## Asphalion End to End support during product lifecycle















## Drug discovery

## **Non-Clinical**

Clinical

# Dossier preparation

Market Authorization

## **Post- Authorization**

- Regulatory Roadmap
- Regulatory strategy
- Feasibility assessments
- Gap analysis

- Support in Scientific Advice meeting (EMA, FDA, NCA)
- SME & Orphan Drug designation process
- Clinical Trial Application & CTIS
- IMPD+IB
- IND, PIPs/iPSPs, GMOs
- Scientific, Medical and Regulatory writing

- Pre-submission meeting
- Medical writing for preclinical and clinical section (M2-5)
- CMC writing (M3)
- eCTD publishing
- Transparency

- EU (CP, DCP, MRP, RUP and NP); MHRA (IRP, NP); US (ANDA, BLA, NDS);
  ACCESS Consortium procedure management
- Translations management
- Life cycle management
- Data management
- Dossier preparation
- Variations management

CMC

Gap Analysis | Feasibility Assessments | Strategic Consulting | CMC writing during development | CMC writing for registration and life-cycle

### **Experience**



Continuous Improvement

Vision

## Pharmacovigilance

Drug Safety for Clinical Trials | EU-QPPV, PSMF, RMP and PV agreements | Case management, safety reports

### **Regulatory Operations**

#### xEVMPD:

- New MAs
- Variations
- Investigational products (IMP)
- SPOR:
- SMS
- OMS Registrations
- QC Eudravigilance registrations

#### **ISO IDMP compliance:**

- Consulting and Strategy
- Data management
- IDMP readiness
- Regulatory intelligence

#### Implementation of:

- CTMS, RIMS, DMS, eCTDtools, etc.
- Overall Data Management: Preparation, Migration, Validation
- Active Regulatory Review