

## **Validation Associate**

### **Essential Duties and Responsibilities**

Performs a wide variety of activities to ensure compliance with applicable regulatory requirements. Specific responsibilities will include:

- Write and execute or oversee execution of equipment and computer IQ/OQ/PQ protocols and reports.
- Assist in validation initiatives including execution of protocols, compiling and reviewing data, writing summary reports and other required activities.
- Assist in other QA Activities, as required.

### **Qualifications**

Requires ability to plan, prioritize and organize diversified workload. Must have excellent technical writing, verbal communication skills, attention to detail and the ability to work on multi-departmental initiatives.

### **Education and/or Experience**

BS experience in a scientific discipline and a minimum of 1-2 years related experience in the biotechnology industry, with working knowledge of cGMP.

Qualified candidates should send resume and cover letter to:

Human Resources  
Insmmed Therapeutic Proteins  
2590 Central Ave.  
Boulder, CO 80301  
[careers@insmed.com](mailto:careers@insmed.com)