export and their relevant legalization

- Regulatory consultancy on parallel imports, for importations due to drug shortage and for direct importation
- Technical and administrative regulatory consultancy: homeopathic products, medical gases, radiodrugs, and allergens
- Advice on new applications, variations and renewals of manufacturing and importation authorizations for active substances or finished products
- Submission of ASMFs of APIs and dossiers of starting materials to the EDQM for the registration,
   renewal and variation of CEPs
- Requests for GMP certificates
- Assistance in the registration and maintenance of EMA accounts (IAM; IRIS i-SPOC, eAF Applicant)
- Assistance in obtaining authorization for the storage of medicines (distributors)
- Assistance in the procedures related to narcotic drugs and drug precursors

Regulatory advice during all the steps for the development of orphan drugs, from the obtaining of the initial orphan designation to the marketing authorization

## Medicinal products for veterinary use

- Feasibility studies for business projects
- Regulatory updates
- Preparation of applications for marketing authorizations for national and European procedures
- Preparation of variations, extensions, transfers of ownership and renewals of marketing authorizations
- Due diligence, gap analysis and preparation of chemical, clinical and pharmaco-toxicological expertise
- Translation and update of Summaries of Product Characteristics (SmPCs), labels and Package Information Leaflets (PILs) in accordance with current regulations
- Preparation of artworks for immediate, secondary packaging and PILs
- Request for GMP certificates, Certificates of Pharmaceutical Product (CPPs) and other documents required for export and the relevant legalization
- Consultancy on new applications, renewals and variations of manufacturing authorizations for active substances or finished products
- Revision of the labels and the composition of animal feed for veterinary use in accordance with current legislation

# Food supplements, Food for Specific Groups (FSG), Foods and Novel foods

- Feasibility studies for business projects
- Preparation of the scientific rationale for supplements containing herbal preparations (botanicals)
- Conformity assessment of labelling and composition
- Assistance with Food for Specific Groups (FSG), including Food for Special Medical Purposes (FSMP)
- Assistance on nutritional and health claims in accordance with the Regulation (EC) no. 1924/2006
- Authorization procedures of novel foods at the European Commission
- Development of artworks for the packaging material
- Notification procedure with the Italian Ministry of Health of food supplements and other foods subject to notification
- Assistance in the notification procedures for the marketing in most EU countries and in some non-EU countries
- Evaluation of advertising material and drafting of brochures and leaflets
- Assistance with biological certification of food supplements
- Assistance with registration of companies as Food Business Operator (FBO)
- Assistance with the upland of products to the main databases, including: Farmadati Italia, Codifa and CSF Sistemi

### Cosmetics

- Feasibility studies for business projects
- Consultancy in order to comply with the provisions of Regulation (EC) no. 1223/2009
- Review of the technical and administrative documentation provided by the company
- Conformity assessment of labelling
- Preparation of artworks for cosmetic packaging
- Preparation of data sheets on the toxicological characteristics of cosmetic ingredients
- Preparation of the safety assessment of cosmetic products
- Preparation of the **Product Information File** (PIF)
- Entering data into the European Cosmetic Products Notification Portal (CPNP)
- Evaluation of suppliers and audits by technicians at manufacturing companies
- Request for Free Sale Certificates (FSCs) and the relevant legalization
- Evaluation of advertising material

# Medical devices (MDs) and in vitro diagnostic medical devices (IVDs)

- Feasibility studies for business projects
- Advice on Italian and European regulations in force in Italy, in the EU, and in some non-EU countries
- Assumption of the role of Authorized representative for extra-EU companies
- Assumption of the role of UK Responsible Person (UKRP)
- Revision and verification of the compliance of technical documentation and requirements for the registration of products in Italy, in the EU, and in some non-EU countries
- Registration of manufacturers, importers, authorized representatives, and assemblers in the EUDAMED form regarding the business operators to obtain the single registration number (SRN)
- Notification of MDs and IVDs in the Database/ Repertorio of the Italian Ministry of Health and in other EU Member States

- Registration of manufacturers of custom-made medical devices
- Preparation of technical files for CE marking
- Drafting of Clinical Evaluation Plan (CEP) and Clinical Evaluation Report (CER)
- Contacts with the Notified Bodies and consultations for obtaining the CE marking
- Contacts with qualified laboratories to conduct tests on products
- Assumption of the role of Quality Assurance (QA)
- Assumption of the role of Person responsible for the regulatory compliance (PRRN)
- Regulatory activities related to the import/ export of MDs and IVDs
- Request for Free Sale Certificates (FSCs) and relevant legalization
- Verification and assistance with the authorization of advertisements

# Presidi Medico Chirurgici (PMC) and biocides

- Feasibility studies for business projects
- Information on the Italian legislation on PMCs
- Advice on the regulations in force and on the evolution of the legislation on biocidal products, in particular on the transition period for PMCs-biocides
- Verification of the inclusion of the active substances in the **Union list** in order to classify the product as a PMC/biocide/product of free sale or otherwise, in accordance with current regulations
- Verification of the requirements for registration in certain EU and non-EU countries
- Assistance in the preparation and submission of the registration dossier to the Italian Competent Authorities and relevant authorization process for a PMC

- Assistance in the preparation of the dossier for the manufacturing authorization for PMCs and the relevant authorization procedure at the Italian Ministry of Health
- Preliminary assessment, preparation and submission of the dossier for the authorization of a biocidal product and assistance throughout the entire authorization process
- Identification of studies to be performed (chemical-physical, toxicological and ecotoxicological and efficacy studies, etc.) in accordance with the product type (PT) of interest, the active ingredient and the intended use of the product
- Contacts with qualified laboratories for product testing

- Preparation and verification of PMC and Biocide labels in compliance with regulatory requirements
- Request for Free Sale Certificates (FSCs) for PMCs and relevant legalization
- Development of mock-ups of labels and logos, design and development of brochures, data sheets and advertising materials
- Preliminary assessment of advertisement material for PMCs and requests for the appropriate authorization from the Italian Ministry of Health
- European notification into the ECHA portal (PCN)



### **VIGILANCE**

# Pharmacovigilance for medicinal products for human use

- Assumption of the role of European Qualified Person for Pharmacovigilance (EU-QPPV) –
  including the availability of a Deputy
- Assumption of the role of Local Contact Point for Italy including the availability of a Deputy
- Management of the entire pharmacovigilance quality system and assumption of the role of pharmacovigilance quality Responsible Person.
- Periodic training for internal staff of MA Holders/Italian Affiliates on pharmacovigilance and operational procedures
- Periodic pharmacovigilance training for medical sales representatives
- Screening for adverse reactions in EudraVigilance and case management
- Conducting pharmacovigilance audits
- Periodic verification of Italian and international scientific literature for medicines and active ingredients
- Medical evaluation of adverse drug reactions (ADRs)
- Follow-up management
- Data Entry and Quality Control of the ICSR in the safety database (SafetyDrugs®)
- Submission of ICSRs to Competent Authorities (Eudravigilance)
- Safety data exchange with business partners and English translation of Italian ICSRs in CIOMS

- and/or in XML format according to the E2B standard
- Insertion and updates of medicines in the EMA database Extended EudraVigilance Medicinal Product Dictionary (XEVMPD)
- Preparation of the Periodic Safety Update Report (PSUR)
- Preparation of the Risk Management Plan (RMP)
- Preparation and maintenance of the Pharmacovigilance System Master File (PSMF)
- Drafting, review and updating of safety agreements for the exchange of information (Safety Data Exchange Agreements - SDEA)
- Periodic review of safety data
- **Signal** Analysis
- Back log of ADR reports and uplaod into the safety database
- Due Diligence of Pharmacovigilance documentation in case of MA transfers

# Surveillance and vigilance for MDs/IVDs

- Devicevigilance activities, with assumption of the role of vigilance responsible person
- Management of incident reports to the Competent Authorities
- Consultancy in the phase of investigation and preparation of corrective actions (FSCA)
- Activities of post marketing surveillance and Post-Market Clinical Follow-up (PMCF): implementation of the MDR and IVDR requirement
- Drafting of PSURs and other PMS and PMCF documents
- Trend analysis



# Foodvigilance and phytosurveillance

Management of post-marketing foodvigilance and of phytosurveillance in Italy and Europe

# Cosmetovigilance

- Assumption of the role of Contact Point for the Italian and European Authorities
- Managing cosmetovigilance and post marketing surveillance in Italy and in the European Union



# Scientific information and advertising

- Assumption of the role of Responsible for the Scientific Information of medicinal products for human use
- Assistance with the Scientific Information of medicines and submission to the Italian Medicine Agency (AIFA)
- Request of authorisation for conferences and congresses
- Assumption of the role of ACC Contact Person (Conferences and Congresses Authorization)

 Request of registration of Pharmaceutical Sales Representatives by the Regions to access health facilities

 Assistance to obtain certification according to the Farmindustria guidelines on Scientific Information

 Evaluation of advertising material to the public related to human OTC medicines, no-prescription veterinary medicines,
 PMCs, medical devices and IVDs, authorization requests to the Ministry of Health and authorization procedures

 Development of artworks of brochures and advertising materials and collaboration in the preparation and maintenance of websites

Assistance with promotional materials in other EU countries

### Clinical trials

- Regulatory consultancy and assistance for the authorization of interventional and observational clinical trials for medicinal products, medical devices and food supplements
- Review of documentation and technical assistance in the submission to the Competent Authorities and Ethics Committees
- Data entry into the Clinical Trials Information System (CTIS) and into the National Register for Observational Studies
- Translation of the dossiers, clinical protocols, informed consent and other documents to be included in the Clinical Trial Application (CTA)



# **Quality Services**

These consulting activities are addressed to companies and institutions wishing to comply with the necessary requirements for the **achievement of ISO 9001** and **ISO 13485**, **ISO 22716**, **GXP** (GMP, GDP and GCP).

GUARANTEE

In this area the following services are provided:

- Preparation of the corporate organization chart
- Preparation of Job Descriptions
- Preparation of the Quality Manual or evaluation of that already in use at the client site
- Preparation of Standard Operating Procedures (SOPs) and optimization of management procedures for all areas of regulatory activities
- Review of **technical agreements** and execution of audits at suppliers
- Assumption of the role of Quality Assurance
- Contacts with Certification Bodies
- Implementation of the quality system in accordance with ISO standards
- Preparation of registration system to ensure the compliance with procedures and manuals

#### **Audits**

#### Audit at the following facilities:

- Manufacturing sites of active pharmaceutical ingredients (APIs) and finished products, in Italy
  and in other European or non-EU countries
- Companies and suppliers of medicines, medical devices, food supplements, PMCs, biocides, cosmetics
- Companies offering services of clinical trials, pharmacovigilance, regulatory affairs
- Warehouses, distributors, wholesalers

### Publications in the Italian Official Journal

• Electronic **publication** services for **listings** in the Official Journal of the Italian Republic



## Legal and notary services

- Legal and Public Notary assistance in the regulatory sector
- Collaboration in the preparation of contracts for the purchase and sale of products
- Elaboration of expertise on legal issues related to regulatory activities

### **Translations**

- Scientific translations from/into the following languages: Italian, English, Spanish, French, German, Russian and other languages
- Sworn translations
- Sworn translation into German of leaflets and labels and upload to the Unifarm database (bilingualism) for the Italian market





#### e g u l a t o r y

#### Rome

Operational Headquarters: Via dell'Arco di Travertino 11 00178 Rome Tel. +39 06 77209020 Fax +39 06 70474067

Registered Office: Viale Manzoni, 59 00185 Rome direnzo@direnzo.biz Skype: di.renzo.regulatory.affairs twitter: @drregulatory

Milan Piazza Luigi di Savoia, 24 20124 Milano Tel. e Fax: +39 02 67380552

London 9 Seagrave Road London SW16 1RP **VISIT** our **WEBSITE** 

www.direnzo.biz



OCT/2023 - V2