



## ONCOLOGY PHYSICIAN

### About us

Advanced Accelerator Applications (AAA) is an innovative radiopharmaceutical company that develops, produces and commercializes Molecular Nuclear Medicine (MNM) products. AAA's lead therapeutic product candidate, Lutathera<sup>®</sup>, is a novel MNM compound that AAA is currently developing for the treatment of neuroendocrine tumors, a significant unmet medical need.

Founded in 2002, AAA has its headquarters in Saint-Genis-Pouilly, France. AAA currently has 21 production and R&D facilities able to manufacture both diagnostics and therapeutic MNM products, and has over 500 employees in 13 countries. AAA is listed on the Nasdaq Global Select Market under the ticker "AAAP". For more information, please visit: [www.adacap.com](http://www.adacap.com).

AAA has a broad pipeline of products in development. In addition to the company lead product, Lutathera<sup>®</sup>, and its Gallium-68 Dotatate product NetSpot recently approved by the FDA, Phase II studies have been initiated with Annexin V-128, a diagnostic 99mTc tracer for apoptosis and necrosis, and the company has other imaging and therapeutic radiopharmaceutical products in development.

We are currently recruiting for an **Oncology Physician** to join our Clinical Development team.

### Core Responsibilities:

- Works within a Global Clinical Development Team with responsibilities across geographies within the assigned clinical trial program
  - Supports the clinical development strategy for one or more indications including successful regulatory submission
  - Designs and optimizes Clinical trial design
  - Leads the Clinical Development Team, including interactions with discovery, safety, marketing, regulatory, research and other functions
  - Designs and assists in authoring protocols and protocol amendments
  - Oversees and is responsible for the quality, coordination and timeliness of protocol development
  - Conducts medical review and interpretation of efficacy and safety data from clinical trials
  - Responsible for the quality, coordination, medical accuracy and timeliness of clinical study reports
  - Works with Head of Clinical and other team members to prepare abstracts, manuscripts and presentations for external meetings as well as author clinical sections of regulatory documents (IB, IND sections)
  - Assists senior management to oversee quality, coordination and timeliness of clinical sections of INDs, Investigator Brochures, CTAs, ISS's, ISE's, and clinical expert reports
  - Presents and discusses data and findings at relevant team, governance, KOL and regulatory meetings
-



- Participates, as required, as a clinical representative on Project Teams
- Establishes and maintains working relationship with Study investigators, key opinion leaders, academicians, and Senior managers and department heads across the corporation
- Ensures adherence to GCP/ICH and AAA SOP standards
- Maintains clinical and scientific awareness in area of expertise
- Guides day to day work of one or more Clinical Development Managers
- Implements lifecycle requirements and plans for designated indications

### **Key Requirements:**

- Minimum M.D. with a specialized knowledge in Solid Tumor Therapeutic or Disease Area (sub specialty, certification a plus)
- Minimum 6 years in Clinical Medicine with at least 4 in the pharmaceutical industry
- Understanding of drug development process
- Proven ability to work with and develop key opinion/thought leader
- Knowledge of Good Clinical Practices (GCP), FDA regulations and guidelines, and applicable international regulatory requirements
- Experience in designing Clinical trial strategies to obtain regulatory approval
- Ability to provide scientific and clinical expertise to a clinical development program and evaluate scientific and clinical strategies for a product
- Ability to develop and evaluate strategies for the clinical development of a designated indication and to critically evaluate outside expert advice
- Experience leading strategic planning for an indication or disease state
- Experience in Clinical project planning
- Experience working on global and complex Clinical trials
- Experience working effectively in a team/matrix environment
- Leadership Competencies
  - Drive Innovation
  - Shape Strategy
  - Promote Open Communication, Foster Teamwork
  - Create Global Mindset
  - Demonstrate Adaptability, Foster Risk Taking

### **Competencies /Skills:**

- Ability to use scientific and clinical knowledge to conceptualize study designs
  - Ability to anticipate and resolve problems
  - Excellent interpersonal skills and ability to communicate effectively with people in different regions and functions
  - Sound organizational skills
-



- Strong scientific/technical skills
- Strong interpersonal capabilities and ability to build networks
- Proven leadership skills
- Proven verbal communication and technical writing skills
- Ability to present clearly in scientific and clinical settings
- Ability to lead cross-functional teams and be a true team player
- Possesses sense of urgency and initiative
- Project management skills and focus on delivery of results

**Submission of your application:**

Please send your application in via email to [recruitment-US@adacap.com](mailto:recruitment-US@adacap.com). Your application shall include all the information you consider relevant, and at a minimum, a motivation letter, your personal data, education, employment history and details on your current position and salary as well as your salary expectations for this position.

Please note that we will only reply to candidates that we wish to bring to interview stage.

---