



# Alexandra TIZON

Freelance Senior Clinical Operations Manager  
DPO Externe

## CONTACT

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## CERTIFICATIONS

### ICH GCP Training E6(R3)

The Global Health Network Certificate

November 2025

### MOOC GDPR

CNIL Certificate

November 2025

### MOOC Cyber Security

ANSSI Certificate

November 2025

## EDUCATION

Clinical Research Associate Certificate

University Diploma Paris, France 1998

Biochemistry BSc Level

Bordeaux University, France 1997

## PROFESSIONAL PROFILE

- Senior Clinical Operations Project Manager with 27 years delivering complex extensive Phase I to III trials across biotech, pharma and CRO environments.
- Expertise in clinical trial strategy from feasibility until archiving, ongoing improvement operational aspects, leadership, regulatory submission (EMA: CTA, CTIS and FDA: pre-IND/IND), risk and quality management, CRO oversight and inspection readiness.
- Proven track record in ensuring high quality execution, robust TMFs and eTMFs and on time on budget delivery on fast many highly regulated environments.
- Maintain registers, draft procedures, conduct DPIAs, and manage data breaches.
- ICH-GCP, GDPR, RIA, personal data protection, cybersecurity, French anti-gift law
- Available as a freelance consultant for short- or long-term assignments.

## KEY ACHIEVEMENTS

- Delivered around 30 national and international clinical studies
- Lead 3 EMA/FDA scientific advice processes with positive regulatory outcomes
- Collaborated 100% regulatory submission approvals
- Achieved 100% inspection readiness across multiple audits
- Successfully planning, organization, recovered at risks tools, improving timelines, analyzing data, managing budget and TMF completeness
- Established operational frameworks for multiple start-up and biotech companies

## CORE SKILLS

- Global CPM (Phase I to III – Non interventional studies - IITs), Cross functional leadership with international study teams
- Proactive, anticipating issues and developing solutions
- CROs and vendors oversight, financial excellence
- Experience with EMA (Scientific advice, CTA/CTIS/IRIS, Eudravigilance) FDA (Scientific advice, Pre-IND, IND)
- Maintain GDPR compliance
- Risk management, mitigation strategies, IMP management and supplies strategies, project plans and SOPs set-up and updates.
- TMF set-up, QC reviews and archiving, inspection readiness
- Communication skills and ability to work on multiple projects, ability to multitask, prioritize with strong attention to detail and accuracy of information.

# Alexandra

## TIZON

Freelance Senior Clinical  
Operations Manager  
DPO externe

### THERAPEUTIC EXPERIENCE

Oncology, Sepsis, Hematology,  
Dermatology, Gastrointestinal  
Diseases, Infectious Diseases,  
Postpartum Hemorrhages, Orphan  
Diseases, Diabetes, Graft Versus  
Host Disease, Microbiome  
Restoration Therapy, Vaccines,  
Phagotherapy

### LANGUAGES

- French: Native
- English: Fluent
- German: Elementary

### SOFTWARE

CTMS, EDC (Rave, Ennov)

eTMF (Veeva, Viedoc, Ennov)

Quality (Ennov)

### EXPERIENCE

#### Freelance Senior Clinical Operations Manager - DPO externe

**BIOXELA**- Since November 2025

##### Clinical Operations Project Manager - April 2021 – November 2025

*Pherecydes Pharma - Phaxiam, Lyon, France*

- Oversee CRO activities, deliverables, audits, site inspection, contract and budget negotiation.
- Management of deadlines, metrics, finances and forecasting
- IMP management, Risk Management, SOPs and project plans set-up and updates, Data Management activities follow-up and participation of data review meeting, TMF set up and quality review
- Regulatory submissions (CTIS and previous regulations)
- Investigator Meetings management, DSMB management, medical writing participation to CSR, abstracts and publication.

##### Senior Clinical Trial Manager - September 2018 – January 2021

*Genkyotex, Archamps, France*

- Overall management of international studies Phase I, Phase II and Investigator Initiated Studies.
- Oversee CROs activities and deadlines, project plans creation and update, TMFs set up, quality check review and archiving.
- Regulatory: FDA and EMA scientific advice, orphan drug designation, IND and CTA/EC submission, Eudravigilance. Inspections and audits readiness.
- Set up and update SOPs, DPO clinical trials expert (record of processing activities, Privacy Impact Assessment)

##### Freelance Senior Clinical Project Manager - December 2017 – August 2018

*Maat Pharma, Elsalys Biotech and Genkyotex, France*

- Overall management of international studies Phase I, Phase II, Phase III and Investigator Initiated Study.
- Oversee CROs activities and deadlines, project plans creation, budget
- Databases creation relating to retrospective studies, data management
- Mentoring clinical project manager
- TMFs set up, quality control and archiving, inspection readiness.

##### Clinical Project Director - May 2017 – July 2017

*DBV Technologies, France*

- Implement a global strategy of a TMF. Set up SOPs. Mentoring.

##### Senior Clinical Development Manager- November 2016 – March 2017

- Overall management of international studies in Oncology.

##### Senior Clinical Project Manager November 2012 – November 2016

*Covance (Labcorp) remote, France*

- Overall management of international studies US, EU, Asia Pac in accordance with local regulations, approved procedures and ICH/GCP guidelines
- Management of deadlines, metrics, audits, resources, finances, contracts and forecasting activities
- Regular and proactive liaison with other internal departments (legal, safety, regulatory, data management, statistics and quality)
- Project plans creation and update, identify risks and find solutions proactively
- Audits and inspection readiness.
- Ensure cross functional teamwork among project team members.

From **January 2008 to October 2012**, **Senior Clinical Project Manager – Quintiles (IQVIA) in house at Pfizer**, France: management of local Pfizer clinical trials only.

From **January 2005 to December 2007**, **Clinical Trial Manager – LFB, Les ULIS**, France: management phase II and III in orphan diseases.

From **March 2002 to December 2004**, **Clinical Research Associate – LFB, Les ULIS**, France: Regulatory submissions and monitoring of a phase I and IV and a non interventional study.

From **February 2001 to February 2002**, **Clinical Research Associate and Data Manager – GERCOR**, France: preparation and management of clinical studies phase II and III in oncology.

From **January 1998 to January 2001**, **Clinical Research Associate – Bordeaux Hospital**, France: Management of a pivotal phase III study with Ribavirin in Chronic Hepatitis C. French Agency inspection.