



ABOUT ME

I'm a resourceful and determined person, driven by a desire to want to actively contribute to scientific progress. My goal is to work in the Clinical Research, aiming to integrate my passion for science and innovation in conducting clinical trials.

SIMONA PELLEGRINO

CLINICAL RESEARCH EDUCATION

June 2024

- **E-learning course ICH GOOD CLINICAL PRACTICE E6 (R2)**

January 2024

- **Clinical Research Academy (CRO ClinOpsHub S.r.l.)**

- Risk Assessment for Clinical Research
- TMF / eTMF: inspection-readiness approach
- Study-specific eCRF validation

November 2023

- **MISSIONE CRA - 50 hours clinical research training course in compliance with the Italian Ministerial Decree 15/11/2011 provided by: CRO ClinOpsHub S.r.l. - CRO Yghea S.p.a.**

Main Topics:

- Methodology and Regulation of Clinical Trials
- Good Clinical Practice (**GCP**)
- Standards for Good Manufacturing Practice (**GMP**)
- Pharmacovigilance
- Quality Systems and Quality Assurance
- Focus on Phase 1 Quality
- Tasks of Monitors as per the paragraph 5.18 of Annex 1 to the Ministerial Decree 07/15/1997
- Tasks of Clinical Research Coordinators
Focus on **Reg. (EU) No.536/2014**
- Workshop on Site Selection, Essential Documents, Drug Accountability
- Site Visits Simulation (PSV, SIV, SMV, COV)

EDUCATION

February 2022 - present

- **Advanced English Language Course**
At My English School, Milan




October 2015 - March 2022

- **Master's degree in Pharmacy**
At University of Salerno




January 2018 - July 2018

- **Erasmus program - Pharmacy course**
At Universitat de València

LANGUAGES

-  Italian - **Native Language**
-  English - **Advanced**
-  Spanish - **Basic**

COMPUTER SKILLS

-  Microsoft Word - **Advanced**
-  Microsoft P. Point - **Advanced**
-  Microsoft Excel - **Intermediate**

OTHER SKILLS

- **Team working**
- **Problem-solving**
- **Flexibility and adaptability**

DRIVE LICENCE B

HOBBIES



WORK EXPERIENCES AND SOFT SKILLS

April 2024 - June 2024

■ **Clinical Trial Assistant (CTA)**

At Medpace Italia - Milan, Italy

- Perform a variety of support tasks for departmental personnel/processes;
- Perform administrative duties in conformity with company policies and procedures;
- Maintain databases/spreadsheets as necessary to facilitate tracking/documentation of departmental activities;
- Conduct quality control reviews of departmental documents as necessary;
- Support other departments on ad hoc projects.

September 2022 - March 2024

■ **Pharmacist in the role of deputy director**

■ **Pharmacist**

At Lloyds Admenta Italia S.p.a. - Milan, Italy

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- During my experiences as a CTA, I fine-tuned **attention to details** through the accurate management of ISF documentation; as a pharmacist I ensured the safety, efficacy and appropriateness of medical prescriptions.
 - I proved the ability to **communicate on multiple levels**, interacting with stakeholders involved in the clinical trial, suppliers and patients, explaining complex informations in an understandable way.
 - I acquired a **critical mindset**, making thoughtful decisions to ensure the safety and well-being of patients.
 - As Clinical Trial Assistant, **time management** was crucial to ensure self-organization during the preparation and updating of Investigator Site File.
 - Both my international experience and practice of team sports allowed me to mature **flexibility and respect for diversity** through building relationships and interactions with people of different cultures and nationalities.

